Intermittent fasting and mental and physical fatigue in obese and non-obese rats

Paige Niepoetter¹, Carolyn Butts-Wilmsmeyer¹², Chaya Gopalan³⁴*

¹ Department of Biological Sciences, Southern Illinois University Edwardsville, Edwardsville, IL, United States of America, ² Center for Predictive Analytics, Southern Illinois University Edwardsville, Edwardsville, IL, United States of America, ³ Department of Applied Health, Southern Illinois University Edwardsville, Edwardsville, IL, United States of America, ⁴ Department of Nurse Anesthesiology, Southern Illinois University Edwardsville, Edwardsville, IL, United States of America

* cgopala@siue.edu

Abstract

Intermittent fasting (IF) is an alternating pattern of restricting eating. This study evaluated mental and physical fatigue secondary to IF (daily 18-hour fast, 7-days-a-week) in the high-fat diet (HFD)-induced male obese Sprague Dawley rats. Fifty-four rats were randomly assigned to a HFD (n = 28) or a standard diet (SD; n = 26). After six weeks, the HFD rats were divided into one of four groups: obese HFD ad libitum (OB-HFD-AL), obese HFD-IF (OB-HFD-IF), obese SD-AL (OB-SD-AL), and obese SD-IF (OB-SD-IF). Similarly, non-obese controls were grouped into HFD-AL (C-HFD-AL), non-obese HFD-IF (C-HFD-IF), non-obese SD-AL (C-SD-AL), and non-obese SD-IF (C-SD-IF). After 2 weeks of IF, mental and physical fatigue were measured using open field (OF) and novel object recognition (NOR) tests. Rats on IF gained weight at a slower pace (p < 0.05) and had lower glucose levels (p < 0.01) compared to the AL group. In non-obese rats, ketone levels were higher in the IF-HFD group than IF-SD (p < 0.05) and AL-SD (p < 0.01) animals. Obese rats exhibited elevated blood ketone levels in IF-SD conditions versus AL-SD rats (p < 0.01). AL-HFD rats had higher ketone levels than AL-SD animals in both obese and non-obese groups (p < 0.05). In conclusion, rats with higher blood ketone levels, whether they were on IF or AL, traveled a greater distance during OF suggesting a lack of physical fatigue. There was no significant difference between IF and AL during NOR indicating a lack of mental fatigue. Thus, IF results in reduced body weight and blood glucose levels but does not induce physical or mental fatigue.

Introduction

Obesity is associated with cardiovascular disease, type 2 diabetes, several types of cancers, mental illnesses, cognition impairment, and chronic neurological degenerative conditions such as Alzheimer’s disease and dementia [1–3]. For example, Simon et al. reported that obesity is associated with an approximately 25% increase in mood and anxiety disorders [4]. A meta-analysis of 17 studies found that individuals with obesity were 1.26 times more likely to experience depression compared to non-obese individuals [2]. Multiple lifestyle modifications have
been studied to combat the adverse effects of obesity, including dietary measures such as intermittent fasting (IF) [5, 6].

IF has been practiced for many decades in several different religions. Followers of Islam engage in IF during the holy month of Ramadan, where fasting takes place every day of the month ranging between 11–22 hour intervals [7]. While Ramadan fasting is one example of time-restricted feeding, other IF regimens include alternate-day fasting switching between consuming no calories and regular food intake every other day. Some follow modified fasting regimens that consume 20% of the average daily caloric intake on fasting days, such as in the case with the popular 5:2 diet, where fasting occurs 2 days a week nonconsecutively, with the other five days consisting of regular food intake. IF, irrespective of the strategy used, is shown to produce beneficial effects, including increased insulin sensitivity, weight loss, and reductions in plasma cholesterol levels [7–9].

The IF regimen is expected to induce certain cellular changes such as depleted levels of glycogen stores and increased blood ketones as a result of higher rate of lipolysis [10, 11]. Weight loss and baseline blood glucose levels are often the byproducts of increased energy expenditure through lipid metabolism [10, 12]. The beneficial effects of IF are demonstrated in a study by Spezani et al. (2020). Twelve-week-old C57BL/6J mice were fed either a control diet (C; 10% kcal fat), a HFD (50% kcal fat), or a high fructose diet (HFru; 50% kcal fructose) for eight weeks. After these eight weeks, half of the rats started on an alternate day IF regimen (IF; 24 h fed, 24 h fasting) while still maintaining their diet type for four weeks. All groups tested benefited from IF with improved glycemic control, reduced insulin resistance, and weight loss [13]. These findings were reiterated by Gotthardt and Bello (2017) using alternate day IF (IMF) in adult obese male C57BL/6 mice [14].

