Energy Expenditure Responses to Fasting and Overfeeding Identify Phenotypes Associated With Weight Change

Because it is unknown whether 24-h energy expenditure (EE) responses to dietary extremes will identify phenotypes associated with weight regulation, the aim of this study was to determine whether such responses to fasting or overfeeding are associated with future weight change. The 24-h EE during energy balance, fasting, and four different overfeeding diets with 200% energy requirements was measured in a metabolic chamber in 37 subjects with normal glucose regulation while they resided on our clinical research unit. Diets were given for 24 h each and included the following: 1) low protein (3%), 2) standard (50% carbohydrate, 20% protein), 3) high fat (60%), and 4) high carbohydrate (75%). Participants returned for follow-up 6 months after the initial measures. The decrease in 24-h EE during fasting and the increase with overfeeding were correlated. A larger reduction in EE during fasting, a smaller EE response to low-protein overfeeding, and a larger response to high-carbohydrate overfeeding all correlated with weight gain. The association of the fasting EE response with weight change was not independent from that of low protein in a multivariate model. We identified the following two independent propensities associated with weight gain: a predilection for conserving energy during caloric and protein deprivation and a profligate response to large amounts of carbohydrates.

Human overfeeding studies (1–5) suggest that there is a considerable interindividual variation in the energy cost of weight gain. In a prior cross-sectional study (5), the increase in energy expenditure (EE) with overfeeding and the decrease with fasting (FST) were found to be correlated in a small group of 14 male subjects. Our group has previously shown that the EE response to overfeeding varies considerably among individuals but is consistent and reproducible within individuals. This individual contribution explains more of the observed variability in the EE changes with overfeeding than changes to the macronutrient content of the diet (6). These studies seem to indicate that phenotypic differences may exist in the EE responses to FST or overfeeding that may affect susceptibility to weight gain. As overeating or caloric restriction are necessary to alter weight, perturbations in energy balance (EB) may be needed to uncover responses that signify an energy-conserving physiology versus a physiology that is better able to resist weight gain. We now extend our previous findings by addressing the question of whether this interindividual variation in EE changes relates to future weight change.

During overfeeding, the metabolic response depends, in part, on the macronutrient composition of the diet in addition to the contribution from interindividual variation (6). Although it has been proposed that low-protein diets might magnify differences in the propensity to obesity (2,7), a recent study (8) has shown that the EE response is smaller and fat mass (FM) gain is similar when overeating low-protein diets compared with normal-protein diets. Further, high-carbohydrate diets have been shown to have a greater EE increase during overfeeding compared with high-fat diets (9). In addition, a single large high-carbohydrate meal has been shown to activate brown adipose tissue (10). Differences in the short-term

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(24-h) EE response to overeating diets varying in macronutrient content may therefore facilitate the identification of human phenotypes with increased susceptibility to future weight gain. We hypothesized that a larger reduction in EE during FST and a smaller increase in EE during 24 h of overfeeding would be associated with weight gain at 6 months in free-living, healthy individuals who had not been counseled on any lifestyle changes. In addition, we hypothesized that varying the macronutrient content of the overfeeding diet might identify macronutrient-specific differences in the EE response to overeating that would be more strongly associated with future weight change.

RESEARCH DESIGN AND METHODS

Subjects
Volunteers were recruited from the Phoenix, AZ, area between 2007 and 2013 and were admitted to our clinical research unit (CRU) for 25 days to participate in an inpatient study exploring the metabolic responses to FST and overfeeding. Among the 59 individuals who completed the baseline CRU admission, 37 had follow-up data for body weight 6 months after CRU discharge and were included in the present analysis (Fig. 1). This report represents a preplanned analysis of an ongoing study when a target sample size of 37 subjects had completed the 6-month follow-up to provide 90% power ($\alpha = 0.05$) to detect a simple correlation of 0.5 between the percentage change in EE with overfeeding or FST and the primary end point of body weight change at follow-up. These 37 individuals did not differ from the larger initial group with regard to demographics, anthropometrics, and 24-h EE measures. All subjects reported a stable weight for at least 6 months and were healthy according to history, physical examination, electrocardiogram, and laboratory test results. None of the subjects had a vegetarian or gluten-free lifestyle, and none had a known food allergy. All women were premenopausal and not pregnant. All volunteers provided informed, written consent. The experimental protocol was approved by the Institutional Review Board of the National Institute of Diabetes and Digestive and Kidney Diseases.

