MiniReview

Weight Cycling and Cancer: Weighing the Evidence of Intermittent Caloric Restriction and Cancer Risk

Henry J. Thompson and Anne McTiernan

Abstract

Overweight and obese individuals frequently restrict caloric intake to lose weight. The resultant weight loss, however, typically is followed by an equal or greater weight gain, a phenomenon called weight cycling. Most attention to weight cycling has focused on identifying its detrimental effects, but preclinical experiments indicating that intermittent caloric restriction or fasting can reduce cancer risk have raised interest in potential benefits of weight cycling. Although hypothesized adverse effects of weight cycling on energy metabolism remain largely unsubstantiated, there is also a lack of epidemiologic evidence that intentional weight loss followed by regain of weight affects chronic-disease risk. In the limited studies of weight cycling and cancer, no independent effect on postmenopausal breast cancer but a modest enhancement of risk for renal cell carcinoma, endometrial cancer, and non-Hodgkin’s lymphoma have been reported. An effect of either intermittent caloric restriction or fasting in protecting against cancer is not supported by the majority of rodent carcinogenesis experiments. Collectively, the data argue against weight cycling and indicate that the objective of energy balance-based approaches to reduce cancer risk should be to strive to prevent adult weight gain and maintain body weight within the normal range defined by body mass index. Cancer Prev Res; 4(11); 1736–42. ©2011 AACR.

Introduction

The prevalence of overweight and obesity, as defined by body mass index (BMI) more than 24.9 (body weight in kg divided by height in m²), has increased at an epidemic rate over the last few decades (1), and an excess of adult body weight for height is associated with increased risk for a number of chronic diseases including certain types of cancer referred to as obesity-associated cancers (2–4). Rising obesity has stimulated increased efforts to lose weight (5, 6), and all approaches for inducing weight loss, in humans or animal models, involve reducing caloric intake relative to caloric need (7). Attempts to lose weight by any approach vary in their results, however, and the weight that is lost is frequently regained (8–10). This weight loss and regain in people indicates an intermittent dietary pattern that provides the link between intermittent caloric restriction in animal studies of energy balance and weight cycling in humans. Historically, repeated cycles of weight loss and regain have been called weight cycling, "yo-yo dieting," or, more generally, weight fluctuation, weight variability, or weight instability (11). Weight cycling is a complex behavior and remains ill-defined, thus making it difficult to study in human populations or to simulate in animal models (12). The several reported patterns of weight cycling include the typical pattern in obese individuals, which is weight loss and then regain in a repetitive cycle (Fig. 1A). Less commonly considered patterns include weight cycling among normal-weight individuals (Fig. 1B) and weight cycling in individuals with a BMI below the normal range (Fig. 1C). These patterns and their many conceivable variants illustrate the complexity of investigating the effects of weight cycling in human populations. Cutter and colleagues proposed that the key elements to consider in characterizing weight cycling are the amplitude of the cycles (the amount of weight gained or lost), the frequency of cycling (the number of cycles experienced), and the duration of the cycles (the timeframe over which cycles occur: days, weeks, months, years; ref. 12). Many other factors also can be considered, including the time during the lifecycle and stage during a disease process that weight cycling occurs (11).

The focus of this review is on the pattern of weight cycling illustrated in Fig. 1A and, furthermore, on the animal-model correlative of this pattern of human weight cycling. Over the last decade, a series of reports using genetically engineered or transplantable tumor models have indicated that intermittent caloric restriction or intermittent fasting, which represents an extreme form of caloric restriction, may exert beneficial effects against cancer, generating scientific interest in this...
Investigations of weight cycling have primarily been based on the assumption that it has negative health consequences. The majority of work has centered on the concern that weight cycling promotes the development of excessive weight gain. Specific hypotheses that have been evaluated include that weight cycling (i) impairs future weight loss and promotes future weight gain, (ii) increases food/caloric efficiency, (iii) increases relative, total, and/or central adiposity, (iv) increases preference for dietary fat, (v) decreases caloric expenditure, (vi) increases lipogenic enzyme activity, and (vii) promotes insulin resistance (22, 23). However, while stimulating considerable investigation, none of these effects has been substantiated in rodent experiments or clinical studies (9, 11, 23, 24).