Obesity is linked to low endurance and increased fatigue in humans [15, 16]. IF may pose a solution to obesity-related fatigue, though assessments of fatigue on this regimen are scarce and the results are often mixed [5, 17, 18]. Self-assessments of fatigue in those participating in Ramadan fasting have reported increased fatigue while on the regimen. A study by Chaouachi et al. (2009) with male judo athletes, during Ramadan, reported increased fatigue, though physical performance was relatively unchanged [18]. A group of nurses completed a similar fatigue self-assessment and reported increased fatigue while fasting, with fatigue increasing as subjective health scores declined [19]. Other studies have shown improvement in subject fatigue scores during times of IF. A study by Bowen et al. (2018) used self-assessments to measure fatigue in obese individuals partaking in a high-protein diet with or without alternate-day fasting. They reported that fatigue decreased overall, but those also participating in alternate-day fasting had even greater levels of improvement [17]. A meta-analysis performed by Abadia, Daab, and Bouzid related physical measures to Ramadan fasting and found that power and sprinting measures were reduced after Ramadan, though aerobic performance fatigue index scores were not influenced [20]. Overall, such mixed results produced by human studies related to IF and fatigue demonstrate the knowledge deficit that must be filled in order to positively state that IF can mitigate the fatigue associated with obesity. It has been suggested that the IF regimen may mitigate physical fatigue by providing another fuel source in ketones such as beta-hydroxybutyrate (BHB), which can be used once carbohydrates have been depleted [21, 22]. Moreover, elevated ketone levels are associated with greater time to fatigue in rodents exposed to forced swim tests and forced walking models [23, 24], though additional literature in rodent models is scarce.

Obesity has also been associated with impairments in cognition, the higher-order process of learning, memory formation, and retrieving information through thought, experience, and the senses [4, 25–27]. Considering the many benefits of IF, some studies have investigated its potential for improving cognitive functions as well, though this information is limited [8, 28,
A study by Li et al. (2013) exposed 7-week-old mice to control, HFD, or alternate-day fasting with SD conditions over 11 months. After this time, the mice underwent a Barnes maze test, which measures spatial working memory (SWM) by recording the amount of time it takes to enter the correct target box that was previously introduced during a habituation phase. The mice in the fasting condition exhibited better memory and cognition during the Barnes maze test than mice in the other conditions [30]. Many studies attribute this improvement in cognition to caloric restriction as the window of food intake is limited [31]. Geng et al. (2007) investigated the cognitive effects of a 60% calorie-restrictive diet in 18-month-old versus ad libitum rats over six months. At the end of this study, a Morris water maze test was performed, where SWM was measured by how long it took for a rat to find an escape platform that they were previously introduced to and found that rats on the calorie-restrictive diet outperformed the ad libitum group in the Morris water maze test [32]. These studies suggest that caloric restriction via IF is responsible for the beneficial cognitive effects experienced, though it is likely that deeper cellular mechanisms are at play. For example, elevated ketone levels may provide alternative energy sources for cognitive functions [33]. A study by Murray et al. (2016) found improved memory during radial arm maze testing in rats fed a high ketone ester diet which resulted in higher plasma beta-hydroxybutyrate levels after 36 days of this dietary intervention [24]. Our previous study using HFD-induced obese rats had increased ketone levels which were associated with greater time spent with novel versus familiar objects in NOR testing, indicating that ketones protected against memory deficits [34]. Ketones may exert these benefits in a number of ways. It is thought that the neuroprotective qualities of ketones stem from increasing energy production by promoting mitochondrial reproduction in neurons and reducing neuronal apoptosis [22, 35]. It is also possible that the metabolic switch from glucose to ketone utilization could increase the expression of brain-derived neurotrophic factor (BDNF), which has been shown to improve cognition by increasing neurogenesis, synaptogenesis, and preventing apoptosis [36–38].

Since the connection between IF and fatigue is not well understood in rodents and is often subjective in humans, additional exploration is needed to better understand the benefits IF has to offer, as well as potential disadvantages [5]. The goal of this study was to utilize behavioral testing after exposure to IF in obese and non-obese rats fed either a SD or HFD in the hopes of gaining a better understanding of the connection between ketones and mental and physical fatigue. It was hypothesized that IF would protect against mental and physical fatigue via increased ketone (BHB) levels [39].

Materials and methods

Animals

Fifty-four male Sprague Dawley rats at 7 weeks of age were received from Envigo Labs, Indianapolis, IN. They were housed individually under controlled laboratory conditions (12-hour light/dark cycle with lights on at 7:00 PM at a room temperature of 20.0–22.2˚C) in solid-bottom cages with aspen chip bedding. All protocols described were approved by the Southern Illinois University Edwardsville Institutional Animal Care and Use Committee (040618-CG2).

Diet

Upon arrival, animals were randomly placed into one of the 2 diet groups. The first 6 weeks of the study consisted of inducing obesity by feeding one group of rats a HFD (n = 28; formula D12492 from Research Diets Inc) while the rest received a SD (n = 26; Mazuri rat chow 5663; Table 1).
Metabolic testing

Capillary blood sampling was used to obtain overnight fasting glucose (mg/dL) and BHB (mmol/L) levels between 7:00 and 9:00 am on day 6 each week. Blood samples for this testing were obtained by pricking the rats’ tail veins using 26-gauge lancets. Results were obtained immediately using a Keto-Mojo (Napa Valley, CA) glucose and ketone meter (model TD-4279). Previous studies have used capillary sampling and ketone test strips to measure circulating levels of BHB [40, 41].

Intermittent fasting

Once diet-induced obesity (DIO) was achieved and behavioral tests were completed, the animals that were fed HFD were referred to as obese (OB), and those that received SD became non-obese controls (C). Both the HFD and the SD groups were divided into four subgroups each (Fig 1; Table 2): obese HFD ad libitum (OB-HFD-AL), obese HFD-IF (OB-HFD-IF), obese SD-AL (OB-SD-AL), obese SD-IF (OB-SD-IF), non-obese HFD-AL (C-HFD-AL), non-obese HFD-IF (C-HFD-IF), non-obese SD-AL (C-SD-AL), and non-obese SD-IF (C-SD-IF). Animals in the IF groups were fasted for 18 hours per day, 7 days a week. Animals were on IF for 2 weeks before behavioral testing was initiated. After 2 weeks on IF, behavioral tests were repeated to evaluate physical and mental fatigue.