Upon CRU admission, volunteers were given a weight-maintaining diet (WMD) consisting of 50% carbohydrates, 30% fats, and 20% proteins, with total caloric content based on previously derived equations specific to our CRU that include weight, BMI, and sex (11). Morning weight was checked daily, and the WMD was adjusted as necessary throughout the CRU stay to maintain a stable weight ($\pm 1\%$). The WMD was given throughout the stay except on the days when subjects had 24-h EE assessments. Volunteers were asked to consume all food given to them, and to engage only in sedentary activities for the duration of their stay on the CRU. Body composition was measured using DXA (DPX-1; Lunar Corp, Madison, WI). After 3 days on the WMD, a 75-g oral glucose tolerance test was performed. Only individuals with normal glucose regulation (12) were eligible to participate. Plasma glucose concentrations were measured using an enzymatic oxygen-rate method (Beckman Glucose Analyzer 2; Beckman Instruments, Brea, CA) ($n = 7$) or the comparable Analox GM9 glucose oxidase method (Analox Instruments USA, Lunenburg, MA) ($n = 30$).

![Figure 1](diabetes.diabetesjournals.org) Schlögl and Associates 3681

Figure 1—Flow diagram of participant progress through the study.
EE Measures and Dietary Interventions

Each volunteer completed seven 24-h EE assessments in a whole-room indirect calorimeter: two eucaloric assessments (the first eucaloric measurement in the metabolic chamber [EB0] and EB) followed by five EE measurements during the dietary interventions described below (Fig. 2). There was a 3-day washout period between each dietary intervention to allow any residual effects of the 24-h dietary intervention to wane. The average coefficient of variation (CV) of the volunteers’ body weight prior to the dietary interventions was 0.94 ± 0.48%, indicating that body weight was stable (<1%) during the admission period.

For all diets, volunteers were given breakfast at 0700 h and entered the calorimeter 1 h later. Further meals were provided inside the calorimeter at 1100, 1600, and 1900 h through a two-door airlock. Total energy intake of the four meals given during EB0 was 80% of the WMD to account for reduced activity in the calorimeter (13). To increase the precision of the EE measure during EB, energy intake during the second eucaloric measurement (EB) was equal to the 24-h EE value measured in EB0. The 24-h EE from this second eucaloric assessment (EB), which was used as the baseline comparator, was then doubled to determine the number of kilocalories given for the subsequent overfeeding diets (200% energy requirements).

Volunteers completed in randomized order five intervention diets, each of which was administered for only 24 h, as follows: fasting (FST); low-protein overfeeding with 51% carbohydrate, 46% fat, and 3% protein (LPF); standard overfeeding with 50% carbohydrate, 30% fat, and 20% protein (SOF); high-fat, normal-protein overfeeding with 20% carbohydrate, 60% fat, and 20% protein; and high-carbohydrate, normal-protein overfeeding with 75% carbohydrate, 5% fat, and 20% protein (CNP) (Fig. 2). The macronutrient composition of each diet was determined using The Food Processor software (ESHA Research, Salem, OR). Subjects returned all uneaten portions to the metabolic kitchen for weighing, so that actual intake by macronutrient could be calculated. Five (2% of total 222 chamber sessions) EE measurements (1 standard, 2 high-fat, and 2 high-carbohydrate diet) were excluded as <95% of food was consumed.

