# Prospective pilot study: evaluation of a new dressing for patients treated in peritoneal dialysis

| Submission date   | Recruitment status              | Prospectively registered    |
|-------------------|---------------------------------|-----------------------------|
| 23/08/2006        | No longer recruiting            | ☐ Protocol                  |
| Registration date | Overall study status            | Statistical analysis plan   |
| 30/10/2006        | Completed                       | Results                     |
| Last Edited       | Condition category              | Individual participant data |
| 19/09/2007        | Urological and Genital Diseases | Record updated in last year |

# Plain English summary of protocol

Not provided at time of registration

# Contact information

## Type(s)

Scientific

#### Contact name

Dr Robert Milongo

#### Contact details

AGDUC Hôpital La Tronche Grenoble France 38 700

# Additional identifiers

Protocol serial number 1445

# Study information

Scientific Title

**Study objectives** 

Evaluation of a new Peritoneal Dialysis (PD) dressing.

#### Ethics approval required

Old ethics approval format

#### Ethics approval(s)

Submitted to CPP Grenoble (reference number: 06-GAMB-1), will receive verdict on 20th September 2006.

#### Study design

Monocentric, open, comparative study

#### Primary study design

Observational

## Study type(s)

Treatment

#### Health condition(s) or problem(s) studied

Chronic renal failure

#### **Interventions**

Evaluation of dressing advantages via questionnaires

#### Intervention Type

Other

#### Phase

**Not Specified** 

#### Primary outcome(s)

Dressing advantages are evaluated via questionnaires, which have to be filled in at specific time schedule. Comfort and safety evaluation will be performed by the patient at the end of each period of product use (end of PD Immo use and end of usual dressing use).

## Key secondary outcome(s))

Handling is also evaluated via the patient questionnaire.

The incidence of Adverse Events (AE) will be followed during the study period.

## Completion date

25/11/2006

# **Eligibility**

#### Key inclusion criteria

- 1. Patients with well healed exit site
- 2. Patients treated for at least three months in peritoneal dialysis irrespective of the treatment mode

- 3. Patients with healthy skin judged by investigator
- 4. Patients aged 18 years and older
- 5. Patients having signed a written consent (informed consent) to participate in the study

#### Participant type(s)

Patient

#### Healthy volunteers allowed

No

#### Age group

Adult

## Lower age limit

18 years

#### Sex

All

#### Key exclusion criteria

- 1. Patients treated in hemodialysis
- 2. Patients with diagnosed abdominal pain not related with PD-solution
- 3. Peritonitis within one month prior to the study
- 4. Presence of exit-site, tunnel infection
- 5. Patients with ongoing peritonitis
- 6. Patients participating in other studies during the period of this study
- 7. Patients practising swimming
- 8. Patients under quardianship
- 9. Pregnancy, lactation

#### Date of first enrolment

25/09/2006

#### Date of final enrolment

25/11/2006

# Locations

#### Countries of recruitment

France

# Study participating centre

**AGDUC** 

Grenoble

France

38 700

# Sponsor information

## Organisation

Gambro Industries (France)

#### **ROR**

https://ror.org/01mgtdr23

# Funder(s)

## Funder type

Industry

#### **Funder Name**

Gambro Industries (France)

# **Results and Publications**

Individual participant data (IPD) sharing plan

# IPD sharing plan summary

Not provided at time of registration