# Home parenteral nutrition: a systematic review 

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# Home parenteral nutrition: a systematic review 

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This report is one of a series covering acute care, diagnostics and imaging, methodology, pharmaceuticals, population screening, and primary and community care. It was identified as a priority by the Pharmaceuticals Panel (see inside back cover).

The views expressed in this publication are those of the authors and not necessarily those of the Standing Group, the Commissioning Board, the Panel members or the Department of Health.

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## Contents

List of abbreviations ..... i
Summary ..... iii
I Background information ..... 1
2 Research questions addressed ..... 3
3 Review methods ..... 5
Section 1 Which patients have received HPN? ..... 5
Section 2 What has been the patient
experience of HPN? ..... 5
Section 3a Organisation ..... 5
Section 3b Comparative data ..... 6
Section 4 Cost-effectiveness of HPN ..... 6
Section 5 What gaps are there in the evidence? ..... 6
Literature search strategy and study retrieval ..... 6
4 Results ..... 9
Located studies ..... 9
Issues of study validity ..... 10
Design of studies included in the review ..... 10
Results for Section 1 Which patients have received HPN? ..... 10
Results for Section 2 Patient experience on HPN ..... 14
Results for Sections 3a and b Organisation and comparative data ..... 27
Results for Section 4 Economic analysis ..... 29
5 Discussion ..... 31
Home parenteral nutrition and intestinal failure ..... 31
The need for the review ..... 32
Finding the research ..... 32
The standards of the literature ..... 32
Trends of use ..... 32
Quality of care ..... 32
Organisation and evidence of effectiveness ..... 33
Economic appraisal ..... 33
6 Conclusions ..... 35
What gaps in the evidence exist? (Section 5 of the review method) ..... 35
Which questions need to be addressed? ..... 35
What methodological issues need to be addressed in future research? ..... 35
7 References ..... 37
Studies satisfying the criteria for inclusion in the review ..... 37
Other references ..... 39
Studies not included in the systematic review of HPN and the reasons why ..... 39
Appendix I ..... 49
The history of HPN ..... 49
Appendix 2 ..... 51
Review protocol ..... 51

## List of abbreviations

| ASPEN | American Society for Parenteral and Enteral Nutrition |
| :--- | :--- |
| BAPEN | British Association for Parenteral and Enteral Nutrition |
| CI | confidence interval |
| ESPEN | European Society for Parenteral and Enteral Nutrition |
| HPN | home parenteral nutrition |
| MVD | mesenteric vascular disease |
| MVO | mesenteric vessel occlusion |
| QALY | quality-adjusted life year |
| QoL | quality-of-life |
| SCI | Science Citation Index |
| TPN | total parenteral nutrition |
|  |  |

# Summary 

## Objectives

The objective of this Review was to locate, appraise and summarise evidence from scientific studies on home parenteral nutrition (HPN) in order to answer specific research questions on the effectiveness of this technology.

The following questions were asked. What patients have received HPN? What has been the experience of patients on HPN programmes? How have HPN programmes been organised, and what techniques and equipment have been used, and to what effect? What comparative information is available on effectiveness? What evidence exists for the costeffectiveness of HPN? What questions about the provision of HPN could be answered with additional research, and what studies would be most suitable?

## Data sources

A comprehensive list of studies was provided by an extensive search of electronic databases (including MEDLINE, Embase, Science Citation Index, Uncover, Cinahl, Caredata, Food Science and Technology Abstracts, NTIS, Pascal, Psychlit, and Economic Literature Index), relevant journals (including Journal of Parenteral and Enteral Nutrition, Clinical Nutrition, American Journal of Clinical Nutrition, Nutrition, Clinical Gastroenterology, Nutrition Reviews, Annals of Nutrition and M etabolism, Nutrition and Cancer, Nutrition and H ealth, and Journal of Paediatric Nutrition and $M$ etabolism), and scanning of reference lists, as well as other search strategies outlined in the protocol.

## Study selection

Studies relevant to the questions were selected. The inclusion criteria were fairly broad because of the quality of the studies located.

## Data extraction

Data extraction forms were used to collect data from studies included in the review. The data was checked by a second researcher to reduce error.

## Data synthesis

Quantitative analysis was difficult owing to the type of studies located. The data is discussed in a qualita-
tive manner. Where complication rates have been given, we have attempted to combine the results in a quantitative manner.

## Results

The age and sex of patients on HPN varies according to the underlying disease but, on the whole, patients are young (see Tables 4a and 4b). There are trends showing an increased use of the technology at the extremes of the age range. There are marked differences between countries on the underlying diseases for which HPN is indicated. For example, many more patients with an underlying malignancy are treated in Italy and the USA than in the UK ( $40-67 \%$ versus $8 \%$ ). Morbidity rates for the majority of patients are acceptable (see Table 8), the complications tend to be related to the central venous catheter. It is fairly clear that a minority of patients are susceptible to recurrent problems and that many patients have very few complications. The mortality rate for HPN patients (see Table 10) was good for those patients with benign underlying disease (for example, $5 \%$ of Crohn's HPN patients die per year), and there are very few reports of patients dying from complications of the technology. The survival of those with malignant disease and AIDS is poor, almost all having died from the underlying disease at one year; despite this, most programme growth worldwide is due to an increase in the numbers of patients with these diagnoses (see Table 5). Quality of life is reasonable for patients with benign disease (see Table 9); no studies were found that examined the quality of life of HPN patients with malignant disease. Economic analysis shows that the cost of HPN treatment is cheaper than the alternative of in-patient care (see Table 18). There is a paucity of comparative studies examining different aspects of the technology, and this accounted for the majority of gaps in the evidence.

## Conclusions

The use of HPN for benign intestinal failure is supported by evidence from the scientific studies located. There are, however, large gaps in the evidence, particularly relating to the use of HPN in malignant disease and AIDS. A programme of research is suggested at the end of this review.

## Chapter I

## Background information

Home parenteral nutrition (HPN) is a complex technology involving the intravenous infusion of liquid nutrition directly into a central vein. The patient, or carer, is taught to manage the complicated sterile routine, enabling transfer of care to the home. It is an expensive and time-consuming routine which may be required for many years or, in some cases, life. It intrudes into the patient's life and any alteration from the daily routine requires planning. This is likely to have an effect on the quality of life experienced by both the patient and his or her family.

The technology is used to treat intestinal failure, defined as an inadequate intestinal function for absorption of fluid electrolyte and nutrient requirements. Intestinal failure can be caused by destruction of the available absorptive surface (for example, Crohn's disease), chronic intestinal obstruction (malignant disease and motility disorders), or by extensive removal of the absorptive surface (following mesenteric artery occlusion or extensive small bowel resection). There is a range of severity of intestinal failure from complete to partial. Complete failure suggests that the patient will require all fluids, nutrients and trace elements to be given parenterally, suggesting that the patient would die quickly of a combination of dehydration and malnutrition without treatment. Partial failure usually means that some parenteral support is required (possibly only fluids) but that intensive enteral support might suffice. Adaptation of the intestinal mucosa may allow 'weaning' of parenteral
support, and is one reason why patients can stop HPN treatment. Recovery from the underlying disease is the other main reason for stopping parenteral support. The liquid nutrition is infused (usually overnight) through a sterile, pemanently in-dwelling, central venous access device. The complex techniques of infusing the nutrition safely are mastered by the patient or a carer, usually before the patient is discharged from hospital.

Patient referral patterns for HPN treatment are inconsistent, some regions in the UK having very few HPN patients. However, there are several large centres in the UK where HPN is considered as an essential, life-saving treatment. The prevalence of HPN patients in the northwest of England and Scotland is approximately $18-20$ per million. This figure is comparable to that from Denmark, a country with a very similar disease profile to the UK. (The UK national average is 2.6 per million, reflecting the fact that higher rates are seen close to major referral centres.)

The NHS Research and Development Programme's Standing Group on Health Technology Assessment prioritised HPN as a technology requiring further assessment because of cost, effect on quality of life, variable referral patterns and uncertainty regarding effectiveness. The aim, therefore, of this systematic review was to assess the extent and quality of evidence on HPN and to identify that which should inform practice.

## Chapter 2

## Research questions addressed

The aim of this review is to locate, acquire and synthesise studies concerning the use, effectiveness and cost-effectiveness of home parenteral nutrition. The review falls into five main sections, illustrated by the research questions below; for each of these the literature on both adults and children is reviewed. The first four sections are included in chapter 4 - Results; the fifth is covered in chapter 6 - Conclusions.

Searching the HPN literature suggests that there are very few comparative studies available. Most of the literature consists of case series. The research questions were formulated bearing this is mind, so as to make the best use of what information is available.

Section 1. Which patients have received H PN ? It is important to identify the types of patients who have received HPN treatment. We aimed to describe the age, sex and diagnostic profiles of patients and to outline trends that may be taking place.

Section 2. What has been the experience of patients on H PN programmes?
We aimed to describe the type and incidence of complications, survival, duration of HPN treatment, quality of life and why HPN treatment was
discontinued, so that an accurate profile of 'life on HPN' could be constructed. We also aimed to identify any moderating factors.

Section 3. (a) H ow haveH PN programmes been organised, and what techniques and equipment have been used, and to what effect? (b) What comparative information is available on effectiveness?
We aimed to find out what methods of organisation of HPN programmes existed and whether any organisational model was superior. We also wanted to identify what comparative evidence exists on different aspects of the technology.

Section 4. What eviden ce exists for the cost-effectiveness of H PN ?
The aim of this section was to compare the costeffectiveness of HPN with any alternative treatments that might be available. In addition, an examination of the costs involved with an HPN programme was undertaken.

Section 5. What questions about the provision of H PN could be answered with additional research, and what studies would be most suitable?
Our aim was to highlight gaps in the research knowledge and also to identify key research questions which need to be answered.

## Chapter 3

## Review methods

Aprotocol was developed for the systematic review of HPN following the NHS Centre for Reviews and Dissemination Guidelines (Deeks \& al, 1996).

## Section I Which patients have received HPN?

A longitudinal inception study would provide actual patient numbers being treated on HPN by subgroup, whilst a cross-sectional sample would only provide a snapshot of the types of patients being treated with HPN. The results will differ according to the length of time spent on HPN. The results of the two types of studies will be interpreted separately. Retrospective data collection is prone to be less complete than prospective data collection.

Complete assessment or random sampling are the best ways of maintaining representativeness. Both rely on the correct identification of a sampling frame. When these are not available, it may be possible to validate sample coverage by taking a sample using a second source and noting the degree of similarity in sample members. Nonresponse and missing data in a survey reduces its validity.

## Inclusion criteria

(a) Surveys of HPN use.
(b) Information from databases of HPN users.
(c) Data from cohorts of HPN patients.

## Points for assessing validity

(a) Are the patient numbers based on new patients in a given period, or from a crosssectional sample?
(b) Was the data collected prospectively or retrospectively?
(c) What proportion of patients were sampled and how was the sample chosen?
(d) From what group was the sample selected and how representative is it?
(e) Was the sample coverage validated and if so was it found to be acceptable?
(f) For what proportion of selected patients was no information available?
(g) How complete was the data that was acquired?

## Section 2 What has been the patient experience of HPN?

Studies which have been included in Section 1 may also be of interest here. Ascertainment of outcomes should be free from bias. Where excessive patient investigations are carried out, a higher number of events may be detected than in case series with less active investigation. The instruments used to measure subjective issues like quality of life need to be validated. A potential problem is this area is the choice of denominators for the calculation of rates, whether they are patient numbers, patient years, or the way in which data has been analysed. When reporting rates it is important to make the distinction between per 1000 per year, which implies a group of patients are all followed up for the same length of time, and per 1000 patient years.

## Inclusion criteria

(a) Studies reporting the experience of inception cohorts of HPN users.
(b) Studies giving information on length of treatment, mortality, complications, or quality of life.

## Points for assessing validity

(a) Is data collected prospectively or retrospectively?
(b) What cohort was recruited?
(c) How much of the cohort was successfully recruited?
(d) How complete was the follow-up?
(e) What procedure was used to detect complications?
(f) What quality-of-life instrument was used and how was it validated?

## Section 3a Organisation

## Inclusion criteria

Surveys assessing issues in the delivery of HPN since 1980.

## Points for assessing validity

Any study which has assessed how HPN programmes have been organised.

## Section 3b Comparative data

The validity of comparisons between, for example, different procedures, will greatly depend on the study design. In addition, different aspects of validity will be important to different designs of study. The studies will be grouped according to their design in the analysis.

## Studies to be included

Any comparative study.

## Points for assessing validity

(a) What was the study design? Descriptive or comparative?
(b) Is data collection prospective or retrospective?
(c) Was a comparison made?
(d) How were allocations to treatment made?
(e) Was follow-up complete?
(f) Were the groups comparable with respect to age and diagnosis?
(g) What outcomes were measured?
(h) Was the length of follow-up more than 3 months in all cases?
(i) How were outcomes assessed?

## Section 4 Cost-effectiveness of HPN

## Inclusion criteria (economic analyses)

Any economic evaluation of an HPN programme.

## Validity

(a) What methodological technique has been applied?
(b) What was the comparison made with?
(c) What perspective was adopted?
(d) Were all costs considered?
(e) Were costs measured appropriately?
(f) Were all outcomes considered?
(g) Were they measured appropriately?
(h) How was quality of life assessed?
(i) Was a marginal analysis performed?
(j) Was the robustness of the result tested in a sensitivity analysis?

## Section 5 What gaps are there in the evidence?

What clinical issues need to be addressed?
What methodological issues need to be addressed?

## Literature search strategy and study retrieval

The aim of the literature search was to provide a comprehensive list of primary studies. It included all types of study design and included all possible aspects of HPN technology.

Before starting the search, advice was sought from an information scientist at the NHS Centre for Reviews and Dissemination, York, and from a senior medical librarian based at Hope Hospital, Salford.

The following possible sources of data were identified.

- Electronic databases
- Hand-searching of relevant journals
- Personal literature collections
- Conference proceedings
- Writing to all major centres in Europe and the USA
- Science Citation Index
- Scanning reference lists of studies located


## Electronic databases

It is well documented in the literature that many studies can be missed if searches are limited to only one database. We searched 11 separate databases from 1968 onwards, that is, from the origins of HPN (see Appendix 2 Review protocol).

## Hand searching

Ten journals were hand searched for the period, January 1980-July 1995 (or for whatever period within this frame they were available).

## Conference proceedings

Proceedings of the annual conferences of the following bodies were obtained.

- ASPEN (American Society for Parenteral and Enteral Nutrition) - 1993, 1994, 1995
- ESPEN (European Society for Parenteral and Enteral Nutrition) - 1993, 1994, 1995
- BAPEN (British Association for Parenteral and Enteral Nutrition) - 1994


## Personal literature collections

We examined the studies collected by Professor Sir Miles Irving who was one of the founders of HPN in the UK.

## Letters to experts

We contacted major European and American centres requesting published and unpublished studies, and any details of on-going work.

## Visits to major meetings

Two members of the project team attended the ESPEN meeting in Rome (September 1995) and the BAPEN meeting (December 1995).

## Selection of eligible studies, checking validity and data extraction

One researcher checked articles for eligibility for each of the research questions, graded their validity, and extracted the necessary data. A second researcher validated these decisions by processing a $10 \%$ random sample of included and excluded studies.
therefore, discussed in a qualitative manner except in the case of the complications where a weighted average is calculated. The qualitative analysis takes into account the magnitude of the results and the size and validity of the studies, together with any moderating factors.

## Peer review

Once completed the manuscript was submitted for peer review to the following experts.

- Professor Bernard Messing, Paris
- Dr Karin Ladefoged, Copenhagen
- Professsor Anne Ferguson/Dr Subrata Ghosh, Edinburgh
- Dr Andre Van Gossum, Brussels
- Professor John Lennard-Jones, London
- Mr Kenneth Fearon, Edinburgh


## Study synthesis

Because of the lack of comparability of data, quantitative synthesis is not appropriate. The results are

## Chapter 4

## Results

## Located studies

A total of 256 studies were located using the methods described. The number of additional articles located by each method is shown.

## Electronic databases

The numbers of additional articles found by searching relevant electronic databases are shown in Table 1.

TABLE 1 Number of additional studies located in each database compared to those in MEDLINE

| Database | No of <br> studies found |
| :--- | :---: |
| MEDLINE (Index Medicus on-line) | 59 |
| EMBASE (Excerpta Medica on-line) | 13 |
| UNCOVER | 10 |
| CINAHL | 0 |
| CAREDATA | 0 |
| Food Science and Technology Abstracts | 0 |
| NTIS | 0 |
| PASCAL | 0 |
| PSYCHLIT | 2 |
| Economic Literature Index | 0 |

## Hand searching of relevant journals

The numbers of additional references found by searching appropriate journals are shown in Table2.

