Higher Heart-Rate Variability Is Associated with Ventromedial Prefrontal Cortex Activity and Increased Resistance to Temptation in Dietary Self-Control Challenges

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Higher levels of self-control in decision making have been linked to better psychosocial and physical health. A similar link to health outcomes has been reported for heart-rate variability (HRV), a marker of physiological flexibility. Here, we sought to link these two, largely separate, research domains by testing the hypothesis that greater HRV would be associated with better dietary self-control in humans. Specifically, we examined whether total HRV at sedentary rest (measured as the SD of normal-to-normal intervals) can serve as a biomarker for the neurophysiological adaptability that putatively underlies self-controlled behavior. We found that HRV explained a significant portion of the individual variability in dietary self-control, with individuals having higher HRV being better able to down-regulate their cravings in the face of taste temptations. Furthermore, HRV was associated with activity patterns in the ventromedial prefrontal cortex (vmPFC), a key node in the brain’s valuation and decision circuitry. Specifically, individuals with higher HRV showed both higher overall vmPFC blood-oxygen-level-dependent activity and attenuated taste representations when presented with a dietary self-control challenge. Last, the behavioral and neural associations with HRV were consistent across both our stress induction and control experimental conditions. The stability of this association across experimental conditions suggests that HRV may serve as both a readily obtainable and robust biomarker for self-control ability across environmental contexts.

Key words: decision making; fMRI; HRV; self-regulation

Significance Statement
Self-control is associated with better health, but behavioral and psychometric self-control measures allow only indirect associations with health outcomes and may be distorted by reporting bias. We tested whether resting heart-rate variability (HRV), a physiological indicator of psychological and physical health, can predict individual differences in dietary self-control in humans. We found that higher HRV was associated with better self-control and improved predictions of choice behavior. Specifically, higher HRV was associated with more effective downregulation of taste temptations, and with a diminished neural representation of taste temptations during self-control challenges. Our results suggest that HRV may serve as an easily acquired, noninvasive, and low-cost biomarker for self-control ability.

Introduction
Self-regulation has been associated with a wide range of life outcomes, from educational achievement and socioeconomic status to mental and physical health (Mischel et al., 1989; Duckworth, 2011; Moffitt et al., 2011). Therefore, accurate predictors of individuals’ self-regulatory abilities are important tools in both basic scientific research as well as applied domains, including education and medicine. Self-regulation is generally assessed in specific domains by psychometric questionnaires or laboratory tasks. Unfortunately, participants can potentially distort these measurements by reporting socially desirable answers or behaving according to the presumed goals of the experimenter. Therefore, measures based on physiological readouts that are easy to obtain, domain independent, and robust to reporting biases could be important tools in the assessment of self-control.

One such readout is heart-rate variability (HRV). Measures of HRV have been linked to self-regulatory capacities and performance in the domain of emotion (Thayer and Lane, 2000, 2009), raising the question of whether they might serve as more general
predictors of self-control. HRV is a well established physiological characteristic of all vertebrates (Grossman and Taylor, 2007): the timing between subsequent heartbeats oscillates on the order of milliseconds and no two neighboring beat pairs (RR intervals) are of exactly the same length (Camm et al., 1996). An animal’s HRV is sensitive to both physical and mental strain (Porges and Raskin, 1969), and differences in resting HRV can distinguish between states of health and disease (Henri et al., 2014, 2015). High resting HRV has been associated with good physical (Masi et al., 2007; Brändle et al., 2015) and mental health (Thayer and Brosschot, 2005), while chronic decreases in HRV indicated disease states and slow recovery from stress (Weber et al., 2010; Stalder et al., 2011).

The polyvagal (Porges, 1995, 2001) and neurovisceral integration (Thayer and Lane, 2000, 2009) theories postulate a mechanistic link between HRV and self-regulation. Both associate CNS regulation of the cardiovascular system, which is necessary to prepare reactions to challenges in the environment, with adaptive behavior at a higher cognitive level. However, we note that it remains unclear to what degree the central versus peripheral nervous system influences HRV. Although we do not yet fully understand all of the physiological and cognitive factors driving HRV (Heathers, 2014), we build on previous proposals (Grossman and Taylor, 2007) and posit that HRV serves as a readout of an individual’s allostatic capacities to integrate behavioral strategies and energy stores in response to demands in the environment.

Higher HRV has been linked to several cognitive processes that support self-regulation, including the following: (1) reallocation of attention (e.g., disengaging from stimuli that are not threatening in the current context), which may reduce allostatic load (McEwen and Wingfield, 2003); (2) persistence (Reynard et al., 2011); and (3) working memory (Gianaros et al., 2004; Hansen et al., 2004). In contrast, low HRV has been associated with disinhibition and dysregulated social conduct (Beauchaine, 2001, 2007). Furthermore, Daly et al. (2014) reported that higher trait self-control, measured using the self-report scale developed by Tangney et al. (2004), correlates with higher resting HRV. However, the links between HRV and self-control at the neural level remain unknown.

In this study, we used fMRI to investigate the relationship between dietary self-control and resting HRV. We hypothesized that better self-control should be associated with higher HRV, and that individual differences in HRV would be associated with neural processing within a self-control network, including the dorsolateral prefrontal cortex (dLPFC) and the ventromedial prefrontal cortex (vmPFC; Hare et al., 2009; Maier et al., 2015). We indeed found that higher resting HRV was associated with better dietary self-control. Furthermore, we found that HRV positively correlated with activity in the vmPFC when individuals faced self-control challenges, and that high-HRV individuals showed a decreased sensitivity to taste attributes in the vmPFC. These vmPFC findings suggest a neural mechanism for the down-regulation of tempting taste attributes that may facilitate dietary self-control.