The perception that weight cycling has negative consequences on human health has also been disseminated by reports that it increases morbidity and mortality (25–28). These early reports of adverse effects, however, have been shown to be due to a failure to account for intentionality of weight loss. For example, intentional weight loss was associated with a nearly 25% reduced all-cause mortality [hazard rate ratio (HRR) = 0.76; 95% CI: 0.60–0.97] compared with a one-third higher such risk for unintentional weight loss (HRR = 1.31; 95% CI: 1.01–1.70; ref. 29). As summarized in ref. 30, when unintentional weight loss studies are excluded, the majority of evidence fails to support an adverse effect of weight cycling on health, with the exception of gallbladder stones, which have a higher frequency in people who weight cycle (31, 32).

Cancer and Weight Cycling

Cancers of the breast (postmenopausal), colon, endometrium, esophagus, kidney (renal cell), and pancreas have been reported to be associated with obesity based on exhaustive reviews of the effects of body weight, adiposity, weight gain, and weight loss on the prevalence of cancer (2, 33), and recent evidence indicates that prostate cancer may be added to this list (3). Despite extensive investigations of factors related to energetics and cancer, the effects of weight cycling have been reported only for breast, endometrium, kidney, and lymphopoetic cancers (Table 1). In a large prospective study of weight change and breast cancer, no evidence was found to support an independent effect of weight cycling, defined as losing 20 pounds or more and gaining at least half of them back within a year (OR = 1.0; 95% CI: 0.9–1.1; ref. 34). Similarly, a case–control study of the effects of body size in relation to postmenopausal breast cancer found a nonsignificant increase in risk (OR = 2.11; 95% CI: 1.00–4.44) among women who exhibited a fluctuating pattern of body size, defined as body weight varying between more or equal to the median of the control group and less than the median, throughout adulthood (35). On the other hand, 2 studies in the Women’s Health Initiative do indicate an association of weight cycling with increased

Figure 1. Patterns of weight cycling. BMI < 18.5 is underweight, 18.5 to 24.9 is considered normal weight, 25 to 29.9 is considered overweight, and > 30 is considered obese. A, patterns of weight regulation involving obese individuals, either consistently in the obese range, individual O or transiently losing weight, regaining the weight, and repeating the cycle, individual T. B, individual X consistently maintains BMI in the normal range (18.5–24.9), with small weight fluctuations; individual Y engages in intermittent caloric restriction to induce weight loss, whereas individual Z periodically fails to regulate body weight and transiently attains a body weight above the normal range for BMI. C, individual U has a BMI that fluctuates entirely below the normal range. This type of underweight pattern can be associated with eating disorders, fad dietary practices, or natural disasters or wars leading to weight cycling that can reach down to starvation/famine levels of BMI.
risk. First, the incidence of renal cell carcinoma was increased in postmenopausal women who experienced intentional weight cycling (10 or more pounds) 10 or more times relative to stable-weight women [relative risk (RR) = 2.6; 95% CI: 1.6–4.2; \( P_{\text{trend}} = 0.0005 \)]. Second, women had an increased risk of developing non-Hodgkin’s lymphoma if they intentionally lost at least 50 pounds 3 or more times (HR = 1.97; 95% CI: 0.93–4.16; \( P_{\text{trend}} = 0.05 \)) or 20–49 pounds 3 or more times (HR = 1.55; 95% CI: 1.00–2.40; \( P_{\text{trend}} = 0.05 \)), but there was a reduced risk of non-Hodgkin’s lymphoma associated with smaller amounts of weight loss (10–19 pounds 3 or more times; HR = 0.78; 95% CI: 0.46–1.33; \( P_{\text{trend}} = 0.40 \), ref. 37). Similar nonstatistically significant, trends of altered risk were seen in association with multiple myeloma and leukemia.

