

Gut-brain interaction: exploring the link between bodily states and decision making

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Abstract

Physiological need states adaptively shape decision-making, yet its effects on distinct behavioural components remain poorly characterised in humans. In particular, behavioural dimensions such as impulsivity and motivation are often treated as stable traits, overlooking how bodily signals dynamically regulate behaviour in response to energetic demands. Here, we aim to clarify the importance of metabolic signalling in adaptive behavioural control of food and non-food behaviour in humans. Following an overnight fast, healthy participants completed tasks probing food- and money-related impulsivity, effort-based motivation, and valuation, alongside assessments of fasting duration and body composition. Fasting selectively increased impulsive responding for food (compared to monetary) rewards, as indexed by longer stop-signal reaction times. This food-specific increase in impulsivity was robust across analyses and was not explained by higher subjective valuation of food; rather, greater willingness to pay for food was associated with improved impulse control. Body fat percentage moderated fasting effects, suggesting an interaction between short-term energy deficit and long-term energy reserves. In contrast, effort exertion in an incentive-motivation task was strongly up-regulated by energy deficit in a domain-general manner, independent of reward type. Effort decreased with higher body fat percentage, but this effect was partially normalised by fasting. A composite measure of relative energy deficit, integrating fasting duration and fat mass, provided a parsimonious account of individual differences in effort spending and outperformed models based on fasting or body composition alone. Valuation measures further revealed a higher willingness to pay for food with increasing body fat percentage, but did not account for behavioural effects observed in impulsivity or effort tasks. Finally, questionnaire-derived trait measures showed limited correspondence with state-dependent behavioural changes, highlighting the dynamic nature of energy-dependent decision processes. Together, these findings demonstrate dissociable and complementary effects of energy status on impulse control and motivated behaviour, showing how adaptive decision making emerges from the interaction between acute metabolic signals and long-term bodily energy reserves.

1 **Introduction**

2 In order to survive, all animals need to continuously adapt their behaviour to the
3 constraints of their environment and the needs of their body (Flavell et al., 2022).
4 Deciding how and when to act, or how much energy to invest, will have different
5 consequences for fitness depending on the context and the energetic state. Yet,
6 behavioural dimensions such as impulsivity and motivation are often conceptualized
7 as static traits, dismissing how bodily signals dynamically shape their highly adaptive
8 nature.

9 Hunger is a perfect example of how caloric needs can control our behaviour. From an
10 ecological perspective, energy deprivation should bias decision-making systems
11 toward strategies that favour the acquisition of food. While the benefit of hunger in
12 motivating eating is evident (it thus satisfies the underlying caloric needs), the
13 influence of hunger might extends, beyond food consumption, to other behavioural
14 dimensions and reward domains. Consistent with this view, there is evidence that,
15 beyond food reward processing (Siep et al., 2009), short term fasting could alter
16 impulse control (Howard et al., 2020; Voigt et al., 2021), risk assessment (van
17 Swieten et al., 2023), temporal discounting (Skrynka & Vincent, 2019), or action
18 vigour (Hanssen et al., 2021; Pirc et al., 2019). Yet, the nature and domain (i.e. food
19 vs. non-food oriented) specificity of hunger's effects are still not well delineated, and
20 the role of energetic need on the flexible adjustment of behaviour beyond food intake
21 remains largely unclear, especially in humans (Bamberg & Moreau, 2025; Benau et
22 al., 2014). One possibility is that fasting induces a domain-general reduction in self-
23 control and an increase in motivation across reward types. Alternatively, fasting may
24 selectively prioritise biologically relevant rewards, such as food, while leaving
25 decision-making for abstract rewards relatively unaffected. Empirical evidence has
26 been limited by a focus on single reward domains, most commonly food, making it
27 difficult to distinguish between these accounts. Clarifying whether hunger produces
28 domain-general or domain-specific changes in behaviour is essential for
29 understanding the adaptive logic of state-dependent decision making.

30 At the neural level, cumulative animal literature shows that metabolic and hormonal
31 signals associated with hunger, such as insulin, ghrelin, or glucagon like peptide-1

(GLP-1), modulate various brain circuits and especially the dopaminergic system (Cassidy & Tong, 2017; Geisler & Hayes, 2023; Palmiter, 2007). Given the importance of midbrain dopamine signalling for reward processing, motivation, and impulsivity, these findings offer a mechanistic pathway to explain the importance metabolic signalling in the control of those behavioural dimensions. Recent studies further support this theory by showing that dopaminergic circuits are also affected by metabolic state in humans (Hanssen et al., 2021; Kullmann et al., 2021). Importantly, the same dopaminergic neurocircuits are involved in both food and non-food reward processing (Oren et al., 2022), suggesting that the behavioural consequences of their state-dependent modulation could generalize to other domains than food reward. Despite those advances, evidence for state-dependent modulation of the neural circuits underling food and non-food behaviour in humans is still lacking, further prompting the need to clarify the importance of metabolic signalling in adaptive behavioural control.

In addition to transient energetic states, individuals differ in their long-term energy reserves. Body fat mass, long term storage of energetic resources in the body, also influences metabolic and hormonal signalling relevant for reward processing and decision making. Notably, leptin, a hormone produced by the adipose tissue, can directly alter dopaminergic function in the brain (Fulton et al., 2006; Opland et al., 2010). Accordingly, obesity have been robustly associated with dopaminergic dysregulations (Kroemer & Small, 2016), supporting the theory that reward processing is affected by metabolic changes. Yet most studies treat adiposity as a static individual characteristic emerging as a *consequence* of food preferences and eating habits: the same dopamine-dependent behavioural dimensions outlined above are classically conceptualised as fixed psychometric traits defining food behaviour, and thus body composition. More precisely, higher body fat is robustly associated with higher impulsivity (Bartholdy et al., 2016; Garcia-Garcia et al., 2022; Mobbs et al., 2010) and, although less conclusively, with lower motivation (Giesen et al., 2010; Hanssen et al., 2022; Mathar et al., 2016). Importantly, those conclusions are derived independently of acute physiological challenges, leaving open the question of possible interaction between acute regulation by hunger and (chronic) body composition on dopaminergic function and thus adaptive behavioural control. In addition, lean body mass is the main driver of resting energy expenditure of an individual (Dulloo et al.,

2017). The ratio between fat and lean mass therefore reflects the relative energy reserve available, defining the urgency of the need to find food, and constraining potential effort expenditures. From an ecological perspective, behavioural responses to fasting should thus depend not only on immediate hunger signals but also on longer-term energy reserves and basal energy consumption, as the same degree of deprivation may carry different biological significance across individuals. The role of body-composition in the modulation of hunger-dependent behavioural adaptation is however, as of today, largely unexplored.

To address this gap, we assess in a series of experiments in humans how an acute energy deficit induced by fasting interacts with long term energy requirements, as reflected by body composition, to modulate impulsivity (inhibitory control) and incentive motivation (willingness to exert physical effort). Additionally, we test the relative influence of food and non-food incentives to explore the domain specificity of those adaptive behavioural modulations.

Results

Based on the hypothesis that metabolic needs adaptively regulates decision making, we aimed to determine the influence of the long- and short-term fluctuations in energy levels on the measured behavioural dimensions, contrasting food and monetary conditions to assess the generality or food specificity of those influences. We invited healthy participants (N=94) to come to the lab after an overnight fast. During the testing session, they had the opportunity to earn food and monetary outcomes in a set of behavioural experiments (see Fig. 1 and Methods) designed to assess their outcome-specific motivation (incentive force task) and impulsivity (stop signal task). After completing an auction task capturing the subjective valuation of the food vs. monetary rewards, they were allowed to consume their wins (eat the food and pocket the money) before filling in various questionnaires related to their drive, impulse control, and food behaviour. In addition to those behavioural and self-report markers, we recorded the body composition and fasting duration of each participant to assess their metabolic status.

Fasting selectively increases impulsivity for food

To quantify the variations in impulse control across our participants, we adapted the classic stop signal task (Verbruggen et al., 2008) as a first behavioural task (Fig. 1 top). Briefly participants were instructed to press a button or refrain from pressing it in response, respectively, to go and stop signals. Trials were organised in alternating blocs rewarding performances with either food or money, as clearly indicated by pictures of the outcomes at the beginning of each bloc and flanking the go/stop cues. For each participant and each bloc type, we computed the stop signal reaction time (SSRT) which captures the relative time needed to stop an ongoing response process, that is a longer SSRT indicates a weaker executive control and therefore a more impulsive responding.

A mixed effect model revealed a strong interaction between fasting duration and reward type ($p < 0.001$, Fig. 2a) which was further modulated by body composition (3 way interaction, $p = 0.044$). This effect was driven by a fasting x fat% in the SSRT in the food condition ($p = 0.040$) which could not be observed in the money condition ($p = 0.362$). To unpack this complex interaction, we calculated the difference in our impulsivity measures between the two conditions, $\Delta SSRT = SSRT_{food} - SSRT_{money}$.

A first analysis suggested a fasting x fat% interaction ($p = 0.022$) which could be understood by the fact that the relative impulsivity for food tend to increase with body fat % (short fast: $r = 0.261$, $p = 0.077$) but fasting mitigates this tendency (long fast: $r = -0.176$, $p = 0.264$; Fig. 2b). Strikingly, the impact of fasting on the $\Delta SSRT$ was stronger for participant with a lower body fat percentage (low body fat: $r = 0.396$, $p = 0.030$; high body fat: $r = 0.308$, $p = 0.098$, Fig. 2b), suggesting that energy reserves modulate the influence of fasting induced energy deficit. The body composition modulatory effect, however, reduced to a simpler but highly significant effect of fasting ($p < 0.001$, Fig. 2d) when confounding factors were included in the linear model, confirming the strong influence of energy deficit on food-specific impulse control. This follow-up analysis also revealed that the relative impulsivity for food decreased with the willingness to pay for food ($p = 0.005$, Fig. 2e). While slightly counterintuitive, as one could expect that a higher subjective valuation of food should yield a more impulsive behaviour, this falls in line with previous results demonstrating that impulse control improves for higher reward prospects (Giuffrida et al., 2023).

Together, these results demonstrate that changes in physiological state induced by fasting modulate impulse control for food rewards. Further, fat reserves moderated this dynamics, hinting at a more complex interplay between short and long term energy status on cognitive control. Critically, those effects could not be explained by an increase in the subjective value of food which, on the contrary, improved performances.

Relative energy deficit drives motivation to effort

In order to assess the role of metabolic state on effort regulation, we adapted also classic incentive motivation task (Pessiglione et al., 2007), Fig. 1 b as our second behavioural measure. Briefly, participants held a dynamometer in their hand which they could squeeze to raise the level of a thermometer-like scale on the screen and thus increase their chances of earning a reward. The colour of the scale indicated the rate at which the thermometer would rise, allowing participant to gauge their behaviour as a function of the effort required to fill the scale up (difficulty level) and therefore the actual cost/benefit ratio at stake. As for the stop signal task, trials were organised in alternating blocs (indicated by food or money pictures displayed next to the scale) prescribing which type of reward performance will be translated to at the end of the session.