Materials and Methods

Participants

Fifty-one men participated in this study. The sample is the same as in the study by Maier et al. (2015), where we reported the effects of stress on behavioral and neural self-control processes, but no heart-rate analyses. We included only male participants to facilitate the collection and analysis of cortisol responses to stress in our previous work. Baseline heart-rate data for two participants were lost due to recording failure. In the present report, we include the subset of participants for whom we have both heart-rate and fMRI data (22 control and 27 stress group participants). The Ethics Committee of the Canton of Zurich approved this study and all participants provided written informed consent on the study day. All participants were right-handed and had normal or corrected to normal vision. None of them reported any history of somatic or psychiatric disease, nor did they take any prescription medication. On average, participants in the sample had a blood pressure in the (high) normal range for their age group (mean + SD: systolic blood pressure, 130 ± 14 mmHg; diastolic blood pressure, 77 ± 9 mmHg).

Participants were excluded during the recruitment stage if they suffered from any allergies, food intolerances, or eating disorders. We also excluded individuals who followed a specific diet (e.g., eating vegetarian, vegan, gluten-free/lactose-free, etc.), or who did not report enjoying and regularly consuming snack foods (regularly was defined as ≥ 2 occasions per week). A final eligibility criterion was that participants had to report that they were trying to maintain a healthy lifestyle, including exercise and an overall balanced diet. Together, these criteria ensured that participants would face a meaningful self-control challenge in the dietary-choice task.

To ensure a homogeneous reaction of the hypothalamic–pituitary–adrenal axis in response to stress induction, participants were asked to abstain from drinking alcoholic or caffeinated beverages in the 18 h before the study, to not exercise in the 6 h before the study, and to come to the laboratory well rested. We only recruited nonsmokers who had no history of drug abuse. We asked participants to go to bed by midnight at the latest on the day before the study and to get a good night’s sleep. We instructed participants to not take any medication that alters the blood flow (e.g., analgesics) in the 72 h before their appointment. To motivate the dietary choices, participants were instructed to eat a small meal (sandwich or salad with ~450 kcal) 3 h before the study and consume nothing but water after that.

Allen et al. (2007) identified age, exercise habits, and obesity as potential confounding factors for heart-rate analyses. Our sample was relatively homogeneous with regard to these factors. The men were on average 21.2 years old (SD, ± 2 years), had a normal body mass index (BMI; mean + SD: 22.7 ± 2.1), trained on average 1.6 (SD, ± 1.4) times per week for building strength, and had completed an average of 1.9 (SD, ± 1.3) cardio training sessions per week during the past 4 weeks before the study, resulting in a combined mean of 3.6 (SD, ± 2.1) weekly training sessions per participant. Other factors identified by Allen and colleagues, including smoking, gender, caffeine and alcohol intake, and circadian rhythm, were controlled for by our study exclusion criteria and design.

Procedure

In the 30–40 min preceding the resting HRV measurement, participants had rated 180 food items for health and taste to create tempting dietary choice pairs. After the heartbeat-interval measurement, a stress induction (Socially Evaluated Cold Pressor Test (SECPPT)) or control procedure was administered. Assignment to the stress induction or control conditions was unknown to both the participant and the experimenter at the time of HRV measurement. However, participants knew they would be randomly assigned to one treatment or the other based on the information provided with the consent forms at the beginning of the study. The SECPPT treatment elicited an acute stress response, as indicated by higher cortical values in the Stress group (mean ± SEM cortisol at max-stress: Mean, 9.64 ± 1.09 nM/l; Control, 6.6 ± 0.67 nM/l), and higher reports of perceiving to be stressed than the Control group (on a visual analog scale from 0 to 100 with “not at all stressed” and 100 “extremely stressed”: mean ± SEM, Stress, 33 ± 4%; Control, 19 ± 5%). Details of the stress induction and behavioral task were reported by Maier et al. (2015). Here, we focus on the relationship between the baseline HRV parameter and dietary self-control success (SCS) and its neural correlates. However, given that the stress treatment is known to change dietary choices (Maier et al., 2015), we included it as a factor in all regression models.

Immediately after the stress induction, participants were scanned with blood-oxygen-level-dependent (BOLD) fMRI while they made choices...
in a dietary self-control task. The screen always depicted two food items, and participants had to choose whether they wanted to eat the item on the right or on the left (Fig. 1a). The binary food choices fell into one of two categories. In the first, choosing the healthier item was trivial because the healthier item was also tastier. We refer to this category as no-challenge trials because there is no self-control challenge. In the second choice category, self-control challenges, the healthier item was the less tasty of the two foods and thus presented a conflict between taste and health attributes. Self-control challenges were presented on approximately half of the 210 trials for each participant. To examine the variability of the challenge that different participants faced, we normalized the ratings to fall between 0 and 100 points, and calculated absolute taste and health differences between every pair of food items each participant faced. On a scale from 0 to 100, the mean health difference was 34 ± 25 points (mean ± SD) and the mean taste difference was 29 ± 21 points.

The behavioral and fMRI analyses in this paper focus on the self-control challenge cases. Before the task started, participants were reminded to choose the healthier item as often as they could, consistent with their healthy lifestyle goals. Participants knew that one of their choices would be realized in the end, and they would have to eat whatever they chose on the trial that was randomly drawn for being paid out.

Psychometric inventories
German versions of the Spielberger State-Trait Anxiety Inventory (Laux et al., 1981), Three Factor Eating Questionnaire (Pudel and Westenhöfer, 1989), and Behavioral Inhibition and Activation Scales (Strobel et al., 2001) were administered at the end of the study. Data for the trait anxiety scale of the State-Trait Anxiety Inventory are missing for one participant, as he failed to complete the second page of the questionnaire.