In a case–control study, the risk for renal cell carcinoma also was elevated in female weight cyclers (OR = 2.31; 95% CI: 1.04–5.12, \( P_{\text{trend}} = 0.0005 \); ref. 38). A population-based case–control study found that a history of weight cycling (greater than 20-pound weight loss with at least half regained within a year) was associated with a modest increase in the risk of endometrial cancer after adjustment for BMI and other factors (OR = 1.27; 95% CI: 1.00–1.61; \( P_{\text{trend}} = 0.05 \); ref. 39).

### Rodent Carcinogenesis Studies

**Chemically induced models**

Kritchevsky and colleagues (40) investigated effects of weight cycling (induced by alternating periods of caloric restriction and ad libitum feeding) on the promotion phase of 7,12-dimethyl[\( \alpha \)]benzanthracene (DMBA)-induced mammary cancer, modeling common patterns of weight regulation via dieting (Table 2). They evaluated effects of 25% caloric restriction relative to ad libitum feeding at different times during the promotion/progression phase of mammary carcinogenesis and for cycles of 4 or 8 weeks. Body weight and tumor latency graphs from that study showed that retardation of weight gain suppressed mammary tumor development. When rats had free access to food and accelerated weight gain, however, the rate of tumor occurrence increased, particularly when short-term (4 weeks) caloric restriction was followed by an extended (12 weeks) period of ad libitum feeding and weight regain (50% tumor incidence, ad libitum alone, vs. 60% tumor incidence in caloric restriction followed by ad libitum feeding). Findings of Sylvester and colleagues (41) on caloric restriction during tumor promotion parallel those of Kritchevsky and colleagues, despite employing 50% caloric restriction relative to ad libitum feeding and imposing a restriction on all components of the diet instead of just on calories, a distinction described in detail in ref. 42. In another study, an effort to control the magnitude of weight cycling involved subjecting DMBA-treated obese adult Wistar rats to 4 cycles of 50% caloric restriction relative to ad libitum fed rats to achieve a 20% weight loss followed by weight regain (43). A reduced tumor incidence occurred in weight-cycled animals, but the difference was not statistically significant (mammary tumor incidence of 18% ad libitum vs. 9% weight-cycled).