Performances were renormalised to each participant's strength before entering a linear model fitted for each cue type and including an intercept, the difficulty level, and the trial number. A group level analysis showed that, unsurprisingly, participants exerted less force and were therefore ready to forego their chances of reward, as difficulty increased ($p < 0.001$; Fig. 3a). While the type of reward at stake also affected the performances (interaction with intercept $p < 0.001$; difficulty: $p < 0.001$; trial: $p = 0.011$), this was mainly driven by the difference in subjective valuation of the outcomes (all $p < 0.033$) and not by an interaction of metabolic factors (all $p > 0.446$). Accordingly, we averaged the two conditions before exploring the influence of the energy status on behaviour. Interestingly, fluctuations in average performances were explained by an interaction between fasting duration and body composition ($p = 0.006$, correcting for sex differences). Indeed, while overall performances drastically declined with body fat percentage ($r = -0.34, p = 0.001$, Fig. 3d), this effect was partially mitigated by fasting (short fast, $p < 0.002$; long fast, $p = 0.9$; Fig. 3c) suggesting that fasting could partly

normalise the negative influence of fat mass on motivation. Looking at the effect of fasting in subgroups of participants split by their body fat % (Fig. 3d) provides a possible explanation for this complex pattern. Indeed, motivation appears to be related to fasting when appraised not by its duration but in terms of the relative energy deficit it induces. To test this hypothesis, we estimated the amount of calories burned during the fasting period relative to the amount of calories stored in the body as fat (see methods). This measure of relative energy deficit (RED) was strikingly similar to our motivation measure (Fig. S1): participants with a high body fat percentage (*i.e.* a slow metabolic rate and large reserves) had a low RED which slowly increased with fasting; in contrast, participants with a lower body fat percentage (*i.e.* burning calories fast, with low energy stocks) had a higher RED which was less consistently affected by fasting as body composition dominated the variations between individuals. Accordingly, RED strongly predicted effort spending ($p < 0.001$, Fig. 3e) and provided a more parsimonious explanation than the fasting x body fat interaction ($\Delta \text{BIC} = 2.1$).

In summary, our results demonstrate that effort spending is powerfully up-regulated by short term energy deficits. In contrast to impulse control measurements reported above, this metabolic effect appears very pervasive and does not depend on outcome quantity or identity which additionally influence motivated behaviour.

Changes in willingness to pay for food does not explain impulsivity nor effort variations

During the first two tasks, capturing respectively impulsivity and motivation, participants were instructed that good performances will earn them food and money tokens, depending on blocs, to be exchanged for actual food items and money at the end of the experiment (Fig. 1 bottom). The subsequent auction task was framed as an opportunity to reallocate the tokens they won by betting on 30 pairs of fortune wheels. On a given trial, each of 10 tokens could be placed either on a wheel associated with one of 30 of the available food items, or on another associated with an equivalent monetary value. As each token granted a 10% chance of the fortune wheel to stop on a win, participants could decide to either place all their bets on one wheel, and thus ensure to win the associated outcome, or split their bets and have a chance to win both rewards.

1 The total amount of tokens allocated to snacks, irrespective of the strategy, reflected
2 the willingness to pay (WTP) for food. While a linear model including fasting
3 duration and body composition did not yield any significant effect, WTP correlated
4 with body fat percentage when tested separately ($p=0.040$, Fig. 4).

5 Importantly, prospects of reward are strong predictors of both motivation and
6 impulsivity, higher incentives being associated with better performances in both of
7 those measures. While WTP for food was not robustly modulated by metabolic state
8 in our data, it could still partially explain the influence of bodily-state on
9 performances. To control for this potential confound, WTP was systematically
10 included as a covariate in the analysis of the stop signal and the incentive motivation
11 tasks. However, adding this control did not affect the results. In conclusion, the
12 influence of metabolic state on behaviour we identified above cannot be simply
13 explained by a change in the subjective value of food rewards, and rather reflect a
14 fundamental regulatory process of action by the physiological state.

15 ***Questionnaires capture static but not adaptive behavioural phenotypes***

16 We performed an exploratory factor analysis to summarize the 15 questionnaires
17 filled in by each participant, yielding five factors (Fig. 5 top). The first two ones
18 related to food behaviour, and more precisely to sensitivity to external food triggers
19 (“uncontrolled eating”, similar to the previously reported factor (Vainik et al., 2015)),
20 and the active tendency to restraint one’s food behaviour (“cognitive restraint”). The
21 next two factors related to more general behaviour, namely sensitivity to rewards
22 vs. punishment (“drive”), and impulsive tendencies (“impulsiveness”). The last factor
23 captured associations between depressive traits and compulsive tendencies along with
24 food coping strategies (“compulsiveness”).

25 To understand the link between each participant’s traits, as captured by the self-report
26 questionnaires, and their actual implementation in actions, we then correlated the
27 factor scores with the task performances (Fig. 5 bottom).

28 Concerning the stop signal task, we found a single correlation between accuracy in the
29 go condition (ie. correctly following the arrow direction) and the impulsiveness
30 factor. While surprising at first, as SSRT is the metric expected to capture impulsivity
31 in this task, this finding is in line with previous reports showing that inaccurate action

1 responses were more predictive of trait impulsivity than action inhibition (Portugal et
2 al., 2018). A more direct explanation for the lack of correlation between trait
3 impulsivity and the SSRT performances is that the latter is a highly dynamic
4 phenotype continuously adapting to physiological state and therefore unlikely to be
5 captured by questions intended to apprehend static qualities. Overall, our results
6 highlight the fundamental shortcomings of questionnaires when quantifying fast
7 fluctuating behaviours such as (fasting-dependent, food-specific) impulsivity.

8 The force task evidenced a simpler, domain general, behavioural marker summarized
9 by the average effort performance. This metric positively correlated with the “drive”
10 factor, reflecting the well-known importance of reward sensitivity in the regulation of
11 effortful actions. Effort was further anti-correlated with the “uncontrolled eating”
12 factor, which could be explained by the fact that this factor also captured a negative
13 drive dimension (ie. sensitivity to punishment) which partially mirrored the “drive”
14 factor. Finally, average effort was negatively associated with our last factor we
15 labelled “compulsiveness”. This factor also loaded depression and stress
16 questionnaires and could reflect a more general mood downregulation that would
17 negatively affect motivation.

18 Finally, in the auction task, proportion of certain bets (the number of snacks secured
19 by placing all tokens on the food wheel) positively correlated with cognitive restraint
20 and negatively with impulsivity factors, suggesting that those traits can translate into
21 risk aversion when implementing actual food choices and are therefore also pertinent
22 to understand weight regulation.

23 Next, we explored the link between trait dimensions and body weight. Almost all
24 factors correlated with body composition, indicating that higher fat percentage is
25 related to a higher sensitivity to food cues along with stronger attempts to restraint
26 such urges, and more compulsiveness. The factors we identified are in line with
27 previous reports, in particular uncontrolled eating (Vainik et al., 2015). Body fat was
28 also associated with a lower drive, hinting that the motivational deficit we measured
29 in the effort task might not be due only to a dampened fasting effect in the more
30 corpulent participants but also to a more general lack of reward sensitivity.

31 Interestingly, “impulsiveness” did not correlate with body composition. While various
32 measures of impulsivity have been associated with body weight before, these

associations originate from clinical populations suffering from food related disorders such as morbid obesity or binge eating disorder and might not hold for the general population (Bartholdy et al., 2016; Lavagnino et al., 2016).

Altogether, our observations show that while self-report questionnaires can capture some behavioural phenotypes and their importance for weight regulation, they also overlook the dynamical aspects of behaviour and therefore fail to fully capture the adaptive nature of metabolic regulation.

Discussion

In this study, we explored in healthy participants the modulatory role of metabolic state on various facets of behaviour. First, we revealed that fasting increases impulsivity selectively for food, an effect which was dampened by body fat percentage. Second, we identified that energy deficit drives a global motivation to exert effort, an effect also dependent on body composition. Together, our results demonstrate that behaviour is highly adaptive and depends on a complex interaction between short-term and long-term energy state variations.

Concerning impulsivity, our results generalise recent findings that identified a difference in inhibitory control between fed and fasted states in a food-related task (Howard et al., 2020) by showing that impulsivity progressively increases with fasting duration. We also demonstrated that the effect of fasting was specific to food rewards, confirming the existence of a domain specific impulsive behaviour (Zhang et al., 2017). Notably, another work, relying on a different measure of impulsivity (information sampling), reported non-food related changes in impulsivity with hunger (Voigt et al., 2021). This discrepancy can be resolved by acknowledging that impulsivity is a multifaceted construct encompassing distinct neurobehavioural mechanisms (Mobbs et al., 2010). In light of this literature, our results suggests that state-dependent modulation of impulsive behaviour might be domain specific, or not, contingent on the underlying process actually measured. Interestingly, the stop signal task we used in this study is recognized to yield highly volatile results within individuals (Thunberg et al., 2024), and to offer only mediocre correlation with obesity (Bartholdy et al., 2016). According to our data, these negative results could be explained by the fact that fasting duration has a strong impact on behaviour, a

1 confounding factor systematically neglected in the literature. Collectively, these
2 results highlight that to understand the exact relation between body composition and
3 impulsivity, future studies should carefully account for the fluctuations (or the lack
4 thereof) induced by hunger state.

5 Concerning motivation, our data confirms previous reports showing an augmentation
6 of effort to obtain food with increasing hunger (Arumäe et al., 2019; Pirc et al., 2019;
7 Ziauddeen et al., 2012). Notably, those studies relied solely on food rewards,
8 suggesting a food specific effect. In contrast, our study revealed that fasting increases
9 vigour regardless of the type of outcome, suggesting a general motivational effect. We
10 also found a negative correlation between body fat and motivation. While this result
11 align with previous studies (Mansur et al., 2019; Mathar et al., 2016), we however
12 suggest a different interpretation to this relation: instead of being a hallmark of
13 adiposity that could arguably be attributed to dopaminergic dysregulation, the
14 reduction in vigour in participants with higher body fat could be due to a reduced
15 influence of fasting. More precisely, we propose that the caloric deficit induced by
16 fasting needs to be put in perspective with the energy reserve of the body: with higher
17 body fat, fasting is less dramatic for survival and thus has a lower impact on
18 behaviour. Mechanistically, this could be explained by the observation that, in
19 rodents, fasting induced increase in motivation depends on the switch to a ketosis
20 metabolic regime, the timing of which depends on fat reserves (Koubi et al., 1991).
21 Contrasting with this conclusion, other studies found an increase in motivation with
22 BMI (Epstein et al., 2007; Giesen et al., 2010; Hanssen et al., 2021). Differences in
23 the experimental design might explain these discrepancies. First, those studies
24 compared lean to obese populations, while we only tested healthy weight participants.
25 Morbid obesity is associated with numerous metabolic and neural changes that could
26 disrupt the normal regulation of motivation by bodily state. Second, they offered to
27 win high-calorie snack foods, while our selection contained healthy options. As
28 motivation seems to be dependent on the type of food at stake (Mathar et al., 2016), it
29 is possible that a more granular approach would reveal distinct effects depending on
30 the macronutrient composition of the food reward. More generally, our work
31 underscore the need to account for variations in energy needs to correctly identify
32 motivational phenotypes.