## Conference proceedings

We found 58 abstracts that were relevant to the review. Attempts to locate completed studies from the authors were successful in eight cases. Abstracts for which no full report was available were not included in the review.

## Personal literature collections

A search of personal literature collections yielded 15 additional papers.

TABLE 2 Number of studies located by hand searching relevant journals

| Journal | No of <br> studies found |
| :--- | :---: |
| American Journal of Clinical Nutrition | 2 |
| Annals of Nutrition and Metabolism | 1 |
| Clinical Nutrition | 15 |
| Clinical Gastroenterology | 4 |
| Journal of Paediatric Gastroenterology |  |
| and Nutrition | 4 |
| Journal of Parenteral and Enteral Nutrition | 28 |
| Nutrition | 7 |
| Nutrition and Cancer | 2 |
| Nutrition and Health | 0 |
| Nutrition Reviews | 0 |

## Letters to experts

Replies were received from seven of the 24 centres contacted. The replies included details of a total of 25 papers, of which eight had not been previously identified. We did not receive any data relating to work in progress.

## Visits to major meetings

The amount of relevant on-going work was disappointing. Experts from centres were contacted and asked to supply further data if they had not already done so. No additional studies were located.

## Scanning of reference lists

Scanning the reference lists of these studies located a further 56 articles. A further scan of the studies located by this method identified an additional 19 studies and a third scan found three more papers.

## Science Citation Index

A search of the Science Citation Index for the following names was carried out; Scribner, Jeejeebhoy, Shils, Wilmore, Rhoads, Vars, Ladefoged, Irving, Messing and Howard. We did not locate any new studies.

## Issues of study validity

## Inclusions and exclusions

A total of 256 studies were located and retrieved, not including abstracts, letters, editorials, case reports and review papers.

There was one disagreement between reviewers in the excluded sample, which resulted in the inclusion of that study. No further discrepancies were found. In all, 191 studies did not satisfy the inclusion criteria. The main reason for their exclusion is given in the reference section. Many of the studies retrieved were only relevant to hospital TPN and included peri-operative feeding, chemotherapy and short-term feeding. Some hospitalbased studies were relevant to HPN, for example, studies looking at care of central lines, but most were not relevant. Another common reason for exclusion was the age of the study. HPN use in the UK was rare prior to 1980 but was used in the USA during the 1970s. During early HPN use, complication rates were relatively high and were probably related to the type of catheters that were available, the quality of the intravenous solutions available, and the protocols that were followed. Elimination of early studies was important, to prevent learning curve bias skewing the results of more recent HPN use. The studies from the 1970s tended to be case series and included only small numbers of patients. No well-designed studies with large numbers of patients were excluded as a result of the age exclusion criteria being applied.

The majority of studies found were case series and the centres producing case series tended to report experience as their HPN population grew. This led to much data duplication; many studies included the same patients as in previous reports plus the new cases. For these studies only, the latest and most comprehensive report was included.

Some studies were relevant to HPN patients but were not specifically related to the questions we had set. These studies tended to relate to the biochemical details of total parenteral nutrition (TPN) and their consequences.

Review papers were excluded when they contained no original empirical data, but they were useful sources of citations. The unreliable nature of data contained in abstracts meant that they were not used for data extraction; letters were excluded for the same reason.

Out of 256 studies, 65 fulfilled the inclusion criteria and were used to provide evidence to answer the
research questions. These studies, which are listed in the references, were subjected to data extraction, and the results of this are outlined below.

## Design of studies included in the review

The types of study design identified in the studies included in the review are shown below. Only one randomised trial was identified. The economic appraisals of HPN were limited to a cost-utility analysis from Canada (Detsky, 1986) and a costutility analysis from the UK (Richards, 1996). In addition, there were several simple cost analyses. There were 54 case series. Only six studies had any form of comparative data.

Section 1 (Which patients have received HPN?) Case series
Section 2 (Patient experience?) Case series Quality-of-life questionnaires 7
Section 3a (Organisation) Case series 1
Section 3b (Comparative data) Randomised controlled trial 1 Prospective controlled 1 Comparative studies 4

Section 4 (Economic analysis) Cost-utility analysis 2 Simple cost analysis 5

The total number is greater than 64 because some studies were relevant to more than one section.

## Results for Section I Which patients have received HPN?

Fifteen studies were relevant to this section. Seven were from the USA and eight were from Europe (see Table 3). All the studies were longitudinal case series and only one was prospective. Patients were often included more than once in the results of a study, as several sources of data were used which overlapped. The sample sizes ranged from nine to more than 9000 . It was often impossible to say what proportion of the total HPN population had been sampled by these studies, and there were no studies that randomly sampled a HPN population. Few studies reported how many patients were lost to follow-up.

It can be seen from Tables $4 a$ and $4 b$ that there is a trend towards the use of HPN in older age groups and the very young. The increasing age of HPN

TABLE 3 Summary of studies selected for Section I: Which patients have received HPN?

| Study (Country) | Date | Study design | Patients included in more than one study | Sample size | Proportion of HPN population sampled (\%) | Sampling frame | Random sampling | \% lost to follow-up |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| $\begin{aligned} & \text { Bisset, } 1992 \\ & \text { (UK) } \end{aligned}$ | 1987-92 | Case series; longitudinal; retrospective | possible | 10 |  | none | no | 0 |
| $\begin{aligned} & \text { DePotter, } 1992 \\ & \text { (France) } \end{aligned}$ | 1981-90 | Case series; longitudinal; retrospective | possible | 156 | ? | none | no | ? 0 |
| Goutebel, 1987 (France) | 1979-85 | Case series; longitudinal; retrospective | yes | 85 | ? | none | no | ? 0 |
| Griffith, 1984 (UK) | 1978-83 | Case series; longitudinal; retrospective | possible | 9 | 100 | none | no | ? 0 |
| Howard, I993 <br> (USA) | 1985-90 | Case series; longitudinal; retrospective | yes | 2275 | 10 (E) | patient <br> registry | no | ? |
| Howard, I99। <br> (USA) | 1984-87 | Case series; longitudinal; retrospective | yes | 1594 | 7.8 (E) | none | no | 7 |
| Howard, 1986 (USA) | 1983-85 | Case series; longitudinal; retrospective | yes | 2556 | ? | five separate sources | no | ? |
| Howard, 1995 <br> (USA) | 1985-92 | Case series; longitudinal; retrospective | yes | 9288 | 5 | two separate sources | no | ? |
| Messing, 1989 (Europe) | 1974-85 | Case series; longitudinal; retrospective | yes | 194 | ? | 27 centres | no | ? |
| Messing, 1995 (France/Belgium) | 1980-89 | Case series; longitudinal; retrospective | yes | 217 | ? | nine centres | no | 0 |
| O'Hanrahan, 1992 (UK) | 1977-91 | Case series; longitudinal; retrospective | yes | 400 | ? | none | no | ? |
| Ralston, I984 (USA) | 1977-82 | Case series; longitudinal; retrospective | possible | 14 | ? | one centre/ $<2$ months of age | no | 35 |
| SchmittSommerfeld, 1990 (USA) | 1980-85 | Case series; longitudinal; retrospective | possible | 35 | ? | none | no | ? 0 |
| Van Gossum, 1995 (Europe) | 1993-94 | Case series; longitudinal; prospective | yes | 496 | 80 (E) | 95 centres | no | ? |
| $\begin{aligned} & \text { Vargas, } 1987 \\ & \text { (USA) } \end{aligned}$ | 1976-86 | Case series; longitudinal; retrospective | possible | 102 | ? | none | no | ? 0 |

TABLE 4a Section I. Results - Age and sex of adult patients commenced on HPN

| Study | Mean age (range) | $\%$ in an age group |  |  | Mean age by diagnostic group |  | Male:Female | Trends |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Goutebel, 1987 | $\begin{gathered} \text { Median } 50 \\ (18-79) \end{gathered}$ |  |  |  |  |  |  |  |
| Griffith, 1984 | 32.8 (22-45) |  |  |  | 5:4 |  |  |  |
| Howard, 1991 |  |  |  |  | Crohn's <br> MVD AIDS | $\begin{aligned} & 36 \\ & 57 \\ & 29 \end{aligned}$ | More than 80\% of those with radiation enteritis were women |  |
| Howard, 1993 |  |  |  |  | Crohn's <br> Radiation <br> enteritis <br> Cancer | $\begin{aligned} & 35 \\ & 57 \\ & 43 \end{aligned}$ | $\begin{aligned} & 3: 5 \\ & 1: 2 \\ & 9: 10 \end{aligned}$ |  |
| Howard, 1995 |  |  |  |  | Crohn's <br> MVD <br> Cancer <br> AIDS | $\begin{aligned} & 36 \\ & 49 \\ & 44 \\ & 33 \end{aligned}$ |  |  |
| Messing, 1989 | 44 |  |  |  |  |  | 1.08 |  |
| Messing, 1995 | $46.5$ | $\begin{aligned} & \text { age }<40 \\ & \% \quad 40 \end{aligned}$ | $\begin{gathered} 40-60 \\ 37 \end{gathered}$ | $\begin{aligned} & 60+ \\ & 23 \end{aligned}$ |  |  |  |  |
| O'Hanrahan, 1992 |  | $\begin{array}{ll} \text { age } & 0-30 \\ \% & 31 \\ \% & 44 \end{array}$ | $\begin{aligned} & 31-50 \\ & 54.5 \\ & 38.5 \end{aligned}$ | $\begin{array}{r} 50+ \\ 14.5 \\ 17.5 \end{array}$ |  |  |  | Slight increases at the extremes for the second 200 patients |
| Van Gossum, 1995 |  | $\begin{array}{lc} \text { age } & \text { 16-40 } \\ \% & 36 \end{array}$ | $\begin{gathered} 41-60 \\ 41 \end{gathered}$ | $\begin{aligned} & 61+ \\ & 23 \end{aligned}$ |  |  |  |  |

TABLE 4b Section I. Results - Age and sex of paediatric HPN patients

| Study | $\mathbf{0 - 2 4}$ months <br> $\mathbf{n}(\%)$ | $\mathbf{2 - 1 0}$ years | II-I8 years | Male:Female | Trends |
| :--- | :---: | :---: | :---: | :---: | :---: |
| DePotter, 1992 | 51 | 25 | 24 | $90: 66$ |  |
| Schmitt-Sommerfeld, 1990 | 17 <br> $(1-12$ months $)$ | 20 <br> $(1-12$ years $)$ | 63 <br> $(12-23$ years $)$ |  |  |
| Vargas, 1987 | 59 <br> $(0-36$ months $)$ |  | 29 |  |  |

patients is almost certainly explained by its increasing use in malignant disease. The use of HPN in Crohn's disease is associated with younger patients and it is used more commonly in females, reflecting the prevalence of Crohn's disease in the UK. The use of HPN in radiation enteritis is naturally associated with older female patients. As the incidence of atherosclerosis increases with age, it is apparent that patients with mesenteric vascular disease (MVD) who require HPN are in the older age range.

In paediatric practice the main use of HPN is in the $0-24$-month age group, with an excess of males.

The diagnostic subgroups in paediatric practice are varied, but Crohn's disease remains common in older children. The small numbers of patients in the studies meant that no firm conclusions could be made regarding the use of HPN in children, particularly in the UK.

In the UK and some European countries, the largest diagnostic group comprises patients with Crohn's disease; the use of HPN for patients with a cancer diagnosis is exceptional (see Tables 5 and 6). Anecdotal evidence suggests that many UK cancer patients receive enteral rather than parenteral nutritional support. In the USA and Italy,

TABLE 5 Patients recruited to adult HPN programmes by country and diagnostic group

| Country (Study) <br> (Study) | Sample size | Crohn's disease (\%) | MVD (\%) | Malignancy <br> (\%) | Radiation enteritis (\%) | Motility disorder <br> (\%) | AIDS <br> (\%) | Others <br> (\%) | Trends |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| USA <br> (Howard, 1986) | $\begin{array}{r} 465 \\ 153 \\ 224 \\ 1351 \\ 268 \\ 89 \end{array}$ | NR 20 <br> OF 40 <br> CR 20 <br> CI 26 <br> CII 12 <br> CIII 9 | $\begin{array}{r} 10 \\ 9 \\ 7 \\ 0 \\ 0 \\ 1 \end{array}$ | $\begin{array}{r} 44 \\ 3 \\ 25 \\ 28 \\ 41 \\ 48 \end{array}$ |  | $\begin{array}{r} 5 \\ 7 \\ 14 \\ 2 \\ 3 \\ 4 \end{array}$ |  | $\begin{aligned} & 21 \\ & 41 \\ & 34 \\ & 44 \\ & 44 \\ & 36 \end{aligned}$ | Increased use in cancer patients. In 1978, $17 \%$ on HPN had cancer. This increased to $44 \%$ in 1983 |
| USA <br> (Howard, 1991) | 1594 | 1984 25 <br> 1985 19 <br> 1986 14 <br> 1987 12 | $\begin{array}{r} 14 \\ 14 \\ 7 \\ 8 \end{array}$ | $\begin{aligned} & 16 \\ & 26 \\ & 30 \\ & 39 \end{aligned}$ | $\begin{array}{r} 11 \\ 6 \\ 4 \\ 4 \end{array}$ | $\begin{array}{r} 13 \\ 9 \\ 8 \\ 8 \end{array}$ | $\begin{aligned} & 1 \\ & 2 \\ & 3 \\ & 2 \end{aligned}$ |  | Increased use in cancer versus benign diagnostic groups |
| USA <br> (Howard, 1993) | 1672 | $\begin{array}{rr} 1985 & 17 \\ 1987 & 11 \\ 1989 & 7 \end{array}$ |  | $\begin{aligned} & 35 \\ & 43 \\ & 46 \end{aligned}$ | $\begin{aligned} & 5 \\ & 4 \\ & 2 \end{aligned}$ |  |  |  | $90 \%$ of programme growth accounted for by new patients with malignant disease |
| USA <br> (Howard, 1995) | 9288 | 11 | 6 | 41 | 3 | 5 | 6 | 22 | Use of HPN doubled between 1989 and 1992 |
| France/ Belgium (Messing, 1995) | 217 | 25 | 27 | 20 | 22 |  | 11 |  |  |
| France (Van Gossum, 1995) | 133 | 15.7 | 16.5 | 21 | 17.2 |  | 6 | 23 |  |
| Italy (Pironi, 1993) | 135 | 2 | 9.7 | 67 | 5.2 |  | 0.7 | 14 |  |
| Scandinavia (Van Gossum, 1995) | 55 | 18 | 6 | 56 | 0 |  | 0 | 18.5 |  |
| Germany (Van Gossum, 1995) | 38 | 10.5 | 2.6 | 81 | 0 |  | 0 | 5 |  |
| Belgium (Van Gossum, 1995) | 25 | 4 | 8 | 45 | 12.5 |  | 25 | 4 |  |
| UK (Van Gossum, 1995) | 56 | 44.5 | 10.5 | 8.9 | 3.5 |  | 3.5 | 28 | Increased use in cancer and AIDS |
| UK (O'Hanrahan 1992) |  | 45.5 | 12 | 5 | 5.5 |  | 5 | 24 (96 patients) | 5-fold programme growth from 1980 to 1990 |

TABLE 6 Patients recruited to paediatric HPN programmes by diagnosis

| Study | Country | Crohn's <br> disease <br> (\%) | Chronic <br> pseudo <br> obstruction <br> (\%) | Intractable <br> diarroea <br> (\%) | Immune <br> deficiency <br> (\%) | Short <br> syndrome <br> bowel <br> (\%) | Malignancy <br> (\%) | Others Trends <br> (\%) |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| DePotter, 1992 | France | 15 | 19 | 5 | 8 | 41 | 12 |  |
| Schmitt- <br> Sommerfeld, 1990 | USA | 57 | 0.3 |  | 23 |  | 17 |  |
| Vargas, 1987 | USA | 22 | 10 | 15 |  | 10 | 44 |  |
| $\dagger$, Not elsewhere classified. |  |  |  |  |  |  |  |  |

the main diagnostic group comprises patients with malignant disease, although there are still many HPN patients with Crohn's disease. Italy has the highest percentage of HPN patients with a cancer diagnosis. Anecdotal evidence suggests this is related to the uncommon use of enteral nutritional support for these patients (there is a low consumption of enteral nutrition products in Italy compared with the rates of consumption in other similar European countries).

