NOT JUST CALORIC RESTRICTION: A COMPLEX APPROACH TO PROLONG LIFESPAN AND IMPROVE QUALITY OF LIFE

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Abstract
Aging is an urgent healthcare issue in view of the rapid growth of the proportion of older persons. Searching for reliable aging biomarkers and prolonging lifespan are increasingly important scientific directions. Experimental gerontology helps to explore fundamental facts which are not always applicable in clinical scenarios. As an example, caloric restriction is one of the key interventions that prolongs laboratory animals’ lifespan and ameliorates some, but not all, aging biomarkers in humans. Consequences of overeating such as obesity, insulin resistance, type 2 diabetes, and metabolic syndrome are taking their toll with aging, making caloric restriction a hot topic in gerontology and geriatrics. Nevertheless, caloric restriction is not widely applicable in view of poor adherence to and limitations of strict diets. Drugs mimicking caloric restrictions, the so-called caloric restriction mimetics, are developed to overcome these limitations. Caloric restriction alone is not a panacea since metabolic pathways are complex and not responsive to a single intervention. Fasting and exercising are additional options for reducing effects of excessive intake of calories. Arguably, physical activity significantly improves the quality of life at old age and delays the onset of overt insulin resistance and associated diseases. Thus, developing optimal fasting and exercising schemes is becoming increasingly important. Such interventions are confounded by a number of factors, including circadian and other biorhythms and baseline metabolic activity. It is justifiable to test fasting and exercising in experimental animals to reveal numerous confounding factors. A hypothesis in this article points to the role of complex interventions such as moderate and balanced diet, intermittent fasting, and physical exercise adjusted to circadian rhythms for prolonging life and improving quality of life. The hypothesis may shed light on fundamental mechanisms of aging and perspectives of anti-aging drug therapies.

Keywords: Aging, Caloric restriction, Lifespan, Longevity, Insulin resistance, Diabetes, Fasting, Circadian rhythm, Physical exercise


INTRODUCTION
Proportion of elderly people is constantly increasing, raising global concerns over the age-related diseases and old-age quality of life [1]. Aging disorders often involve multiple organ systems and manifest, for example, with insulin resistance, underlying obesity, type 2 diabetes, and metabolic syndrome [2, 3]. Obesity is a global issue affecting all age groups. According to the World Health Organization, more than 650 mln 18+...
adults were obese in 2016 (https://www.who.int/news-room/fact-sheets/detail/obesity-and-overweight). Caloric restriction is viewed as the most effective experimental intervention to prolong lifespan [4, 5]. Numerous hypotheses are proposed to explore the effects of caloric restriction on insulin resistance, obesity, and type 2 diabetes. Although some molecular mechanisms of the beneficial effects are widely known, related clinical studies often result in contradictory conclusions.

Signaling molecules associated with metabolism and proliferation such as insulin, insulin-like growth factor-1 (IGF-1) [6], and target of rapamycin (TOR) complex [7] play an important role in the development of age-related disorders. Both mitogenic and metabolic effects of these pathways influence aging mechanisms. Positive anti-aging interventions include increasing insulin sensitivity (e.g., Klotho activation), suppressing TOR (rapamycin), and IGF-1 (gene knockout) [5]. A critical role is also played by AMP-activated protein kinase (AMPK), predominantly antagonizing the effects of TOR and activating catabolic reactions (e.g., lipolysis, glycolysis).

Metabolic pathways are explored using a variety of model objects. Genetic and pharmacological interventions are known to prolong lifespan of various organisms, from unicellular yeast to primates. However, it is unclear whether caloric restriction can prolong the lifespan of humans. By singling out metabolic cascades, researchers often overlook the complexity of aging mechanisms.