1 While we can only postulate about the neurobiological mechanisms implementing the
2 adaptive behavioural regulation we identified, our findings are consistent with the
3 metabolic regulation of dopaminergic circuits (Hsu et al., 2018), critical for the
4 control of the willingness to exert effort (Hsu et al., 2018) and impulse control (Eagle
5 & Baunez, 2010; Winstanley, 2011). Indeed, hunger related hormones, such as
6 ghrelin, potentiate dopaminergic activity and thus motivation, while leptin, produced
7 by the adipose tissue, tend to dampen it (see Geisler & Hayes (2023) for a review).
8 Furthermore, obesity has been robustly associated with dopaminergic dysregulations,
9 although the exact relation with body weight regulation is still unclear (Janssen &
10 Horstmann, 2022). A better understanding how the various metabolic signals interact,
11 at their respective timescales, to control dopaminergic function is needed to decipher
12 state-dependent regulation of behaviour.

13 From an ecological perspective, our data are in line the hypothesis that energy
14 requirements shift behaviour toward food acquisition by increasing effort expenditure
15 and biasing decisions toward more immediate actions. While we also found that
16 adiposity was associated with a lower motivation, we proposed that this only reflect a
17 blunted or delayed effect of fasting in participant with higher energy reserves. Our
18 interpretation is thus at odds with the classical view in the literature, associating
19 obesity with a set of “traits” or cognitive phenotypes assessed by psychometric
20 questionnaires (Gerlach et al., 2014; Robinson et al., 2020). The importance of fast
21 metabolic influences, overlooked by this approach, could explains why the
22 multifaceted relation between body composition and cognitive dimensions still
23 remains poorly predictive (Vainik et al., 2019). Here, we challenge the idea that body
24 weight management is caused by a static behavioural phenotype and suggest a reverse
25 causality: higher body fat weakens the ability to adapt behaviour to rapid metabolic
26 fluctuations. We further argue that behaviour needs to be approached as a dynamical
27 process tightly regulated by physiological states such as energy levels. Follow-up
28 studies could explore how environmental factors, such as food availability, come in to
29 play to affect decision making and the resulting metabolic trajectory.

1 **Methods**

2 ***Participants***

3 A total of 116 volunteers were recruited from a local database. A first screening for
4 exclusion criteria allowed us to identify volunteers either being underweight (BMI <
5 18.5, n = 2), obese (BMI > 30, n = 6), following a restrictive diet (n = 1), having a
6 physiological condition that could affect their food behaviour (n = 7), suffering or
7 having a history of psychiatric disorder(s) (n = 3), scoring high on depression scale
8 (BDI>17, n = 3), or being pregnant (n = 1). One participant was further excluded for
9 failing to come fasted the day of the experiment. In total, 94 healthy participants
10 underwent the full experimental protocol.

11 ***Experimental design: behavioural***

12 Before being invited, participants first had to fill in all questionnaires related to food
13 or used for exclusion utilising a dedicated online platform (LimeSurvey) hosted in our
14 institute. They were then requested to refrain from consuming any food or caloric
15 beverages after 10 PM the day before coming to the lab for a single testing session
16 starting at 8AM. Upon arrival, participants were familiarized with the general course
17 of experiment. In particular, they were told they will need to perform two behavioural
18 tasks (a stop signal task and incentive motivation task, order counterbalanced across
19 participants) allowing them to earn, depending on the experimental block, “food
20 tokens” or “money tokens”. Critically, we informed them that those tokens could
21 respectively be traded afterwards for actual food items and cash from a selection of
22 snacks and a cashbox on display in the room. In addition, participants had to rate their
23 liking of each of the 30 food items using a visual analogue scale before starting the
24 behavioural tasks. This setup ensured that the framing of the tasks in “food” and
25 “money” blocs was clearly mapped onto real and concrete outcomes of different
26 nature. Moreover, we explicated that as they filled in the remaining questionnaires
27 afterwards they would be allowed to consume the food they won. In addition, they
28 would have to stay in the lab until the end of the experiment, at 11 AM, with no
29 access to any other food. This helped preventing strategies based on the
30 exchangeability of the outcomes, *e.g.* buying food with the money earned or stashing
31 the snack for later use or trade. Unbeknownst to the participant before the end of the
32 effort and stop signal tasks, the conversion of the collected tokens into actual rewards

was carried out using a “fortune wheel” auction task. This procedure allowed us to measure the willingness of the participants to pay for food, i.e. to quantify the subjective value of the two reward types relative to each other.

Throughout the session, participants had to rate on a visual analogue scale their level of hunger, thirst, and satiety for a total of seven rating blocks. Finally, the session ended with a series of anthropometric measurements to assess the body composition of the participants and estimate the muscle size of their forearm.

All procedures were approved by the ethics committee of the University of Cologne and we obtained written informed consent from all participants prior to the experiment.

Stop signal task

The stop signal task is adapted from the “STOP-IT” open-source software developed by Verbruggen et al. (2008). Participants were seated in front of a computer screen and were asked to keep their index finger in the centre of the left and right arrow keys of the keyboard until a white arrow appears (go-signal). In this case, they were instructed to quickly indicate the direction of the arrow with their index finger by pressing the appropriate arrow key (go-trials). In 25% of the trials, the white arrow turned blue (stop-signal) and participants were told to withhold their response (stop-trials). After a short practice block (32 trials), participants completed six experimental blocks with 96 trials each. Every new block was initiated by the presentation of a stimulus picture announcing the incentive to play for in this block. The incentive type varied from block to block. The incentive type of the first block (food tokens or money tokens) was counterbalanced across subjects. In between blocks, the word “pause” was centred in white letters on a black screen for 15 seconds.

Every experimental trial started with the presentation of a small white dot (fixation sign) in the centre of a black screen. On the left and right side of the fixation sign two small stimulus pictures of the incentive were displayed. After a jittered delay between 850 ms, the white dot was replaced by a white arrow after a random time of at least 500 ms (intertrial interval) and at most 1350 ms. The visual stimuli of the incentive were not displayed in the practice part. While there is no feedback presentation in the experimental part, participants were offered the information about the success of their responses in the practice part to increase their consciousness of performance and

1 improve learning. In total the go stimulus and the possibly following stop-signal were
2 displayed for a maximum time of 1500 ms (maximal reaction time) or until the
3 response occurred.

4 The delay of a stop-signal (SSD) was adapted according to the participant's
5 performance by using the staircase tracking procedure (i.e., Jahfari et al, 2011,
6 p. 6892). Each incentive type had its own staircase: For example, the second food-
7 token block began with the SSD of the first food-token block. The practice and the
8 first experimental block were initialized by a SSD of 250 ms, but the first SSD of the
9 second experimental block was given by the last SSD of the first experimental block.
10 In the first two experimental blocks of each incentive type, the initial SSD (i.e., 250
11 ms) was increased by 50 ms, when successfully withholding a response in a stop-trial,
12 and decreased by 50 ms when responding to a go stimulus in a stop-trial. In the last
13 block of each incentive type, the SSD increased or decreased by one thirtieth of a
14 second. By using this staircase procedure, the probability of successful stop
15 performance was about 50% and led to a maximal competition of go and stop
16 processes. The idea of a competition between the terminations of the respective two
17 processes is called horse race model (Logan & Cowan, 1984).

18 **Effort task**

19 The effort incentive motivation task is adapted from (Pessiglione et al., 2007). First,
20 the participants are given a hand dynamometer (Vernier Software & Technology) in
21 their dominant hand. During an initial calibration phase, they are given three attempts
22 (4 seconds each) to squeeze the device as hard as he can. The maximal force is then
23 used to define the difficulty of the effort task as describe below. Each trial started with
24 the display of the type of outcome at stake (food or money) and a thermometer-like
25 scale. By squeezing the handle (within 3s), the participant could then fill up the
26 thermometer, the “mercury” height being proportional to the exerted force. Critically,
27 we instructed the participants that the higher the mercury, the more tokens, and
28 therefore the more food of money, depending on the cue, they would obtain at the
29 end. In order to assess the subjective motivation to effort, the reward cue (picture of
30 food items of cash) was kept constant but the amount of force required to fill the
31 thermometer up the difficulty was systematically varied from trial to trial. More
32 precisely, reaching the top of the scale required 90% (easy), 115% (medium), or
33 140% (hard) of the calibration force. This difficulty level was indicated by the colour

1 of the “mercury” (respectively green, orange, and dark red) to allow the participants
2 to plan their effort before pressing. While this implementation contrasts with the
3 original experiment, where the reward at stake rather than the force scaling was
4 altered, both task variations effectively modulate the conversion rate between the
5 exerted force and the amount of reward earned which is the main determinant of
6 motivation to effort behaviour. This modified design prevented the use of explicit
7 quantities of tokens as cues, making it more similar to the stop signal task and
8 allowing us to give the same amount of tokens to all participants in the auction task
9 without arousing too much suspicion in the participants.

10 Similarly to the stop signal task, the complete task consisted of one practice bloc (12
11 trials) followed by an alternation of 20 food and money blocs (12 trials each), each
12 starting with a full screen picture of the outcome to come, for a total of 240 trials. The
13 bloc order was counterbalanced across subjects.

14 **Auction task**

15 The auction task was run last and allowed the participants trade the tokens they earned
16 for actual food snacks and cash.

17 We first informed all participants that they won 300 tokens during the force and stop-
18 signal tasks and that they now had the opportunity to bid on food and monetary items
19 in a sequence of 30 lotteries. Each lottery consisted of two independent fortune
20 wheels, one associated with a fixed amount of money (0.70€), the other to a snack of
21 equivalent value changing in each trial (so 30 different snacks in total). The position
22 and order of the snacks were randomized across trials and participants. In a given
23 trial, participants had to allocate 10 tokens between the wheels in order to increase the
24 probability of winning the associated outcome using the rule one token = 10% chance.
25 Tokens could be moved to and between the wheels using the arrows of the keyboard,
26 each bet being displayed as a slice (1/10th) of the wheel being coloured. Therefore,
27 the participants had the choice between securing one of the option (put all tokens on
28 one wheel and none on the other, outcome probability = 100% / 0%) or try and win
29 both outcomes with a risk of winning nothing (e.g. put 7 tokens on the food wheel and
30 get a 70% chance of getting the snack and 30% of earning the cash). After bidding on
31 all the lotteries, 6 out of 30 were actually implemented by spinning the wheels on the
32 screen. Although the outcome appeared random, the result was biased to ensure that

1 the participant won its 3 most desired snacks in order to observe the following food
2 consumption, during the questionnaires.

3 The rationale for this task is two-fold. First, by letting participant bet their tokens
4 concurrently on food or money rewards, we could measure the relative preference of
5 the participants for the two types of outcome (*i.e.* their willingness to pay for food, or
6 relinquish food for money). Second, the lottery implemented here allowed to
7 indirectly assess the risk aversion or seeking profile of the participant. Critically, the
8 two dimensions were relatively independent: the preference for one outcome could be
9 expressed either by balancing the bets within trial (risky behaviour), or betting all
10 tokens on one of the outcome at each trial (risk averse behaviour) but alternating
11 across trials according to their inclinations.

12 **Questionnaires**

13 Participants filled in a total of 15 questionnaires relating to impulsivity, compulsivity,
14 control, drive, and eating behaviour (see Tbl. 1 for details). Questionnaires related to
15 disorders or food behaviour were completed online to respectively allow for exclusion
16 before coming to the lab and avoid making participants too self-conscious about their
17 food behaviour as we observed their snack consumption. Other questionnaires were
18 filled on paper on the testing day.