Behavioral testing

Baseline behavioral testing occurred during the last 2 weeks of inducing obesity. The common definition of physical fatigue is the inability of muscles to maintain a needed level of power during and after physical activity, which is measured in a variety of ways in humans, but is

| Table 1. Macronutrient composition (Niepoetter et al. 2021). |
|------------------|---------|---------|
|                | HFD    | SD      |
| Fat (kcal)      | 60%    | 17%     |
| Carbohydrate (kcal) | 20%    | 56%     |
| Protein (kcal)  | 20%    | 27%     |
| Energy Density (kcal/g) | 5.21   | 3.41    |
| Fat Source      | lard, soybean oil | flaxseed oil, polyunsaturated fatty acids |

HFD: high-fat diet; SD: standard diet; kcal: kilocalories; kcal/g: kilocalories per gram of food.

https://doi.org/10.1371/journal.pone.0275684.t001

Fig 1. Illustrates the groups and their treatments. High-fat diet (HFD); Standard diet (SD); Intermittent fasting (IF); Ad libitum (AL).

https://doi.org/10.1371/journal.pone.0275684.g001
commonly assessed by OF in rodents [16, 29]. During OF, movements such as the number of line crossings, distance/time moving, and motion freezing are recorded [29]. OF measures exploratory behavior and movement to indicate the physical health of rodents. The OF apparatus utilized a 100 cm x 100 cm opaque plexiglass arena with central and peripheral zones. The animal was placed into the arena to explore freely for 6 minutes. The movements were tracked using ANY-maze video tracking system (Stoelting, Wood Dale, Illinois). The total distance traveled was recorded to examine the locomotor activity of the animal. OF was performed for eight consecutive days in the dark under red light conditions during their night cycle. Mental fatigue, the impairment of cognitive performance due to the reduced mental alertness or the feeling of absence of energy, is evaluated by NOR testing in rodents [42, 43]. NOR testing measuring recognition memory provides an indirect assessment of cognition [42]. After the completion of OF testing over 8 consecutive days, the animals underwent a NOR study. Five objects of different shapes, colors, and dimensions were utilized for this study. During the familiarization phase, one object was placed in the same opaque plexiglass arena as used in OF, and the rats were then allowed 5 minutes to investigate the object within the arena. The familiarization phase was immediately followed by the first IEI called the 0-hour test. The remaining IEIs occurred at 24-hours (one day), 72 hours (3 days), and 168-hours (7 days) after the initial 0-hour test. The duration spent by the rats investigating the new object compared to the familiar object translates to the animal's recognition and provides a numerical measurement for memory [34, 42].

### Statistical analysis

**A priori power analysis and data quality assurance.** G*Power (version 3.1.9.4) was used to calculate the sample size needed to obtain a power of at least 0.8 at an α = 0.05. Power calculations were based on a moderate correlation among repeated measurements (r = 0.5) and a moderate effect size (η² = 0.15). If rats did not gain at least 10% on the HFD relative to the mean of the SD control, these animals were excluded from the analysis. As such, the initial calculated sample size of 44 individuals was increased by a factor of 22.7% (10 individuals) to buffer against non-responders.

**ANOVA of individual variables.** Measurements of body weight, blood glucose and ketones as well as the behavioral data analysis during DIO is previously published [34]. In this study, body weight, blood glucose, and blood ketones were measured once per week and the averages of each of these variables analyzed using a repeated measures analysis of variance (ANOVA) in PROC MIXED of SAS (version 9.4). Behavioral measurements (i.e., total distance traveled and time spent with novel versus familiar objects) were measured on a daily basis.
timescale after 2 weeks of IF treatment. Specifically, distance traveled was measured daily for 8
days and the variables in NOR testing were collected at days 31 (0), 32 (1), 34 (3), and 38 (7).
As such, a repeated measures ANOVA was also used, but the frequency of the repeated mea-
surement differed between these and the weekly model. Additionally, the novel preference was
first calculated with the total amount of time spent with the novel object, then as the natural
log of the ratio of time spent with the novel object to the time spent with the familiar object. A
correlation analysis of ketone levels and behavioral measurements during OF and NOR was
performed using the cor.test function in R (version 4.0.4).

Results

Physiological measurements

Both non-obese \( (p<0.01; \text{Fig 2}) \) and obese \( (p<0.05; \text{Fig 3}) \) rats on IF weighed an average of 22g
less than AL rats. Diet type had a significant effect on glucose levels in both non-obese and
Obese rats, with HFD exhibiting higher glucose levels than SD-fed rats. Glucose levels were significantly lower in the non-obese rats undergoing IF, when compared to AL rats (p<0.01; Fig 2). Obese rats on IF also had lower glucose levels compared to AL rats, reaching significant levels at weeks 2 and 3 (p<0.01; Fig 3).

In non-obese rats, ketone levels were higher in the IF-HFD group compared to the IF-SD (p<0.05) and AL-HFD (p<0.01) groups (Table 3). Obese rats exhibited higher blood ketone levels in IF-SD conditions versus AL-SD rats (p<0.01; Table 4). In both obese and non-obese groups, AL-HFD animals had higher ketone levels than AL-SD rats (p<0.05; Tables 3 and 4).