Statistical analyses
All behavioral data were analyzed using either the Matlab (Release 2014b, version 8.4.0.150421, MathWorks, RRID:SCR_001905) or R (Version 3.2.1; R Core Team, RRID:SCR_001905) statistical software packages. The fMRI results were depicted using the MRIcron software package (http://www.slicer.org/mricron/micro-micronet, RRID:SCR_002403). All correlations reported in this paper were assessed with a non-parametric bootstrap method. Two-tailed $p$ values for correlations were obtained by testing the Pearson’s correlation coefficients ($r$) against a null distribution generated from 5000 permutations of the data. The 95% confidence intervals (CIs) for the correlations were computed from 5000 bootstrapped samples of the data. The multiple regression model in Equation 2 below was fit using the “lm” function in R. To visualize the HRV by taste and health difference interactions, we plot the estimated SCS levels at specific combinations of HRV and taste or health differences.

Heart-rate data acquisition
We measured baseline heart rate at rest with the Polar RS 800 CX system [for a cross-validation of this method with echocardiogram (ECG), see Quintana et al., 2012]. All measurements were collected between 1:30 P.M. and 5:00 P.M. to control for circadian rhythms (Heathers, 2014). The baseline measurement was always taken in a single session before any stress or control treatment was administered. Participants were seated in a quiet room and instructed that upon mounting the Polar watch and pressing start, they would need to sit upright and remain quiet and calm during the subsequent baseline-recording interval. A baseline recording was taken for 6 min. The first 3 min of the recording were discarded from the analysis to yield a set of data that were less affected by such factors as initial motion while acclimatizing to the recording environment (Quintana et al., 2016). We focused on baseline (i.e., resting) heart-rate measures to obtain a domain-general index of HRV.

Heart-rate data analyses
HRV can be calculated in two different domains: time and frequency. The full range of measures is discussed in the guidelines by the Task Force on HRV (Campp et al., 1996). Time-domain measures have the advantage of being more robust than frequency measures. Two different time-domain measures are commonly used and both characterize the distribution of interbeat intervals, which are defined as the time between two subsequent heart beats [i.e., the difference between two R peaks in the ECG (Guyton and Hall, 2006), hence also called the “RR interval”]; Fig. 1b]. The SD of all RR (also “NN” for “normal-to-normal”) intervals (SDNN) describes the total HRV within a given period (Eq. 1). The root mean square of successive differences (RMSSD) calculated between adjacent RR intervals is more sensitive to influences of short-term regulation of the heartbeat. Here we focus on HRV at rest (i.e., in the absence of specific, discrete input stimuli), and thus take SDNN as our primary measure of variability.

We chose total HRV (measured as SDNN) as our biomarker for two reasons. First, SDNN is deemed to be the most robust measure of HRV. Among all commonly computed HRV measures, it has been reported to be least compromised by different data preprocessing pipelines, especially the application of artifact correction (Salo et al., 2001). Second, the process of dietary choice is a complex behavioral outcome that may not only depend on a capacity for effective cognitive regulation that helps to achieve self-control goals, but may also be influenced by peripheral factors (e.g., endocrine status, metabolism, and energy expenditure) that are indicative of the current state of the organism. SDNN reflects all influences on the RR interval series, while it is known to correlate highly, although not perfectly, with measures that putatively reflect phasic vagal control of cardiac variability in measures taken under sedentary resting conditions (Allen et al., 2007).

The complete recording of RR intervals for each participant was extracted using the Polar software, without any transformations of the data. Three-minute intervals of the raw data were then preprocessed with the Artifact toolbox (Version 2.08, 64-bit; Kaufmann et al., 2011), which has a better artifact detection rate and shows fewer false detections than the commonly used Kubios HRV toolbox. To identify artifacts, the Artifact toolbox implements the algorithm of Berntson and Stowell (1998), which aims to exclude any potential artifacts before computing the criterion for identifying true artifacts. Based on the report of Salo and colleagues (2001), who compared editing procedures for correcting single RR artifacts, the identified artifacts were deleted from the RR sequence to obtain the cleanest estimate for SDNN. On average, we corrected 2.1% (SD, ±3.1%) of the RR intervals in each sample. Apart from two datasets that had a high number of artifacts requiring correction (12.6 and 10.5% RR intervals removed), all other datasets had between 0 and 6% artifacts corrected (21 datasets were diagnosed as free of artifacts). As a high number of corrected artifacts might be a concern for interpreting our findings, we checked all models for robustness with regard to the number of corrected artifacts.

SDNN was calculated as follows (Eq. 1):

$$SDNN = \sqrt{1 \over N-1} \sum_{j=1}^{N} (R_j - \bar{R})^2$$

where $R_j$ represents each RR interval, $N$ is the number of RR intervals, and $\bar{R}$ is the average RR interval.
HRV was calculated with the Artiifact software suite, using fast Fourier transforms (Bernston and Stowell, 1998; Kaufmann et al., 2011) with an interpolation rate of 4 Hz (spline interpolation) and a Hanning window width that matched the total length of the edited recording (180 s or slightly less in case of deletion correction). Frequency bands were bounded between 0.003 and 0.04 Hz for the very low frequency band, 0.04 and 0.15 Hz for the low-frequency band, and 0.15 and 0.4 Hz for the high-frequency band.

**fMRI data acquisition**

Images were acquired using a Philips Achieva 3 T whole-body scanner with an eight-channel sensitivity-encoding head coil (Philips Medical Systems) at the Laboratory for Social and Neural Systems Research, University Hospital Zurich. Stimulus presentation was controlled with the Psychophysics Toolbox Software [Psychtoolbox 3.0 (Brainard, 1997), RRID:SCR_002881]; the paradigm was presented via a back-projection system to a mirror mounted on the head coil.