The trends indicated by the studies show that there is an increasing use of HPN in cancer patients and that this increase is responsible for the majority of programme growth ( $90 \%$ in the USA) in those countries where the use of HPN for malignant disease is high. In the UK, the number of HPN patients with malignant disease is small but has increased from $5 \%$ in 1992 to $8.9 \%$ in 1994. The wasting associated with AIDS is also becoming an increasingly common reason for HPN therapy and, in a recent paper from the USA (Howard, 1995), $6 \%$ of more than 9000 patients had this diagnosis. In the UK, AIDS was the reason for HPN being used in two of 53 patients registered during 1993.

## Results for Section 2 Patient experience on HPN

A total of 56 studies were relevant to Section 2 and these are summarised in Table 7. Of these, 37 studies were from the USA and the rest were from Europe. In 41 studies only adults were included, seven looked at a mixture of adults and children, and seven looked at children only; one study did not give ages. The sample sizes were similar to those for Section 1. Retrospective case series predominated, the only other study design being a quality-of-life assessment. As with Section 1 studies, it was unusual to see the numbers lost to follow-up being reported.
studies examined the quality of life of HPN patients; ten studies reported survival; 26 studies recorded the duration of HPN, and 28 studies reported the reasons for discontinuing HPN.

One of the main complications of HPN is sepsis (see Table 8 and Figure 1 ) and the most common focus for sepsis is the central venous catheter. Catheter sepsis is to some extent related to how well patients are trained in HPN techniques and, in turn, patient training is related to the skill and experience of the nutrition nurse. The larger series show a narrow band of episodes per catheter year (0.38-0.50). Smaller series seem to have fewer episodes of catheter sepsis; however, the confidence intervals are much wider. A weighted average of the rate of catheter sepsis was 0.34 ( $95 \% \mathrm{CI}, 0.32$, 0.37 ) episodes per catheter year. The number of patients experiencing an episode of sepsis per catheter year indicates that this complication occurs several times in a minority of patients, and that many patients remain sepsis free. Two studies have a much higher rate of catheter sepsis; these rates are probably explained by one patient group being immunosuppressed and another group including an excess of paediatric patients.

The weighted average rate for catheter occlusion was 0.071 ( $95 \% \mathrm{CI}, 0.059,0.083$ ) episodes per catheter year (see Figure 2). Catheter occlusion might be caused by faulty catheter care or by an inappropriate infusion regimen. Central lines which can not be cleared by thrombolysis require removal and replacement.

The overall rate for central vein thrombosis was 0.027 ( $95 \% \mathrm{CI}, 0.02,0.034$ ) episodes per catheter year (see Figure 3). Thrombosis is associated with difficulties with venous access in the future, and may also be related to an inappropriate infusion regimen or faulty catheter placement.

TABLE 7 Summary of studies relevant to Section 2: the experience of patients on HPN

| Study: Country (Time frame) | Sample size (A/P) | Inception cohort | \% of HPN pop inc. | Study design | lost <br> to followup | Complications reported | Quality of life measured | Survival measured (months) | Duration <br> of HPN use measured (months) | Reported reason for stopping HPN |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| August, 1991: <br> USA (1980-89) | $17$ (A) | yes |  | Case series; retrospective | 0 | no | no | yes | yes | yes |
| Beers, I990: <br> USA (1975-88) | $\begin{aligned} & 107 \\ & (\mathrm{~A}) \end{aligned}$ | yes | $100$ | Case series; retrospective | 0 | yes | no | no | yes | no |
| Bisset, I992: <br> UK (1987-92) | 10 <br> (P) | yes | $100$ | Case series; retrospective | 0 | yes | no | no | no | yes |
| Bowyer, 1985: USA (1975-82) | $\begin{gathered} 9 \\ (\mathrm{~A}) \end{gathered}$ | yes |  | Case series; retrospective | $?$ | yes | no | no | yes | no |
| Buchman, I993: USA (?) | 41 <br> (A) | yes |  | Case series; retrospective | ? | yes | no | no | no | no |
| Buchman, I993: USA (15 years) | 33 <br> (A) | yes |  | Case series; retrospective | 0 | yes | no | no | no | no |
| Buchman, 1994(b) USA (I973-91) | $\begin{gathered} 527 \\ (A / P) \end{gathered}$ | yes | ? | Case series; retrospective | ? | yes | no | no | yes | no |
| Buchman, 1994(a): <br> USA (I973-9I) | $\begin{aligned} & 527 \\ & (A / P) \end{aligned}$ | yes | ? | Case series; retrospective | $2 \%$ of inf. | yes | no | no | yes | no |
| Burnes, 1992: <br> USA (1986-89) | 63 <br> (A) | yes |  | Case series; retrospective | ? | yes | no | no | no | yes |
| Byrne, 1979: <br> USA (?) | $\begin{gathered} 106 \\ (\mathrm{~A} / \mathrm{P}) \end{gathered}$ | yes |  | Case series; retrospective | ? | yes | no | no | no | yes |
| Carlson, 1996: UK (I992) | 73 <br> (A) | no | 93 | Q-o-L <br> interviews; retrospective | 0 | no | P | no | no | no |
| DePotter, 1992: <br> France (1981-90) | 156 <br> (P) | yes |  | Case series; retrospective | ? | yes | no | no | yes | yes |
| Detsky, I986: <br> Canada (1970-82) | $\begin{array}{r} 37 \\ 2) \quad(\mathrm{A}) \end{array}$ | yes | 51 | Q-o-L interviews; prospective | $49$ | no | P | no | no | no |
| Dollery, 1994: <br> UK (1983-93) | 34 <br> (P) | yes |  | Case series; retrospective | $?$ | yes | no | yes | no | no |
| Duclaux, I993: <br> France (?) | 44 <br> (P) | yes | ? | Case series; retrospective |  | no | P | no | no | no |
| Dudrick, 1984: <br> USA (1974-83) | $\begin{gathered} 133 \\ (\mathrm{~A} / \mathrm{P}) \end{gathered}$ | yes |  | Case series; retrospective |  | yes | no | no | yes | no |
| Foldes, 1990: USA (19 months) | 10 <br> (A) | yes | ? | Case series; retrospective | $100$ | yes | no | no | no | no |
| Galandiuk, 1990: USA (1976-87) | $\begin{aligned} & 39 \\ & \text { (A) } \end{aligned}$ | yes | $21$ | Case series; retrospective | $?$ | yes | P | no | yes | yes |
| Gouttebel, I987: <br> France (1979-85) | 85 <br> (A) | yes | ? | Case series; retrospective | ? | yes | no | no | yes | yes continued |
| ?, Not stated or not known; Q-o-L, quality of life; A, adult; P, paediatric; p, patient centred assessment of Q-o-L; f, functional assessment; E, estimated. |  |  |  |  |  |  |  |  |  |  |

TABLE 7 contd Summary of studies relevant to Section 2: the experience of patients on HPN

| Study: <br> Country <br> (Time frame) | Sample size (A/P) | Inception cohort | \% of <br> HPN <br> pop inc. | Study design | \% <br> lost <br> to followup | Complications reported | Quality of life measured | Survival measured (months) | Duration <br> d of HPN <br> use measured (months) | Reported reason for stopping HPN |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Grabowski, I989: USA (?) | $\begin{gathered} 4 \\ (\mathrm{~A}) \end{gathered}$ | yes |  | Case series; retrospective | 0 | no | no | yes | yes | yes |
| Griffith, 1984: UK (1978-83) | $\begin{gathered} 9 \\ (\mathrm{~A}) \end{gathered}$ | yes |  | Case series; retrospective | 0 | yes | no | no | yes | yes |
| Herfindal, 1992: <br> USA (1980-87) | $\begin{gathered} 56 \\ (\mathrm{~A} / \mathrm{P}) \end{gathered}$ | yes |  | Case series; retrospective | 10 | yes | no | no | yes | no |
| Herfindal, I989: USA (?) | $\begin{aligned} & 347 \\ & \text { (A) } \end{aligned}$ | no |  | Q-o-L interviews; prospective | ? | no | P | no | no | no |
| Howard, 1993: USA (1985-90) | $2275$ <br> (A) | mix | $10 \text { (E) }$ | Case series; retrospective | ? | yes | no | yes | no | yes |
| Howard, I991: USA (1984-87) | 2916 <br> (A) | mix | $7.9 \text { (E) }$ | Case series; retrospective | 7 | yes | no | yes | no | yes |
| Howard, I986: USA (1983-85) | $\begin{gathered} 2550 \\ \text { (A) } \end{gathered}$ | mix |  | Case series; retrospective | ? | yes | no | yes | yes | yes |
| Howard et al, 1995: USA <br> (1985-92) | $9288$ <br> (A) | mix | $5 \text { (E) }$ | Case series; retrospective | ? | yes | no | yes | no | yes |
| Hurley, 1990: USA (?) | $\begin{aligned} & 23 \\ & (\mathrm{~A}) \end{aligned}$ | yes |  | Case series; retrospective | ? | yes | no | no | no | no |
| Johnston, I994: <br> UK (1980-93) | $34$ <br> (A) | yes |  | Case series; retrospective | 0 | yes | no | no | no | no |
| Johnston, 1993: <br> Scotland <br> (1980-92) | $\begin{gathered} 30 \\ (\mathrm{~A} / \mathrm{P}) \end{gathered}$ | yes |  | Case series; retrospective | 0 | yes | no | no | yes | yes |
| $\begin{aligned} & \text { King, 1993: } \\ & \text { USA (1981-90) } \end{aligned}$ | $\begin{aligned} & 61 \\ & (\mathrm{~A}) \end{aligned}$ | yes |  | Case series; retrospective | ? | yes | P | yes | yes | yes |
| Ladefoged, 198I: <br> Denmark <br> (1978-79) | $\begin{aligned} & 13 \\ & \text { (A) } \end{aligned}$ | yes |  | Q-o-L interviews; prospective | 0 | no | P | no | no | no |
| Manji, I989: <br> USA (1989) | $\begin{gathered} 5 \\ (\mathrm{~A}) \end{gathered}$ | yes |  | Case series; retrospective | 0 | yes | no | no | yes | no |
| Mercier, I995: <br> Canada (1992-95) | 16 $(?)$ | yes |  | Case series; retrospective | ? | no | no | no | no | yes |
| Messing, 1995: <br> France/Belgium <br> (1980-89) | $217$ <br> (A) | yes |  | Case series; retrospective | 0 | no | no | yes | yes | yes |
| Messing, I989: <br> Europe (1974-85) | $194$ (A) | yes |  | Case series; retrospective |  | yes | f | no | yes | yes |
| $\begin{aligned} & \text { Miller 1979: } \\ & \text { USA (1970-78) } \end{aligned}$ | $\begin{aligned} & 10 \\ & \text { (A) } \end{aligned}$ | yes |  | Case series; retrospective | 0 | no | no | no | yes | yes <br> continued |
| ?, Not stated or not known; Q-o-L, quality of life; A, adult; P, paediatric; p, patient centred assessment of Q-o-L; f, functional assessment; E, estimated. |  |  |  |  |  |  |  |  |  |  |

TABLE 7 contd Summary of studies relevant to Section 2: the experience of patients on HPN

| Study: <br> Country <br> (Time frame) | Sample size (A/P) | Inception cohort | \% of <br> HPN <br> pop <br> inc. | Study design | \% <br> lost <br> to follow- <br> up | Complications reported | Quality of life measured | Survival measured (months) | Duration d of HPN use measured (months) | Reported reason for stopping HPN |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Mukau, I992: <br> USA (1988-90) | $\begin{aligned} & 50 \\ & \text { (A) } \end{aligned}$ | yes |  | Case series; retrospective | ? | yes | no | no | no | no |
| Nightingale, 1995: <br> UK (1984-92) | 17 <br> (A) | no |  | Case series; retrospective | 0 | yes | no | no | no | no |
| $\begin{aligned} & \text { O'Hanrahan, } \\ & \text { I992: UK } \\ & \text { (1977-91) } \end{aligned}$ | $\begin{gathered} 400 \\ (\mathrm{~A} / \mathrm{P}) \end{gathered}$ | yes |  | Case series; retrospective | ? | yes | f | no | yes | yes |
| Perl, I98I: USA (I year) | 10 <br> (A) | no |  | Case series; prospective | 0 | yes | no | no | no | no |
| Pironi, I993: <br> Italy (1986-93) | $\begin{aligned} & 18 \\ & \text { (A) } \end{aligned}$ | yes |  | Case series; retrospective | 0 | yes | f | no | yes | yes |
| Ralston, 1984: <br> USA (1977-82) | $\begin{gathered} 9 \\ (\mathrm{P}) \end{gathered}$ | yes |  | Case series; retrospective | 33 | yes | no | no | no | yes |
| Richards, I995: UK (I995) | $\begin{aligned} & 51 \\ & \text { (A) } \end{aligned}$ | no |  | Q-o-L interviews; prospective | 0 | no | P | no | no | no |
| Robb, I983: USA (?) | $\begin{aligned} & 42 \\ & \text { (A) } \end{aligned}$ | yes |  | Q-o-L interviews; prospective | ? | yes | yes | no | yes | no |
| Roslyn, 1983: <br> USA (1976-80) | $128$ <br> (A) | yes |  | Case series; retrospective | ? | yes | no | no | no | no |
| Schmidt- <br> Sommerfeld, 1990 <br> USA (1980-85) | $\begin{aligned} & 35 \\ & \text { (P) } \end{aligned}$ | yes |  | Case series retrospective | ? | yes | no | no | yes | yes |
| Shike, 1980: USA (?) | $\begin{aligned} & 16 \\ & \text { (A) } \end{aligned}$ | yes |  | Case series; prospective | 0 | yes | no | no | no | no |
| Shike, 1986: USA (?) | $\begin{aligned} & 12 \\ & \text { (A) } \end{aligned}$ | yes | 57 | Case series; prospective | 0 | yes | no | no | no | no |
| Singer, I991: <br> USA (1987-88) | $\begin{aligned} & 22 \\ & \text { (A) } \end{aligned}$ | yes |  | Case series; retrospective | 9 | yes | no | no | yes | yes |
| Smith, 1993: <br> USA (?) | $\begin{aligned} & 116 \\ & \text { (A) } \end{aligned}$ | no |  | Q-o-L interviews; prospective | ? | no | P | no | no | no |
| Staun USA (?) | $\begin{aligned} & 15 \\ & \text { (A) } \end{aligned}$ | yes |  | Case series; prospective | ? | yes | no | no | no | no |
| Steiger, 1983: <br> USA (I976-8I) | $\begin{aligned} & 39 \\ & \text { (A) } \end{aligned}$ | yes | 78 | Case series; retrospective | ? | yes | no | no | yes | yes |
| Van Gossum, 1995: Europe (1993-94) | $211$ <br> (A) | yes | $80 \text { (E) }$ | Case series; prospective | ? | no | no | yes | no | yes |
| Vargas, 1987: <br> USA (1976-86) | $\begin{aligned} & 102 \\ & (\mathrm{P}) \end{aligned}$ | yes |  | Case series; retrospective | ? | yes | no | no | yes | yes |
| Weiss, 1982: <br> USA (1978-80) | $\begin{gathered} 9 \\ (\mathrm{~A}) \end{gathered}$ | yes |  | Case series; retrospective | 0 | yes | no | no | yes | yes |

?, Not stated or not known; Q-o-L, quality of life; A, adult; P, paediatric; p, patient centred assessment of Q-o-L; f, functional assessment; $E$, estimated.