19 **Hedonic ratings**

20 A picture of each snack was displayed in the centre of the screen above the question
21 “How strongly do you like or dislike this item?” (“Wie stark ist Ihre Vorliebe bzw.
22 Abneigung”). On the left side was a labelled hedonic scale (Lim et al., 2009) ranging
23 from “greatest imaginable dislike” (“Stärkste Abneigung, die vorstellbar ist”) at the
24 bottom to “greatest imaginable like” (“Stärkste Vorliebe, die vorstellbar ist”) at the
25 top. Participants moved the cursor to indicate their preference on the scale and clicked
26 to validate their response, and so on until all items were rated.

27 **State ratings**

28 Subjective state (hunger, satiety, and thirst) was measured using a visual analogue
29 scale. For each dimension, a question on the screen (“How hungry/sated/thirsty are you
30 at the moment?”; “Wie hungrig/satt/durstig sind Sie momentan?”) prompted the
31 participant to rate their current state. To this end, they used the mouse to move a
32 cursor to any point between the left (“not hungry/sated/thirsty at all”; “gar nicht

1 hungri/satt/durstig”) and right (“very hungry/sated/thirsty”; “sehr
2 hungri/satt/durstig”) anchors which best reflected their feeling, and validated their
3 response using a left click.

4 **Software**

5 All experiments were run using the Psychtoolbox 3.0 (<http://psychtoolbox.org>) on
6 Matlab (The Mathworks Inc.). The measurements from the hand dynamometer were
7 captured using a homemade Matlab code ([https://github.com/lionel-rigoux/vernier-](https://github.com/lionel-rigoux/vernier-toolbox)
8 toolbox).

9 **Body composition**

10 We first measured body composition using a SECA mBCA 515/514 impedance scale
11 which provided, in addition to the total body weight, the absolute fat mass, fat
12 percentage, fat-free mass, and skeletal muscle mass (full body and limb by limb) of
13 the participant. We also measured the participant’s height to compute their Body
14 Mass Index (BMI) according to the formula $BMI = bodyweight/height^2$ (all
15 measurements in SI units). To control for the natural difference in adiposity between
16 males and females, we also computed a “normalized fat percentage” score by
17 demeaning the fat percentage within each gender group.

18 ***Statistical analyses***

19 **Maximal Physiological Force**

20 The maximal force a muscle can generate is directly proportional to the number of
21 fibres it contains, which can be approximated by the muscle cross sectional area
22 (CSA). In other words, we can predict the maximal physiological force (MPF) of a
23 participant by approximating, using anthropometric measurements, the CSA of their
24 muscles (Maughan et al., 1983) — in our case, the muscles of the forearm in charge
25 of gripping. Practically, we first measured the length (L , between the ulna’s head and
26 the styloid process) and maximum circumference (C , 1/3rd from the ulna’s head) of
27 the forearm. Then, we used calipers to measure the skinfold (S , skin + fat layers) of
28 the interior and exterior sides of the forearm. Approximating the forearm geometry
29 with a cylinder, the CSA can then be computed as the total area of the limb section
30 minus the fat + bone area (Heymsfield et al., 1982):

$$1 \quad CSA = \frac{(C - \pi S)^2}{4\pi} - B$$

2 Where C and S are in cm, and B , the bone area, in cm^2 (set to 1.8 according to E. S.
3 Hsu et al. (1993)). Finally, the MPF can be calculated by simply scaling the CSA:

$$4 \quad MPF = CSA \times F$$

5 where $F = 2.45 + 0.288 \times L$ was previously measured in a large cohort of adults
6 (Neu et al., 2002).

7 We validated our measure by regressing the MPF on the body composition measures.
8 As expected, the MPF was highly predicted by the participants' fat-free mass, which
9 is mainly composed by skeletal muscles (main effect: $p < 0.001$). Critically, this
10 relation was not affected by the fat mass (interaction term: $p = 0.398$) nor,
11 alternatively, by the fat mass percentage (interaction term: $p = 0.545$).

12 **Relative energy deficit**

13 In order to estimate the relative energy deficit induced by fasting in our participant,
14 we first estimated the basal metabolic rate (BMR), which represent the number of
15 calories burned by the body, at rest, during a day. Using the Mifflin-St Jeor equation,
16 the BMR can be derived from the lean mass (LM) measured with the impedance scale:

$$17 \quad BMR = 370 + (21.6 \times LM)$$

18 From this, we computed the caloric deficit ($E_deficit$) induced by a fast of T hours:

$$19 \quad E_deficit = BMR \times T/24$$

20 Assuming that a gram of body fat stores around 9 kcal, and using the absolute fat mass
21 (FM) measured with the impedance scale, the total energy stored in the body fat
22 ($E_available$) is given by:

$$23 \quad E_available = 9 \times 1000 \times FM$$

24 Finally, the relative energy deficit (RED) can be expressed as the ratio between the
25 caloric deficit induced by fasting and the calories available in the fat storage:

$$26 \quad RED = E_deficit / E_available$$

1 **Stop signal reaction time**

2 We first computed the average reaction time (RT) in the GO condition (RT_{GO}) as an
3 indicator of the response process efficiency. We used a geometric mean to
4 counterweight the heavy tail of the RT distribution and therefore avoid an
5 overestimation. To obtain a signature of the relative efficiency of the inhibition
6 process, we estimated the Stop Signal Reaction Time (SSRT; see Matzke et al. (2018)
7 for a review) from the STOP trial behaviours as follow: We started by performing a
8 logistic regression to predict correct STOP responses as a function of the SSD. To this
9 end, we located the inflection point of the fitted logistic function to obtain the SSD for
10 which the participant had a 50% chance of stopping, known as the critical SSD
11 (SSD_{crit}). Finally, SSRT was computed as the difference between the inhibitory and
12 response latencies: $SSRT = SSD_{crit} - RT_{GO}$. Note that a shorter SSRT correspond to
13 a relatively faster suppression of the action and therefore a more efficient inhibitory
14 process.

15 We checked the performances of all subjects for signs of failures of the experimental
16 procedure. First, performances in this task are usually close to nominal and an error
17 rate higher than a few percents indicates that the participant did not fulfilled the task
18 properly. Accordingly, we excluded participants who, in the GO condition, responded
19 incorrectly in more than 10% of the trials, or failed to respond at all in more than 25%
20 of the trials. We also excluded participants who achieved a proportion of correct
21 inhibition outside of the [40% - 60%] range, as it indicated that the staircase did not
22 converge. In total, 7 participants were excluded from any further analyses relying on
23 the SSRT measurements.

24 **Effort task**

25 The maximum of the force profile was extracted for each trial. For each subject and
26 cue, this peak force was fitted with a linear model to capture the average force
27 (intercept) and the incentive effect. The respective beta estimates where then entered
28 in follow-up linear models to infer group-level statistics related to between subject
29 effects. When appropriate, the subject-level statistics were averaged across cues to
30 derive domain general influences.

31 One subject was excluded from the analyses relying on the effort task due to a
32 calibration error.

1 Auction task

2 For each of the 30 pictures, the response was measured as the relative number of
3 tokens (between 0 and 1) bet on the food item. We then computed the mean and
4 variance of the responses across all trials reflecting respectively the general
5 preference for food and the strategy used for betting. Indeed, on the one hand, the
6 most risky strategy is to always split the bet proportionally to one's mean preference
7 on each trial, e.g. to bet 5 tokens on food and 5 tokens on money if the participant has
8 no bias toward one type of reward. In that case, the variance across trials will be 0 (in
9 our example, the response is always 0.5). On the other hand, the safe strategy is to bet
10 all the tokens either on the food item or on the money wheel in a given trial and to
11 alternate the type of reward thus 'secured' across trials. In our example above, a risk
12 averse participant would bet all tokens on the food wheel in 15 trials and on the food
13 item in also 15 trials, effectively expecting to receive a balanced amount of rewards of
14 both types. In that case, the response will follow a binomial distribution (response is
15 always be 0 or 1) with a variance $mean(response) \times (1 - mean(response))$. By
16 dividing the measured response variance by this maximal theoretical variance given
17 the empirical reward bias, we obtain a standardized measure (between 0 and 1) of
18 risk aversion that is independent on the reward preference:

$$19 \quad riskaversion = var(response) / (mean(response) \times (1 - mean(response)))$$

20 Finally, for each participant, the responses were entered in a linear model that
21 included as regressors the hedonic and familiarity ratings of the participant as well as
22 the price and caloric content of each picture. The regression weights therefore
23 indicated the individual drivers of the preference for the food bets.

References

- Arnau, R. C., Meagher, M. W., Norris, M. P., & Bramson, R. (2001). Psychometric evaluation of the Beck Depression Inventory-II with primary care medical patients. *Health Psychology, 20*(2), 112–119. <https://doi.org/10.1037/0278-6133.20.2.112>
- Arumäe, K., Kreegipuu, F., & Vainik, U. (2019). Assessing the Overlap Between Three Measures of Food Reward. *Frontiers in Psychology, 10*, 883. <https://doi.org/10.3389/fpsyg.2019.00883>
- Bamberg, C., & Moreau, D. (2025). Acute effects of fasting on cognitive performance: A systematic review and meta-analysis. *Psychological Bulletin, 151*(9), 1147–1169. <https://doi.org/10.1037/bul0000492>
- Bartholdy, S., Dalton, B., O'Daly, O. G., Campbell, I. C., & Schmidt, U. (2016). A systematic review of the relationship between eating, weight and inhibitory control using the stop signal task. *Neuroscience & Biobehavioral Reviews, 64*, 35–62. <https://doi.org/10.1016/j.neubiorev.2016.02.010>
- Benau, E. M., Orloff, N. C., Janke, E. A., Serpell, L., & Timko, C. A. (2014). A systematic review of the effects of experimental fasting on cognition☆. *Appetite, 77*, 52–61. <https://doi.org/10.1016/j.appet.2014.02.014>
- Burger, J. M., & Cooper, H. M. (1979). The desirability of control. *Motivation and Emotion, 3*(4), 381–393. <https://doi.org/10.1007/BF00994052>
- Burns, G. L., Keortge, S. G., Formea, G. M., & Sternberger, L. G. (1996). Revision of the Padua Inventory of obsessive compulsive disorder symptoms: Distinctions between worry, obsessions, and compulsions. *Behaviour Research and Therapy, 34*(2), 163–173.
- Cappelleri, J. C., Bushmakina, A. G., Gerber, R. A., Leidy, N. K., Sexton, C. C., Karlsson, J., & Lowe, M. R. (2009). Evaluating the Power of Food Scale in obese subjects and a general sample of individuals: Development and measurement properties. *International Journal of Obesity, 33*(8), 913–922. <https://doi.org/10.1038/ijo.2009.107>
- Carver, C. S., & White, T. L. (1994). Behavioral inhibition, behavioral activation, and affective responses to impending reward and punishment: The BIS/BAS scales. *Journal of Personality and Social Psychology, 67*(2), 319.
- Cassidy, R. M., & Tong, Q. (2017). Hunger and Satiety Gauge Reward Sensitivity. *Frontiers in Endocrinology, 8*. <https://doi.org/10.3389/fendo.2017.00104>
- Dulloo, A. G., Jacquet, J., Miles-Chan, J. L., & Schutz, Y. (2017). Passive and active roles of fat-free mass in the control of energy intake and body composition regulation. *European Journal of Clinical Nutrition, 71*(3), 353–357. <https://doi.org/10.1038/ejcn.2016.256>
- Eagle, D. M., & Baunez, C. (2010). Is there an inhibitory-response-control system in the rat? Evidence from anatomical and pharmacological studies of behavioral inhibition. *Neuroscience and Biobehavioral Reviews, 34*(1–3), 50–72. <https://doi.org/10.1016/j.neubiorev.2009.07.003>