Ambient temperature averaged 23.6 ± 1.4°C. The average O₂ consumption and CO₂ production were used to calculate the 24-h EE and respiratory quotient (RQ), as previously described (6). The RQ was used as a proxy for the carbohydrate-to-fat oxidation ratio. Quality control tests were performed monthly, and demonstrated mean recoveries of 99 ± 3% (CV 3.6%) and 98 ± 3% (CV 3.4%) for O₂ and CO₂, respectively. EB was the difference between caloric intake and 24-h EE. Spontaneous physical activity (SPA) was detected by radar sensors and WAS expressed as the percentage of time in which motion was detected.

Follow-up Visit

Upon completion of the EE assessments, participants were not provided with any lifestyle counseling and were advised to return to their usual habits. They were, however, provided with the results of their DXA scan and oral glucose tolerance test. Participants were discharged from the CRU and were asked to return at a scheduled 6-month follow-up visit for the measurement of weight and body composition.

Statistical Analysis

Statistical analyses were performed using the SAS version 9.2 (SAS Institute, Cary, NC). The α value was set at 0.05. Data are presented as the mean ± SD. The Shapiro-Wilks test was used to assess the normality of the data. Data

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**Figure 2** — Study diagram of the clinical study.

EB0 and EB: energy balance diet with 50% carbohydrate, 30% fat, 20% protein
FST: fasting
LPF: low-protein overfeeding with 51% carbohydrate, 46% fat, 3% protein
SOF: standard overfeeding with 50% carbohydrate, 30% fat and 20% protein
FNP: high-fat, normal-protein overfeeding with 20% carbohydrate, 60% fat and 20% protein
CNP: high-carbohydrate, normal-protein overfeeding with 75% carbohydrate, 5% fat and 20% protein

FST, LPF, SOF, FNP and CNP chambers were done in a random order.
were scanned for potential outliers using the methods of Grubbs (14) and Tukey (15), and the generalized extreme Studentized deviate test (16). No outliers were identified. Differences between groups were evaluated using Student t test or \( \chi^2 \) analyses for continuous and categorical variables, respectively. Ethnic differences were assessed by one-way ANOVA. To normalize the EE response to body size, the percentage change in 24-h EE (%EE) during each dietary intervention was calculated as the difference divided by the 24-h EE during EB and expressed as a percentage, as follows:

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\text{%EE response}_d = \left( \frac{24 \text{ h EE}_d - 24 \text{ h EE}_{\text{energy balance}}}{24 \text{ h EE}_{\text{energy balance}}} \right) \times 100.
\]

Pearson correlations were used to determine the correlations between normally distributed continuous variables, and Spearman correlations were used for non-normally distributed variables. For some analyses, the EE responses to the four overfeeding diets were averaged per person to understand the general effects of overfeeding. Associations with the response to overfeeding were determined from mixed models, accounting for repeated measures and including the variables age, sex, ethnicity, percentage of body fat, and diet. Differences between diets were adjusted for multiple comparisons using the Tukey range test.

Significant correlations between EE responses to FST and overfeeding with weight change were followed-up with regression models to adjust for age, sex, ethnicity, and baseline weight. All results were confirmed using the percentage weight change per month in place of absolute weight change. Similar models were calculated for the absolute changes in FM and fat-free mass (FFM), including initial baseline measures as covariates. Multivariate regression models were created to determine the independence of the identified associations. Adjusting for SPA did not substantially change any results; thus, only findings using unaltered 24-h EE are reported.

**RESULTS**

**Subjects Characteristics**

General, anthropometric, and EE characteristics of the study population during EB are shown in Table 1. Body composition, 24-h EE, and the percentage change in EE with FST or overfeeding did not differ between ethnic groups.