TABLE 8 Complications of HPN (episodes per catheter year, unless indicated)

| Study | Catheter sepsis (95\% CI) | Catheter sepsis (patients per catheter year) $(95 \% \mathrm{Cl})$ | Catheter occlusion (95\% CI) | Central vein thrombosis (95\% CI) | Liver/ biliary problems (95\% CI) | Metabolic bone disease (95\% CI) | Other |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Beers, 1990 | - |  | - | $\begin{gathered} 0.04 \\ (0.02,0.07) \end{gathered}$ | - | - | - |
| Bisset, 1992 | - |  | - | - | - | - | Sepsis 0.73 |
| Bowyer, 1985 | - |  | - | - | $15 \%(7 \%, 27 \%)$ <br> liver problems. $\begin{gathered} 3 \%(0.4 \%, 12 \%) \\ \text { deaths } \end{gathered}$ | - | - |
| Buchman, 1994(a) | - |  | $\begin{gathered} 0.07 \\ (0.06,0.09) \end{gathered}$ | $\begin{gathered} 0.02 \\ (0.01,0.03) \end{gathered}$ | - | - | - |
| Buchman, I994(b) | $\begin{gathered} 0.23 \\ (0.2,0.27) \end{gathered}$ <br> not possible to calculate rates for children | $\begin{gathered} 0.23 \\ (0.2,0.26) \end{gathered}$ | - | - | - | - | - |
| Buchman, 1993 |  |  |  |  |  |  | Low plasma-free choline levels are prevalent, associated with elevated serum aminotransferases |
| Buchman, 1993 |  |  |  |  |  |  | Fall in renal function of $3.5 \pm 6.3 \%$ per year |
| Burnes, 1992 | $\begin{gathered} 0.27 \\ (0.2,0.35) \end{gathered}$ |  | - | - | - | - | - |
| DePotter, 1992 | $\begin{gathered} 0.40 \\ (0.33,0.49) \end{gathered}$ | 0.23 | $\begin{gathered} 0.04 \\ (0.02,0.07) \end{gathered}$ | - | $\begin{gathered} 0.03 \\ (0.01,0.05) \end{gathered}$ | - | - |
| Dollery, 1994 | - |  | - | 16 episodes of major thrombosis in 12 of 34 patients | - | - | - |
| Dudrick, 1984 | $\begin{gathered} 4.39 \\ (0.26,0.54) \end{gathered}$ | $\begin{gathered} 0.15 \\ (0.08,0.26) \end{gathered}$ | - | - | - | - | - |
| Foldes, 1990 | - |  | - | - | - | $\begin{gathered} 90 \% \\ (56 \%, 99 \%) \end{gathered}$ | - |
| Galandiuk, 1990 | $\begin{gathered} 0.27 \\ (0.19,0.38) \end{gathered}$ |  | - | - | - | - | - |
| Gouttebel, 1987 | $\begin{gathered} 0.70 \\ (0.49,0.97) \end{gathered}$ | $\begin{gathered} 0.42 \\ (0.26,0.4) \end{gathered}$ | - | - | - | - | continued |

TABLE 8 contd Complications of HPN (episodes per catheter year, unless indicated)

| Study | Catheter sepsis (95\% CI) | Catheter sepsis (patients per catheter year) <br> (95\% CI) | Catheter occlusion (95\% CI) | Central vein thrombosis (95\% CI) | Liver/ biliary problems (95\% CI) | Metabolic bone disease (95\% CI) | Other |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Herfindal, 1992 | $\begin{gathered} 0.46 \\ (0.3,0.7) \end{gathered}$ | $\begin{gathered} 0.14 \\ (0.06,0.28) \end{gathered}$ | $\begin{gathered} 0.22 \\ (0.12,0.39) \end{gathered}$ | - | $\begin{gathered} 0.42 \\ (0.27,0.63) \end{gathered}$ | $\begin{gathered} 0.05 \\ (0.01,0.15) \end{gathered}$ | Metabolic complications 0.61 |
| Howard, 1993 |  |  | - | - | - |  | Total complication rate is higher for those under 18 years |
| Howard, 1986 | $\begin{gathered} 6.37 \\ (0.33,0.42) \end{gathered}$ |  | - | - | - | $\begin{gathered} 0.013 \\ (0.005,0.025) \end{gathered}$ |  |
| Hurley, 1990 | $\begin{gathered} 0.30 \\ (0.17,0.49) \end{gathered}$ |  | $\begin{gathered} 0.20 \\ (0.06,0.47) \end{gathered}$ | - | - |  | Total complications <br> Cancer $2.22(1.4,3.4)$ <br> Benign 0.89 ( $0.64, \mathrm{I} .2$ ) <br> $p<0.01$ |
| Johnston, 1993 | $\begin{gathered} 30.16 \\ (0.05,0.47) \end{gathered}$ |  | - | $\begin{gathered} 0.28 \\ (0.15,0.47) \end{gathered}$ | - | - | - |
| King, 1993 | $\begin{gathered} 0.54 \\ (0.22, \mathrm{I} . \mathrm{II}) \end{gathered}$ | ? | - | - | - | - | - |
| Manji, 1989 | - |  | - | - | Symptomatic gallstones in 100\% | - | - |
| Messing, 1989 | $\begin{gathered} 0.38 \\ (0.30,0.48) \end{gathered}$ | ? | 0.18 | 0.07 | - | - | - |
| Mukau, 1992 | $\begin{gathered} 0.2 \\ (0.1,0.35) \end{gathered}$ | ? | - | $\begin{gathered} 0.07 \\ (0.02,0.17) \end{gathered}$ | - | - | - |
| Nightingale, 1995 | 24 fungal infections. Total no of ines not given |  | - | - | - | - | Four developed eye infections. Two had recurrent infection |
| O'Hanrahan, 1992 | $\begin{gathered} 0.47 \\ (0.38,0.58) \end{gathered}$ | ? | $\begin{gathered} 0.44 \\ (0.36,0.55) \end{gathered}$ | $\begin{gathered} 0.06 \\ (0.03,0.11) \end{gathered}$ | - | - | $\begin{gathered} \text { Metabolic } \\ \text { complications } \\ 0.12(0.08,0.18) \end{gathered}$ |
| Perl, 1981 | - |  | - | - | - | - | Depression 80\% <br> $(44,98)$ |
| Pironi, 1993 | $\begin{gathered} 0.12 \\ (0.03,0.3) \end{gathered}$ |  | $\begin{gathered} 0.03 \\ (0,0.16) \end{gathered}$ | $\begin{gathered} 0.09 \\ (0.02,0.26) \end{gathered}$ | $\begin{gathered} 0.15 \\ (0.05,0.34) \end{gathered}$ | - | - |
| Robb, 1983 | $\begin{gathered} 0.42 \\ (0.25,0.68) \end{gathered}$ |  | $\begin{gathered} 0.1 \\ (0.03,0.24) \end{gathered}$ | - | - | - | - |
| Roslyn, I983 | - |  | - | - | $\begin{gathered} \text { Symptomatic } \\ \text { gallstones in } \\ 23 \%(15 \%, 32 \%) \end{gathered}$ | - | continued |
| ?, Data in the study not sufficient for calculation of rates;A, AIDS; C, cancer; H, HPN. |  |  |  |  |  |  |  |

TABLE 8 contd Complications of HPN (episodes per catheter year, unless indicated)

| Study | $\begin{aligned} & \text { Catheter } \\ & \text { sepsis } \\ & (95 \% \mathrm{CI}) \end{aligned}$ | Catheter sepsis (patients per catheter year) (95\% CI) | Catheter occlusion (95\% CI) | Central vein thrombosis (95\% CI) | Liver/ biliary problems (95\% CI) | Metabolic bone disease (95\% CI) | Other |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Schmidt- <br> Sommerfeld, $1990$ | $\begin{gathered} 0.71 \\ (0.5,0.97) \end{gathered}$ |  | $\begin{gathered} 0.29 \\ (0.17,0.47) \end{gathered}$ | $\begin{gathered} 0.07 \\ (0.02,0.19) \end{gathered}$ | - | - | - |
| Shike, 1986 | - |  | - | - | - | $\begin{gathered} 67 \% \\ (35 \%, 90 \%) \end{gathered}$ | - |
| Shike, 1980 | - |  | - | - | - | $\begin{gathered} 75 \% \\ (48 \%, 93 \%) \end{gathered}$ | - |
| Singer, 1991 | $\begin{gathered} \text { A } 0.43 \\ (0.05,1.55) \\ \text { C } 0.2 \\ (0.07,1.43) \\ \text { H } 0.1 \\ (0.04,0.22) \end{gathered}$ |  | $\begin{gathered} \text { A } 0.21 \\ (0.05,1.2) \\ \text { C } 0.03 \\ (0,0.18) \\ \text { H } 0.06 \\ (0.02,0.16) \end{gathered}$ | - | - | - | Metabolic disturbance A $0.43(0.05,1.55)$ C $0.49(0.28,0.81)$ H $0.17(0.08,0.3)$ |
| Staun | - |  | - | - | - | 4\% decrease in bone mineral content per year | - - |
| Steiger, 1983 | - |  | - | - | - |  | \% of hospitalised days <br> Crohn's 24\% <br> MVD 58\% <br> Radiation enteritis 13\% |
| Vargas, 1987 | $\begin{gathered} 0.37 \\ (0.29,0.46) \end{gathered}$ | $\begin{gathered} 0.20 \\ (0.15,0.28) \end{gathered}$ | - |  | $\begin{gathered} \text { Any } 0.06 \\ (0.03,0.1) \\ \text { Severe } 0.024 \\ (0.008,0.057) \end{gathered}$ | - | - |
| Weiss, 1982 | $\begin{gathered} 0.2 \\ (0.01,1.1 \mathrm{I}) \end{gathered}$ |  | $\begin{gathered} 0.2 \\ (0.01, \mathrm{I} .1 \mathrm{I}) \end{gathered}$ |  | - | - | - |
| ?, Data in the study not sufficient for calculation of rates;A,AIDS; C, cancer; H, HPN. |  |  |  |  |  |  |  |

The incidence of liver abnormalities was difficult to assess from the evidence located. Severe problems were rare ( 0.025 episodes/catheter year: Vargas, 1987) but minor abnormalities are probably very common. To some extent the incidence of liver abnormalities will vary according to how thoroughly the patient is investigated.

Metabolic bone disease can cause severe, debilitating illness but incidence rates are difficult to determine. Metabolic bone disease in mild forms is probably very common if it is looked for carefully.

Metabolic complications such as fluid and electrolyte imbalance are also probably very common.
are probably not recorded. The studies we located rarely gave details of more severe abnormalities. When metabolic problems were reported they were fairly common (range, 0.12-0.61 episodes/ catheter year).

There were many other rare complications of HPN, which were usually reported as interesting cases. These were wide-ranging and often related to catheter sepsis, such as subacute bacterial endocarditis, septic thromboembolism, or candida endopthalmitis. Depression has been commonly reported, and is one of a range of psychiatric illnesses that have been noted in HPN patients. The severity of the underlying condition and the dependence on a machine are amongst the commonest underlying causes of depression.


FIGURE I Incidence of catheter sepsis

Renal function is thought to deteriorate by approximately $5 \%$ per year. There is usually a multifac-torial pathophysiology behind the deterioration, hence the need to avoid dehydration and nephrotoxic drugs and to monitor renal function on a regular basis.

Only one study (Hurley, 1990) described the differences in total complication rates between a group of patients with malignant disease and a benign group. There were an excess of complications in the malignant group ( 2.2 versus 0.9 events/catheter year).

Quality of life (see Table 9) was measured using validated instruments in five studies. Index scores on a $0-1$ scale ( $0=$ death, $1=$ best possible) varied from 0.51 to 0.73 . Detsky (1995) recently admitted that the score he produced in 1986 (0.73) was probably not correct because the methods he used to measure quality of life overestimated the true
value, which was somewhat less. The best quality of life was seen with young patients, longer duration of treatment, high self-esteem, a good relationship with a partner, wealth and employment. The worst quality of life was experienced by patients who were older, addicted to narcotics, poor, single, unemployed, had a short duration of treatment and few family coping skills.

Functional assessments have been used to assess outcomes in several series and they are estimated by the physician. The best functional outcomes are seen in younger patients with Crohn's disease. The worst functional outcomes are seen in older patients with an underlying malignancy, pseudoobstruction or Crohn's disease. Only one study (Galandiuk, 1990) has examined quality of life before and after HPN was started. This study, which was carried out only on patients with Crohn's disease, showed that quality of life improved on HPN.


FIGURE 2 Incidence of catheter occlusions


FIGURE 3 Incidence of central vein thrombosis

Survival on HPN (Table 10) is best for patients with benign disease. Of the subgroups, Crohn's disease seems to have the best survival rate, with several series reporting a better than $90 \%$ 1 -year survival. The reported 1-year survival for HPN patients with malignant disease varies from $15 \%$ to $30 \%$. Carcinoma of the ovary seems to have a particularly poor outcome, with one study measuring the mean survival as only 30 days.

Patients with AIDS have a particularly poor outcome, with only $7-12 \%$ surviving 1 year. This statement on survival obviously reflects the natural history of the underlying disease and is not meant to reflect the effectiveness of HPN therapy. It does serve to illustrate that as patients with malignant disease and AIDS usually only survive for short periods, patient selection for HPN is very important.

TABLE 9 Quality of life on HPN

| Study | Whose values? | Instrument used | Profile or index | Index scores | Best <br> Q-o-L or outcome | Worst Q-o-L or outcome | Comments |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Carlson, 1995 | Patient | Non-validated questionnaire | Index | $\begin{gathered} 0.64 \\ (0-1 \text { scale } \end{gathered}$ | - | - | Q-o-L independent of variables tested. Younger patients keen on intestinal transplantation. |
| Detsky, 1986 | Patient | Category scaling, time trade-off + one other | Index | $\begin{gathered} 0.73 \\ \left(0-1 \text { scale }^{* *}\right) \end{gathered}$ | Scores improve with time and peak at 4-5 years | Lowest scores seen in the first year of HPN | Scores were measured for 37 and estimated for 36 . No subgroup analysis was performed. |
| Duclaux, 1993 | Doctor | Non-validated simple questionnaire | Profile |  |  |  | Q-o-L much improved at home. Development and psychological well-being much improved. |
| Galandiuk, 1990 | Patient plus doctor | Q-o-L score, social activity score, psychological score | Index | Pre-HPN, 7.1 On HPN, $5.3^{\#}$ | - | - | Index scores were better on HPN; pre-HPN Q-o-L was significantly worse ( $p<0.0 \mathrm{I}$ ). All patients in this study had Crohn's disease. |
| Herfindal, 1989 | Patient | Multiplevalidated instruments | Profile |  | Long duration (> 6 months) | Duration $<6$ months | HPN patients had lower (worse) scores than renal transplant recipients and normal US population. |
| King, 1993 | Doctor | Q-o-L assessed by retrospective case note review | Profile |  | - | - | All patients with gynaecological malignancy. Improvements noted in pain, vomiting, fatigue, morale and social interactions ( $p<0.05$ ) compared with pre-HPN status. |
| Ladefoged, \|98| | Patient | Non-validated questionnaire | Profile | - | Acceptable <br> in $2 / 3$ <br> of cases | - | Q-o-L parameters were independent of all variables. BUT, not enough data to test. |
| Messing, 1989 | Doctor | Functional assessment | - | - | $\begin{aligned} & \text { Age }<65 \\ & \text { years, } \\ & \text { benign } \end{aligned}$ | Age > 65 <br> years, <br> malignancy, <br> pseudo- <br> obstruction | Simple 4-stage rehabilitation profile. Stage decided by physician, not the patient. |
| O'Hanrahan, 1992 | Doctor | Functional assessment | Profile | - | Crohn's disease | All other diagnostic groups | Data overlap with Messing (1989). <br> Same 4-point scale used. |
| Pironi, 1993 | Doctor | Functional assessment | Profile | - | - | - | Same 4-point scale as Messing and O'Hanrahan. Two-thirds in the upper two groups. |

\# Scale 3 to 9,9 severe disablement and $3=$ best possible $Q$-o-L. ** Scale 0 to $I, 0=$ death, $I=$ best possible $Q$-o-L.