- 1 Epstein, L. H., Temple, J. L., Neaderhiser, B. J., Salis, R. J., Erbe, R. W., & Leddy, J.
2 J. (2007). Food reinforcement, the dopamine D2 receptor genotype, and energy intake
3 in obese and nonobese humans. *Behavioral Neuroscience*, 121(5), 877–886.
4 <https://doi.org/10.1037/0735-7044.121.5.877>
- 5 Ettelt, S. (2005). *Impulsivität und zwangsstörung und deren familiäre komorbidität*
6 *und kosegregation*.
- 7 Flavell, S. W., Gogolla, N., Lovett-Barron, M., & Zelikowsky, M. (2022). The
8 emergence and influence of internal states. *Neuron*, 110(16), S089662732200407X.
9 <https://doi.org/10.1016/j.neuron.2022.04.030>
- 10 Francis, H., & Stevenson, R. (2013). Validity and test-retest reliability of a short
11 dietary questionnaire to assess intake of saturated fat and free sugars: A preliminary
12 study. *Journal of Human Nutrition and Dietetics*, 26(3), 234–242.
13 <https://doi.org/10.1111/jhn.12008>
- 14 Fulton, S., Pissios, P., Manchon, R. P., Stiles, L., Frank, L., Pothos, E. N., Maratos-
15 Flier, E., & Flier, J. S. (2006). Leptin Regulation of the Mesoaccumbens Dopamine
16 Pathway. *Neuron*, 51(6), 811–822. <https://doi.org/10.1016/j.neuron.2006.09.006>
- 17 Garcia-Garcia, I., Neseliler, S., Morys, F., Dadar, M., Yau, Y. H. C., Scala, S. G.,
18 Zeighami, Y., Sun, N., Collins, D. L., Vainik, U., & Dagher, A. (2022). Relationship
19 between impulsivity, uncontrolled eating and body mass index: A hierarchical model.
20 *International Journal of Obesity*, 46(1, 1), 129–136. [https://doi.org/10.1038/s41366-](https://doi.org/10.1038/s41366-021-00966-4)
21 [021-00966-4](https://doi.org/10.1038/s41366-021-00966-4)
- 22 Garner, D. M., Olmstead, M. P., & Polivy, J. (1983). Development and validation of a
23 multidimensional eating disorder inventory for anorexia-nervosa and bulimia.
24 *International Journal of Eating Disorders*, 2(2), 15–34. [https://doi.org/10.1002/1098-](https://doi.org/10.1002/1098-108X(198321)2:2<15::AID-EAT2260020203>3.0.CO;2-6)
25 [108X\(198321\)2:2<15::AID-EAT2260020203>3.0.CO;2-6](https://doi.org/10.1002/1098-108X(198321)2:2<15::AID-EAT2260020203>3.0.CO;2-6)
- 26 Geisler, C. E., & Hayes, M. R. (2023). Metabolic hormone action in the VTA:
27 Reward-directed behavior and mechanistic insights. *Physiology & Behavior*, 268,
28 114236. <https://doi.org/10.1016/j.physbeh.2023.114236>
- 29 Gerlach, G., Herpertz, S., & Loeber, S. (2014). Personality traits and obesity: A
30 systematic review. *Obesity Reviews*, 16(1), 32–63. <https://doi.org/10.1111/obr.12235>
- 31 Giesen, J. C. A. H., Havermans, R. C., Douven, A., Tekelenburg, M., & Jansen, A.
32 (2010). Will Work for Snack Food: The Association of BMI and Snack
33 Reinforcement. *Obesity*, 18(5), 966–970. <https://doi.org/10.1038/oby.2010.20>
- 34 Giuffrida, V., Marc, I. B., Ramawat, S., Fontana, R., Fiori, L., Bardella, G., Fagioli,
35 S., Ferraina, S., Brunamonti, E., & Pani, P. (2023). Reward prospect affects strategic
36 adjustments in stop signal task. *Frontiers in Psychology*, 14.
37 <https://doi.org/10.3389/fpsyg.2023.1125066>
- 38 Grunert, S. (1989). Ein inventar zur erfassung von selbstaussagen zum
39 ernährungsverhalten. *Diagnostica*, 35(2), 167–179.
40 <https://psycnet.apa.org/record/1991-73148-001>

- 1 Hanssen, R., Kretschmer, A. C., Rigoux, L., Albus, K., Edwin Thanarajah, S.,
2 Sitnikow, T., Melzer, C., Cornely, O. A., Brüning, J. C., & Tittgemeyer, M. (2021).
3 GLP-1 and hunger modulate incentive motivation depending on insulin sensitivity in
4 humans. *Molecular Metabolism*, 45, 101163.
5 <https://doi.org/10.1016/j.molmet.2021.101163>
- 6 Hanssen, R., Thanarajah, S. E., Tittgemeyer, M., & Brüning, J. C. (2022). Obesity – A
7 Matter of Motivation? *Experimental and Clinical Endocrinology & Diabetes*, 130(5),
8 290–295. <https://doi.org/10.1055/a-1749-4852>
- 9 Hartmann, A. S., Rief, W., & Hilbert, A. (2011). Psychometric properties of the
10 german version of the barratt impulsiveness Scale, version 11 (BIS-11) for
11 adolescents. *Perceptual and Motor Skills*, 112(2), 353–368.
12 <https://doi.org/10.2466/08.09.10.PMS.112.2.353-368>
- 13 Heymsfield, S. B., McManus, C., Smith, J., Stevens, V., & Nixon, D. W. (1982).
14 Anthropometric measurement of muscle mass: Revised equations for calculating
15 bone-free arm muscle area. *The American Journal of Clinical Nutrition*, 36(4), 680–
16 690.
- 17 Holt, C. L., Clark, E. M., & Kreuter, M. W. (2001). Weight locus of control and
18 weight-related attitudes and behaviors in an overweight population. *Addictive*
19 *Behaviors*, 26(3), 329–340. [https://doi.org/10.1016/s0306-4603\(00\)00108-8](https://doi.org/10.1016/s0306-4603(00)00108-8)
- 20 Howard, M., Roiser, J. P., Gilbert, S. J., Burgess, P. W., Dayan, P., & Serpell, L.
21 (2020). Short-Term Fasting Selectively Influences Impulsivity in Healthy Individuals.
22 *Frontiers in Psychology*, 11. <https://doi.org/10.3389/fpsyg.2020.01644>
- 23 Hsu, E. S., Patwardhan, A. G., Meade, K. P., Light, T. R., & Martin, W. R. (1993).
24 Cross-sectional geometrical properties and bone mineral contents of the human radius
25 and ulna. *Journal of Biomechanics*, 26(11), 1307–1318. [https://doi.org/10.1016/0021-9290\(93\)90354-H](https://doi.org/10.1016/0021-9290(93)90354-H)
- 27 Hsu, T. M., McCutcheon, J. E., & Roitman, M. F. (2018). Parallels and Overlap: The
28 Integration of Homeostatic Signals by Mesolimbic Dopamine Neurons. *Frontiers in*
29 *Psychiatry*, 9. <https://doi.org/10.3389/fpsyg.2018.00410>
- 30 Janssen, L. K., & Horstmann, A. (2022). Molecular Imaging of Central Dopamine in
31 Obesity: A Qualitative Review across Substrates and Radiotracers. *Brain Sciences*,
32 12(4), 486. <https://doi.org/10.3390/brainsci12040486>
- 33 Koubi, H. E., Robin, J. P., Dewasmes, G., Le Maho, Y., Frutoso, J., & Minaire, Y.
34 (1991). Fasting-induced rise in locomotor activity in rats coincides with increased
35 protein utilization. *Physiology & Behavior*, 50(2), 337–343.
36 [https://doi.org/10.1016/0031-9384\(91\)90075-Y](https://doi.org/10.1016/0031-9384(91)90075-Y)
- 37 Kroemer, N. B., & Small, D. M. (2016). Fuel not fun: Reinterpreting attenuated brain
38 responses to reward in obesity. *Physiology & Behavior*, 162, 37–45.
39 <https://doi.org/10.1016/j.physbeh.2016.04.020>
- 40 Kühner, C., Bürger, C., Keller, F., & Hautzinger, M. (2007). Reliability and validity
41 of the revised Beck Depression Inventory (BDI-II). Results from German samples.
42 *Der Nervenarzt*, 78(6), 651–656. <https://doi.org/10.1007/s00115-006-2098-7>

- 1 Kullmann, S., Blum, D., Jaghutriz, B. A., Gassenmaier, C., Bender, B., Häring, H.-U.,
2 Reischl, G., Preissl, H., la Fougère, C., Fritsche, A., Reimold, M., & Heni, M. (2021).
3 Central Insulin Modulates Dopamine Signaling in the Human Striatum. *The Journal*
4 *of Clinical Endocrinology & Metabolism*, 106(10), 2949–2961.
5 <https://doi.org/10.1210/clinem/dgab410>
- 6 Lavagnino, L., Arnone, D., Cao, B., Soares, J. C., & Selvaraj, S. (2016). Inhibitory
7 control in obesity and binge eating disorder: A systematic review and meta-analysis of
8 neurocognitive and neuroimaging studies. *Neuroscience & Biobehavioral Reviews*,
9 68, 714–726. <https://doi.org/10.1016/j.neubiorev.2016.06.041>
- 10 Lim, J., Wood, A., & Green, B. G. (2009). Derivation and Evaluation of a Labeled
11 Hedonic Scale. *Chemical Senses*, 34(9), 739–751.
12 <https://doi.org/10.1093/chemse/bjp054>
- 13 Lowe, M. R., Butryn, M. L., Didie, E. R., Annunziato, R. A., Thomas, J. G., Crerand,
14 C. E., Ochner, C. N., Coletta, M. C., Bellace, D., Wallaert, M., & Halford, J. (2009).
15 The Power of Food Scale. A new measure of the psychological influence of the food
16 environment. *Appetite*, 53(1), 114–118. <https://doi.org/10.1016/j.appet.2009.05.016>
- 17 Mansur, R. B., Subramaniapillai, M., Zuckerman, H., Park, C., Iacobucci, M., Lee,
18 Y., Tuineag, M., Hawco, C., Frey, B. N., Rasgon, N., Brietzke, E., & McIntyre, R. S.
19 (2019). Effort-based decision-making is affected by overweight/obesity in major
20 depressive disorder. *Journal of Affective Disorders*, 256, 221–227.
21 <https://doi.org/10.1016/j.jad.2019.06.002>
- 22 Mathar, D., Horstmann, A., Pleger, B., Villringer, A., & Neumann, J. (2016). Is it
23 Worth the Effort? Novel Insights into Obesity-Associated Alterations in Cost-Benefit
24 Decision-Making. *Frontiers in Behavioral Neuroscience*, 9.
25 <https://doi.org/10.3389/fnbeh.2015.00360>
- 26 Matzke, D., Verbruggen, F., & Logan, G. (2018). The stop-signal paradigm. *Stevens’*
27 *Handbook of Experimental Psychology and Cognitive Neuroscience*, 5, 383–427.
- 28 Maughan, R. J., Watson, J. S., & Weir, J. (1983). Strength and cross-sectional area of
29 human skeletal muscle. *The Journal of Physiology*, 338(1), 37–49.
30 <https://doi.org/10.1113/jphysiol.1983.sp014658>
- 31 Mobbs, O., Crépin, C., Thiéry, C., Golay, A., & Van der Linden, M. (2010). Obesity
32 and the four facets of impulsivity. *Patient Education and Counseling*, 79(3), 372–377.
33 <https://doi.org/10.1016/j.pec.2010.03.003>
- 34 Neu, C. M., Rauch, F., Rittweger, J., Manz, F., & Schoenau, E. (2002). Influence of
35 puberty on muscle development at the forearm. *American Journal of Physiology -*
36 *Endocrinology And Metabolism*, 283(1), E103–E107.
37 <https://doi.org/10.1152/ajpendo.00445.2001>
- 38 Opland, D. M., Leininger, G. M., & Myers, M. G. (2010). Modulation of the
39 mesolimbic dopamine system by leptin. *Brain Research*, 1350, 65–70.
40 <https://doi.org/10.1016/j.brainres.2010.04.028>