**24-h EE Response to FST or Overfeeding**

Compared with EB, the %EE decreased with FST \(( -8.5 \pm 5.0\% ; P < 0.001)\) and increased with overfeeding (Table 2, Fig. 3B). The average percentage increase in 24-h EE during the four overfeeding diets \((9.0 \pm 4.0\%)\) correlated with the percentage decrease in 24-h EE with FST \((r = 0.55, P = 0.001)\) (Fig. 4A). Individually, the percentage decrease in 24-h EE with FST correlated...
with the %EE during low-protein overfeeding ($r = 0.46$, $P = 0.006$) (Fig. 4B). The percentage increase in 24-h EE during high-carbohydrate overfeeding correlated with the %EE response to the high-fat, normal-protein ($r = 0.53$, $P = 0.002$) and standard overfeeding ($r = 0.38$, $P = 0.02$) diets, as well as with the percentage decrease in 24-h EE with FST ($r = 0.40$, $P = 0.02$). The mean %EE response to overfeeding (the average of all four diets) was inversely related to the percentage of body fat ($r = -0.43$, $P = 0.008$).

In a mixed model accounting for repeated measures, adjusting for age, sex, and ethnicity, and including only the four overfeeding diets, diet ($P < 0.001$) and the percentage of fat ($\beta = -0.12\%$, $P = 0.03$) were independent determinants of the %EE.

**Determinants of Future Weight Change**

Changes in body weight and body composition at follow-up (6.5 ± 0.9 months, range 5.2–9.2 months) are shown in Table 1. The variance in weight change at 6 months was normally distributed ($P = 0.44$, Shapiro-Wilks test) around a mean increase of 1.2 ± 4.4 kg (range −6.1 to 11.2 kg) without any suspected outliers. There was no difference between sexes or ethnicities in body weight change.

A greater reduction in 24-h EE during FST was associated with weight gain at 6 months ($r = -0.35$, $P = 0.04$) (Fig. 5A), and this was still true after adjustment for age, sex, ethnicity, and baseline weight ($\beta = -0.32$ kg per 1% difference in 24-h EE response, $P = 0.05$). Similarly, the %EE response during low-protein overfeeding at baseline was negatively associated with absolute body weight change ($r = -0.55$, $P = 0.001$) (Fig. 5B), and this held true after adjustment for age, sex, ethnicity, and baseline weight ($\beta = -0.42$ kg per 1% increase in 24-h EE response, $P = 0.01$). There was no association between the change in body weight and the average %EE during the three overfeeding diets with 20% protein content ($r = 0.16$, $P = 0.35$), and the EE responses to standard ($r = 0.03$, $P = 0.86$) or high-fat, normal-protein ($r = 0.06$, $P = 0.75$) overfeeding were also not associated with weight change. The EE response to high-carbohydrate, normal-protein overfeeding was positively associated with weight change at follow-up ($r = 0.33$, $P = 0.05$, $\beta = 0.41$ kg per 1% increase in 24-h EE, $P = 0.009$ adjusted for age, sex, ethnicity, and baseline weight) (Fig. 5C). In a multivariate model, both the 24-h EE responses to low-protein ($\beta = -0.44$ kg per 1% difference in 24-h EE response, $P = 0.004$) and high-carbohydrate, normal-protein ($\beta = 0.38$ kg per 1% difference in 24-h EE response, $P = 0.003$) overfeeding, but not the 24-h EE

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**Table 2—Extent of 24-h EE responses during eucaloric feeding, 200% overfeeding with diets varying in macronutrient content, and FST**