TABLE 9 contd Quality of life on HPN

| Study | Whose <br> values? <br> Instrument | Profile <br> or index | Index <br> scores | Best <br> Q-o-L or <br> outcome | Worst <br> Q-o-L or <br> outcome | Comments |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |

TABLE 10 Survival on HPN

| Study | Benign underlying disease | Malignant underlying disease (including AIDS) |
| :---: | :---: | :---: |
| August, 1991 | - | Average months survived. Cancer ovary 1.3 <br>  Cancer colon 3 <br>  Cancer appendix 6 |
| Grabowski, 1989 | Scleroderma. 3 (of 4) died at $12,14,17$ months. | - |
| Howard, 1991 | I-year mortality rates:  <br> Crohn's disease $5 \%$ <br> MVO $20 \%$ (4\% thereafter) <br> Pseudo-obstruction $20 \%$ | I-year mortalityrates: <br> Cancer$\quad 75 \%$  <br> AIDS $93 \%$ |
| Howard, 1993 | I-year survival:  <br> Crohn's disease $95 \%$ <br> Radiation enteritis $76 \%$ | I-year survival: <br> Cancer 30\% |
| Howard, 1986 | $50 \%$ survival at 36 months $15 \%$ survival at year 8 | $50 \%$ survival at 6 months $15 \%$ at I year. All dead by 23 months |
| Howard, 1995 | I-year survival >90\% (age 0-55) <br> I-year survival $\sim 65 \%$ (age $>55$ ) |  |
| King, 1993 | - | Gynaecological malignancy <br> Median survival 2 months (range 0-26) |
| Messing, 1995 | 1-year survival $91 \%$ <br> 2-year survival $70 \%$ <br> 3-year survival $62 \%$ | - |
| Van Gossum, 1995 | 6-month mortality rates:  <br> Crohn's disease $0 \%$ <br> MVD $8 \%$ <br> Miscellaneous $13 \%$ <br> Radiation enteritis $7 \%$ | 6-month mortality rates:  <br> Cancer $71 \%$ <br> AIDS $88 \%$ |

The duration of HPN use (Table 11 and Figure 4) reflects the survival and the disease activity. In Crohn's disease there are two peaks; the first is from 0 to 6 months, and reflects the use of HPN
exacerbation of the disease. Longer duration of use (more than 2 years) reflects the use of HPN for established short bowel syndrome. More than $50 \%$ of Crohn's patients are on HPN for more than 2 years. The duration of use in malignant disease is

TABLE 11 Duration of HPN use

| Study | Duration of HPN use (by diag | nostic subgroup where possible) | Length of follow-up |
| :---: | :---: | :---: | :---: |
| Beers, 1990 | 3.54 years | (all benign) | 6 years |
| Bowyer, 1985 | 8-95 months | (all benign) | 7 years |
| DePotter, 1992 | 615 days (range 30-3532 days) | (all benign) | 9 years |
| Galandiuk, 1990 | 1083 days (range 33-3258) | (all Crohn's) $51 \%$ on HPN for > 2 years | 11 years |
| Grabowski, 1989 | 12-86 months | (all scleroderma) | ? |
| Griffith, 1984 | Average 9.6 months | (all benign) | 5 years |
| Messing, 1995 | Median 19 months (range I-137) | (all benign) | 9 years |
| Pironi, 1993 | $22 \pm 22$ months | (all benign) | 7 years |
| Robb, 1983 | 42.7 months (range 6-114) | (all benign) | ? |
| Schmidt-Sommerfeld, 1990 | 577 days (range 58-2633) | (all benign) | 5 years |
| Steiger, 1983 | Crohn's disease MVD <br> Radiation enteritis | $\begin{aligned} & 842 \text { days (mean) } \\ & 884 \text { days } \\ & 494 \text { days } \end{aligned}$ | 5 years |
| Miller, 1979 | 15.7 months (range I-52) | (all radiation enteritis) | 8 years |
| Gouttebel, 1987 | Benign group Malignant group | $\begin{array}{ll} 357 \text { days } & (30-4155) \\ 93 \text { days } & (30-421) \end{array}$ | 6 years |
| August, 1991 | 53 days (range 5-208) | (malignant) | 9 years |
| King, 1993 | 75 days (range 2-414; median 28) | (malignant) | 9 years |
| Singer, 1991 | 2.5 months (range 0.7-10.8) | (all AIDS) | 1 year |
| Weiss, 1982 | 6.2 months (range 0.5-19) | (malignant) | 2 years |



FIGURE 4 Mean duration of HPN use (■, malignant; $\square$, benign)
generally very short, with only a minority of patients continuing on HPN for more than 1 year. More accurate analysis of this data is impossible because the reported duration of HPN use is affected by the length of the study follow-up. For example, a short period of observation would show more patients continuing treatment and survivors than a long study.

It is difficult to generate accurate figures from the available evidence concerning the reasons for
discontinuing HPN (Table 12). This is because there are wide variations in the follow-up times; we have, therefore, only included studies which compare different disease subgroups. However, it is clear that the main reason for discontinuing HPN for patients with malignant disease is death (range, 60-100\%). The patients with benign disease tend to either recover (range, 40-70\%) or continue on HPN for long periods (range, $25-50 \%$ ); only a minority are discontinued because of death (range, $2-30 \%$ ).

TABLE 12 Reasons for discontinuing HPN
The length of follow-up in each study varied considerably (see Table 7), and it was not, therefore, possible to compare individual studies. The studies below reported reasons for discontinuing HPN for subgroups as indicated; studies which did not report subgroup outcomes were not included.

| Study | Diagnosis | Still on HPN <br> (\%) | Recovered <br> (\%) | Dead <br> (\%) | Lost <br> (\%) |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Byrne, 1979 | Benign | 45 | 39 | 16 | 0 |
|  | Cancer | 0 | 0 | 100 | 0 |
| Gouttebel, 1987 | Crohn's disease | ? | ? | 28 | ? |
|  | Cancer | 9 | 0 | 91 | 0 |
| Howard, 1993 | Crohn's disease | 25 | 70 | < 5 | ? |
|  | MVD | 50 | 25 | 25 |  |
|  | Cancer | 10 | 25 | 65 |  |
| Howard, 1991 | Crohn's disease | 47 | 38 | 5 | 7 |
|  | MVD | ? | 16 | 20 |  |
|  | Pseudo-obstruction | 48 | 21 | 20 |  |
|  | Cancer | 25 | ? | 75 |  |
|  | AIDS | ? | ? | 93 |  |
| Howard, 1986 | Crohn's disease | ? | ? | 28 | ? |
|  | Cancer | ? | ? | 100 | ? |
| Howard, 1995 | Crohn's disease | 25 | 70 | 2 |  |
|  | MVD | 48 | 27 | 19 |  |
|  | Cancer | 8 | 26 | 63 |  |
|  | AIDS | 6 | 13 | 73 |  |
| Messing, 1989 | Crohn's disease | 35 | 57 | 8 | ? |
|  | MVD | 32 | 40 | 28 |  |
|  | Pseudo-obstruction | 20 | 40 | 40 |  |
|  | Cancer | 26 | 13 | 60 |  |
|  | Radiation enteritis | 26 | 26 | 47 |  |
| Steiger, 1983 | Crohn's disease | ? | ? | 26 | ? |
|  | MVD |  |  | 43 |  |
|  | Radiation enteritis |  |  | 38 |  |
| Van Gossum, 1995 | Crohn's disease | 75 | 25 | 0 | ? |
|  | MVD | 65 | 27 | 7 |  |
|  | Cancer | 14 | 15 | 71 |  |
|  | AIDS | 0 | 12 | 88 |  |
|  | Radiation enteritis | 81 | 12 | 6 |  |
| Vargas, 1987 | Benign | 21 | 50 | 30 | 0 |
|  | Cancer | 0 | 0 | 100 |  |

TABLE 13 Summary of studies selected for Section 3 How have HPN programmes been organised? What comparative evidence is available on effectiveness?

| Studies - <br> Section 3a | Time frame | Study design | Groups compared | Sample size | Comparable groups | Length of follow-up | \% lost to follow-up | Outcomes assessed? |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Van Gossum, 1995 | $\begin{aligned} & 1993- \\ & 94 \end{aligned}$ | Case series; retrospective | Different countries | 211 | N/A | 6-12 months | ? | Organisation, survival, catheter type |
| Studies - <br> Section 3b | Time frame | Study design/ Grade of evidence | Groups compared | Sample size | Comparable groups | Length of follow-up | \% lost to follow-up | Outcomes assessed? |
| Howard, 1989 | $\begin{aligned} & 1983- \\ & 88 \end{aligned}$ | Case series retrospective Grade C | Reservoir vs external catheter | 58 | No | I month4 years | ? | Catheter complications |
| Hyltander, I99I | 10 weeks | Randomised controlled trial; prospective Grade R | HPN vs no HPN during chemotherapy | 33 | Yes | 10 weeks | 0 | Nutritional status |
| Jarrard, 1980 | 1978 | Prospective controlled, nonrandomised; Grade C | Daily vs alternate day dressings | 38 | No | \| $\|-3\|$ days | ? | Catheter colonisation |
| Johnston, I994 | $\begin{aligned} & 1980- \\ & 93 \end{aligned}$ | Case series; retrospective Grade H | Effect of unit experience on complication rate | 34 | N/A | 1.7 years | 0 | Complication rates |
| Pithie, I988 | $\begin{aligned} & 6 \text { years } \\ & \text { (1980s) } \end{aligned}$ | Case series; retrospective Grade C | Catheter tip position | 69 | N/A | NS | 0 | Superior vena cava thrombosis |
| Rannem, I990 | $\begin{aligned} & 1976- \\ & 88 \end{aligned}$ | Case series; retrospective Grade H | Before and after use of disinfectants | $58$ | $\mathrm{N} / \mathrm{A}$ | 2 months11.5 years | ? | Catheter sepsis |
| N/A, not applicable; ?, not known or not stated; Grade R, evidence from randomised comparison; Grade $C$, evidence from concurrent non-randomised comparison; Grade H, evidence from historical non-randomised comparison. |  |  |  |  |  |  |  |  |

## Results for Section 3a and b Organisation and comparative data

## Organisation

Only one study has reported data on the organisation of HPN programmes in different countries (Van Gossum, 1996) (see Table 13). The study was a multicentre collection of all new patients commenced on HPN in Europe, between 1993 and 1994, and examined organisation, outcomes in terms of survival, and types of catheters used. Since
the experience of the major European centres was pooled, the variation between countries was not reported. The tables are, therefore, a general profile of European HPN activity.

The organisation of HPN programmes is poorly documented in the literature. In Europe, the prescribing hospital and commercial supply companies supply most HPN (see Table 14). Most patients in Europe are trained in hospital (for an average of 14.2 days) and according to protocols in $63 \%$ (see Table 15). Surprisingly, only $43 \%$ were self-caring after training (Van Gossum, 1996).

TABLE 14 Delivery of solutions, disposables and pumps

| Supplier | TPN <br> solutions | Disposables | Pumps |
| :--- | :---: | :---: | :---: |
| Prescribing <br> hospital | $57 \%$ | $58 \%$ | $53 \%$ |
| Local hospital | $1.4 \%$ | $1.6 \%$ | $1.2 \%$ |
| Local pharmacy | $2.4 \%$ | $7 \%$ | $4 \%$ |
| Delivery <br> company | $37 \%$ | $31 \%$ | $18 \%$ |
| Others | $1.2 \%$ | $1.4 \%$ | $10 \%$ |

## Central venous catheters

The commonest central venous access system used in Europe is an external catheter, such as a Broviac or Hickman line, used in $74 \%$ of cases. Implanted reservoir type catheters, which have no external components when not in use, (such as the Portacath) were used less frequently ( $26 \%$ of cases). The external catheters have a section which lies permanently outside the skin to which the infusion is attached.

## Comparative data

Six studies were included in Section 3b on comparative data (see Table 13). One was a randomised, controlled trial (Hyltander, 1991), one was a prospective non-randomised trial (Jarrard, 1980), three were case series (Howard, 1989; Johnston, 1994; Pithie, 1988) and one used historical cohorts (Rannem, 1990). The sample sizes were small (33-69) and were not well matched in some studies. The length of follow-up was often short and may be considered insufficient for technologies

TABLE 15 Training

| Trained in hospital | $83 \%$ |
| :--- | :--- |
| Trained outside hospital | $15 \%$ |
| Training protocol used | $63 \%$ |
| Training manual supplied | $63 \%$ |
| Self-caring after training | $43 \%$ |
| Cared for by relatives | $29 \%$ |
| Community nurse carer | $26 \%$ |
| Average training time | 14.2 days |

such as HPN, which are often used for years. Comparative data on aspects of HPN technology was available in only six $(7.5 \%)$ of the 79 studies which satisfied the inclusion criteria. The randomised, controlled trial of patients with testicular tumours having chemotherapy showed that HPN did not affect the nitrogen balance but did maintain weight owing to fat accumulation. The short duration of the study did not allow important outcomes such as survival to be measured. The other comparative studies suggested that patients would benefit if iodine tincture or chlorhexidine disinfectants and alternate day dressings were used. One study showed that catheter tip position in the right atrium was associated with fewer complications than if the tip was in the superior vena cava; however, this was a small retrospective study and guidelines on ideal catheter tip position cannot be confidently made from such a study. The results of these studies are summarised in Table 16.

TABLE 16 Comparative studies examining aspects of the process of HPN therapy

| Study | Time frame | Study design | Groups compared | Sample size | Outcomes assessed? | Findings |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Howard, 1989 | 1983-88 | Comparison of experience; non-randomised | Reservoir vs external catheter | 58 | Catheter complications | The implanted reservoir was associated with significantly fewer ( $p<0.05$ ) complications than the external catheter. |
| Hyltander, \|99| | 10 weeks | Randomised controlled trial | TPN vs no TPN in chemotherapy | 33 | Nutritional status | Body weight was preserved but this was simply fat accumulation. Nitrogen balance was not maintained. Exercise tolerance was not improved by HPN. |
| Jarrard, 1980 | 1978 | Prospective controlled | Daily vs alternate day dressings | 38 | Catheter colonisation | Daily dressing changes reduced catheter colonisation but this was not significant. Daily dressing was expensive and time-consuming. continued |

TABLE 16 contd Comparative studies examining aspects of the process of HPN therapy

| Study | Time frame | Study design | Groups compared | Sample size | Outcomes assessed? | Findings |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Johnston, 1994 | 1980-93 | Case review plus before and after | Current patients compared with historical controls | 34 | Complication rates | Increasing experience was associated with a reduction in the complication rate ( $p<0.000$ I). Loss of an experienced nutrition nurse did not change the complication rate. |
| Pithie, 1988 | 6 years (1980s) | Case review | Catheter tip position | 69 | Superior vena cava thrombosis | Positioning of the catheter tip in the right atrium reduced the incidence of superior vena cava thrombosis compared with catheters placed in the superior vena cava ( $p<0.01$ ) using glucose as the energy source. |
| Rannem, 1990 | 1976-88 | Case series; non-randomised | Catheter sepsis using various disinfectants | 58 | Catheter sepsis | Incidence of catheter sepsis was significantly higher ( $p<0.05$ ) when povidone iodine was used as a disinfectant rather than iodine tincture or chlorhexidine. |

## Results for Section 4 Economic analysis

There were seven studies that examined economic aspects of HPN therapy, and these are summarised in Table 17. Two studies were from the UK, four from the USA and one from Canada. All examined the costs of HPN from the health service perspective and ignored patient costs. Two studies
examined costs and benefits as part of a formal cost-utility analysis and these studies included marginal quality-of-life and sensitivity analyses. The evidence from these studies for the costeffectiveness of HPN is presented in Table 18.

Two cost-utility analyses were located. The marginal cost per quality-adjusted life year (QALY) varied from Canadian \$14,600 (Detsky,

TABLE 17 Section 4. What evidence exists on the cost-effectiveness of HPN? Summary of studies

| Study: <br> Country <br> (Period) | Case mix | Perspective | Methodology | Costs | Benefits measured | Q-o-L assessment | Marginal analysis | Sensitivity analysis |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Detsky, I986: <br> Canada (1970-82) | Benign | Hospital | Cost-utility analysis | Hospital | Patient | yes | yes | yes |
| Richards, 1996: UK (I995) | Benign | Hospital | Cost-utility analysis | Hospital | Patient | yes | yes | yes |
| Bisset, I992: <br> UK (I992) | Benign | Hospital | Cost analysis | Some | None | no | no | no |
| Wesley, 1983: USA (1983) | Benign | Hospital | Cost analysis | Some | None | no | no | no |
| Wateska, 1980: USA (1980) | Benign | Hospital | Cost analysis | Hospital | None | no | no | no |
| Dzierba, I984: <br> USA (1982-83) | Benign | Hospital | Cost analysis | Hospital | None | no | no | no |
| Baptista, 1984: USA (1984) | Benign | Hospital | Cost analysis | Hospital | None | no | no | no |

TABLE 18 Section 4 Results. Economic evaluations of HPN. Evidence of cost-effectiveness of HPN

| Study | Country (Year) | Perspective | Methodology | Findings (costs are given as reported and are not adjusted to 1995 values) | Sensitivity analysis | \% difference between hospital TPN and HPN |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Baptista, 1984 | USA (1984) | Health service | Cost analysis | Regular assessment of all aspects of patient care can result in significant fiscal savings. | - | - |
| Bisset, 1992 | UK (1992) | Health service | Cost analysis | HPN solutions, pump and consumables cost $£ 23,000-30,000$ per year. | - | - |
| Dzierba, 1984 | USA (1982-83) | Health service | Cost analysis | Hospital TPN more expensive than HPN. $\$ 32,850$ per year for HPN. Approx $\$ 57,000$ for hospital TPN. | - | 72 |
| Wateska, 1980 | USA <br> (1980) | Health service | Cost analysis | First year cost of HPN $\$ 21,465$, thereafter $\$ 19,700$ per year. Hospital TPN costs $\$ 73,720$. | - | 73 |
| Wesley, 1983 | USA <br> (1983) | Health service | Cost analysis | \$33,000-36,000 per year for HPN; \$182,000 for hospital TPN. | - | 81 |
| Richards, 1996 | UK (1995) | Health service | Cost-utility analysis | First year cost of HPN is $£ 44,288$. The marginal cost per QALY was $£ 69,000$. One year of hospital TPN costs $£ 93,000$. | Sensitive to the age of the patient. | 65 |
| Detsky, 1986 | Canada (1970-82) | Health service | Cost-utility analysis | Marginal cost per QALY $\$ 14,600$. Increase of 3.3 years of quality-adjusted survival compared with the alternative of intermittent hospital nutritional support. Cost-utility compares favourably with other health care programmes when used for benign diseases. | Sensitive to the assumptions made regarding the costs of alternative treatments. |  |

1986) to UK $£ 69,000$ (Richards, 1996). Reflation of the 1986 Canadian value equates to approximately $£ 12,000$ per QALY. The most recent estimate of costs to the NHS were $£ 45,000$ for the first year and $£ 36,000$ for subsequent years.