### Behavioral data

Although there was no significant difference in the time spent with novel versus familiar objects in NOR testing or in the OF measurements between IF and AL groups in non-obese or obese rats (Tables 5 and 6; Fig 4), higher blood ketone levels correlated with greater distance traveled in both IF and AL groups (p<0.05; Table 7).

### Discussion

This study utilized a DIO model to examine whether IF would result in mental and physical fatigue in obese and non-obese rats. It was found that IF, regardless of diet, led to decreased weight gain and lower blood glucose levels [13, 44, 45]. These findings are supported by Bhoumik et al. (2020), which used Wistar rats to evaluate the effects of a time-restricted feeding model of IF (18-hour fast, 6-hour fed) or alternate day fasting (24 hours fed, 24 hours fasted) compared to AL controls over a 4-week time period. At the end of the study, it was found that both time-restricted feeding and alternate-day fasting resulted in decreased body weight and lower fasting blood glucose levels compared to AL rats [44]. In a study by Spezani et al. (2020),

### Table 3. Blood ketone levels in non-obese control rats.

| Group    | Experimental Regimen | Mean ± Standard Error |
|----------|----------------------|-----------------------|
| Non-obese| IF-HFD               | 1.2± 0.1†             |
|          | AL-HFD               | 1.1± 0.1†             |
|          | IF-SD                | 0.8± 0.1              |
|          | AL-SD                | 0.7± 0.1              |

* Significant difference of p<0.05 compared to IF-SD
† Significant difference of p<0.01 compared to AL-SD
‡ Significant difference of p<0.05 compared to AL-SD; Intermittent fasting (IF); ad libitum (AL); high-fat diet (HFD); standard diet (SD).

https://doi.org/10.1371/journal.pone.0275684.t003

### Table 4. Blood ketone levels in obese rats.

| Group | Experimental Regimen | Mean ± Standard Error |
|-------|----------------------|-----------------------|
| Obese | IF-HFD               | 0.5± 0.1              |
|       | AL-HFD               | 0.8± 0.1†             |
|       | IF-SD                | 0.8± 0.1†             |
|       | AL-SD                | 0.3± 0.2              |

‡ = Significant difference of p<0.05 compared to AL-SD
† = Significant difference of p<0.01 compared to AL-SD; Intermittent fasting (IF); ad libitum (AL); high-fat diet (HFD); standard diet (SD).

https://doi.org/10.1371/journal.pone.0275684.t004
male C57BL/6 mice were exposed to IF (alternating between 24-hour access to food and 24-hour without access to food) for a period of 4 weeks while being fed either a standard (10% kcal fat), high-fat (50% kcal fat), or high-sucrose (50% kcal sucrose) diet. After 4 weeks on the IF regimen with these diets, all the mice exhibited weight loss and lower fasting glucose levels [13].

Weight loss and lower glucose levels are often associated with the metabolic switch that occurs during an IF regimen, which results in an increase in BHB levels due to lipid metabolism [10, 12]. In this study, obese rats fed SD on IF had higher ketone levels than their AL-SD counterparts. The group that exhibited the greatest difference in blood ketone levels were the C-AL-HFD and C-IF-HFD groups. In a study by Dedaul et al. (2019) using non-obese male C57BL/6 mice that were fed HFD fasted for 8 hours a day for 4 days or fed AL, IF alone increased beta-oxidation but HFD paired with IF further increased this process, thus resulting in higher ketone levels [46]. It was shown that mice fed HFD-IF had greater metabolic flexibility as evidenced by the increase in phosphorylation of lipid metabolism regulators and greater ability to activate lipolysis in white adipose tissue [46].

While differences in ketone levels were not significantly different between all of the IF groups compared to AL groups (Fig 1), higher ketone levels, whether induced by diet or the IF regimen, were correlated with increased distance traveled during OF, indicating a resistance to physical fatigue. Ketones provide an alternative fuel for oxidative phosphorylation and makes oxidation a preferential process, which minimizes glycolysis [47]. Increased ketone levels have

Table 5. Open field measurements in Non-obese control rats.

|                          | Non-obese Rats |                     |
|--------------------------|----------------|---------------------|
|                          | AL             | IF                  |
|                          | Mean ± Standard Error | Mean ± Standard Error |
| Distance                 | 4.2± 0.2       | 4.0± 0.2            |
| Line Crossings           | 240.6 ± 11.5   | 224.9 ± 8.0         |
| Mean Speed               | 0.09 ± 0.01    | 0.08 ± 0.01         |
| Middle Zone Distance Traveled | 0.8 ± 0.1     | 0.7 ± 0.05          |
| Time Spent in Middle Zone| 41.5 ± 4.0     | 41.8 ± 3.3          |
| Wall Zone Distance Traveled | 2.9 ± 0.2     | 2.7 ± 0.1           |
| Time Spent in Wall Zone  | 291.1 ± 4.9    | 292 ± 4.9           |

Ad libitum (AL); Intermittent fasting (IF).

https://doi.org/10.1371/journal.pone.0275684.t005

Table 6. Open field measurements in obese rats.

|                          | Obese Rats |                     |
|--------------------------|------------|---------------------|
|                          | AL         | IF                  |
|                          | Mean ± Standard Error | Mean ± Standard Error |
| Distance                 | 3.6 ± 0.13 | 3.7 ± 0.15          |
| Line Crossings           | 228.9 ± 19.5 | 222.6 ± 5.8       |
| Mean Speed               | 0.06 ± 0.01 | 0.06 ± 0.01         |
| Middle Zone Distance Traveled | 0.7 ± 0.1    | 0.7 ± 0.06          |
| Time Spent in Middle Zone| 40.5 ± 6.2  | 41.8 ± 4.6          |
| Wall Zone Distance Traveled | 2.7 ± 0.2    | 2.6 ± 0.1           |
| Time Spent in Wall Zone  | 291.2 ± 8   | 287.7 ± 5.8         |