We acquired gradient echo T2*-weighted echo-planar images (EPIs) with BOLD contrast (41 slices per volume; field of view, 200 × 200 mm; slice thickness, 2.5 mm; 0.6 mm gap; in-plane resolution, 2.5 × 2.5 mm; matrix, 80 × 80; repetition time, 2460 ms; echo time, 30 ms; flip angle, 77°) and a SENSE (sensitivity-encoding for fast MRI) reduction factor of 2. Volumes were acquired in axial orientation at a +15° tilt to the anterior commissure–posterior commissure line. We collected 161 volumes in ascending order during each of the three experimental runs, together with five “dummy” volumes at the start and end of each run. A T1-weighted turbo field echo structural image was acquired in sagittal orientation for each participant at the end of the scanning session with the same angulation that applied to the functional scans (181 slices; field of view, 256 × 256 × 181 mm; slice thickness, 1 mm; no gap; in-plane resolution, 1 × 1 mm; matrix, 256 × 256; repetition time, 8.4 ms; echo time, 3.89 ms; flip angle, 8°). To measure the homogeneity of the magnetic field, we collected B0/B1 maps before the first and second run and before acquiring the structural scan (short echo time, 4.29 ms; long echo time, 7.4 ms). We measured breathing frequency and took an ECG with an eight-channel sensitivity-encoding head coil (Philips Medical Systems) at the Laboratory for Social and Neural Systems Research, University Hospital Zurich. Stimulus presentation was controlled with the Psychophysics Toolbox Software [Psychtoolbox 3.0 (Brainard, 1997)], RRID:SCR_002881; the paradigm was presented via a back-projection system to a mirror mounted on the head coil.

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**fMRI preprocessing**

Functional data were spatially realigned and unwarped with statistical parametric mapping software (SPM8, Update Rev. Nr. 5236; Functional Imaging Laboratory, University College London, RRID:SCR_007037), segmented according to the participant’s T1-weighted high-resolution structural image and normalized to the individual mean EPI template before smoothing with an isotropic Gaussian kernel (4 mm full width at half maximum). As a last step in preprocessing, we used RETROICOR, as implemented in the PhysIO toolbox, to model respiration and heartbeat (Glover et al., 2000) to account for fluctuations in the BOLD signal due to physiological noise. The PhysIO Toolbox by Kasper et al. (2009) is distributed as open source code as part of the TAPAS (Translational Algorithms for Psychiatry-Advancing Science) software collection (www.translationalneuromodeling.org/tapas/). Following Harvey et al. (2008), the PhysIO algorithm uses Fourier expansions of different order to estimate the phases of cardiac pulsation (third order), respiration (fourth order), and cardiorespiratory interactions (first order). For two participants, the scanner could not save physiological data due to a technical problem. For these participants, only the standard motion-correction procedure was applied.

**fMRI analyses**

General linear models. In all fMRI analyses, regressors in the models were defined as boxcar functions with durations equal to the reaction time on each trial. All three fMRI models also included regressors for head-motion, respiratory, and cardiac effects on each trial to account for variance in the BOLD signal associated with these sources of noise.

Our primary general linear model (GLM), GLM-CH, tested for regions that correlated with HRV during self-control challenges. The regression modeled as events of interest all trials that contained (1) a challenge, (2) no challenge, while controlling for (3) healthier experimenter recommendations and (4) less healthy experimenter recommendations. Note that the experimenter recommendations were included in the choice task to test a separate hypothesis unrelated to the current report. Those recommendations are not discussed here. Self-control challenge and no-challenge trials included parametric modulators for relative health and taste differences. We computed a first-level contrast of the regions that correlated with HRV during self-control challenges. The regions were compared against the null hypothesis of no correlation.

No Challenge trials. At the second (group) level, we performed group-level analyses because the TFCE and permutation algorithms are designed to be more powerful than the randomization test and were implemented in FSL for all group-level analyses.

For these participants, only the standard motion-correction procedure was applied.
A second, separate GLM, GLM-SV, was computed to determine whether BOLD activity was related to the integrated value of the chosen food. This GLM included parametric regressors for the integrated subjective value of the chosen and nonchosen food items on every choice onset. Once again, additional regressors controlling for the impact of the experimenter recommendations were included in the model with separate regressors for events in which participants chose based on the recommendation, and in which they did not follow the recommendation. We modeled each participant’s subjective value of food items on every trial by combining the weighted values for the taste and health of each food. The weights were derived from individual logistic regressions on the participant’s choices (identical to Maier et al., 2015). Briefly, for each participant, a logistic regression estimated the probability of choosing the left item as a function of the taste and health of the left and right item, with all ratings z-scored within participant before entering them in the model. Two additional binary regressors indicated whether the left or right item had been recommended. These regressors for left and right item recommendations took the value of 1 when the item was recommended, and 0 when it was not recommended. When no recommendation was given on a trial, both regressors had a value of zero. Note that the spatial presentation of the items was completely randomized, so that the left item was equally likely to be the healthier or the tastier of both options. We took the mean of the taste betas for the left and right item obtained for each participant, averaging them into a common taste weight for this individual. The same was done for health. These averaged taste and health weights were then used to multiply the z-scored taste and health values of each item presented in the choice paradigm. To obtain the subjective value for each food, the weighted taste and health values were added up to a weighted subjective value separately for the left and right food items.

We computed a first-level contrast for the chosen food value for each participant and extracted betas from this contrast within our functional ROI of the vmPFC.