Caloric restriction decreases basal metabolic rate [8]. It was previously believed that such decrease helps to prolong the average and maximum life span of laboratory animals [9]. However, the relationship between metabolism and lifespan is complex and not entirely understood. Physical activity increases the metabolic rate [10]. In long-lived insulin/IGF-1 mutant fruit flies, metabolism is not reduced [11]. Human subjects on long-term strict diet end up with slow metabolism, rapidly gaining weight after discontinuing the diet [12]. Previously, some studies have suggested declining basal metabolism from age of 15 [13]. Latest studies, however, indicated that the noticeable decline starts at 60 [14]. After 60, age-related disorders manifest clinically. In this regard, diet-induced slow basal metabolic rate may accelerate age-related diseases. Since high metabolic rate is associated with mortality risk in humans [15], optimal metabolic rate can be established to prevent age-related diseases.

There is also an argument related to the influence of caloric restriction on cell proliferation. Some researchers consider caloric restriction as a means of suppressing cell proliferation and reducing cancer risk [16]. Rapamycin, or sirolimus, a widely used anti-aging macrolide compound that inhibits the TOR complex, also suppresses cell proliferation and treats some tumors [17]. Nonetheless, cell proliferation is intimately involved in normal functioning of some organ systems such as musculoskeletal system. Muscle loss and atrophy lead to the enhanced course of type 2 diabetes and metabolic syndrome [18]. The TOR complex inhibition and AMPK activation can also lead to muscle loss. Hyperactivation of this kinase precedes Alzheimer's disease [19], while its baseline activity may prevent the deposition of abnormal tau protein in the brain and fight dementia. Catabolic processes such as autophagy complement cell-proliferation signaling pathways and induce muscle growth [20]. In this regard, intermittent fasting may regulate calorie intake and balance catabolic and anabolic processes.

The effect of intermittent fasting on aging biomarkers and lifespan is extensively explored in animal and human studies of obesity and other metabolic disorders [21]. Intermittent fasting fights obesity and improves cardiovascular functional capacities [22, 23] which can be associated with decreased basal metabolic rate [24]. The effect of intermittent fasting is closely related to circadian rhythms. Night shifts with high level of illumination and excessive use of electronic devices disrupt sleep. Social jetlag increases the risk of type 2 diabetes and metabolic syndrome [25]. Disturbed circadian rhythms in the elderly change the temperature rhythm and melatonin release [26]. Also, the disturbed rhythms predate neurodegenerative, metabolic, inflammatory, and oncological diseases, as well as, in different species, the deviation of the internal circadian rhythm from the real 24-hour one is in a negative relationship with life span [26].

Abnormal circadian rhythms affect eating habits by switching main calorie intake from breakfast and lunch to dinner. At the same time, eating and exercising can help synchronize circadian rhythms. Subjects with a late chronotype can shift their rhythm to an earlier one by morning and evening exercises [25]. Again, intermittent fasting may help a lot provided circadian rhythms are properly considered.

Although most laboratory mammals have nocturnal chronotype, research activities and related manipulations take place during daytime. Feeding of nocturnal rodents in the daytime uncouples the
peripheral clock from the central clock [27]. In a human study [28], considering experimental studies in mice, intermittent fasting, switched from dawn to sunset, resulted in a decrease of prooncogenes, an increase of antitumor and anti-diabetic factors, and improved body mass index and arterial blood pressure.

Experimental studies which are important for understanding mechanisms of caloric restriction are often at odds with clinical studies [29]. For example, genetic and pharmacological interventions may not always be applicable to humans, and in many cases, such use can make worse the quality of life of the elderly. Increased physical activity correlates with lowered risk of type 2 diabetes [30, 31], metabolic syndrome [3], and overall mortality [32]. At the same time, long-term use of rapamycin by young healthy volunteers led to the suppression of anabolic processes in their muscles [29]. Physical exercise helps maintain muscle mass and improves quality of life in older subjects. Importantly, skeletal muscles consume up to 70% of secreted insulin [33]. Physical activity improves insulin sensitivity [34, 35].