Ad libitum (AL); Intermittent fasting (IF).

https://doi.org/10.1371/journal.pone.0275684.t006
been associated with improved physical performance and decreased fatigue in previous studies [23, 24]. A study by Nozawa et al. (2009) demonstrated the benefits of increased blood ketone levels in combating physical fatigue with mice exposed to bonito extract, an agent that increases ketone levels. Mice exposed to bonito extract were put through a forced swimming test and forced walking model to test for physical fatigue. Mice on bonito extract exhibited increased ketone levels as expected and resistance to physical fatigue [23]. This finding is further supported by a study by Murray et al. (2016) where rats fed a 30% ketone diet produced higher ketone levels and ran 32% further than control rats during a treadmill walking test [24].

Similar to OF testing results, the NOR testing results showed that IF had no negative impact on recognition memory which is an indirect measure of cognition. While many studies have reported improved cognition with IF regimens, these studies utilized longer duration of IF [30, 48, 49]. A study by Elesawy et al. (2021) saw improvements in cognition via elevated plus maze testing after 12 weeks of IF (16-hour daily fast). It was reported that IF rats had an increase in BDNF and neurotrophin-3, which they contributed as the factors improving cognition [49]. A study by Anson et al. (2003) utilized an alternate day fasting model in C57BL/6 male mice over a period of 20 weeks and measured IGF-1 signaling as evidence of neuroprotective qualities. It was found that IF rats had higher levels of IGF-1 signaling compared to AL groups, suggesting

---

**Table 7. Ketone correlation matrix including all rat groups.**

| Ketone (mmol/L) | Distance Traveled (m) | Time with A | Time with X |
|-----------------|-----------------------|-------------|-------------|
| Correlation     | 0.215                 | -0.089      | -0.120      |
| p-value         | 0.026*                | 0.546       | 0.415       |

* = p<0.05. Time with A: time with familiar object; Time with X: time with novel object.

---

https://doi.org/10.1371/journal.pone.0275684.t007
that IF may have a beneficial effect on cognition [48]. An improvement in cognition was also demonstrated in a study by Li et al. (2013). After exposing 7-week-old mice to control, HFD, or alternate-day fasting with SD conditions over 11 months, the animals underwent a Barnes maze test to measure spatial working memory by recording the amount of time it took to enter the correct target box that was previously introduced during a habituation phase. The mice in the fasting condition outperformed mice in the other groups [30].

One limitation of this study was the shortened duration of exposure to IF. Though 2 weeks of exposure produced noticeable differences in body weight and glucose levels, it is possible that extending the duration of IF could enhance the results of behavioral tests as well as ketone levels. The IF regimen in our study did not cause a significant increase in the ketone levels in the OB-IF-HFD versus OB-AL-HFD group or the C-IF-SD versus C-AL-SD groups, potentially due to the shorter duration of IF. Other studies have shown increases in ketone levels when exposing rodents to IF over a long-term duration [48, 50]. A study by Anson et al. (2003) utilized an alternate day fasting regimen in male C57BL/6 mice for a period of 20 weeks. By the end of the study, these mice exhibited decreased glucose levels and increased plasma ketone levels while the body weight was maintained throughout the study [48]. These findings were reiterated in a study by Park et al. (2020) in an 8-week study of rats fed either a ketogenic diet, 30% HFD, IF (24 hours fed, 24 hours fasted), high carbohydrate, or control diet. Rats on IF had higher plasma ketone levels and decreased body weight, but did not exhibit a difference in fasting glucose levels [50]. Additionally, measuring food consumption and calculating caloric intake could be used to further explain the observations from this study. Future studies should include food intake measurements and an extended timeline to overcome these limitations.

In conclusion, this study validates the use of IF for improved fasting glucose levels and decreased weight gain irrespective of the nature of the diet in both non-obese and obese groups. IF for 2 weeks does not contribute to mental or physical fatigue but longer duration may offer more benefits. Furthermore, increased ketone levels were correlated with increased physical activity, suggesting a protective role of ketones against physical fatigue.

Supporting information
S1 File. (RTF)
S1 Graphical abstract. (TXT)

Author Contributions
Conceptualization: Chaya Gopalan.
Data curation: Carolyn Butts-Wilmsmeyer, Chaya Gopalan.
Formal analysis: Carolyn Butts-Wilmsmeyer.
Funding acquisition: Chaya Gopalan.
Investigation: Paige Niepoetter, Chaya Gopalan.
Methodology: Paige Niepoetter, Chaya Gopalan.
Project administration: Chaya Gopalan.
Resources: Chaya Gopalan.
Software: Chaya Gopalan.

Supervision: Chaya Gopalan.

Validation: Chaya Gopalan.

Visualization: Chaya Gopalan.

Writing – original draft: Paige Niepoetter, Chaya Gopalan.

Writing – review & editing: Paige Niepoetter, Chaya Gopalan.

References

1. Apovian CM. Obesity: Definition, Comorbidities, Causes, and Burden. THE AMERICAN JOURNAL OF MANAGED CARE. 2016; 22(7):10.

