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The authors of the above manuscript would like to apologize and retract it because in some paragraphs there are verbatim and unquoted sentences from others texts, although most of them, but not all, have been referenced. A corrected version of this review will be available in the next volume of Arq Bras Endocrinol Metab.

This retraction confirm the integrity of papers published in Arq Bras Endocrinol Metab.
Translational research into gut microbiota: new horizons in obesity treatment

Pesquisa translacional em microbiota intestinal: novos horizontes no tratamento da obesidade

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ABSTRACT

Obesity is a pandemic which has been rapidly developing for three decades. When a population is submitted to the same nutritional stress, some individuals are less susceptible to diet-induced weight gain and hyperglycemia. This observation suggests that other mechanisms are involved which are not directly related to the human genome. The human gut contains an immense number of microorganisms, collectively known as the microbiota. Evidence that gut microbiota composition can differ between obese and lean humans has led to the speculation that gut microbiota can participate in the pathophysiology of obesity. Different mechanisms have been proposed to explain the link between gut flora and obesity. The first mechanism consists in the role of the gut microbiota to increase energy extraction from indigestible dietary polysaccharides. The second, consists in the role of gut flora to modulate plasma lipopolysaccharide levels which triggers chronic low-grade inflammation leading to obesity and diabetes. A third mechanism proposes that gut microbiota may induce regulation of host genes that modulate how energy is expended and stored. However, further studies are needed to clarify a number of issues related to the relationship between the gut microbiota and obesity.

Keywords
Obesity; gut flora; energy extraction; lipopolysaccharide

RESUMO

A obesidade é uma pandemia que afeta milhões de pessoas em todo o mundo. Quando uma população é submetida ao mesmo estresse nutricional, alguns indivíduos são menos suscetíveis ao ganho de peso induzido pela dieta e à hiperglicemia. Essa observação sugere que outros mecanismos não diretamente relacionados ao genoma humano estejam envolvidos. O intestino humano é colonizado por milhões de bactérias, que coletivamente constituem a flora comensal normal. A evidência de que a composição da flora intestinal pode ser diferente em humanos magros e obesos levou à especulação de que a flora intestinal pode participar na fisiopatologia da obesidade. Diferentes mecanismos foram propostos para tentar explicar a correlação entre flora intestinal e obesidade. O primeiro mecanismo consiste no papel da flora intestinal na extração de energia de polissacarídeos não digeríveis. O segundo mecanismo envolve a modulação dos níveis de lipopolissacarídeo pela flora intestinal, o que desencadeia uma inflamação crônica subclínica que acarreta obesidade e diabetes. Um terceiro mecanismo propõe que a flora intestinal pode induzir a regulação de genes do hospedeiro que modulam como a energia é gasta e armazenada. Entretanto, estudos adicionais são necessários para estabelecer o papel da flora intestinal no desenvolvimento da obesidade.

Descritores
Obesidade; flora intestinal; extração de energia; lipopolissacarídeo
INTRODUCTION

Obesity is increasing among humans and is becoming a pandemic which has been rapidly developing for three decades (1). Obesity results from alterations in the body’s regulation of energy intake, expenditure, and storage. An increase in intake of energy-dense foods, specially combined with reduced physical activity, surely contributes to the high prevalence of obesity (2). However, when a population is submitted to the same nutritional stress, some individuals are less susceptible to diet-induced weight gain and hyperglycemia (3,4). This observation suggests that other mechanisms are involved which are not directly related to the human genome.

The human gut contains an immense number of microorganisms, collectively known as the microbiota. This community is dominated by anaerobic bacteria and includes ~500-1,000 species whose collective genomes are estimated to contain 100 times more genes than our own human genome (5,6). Our gut microbiota can be pictured as a microbial organ which contributes to our homeostasis (7); its functions are multiple and largely diverse.

Recently, gut microbiota has been shown to affect fat storage and energy harvesting (8), which suggests that intestinal microorganisms may play a direct role in the development of obesity. In this article, we review the published evidence supporting the potential role of the gut microbiota in the development of obesity.

NORMAL GUT MICROBIOTA

Commensal bacteria are present in vast number in the gastrointestinal tract. Resident bacteria outnumber human somatic and germ cells ten-fold, and represent a combined microbial genome well in excess of human genome (9).

Acid bile and pancreatic secretions hinder the colonization of the stomach and proximal intestine by most bacteria. The colon is colonized with approximately 10^{12} organisms/g intestinal content (Figure 1). Recently, it was demonstrated that Bacteroidetes and Firmicutes account for more than 90% of all phylotypes of Bacteria. More than 90% of the bacterial population are obligate anaerobes, predominant species being: Bacteroides, Enterobacterium, Bifidobacterium, Fusobacterium, Peptostreptococcus among other.