To examine the impact of health and taste attributes on the BOLD signal, we used a third GLM, GLM-HT, that modeled the following five events: (1) all choices, (2) trials on which the healthier food was recommended and chosen, (3) trials on which the healthier food was recommended and not chosen, (4) trials on which the less healthy food was recommended and chosen, and (5) trials on which the less healthy food was recommended and not chosen. Note that the 30 baseline trials did not contain any recommendation and, therefore, the sum of regressors 2–5 does not equal regressor 1. The first regressor for all choices included the following four parametric modulators: (1) health of the chosen item (Hc), (2) taste of the chosen item (Tc), (3) health of the nonchosen item (Hnc), and (4) taste of nonchosen item (Tnc). These parametric regressors were not orthogonalized with respect to one another. We computed first-level contrasts for Tc–Tnc and Hc–Hnc. We then extracted the betas for these contrasts from our functional vmPFC ROI. The significance of the correlations between BOLD sensitivity to taste and health differences in the vmPFC and HRV were determined from 5000 permutations of the data.

Anatomical masks. The combined anatomical mask for the vmPFC was constructed from a conjunction of the bilateral frontal pole, frontal medial, paracingulate, and subcallosal cortex areas that exceeded 20% probability of belonging to the respective structure in the Harvard–Oxford Cortical Atlas (HOA; Desikan et al., 2006). To limit the mask to our ROI along the medial wall, the HOA-derived anatomical mask was intersected with a rectangular box around the midline (coordinates in mm: x = [−22, 21], y = [−110, 73], z = [−35, 91]). The anatomical mask of the left dlPFC was constructed from a conjunction of the left inferior frontal gyrus (pars opercularis and reticularis) and left superior frontal gyrus areas that exceeded 20% probability of belonging to these structures, according to the HOA.

Because we tested two separate regions of interest (vmPFC and dlPFC), we used a critical value of $p < 0.025$ (i.e., 0.05/2) for small-volume correction at the voxel level.

### Results

**HRV**

The mean duration of RR intervals across all participants was 929.3 ± 136.3 ms (sample median of the median duration of RR intervals: 947 ± 115 ms), resulting in a mean heart rate of 66 ± 10 beats per minute in our sample (values are derived after deletion of artifacts). Our participants expressed a median total HRV (measured as SDNN) of 98.7 ± 50.1 ms MAD within our 3 min baseline measurement. Consistent with previous reports (Tsuji et al., 1996), HRV was inversely related to average heart rate ($p = −1.17 ± 0.47$, $T = −2.48$, $p = 0.017$; see Table 1). However, total HRV did not differ between participants later assigned to the Stress or Control groups (Stress, 98.7 ± 29.6 ms; Control, 97.7 ± 30.9 ms; $p = 0.93$; $Z = 0.09$; Wilcoxon rank-sum test). Regarding biological and psychological markers of the stress reaction, baseline SDNN and cortisol reactions (area under the curve with respect to ground; Pruessner et al., 2003) were not significantly correlated ($r = −0.19$, $p = 0.18$), nor were baseline SDNN and perceived stress ($r = −0.10$, $p = 0.49$). Visual inspection of a scatter plot revealed one outlier in the SDNN measure: the value for this participant fell between 2 and 3 SDs from the sample mean. Therefore, we checked our results for robustness with and without this participant’s data and found that all results remained significant in both cases.

One concern in the evaluation of HRV is that applying artifact correction might inflate indices of HRV (Heathers, 2014; Quintana and Heathers, 2014) and as few as one edited artifact in the RR interval series may do so (Berntson and Stowell, 1998). Indeed, we observed a significant positive correlation between the number of artifacts that we corrected per dataset and the SDNN ($r = 0.44$, $p = 0.003$; CI [0.26, 0.61]). Thus, we included the number of corrected artifacts and mean heart rate as additional covariates in our regression model to test whether HRV would be predictive beyond these influences.

**HRV and self-control behavior**

We originally defined SCS as choosing the healthier, but less tasty, of two food items in challenging trials in which health and taste conflicted, meaning that the participant had to overcome his own taste preferences to choose the healthier option. We initially

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### Table 1. GLM predicting total HRV (GLM-HRV)

<table>
<thead>
<tr>
<th>Regressor</th>
<th>β Estimate</th>
<th>SEM</th>
<th>t</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intercept</td>
<td>189.41</td>
<td>31.17</td>
<td>6.08</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Number of artifacts</td>
<td>5.07</td>
<td>1.49</td>
<td>3.41</td>
<td>0.0014</td>
</tr>
<tr>
<td>Mean heart rate</td>
<td>−1.17</td>
<td>0.47</td>
<td>−2.48</td>
<td>0.017</td>
</tr>
<tr>
<td>Trait anxiety</td>
<td>−0.57</td>
<td>0.63</td>
<td>−0.91</td>
<td>0.37</td>
</tr>
</tbody>
</table>

Results from a GLM with possible determinants of HRV (represented as untransformed values of SDNN in millimeters): the number of artifacts corrected in the dataset and mean heart rate (after artifact correction by deletion). To assess whether trait anxiety explains additional variance beyond an increase in mean heart rate, we added a regressor with the trait anxiety score as measured by the Spielberger State-Trait-Anxiety Inventory, so that mean heart rate and trait anxiety compete for variance in the same model. When doing so, mean heart rate accounts for a decrease in HRV, but trait anxiety does not explain further variance. The results above hold when excluding the HRV outlier from this model.
tested the relationship between total HRV (i.e., SDNN) and SCS in a bivariate correlation analysis. Total HRV was associated with the frequency of SCS in the dietary-choice task over all participants (Pearson’s $r = 0.36$, $p = 0.01$; CI [0.07, 0.59]; all $p$ values are derived from 5000 permutations of the data; excluding the HRV outlier: $r = 0.33$, $p = 0.02$, CI [0.03, 0.58]). For comparison, the correlation between SCS and the cognitive restraint in eating score (RE) obtained from the Three Factor Eating Questionnaire (Pudel and Westenhöfer, 1989) was $r = 0.35$ ($p = 0.01$; CI [0.11, 0.55]). This restraint score captures the degree to which individuals use cognitive strategies to limit calorie intake, for example by counting calories, deliberately picking small portions of food, or consuming foods with lower calorie content. Thus, as a biomarker of dietary self-control, HRV explains approximately the same amount of individual variance in choice behavior as an established psychometric index of eating behavior. Restraint eating scores were not significantly associated with HRV ($r = 0.14$, $p = 0.35$; CI $[-0.28, 0.42]$), suggesting that the two measures could be readily combined to explain additional variation in self-control behavior.