Mehta and colleagues investigated cycles of 48 hours 40% caloric restriction followed by 48 hours during which rats were fed the same diet as age-matched, ad libitum–fed animals to prevent overeating (also termed rebound eating) relative to the ad libitum–fed control rats (44). This pattern of feeding and weight cycling virtually eliminated the
<table>
<thead>
<tr>
<th>Cancer site</th>
<th>Risk assessment</th>
<th>Amplitude</th>
<th>Frequency</th>
<th>Duration</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mammary, DMBA-induced, rat</td>
<td>RR = 1.20; 95% CI: 0.68–2.11, NS</td>
<td>25% CR</td>
<td>1 cycle</td>
<td>4 wk CR; 12 wk AL</td>
<td>(40)</td>
</tr>
<tr>
<td>Mammary, DMBA-induced, rat</td>
<td>RR = 0.90; 95% CI: 0.47–1.73, NS</td>
<td>25% CR</td>
<td>1 cycle</td>
<td>4 wk AL, 8 wk CR, 4 wk AL</td>
<td>(41)</td>
</tr>
<tr>
<td>Mammary, DMBA-induced, rat</td>
<td>RR = 0.74; 95% CI: 0.49–1.12, NS</td>
<td>50% CR</td>
<td>1 cycle</td>
<td>5 wk AL, 4 wk CR, 12 wk AL</td>
<td>(42)</td>
</tr>
<tr>
<td>Mammary, DMBA-induced, rat</td>
<td>RR = 0.50; 95% CI: 0.14–1.84, NS</td>
<td>50% CR</td>
<td>4 cycles</td>
<td>Loss of 20% body weight and then regained</td>
<td>(43)</td>
</tr>
<tr>
<td>Mammary, DMBA-induced, rat</td>
<td>RR = 0.89; 95% CI: 0.59–1.35, NS</td>
<td>40% CR</td>
<td>15 cycles</td>
<td>48 h AL, 48 h CR for 9 wk</td>
<td>(44)</td>
</tr>
<tr>
<td>Mammary, DMBA-induced, rat</td>
<td>RR = 0.90; 95% CI: 0.63–1.30, NS</td>
<td>40% CR</td>
<td>16 cycles</td>
<td>48 h AL, 48 h CR for 10 wk</td>
<td>(45)</td>
</tr>
<tr>
<td>Mammary, MNU-induced, rat</td>
<td>RR = 1.22; 95% CI: 0.89–1.68, NS</td>
<td>33% CR</td>
<td>4 cycles</td>
<td>1 wk CR, 3 wk AL</td>
<td>(46)</td>
</tr>
<tr>
<td>Mammary, MMTV-TGF-α (lep&lt;sup&gt;db/db&lt;/sup&gt;) transgenic mouse</td>
<td>RR = 0.04; 95% CI: 0.01–0.30, P &lt; 0.05</td>
<td>50% CR</td>
<td>12 cycles</td>
<td>3 wk CR, 3 wk, AL</td>
<td>(13)</td>
</tr>
<tr>
<td>Mammary, MMTV-TGF-α/Lepr&lt;sup&gt;−/−&lt;/sup&gt; transgenic mouse</td>
<td>RR = 0.18; 95% CI: 0.09–0.40, P &lt; 0.001</td>
<td>50% CR</td>
<td>12 cycles</td>
<td>3 wk CR, 3 wk, AL</td>
<td>(14)</td>
</tr>
<tr>
<td>Mammary, MMTV-TGF-α transgenic mouse</td>
<td>RR = 0.13; 95% CI: 0.06–0.28, P &lt; 0.05</td>
<td>50% CR</td>
<td>12 cycles</td>
<td>3 wk CR, 3 wk, AL</td>
<td>(15)</td>
</tr>
<tr>
<td>Mammary, MMTV-Neu transgenic mouse</td>
<td>RR = 0.60; 95% CI: 0.27–1.33, NS</td>
<td>50% CR</td>
<td>12 cycles</td>
<td>3 wk CR, 3 wk, AL</td>
<td>(16)</td>
</tr>
<tr>
<td>Multiple tumor types, p53-deficient (p53&lt;sup&gt;−/−&lt;/sup&gt;) mice</td>
<td>RR = 0.64; 95% CI: 0.31–1.32, NS</td>
<td>24-hour fast</td>
<td>28–40 cycles</td>
<td>1 d/wk for 7–10 mo</td>
<td>(17)</td>
</tr>
<tr>
<td>Mammary, spontaneous in DBA inbred mice</td>
<td>RR = 1.12; 95% CI: 0.94–1.34, NS</td>
<td>24-hour fast</td>
<td>220 cycles</td>
<td>24-hour fast, 2 times/wk for 110 wk</td>
<td>(54)</td>
</tr>
</tbody>
</table>

Abbreviations: CR, caloric restriction; AL, ad libitum; NS, not significant; MNU, 1-methyl-1-nitrosourea. DBA refers to a specific inbred mouse strain.

<sup>a</sup>Information is provided on the cancer site, agent used to induce cancer, and the rodent species used.

<sup>b</sup>The experimental approaches and number of rodents per treatment group varied markedly among the reported experiments. To provide a more uniform format in which to view and interpret experimental findings, risk assessment was limited to cancer incidence, and RR estimates were calculated using the cancer incidence and N for each group. Ad libitum–fed rodents were used as the referent group, and intermittently calorically restricted rodents were used as the experimental group. RR and CIs for the RR were calculated using the method described in ref. 56.

<sup>c</sup>Classification of weight cycling as defined in ref. 12.