The intestinal flora plays an important role in normal gut function and maintenance of the host’s health. The benefits of commensal bacteria are well known:

help us to digest cellulose and salvage energy, form a natural defense barrier which is considered to be essential in the development, and maturation of mucosal and systemic immune systems. The gut microbiota is composed of potentially pathogenic bacteria besides numerous health-promoting nonpathogenic microorganisms.

Despite our limited understanding of the composition of the indigenous gut microbiota, evidence suggests that it is established within the first year of life. The fetal gut is sterile but colonization begins immediately after birth and is influenced by the mode of delivery, infant diet, hygiene levels and medication (10). Intestinal colonization begins at delivery, when the infant is exposed to vaginal flora, maternal fecal bacteria and environmental bacteria, and is likely to continue beyond the perinatal period, at least up to the age of 12-24 months, before a stable unique flora is established (11-13). Studies have shown that the adopted delivery method can have an influence. Infants delivered vaginally have earlier colonization with both Bifidobacterium and Lactobacillus, whereas infants delivered by cesarean section can have colonization with these beneficial organisms delayed by up 30 days (10).

As indicated above, the assembly of the gut microflora commences at birth, but its composition will undergo dramatic changes during postnatal development. The diet is clearly a key factor which regulates the sequence and the nature of colonization. Enterobacteria and bifidobacteria represent early colonizers, although differences in intestinal flora composition occur between breast- and formula-fed infants (14). In breast-fed infants, Bifidobacterium is a primary organism and the microflora produces high amounts of acetate and lac-
tate restricting the growth of potential pathogens such as *Escherichia coli* and *Clostridium perfringens*. In addition, colonization is affected by gestational age. Premature infants in the neonatal intensive care unit undergo a delayed colonization with a limited number of bacterial species that tend to be more virulent (15,16).

Although the composition of microbiota varies along the length of the gut and during the life of the host, it is quite stable during a considerable part of normal human lifespan. The transformation to adult-type microbiota is likely triggered by multiple host and external factors (17,18), including the effects of the microbiota itself, developmental changes in the gut environment, and transition to an adult diet.

A limited number of studies have indicated that the colonic microflora changes with ageing. The principal microbiological differences between adults and elderly were the occurrence of higher numbers of enterobacteria and a lower number of anaerobe populations in the elderly group (19). Bifidobacterial species, which are viewed as being protective, dramatically decreases with age, whereas clostridia and enterobacterial, which are viewed as being detrimental to health, increase (20,21).

The gut microflora of one person can differ markedly from another and comparative studies of adults with varying degrees of relatedness have shown that the host genotype is more important than diet, age, and lifespan in determining the composition of the gut microbiota (19,22).

Studies using germ-free mice have shown that gut microbiota is critical for maintaining normal gastrointestinal and immune function and normal digestion of nutrients. Germ-free animals are more susceptible to infection and have reduced vascularity, digestive enzyme activity, muscle wall thickness, cytokine production, mucosal-associated lymphoid tissue, motility and serum immunoglobulin levels (9).

**GUT MICROBIOTA AND OBESITY**

**Extraction of additional calories from ingested food**

Given the worldwide epidemic of obesity, there is interest in how interactions between human and gut microbiota may contribute to the development of obesity. Recent evidence suggests that the trillions of bacteria that normally reside within the human gut affect nutrient acquisition and energy regulation. Furthermore, bacterial lypopolysaccharide derived from intestinal microbiota may act as a triggering factor linking inflammation to high-fat diet-induced obesity (Figure 2).

Backhed and cols. demonstrated that conventionally reared mice have a higher body fat content than germ-free mice even though they consume less food than their germ-free counterparts (23). Along the same line, it was later shown that colonization of germ-free mice with an obese microbiota resulted in a significant greater increase in total body fat than colonization with lean microbiota (24). This study also characterized the distal gut microbiomes of obese leptin-deficient (*ob/ob*) mice and their lean (*ob/+* and *+/+*) littermates and it was found that obese mice have a higher proportion of intestinal Firmicutes with a corresponding reduction in the numbers of Bacteroidetes. These investigators showed that the microbiota of obese mice was rich in genes encoding enzymes that break down otherwise indigestible dietary polysaccharides. They also found more end products of fermentation and fewer calories in the feces of the obese mice, leading them to speculate that the gut microbiota in these mice facilitate the extraction of additional calories from ingested food.

Ley and cols. also demonstrated that obesity can be associated with altered gut microflora in a rodent model (25). The comparison of the gut microbiota of...
ob/ob mice versus lean mice showed that the relative abundance of the Bacteroidetes in ob/ob mice was 50% lower, whereas that of the Firmicutes was 50% higher (Figure 3). In obese humans, the same team of investigators reported a decrease in the relative proportion of Bacteroidetes when compared to lean individuals (26). Additionally, when obese patients lost weight over a one-year period, the proportion of Firmicutes became similar to that of lean individuals.