| Diet     | 24-h FQ (ratio) | 24-h RQ (ratio) | 24-h EE (kcal/day) | Change in 24-h EE (%) | TEF (%) | 24-h SPA (%) |
|----------|-----------------|-----------------|--------------------|-----------------------|---------|--------------|
| EB       | 0.86            | 0.87 ± 0.03     | 2,036 ± 281        | N/A                   | 8.4 ± 4.9† | 5.4 ± 3.3     |
| FST      | 0.71            | 0.79 ± 0.03     | 1,857 ± 224†       | -8.5 ± 5.0‡          | N/A     | 5.0 ± 3.7     |
| LPF      | 0.85            | 0.91 ± 0.05†    | 2,093 ± 299†       | 2.8 ± 4.9†           | 5.7 ± 2.7‡ | 5.7 ± 4.1     |
| SOF      | 0.86            | 0.89 ± 0.04*    | 2,251 ± 339*       | 10.9 ± 5.7‡          | 9.9 ± 3.3‡ | 5.9 ± 3.3     |
| HPF      | 0.78            | 0.83 ± 0.04*    | 2,186 ± 319*       | 8.7 ± 4.9†           | 8.7 ± 3.0‡ | 5.6 ± 3.7     |
| CNP      | 0.93            | 0.94 ± 0.05*    | 2,330 ± 321*       | 14.4 ± 5.3‡          | 11.8 ± 3.5† | 6.5 ± 4.2     |

Data are presented as the mean ± SD, unless otherwise indicated. FQ, food quotient; HPF, high-fat overfeeding; N/A, not applicable; TEF, thermic effect of food. The FQ (i.e., the expected 24-h RQ based on the macronutrients in each diet) was calculated from published equations (6). The %EE was calculated with respect to the 24-h EE during EB. The TEF of each diet was calculated by subtracting the 24-h EE during FST from the 24-h EE during the relevant dietary intervention, and then was expressed as a percentage of the corresponding total caloric intake. †$P < 0.05$ by Tukey range test compared with EB. ‡$P < 0.05$ vs. 0. §$P < 0.0001$ vs. 0.

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**Figure 3**—Macronutrient composition of the dietary interventions (A) and related 24-h EE response (B). Protein, carbohydrate, and fat content of the diets are expressed in grams based on a representative diet for an individual requiring 2,000 kcal for EB and 4,000 kcal for overfeeding (A). The 24-h EE response to each dietary intervention is expressed as the percentage change compared with the 24-h EE measured during EB (B). Error bars represent the mean with SD.
response to FST ($\beta = -0.15$ kg per 1% difference in the 24-h EE response, $P = 0.18$), were independently associated with weight change at follow-up. Results did not change with serial adjustment for age, sex, ethnicity, or baseline weight. The EE response to FST was only significantly associated with weight change when the 24-h EE response to low-protein overfeeding was removed from the multivariate model.

To further illustrate the independent effects of the EE response to low-protein and high-carbohydrate overfeeding on weight change, we categorized subjects in four subgroups according to the median %EE during these two overfeeding diets (Fig. 6). Subjects with a higher-than-median EE response during high-carbohydrate overfeeding and a lower-than-median EE response during low-protein overfeeding ($n = 7$) gained more weight compared with those with the opposing EE responses ($n = 6$) (mean difference 7% increase in their baseline weight, $P = 0.007$), despite similar baseline body weight ($P = 0.80$). The 24-h EE response to low-protein overfeeding was associated with changes in both FM ($r = -0.48$, $P = 0.004$) and FFM ($r = -0.36$, $P = 0.04$) at 6 months; however, the EE response to high-carbohydrate overfeeding was associated only with the FM change ($r = 0.37$, $P = 0.04$), but not with the FFM change ($P = 0.6$).

The 24-h RQ during FST ($r = -0.41$, $P = 0.01$), but not during any overfeeding diet ($P > 0.2$), was negatively associated with weight change (Fig. 5D), and this was still true after adjustment for age, sex, ethnicity, and baseline weight ($\beta = -0.56$ kg per 0.01 change in FST RQ, $P = 0.01$). The FST RQ was associated with the FFM change at 6 months ($r = -0.34$, $P = 0.05$), but not with the change in FM ($r = -0.26$, $P = 0.14$). There was no association between the 24-h RQ and either the %EE ($P = 0.14$) or the absolute 24-h EE ($P = 0.23$) during FST. Neither the EE response to low-protein overfeeding diet nor the response to the high-carbohydrate overfeeding diet was correlated with the FST RQ. In a full model including all observed associations with weight change, only the percentage changes in 24-h EE during low-protein overfeeding ($\beta = -0.46$, $P = 0.002$) and high-carbohydrate overfeeding ($\beta = 0.39$, $P = 0.006$) remained independent predictors of weight change at 6 months. All results were similar if the data set was limited to men only. All longitudinal results were similar, and often slightly stronger, if the percentage weight change from baseline weight was substituted for the absolute weight change (data not shown).