2. de Wit L, Luppino F, van Straten A, Penninx B, Zitman F, Cuipers P. Depression and obesity: A meta-analysis of community-based studies. Psychiatry Research. 2010 Jul; 178(2):230–5.

3. Solas M, Milagro FI, Ramírez MJ, Martínez JA. Inflammation and gut-brain axis link obesity to cognitive dysfunction: plausible pharmacological interventions. Current Opinion in Pharmacology. 2017 Dec; 37:87–92. https://doi.org/10.1016/j.coph.2017.10.005 PMID: 29107872

4. Simon GE, Von Korff M, Saunders K, Miglioretti DL, Crane PK, van Belle G, et al. Association Between Obesity and Psychiatric Disorders in the US Adult Population. Arch Gen Psychiatry. 2006 Jul 1; 63(7):824. https://doi.org/10.1001/archpsyc.63.7.824 PMID: 16818872

5. Halpern B, Mendes TB. Intermittent fasting for obesity and related disorders: unveiling myths, facts, and presumptions. Archives of Endocrinology and Metabolism [Internet]. 2021 Jan 18 [cited 2021 Oct 8]; Available from: https://www.scielo.br/scielo.php?script=sci_arttext&pid=S2359-39972021005001204&lng=en&nrm=iso

6. Harris L, Hamilton S, Azevedo LB, Olajide J, De Brún C, Waller G, et al. Intermittent fasting interventions for treatment of overweight and obesity in adults: a systematic review and meta-analysis. JBI Database of Systematic Reviews and Implementation Reports. 2018 Feb; 16(2):507–47. https://doi.org/10.11124/JBRIR-2016-003248 PMID: 29419624

7. Patterson RE, Laughlin GA, LaCroix AZ, Hartman SJ, Natarajan L, Senger CM, et al. Intermittent Fasting and Human Metabolic Health. Journal of the Academy of Nutrition and Dietetics. 2015 Aug; 115(8):1203–12. https://doi.org/10.1016/j.jand.2015.02.018 PMID: 25857868

8. Moro T, Tinsley G, Bianco A, Marcolin G, Pacelli QF, Battaglia G, et al. Effects of eight weeks of time-restricted feeding (16/8) on basal metabolism, maximal strength, body composition, inflammation, and cardiovascular risk factors in resistance-trained males. J Transl Med. 2016 Dec; 14(1):290. https://doi.org/10.1186/s12967-016-1044-0 PMID: 27737674

9. Rynders CA, Thomas EA, Zaman A, Pan Z, Catenacci VA, Melanson EL. Effectiveness of Intermittent Fasting and Time-Restricted Feeding Compared to Continuous Energy Restriction for Weight Loss. Nutrients. 2019 Oct 14; 11(10):2442. https://doi.org/10.3390/nu11102442 PMID: 31614992

10. Anton SD, Moehl K, Donahoo WT, Marosi K, Lee SA, Mainous AG, et al. Flipping the Metabolic Switch: Understanding and Applying the Health Benefits of Fasting: Flipping the Metabolic Switch. Obesity. 2018 Feb; 26(2):254–68.

11. Rui L. Energy Metabolism in the Liver. In: Terjung R, editor. Comprehensive Physiology [Internet]. 1st ed. Wiley; 2014 [cited 2022 Feb 2]. p. 177–97. Available from: https://onlinelibrary.wiley.com/doi/10.1002/cphy.c130024

12. Dong TA, Sandesara PB, Dhindsa DS, Mehta A, Arneson LC, Dollar AL, et al. Intermittent Fasting: A Heart Healthy Dietary Pattern? The American Journal of Medicine. 2020 Aug; 133(8):901–7. https://doi.org/10.1016/j.amjmed.2020.03.030 PMID: 32330491

13. Spezani R, da Silva RR, Martins FF, de Souza Marinho T, Aguilá MB, Mandarim-de-Lacerda CA. Intermittent fasting, adipokines, insulin sensitivity, and hypothalamic neuropeptides in a dietary overload with high-fat or high-fructose diet in mice. The Journal of Nutritional Biochemistry. 2020 Sep; 83:108419. https://doi.org/10.1016/j.jnutbio.2020.108419 PMID: 32580132

14. Gotthardt JD, Bello NT. Meal pattern alterations associated with intermittent fasting for weight loss are normalized after high-fat diet re-feeding. Physiology & Behavior. 2017 May; 174:49–56. https://doi.org/10.1016/j.physbeh.2017.02.046 PMID: 28263771

15. Cavuoto LA, Nussbaum MA. The Influences of Obesity and Age on Functional Performance During Intermittent Upper Extremity Tasks. Journal of Occupational and Environmental Hygiene. 2014 Sep 2; 11(9):589–90. https://doi.org/10.1080/15459624.2014.887848 PMID: 24484265
16. Mehta RK. Impacts of obesity and stress on neuromuscular fatigue development and associated heart rate variability. International Journal of Obesity. 2015; 7. https://doi.org/10.1038/ijo.2014.127 PMID: 25042859

17. Bowen J, Brindal E, James-Martin G, Noakes M. Randomized Trial of a High Protein, Partial Meal Replacement Program with or without Alternate Day Fasting: Similar Effects on Weight Loss, Retention Status, Nutritional, Metabolic, and Behavioral Outcomes. Nutrients. 2018 Aug 23; 10(9):1145. https://doi.org/10.1519/JSC.0b013e3181bc17fc PMID: 19910805

