

How anxiety and body signals lead to binge eating in daily life

Submission date 06/11/2025	Recruitment status No longer recruiting	<input type="checkbox"/> Prospectively registered
Registration date 10/11/2025	Overall study status Completed	<input type="checkbox"/> Protocol
Last Edited 07/11/2025	Condition category Mental and Behavioural Disorders	<input type="checkbox"/> Statistical analysis plan
		<input type="checkbox"/> Results
		<input type="checkbox"/> Individual participant data
		<input checked="" type="checkbox"/> Record updated in last year

Plain English summary of protocol

Background and study aims

This study explored how anxiety and the body's physical stress signals interact to trigger binge-eating episodes in daily life. The aim was to understand, in real time, how changes in anxiety, heart rate, and stress hormones are linked to urges to binge eat. By identifying these moment-clip patterns, researchers hope to inform new ways of preventing binge-eating episodes before they occur.

Who can participate?

Adults aged 18–55 years living in Germany were invited to take part. Participants were placed into one of four groups:

1. People with both Binge-Eating Disorder (BED) and Generalized Anxiety Disorder (GAD)
2. People with BED only
3. People with GAD only
4. Healthy volunteers with no mental health diagnoses

All participants needed to have a body mass index (BMI) between 18.5 and 40 kg/m² and be fluent in German. People were excluded if they had current substance misuse, psychosis, neurological illness, active suicidal thoughts, were pregnant, or were taking medication that could affect stress hormones or heart rate.

What does the study involve?

The study was observational, meaning no experimental treatment was given. Participants first visited the laboratory for a standard stress test to measure their body's response under pressure. They then took part in a 14-day monitoring period in their everyday environment. During this time, participants:

1. Received short smartphone surveys about their feelings and urges to binge eat several times a day
2. Wore a heart rate monitor to track changes in stress and relaxation
3. Collected saliva samples four times daily to measure stress hormone (cortisol) levels
4. Allowed their smartphone to record information such as movement and screen use

This combination of data helped researchers understand how anxiety and physical stress responses interact in real-world settings.

What are the possible benefits and risks of participating?

There were no direct benefits to participants, but the findings will help improve understanding of how anxiety contributes to binge eating, which could lead to better prevention and treatment approaches in the future. Risks were minimal and mainly related to mild discomfort or stress during the laboratory test, and the time commitment required for daily surveys and sample collection.

Where is the study run from?

The study was coordinated and conducted by Freie Universität Berlin, Germany, under the ethical oversight of the Charité University Medical Center's ethics commission.

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

The study began on 15 December 2024 and was completed on 12 September 2025.

Who is funding the study?

The study was funded by the Deutschlandstipendium National Scholarship Program, a government-funded initiative.

Who is the main contact?

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Contact information

Type(s)

Public, Scientific, Principal investigator

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Additional identifiers

Clinical Trials Information System (CTIS)

Nil known

ClinicalTrials.gov (NCT)

Nil known

Protocol serial number

6734

Study information

Scientific Title

Physiological correlates of anxiety-driven binge eating in a naturalistic setting

Acronym

ADBE

Study objectives

To investigate the real-time psychophysiological mechanisms (specifically, heart rate variability and cortisol) through which anxiety triggers binge urges and episodes in the natural environment, and to determine how these mechanisms are amplified in individuals with comorbid Generalized Anxiety Disorder (GAD) and Binge-Eating Disorder (BED).

Ethics approval required

Ethics approval required

Ethics approval(s)

approved 10/12/2024, Ethics Committee of Freie Universität Berlin (Kaiserswerther Str. 16-18, Berlin, 14195, Germany; +49 (0)17668328908; ethikkommission@charite-medizin.org), ref: 6734

Study design

Prospective observational longitudinal cohort study

Primary study design

Observational

Study type(s)

Other

Health condition(s) or problem(s) studied

Binge-Eating Disorder (BED), Generalized Anxiety Disorder (GAD)

Interventions

This study employed a prospective, observational longitudinal cohort design with four distinct arms: a comorbid Binge-Eating Disorder and Generalized Anxiety Disorder group (BED + GAD, n = 40), a BED-only control group (n = 20), a GAD-only control group (n = 20), and a healthy control group (n = 40). The primary purpose was etiological, aiming to understand the basic psychophysiological mechanisms of anxiety-driven binge eating. The design was non-randomized and unblinded, featuring a 14-day ecological ambulatory assessment period for real-time data collection, preceded by a laboratory-based stress reactivity assessment. The study did not involve an interventional treatment or investigational medicinal product.

All participants underwent the same assessment procedures, including: Diagnostic Clinical Interview (SCID-5), Laboratory Stress Test (Trier Social Stress Test), and a 14-day Ambulatory Monitoring Period (Ecological Momentary Assessment, continuous heart rate monitoring, passive smartphone sensing, and salivary cortisol sampling).

Intervention Type

Other

Primary outcome(s)

1. Momentary urge to binge (measured on a 0-100 visual analogue scale 5 times daily during the 14-day ambulatory period)
2. Binge episode occurrence (binary yes/no from event-contingent reports during the 14-day ambulatory period)

Key secondary outcome(s)

1. Heart rate variability (Root Mean Square of Successive Differences [RMSSD] in milliseconds), measured continuously during the 14-day ambulatory period and in response to the laboratory stress test
2. Salivary cortisol level (nmol/L), measured at four fixed times daily during the 14-day period and in response to the laboratory stress test
3. Contextual/behavioral markers (location variance, screen time) derived from passive smartphone sensing during the 14-day period

Completion date

12/09/2025

Eligibility

Key inclusion criteria

1. Adults aged 18–55 years
2. Body mass index (BMI) between 18.5 and 40 kg/m²
3. Sufficient fluency in German to complete study procedures
4. Meeting DSM-5 diagnostic criteria for one of the four defined groups:
 - 4.1. Comorbid Group (GAD+BED): Co-occurring Generalized Anxiety Disorder and Binge-Eating Disorder
 - 4.2. BED Control Group (BED-only): Binge-Eating Disorder and no lifetime anxiety disorder
 - 4.3. Anxiety Control Group (GAD-only): Generalized Anxiety Disorder and no lifetime eating disorder
 - 4.4. Healthy Control Group (HC): No current or lifetime psychiatric diagnosis

Participant type(s)

Healthy volunteer, Patient

Healthy volunteers allowed

No

Age group

Adult

Lower age limit

18 years

Upper age limit

55 years

Sex

All

Total final enrolment

120

Key exclusion criteria

1. Current substance use disorder, psychosis, or neurological disease
2. Active suicidality
3. Pregnancy
4. Current use of medications known to influence cardiac or endocrine function (e.g., beta-blockers, corticosteroids)

Date of first enrolment

15/12/2024

Date of final enrolment

01/09/2025

Locations**Countries of recruitment**

Germany

Study participating centre

Freie Universität Berlin

Kaiserswerther Str. 16-18

Berlin

Germany

14195

Sponsor information**Organisation**

Freie Universität Berlin

ROR

<https://ror.org/046ak2485>

Funder(s)**Funder type**

Government

Funder Name

Results and Publications

Individual participant data (IPD) sharing plan

Yes, there is a plan to share fully anonymized and de-identified individual participant data underlying the results reported in the final publication. This data will be made available to researchers beginning 6 months after the main findings are published. Access will be granted to researchers who provide a methodologically sound proposal for use in approved research. Proposals should be directed to the corresponding author via email, and requestors will be required to sign a data sharing agreement before the data is released.

IPD sharing plan summary

Available on request

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Participant information sheet	Participant information sheet	11/11/2025	11/11/2025	No	Yes