The studies showed that HPN was 65-80\% cheaper than the alternative hospital treatment. The cost-utility analyses suggested that the cost-utility of treating younger patients was more favourable than older patients.

## Chapter 5

## Discussion

## Home parenteral nutrition and intestinal failure

Home parenteral nutrition has been used for the treatment of varying degrees of intestinal failure on both a short and long term basis. Intestinal failure is defined as an inability of the gastrointestinal system to absorb sufficient fluid, electrolyte and/or nutrients for metabolic requirements. Intestinal failure is one aspect of the short bowel syndrome. The clinical features of this syndrome are intractable diarrhoea, weight loss, dehydration, steatorrhoea, malnutrition, and vitamin and mineral deficiency. If the fluids, electrolytes and nutrients are not replaced then there is a progression from dehydration and malnutrition to death. The speed at which this condition progresses is dependent on the degree of intestinal failure. The degree of intestinal failure depends in turn on the length and function of the intestinal remnant. There are other factors which will affect how good the function will be; these include: the pre-existing disease; the amount of residual disease; the adaptive capacity of the bowel; the site of resection, and the presence or abscence of the ileo-caecal valve.

There are many disease processes which can result in intestinal failure. Congenital problems are uncommon and usually result from intestinal atresia or malrotation of the gut around a congenital band. Another common subgroup in paediatric practice are patients with intractable diarrhoea. Conditions which interrupt the vascular supply of the intestine are common and include mesenteric thromboses or emboli, various coagulopathies, intestinal malrotation or volvulus, and intestinal strangulation. Crohn's disease can result in a short bowel syndrome as a result of extensive disease activity or through multiple intestinal resections. Internal or external enteric fistulation can also result in temporary or permanent intestinal failure. Malignant disease can result in intestinal failure as a result of disease extent or as a result of treatment (for example, extensive removal of bowel or irradiation of parts of the abdominal cavity).

The treatment of intestinal failure is complex. The treatment options include parenteral nutrition, enteral nutrition and small bowel transplantation.

Small bowel transplantation is not widely available at the present time in the UK. Most of the world's experience with this procedure is in the USA. Successful transplantation has been hampered by problems with immunosuppression and infection by viral agents. In the UK, patients who have tolerated HPN very well have been advised to continue with HPN treatment until the morbidity and mortality associated with small bowel transplantation improves. While awaiting a suitable donor organ, the patient will require parenteral nutrition.

Enteral nutrition is suitable for patients with lesser degrees of intestinal failure. It has certain advantages over parenteral nutrition in that it is physiologically more acceptable, villus height is maintained, bacterial translocation is reduced, it is simple to administer, and it is much cheaper than parenteral nutrition. However, there are disadvantages in that it can not be used as the sole means of nutritional support in patients with severe short bowel syndrome. This is because there is simply insufficient absorption due to inadequate function or length of bowel. The large volume of enteral fluid required in the severe cases would exacerbate the enteric fluid loss, and lead to dehydration, malnutrition and electrolyte loss. There is scanty evidence describing the use of enteral feeding for short bowel syndrome and, in particular, we are not aware of any studies comparing the use of enteral with parenteral nutrition. It is difficult to predict which patients with intestinal failure will manage with enteral nutrition alone. Absolute bowel length is not a good predictive factor and patients often have to be monitored over a period of weeks and months to ensure that the nutritional support is adequate.

For severe cases of intestinal failure the mainstay of treatment is parenteral nutrition. As the intestine adapts over a period of up to 2 years, it may be possible to wean the patient from parenteral nutrition to enteral nutrition and, possibly, to a normal diet. Until weaning is completed, the patient is dependent on the parenteral administration of fluid and nutrient requirements. A proportion of patients will never have sufficient bowel left to allow adaptation to take place, and will require parenteral nutrition in order to survive and consideration for a small bowel transplant. Any patient
requiring parenteral nutrition for a significant length of time is a candidate for home therapy. When compared to the alternative of in-patient care, HPN is thought to be more acceptable for the patient and cheaper, and it releases beds for the treatment of others.

## The need for the review

Expansion of the technology into areas where its effectiveness has been questioned, the need for data on use of HPN in the UK, the effect on the quality of life of patients, variable referral patterns, and the cost of treatment, led to the prioritisation of HPN as a technology in need of assessment.

The research questions were designed to outline the important aspects of current practice. The review also attempted to identify areas of current practice that were backed by good evidence of effectiveness and those areas that were not.

## Finding the research

The literature search revealed that evidence was restricted to data from case series, there being very few comparative trials. Although the MEDLINE database was the most productive to search, almost $80 \%$ of the HPN research literature would have been missed by a search limited to MEDLINE. Other databases fell a long way behind. Hand searching of the main European journal (Clinical Nutrition) and the main American journal (Journal of Parenteral and Enteral Nutrition) was fruitful but searching the less well-known journals was not. It would appear that most centres publish their HPN work in these two journals.

Locating abstracts from meetings was useful, as it indicated the type and scope of studies which are on-going. The response to the letters we sent to 24 major centres in Europe and America was disappointing (seven replies containing eight new studies). The poor response was possibly due to the timing of the letters, which were sent out in mid-summer when many people may have been on vacation. We did not receive any details of unpublished or on-going research.

Repeated scanning of reference lists was the most successful method of locating additional studies. The Science Citation Index (SCI) was also a useful method of locating studies; however, we had already collected a large number of studies before trying out the SCI, and no additional studies were found.

## The standards of the literature

Of the 256 studies identified, only 65 satisfied the inclusion criteria. The reasons for exclusion were mainly due to the study being hospital based or pre-1980. Only one randomised, controlled trial was identified. Many studies contained data on patients that may have been included more than once. This was due to sampling patients from different sources and including patients who had been entered on to a national database or registry. This is a key issue when pooling data from several sources and may be a source of bias in reporting.

## Trends of use

The quality of the studies satisfying the inclusion criteria for describing use was generally poor. The main problems being non-random samples of the HPN population, retrospective case series and the possibility of patients being included more than once in a study because of data pooling. There was a general increase in the number of patients with malignant disease being entered into HPN programmes. This accounted for $90 \%$ of programme growth in the USA and an increase from $5 \%$ to $8.9 \%$ in the UK recently. In the UK, the trend is opposite to this, with many patients with malignant disease being supported with enteral nutrition. The use of HPN for the treatment of wasting associated with AIDS is also becoming more common, despite the lack of evidence of effectiveness in this disease.

The use of HPN in paediatric practice is uncommon in this country, and this was reflected in the small number of studies located. The larger series originate from the USA and France but the small number of patients in these studies prevent firm conclusions being drawn regarding HPN use in paediatrics.

## Quality of care

Many of the problems with the quality of the studies also applies to those describing patient experience of HPN. Sepsis arising from the central venous catheter was the most common serious complication associated with HPN treatment. The infections were limited to a minority of patients who had recurrent episodes of sepsis. Thus, the incidence of patients with sepsis per catheter year is low. The incidence of infections varied from 0.11 to 0.71 episodes per catheter year (see Figure 1). It can be assumed that centres in different countries will have, at least, some differences in training techniques and catheter care protocols and, given the range in incidence of
sepsis, these differences may be important. It has been suggested that the type of catheter used affects sepsis rates and that there is a lower incidence of infection if reservoir catheters are used. We did not locate any convincing evidence to support this view. There is evidence from the study by Singer (1991) which shows that episodes of catheter sepsis are more common in the patients with AIDS and cancer than in patients on HPN with benign diseases (see Table 8).

Episodes of central line sepsis, occlusion and central vein thrombosis often require catheter removal. This raises problems with venous access for patients on long-term treatment and can be associated with significant morbidity and mortality. The evidence collected does not allow firm guidelines to be developed in order to minimise these complications. There is a need to know which catheters should be placed where and which nutrient solutions are associated with fewest complications. There is anecdotal evidence which suggests that more reservoir catheters are being used for long-term HPN. The evidence to support this change is based on one study (Howard, 1989), which showed that fewer infections occurred when reservoir catheters were used. However, the study design was weak and based on a comparison of non-randomised groups.

The relationship between complication rates and study size is unclear. Experienced centres that have dealt with large cohorts of patients seem to have similar results to smaller units. There does seem to be a learning curve for new centres and early years can be marred by unacceptable complication rates (Johnston, 1994). It is important for new centres to build on the experience of established centres, in order to avoid the problems of the learning curve.

The quality of life experienced by patients with benign disease is reasonable, considering that this treatment is life-saving and the alternative for many patients would be death. There is no evidence describing the quality of life experienced by those patients with malignant disease or AIDS. There is a clear need for this type of information for this patient group, where the emphasis should be on adding quality as well as quantity of life.

The survival figures reflect the underlying disease and it can be seen that the survival figures for patients with benign disease are good. The survival statistics for patients with malignant disease and AIDS are poor, and this is further evidence of the need to assess these subgroups more thoroughly than in the past, with particular emphasis on the best way of providing nutrition (parenteral or enteral). We do not know how survival has been affected by giving HPN to patients with terminal malignancy or AIDS, and this requires further investigation.

## Organisation and evidence of effectiveness

The organisation of HPN patients has not been examined by comparative study. The remarkable similarity in complication rates shown in larger studies suggests that experience is more important than organisation. There is a need to demonstrate more clearly the role of small units and to assess whether larger units are more effective. The question of where patients should be trained is largely an economic one but patients may benefit from the reassurance of hospital surroundings during early training (according to anecdotal evidence) and this should be taken into consideration.

There were few comparative data on aspects of HPN and the studies were of poor design (such as lack of compatability) and small sample size. These factors prevent any firm conclusions being made regarding most of the aspects that were examined (such as which catheter to use, or the ideal site for the catheter tip).

## Economic appraisal

All the evidence found examining the economic aspects of HPN treatment demonstrated that it is cheaper than in-patient treatment. The cost per QALY measured in two studies was reasonable for benign disease, particularly in young patients, and especially considering the life-saving nature of the treatment. There is a complete lack of economic appraisal of HPN for malignant disease and AIDS.

## Chapter 6

## Conclusions

## What gaps in the evidence exist? (Section 5 of the review method)

The quality and range of evidence of effectiveness was disappointing. The technology of HPN has been present for almost 30 years and yet there is still very little good quality evidence to support many aspects of it.

## Section I

The type of patient who has received HPN has been fairly well documented. There is evidence that, in the UK, there is an increase in the number of those with terminal malignant disease and wasting due to AIDS being treated with HPN. It is hoped that accurate data concerning those patients entered into HPN programmes will continue to be collected as part of a national register, administered by the British Association for Parenteral and Enteral Nutrition Council. Trends in the UK could then be monitored more efficiently.

## Section 2

The complications, survival, duration of treatment, and reasons for discontinuing treatment are fairly well documented. The quality of life of patients on HPN has been poorly assessed in the past particularly those with malignant disease and AIDS. A clear survival advantage has been demonstrated for those with a benign underlying disease. However, there is less evidence to indicate whether the complication rates differ for the disease subgroups.

## Section 3

Organisational models for HPN programmes have been poorly assessed and there are no comparative data that we could locate looking at this aspect of the technology; for example, who should deliver the training and where should patients be trained? Comparative data on many aspects of the technology are completely absent, and those which do exist are marred by non-randomised, poorlydesigned, retrospective investigations performed on small samples.

## Section 4

There is some up-to-date evidence looking at the cost of HPN to the health service. Patient and community costs have not been measured. Only two studies have used a formal methodology for
economic appraisal (cost-utility analysis) and these were performed in 1986 (Canada) and 1995 (UK). Comparisons with other technologies have not been made. There are no economic appraisals of HPN used for malignant disease.

## Which questions need to be addressed?

- What is the cost per QALY of HPN for subgroups of patients to determine, for example, if it is costeffective to use HPN in AIDS and cancer patients and other subgroups where the underlying condition is terminal; that is, is HPN of use in palliative care? As part of such a study it is necessary to calculate the typical quality-of-life profile (measured by repeated assessments using a set of validated health status instruments) of patients before, during and after HPN treatments, and to identify moderating factors such as underlying disease. Also, what is the expected survival for patients with terminal malignant disease and AIDS on HPN, and can 'long survivors' be identified?
- What are the most cost-effective organisational models for HPN programmes and does any one model contribute to an improved outcome (for example, small versus large units)?
- What is the best method for training patients for HPN, and should the training be done at home or in hospital?
- Are reservoir catheters associated with less septic episodes than traditional external catheters? Who should then insert central venous catheters, surgeons or interventional radiologists, and what is the ideal position of the catheter tip?
- How cost-effective is HPN compared with other expensive but life-saving technologies?


## What methodological issues need to be addressed in future research?

As mentioned previously, study design has been the downfall of many investigations into the
effectiveness of HPN. Larger, multicentre, studies should be performed. They should be prospective with a clearly defined aim. Comparative studies should have a control group and be randomised. Quality-of-life assessments and economic analyses should follow validated methodologies.

- It is important to have complete up-to-date registries measuring patient characteristics and experience. Collaboration and adequate funding is essential.
- Episodes of catheter sepsis, occlusion, central vein thrombosis and metabolic imbalance
should be documented as part of centre audit. Standards of care should be compared and maintained.
- Patients should be monitored for the development of liver and bone disease, and these should be recorded as part of the 'total patient experience'.
- All changes in the delivery and management of HPN should be properly evaluated.
Comparisons of alternative modes of delivery should preferably be assessed by randomised, controlled trial.