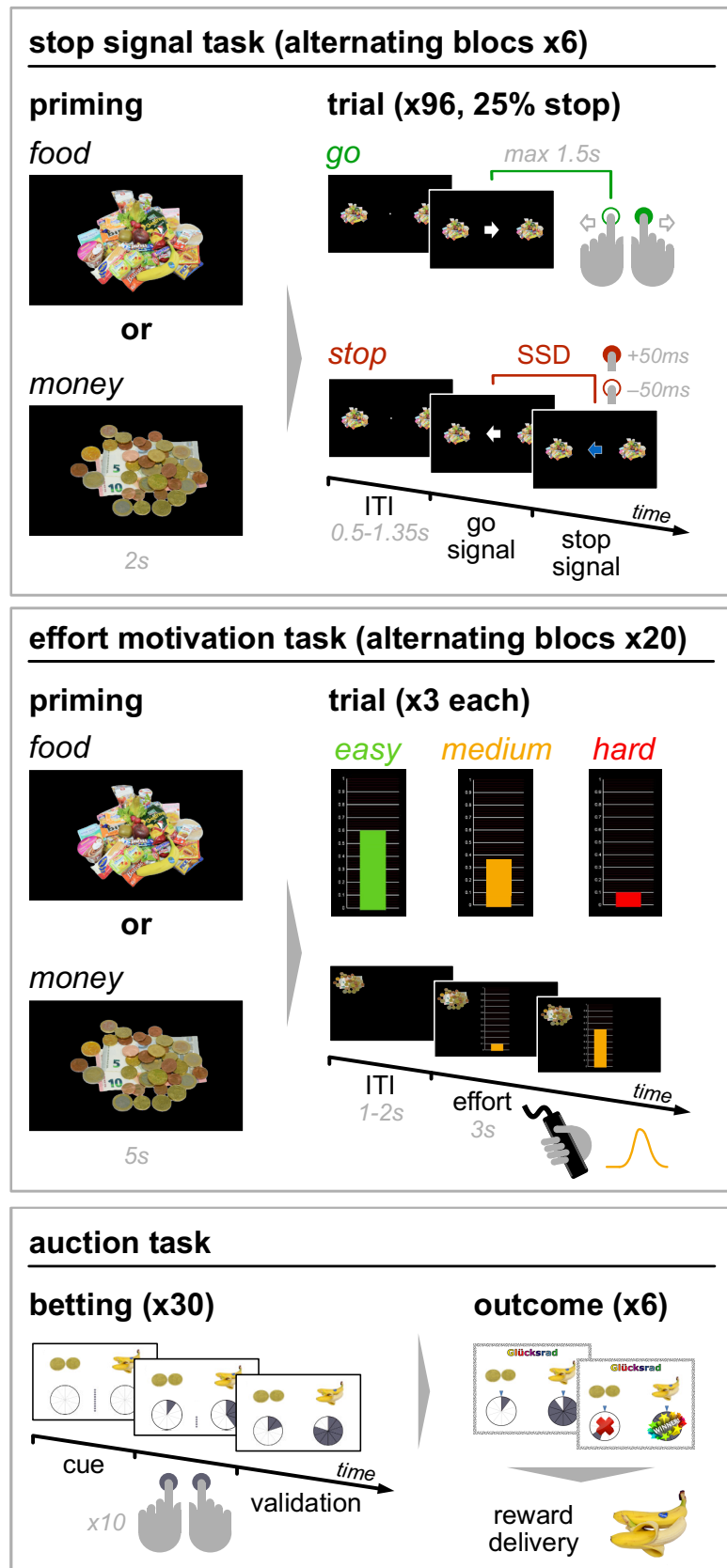
- 1 Oren, S., Tittgemeyer, M., Rigoux, L., Schlamann, M., Schonberg, T., &
2 Kuzmanovic, B. (2022). Neural encoding of food and monetary reward delivery.
3 *NeuroImage*, 257, 119335. <https://doi.org/10.1016/j.neuroimage.2022.119335>
- 4 Palmiter, R. D. (2007). Is dopamine a physiologically relevant mediator of feeding
5 behavior? *Trends in Neurosciences*, 30(8), 375–381.
6 <https://doi.org/10.1016/j.tins.2007.06.004>
- 7 Pessiglione, M., Schmidt, L., Draganski, B., Kalisch, R., Lau, H., Dolan, R. J., &
8 Frith, C. D. (2007). How the brain translates money into force: A neuroimaging study
9 of subliminal motivation. *Science*, 316(5826), 904–906.
10 <https://doi.org/10.1126/science.1140459>
- 11 Pirc, M., Čad, E. M., Jager, G., & Smeets, P. A. M. (2019). Grab to eat! Eating
12 motivation dynamics measured by effort exertion depend on hunger state. *Food*
13 *Quality and Preference*, 78, 103741. <https://doi.org/10.1016/j.foodqual.2019.103741>
- 14 Portugal, A. C. A., Afonso Jr, A. S., Caldas, A. L., Maturana, W., Mocaiber, I., &
15 Machado-Pinheiro, W. (2018). Inhibitory mechanisms involved in Stroop-matching
16 and stop-signal tasks and the role of impulsivity. *Acta Psychologica*, 191, 234–243.
17 <https://doi.org/10.1016/j.actpsy.2018.10.003>
- 18 Probst, C. C., Winter, L. M., Möller, B., Weber, H., Weintraub, D., Witt, K., Deuschl,
19 G., Katzenschlager, R., & van Eimeren, T. (2014). Validation of the questionnaire for
20 impulsive-compulsive disorders in Parkinson’s disease (QUIP) and the QUIP-rating
21 scale in a German speaking sample. *Journal of Neurology*, 261(5), 936–942.
- 22 Pudel, V., & Westenhöfer, J. (1989). *Fragebogen zum essverhalten (FEV):*
23 *handanweisung*. Verlag für Psychologie Hogrefe; Verlag für Psychologie Hogrefe.
24 [https://scholar.google.com/javascript:void\(0\)](https://scholar.google.com/javascript:void(0))
- 25 Robinson, E., Roberts, C., Vainik, U., & Jones, A. (2020). The psychology of obesity:
26 An umbrella review and evidence-based map of the psychological correlates of
27 heavier body weight. *Neuroscience & Biobehavioral Reviews*, 119, 468–480.
28 <https://doi.org/10.1016/j.neubiorev.2020.10.009>
- 29 Saltzer, E. B. (1982). The weight locus of control (WLOC) scale: A specific measure
30 for obesity research. *Journal of Personality Assessment*, 46(6), 620–628.
31 https://doi.org/10.1207/s15327752jpa4606_11
- 32 Satow, L. (2012). *Stress-und coping-inventar (SCI): Test-und skalendokumentation*.
- 33 Siep, N., Roefs, A., Roebroek, A., Havermans, R., Bonte, M. L., & Jansen, A.
34 (2009). Hunger is the best spice: An fMRI study of the effects of attention, hunger
35 and calorie content on food reward processing in the amygdala and orbitofrontal
36 cortex. *Behavioural Brain Research*, 198(1), 149–158.
37 <https://doi.org/10.1016/j.bbr.2008.10.035>
- 38 Skrynka, J., & Vincent, B. T. (2019). Hunger increases delay discounting of food and
39 non-food rewards. *Psychonomic Bulletin & Review*, 26(5), 1729–1737.
40 <https://doi.org/10.3758/s13423-019-01655-0>

- 1 Stanford, M. S., Mathias, C. W., Dougherty, D. M., Lake, S. L., Anderson, N. E., &
2 Patton, J. H. (2009). Fifty years of the Barratt Impulsiveness Scale: An update and
3 review. *Personality and Individual Differences*, 47(5), 385–395.
4 <https://doi.org/10.1016/j.paid.2009.04.008>
- 5 Strobel, A., Beauducel, A., Debener, S., & Brocke, B. (2003). Psychometrische
6 eigenschaften und normen einer deutschsprachigen fassung der sensation seeking-
7 skalen, form V. *Diagnostica*.
- 8 Stunkard, A. J., & Messick, S. (1985). The three-factor eating questionnaire to
9 measure dietary restraint, disinhibition and hunger. *Journal of Psychosomatic*
10 *Research*, 29(1), 71–83. [https://doi.org/10.1016/0022-3999\(85\)90010-8](https://doi.org/10.1016/0022-3999(85)90010-8)
- 11 Thiel, A., & Paul, T. (2006). Test-retest reliability of the eating disorder inventory 2.
12 *Journal of Psychosomatic Research*, 61(4), 567–569.
13 <https://doi.org/10.1016/j.jpsychores.2006.02.015>
- 14 Thunberg, C., Wiker, T., Bundt, C., & Huster, R. J. (2024). On the (un)reliability of
15 common behavioral and electrophysiological measures from the stop signal task:
16 Measures of inhibition lack stability over time. *Cortex*, 175, 81–105.
17 <https://doi.org/10.1016/j.cortex.2024.02.008>
- 18 Torrubia, R., Ávila, C., Moltó, J., & Caseras, X. (2001). The sensitivity to punishment
19 and sensitivity to reward questionnaire (SPSRQ) as a measure of gray's anxiety and
20 impulsivity dimensions. *Personality and Individual Differences*, 31(6), 837–862.
21 [https://doi.org/10.1016/S0191-8869\(00\)00183-5](https://doi.org/10.1016/S0191-8869(00)00183-5)
- 22 Vainik, U., Dagher, A., Realo, A., Colodro-Conde, L., Mortensen, E. L., Jang, K.,
23 Juko, A., Kandler, C., Sørensen, T. I. A., & Möttus, R. (2019). Personality-obesity
24 associations are driven by narrow traits: A meta-analysis. *Obesity Reviews*, 20(8),
25 1121–1131. <https://doi.org/10.1111/obr.12856>
- 26 Vainik, U., Neseliler, S., Konstabel, K., Fellows, L. K., & Dagher, A. (2015). Eating
27 traits questionnaires as a continuum of a single concept. *Uncontrolled eating*.
28 *Appetite*, 90, 229–239. <https://doi.org/10.1016/j.appet.2015.03.004>
- 29 van Swieten, M. M. H., Bogacz, R., & Manohar, S. G. (2023). Gambling on an empty
30 stomach: Hunger modulates preferences for learned but not described risks. *Brain and*
31 *Behavior*, 13(5), e2978. <https://doi.org/10.1002/brb3.2978>
- 32 Vanstrien, T., Frijters, J., Bergers, G., & Defares, P. B. (1986). The dutch eating
33 behavior questionnaire (DEBQ) for assessment of restrained, emotional, and external
34 eating behavior. *International Journal of Eating Disorders*, 5(2), 295–315.
35 [https://doi.org/10.1002/1098-108X\(198602\)5:2<295::AID-](https://doi.org/10.1002/1098-108X(198602)5:2<295::AID-EAT2260050209>3.0.CO;2-T)
36 [EAT2260050209>3.0.CO;2-T](https://doi.org/10.1002/1098-108X(198602)5:2<295::AID-EAT2260050209>3.0.CO;2-T)
- 37 Verbruggen, F., Logan, G. D., & Stevens, M. A. (2008). STOP-IT: Windows
38 executable software for the stop-signal paradigm. *Behavior Research Methods*, 40(2),
39 479–483. <https://doi.org/10.3758/BRM.40.2.479>
- 40 Voigt, K., Giddens, E., Stark, R., Frisch, E., Moskovsky, N., Kakoschke, N., Stout, J.
41 C., Bellgrove, M. A., Andrews, Z. B., & Verdejo-Garcia, A. (2021). The Hunger
42 Games: Homeostatic State-Dependent Fluctuations in Disinhibition Measured with a