Therefore, we modeled SCS (i.e., choosing a healthier, less tasty item) in a multiple regression that included both HRV and RE together (Eq. 2). Beyond testing whether HRV and RE could be combined to explain additional variance in self-control, we included RE in the model because it is a widely used, validated measure for dietary SCS (Laesse et al., 1989; Allison et al., 1992; Williamson et al., 2007). As such, it serves as a benchmark for judging the utility of HRV as a biomarker for dietary self-control. When including both RE and HRV into the same model, we can assess whether HRV is predictive beyond a known trait characteristic of dietary SCS. SCS was calculated according to the following equation (Eq. 2):

$$(\text{Stress} + \text{HRV} + \text{RE}) \times (\text{Hdiff} + \text{Tdiff}) + \text{error}.$$  

This regression allowed us to examine potential interactions between the individual characteristics of HRV and RE and task features, such as the stress manipulation (Stress), as well as the average health (Hdiff) and taste differences (Tdiff) a participant faced within the food-choice task. Note that although all participants faced self-control challenges in the food-choice task, the degree of the challenges depended on each individual’s opinions on the taste and healthiness of the various foods.

Higher HRV and RE characteristics reduced the influence of taste temptations in self-control dilemmas. All results from the regression in Equation 2 are listed in Table 2. Higher HRV levels increased the degree to which high taste temptations (i.e., taste differences) were overcome, leading to greater SCS (Fig. 2b). Moreover, unlike low-HRV participants, those with high HRV successfully used self-control regardless of the average health difference between the two options, suggesting that they engaged self-control even when the benefit of doing so (i.e., the increase in healthiness) was relatively small (Fig. 2c). A similar interaction was observed between RE and taste differences. As previously reported (Maier et al., 2015), acute stress reduced the use of self-control in dietary choice. To determine whether HRV could act as a buffer against acute stress, we computed an extended version of the model in Eq. 2 that also included interactions between stress and HRV and between stress and RE. However, we did not observe significant interactions between Stress and HRV or RE, indicating that the relationship between HRV and dietary self-control persisted in both the Stress and Control groups, but that HRV was not associated with resilience to acute stress. For simplicity and ease of interpretation, we report the reduced model without interaction terms in Table 2.

As a robustness check, we controlled for the influences of the HRV outlier and several other factors that might relate to HRV. We estimated the basic model (Eq. 2) without the HRV outlier, and added the following: age, the combined number of cardio and strength exercise sessions per week, BMI, hunger level, trait anxiety score, mean heart rate, and number of corrected artifacts in the recording as nuisance regressors. The results were qualitatively unchanged and we still observed a significant relationship between taste and HRV as described above ($T = 4.77$, $p = 4.85e^{-05}$). Note that we included trait anxiety in this robustness check because previous work has shown that trait anxiety is also correlated with HRV (Gaburro et al., 2011; Verkuil et al., 2014), and therefore we tested for this relationship in our data as well. We used the trait scale of the Spielberger State-Trait Anxiety Inventory as our measure of anxiety (median score for our sample: 34.5 ± 5.9 out of possible score ranging from 20 to 80). Consistent with previous reports, we observed a negative correlation between HRV and trait anxiety scores (correlation $r = -0.28$, $p = 0.0488$; CI $[-0.48, -0.06]$). Trait anxiety scores were also positively correlated with mean resting heart rates across participants ($r = 0.39$, $p = 0.01$; CI [0.11, 0.60]).

To assess the predictive qualities of RE and HRV with regard to self-control in a more robust way, we predicted self-control levels out-of-sample using the leave-one-subject-out (LOSO) method. After taking one participant’s data out of the sample, we fit the model in Eq. 2 to explain the variance in self-control levels of the remaining participants. Using the $\beta$ coefficients from the training set, we then predicted the self-control level of the left-out participant. Squaring the obtained correlation coefficient for the true and predicted self-control levels in the full model ($r = 0.88$, $p < 0.0001$, CI [0.81, 0.92]) yielded the coefficient of determination for the model with combined predictors of RE and HRV, $R^2 = 0.77$ (Fig. 2a). For comparison, fitting the model without the predictors for HRV and its interactions yielded a lower correlation between true and predicted self-control levels ($r = 0.81$, $p < 0.0001$, CI [0.63, 0.88]), leading to an observed $R^2 = 0.65$. That is, by combining RE and HRV, we could significantly explain an extra 12% of the variation in out-of-sample self-control rates. Using a “split half” instead of an LOSO procedure (i.e., randomly sampling half the dataset for model fitting and using

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Results from a GLM of SCS. SCS was computed as the mean number of trials in which participants chose the healthier, less tasty item in trials in which health and taste were not aligned (challenge trials). Resting HRV was defined as the SD of all RR intervals over a 3 min period. The cognitive restraint in eating score (RE) was obtained from the Three Factor Eating Questionnaire (Pudel and Westenhöfer, 1989). The regresses, Stress, is a binary factor indicating that the participant underwent the stress manipulation. The average health (Hdiff) and taste difference (Tdiff) regressors represent the mean taste temptation and health gain that participants were faced with during the dietary choice task. All estimates are reported with their SEM.
the remaining half for predicting out of sample) yielded a similar benefit (16%) for including HRV in the predictive model (full model: $r = 0.81, p < 0.0001, CI [0.55, 0.92], R^2 = 0.66$; model without HRV: $r = 0.71, p = 0.00004, CI [0.26, 0.89], R^2 = 0.50$).