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## Studies not included in the systematic review of HPN and the reasons why

| Study | Reason for exclusion |
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| Buchman AL, et al, 1994. Choline pharmacokinetics during intermittent intra- <br> venous choline infusion in human subjects. Clin Pharmacol Ther;55:277-83. | No relevant outcomes for <br> this review. |
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| Study | Reason for exclusion |
| :---: | :---: |
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| Study | Reason for exclusion |
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| Study | Reason for exclusion |
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| Study | Reason for exclusion |
| :---: | :---: |
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| Passaro M, et al, 1994. Long term silastic catheters and chest pain. J Parent Ent Nutr;18:240-2. | Not an inception cohort. Case report. |
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| Pennington CR, 1992. HPN; an appraisal. Scot M ed J;37:69-70. | No new data. |


| Study | Reason for exclusion |
| :---: | :---: |
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| Perl M, 1987. HPN and the family. Psych Clinics N orth Am;10:121-7. | No empirical data. |
| Perl M, et al, 1980. Psychological aspects of long term home hyperalimentation. J Parent Ent Nutr;4:554-60. | Duplicate publication. |
| Pironi L, et al, 1994. Morphologic and cytoproliferative patterns of duodenal mucosa in two patients after long term TPN. J Parent Ent N utr;18:351-4. | Case reports only. |
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| Pitt HA, et al, 1983. Increased risk of cholelithiasis with prolonged total parenteral nutrition. Am J Surgery;145:106-12. | Duplicate data. |
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| Schropp K, et al, 1988. Catheter related sepsis; a review of experience with Broviac and Hickman catheters. Nutrition;4:195-200. | Data included in other papers. |
| Scott NA, et al, 1991. Spectrum of intestinal failure in a specialised unit. L an cet;337:471-3. | Hospital based. |
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| Study | Reason for exclusion |
| :---: | :---: |
| Scribner BH, Cole JJ, 1979. Evolution of the technique of home parenteral nutrition. J Parent Ent Nutr;3:58-61. | Historical. Pre-1980. |
| Scribner BH, et al, 1970. Long term total parenteral nutrition. The concept of an artificial gut. JAM A;212:457-63. | Historical. Pre-1980. |
| Shanbhogue LKR, Molenaar JC, 1994. Short bowel syndrome; Metabolic and surgical management. Br J Surgery;81:486-99. | Review - no new empirical data. |
| Shapiro RS, 1990. Ethical and legal issues in the use of TPN. Nutrition;6:397-401. | No empirical data. |
| Sharp JW, Roncagli T, 1992. HPN in advanced malignancies. J Parent Ent Nutr;16:190-1. | No empirical data. |
| Sharp JW, Roncagli T, 1993. HPN in advanced cancer. Cancer Pract;1:119-24. | No empirical data. |
| Shenkin A, et al, 1986. Essential trace element provision to patients receiving home intravenous nutrition in the UK. Clin Nutr;5:91-7. | Data not relevant to the review. |
| Shike M, et al, 1981. A possible role of vitamin D in the genesis of parenteral nutrition induced metabolic bone disease. Ann Internal $\mathrm{M} \mathrm{ed} ; 95: 560-8$. | Not an inception cohort. Old data. Bulk of the paper not relevant. |
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| Shils ME, 1984. Historical aspects of minerals and vitamins in parenteral nutrition. Federation Proc;43:1412-16. | Historical. Pre-1980. |
| Silk D, 1995. Malnutrition in hospital. H osp Update; February:55-61. | No new data. |
| Singer P, et al, 1992. Clinical and immunological effects of lipid based parenteral nutrition in AIDS. J Parent Ent Nutr;16:165-7. | ? repetitive data. Also ? relevance. |
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| Stephens L, et al, 1995. Are clinical signs accurate indicators of the cause of central venous catheter occlusion. J Parent Ent Nutr;19:75-9. | Hospital based. |
| Stokes MA, Irving MH, 1988. How do patients with Crohn's disease fare on HPN? Dis Colon Rectum; June:454-8. | Data presented elsewhere. |
| Stokes MA, 1988. HPN; a review of 100 patient years of treatment in 76 consecutive cases. Br J Surgery;75:481-3. | Data included in other papers. |
| Stokes MA, Irving MH, 1989. Mortality in patients on HPN. J Parent Ent Nutr;13:172-5. | Data included elsewhere. |
| Storch K, 1992. Home parenteral nutrition. N Jersey M ed;89:36-40. | No emprirical data. |
| Stuart R, et al, 1990. Perioperative nutrition in cancer patients. Nutrition;6:4S-7S. | Hospital based. |
| Treasadern JC, et al, 1984. Maintainance of pregnancy in a HPN patient. J Parent Ent Nutr;8:199-202. | Not relevant. |
| Twomey PL, Patching SC, 1985. Cost effectiveness of nutritional support. J Parent Ent Nutr;9:3-10. | Perioperative TPN. |
| Vars HM, 1980. Early research in parenteral nutrition. J Parent Ent N utr;4:467-8. | Historical interest only. |
| Veleisis RA, et al, 1980. Prospective controlled trial of parenteral nutrition associated cholestatic jaundice: effect of protein intake. J Paed;96:893-7. | Hospital based. |
| Watters DAK, et al, 1984. Changes in liver function tests associated with parenteral nutrition. J R oy Coll Surgeons Edin;29:339-44. | Hospital based. |
| Wilcock H, et al, 1991. Artificial nutrition support for patients in the Cambridge health district. H ellth Trends;23:93-100. | No outcomes for the 3 HPN patients included in this study. |


| Study | Reason for exclusion |
| :--- | :--- |
| Wilkinson AW, 1963. Historical background of intravenous feeding. Nutr Dieta;5:295-7. | Historical. Pre-1980. |
| Williams N, et al, 1994. Incidence and management of catheter related sepsis in <br> patients on HPN. Br J Surgery;81:392-4. | Not an inception cohort. |
| Williams N, et al, 1993. The incidence and management of catheter occlusion in <br> patients on HPN. Clin Nutr;12:344-9. | Not an inception cohort. |
| Wilmore D, Dudrick SJ, 1968. Growth and development of an infant receiving <br> all nutrients exclusively by vein. JAM A;203:140-4. | Historical interest only. |
| Winters RW, et al, 1984. History of parenteral nutrition in paediatrics with <br> emphasis on amino acids. Federation Proc;43:1407-11. | Historical. Pre-1980. |
| Wood RJ, 1985. Calciuretic effect of cyclic versus continuous TPN. <br> Am J Clin Nutr;41:614-9. | Small number of <br> patients. No controls, <br> no randomisation. <br> Short period of TPN. |
| Woolman SL, et al, 1979. Zinc in TPN; requirements and metabolic effects. <br> Gastroenterology;76:458-67. | Not relevant to the review. |

## Appendix 1

## The history of HPN

Home parenteral nutrition is a life-saving technology which was developed in the USA during the late 1960s (Scribner,1970; Dudrick, 1968; Wilmore, 1968). It became possible due to the production of safe, stable solutions of protein, fat and glucose. Dudrick and colleagues (1967) showed that puppies fed by intravenous nutrition, developed and grew normally. Wilmore and Dudrick (1968) described their initial efforts to provide intraven-ous nutrition for a neonate with intestinal atresia. A solution containing the required nitrogen, calories and trace elements, was infused into the superior vena cava for a period of 44 days. Normal growth and development was seen to occur. This would appear to be the first well-documented case of a human patient not only gaining weight, but passing into a significant and continuing anabolic state on the basis of parenteral nutrition alone. Using this technique, Dudrick and colleagues (1969) reported the progress of 30 adult patients; they noted good wound healing, weight gain, increased strength and fistula closure in some patients.

Scribner (1970) described the concept of an artificial gut which could provide prolonged nutritional support for a patient incapable of enteric feeding. In Scribner's system, an arterio-venous shunt was used for venous access and nutrients were delivered by gravity or pump. The external arterio-venous shunt was created using a silicone rubber tube with a side arm for the infusion of nutrients. Scribner postulated that once the system was up and running it should be possible to discharge the patient home and with appropriate training he or she should be self caring. This is the first mention of HPN in the literature. The likely costs involved in maintaining a patient on prolonged nutrition at that time were $\$ 5$ per day or $\$ 1800$ per year, with an initial outlay of approximately $\$ 1000$ for pumps, etc. Scribner and Cole (1979) subsequently criticised their study (1970) as being rather premature. The system which they described worked well in the uraemic patients whom they used as controls; however, when it was tried in malnourished patients the standard arterio-venous shunts clotted in almost $100 \%$ of cases. The adverse effects of poor quality veins and normal clotting parameters on graft function had not been anticipated. When they
realised that the shunts were not going to be suitable they were forced into trying a new technique which involved inserting a Tenckhoff catheter via the subclavian route into the right atrium. This catheter initially worked very well and adequate nutrition was restored. Unfortunately, the mechanical trauma suffered by the superior vena cava resulted in thrombosis, obstruction and failure of the catheter. The stiff Tenckhoff catheters were replaced by a newly-developed flexible, soft, silicone rubber tube. This basic change in the design meant that vascular trauma was minimised, resulting in successful long-term venous access.

Broviac (1973) reported experience with a silicone rubber right atrial catheter. The thin intravascular portion was positioned in the right atrium for maximal dilutional effect and the thicker extravascular portion is brought out via a long tunnel on the anterior chest wall. A Dacron ${ }^{\circledR}$ cuff was positioned beneath the skin and, after about 3 weeks, the ingrowth of collagen fibres led to a firm anchor being created. The catheter was flexible, inert and anti-thrombogenic, allowing it to move with each heartbeat, ensuring that the tip did not irritate one particular portion of endocardium. Broviac reported local and generalised infection as the main complication and a mean catheter life of 144 days per patient. This compared favourably with the previously reported average catheter lifespan of 24 days (Wilmore, 1969).

The gravity system of infusion was found to be unreliable when used overnight and required constant vigilance. A powered portable device, contained in a specially desiged vest, was developed which eliminated this problem. Some of the patients criticised the 'wearable' infusion device which delivered the fluids during the day. It was seen as cumbersome and unnecessary and was soon abandoned, being replaced by portable stands and, later, by cyclical night-time infusion.

Jeejeebhoy (1973) reported the experience of a patient who had received HPN for 23 months without complications and with good rehabilitation. The lipid infusions were given separately, as they could not pass through the filters. The lipid infusion provided half of the required calories and was regarded as an absolute requirement if
essential fatty acid deficiency was to be avoided. Jeejeebhoy also suggested that the use of fat prevented the development of a fatty liver.

As experience grew with the technique of home parenteral nutrition further favourable reports appeared (Ivey, 1975; Bordos, 1975; Shils, 1975; Heizer, 1977). An estimate of costs in 1975 revealed that the initial basic costs were about $\$ 700$ but the costs of the infusions had increased to $\$ 7200-\$ 12,000$ per year depending on requirements and on the type of amino acid infusion used. Scribner treated 40 patients in this manner; five patients died but four deaths were as a result of the underlying disease. The average length of treatment was 11 months. The commonest complications were sepsis, thrombo-embolism, metabolic imbalance and fat infiltration of the liver. The choice of patients in Scribner's series was of interest as the diagnostic subgroups were very similar to current practice in the UK. The main indication for HPN being benign disease (Crohn's and MVD). Current practice in the USA has changed significantly and now includes large numbers of patients with malignant disease.

Cases of intestinal failure which were previously beyond the help of medical technology became 'treatable' (Bordos, 1975). Shils (1975) reported his experience with 11 patients maintained at home on intravenous nutrition. He used a standard portable pump system which was equipped with infusion rate monitors. This allowed safe administration overnight, thereby allowing patient freedom during the day and improving the patient's quality of life. Shils noted that most patients were able to learn the necessary techniques within a few weeks if they received daily training sessions. The more controversial aspect of this paper deals with the selection of patients suffering from terminal malignancy; even though this paper was published 20 years ago, controversy still exists. The author justifies his decision to use HPN for these patients with the statement that they are often able to spend a rewarding last few months (or even years, in some cases) at home.

HPN technology diffused to Europe from the USA in the late 1970s and this systematic review examines the world experience from 1980 onwards.

## Appendix 2

## Review protocol

## Introduction

The aim of this review is to locate, acquire and synthesise studies concerning the use, effectiveness and costeffectiveness of home parenteral nutrition. The review will fall into five main sections:

Section 1. What patients have received HPN?
Section 2. What has been the experience of patients on HPN programmes?
Section 3. (a) How have HPN programmes been organised, and what techniques and equipment have been used, and to what effect?
(b) What comparative information is available on effectiveness?

Section 4. What evidence exists on the cost-effectiveness of HPN?
Section 5. What questions about the provision of HPN could be answered with additional research, and what design of study would be most suitable?

For each of the above sections we will consider both adults and children.
Searching the HPN literature suggests that there are very few comparative studies available. Most of the literature consists of case series. The questions above have been formulated bearing this is mind, so as to make best use of what information is available.

## Section I

## Research question

What patients have been entered into HPN programmes with respect to numbers, age, sex, diagnoses, setting (country) and what trends exist?

## Studies to be included

1. Surveys of HPN use.
2. Information from databases of HPN users.
3. Data from cohorts of HPN patients.

## Points for assessing validity

1. Are the patient numbers based on new patients in a given period of time, or from a crosssectional sample?
2. Was the data collected prospectively or retrospectively?
[A longitudinal inception study will provide actual patient numbers being treated on HPN by subgroup, whilst a cross-sectional sample will only provide a snapshot of the sort of patients being treated with HPN. The results will differ according to the length of time spent on HPN. The results of the two types of studies will be interpreted separately. Retrospective data collection is prone to be less complete than prospective data collection.]
3. What proportion of people were sampled and how was the sample chosen?
4. What group was the sample selected from and how representative is the sample?
5. Was the sample coverage validated, and if so was it found to be acceptable?
[Complete assessment or random sampling are the best ways of maintaining representativeness. Both rely on the correct identification of a sampling frame. When these are not available it may be possible to validate sample coverage by taking a sample using a second source and noting the degree of similarity in sample members.]
6. For what proportion of selected people was no information available?
7. How complete was the data that was acquired?
[Non-response and missing data in a survey reduces validity.]

## Data extraction (Section 1)

Study (Code Number)
Title
Journal
Author
Country

Setting

Unit size
(new cases per year)

## Eligibility

Does it include HPN patient numbers?
Is it a survey, database or cohort of HPN-treated patients?

## Validity

Is it a cross-sectional or longitudinal study?
Is it retrospective or prospective?
What were the dates covered?
What was the source of data?
Who was eligible to be in the sample? (What was the sampling frame?)
What sampling fraction was used and was it random?
How was the sampling frame validated?

Patient numbers

| Year | Crohn's | MVD | Pseud | Cancer | AIDS | Other | Mean <br> age | Age <br> range | M:F |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
|  |  |  |  |  |  |  |  |  |  |
|  |  |  |  |  |  |  |  |  |  |
|  |  |  |  |  |  |  |  |  |  |

## Section 2

## Research question

What has been the experience of patients on HPN programmes by diagnosis?
(i) What is the duration of HPN use?
(ii) What is the expected survival of HPN users?
(iii) What are the HPN discontinuation rates, and for what reasons?
(iv) What complications occur, and how often?
(v) What quality of life is experienced by patients receiving HPN?

Studies which have been used in Section 1 may also be of use in this section.

## Inclusion criteria

1. Studies reporting the experience of inception cohorts of HPN users.
2. Studies giving information on length of treatment, mortality, complications, or quality of life.

## Validity

1. Is data collected prospectively or retrospectively?
2. What cohort was recruited?
3. How much of the cohort was successfully recruited?
4. How complete was the follow-up?
5. What procedure was used to detect complications?
6. What quality-of-life instrument was used and how was it validated?

Ascertainment of outcomes should be free from bias. Where excessive patient investigations are carried out this may detect a higher number of events than in case series with less active investigation. The instruments used to measure subjective issues like quality of life need

A potential problem in this area is the choice of denominators for the calculation of rates, whether they are patient numbers, patient years, and the way in which data has been analysed. When reporting rates, it is important to make the distinction between per 1000 per year, which implies a group of patients are all followed-up for the same length of time, and per 1000 patient years.

## Data Extraction (Section 2)

Study (Code Number)
Title
Journal
Author
Country
Setting
Unit size (new cases per year)

## Eligibility

Is a cohort of new HPN patients identified, recruited and followed?
What outcome information is contained?

## Validity

Is data collected prospectively or retrospectively?
What proportion of the cohort was recruited?
How complete was the follow-up?
What procedure was used to detect complications?
What quality-of-life instrument was used and how was it validated?
What denominator is used to calculate the rates?

## Type of study

Results

## Period

Sample size
Age description at recruitment.
Male:Female

Complications

|  | Crohn's | MVD | Pseud | Cancer | AIDS | Other |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
|  |  |  |  |  |  |  |  |
|  |  |  |  |  |  |  |  |
|  |  |  |  |  |  |  |  |

Quality of life

|  | Crohn's | MVD | Pseud | Cancer | AIDS | Other |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
|  |  |  |  |  |  |  |  |
|  |  |  |  |  |  |  |  |
|  |  |  |  |  |  |  |  |

Survival

|  | Crohn's | MVD | Pseud | Cancer | AIDS | Other |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| Year |  |  |  |  |  |  |  |
| Year |  |  |  |  |  |  |  |
| Year |  |  |  |  |  |  |  |

Duration of HPN use

|  | Crohn's | MVD | Pseud | Cancer | AIDS | Other |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
|  |  |  |  |  |  |  |  |
|  |  |  |  |  |  |  |  |
|  |  |  |  |  |  |  |  |

Reasons for stopping HPN

|  | Crohn's | MVD | Pseud | Cancer | AIDS | Other |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| Sample |  |  |  |  |  |  |  |
| Still on |  |  |  |  |  |  |  |
| Recover |  |  |  |  |  |  |  |
| Dead |  |  |  |  |  |  |  |
| Lost |  |  |  |  |  |  |  |

## Section 3a

## Research question

How have HPN programmes been organised, and what techniques and equipment have been used, and to what effect?