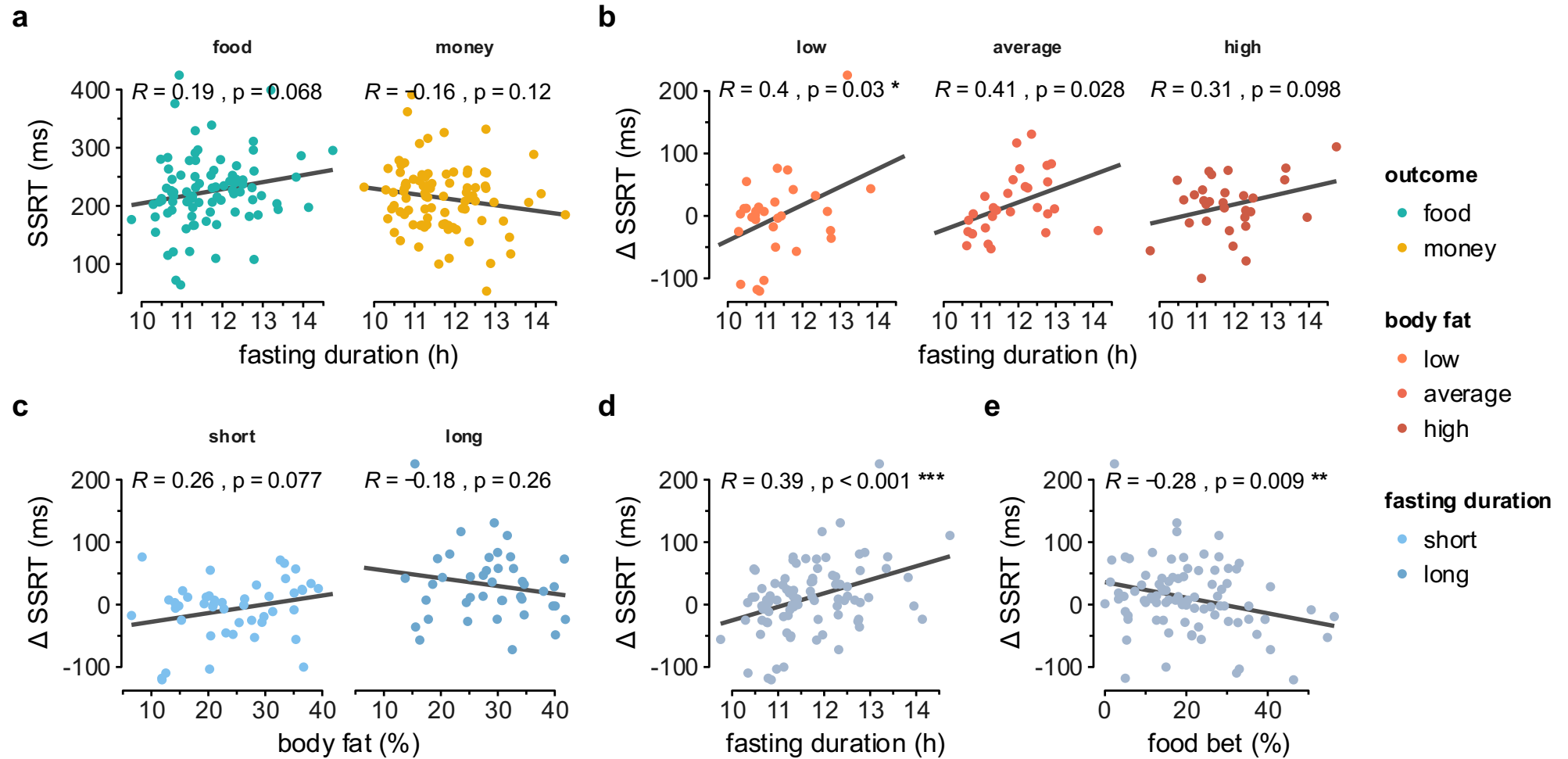
- 1 Novel Gamified Test Battery. *Nutrients*, 13(6), 2001.
2 <https://doi.org/10.3390/nu13062001>
- 3 Weintraub, D., Hoops, S., Shea, J. A., Lyons, K. E., Pahwa, R., Driver-Dunckley, E.
4 D., Adler, C. H., Potenza, M. N., Miyasaki, J., Siderowf, A. D., et al. (2009).
5 Validation of the questionnaire for impulsive-compulsive disorders in Parkinson's
6 disease. *Movement Disorders: Official Journal of the Movement Disorder Society*,
7 24(10), 1461–1467.
- 8 Winstanley, C. A. (2011). The utility of rat models of impulsivity in developing
9 pharmacotherapies for impulse control disorders. *British Journal of Pharmacology*,
10 164(4), 1301–1321. <https://doi.org/10.1111/j.1476-5381.2011.01323.x>
- 11 Zhang, X., Chen, S., Chen, H., Gu, Y., & Xu, W. (2017). General and Food-Specific
12 Inhibitory Control As Moderators of the Effects of the Impulsive Systems on Food
13 Choices. *Frontiers in Psychology*, 8. <https://doi.org/10.3389/fpsyg.2017.00802>
- 14 Ziauddeen, H., Subramaniam, N., Gaillard, R., Burke, L. K., Farooqi, I. S., &
15 Fletcher, P. C. (2012). Food images engage subliminal motivation to seek food.
16 *International Journal of Obesity*, 36(9), 1245–1247.
17 <https://doi.org/10.1038/ijo.2011.239>
- 18 Zuckerman, M., Kolin, E. A., Price, L., & Zoob, I. (1964). Development of a
19 sensation-seeking scale. *Journal of Consulting Psychology*, 28(6), 477.