<sup>d</sup>Two studies discussed in the text are not included in the Table (refs. 16, 17) because final aggregated cancer incidence data could not be accurately ascertained from the reported data. Similarly, refs. 21, 55 are not included because the primary endpoint of these prostate (LAPC-4 cells) xenograft studies was survival; the effect of intermittent caloric restriction was not statistically significant in either study.
protective effect on carcinogenesis associated with chronic caloric restriction (tumor incidence of 63% ad libitum vs. 23% chronic caloric restriction vs. 57% intermittent caloric restriction), a finding that was duplicated (45). Tagliaferro and colleagues had similar findings with intermittent caloric restriction during the promotion/progression phase of 1-methyl-1-nitrosourea–induced rat mammary carcinogenesis; 1 week of 33% caloric restriction followed by 3 weeks of paired refeeding (to prevent rebound eating) for 16 weeks (4 cycles) did not protect against mammary carcinogenesis (tumor incidence of 54% ad libitum vs. 66% intermittent caloric restriction; ref. 46).

Collectively, these reports indicate that short-term bouts of reduced caloric intake do not offer sustained protection against mammary cancer; whereas, maintenance of a lower body weight or weight loss to a maintained lower body weight is protective despite how the lower body weight was achieved. These findings parallel the epidemiologic evidence that adult weight gain is associated with an increased risk for postmenopausal breast cancer and that loss of excess weight for height is accompanied by a reduction in this risk (35, 47–49). The data from chemically induced cancer models are also consistent with the epidemiologic observation that weight cycling does not exert an independent effect on breast-cancer risk (34).

Genetically engineered mouse models of mammary and prostate cancer and lymphoma

Intermittent and chronic caloric restriction. Weight cycling has been studied in various genetically engineered mouse models of both nonobesity and obesity-associated cancer. A series of papers by Cleary and colleagues reported results of intermittent caloric restriction involving 3 weeks of 50% caloric restriction followed by 3 weeks of feeding matched (to prevent overeating) to the intake of ad libitum–fed mice in a mouse model of breast cancer induced by mouse mammary tumor virus (MMTV)-driven overexpression of TGF-α. They conducted weight-cycling experiments in mice that were heterozygous for a defect in leptin (lep<sup>ob/+</sup>), creating a predisposition toward obesity (13, 14), and in mice that were not (15). Intermittent caloric restriction protected against mammary cancer in both models (average incidence across studies: 75.4% for ad libitum–fed mice vs. 9.6% for mice on intermittent caloric restriction). At first glance it seems that these results are at odds with the work in chemically induced models and with the epidemiologic findings summarized earlier. Although it is difficult to reconcile the results of one of these genetically engineered model studies (13) with the other published work, the other 2 studies in this series (14, 15) indicated that growth curves and final body weights were significantly lower in intermittently caloric-restricted mice than in ad libitum–fed mice. When viewed in this light, the results are consistent with the findings of Kritchevsky and colleagues (40) and Sylvester and colleagues (41).

The experiments in the MMTV-TGF-α model also found that the effect of 50% intermittent caloric restriction, the type of weight cycling illustrated in Fig. 1B, was statistically significant and greater than that of 25% chronic caloric restriction in reducing cancer incidence, despite a similar overall intake of calories in both restricted groups. It is not clear, however, if this difference in effect on carcinogenic response was due to direct effects of the magnitude of caloric restriction (50% intermittent vs. 25% chronic) on host systemic factors such as insulin-like growth factor-1, leptin, or adiponectin and/or to direct effects on cell autonomous factors, such as the activity of the signaling network involving mammalian target of rapamycin, that modulate the carcinogenic process (19, 50). Alternatively, the differential effects of 25% chronically imposed and 50% intermittently imposed caloric restriction could have been a consequence of indirect effects on MMTV-driven transgene expression because activity of the MMTV promoter has been reported to be inhibited by dietary restriction (51–53). Other studies suggest that the observations from the MMTV-TGF-α model may not be generalizable. Neither intermittent nor chronic caloric restriction statistically significantly reduced tumor response in an MMTV-driven Her-2/Neu-overexpression model of breast cancer (mammary tumor incidences of 37.5% ad libitum, 33% chronic caloric restriction, and 22.5% intermittent caloric restriction; ref. 18), and intermittent caloric restriction caused only a modest and transient prolongation of tumor latency in the transgenic adenocarcinoma of the mouse prostate model of prostate cancer (16, 17).