## Studies to be included.

Surveys assessing issues in the delivery of HPN since 1980.

## Issues

Only one study is known of which has assessed how HPN programmes have been organised.

## Section 3b

## Research question

Has HPN been compared with any alternative therapies? Have larger centres been compared with smaller centres? Have different techniques used in HPN been compared? What comparative information is available?

## Studies to be included

1. Does the study look at: Alternatives?

Techniques or equipment?
The size of centres?
2. Does the study report the experience of patients on HPN?

## Validity

1. What was the study design? Descriptive/Comparative?
2. Is data collection prospective or retrospective?
3. Was a comparison made?
4. How were allocations to treatment made?
5. Was follow-up complete?
6. Were the groups comparable with respect to age and diagnosis?
7. What outcomes were measured?
8. Was the length of follow-up more than 3 months in all cases?
9. How were outcomes assessed?

The validity of comparisons between different procedures, etc., will greatly depend on the study design.
according to design in the analysis.

## Data Extraction (Section 3)

## Study (Code Number)

Title
Journal
Author
Country
Setting
Se
Unit size
(new cases per year)

## Eligibility

[This has been kept vague because of the lack of relevant studies.]
Does the study evaluate patient experience on HPN?
What outcomes does it measure?
What technology does the paper assess?
Organisation, management or delivery?
Techniques or equipment?
Patient education or nutrition?

## Validity

What was the study design?
Is a comparison made, if so between what?
How were allocations to the different groups made?
What factors of comparability were checked at inception?
How complete was follow-up?
Was follow-up long enough for morbidity and mortality to occur?
What outcomes were assessed?
Was there potential for bias in outcome assessment (masked assessment, etc.)?

## Patients included

Sample size
Description of age
Male:Female
Diagnoses

## Interventions

How was the programme organised?
Nutrition team involvement (advisory, direct care, monitoring)
Home care company
Local pharmacy
Trained at home or in hospital

## 0 utcomes

Outcomes measured and results
Catheters

| Type | Sample | Sepsis | Thrombosis | Dislodge | Other | Lost |
| :--- | :--- | :--- | :--- | :--- | :--- | :---: |
| Reservoir |  |  |  |  |  |  |
| Brov/Hick |  |  |  |  |  |  |

## Results for comparisons

Observed difference and confidence interval
Statistical significance
Authors' conclusion

## Section 4

## Research question

What evidence exists on the cost-effectiveness of HPN? What alternatives to HPN have been considered?

## Inclusion criteria

Any economic evaluation of an HPN programme
[Very few economic analyses are known.]

## Validity

1. What methodological technique has been applied?
2. What was the comparison made with?
3. What perspective was adopted?
4. Were all costs considered?
5. Were costs measured appropriately?
6. Were all outcomes considered?
7. Were they measured appropriately?
8. How was quality of life assessed?
9. Was a marginal analysis performed?
10. Was the robustness of the result tested in a sensitivity analysis?

## Data extraction (Section 4)

## Study (Code Number)

Title
Journal
Author

Country $\quad$ Setting | Unit size |
| :---: |
| (new cases per year) |

## Eligibility

Does the study evaluate both costs and benefits of HPN programmes?

## Validity

What methodological technique has been applied?
What was the comparison made with?
What perspective was adopted?
What costs were considered?
Were all important costs considered?
Were costs measured appropriately?
Were all outcomes considered?
Were they measured appropriately?
How was quality of life assessed?
Was a marginal analysis performed?
Were assumptions tested in a sensitivity analysis?

## Results

What did the study find?

## Section 5

Which questions remain unanswered?
What gaps in knowledge exist?
What clinical issues need to be addressed?
What methodological issues need to be addressed?

## Literature search strategy and study retrieval

The aim of this search is to provide a comprehensive list of primary studies. The field of home parenteral nutrition is fairly well-defined and indexed. This means that it is relatively easy to search electronic
some aspects of HPN technology. HPN is often life-saving which makes comparative investigation very difficult, especially given that there are few alternative treatment strategies. For this reason it was decided that we should attempt to collect all literature concerning the technology of HPN. The search would therefore include all types of study design and include all possible aspects of HPN technology.

The search will not be confined to studies published in English. There are only a small number of centres in Europe who regularly publish HPN data. These centres will be asked to supply any data published in a foreign language and these studies will be translated. It is unlikely that any significant foreign language study will be missed using this policy.

Before starting the search, advice will be sought from an information scientist based at the NHS Centre for Reviews and Dissemination, York, and from a senior medical librarian based at Hope Hospital, Salford.

The following possible sources of data were identified.

- Electronic databases
- Hand searching of relevant journals
- Personal literature collections
- Conference proceedings
- Writing to all major centres in Europe and the USA
- Science citation database
- Scanning reference lists of studies located


## Electronic databases

The following key words will be used to search the databases.
Home care services/Economics, Hospital-based economics, Organisation, Statistical.
Home infusion therapy/Economics, Methods, Nursing.
Home parenteral nutrition, Home total parenteral nutrition, Home ambulatory nutrition/Therapy,
Organisation, Economics, Complications.
Total parenteral nutrition/Home.
Total Parenteral/Nutrition, Home.
Nutritional support.
Nutrition disorders/Therapy.
Short bowel syndrome.
Intestinal failure.
Intestinal fistulas/Therapy.
Crohn's disease/Therapy.
Inflammatory bowel disease/Therapy.
Malignant bowel obstruction.
Intestinal obstruction/Therapy.
Mesenteric vessel occlusion/Thrombosis/Embolisation.
Mesenteric vascular disease/Thrombosis/Embolisation.
Mesenteric artery occlusion/Thrombosis/Embolisation.
Pseudo-obstruction.
Radiation enteritis.
Intestinal radiation damage.
Catheters indwelling.
Central venous access/Devices.
Catheters implantable.
Subcutaneous reservoirs.
Vascular access/Devices.
Quality of life/Home parenteral nutrition.
Economics/Home parenteral nutrition.
It is well-documented in the literature that many studies can be missed if searches are limited to only one database. We will search a number of databases as outlined below. The databases will be searched from 1968 onwards, i.e. from the origins of HPN.

1. MEDLINE (Index M edicus on-line)
2. EMBASE (Excerpta M edica on-line)
3. Science Citation Index
4. UNCOVER
5. CINAHL (US database, mainly nursing based)
6. CAREDATA
7. Food Science and Technology Abstracts
8. NTIS (US Research Reports)
9. PASCAL (French, scientific database which covers medicine)
10. PSYCHLIT
11. Economic Literature Index

## Hand searching

The following journals will be hand searched for the following years; January 1980-July 1995 (if available during these years)

- Journal of Parenteral and Enteral Nutrition
- Clinical Nutrition
- American Journal of Clinical Nutrition
- Nutrition
- Clinical Gastroenterology
- Nutrition Reviews
- Annals of Nutrition and M etabolism
- Nutrition and Cancer
- Nutrition and Health
- Journal of Paediatric Gastroenterology and Nutrition


## Conference proceedings

Proceedings will be obtained for the annual conferences of the following bodies.

- ASPEN (American Society for Parenteral and Enteral Nutrition) 1993, 1994, 1995
- ESPEN (European Society for Parenteral and Enteral Nutrition) 1993, 1994, 1995
- BAPEN (British Association for Parenteral and Enteral Nutrition) 1994

The published abstracts for these meetings will be examined and, if the abstracts are of specific relevance to the review, attempts will be made to obtain papers from the authors. The notoriously unreliable nature of data contained in abstracts means that we do not plan to use abstracts for data extraction.

## Personal literature collections

We will examine the files of Professor Sir Miles Irving who was one of the founders of HPN in the UK. Over the past 20 years, he and his research staff have published widely on the technology of HPN, and this has led to the accumulation of many relevant papers.

## Letters to experts

We will contact major European and American centres and explain the basis of the systematic review. The centres and experts were chosen because they publish regularly on aspects of home parenteral nutrition or they were members of the ESPEN Home Artificial Nutrition Study Group. The systematic review will be explained and the research questions we hope to answer will be included. We will request any relevant literature, published, unpublished and in progress. Permission will be sought if we need to include any unpublished work.

## Visits to major meetings

Two members of the project team (DMR and JLS) will attend the ESPEN meeting in Rome (September 1995) and the BAPEN meeting (December 1995). Attempts will be made to meet with experts from Europe and the USA. The basis of the systematic review will be explained and comments invited. Those experts that we meet will be asked to supply any literature relating to HPN if they have not already done so.

## Selection of eligible studies, checking validity and data extraction

One researcher (DR) will initially check articles for eligibility for each of the research questions, grade their validity, and extract the necessary data. A second researcher (JJD) will validate these decisions by processing a random sample of studies that are suitable and studies that the first researcher deemed to be unsuitable.

Where any researcher finds ambiguity or is unsure in any aspect of these selection procedures he will obtain independent advice from a third researcher.

## Study synthesis

It is unlikely that it will be possible to combine the results of the research in a quantitative manner, it is anticipated that there are very few randomised, controlled trials and a lack of comparative studies, or any studies of similar design. Therefore, the results will be discussed in a qualitative manner. This will take into account the magnitude of the results, the size and validity of the studies together with any moderating factors.

## Protocol modifications

Any further questions that arise from the review will be addressed. This protocol will be adjusted accordingly and the additional questions will be highlighted as post-hoc hypotheses generated by the review.

## Acute Sector Panel

Professor Senga Bond, University of Newcastle-upon-Tyne ${ }^{\dagger}$
Professor Ian Cameron, SE Thames RHA
Ms Lynne Clemence, MidKent Health Care Trust ${ }^{\dagger}$
Professor Cam Donaldson, University of Aberdeen ${ }^{\dagger}$

| Chair: Professor John Farndon, Un |  |
| :--- | :--- |
| Professor Richard Ellis, St | Dr Chris McCall, |
| James's University Hospital, | General Practitioner, <br> Dorset |
| Leeds $\dagger$ |  |
| Dr David Field, Leicester | Professor Alan McGregor, |
| Royal Infirmary NHS Trust $\dagger$ | St Thomas's Hospital, |
| Mr Ian Hammond, | London |
| Hillingdon HA $\dagger$ | Mrs Wilma MacPherson, |
| Professor Adrian Harris, | St Thomas's \& Guy's |
| Churchill Hospital, Oxford | Hospitals, London |

Dr Chris McCall, Practitioner,

Professor Alan McGregor, thomas's Hospital Mrs Wilma MacPherson, Hospitals, London

Professor Jon Nicoll, University of Sheffield $\dagger$ Professor John Norman, Southampton University Professor Gordon Stirrat, Professor Gordon Stirrat,
St Michael's Hospital, Bristol Professor Michael Sheppard, Queen Elizabeth Hospital, Birmingham ${ }^{\dagger}$

Dr William Tarnow-Mordi,
University of Dundee
Professor Kenneth Taylor, Hammersmith Hospital, London ${ }^{\dagger}$

Diagnostics and Imaging Panel
Chair: Professor Mike Smith, University of Leeds ${ }^{\dagger}$

Professor Michael Maisey, Guy's \& St Thomas's Hospitals, London* Professor Andrew Adam, UMDS, London $\dagger$ Dr Pat Cooke, RDRD, Trent RHA Ms Julia Davison, St Bartholomew's Hospital, London ${ }^{\dagger}$

Professor Donald Jeffries, St Bartholomew's Hospital, London ${ }^{\dagger}$
Dr Andrew Moore, Editor, Bandolier ${ }^{\dagger}$
Professor Chris Price, London Hospital Medical School ${ }^{\dagger}$

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Professor Colin Roberts, University of Wales College of Medicine ${ }^{\dagger}$
Miss Annette Sergeant, Chase Farm Hospital, Enfield
Professor John Stuart, University of Birmingham
Dr Ala Szczepura, University of Warwick $\dagger$

Mr Stephen Thornton, Cambridge \& Huntingdon Health Commission Dr Gillian Vivian, Royal Cornwall Hospitals Trust $\dagger$ Dr Jo Walsworth-Bell, South Staffordshire Health Authority ${ }^{\dagger}$ Dr Greg Warner, General Practitioner, Hampshire ${ }^{\dagger}$

## Methodology Panel

Chair: Professor Anthony Culyer, University of York ${ }^{\dagger}$

| Mr Doug Altman, Institute of Health Sciences, Oxford ${ }^{\dagger}$ | University of Oxford | University of Leeds | Professor Ian Russell, University of York ${ }^{\dagger}$ |
| :---: | :---: | :---: | :---: |
| Professor Michael Baum, Royal Marsden Hospital | ofessor George Daveynith, University of Bristol | $\begin{aligned} & \mathrm{Mr} \\ & \text { Lee } \end{aligned}$ |  |
| Professor Nick Black, London School of Hygiene \& Tropical Medicine ${ }^{\dagger}$ | University of Oxford ${ }^{\dagger}$ | Professor Richard Lilford, Regional Director, R\&D, West Midlands ${ }^{\dagger}$ | Mes |
| Professor Martin Buxton Brunel University $\dagger$ | Professor Stephen Frankel, University of Bristol | Mr Nick Mays, Kings Fund Institute, London $\dagger$ | St Bartholomew's London |

Dr David Spiegelhalter, Institute of Public Health, Cambridge ${ }^{\dagger}$
Professor Charles Warlow, Western General Hospital, Edinburgh ${ }^{\dagger}$

## Pharmaceutical Panel

## Chair: Professor Tom Walley, University of Liverpool ${ }^{\dagger}$

Professor Michael Rawlins, University of Newcastle-upon-Tyne*

Dr Colin Bradley, University of Birmingham

Professor Alasdair Breckenridge, RDRD, Northwest RHA

Ms Christine Clarke, Hope Hospital, Salford ${ }^{\dagger}$
Mrs Julie Dent, Ealing, Hammersmith and Hounslow HA, London ${ }^{\dagger}$

Mr Barrie Dowdeswell, Royal Victoria Infirmary, Newcastle-upon-Tyne

Dr Desmond Fitzgerald, Mere, Bucklow Hill, Cheshire ${ }^{\dagger}$
Dr Alistair Gray, Wolfson College, Oxford ${ }^{\dagger}$
Professor Keith Gull, University of Manchester Dr Keith Jones, Medicines Control Agency

Professor Trevor Jones, ABPI, London ${ }^{\dagger}$ Dr Andrew Mortimore, Southampton \& SW Hants Health Authority ${ }^{\dagger}$ Dr John Posnett, University of York
Dr Frances Rotblat, Medicines Control Agency ${ }^{\dagger}$

Dr Ross Taylor,
University of Aberdeen $\dagger$
Dr Tim van Zwanenberg, Northern RHA
Dr Kent Woods, RDRD,
Trent RO, Sheffield ${ }^{\dagger}$

## Population Screening Panel

Chair: Professor Sir John Grimley Evans, Radcliffe Infirmary, Oxford ${ }^{\dagger}$

Dr Sheila Adam, Department of Health* Dr Anne Dixon Brown, NHS Executive, Anglia \& Oxford ${ }^{\dagger}$ Professor Dian Donnai, St Mary's Hospital, Manchester ${ }^{\dagger}$

Professor George Freeman, Charing Cross \& Westminster Medical School, London Dr Mike Gill, Brent \& Harrow Health Authority ${ }^{\dagger}$ Dr JA Muir Gray, RDRD, Anglia \& Oxford RO ${ }^{\dagger}$

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Professor Catherine Professor Nick Wald, Peckham, Institute of Child University of London $\dagger$ Health, London ${ }^{\dagger}$ Dr Connie Smith Parkside NHS Trust, London ${ }^{\dagger}$
Dr Sarah Stewart-Brown,
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Professor Ciaran Woodman, Centre for Cancer Epidemiology, Manchester ${ }^{\dagger}$

## Primary and Community Care Panel

Chair: Professor Angela Coulter, Kings Fund Centre for Health Services Development, London ${ }^{\dagger}$

Professor Martin Roland, University of Manchester* Dr Simon Allison, University of Nottingham Mr Kevin Barton, Bromley Health Authority ${ }^{\dagger}$ Professor John Bond, University of Newcastle-upon-Tyne ${ }^{\dagger}$
Professor Shah Ebrahim, Royal Free Hospital, London

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Mr Lionel Joyce, Chief Executive, Newcastle City Health NHS Trust ${ }^{\dagger}$
Professor Martin Knapp,
London School of Economics \& Political Science ${ }^{\dagger}$
Professor Karen Luker
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Dr Fiona Moss, North Thames British Postgraduate Medical Federation ${ }^{\dagger}$

Professor Dianne Newham, Kings College, London

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