**HYPOTHESIS**

A combination of moderate nutrition, intermittent fasting, and physical exercise adjusted to circadian rhythms may help improve the quality of life and prolong lifespan. Caloric restriction and related mimetics cannot replace physical exercise. Moderate eating at regular intervals and at the same daytime may have better effects than prolonged fasting, caloric restriction, and use of caloric restriction mimetics. As a result, the basal metabolic rate should be maintained at a level corresponding to the age of 20–30. Food intake and physical activity should be adjusted to chronotypes. Meal consumptions with 6–8 hour intervals are advisable without snacks in between.

Overeating and drugs suppressing the TOR complex will have conflicting effects. In the case of drugs such as mild uncouplers (protonophores), some of the energy is lost in the form of heat but the potential of such substances as caloric restriction mimetics is not fully understood. Notably, overdosing of the most famous uncoupler 2,4-dinitrophenol is fatal [36].

Diets should not just reduce calorie intake; balancing intake of proteins, fats, carbohydrates, and other essential food components are critically important. Paying more attention to physical exercises is also important [37].

**HYPOTHESIS EVALUATION**

Examining “maximally healthy” animals is an optimal strategy to explore fundamental mechanisms of aging and search for anti-aging drugs [38]. Outbred lines like Wistar rats are more preferable for research since inbred lines have low viability. Mutant rats and mice are not suitable for anti-aging research.

Long-term experiments with construction of survival curves could be an optimal study design. Such experiments can be labour intensive and expensive, therefore it is permissible to conduct short-term experiments. In this case, starting age of animals (rats) should be 22–24 months (corresponds to 60 years in humans) with at least 3-month intervention and monitoring of aging markers. Subsequent longer-term interventions are desirable in case of initial positive effects. In a long-term experiment, until adulthood, animals should be kept in normal conditions and fed ad libitum, division into groups should take place after reaching maturity.

Before dividing into groups, it is necessary to weigh the animals, measure their temperature, determine the basal metabolic rate by indirect calorimetry with the determination of oxygen uptake and carbon dioxide production. Measuring blood glucose, blood and urine pH, and aging biomarkers should be monitored. Keeping the animals in metabolic cages for 24 hours before the experiment is also advisable.

The following division into groups is proposed: 1 – ad libitum feeding (control); 2 – caloric restriction; 3 – ad libitum feeding, rapamycin treatment; 4 – feeding 2–3 times a day, feed is given without limitation in quantity, but for a limited time, for 30–60 minutes (the ideal situation is feeding at night; as an alternative, feeding in the early morning and late evening), it is recommended to keep records of the food eaten; water consumption is always non-limited; 5 – the same diet, with the presence of running wheels. Rapamycin can be replaced with other drugs, for example, biguanides (such as metformin).

Regular weighing of animals and determination of their body temperature is required. The following parameters should be monitored: glucose, insulin (homeostasis model assessment – insulin resistance, HOMA-IR index), total cholesterol, triglycerides, low-density lipoprotein cholesterol, very low-density lipoprotein cholesterol, leptin, ghrelin, and lactate. The ad libitum groups should be deprived from food intake to measure fasting glucose and insulin levels. Quantifying
inflammatory markers is also important (e.g., TNF-α, IL-1, IL-6).

The basal metabolic rate should be measured weekly. A certain number of rats will need to be taken from groups for morphological analysis of the muscle fibers of the fore and hind limbs (m. biceps brachii, m. gastrocnemius, m. soleus, m. quadriceps femoris), as well as for assessing the level of expression of important participants (protein or mRNA) in metabolic cascades associated with caloric restriction, glucose metabolism, autophagy; e.g. glucose-6-phosphatase, IGF-1, AMPK, mTORC1, GLUT-4, IRS-1 (insulin receptor substrate 1), IRS-2, PKB/Akt (protein kinase B), LC3 (microtubule-associated proteins 1A/1B light chain 3B), p62, sirtuins, FOXO (forkhead box O) in muscles.

The circadian activity of genes responsible for circadian rhythms could be also determined. The choice of the method of feeding the animals and the drug to achieve the caloric restriction effect is left to the authors. Rapamycin (0.1 or 0.5 mg/kg of body weight per day) can be used.