During the initial review of this manuscript, an anonymous reviewer made the insightful suggestion that we could also test the relationship between HRV and another form of self-control in our dataset. Our food-choice paradigm included recommendations about which food to choose, and in some cases these recommendations were in favor of the unhealthy food (participants were told that recommendations were usually, but not always in favor of the healthier item). This feature enables us to test whether HRV is associated with the fraction of trials in which participants overrode this unhealthy recommendation and still chose the healthier food. We found that there was also a positive correlation between HRV and this alternative measure of control ($r = 0.31, p = 0.01, CI [0.01, 0.56]$). These two measures of self-control are not independent because both are based, in part, on the incorporation of health attributes into the choice process (correlation between self-control measures: $r = 0.77, p < 0.0001, CI [0.62, 0.87]$). Even so, this post hoc finding is consistent with the idea that HRV is a domain general marker of self-regulatory ability or efficiency as outlined in the introduction section.

**HRV and BOLD activity during self-control**

To investigate whether HRV could serve as a biomarker of changes in the brain’s decision circuitry in the food self-control paradigm, we analyzed BOLD activity measured during the choice task. Our primary general linear model (GLM-CH) tested for regions that correlated with HRV during self-control challenges.

HRV positively correlated with self-control at the level of observed choices. We hypothesized, therefore, that HRV would be associated with BOLD activity in regions known to be involved in the value computation process during self-controlled choices, namely the vmPFC and dlPFC. We tested this hypothesis in anatomical masks of the vmPFC and dlPFC based on the Harvard-Oxford Cortical Atlas (Desikan et al., 2006). The vmPFC mask comprised the bilateral vmPFC that is part of the brain’s valuation system (Bartra et al., 2013; Clithero and Rangel, 2014; Abitbol et al., 2015; Pessiglione and Delgado, 2015) and has been shown to integrate taste and health values in the dietary self-control paradigm (Hare et al., 2009, 2011, 2014; Foerde et al., 2015; Maier et al., 2015) as well as the separate characteristics of multiattribute choices in other, nonfood domains (Kahnt et al., 2011; Rudorf and Hare, 2014). Our second anatomical mask included the region of the left dlPFC that has been presumed to modulate activity in the vmPFC during self-control choices (Hare et al., 2009, 2011, 2014). We found that BOLD activity in portions of the vmPFC increased as a function of baseline HRV in Challenge > No Challenge trials [MNI peak: [1 46 0] in the paracingulate/cingulate gyrus, small-volume-corrected (SVC) $p = 0.004, T = 4.8$, and a separate, more dorsal cluster in the cingulate gyrus at [21 41 9], SVC $p = 0.038, T = 3.86$; Fig. 3a]. However, we found no association between HRV and BOLD activity in the left dlPFC for the Challenge > No Challenge trials that survived SVC within the anatomical region of the left dlPFC. Exploratory whole-brain analyses yielded no other regions that survived correcting for multiple comparisons. The results of these exploratory analyses can be accessed in a Neurovault repository under the following link: http://www.neurovault.org/collections/DNXFVQPJ/.

To establish that activity in this vmPFC region (the larger cluster with a peak at $x, y, z = 1 46 0$) was relevant to the participants’ choices, we tested whether the chosen food values were represented in the functional region of interest (ROI) correlating with HRV. An integrated value of the chosen food was calculated in a separate GLM (GLM-SV) and we extracted the betas for this chosen food value in the vmPFC ROI. We found that activity in this ROI encoded the integrated value of the chosen food ($T_{(46)} = 3.52, p < 0.001$).

Given our behavioral data linking HRV to the relative influence of taste on dietary choices, we tested whether HRV was also correlated with the degree to which BOLD activity in the vmPFC ROI represented taste attributes (Taste chosen − Nonchosen from GLM-HT). We found that HRV was negatively correlated with the relative taste value representation (Pearson’s $r = -0.42, p = 0.002, CI [-0.60, -0.19]$; Fig. 3b), but not correlated with the relative health value ($r = -0.12, p = 0.42, CI [-0.43, 0.21]$; Fig. 3c). Excluding the HRV outlier did not change the results (taste $r = -0.43, p = 0.004, CI [-0.63, -0.17]$; health $r = -0.09, p = 0.56, CI [-0.43, 0.23]$).

**Discussion**

We found that higher HRV is associated with better self-control in the face of dietary challenges. More specifically, our results show that the choices of individuals with higher HRV are less affected by tempting taste attributes than choices of participants with lower HRV. In parallel, at the neural level, higher HRV correlated with a decreased representation of taste attributes in the vmPFC, a brain region that has been associated with both regulating autonomic responses (Benarroch, 1993) and calculating subjective values of choice options (Bartra et al., 2013; Clithero and Rangel, 2014; Abitbol et al., 2015; Pessiglione and Delgado, 2015). HRV is a measure of physiological fitness that
relates to the integrated functioning of the nervous and cardiac systems. Similarly, successful self-control relies on the integration, and potentially modified evaluation, of actions in the context of higher-order goal attainment. Our data indicate a significant association between these integration processes at the basic physiological (i.e., HRV) and cognitive (i.e., self-control) levels, suggesting that HRV measures may serve as a useful and readily obtainable biomarker for self-control abilities.

