

Trial to investigate the maintenance effects of yawning on salivary cortisol levels

Submission date 19/04/2012	Recruitment status No longer recruiting	<input checked="" type="checkbox"/> Prospectively registered <input type="checkbox"/> Protocol
Registration date 26/04/2012	Overall study status Completed	<input type="checkbox"/> Statistical analysis plan <input checked="" type="checkbox"/> Results
Last Edited 23/09/2015	Condition category Other	<input type="checkbox"/> Individual participant data

Plain English summary of protocol

Background and study aims

There is considerable debate among scientists over why we yawn, and the mechanism of yawning is still not fully understood. About half of adults yawn after someone else yawns due to a phenomenon called contagious yawning. The aim of this study is to test whether levels of the hormone cortisol are higher during yawning and contagious yawning, just as cortisol levels are raised during stress and fatigue.

Who can participate?

Volunteers aged 18-65 will be recruited from students at Bournemouth University.

What does the study involve?

Participants are exposed to three conditions intended to provoke yawning – photos of people yawning, boring text about yawning, and a short video of a person yawning. We collect saliva samples from participants at the start and after yawning to measure saliva cortisol levels, and record the electrical activity of their jaw muscles via surface-placed electrodes. Questionnaires about yawning, anxiety and depression, general health, and demographic and health details are also collected from each participant.

What are the possible benefits and risks of participating?

The results of this study may improve our understanding of yawning and its role in many neurological disorders, and allow us to develop a diagnostic tool for neurological disorders. There are no risks involved. All participants will have the right to withdraw at any time and will be debriefed. All data is anonymised.

Where is the study run from?

Bournemouth University (UK).

When is the study starting and how long is it expected to run for?

June 2012 to December 2012.

Who is funding the study?

Bournemouth University and Santander plc (UK).

Who is the main contact?
Dr Simon Thompson
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Contact information

Type(s)
Scientific

Contact name
Dr Simon Thompson

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Additional identifiers

EudraCT/CTIS number

IRAS number

ClinicalTrials.gov number

Secondary identifying numbers
BU-PS5/10/11-PS1/3/12

Study information

Scientific Title
Thompson Cortisol Hypothesis: trial to investigate the maintenance effects of Yawning on salivary Cortisol levels - an observational study

Acronym
TCH-YawnCort

Study objectives
Cortisol levels rise during the yawning episode

Ethics approval required
Old ethics approval format

Ethics approval(s)

Bournemouth University Research & Ethics Committee, 05/10/2011, ref: BU-PS5/10/11-PS1/3/12
Amendments approved 01/03/2012

Study design

Observational study

Primary study design

Observational

Secondary study design

Cross sectional study

Study setting(s)

Other

Study type(s)

Other

Participant information sheet

Not available in web format, please use the contact details below to request a patient information sheet

Health condition(s) or problem(s) studied

Salivary cortisol levels

Interventions

Collection of saliva samples; collection of non-invasive surface-placed electrode data of electrical nerve impulses around jaw-line during yawning.

Observing participants' yawning, their level of saliva cortisol before and after yawning, and their electrical (nerve) activity around the jaw muscles at rest and during yawning. The duration of observation is determined by the time it takes to view the stimuli (about 20 minutes) and to yawn. If yawning occurs before all the stimuli have been viewed, then the observation period is shorter for the participant as the stimuli merely serve to elicit a yawn. We aim to conduct a longitudinal study at a later date once data has been analysed and results are known.

Intervention Type

Other

Phase

Not Applicable

Primary outcome measure

1. Salivary cortisol levels - collection of saliva deposited (spit into tube) by each participant into sample bottle at start of study then again immediately after yawning, or if no yawn elicited, at end of last stimuli presentation
2. Electrical nerve activity - non-invasive surface-placed electrodes around the jaw line receive data throughout the study (20 mins maximum), or only until yawning is elicited, whichever happens first

Secondary outcome measures

Demographic and descriptive details of participants age, gender, ethnicity

Overall study start date

01/06/2012

Completion date

31/12/2012

Eligibility**Key inclusion criteria**

1. 100 male and female volunteers aged between 18-65 years will be recruited from students at Bournemouth University using the computerised recruitment system (SONA), and Facebook
2. All participants will be properly consented according to code of conduct and research guidelines

Participant type(s)

Patient

Age group

Adult

Lower age limit

18 Years

Upper age limit

65 Years

Sex

Both

Target number of participants

100

Key exclusion criteria

1. Chronic fatigue
2. Diabetes
3. Fibromyalgia
4. Heart condition
5. High blood pressure
6. Hormone replacement therapy
7. Multiple sclerosis
8. Stroke

Date of first enrolment

01/06/2012

Date of final enrolment

31/12/2012

Locations

Countries of recruitment

England

United Kingdom

Study participating centre

Bournemouth University

Poole

United Kingdom

BH12 5BB

Sponsor information

Organisation

Bournemouth University (UK)

Sponsor details

c/o Dr Simon Thompson

Associate Professor

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Poole House (P315)

Fern Barrow

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England

United Kingdom

TS17 6QQ

Sponsor type

University/education

Website

<http://home.bournemouth.ac.uk/>

ROR

<https://ror.org/05wwcw481>

Funder(s)

Funder type

University/education

Funder Name

Bournemouth University (UK) ref: BU-26.08.11

Alternative Name(s)

BU

Funding Body Type

Government organisation

Funding Body Subtype

Universities (academic only)

Location

United Kingdom

Funder Name

Santander plc (UK) ref: SANTANDER-30.09

Results and Publications

Publication and dissemination plan

Not provided at time of registration

Intention to publish date**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	20/09/2012		Yes	No