18. Chauouachi A, Coutts AJ, Chamari K, Wong DP, Chauouachi M, Chatare M, et al. Effect of Ramadan Intermittent Fasting on Aerobic and Anaerobic Performance and Perception of Fatigue in Male Elite Judo Athletes. Journal of Strength and Conditioning Research. 2009 Dec; 23(9):2702–9. https://doi.org/10.1519/JSC.0b013e3181bc17fc PMID: 19910805

19. Ovayolü Ö, Ovayolü N, Taşan E. Does Ramadan Fasting Affect Fatigue in Nurses? Holistic Nursing Practice. 2016 Jul; 30(4):222–6. https://doi.org/10.1097/HNP.0000000000000141 PMID: 27223619

20. Abiad AE, Daab W, Bouzid MA. Effects of Ramadan Fasting on Physical Performance: A Systematic Review with Meta-analysis. Nutrients. 2018 Aug 23; 10(9):1145. https://doi.org/10.3390/nu10091145 PMID: 30142886

21. Chaouachi A, Coutts AJ, Chamari K, Wong DP, Chauouachi M, Chatare M, et al. Effect of Ramadan Intermittent Fasting on Aerobic and Anaerobic Performance and Perception of Fatigue in Male Elite Judo Athletes. Journal of Strength and Conditioning Research. 2009 Dec; 23(9):2702–9. https://doi.org/10.1519/JSC.0b013e3181bc17fc PMID: 19910805

22. Evans M, Cogan KE, Egan B. Metabolism of ketone bodies during exercise and training: physiological basis for exogenous supplementation: Ketone bodies and exercise. J Physiol. 2017 May 1; 595 (9):2857–71.

23. Maalouf M, Rho JM, Mattson MP. The neuroprotective properties of calorie restriction, the ketogenic diet, and ketone bodies. Brain Research Reviews. 2009 Mar; 59(2):293–315. https://doi.org/10.1016/j.brainresrev.2008.09.002 PMID: 18845187

24. Nozawa Y, Yamada K, Okabe Y, Ishizaki T, Kuroda M. The Anti-fatigue Effects of the Low-Molecular-Weight Fraction of Bonito Extract in Mice. Biological & Pharmaceutical Bulletin. 2009; 32(3):468–74. https://doi.org/10.1248/bpb.32.468 PMID: 19252297

25. Murray AJ, Knight NS, Cole MA, Cochlin LE, Carter E, Tchabane K, et al. Novel ketone diet enhances physical and cognitive performance. FASEB J. 2016 Dec; 30(12):4021–32. https://doi.org/10.1007/s40279-016-0077-0 PMID: 27528626

26. Buie JJ, Watson LS, Smith CJ, Sims-Robinson C. Obesity-related cognitive impairment: The role of endothelial dysfunction. Neurobiology of Disease. 2019 Dec; 132:104580. https://doi.org/10.1016/j.nbd.2019.104580 PMID: 31454547

27. Dye L, Boyle NB, Champ C, Lawton C. The relationship between obesity and cognitive health and decline. Proc Nutr Soc. 2017 Nov; 76(4):443–54. https://doi.org/10.1017/S0029665117002014 PMID: 28889822

28. Cifre M, Palou A, Oliver P. Cognitive impairment in metabolically-obese, normal-weight rats: identification of early biomarkers in peripheral blood mononuclear cells. Mol Neurodegeneration. 2018 Dec; 13(1):14. https://doi.org/10.1186/s13024-018-0246-6 PMID: 29566703

29. Barmosky AR. Intermittent fasting vs daily calorie restriction for type 2 diabetes prevention: a review of human findings. Translational Research. 2014; 164(4):10. https://doi.org/10.1016/j.trsl.2014.05.013 PMID: 24993615

30. Seibenhener ML, Wooten MC. Use of the Open Field Maze to Measure Locomotor and Anxiety-like Behavior in Mice. JoVE. 2015 Feb 6;(96):52434. https://doi.org/10.3791/52434 PMID: 25742564

31. Kim C, Pinto AM, Bordoli C, Buckner LP, Kaplan PC, del Arenal IM, et al. Energy Restriction Enhances Adult Hippocampal Neurogenesis-Associated Memory after Four Weeks in an Adult Human Population with Central Obesity; a Randomized Controlled Trial. Nutrients. 2020 Feb 28; 12(3):638.

32. Geng YQ, Guan JT, Xu MY, Xu XH, Fu YC. Behavioral Study of Calorie-restricted Rats from Early Old Age. In: 2007 29th Annual International Conference of the IEEE Engineering in Medicine and Biology Society [Internet] . Lyon, France: IEEE; 2007 [cited 2021 Oct 18]. p. 2393–5. Available from: http://ieeexplore.ieee.org/document/4352809/

33. Newman JC, Covarrubias AJ, Zhao M, Yu X, Gut P, Ng CP, et al. Ketogenic Diet Reduces Midlife Mortality and Improves Memory in Aging Mice. Cell Metabolism. 2017 Sep; 26(3):547–557.e6. https://doi.org/10.1016/j.cmet.2017.08.004 PMID: 28977458