Resting HRV measured over a few minutes with relatively inexpensive and commercially available equipment predicted subsequent self-control in a dietary-choice task as well as a validated psychometric index of dietary behavior [restrained eating scale of the Three Factor Eating Questionnaire (RE)]. Moreover, as a physiological measure presumably outside the domain of conscious control, HRV also has the advantage of being immune to socially desirable reporting (Logan et al., 2008; Devylder and Hilimire, 2015) or memory errors that can affect the accuracy of self-reports. However, when entered into a joint model, both HRV and RE were significantly related to dietary self-control, suggesting that they explained separate components of the variance in dietary choice. Thus, it is possible to combine biomarkers, such as HRV, with behavioral and self-report measures (e.g., the Three Factor Eating Questionnaire) to predict future self-control more accurately.

The fact that such biomarkers as HRV can be easily acquired and readily combined with other survey or task-based measures of self-control is important because, taken in isolation, any single measure is likely to reveal only a partial picture of self-control abilities or proclivities. In a recent meta-analysis by Duckworth and Kern (2011), informant-report and self-report questionnaires, behavioral readouts of executive function, and delay-of-gratification measures showed only moderate convergent validity. In other words, self-control assessed in one fashion was only moderately related to self-control measured in another manner. In agreement with those authors, we believe that self-control is a multidimensional construct best assessed, and potentially forecasted, by combining measures taken across multiple domains, including behavior, self and informant report, and both neural and more general physiological markers, such as HRV.

HRV can explain individual differences in self-control that are robust to changes in environmental context. We have previously shown that experiencing an acute stressor results in diminished self-control in the 45 min period following stressor onset (Maier et al., 2015). In the same sample, we find that resting HRV before stressor onset predicts the level of self-control following stress as well as it predicts choice in the control (i.e., not stressed) participants. Thus, the association between HRV and self-controlled behavior is maintained under the influence of acute stress, suggesting that the association between HRV and self-control may be context-independent.

Our current findings linking resting HRV to subsequent self-control performance extend previous work reporting correlations between HRV and neural activity measured simultaneously during affective and cognitive tasks. In a study using positron emission tomography (PET) and an emotion task, Lane et al. (2009) measured regional cerebral blood flow (rCBF) when participants immersed themselves during 1 min blocks into positive, negative, and neutral emotions (evoked by film clips and vignettes of personal emotional memories) while parallel PET and HRV were recorded. During the presentation of emotional (vs neutral) stimuli, HRV correlated with rCBF in the caudate, midbrain, left insula, and medial prefrontal cortex. When excluding all emotion-specific activation, HRV correlated with rCBF in the right dlPFC, the bilateral parietal cortex, and the left rostral anterior cingulate cortex (ACC) with high-frequency (parasympathetic) components of HRV. Similarly, Gianaros et al. (2004) used PET to correlate HRV with changes in rCBF in medial orbitofrontal cortex (OFC), insula, ACC, amygdala, hippocampus, and cerebellum as a function of task demand in a working-memory paradigm. A study by Nugent et al. (2011) found HRV to be correlated with rCBF in lateral and medial OFC when participants had to achieve different levels of strength in a handgrip task.

In contrast to Lane and colleagues’ (2009) emotion task results and our a priori predictions, we did not observe any significant correlations between HRV and activity in the dlPFC during dietary choices. However, the differences in HRV indices and measurement times (resting vs task) preclude direct comparisons between the previous emotion regulation and current dietary self-control results. It is possible that HRV measures collected during the self-control task would tie in more closely with regulation processes in the dlPFC. However, our goal in the current study was to test whether simple, task-independent measures of HRV are associated with dietary self-control. What is consistent across studies is that individual differences in HRV are correlated with activity in neural regions linked to task performance across several domains (e.g., emotion regulation, working memory, physical effort, and dietary self-control). Together these results indicate that efforts to link cognition with central and peripheral neurophysiology may promote a better understanding of the nature of individual differences in health and cognitive behaviors, and provide opportunities for prediction and early intervention against dysfunctions (Sokol-Hessner et al., 2009; Raio et al., 2013).

Our study represents an important initial step in linking total HRV to self-control ability. This result suggests total HRV should be considered when investigating links between self-control and allostatic capacity in addition to more direct indices of phasic vagal cardiac control of HRV, such as RMSSD and high-frequency HRV. One rationale for doing so in the domain of dietary self-control is that frequency components outside the high-frequency spectrum may include information on metabolic and endocrine processes that are directly relevant to dietary decisions. These components contribute to the total HRV, but oscillate on a slower time scale (Berntson et al., 1997).

Further progress could be made by addressing this question with causal manipulations, for example by inducing endocrine signals of hunger and satiety and investigating whether the association between total HRV and self-control success varies during these states. Another interesting avenue to pursue is whether plasticity-induced changes that enable better regulation, for example through transcranial electrical or magnetic stimulation of the dlPFC, might also lead to an increase in HRV. A study in autistic children suggests this might be the case: Wang et al. (2016) found that weekly treatment with low-frequency repetitive transcranial magnetic stimulation for 3 months both improved chronic autonomic imbalance (i.e., higher low-frequency and lower high-frequency contributions to total HRV, putatively reflecting a tonically high arousal level due to activation of the sympathetic nervous system) and reduced the tonically elevated skin conductance levels commonly seen in autism. This change was accompanied by decreased irritability, reduced hyperactivity, and less stereotyped and compulsive behavior in the autistic children. Future work could therefore address this regulatory mechanism in a healthy population with a similar causal manipulation by stimulation techniques to further explore the nature of the link.
between neural correlates of self-regulation and physiological markers of allostatic capacity.

In conclusion, HRV is a marker of cardiovascular and mental health. Our results indicate that HRV also explains significant variation in self-control during dietary choice. Moreover, both HRV and self-control are fundamental psychometric scale of restrained eating contributted independently to explaining variance in our behavioral model of self-control and could be used in combination to better predict dietary self-control levels.

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