Xylitol clinical studies for prevention: In-school xylitol gummy bear snack study

Prospectively registered Submission date Recruitment status 07/02/2008 No longer recruiting [] Protocol [] Statistical analysis plan Registration date Overall study status 26/02/2008 Completed [X] Results Individual participant data **Last Edited** Condition category 30/07/2008 Oral Health

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 U54 DE14254

Study information

Scientific Title

School-based xylitol gummy bears snacks: a randomized controlled trial

Study objectives

Consumption of xylitol gummy snacks reduces S. mutans and Lactobacillus spp. levels

Ethics approval required

Old ethics approval format

Ethics approval(s)

University of Washington Internal Review Board. Date of approval: 27 March 2007 (ref: HS # 07-4857-B 01)

Study design

Randmised controlled trial

Primary study design

Interventional

Study type(s)

Prevention

Health condition(s) or problem(s) studied

Dental caries

Interventions

Randomisation was carried out at the level of individuals. Gummy bear snacks were packaged by the unit-dose and labeled specific to the students ID number which corresponded with the individual randomization.

Three-group design:

Group 1: Xylitol gummy bears, 11.7 g/day xylitol Group 2: Xylitol gummy bears, 15.6 g/day xylitol

Group 3: Maltitol gummy bear (control)

All groups consumed gummy bears 3 times/day for 6 weeks in the classroom during school hours only.

Intervention Type

Drug

Phase

Not Specified

Drug/device/biological/vaccine name(s)

xylitol

Primary outcome(s)

S. mutans and Lactobacillus spp. levels in plaque samples at enrollment (baseline) and 6 weeks (end of study).

Key secondary outcome(s))

No secondary outcome measures

Completion date

01/06/2007

Eligibility

Key inclusion criteria

Children attending First through Fifth grade at two elementary schools in rural Washington State, USA

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Child

Sex

All

Key exclusion criteria

Children with gastrointestinal problems

Date of first enrolment

01/04/2007

Date of final enrolment

01/06/2007

Locations

Countries of recruitment

United States of America

Study participating centre Dental Public Health Sciences

Seattle United States of America 98195

Sponsor information

Organisation

University of Washington (USA)

ROR

https://ror.org/00cvxb145

Funder(s)

Funder type

Research organisation

Funder Name

National Institute of Dental and Cranio-facial Research (NIDCR, USA; ref: U54 DE14254)

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 | 25/07/2008 | | Yes | No |