Dietary oligofructose in the prevention of antibiotic associated diarrhoea in elderly patients

Submission date	Recruitment status No longer recruiting	Prospectively registered		
12/09/2003		☐ Protocol		
Registration date	Overall study status	Statistical analysis plan		
12/09/2003	Completed	[X] Results		
Last Edited	Condition category	[] Individual participant data		
06/01/2009	Digestive System			

Plain English summary of protocol

Not provided at time of registration

Contact information

Type(s)

Scientific

Contact name

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Contact details

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Additional identifiers

Protocol serial number N0544103828

Study information

Scientific Title

Study objectives

The purpose of this study is to see if the incidence of antibiotic related diarrhoea in elderly patients can be reduced by increasing colonic bifidobacteria concentrations with the concomitant prescription of oligofructose. Patients for the study will be identified by asking the Medical and Department for the Elderly teams if they had admitted any suitable patients. A brief clinical and drug history would be taken.

Ethics approval required

Old ethics approval format

Ethics approval(s)

Not provided at time of registration

Study design

Randomised controlled trial

Primary study design

Interventional

Study type(s)

Not Specified

Health condition(s) or problem(s) studied

Digestive System: Diarrhoea

Interventions

The volunteers would then be randomly allocated to one of two groups.

Group A will take a placebo (10 g/day of sucrose powder) or group B 10 g/day of oligofructose. The trial powders will be administered by the ward nursing staff along with the volunteers' normal medication. Volunteers will take the trial powders for the length of time they are taking antibiotics and for a further 7 days after stopping antibiotics. Volunteers will be assessed continuously by recording on a stool form using a four point scale (1 hard, 2 lumpy, 3 mushy and 4 loose) and interdefecatory intervals. Both give an indication of intestinal transit rate. Recordings will be done by the patient or nursing staff as appropriate. The volunteers will be followed up for 7 days after stopping the oligofructose or placebo. The volunteer will have further stool samples sent to be analysed for the presence of Clostridium difficile toxin if they develop diarrhoea during the study. The study's end point is the occurrence of diarrhoea due to Clostridium difficile.

Intervention Type

Other

Phase

Not Specified

Primary outcome(s)

Not provided at time of registration

Key secondary outcome(s))

Not provided at time of registration

Completion date

15/06/2003

Eligibility

Key inclusion criteria

250 volunteers >65 years old.

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Senior

Sex

Not Specified

Key exclusion criteria

Not provided at time of registration

Date of first enrolment

16/06/2000

Date of final enrolment

15/06/2003

Locations

Countries of recruitment

United Kingdom

England

Study participating centre Box No 201A

Cambridge United Kingdom CB2 2QQ

Sponsor information

Organisation

Department of Health (UK)

Funder(s)

Funder type

Government

Funder Name

Cambridge Consortium - Addenbrooke's (UK)

Results and Publications

Individual participant data (IPD) sharing plan

IPD sharing plan summary

Not provided at time of registration

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Results article	results	15/02/2005		Yes	No