34. Niepoetter P, Butts-Wilmsmeyer C, Kaviani S, Viernow C, Ruholl H, Gopalan C. Correlation between ketones and mental fatigue in high fat-induced obese and non-obese rats. Physiol Rep [Internet]. 2021 Jul [cited 2021 Nov 3];9(13). Available from: https://onlinelibrary.wiley.com/doi/ https://doi.org/10.14814/phy2.14930 PMID: 34197701
35. Cunnane SC, Courchesne-Loyer A, Vandenbergh C, St-Pierre V, Fortier M, Hennebelle M, et al. Can Ketones Help Rescue Brain Fuel Supply in Later Life? Implications for Cognitive Health during Aging and the Treatment of Alzheimer’s Disease. Front Mol Neurosci [Internet]. 2016 Jul 8 [cited 2021 Oct 18];9. Available from: http://journal.frontiersin.org/Article/https://doi.org/10.3389/fnmol.2016.00053 PMID: 27458340

36. Marosi K, Kim SW, Moehl K, Scheibye-Knudsen M, Cheng A, Cutler R, et al. 3-Hydroxybutyrate regulates energy metabolism and induces BDNF expression in cerebral cortical neurons. J Neurochem. 2016 Dec; 139(5):769–81. https://doi.org/10.1111/jnc.13868 PMID: 27739595

37. Lee M, Soya H. Effects of acute voluntary loaded wheel running on BDNF expression in the rat hippocampus. JENB. 2017 Dec 31; 21(4):52–7. https://doi.org/10.20463/jenb.2017.0034 PMID: 29370674

38. Sleiman SF, Henry J, Al-Haddad R, El Hayek L, Abou Haidar E, Stringer T, et al. Exercise promotes the expression of brain derived neurotrophic factor (BDNF) through the action of the ketone body β-hydroxybutyrate. eLife. 2016 Jun 2; 5:e15092. https://doi.org/10.7554/eLife.15092 PMID: 27253067

39. Mattson MP, Longo VD, Harvie M. Impact of intermittent fasting on health and disease processes. Ageing Research Reviews. 2017 Oct; 39:46–58. https://doi.org/10.1016/j.arr.2016.10.005 PMID: 28555095

40. Bentourkia M, Tremblay S, Pifferi F, Rousseau J, Cunnane S. PET study of 11C-acetacetate kinetics in rat brain during dietary treatments affecting ketosis. American Journal of Physiology-Endocrinology and Metabolism. 2009 Apr; 296(4):E796–801.

41. Brownlow ML, Jung SH, Moore RJ, Bechmann N, Jankord R. Nutritional Ketosis Affects Metabolism and Behavior in Sprague-Dawley Rats in Both Control and Chronic Stress Environments. Front Mol Neurosci. 2017 May 15; 10:129. https://doi.org/10.3389/fnmol.2017.00129 PMID: 28555095

42. Antunes M, Biala G. The novel object recognition memory: neurobiology, test procedure, and its modifications. Cogn Process. 2012 May; 13(2):93–110. https://doi.org/10.1007/s10339-011-0430-z PMID: 22160349

43. Boksem MAS, Tops M. Mental fatigue: Costs and benefits. Brain Research Reviews. 2008 Nov; 59 (1):125–39. https://doi.org/10.1016/j.brainresrev.2008.07.001 PMID: 18652844

44. Bhoumik S, Kumar R, Rizvi SI. Time restricted feeding provides a viable alternative to alternate day fasting when evaluated in terms of redox homeostasis in rats. Archives of Gerontology and Geriatrics. 2020 Nov; 91:104188. https://doi.org/10.1016/j.archger.2020.104188 PMID: 32717588

45. Hazzaa SM, Eldaim MAA, Fouda AA, Mohamed ASED, Soliman MM, Elgizawy EI. Intermittent Fasting Ameliorated High-Fat Diet-Induced Memory Impairment in Rats via Reducing Oxidative Stress and Glial Fibrillary Acidic Protein Expression in Brain. Nutrients. 2020 Dec 22; 13(1):10. https://doi.org/10.3390/nu13010010 PMID: 33375195

46. Dedual MA, Wueest S, Borsigova M, Konrad D. Intermittent fasting improves metabolic flexibility in short-term high-fat diet-fed mice. American Journal of Physiology-Endocrinology and Metabolism. 2019 Nov 1; 317(5):E773–82. https://doi.org/10.1152/ajpendo.2019.31503513

47. Cox PJ, Kirk T, Ashmore T, Willerton K, Evans R, Smith A, et al. Nutritional Ketosis Alters Fuel Preference and Thereby Endurance Performance in Athletes. Cell Metabolism. 2016 Aug; 24(2):256–68. https://doi.org/10.1016/j.cmet.2016.07.010 PMID: 27475046

48. Anson RM, Guo Z, de Cabo R, Iyun T, Rios M, Hagepanos A, et al. Intermittent fasting dissociates beneficial effects of dietary restriction on glucose metabolism and neuronal resistance to injury from calorie intake. Proceedings of the National Academy of Sciences. 2003 May 13; 100(10):6216–20. https://doi.org/10.1073/pnas.0357201100 PMID: 12724520

49. Elesawy BH, Raafat BM, Muqbalii AA, Abbas AM, Sakr HF. The Impact of Intermittent Fasting on Brain-Derived Neurotrophic Factor, Neurotrophin 3, and Rat Behavior in a Rat Model of Type 2 Diabetes Mellitus. Brain Sciences. 2021 Feb 15; 11(2):242. https://doi.org/10.3390/brainsci11020242 PMID: 33671898

50. Park S, Zhang T, Wu X, Yi Qiu J. Ketone production by ketogenic diet and by intermittent fasting has different effects on the gut microbiota and disease progression in an Alzheimer’s disease rat model. J Clin Biochem Nutr. 2020; 67(2):188–98. https://doi.org/10.3161/jcbn.19-87 PMID: 33041517