**DISCUSSION**

Our results confirm that humans have the ability to respond to overfeeding and FST with an increase and a decrease in EE, respectively, and that these responses are directly correlated. At baseline, body adiposity was inversely related to the EE response to overfeeding. Individually, four variables related to FST and overfeeding, including a greater decrease in EE with FST, a smaller response to low-protein overfeeding, a greater EE response to high-carbohydrate overfeeding, and a lower FST RQ, were associated with weight gain at 6 months in free-living adults who were eating an ad libitum diet. However, only two independent phenotypes associated with future weight gain emerged, including a more energy-conserving response to low-protein feeding, during both calorie deficit and caloric excess, and separately, a larger EE response to high-carbohydrate overfeeding.

It is well recognized that EE increases with overfeeding and decreases with FST (5,6,17). In a prior cross-sectional study (5), these responses to overfeeding and FST were correlated within individuals, implying the possibility of
“thrifty” and “spendthrift” phenotypes within the population. Recent work from our group (18) found that obese individuals with a more thrifty phenotype, defined by the %EE response to FST, lost less weight in a carefully controlled inpatient weight loss study with 6 weeks of 50% caloric restriction. We have now confirmed that these %EE responses are correlated, and have shown that it is not so much the response to caloric restriction but, rather, the response to protein restriction that defines a thrifty phenotype. Consistent with the finding that more thrifty individuals lose less weight during caloric restriction (18), we observed that free-living individuals with a thrifty phenotype are more likely to gain weight over time. Contrary to expectations, a greater EE response to overconsuming large amounts of carbohydrates, an effect that might be expected to attenuate weight change, was associated with weight gain. Our study differs from many prior studies that have assessed the impact of long-term underfeeding (19–22) or overfeeding (8,17,23–25) with specific diets, in that we were assessing differences in baseline physiology and how such interindividual differences might interact with typical dietary patterns to influence weight change.

It is known from studies such as the Minnesota experiment and the Biosphere 2 project that prolonged energy restriction leads to adaptive reductions in EE (5,19–21). Of note, the diets in both of these studies also had a relatively low proportion of protein (<12%). A more recent study investigating the effects of long-term overconsumption of low-, normal-, and high-protein diets (8) found that FM gain was similar in all three groups, although low-protein diets led to smaller changes in overall weight due to differences in FFM. In our study, both a larger reduction in EE with FST and a smaller EE response
to the low-protein diet despite caloric excess were associated with future weight gain. These responses were correlated, and, in a multivariate model, only the low-protein response remained associated with weight change, indicating a potential similar underlying physiology. A candidate pathway that might explain these findings is the hepatic response to amino acid deprivation that leads to secretion of fibroblast growth factor 21 (FGF21) (26,27). Although FGF21 was originally reported to increase with FST (28–30), a recent study (27) has demonstrated that it is protein restriction, not caloric restriction, that induces increases in circulating FGF21 levels in rodents and in humans. This study (27) also found that FGF21 is required for the EE response to low protein. We observed that the low-protein diet led to the smallest increases, and even decreases, in EE with overfeeding. Other studies (17,24,27) have reported that long-term overfeeding is required for any increased, potentially adaptive, EE response to a low-protein diet. As the sustained intake of a low-protein diet would be unusual in modern society (31), our results may reflect that those individuals able to increase EE more quickly during even short periods of protein restriction are better able to prevent weight gain.