Intermittent fasting. The effect of fasting 1 day per week and feeding matched to the diet of ad libitum–fed mice the other 6 days (to prevent rebound eating) versus the effect of chronic caloric restriction (i.e., 40% restriction relative to ad libitum–fed mice each day of the week) was studied in genetically engineered p53-deficient mice, where cancer development is considered inevitable (20). Although multiple tumor burden was reduced by either fasting or chronic caloric restriction, none of the differences were statistically significant (tumor incidence 40% ad libitum, 23% chronic caloric restriction, and 26% fasting). Although not in genetically engineered models, other studies support the genetically engineered findings. Tannenbaum and Silverstone found that intermittent fasting (24-hour fast 2 times per week) failed to inhibit spontaneous mammary cancer in the inbred (DBA)-mouse strain (tumor incidences of 80% ad libitum and 89% intermittent fasting; ref. 54). The effects of intermittent fasting were also investigated in a xenograft model of prostate cancer. An initial report indicated that intermittent fasting produced a nonsignificant trend toward improved survival following transplantation of LAPC-4 human prostate cancer cells into severe combined immunodeficiency (SCID) mice (21), but a larger follow-up study failed to detect a protective effect (55). Given that fasting is generally not recommended for weight control, these negative findings provide no support for considering the extreme method of caloric control by fasting for reducing cancer risk.
Conclusions

An individual’s body weight depends on the balance between caloric intake and caloric expenditure. In adults, small body-weight fluctuations occur throughout the day and during the course of a week. Overall, trends in energy balance (positive, negative, or equilibrium) result in healthy or unhealthy weight for height. Available data fail to make a compelling case that weight cycling exerts either beneficial or detrimental effects on health independent of effects associated with BMI. Rather, the weight of evidence reinforces the current public health recommendations regarding weight management: (i) maintain adult BMI in the target range of 18.5 to 24.9 (this range may differ depending on race) by preventing weight gain (the major cause of departing the range), and (ii) monitor and correct BMI above 24.9 by initiating weight loss to return to the target range. It is clear that weight cycling is an undesirable public health goal because health benefits of maintaining adult weight in the desirable range for height are well documented. The substantial evidence reviewed here shows that the cycle of weight regain following intentional weight loss generally does not reduce cancer risk, and the focus of weight-cycling research should be on ways to break this cycle.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

Acknowledgments

The authors thank Mary C. Playdon, Shawna B. Matthews, and John N. McGinley for their technical assistance in the preparation of this manuscript.

Grant Support

This work was supported in part by U.S. Public Health Service Grant R01 CA126704 from the National Cancer Institute.

Received March 15, 2011; revised June 25, 2011; accepted September 28, 2011; published OnlineFirst October 7, 2011.

References


Weight Cycling and Cancer: Weighing the Evidence of Intermittent Caloric Restriction and Cancer Risk

Henry J. Thompson and Anne McTiernan


Updated version
Access the most recent version of this article at:
doi:10.1158/1940-6207.CAPR-11-0133

Supplementary Material
Access the most recent supplemental material at:
http://cancerpreventionresearch.aacrjournals.org/content/suppl/2011/10/05/1940-6207.CAPR-11-0133.DC1

Cited articles
This article cites 52 articles, 12 of which you can access for free at:
http://cancerpreventionresearch.aacrjournals.org/content/4/11/1736.full#ref-list-1

Citing articles
This article has been cited by 2 HighWire-hosted articles. Access the articles at:
http://cancerpreventionresearch.aacrjournals.org/content/4/11/1736.full#related-urls

E-mail alerts
Sign up to receive free email-alerts related to this article or journal.

Reprints and Subscriptions
To order reprints of this article or to subscribe to the journal, contact the AACR Publications Department at pubs@aacr.org.

Permissions
To request permission to re-use all or part of this article, contact the AACR Publications Department at permissions@aacr.org.