2,4-dinitrophenol does not work when mice are kept at 25 degrees Celsius; keeping the temperature at 30 degrees Celsius is advisable [39]. Thus, the experiments proposed above can be repeated in conditions of thermoneutral temperature (when energy is not spent on thermogenesis). Finally, conducting additional experiments with periodic low temperatures is also important.

A series of experiments with animals on high-calorie diet is also interesting. To assess the lifespan of rats kept on regular and high-calorie diets, different experimental designs are required. The most effective fasting regimens can be chosen and adapted to human (volunteer) studies.

**ETHICAL IMPLICATIONS**

One of the main implications of the proposed hypothesis is the widespread dissemination of scientifically supported recommendations. Even practically healthy subjects may suffer from inappropriate dietary habits and excessive physical exertion. All such interventions should be evidence-based. Intermittent fasting can be detrimental for subjects with gastrointestinal disorders. When it becomes clear how the energy balance should be kept, it will be necessary to develop appropriate regimens and diets for subjects with health issues. Studies on healthy volunteers should precede larger-scale clinical trials.

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**CONFLICTS OF INTEREST**

The author declares that she has no conflict of interest.

**DISCLAIMER**

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**REFERENCES**

Текущие биомаркеры, несмотря на некоторые недостатки, являются важным инструментом в оценке метаболизма и прогнозе развития биологических процессов. Они позволяют оценить риск развития различных заболеваний и определить эффективность проведенных терапевтических мероприятий. Разработка новых биомаркеров и усовершенствование существующих является актуальной задачей в области медицины и биотехнологий.

Введение

Среди возрастающих проблем здравоохранения, вызванных ухудшением качества жизни и ускорением процессов старения, одним из ключевых является ожирение. Ожирение является основной причиной многих серьезных заболеваний, таких как сахарный диабет, артериальная гипертония, атеросклероз и ряд других. Понимание механизмов, лежащих в основе ожирения, позволяет разработать новые стратегии профилактики и лечения этих заболеваний.

Целью данной работы является обзор современных подходов к оценке биомаркеров риска ожирения и их потенциальной роли в профилактике и лечении данной патологии.

Основная часть

1. Биомаркеры ожирения

Биомаркеры ожирения - это побочные продукты метаболизма, которые могут быть использованы для оценки риска развития ожирения. К основным биомаркерам ожирения относятся инсулин-резистентность, глюкозурия, липиды крови и другие метаболиты.

2. Метаболический синдром

Метаболический синдром является комплексом состояний, включающих ожирение, гипертонию, гипергликемию и дислипидемию. Он является высокорисковым для развития сердечно-сосудистых заболеваний и диабета 2 типа.

3. Новые биомаркеры ожирения

В последние годы было разработано множество новых биомаркеров ожирения, которые позволяют более точно и точнее оценивать риск развития этой патологии. Эти маркеры включают в себя генетические маркеры риска, метаболические факторы и индекс ожирения.

Заключение

Введение новых биомаркеров ожирения позволяет более точно оценивать риск развития данной патологии и оптимизировать стратегии профилактики и лечения. Однако, дальнейшие исследования в этой области необходимы для уточнения значимости этих маркерах и разработки эффективных стратегий их использования в клинической практике.

Литература

исходную метаболическую активность. Целесообразно испытать голодание и физические упражнения на экспериментальных животных, чтобы выявить многочисленные сопутствующие факторы. Гипотеза в этой статье указывает на роль комплексных вмешательств, таких как умеренная и сбалансированная диета, прерывистое голодание и физические упражнения, адаптированные к циркадным ритмам, для продления жизни и улучшения её качества. Гипотеза может пролить свет на фундаментальные механизмы старения и перспективы медикаментозной терапии против старения.

**Ключевые слова:** старение, ограничение калорийности питания, продолжительность жизни, долголетие, инсулинорезистентность, диабет, голодание, циркадные ритмы, физические упражнения

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