The increase in EE with overfeeding was greatest with the high-carbohydrate diet, but, surprisingly, a larger EE increase with this diet was associated with more weight gain. The underlying physiology behind the larger increase in EE with carbohydrate intake is unknown and may be related to genetic differences, alterations induced by prior dietary choices (i.e., a long-term high-carbohydrate diet prior to CRU admission), or a robust inflammatory response to carbohydrates (32). When subjects are fed an isocaloric high-carbohydrate diet for 2 weeks, those individuals who are more likely to store carbohydrates, rather than oxidize them, gain less FM over time (33), and we may be observing a similar phenotype. Alternatively, a high-carbohydrate meal has been reported to increase brown adipose tissue activity (10), which would lead to increased EE. As a higher EE during EB has been associated with greater subsequent ad libitum food intake (34,35), the availability of high-carbohydrate foods in a free-living condition may increase EE and subsequently drive further energy intake in the absence of dietary restraint.

FST RQ was no longer associated with weight change after accounting for the EE responses to low-protein and high-carbohydrate overfeeding. Thus, the initial simple correlation may be due to confounding or may indicate similar, overlapping physiologic mechanisms with the overfeeding results. The association of greater lipid oxidation with FST (i.e., a lower RQ) with future weight gain might suggest that a greater reliance on lipid stores during energy restriction is involved in body weight regulation. This finding may be consistent with a phenotype that preferentially oxidizes rather than stores carbohydrates (33), as the increased lipid oxidation during FST may reflect smaller amounts of glycogen stores. The previously reported associations of higher carbohydrate oxidation during EB with both subsequent increased food intake (36) as well as weight gain (36,37) are further evidence that phenotypic differences that indicate a preference to oxidize, rather than store, ingested carbohydrates are related to weight gain.

A limitation of our study is the lack of hormonal measures that might explain the underlying mechanisms of the EE changes. Nevertheless, prior results from a subset of these subjects (6) demonstrated that catecholamine responses were similar for both FST and the low-protein diet. Additional long-term follow-up is needed to determine whether the baseline measures of EE are associated with weight changes over longer periods of time (38–41). Subjects were asked to resume their previous lifestyle upon CRU discharge, and none of the subjects reported substantially changing their diet in the intervening period; however, formal assessments of diet or physical exercise during the follow-up period were not performed. This was purposeful, as we wanted to examine the relationship of baseline EE physiology with spontaneous short-term weight change under free-living, unencumbered conditions. In addition, it is possible that the level of physical fitness prior to CRU admission may have contributed to the EE response to overfeeding; however, all subjects were admitted to the CRU at the time of the initial assessment and had similar levels of physical activity.
activity during the inpatient stay. Further, adjusting for SPA in our analyses did not impact the results. Although the study includes a small proportion of lean women relative to women classified as obese, all results were similar if the data set was limited only to men. Even in this relatively small study group, we were able to identify subjects with differing phenotypes defined by their EE response to low-protein and high-carbohydrate overfeeding, and people with these phenotypes had substantially different changes in body weight at follow-up. Nevertheless, future studies with larger study populations are warranted to replicate and confirm our results.

In summary, we identified a number of metabolic phenotypes correlated with subsequent weight change that condensed into two independent phenotypes: a smaller EE response to low-protein intake and a greater EE increase with high-carbohydrate intake. Based on these results, it is reasonable to hypothesize that the observed interindividual variation in the EE response to protein restriction constitutes the long-sought, but previously unidentified, "thrift phenotype" that accounts, in part, for the observed interindividual variation in weight loss during similarly calorically restricted diets. Further, the interindividual variation in the EE responses to high-carbohydrate intake may account, in part, for the utility that some individuals find in eating a carbohydrate-restricted diet to limit weight gain. To conclude, an increased understanding of the phenotypic differences between people in response to overeating or underfeeding may lead to new strategies to prevent weight gain.

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