1 Figures

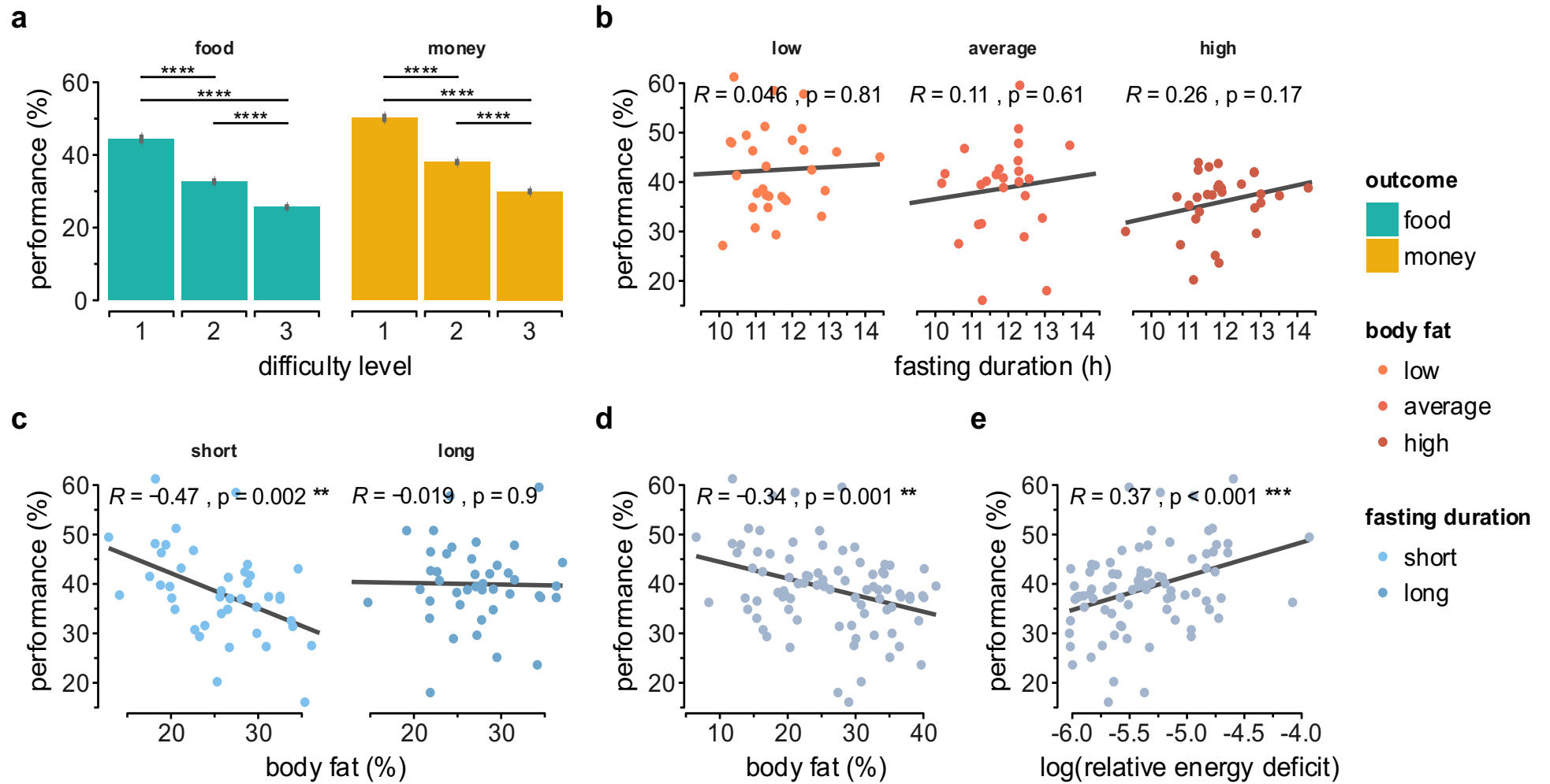
2 Figure 1



1 *Figure 2*

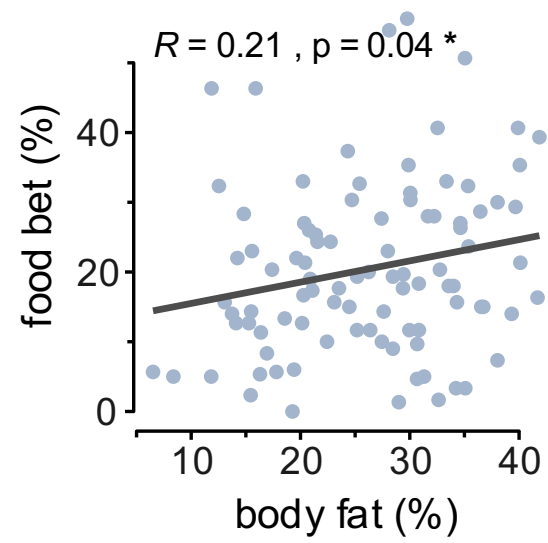


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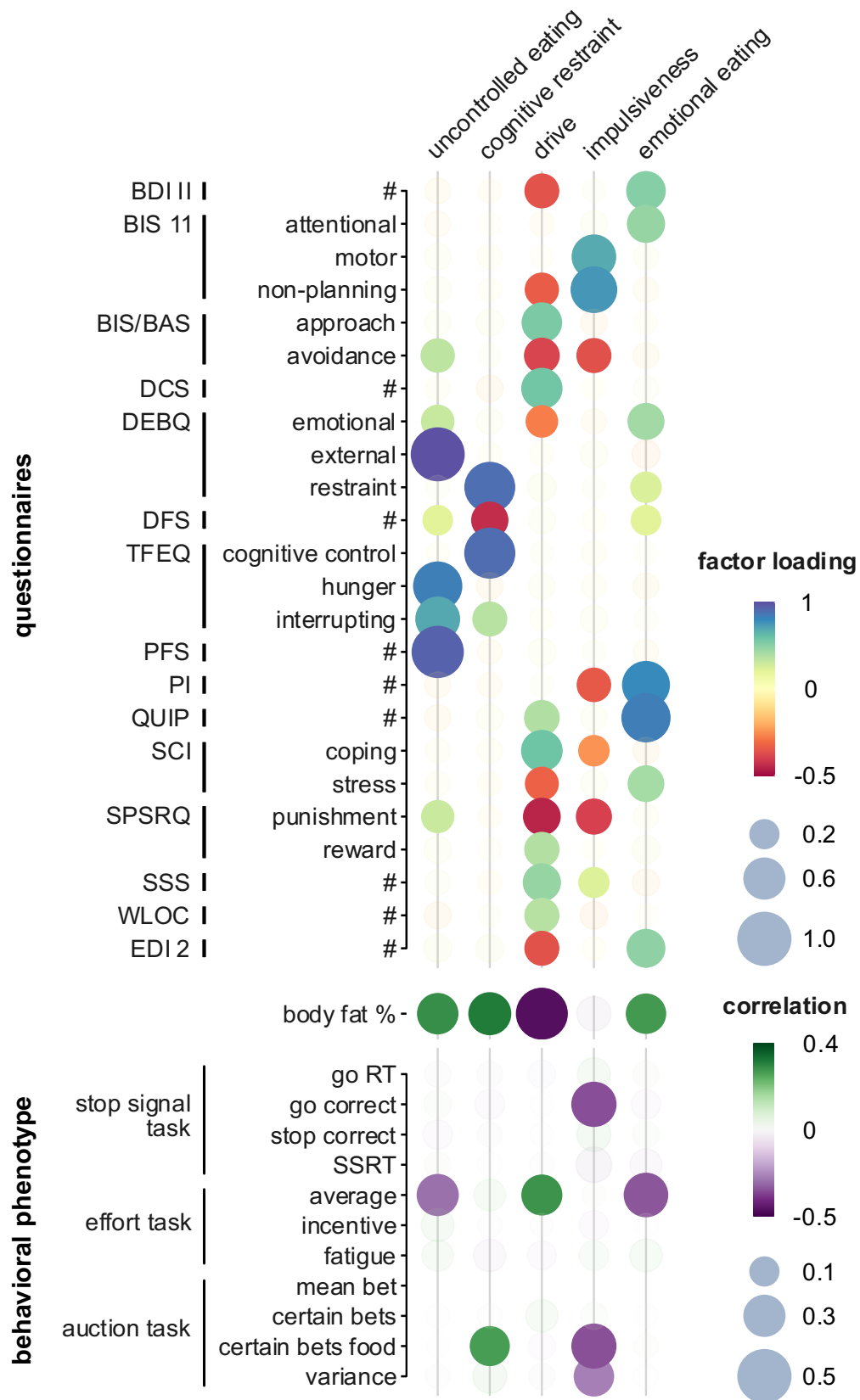


1 *Figure 4*

2



1 *Figure 5*



1 **Figure legends**

2 **Figure 1: Experimental design** *Top: Stop Signal task* Each bloc of 96 trials started
3 with a full screen picture of the type of reward at stake (food or money). On each trial,
4 a fixation cross first appeared in the centre of the screen, flanked by reward cues.
5 Then an white arrow was displayed, prompting the participants to indicate the
6 orientation of the arrow with button presses (“go” condition). In 1/4 of the trials, the
7 arrow turned blue after a variable stop signal delay (SSD). In this “stop” condition,
8 participants were instructed to refrain from responding. The SSD was continuously
9 adjusted to induce a 50% chance of correct response inhibition. *Middle: Effort*
10 *Motivation task* Again, each bloc started with a full screen display of the reward at
11 stake. On each trial, participant could press a hand held dynamometer to raise the
12 level of a gauge on the screen and thus increase their chances of earning the reward.
13 The color of the gauge indicated how hard they had to press to fill the gauge
14 completely (difficulty level). *Bottom: Auction task* On each trial, participant had to
15 split their bet by placing total of 10 tokens on either a monetary reward or a food item
16 of equal value (more tokens = higher chances of winning, 10 token max per option).
17 The outcome was displayed once all the bet were set, and both the monetary and food
18 (snacks) rewards were given to participant to consume.

19 **Figure 2: Impulsive behaviour as quantified by the SSRT measured in the stop**
20 **signal task.** *a)* SSRT as a function of fasting duration for food (green) and monetary
21 (yellow) blocs. *b-e)* Relative impulsivity for food relative to money measured by the
22 difference in SSRT between the two conditions (Δ SSRT). *b)* Δ SSRT increases with
23 fasting duration especially for the lowest (left, light orange), and central (middle, dark
24 orange), compared to the highest (right, red) tercile of body fat percentage. *c)* Δ SSRT
25 increases with body fat percentage for short (left, light blue) but not for long (right,
26 dark blue) fasting duration (duration split at the median for plotting only). *d)* Δ SSRT
27 increases with fasting duration (same as in b, collapsed across body composition). *e)*
28 Δ SSRT decreases with larger bets toward food items (as opposed to monetary item) in
29 the auction task. All statistics are Pearson correlations and lines best fit linear
30 regression computed for the data showed in each plot.

31 **Figure 3: Results of the incentive motivation task.** *a)* Average performance (gauge
32 level) for each difficulty level for food (green) and monetary (yellow) rewards. Bar

1 and error bars represent the group average and standard errors. *b)* Effort increases
2 with fasting duration, especially as body fat increased (here split in tercile from left to
3 right, darker colours means higher body fat). However, higher body fat was also
4 associated with lower effort on average.

5 *c)* This effect of body composition is driven by participant with shorter fasting
6 duration (left, light blue) and normalises with longer fasting (right, dark blue). *d)*
7 Overall, effort decreased with body fat %. *e)* The body composition x fasting
8 interaction is summarized as an effect of relative energy deficit induced by fasting.
9 All statistics are Pearson correlations and lines best fit linear regression computed for
10 the data showed in each plot.

11 **Figure 4: Results of the auction task.** Willingness to pay for food increases with
12 body fat percentage.

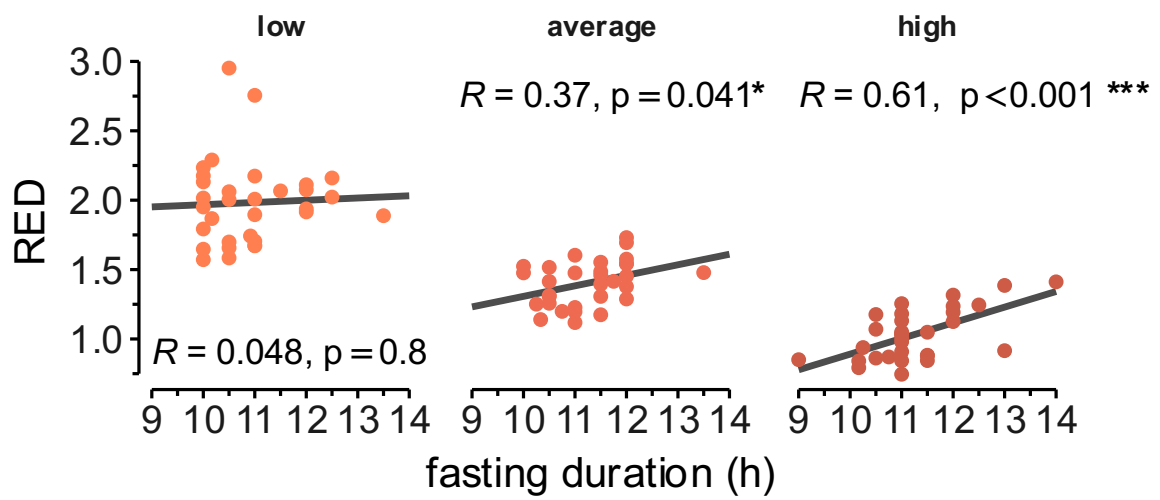
13 **Figure 5: Factor analysis of the questionnaires.** *Top:* Each dot represent the loading
14 of each questionnaire scale (lines) on the five identified factors (column). *Bottom:*
15 Each dot represent the correlation between empirical measures of body composition
16 or behavioural metrics with individual factor scores.

1 Tables

2 Table 1: Questionnaires

Label	Questionnaire	Type	Reference
BIS-11	Barrat Impulsiveness Scale	online	(Hartmann et al., 2011; Stanford et al., 2009)
BDI-2	Beck Depression Inventory	online	(Arnau et al., 2001; Kühner et al., 2007)
BIS/BAS	Behavioral Inhibition and Activation Systems Scales	paper	(Carver & White, 1994; Strobel et al., 2003)
DCS	Desirability of Control Scale	paper	(Burger & Cooper, 1979)
DEBQ	Dutch Eating Behavior Questionnaire	online	(Grunert, 1989; Vanstrien et al., 1986)
DFS	Dietary Fat and free Sugar	online	(Francis & Stevenson, 2013)
EDI2	Eating Disorder Inventroy 2	online	(Garner et al., 1983; Thiel & Paul, 2006)
PFS	Power of Food Scale	online	(Cappelleri et al., 2009; Lowe et al., 2009)
PI-WSUR	Padua Inventory - Washington State University Revision	paper	(Burns et al., 1996; Ettelt, 2005)
QUIP-RS	Questionnaire for Impulsive-Compulsive Disorders - Rating Scale	paper	(Probst et al., 2014; Weintraub et al., 2009)
SCI	Stress and Coping Inventory	online	(Satow, 2012)
SPSRQ	Sensitivity to Punishment and Sensitivity to Reward Questionnaire	online	(Torrubia et al., 2001)
SSS-V	Sensation Seeking Scale	paper	(Strobel et al., 2003; Zuckerman et al., 1964)
TFEQ	Three Factors Eating Questionnaire	online	(Pudel & Westenhöfer, 1989; Stunkard & Messick, 1985)
WLOC	Weight of Locus of Control	online	(Holt et al., 2001; Saltzer, 1982)

1 Supplementary Material



2

3 *Figure S1: **Relative energy deficit*** Relative energy deficit as a function of fasting
4 duration for the low (left), average (middle), and high (right) body fat participants.