Further studies are needed to clarify a number of issues related to the relationship between gut microbiota and obesity. It is not clear whether the small increase of energy extraction can actually lead to a meaningful body weight gain within a short period of time, as suggested in the gut flora transplantation studies. Furthermore, it has been reported in other studies that a diet rich in non-digestible fibers decreases body weight, fat mass and the severity of diabetes (27,28).

Additionally, important questions remain unanswered. The first is why and how the make-up of the microbiota is shifted by differences in body weight. If a host organism had the ability to change its microbiota so as to increase caloric extraction, it would seem most adaptive to do so when facing famine conditions and losing weight. However, exactly the opposite was demonstrated as the microbiota seems to be more efficient in obese humans who already have the most stored energy (26). Moreover, there is also the question of how conditions in the host organism could change the make-up of the microbiota.

**Induction of subclinical inflammation**

Recently, a new hypothesis linking gut microflora to metabolic homeostasis have been proposed (Figure 2). On the basis of the recent demonstration, that obesity and type 2 diabetes are associated with low-grade chronic systemic inflammation (29-33) in the liver, adipose tissue and hypothalamus, Cani and cols. (34) hypothesized that bacterial lipopolysaccharide (LPS) derived from gram-negative bacteria residing in the gut microbiota acts as a triggering factor linking inflammation to high-fat diet-induced diabetes and obesity. They found that high-fat diet feeding resulted in a significant modulation of the dominant bacterial populations within the gut microflora. A reduction in the number of bifidobacteria, *Eubacterium rectale-Clostridium cocoides* group and *Bacteroides*, favoring an increase in the gram-negative to gram-positive ratio was observed. This modulation of gut microflora was associated with a significant increase in plasma lipopolysaccharide, fat mass, body weight gain, liver hepatic triglyceride accumulation, insulin resistance and diabetes. Another study has shown that treatment of rats with polymyxin B, an antibiotic that specifically targets gram-negative organisms, reduced LPS expression and hepatic steatosis (35).

Human studies have provided support for these findings. It was found that plasma lipopolysaccharide levels were significantly higher in the age-matched type 2 diabetes patient group than in individuals without diabetes (36). This study reinforced the hypothesis that lipopolysaccharide may act as a gut microbiota-related factor involved in the development of type 2 diabetes and obesity in humans.

A more recent study reported that modification of gut microbiota by norfloxacin and ampicillin, two antibiotics, ameliorated oral glucose tolerance and reduced hepatic steatosis in ob/ob mice (37). Cani and cols. (38) have also demonstrated that the modulation of gut flora by antibiotic treatment lowers plasma LPS levels, gut permeability, and the occurrence of visceral adipose tissue inflammation and macrophage infiltration in high-fat-fed mice. This effect was correlated with reduced glucose intolerance and body weight gain.
Regulation of host genes that modulate how energy is expended and stored

As described above, it has been recently demonstrated that control mice have a 40% higher body fat content and ~50% higher gonadal fat content than germ-free mice (23). In this study, when the distal gut microbiota from the normal mice was then transplanted into the gnotobiotic mice, there was a 60% increase in body fat within 2 weeks. To clarify possible mechanisms of this effect, the authors showed that the microbiota promoted absorption of monosaccharides from the gut and induced hepatic lipogenesis in the host, responses mediated by 2 proteins: carbohydrate response element-binding protein (ChREBP) and liver sterol response element-binding protein type-1 (SREBP-1) (23).

In an interesting experiment, using genetically modified (fasting-induced adipocyte factor \[Fiaf\]–knockout) mice, the same authors showed that gut microbes suppress intestinal Fiaf, also known as angio- poietin-like protein 4 (Figure 2). Fasting-induced adipocyte factor inhibits lipoprotein lipase activity, thereby catalyzing the release of fatty acids from lipoprotein-associated triacylglycerols, which are then taken up by muscle and adipose tissue. In the study, Fiaf suppression resulted in increased lipoprotein lipase activity in adipocytes and promoted storage of calories as fat, leading the authors to postulate that energy regulation by the gut microbiota occurs through at least three interrelated microbial mechanisms: a) fermentation of indigestible dietary polysaccharides to absorbable forms; b) intestinal absorption of monosaccharides and short-chain fatty acids with their subsequent conversion to fat within the liver; and c) regulation of host genes that promote deposition of fat in lipocytes (23).

In addition, Backhed and cols. (39) have investigated the mechanism(s) underlying resistance to obesity in germ-free mice, by studying germ-free mice consuming a high-fat and sugar-rich diet. They determined that germ-free animals were protected from diet-induced obesity by 2 complementary but independent mechanisms that result in increased fatty acid metabolism: (1) elevated levels of Fiaf trigger the production of peroxisome proliferator-activated receptor gamma coactivator, which is known to increase the expression of genes encoding regulators of mitochondrial fatty acid oxidation; and (2) the activity of AMPK (adenosine monophosphate–activated protein kinase), an enzyme that monitors cellular energy status, is increased. These findings suggest that the gut microbiota can affect both sides of the energy balance equation, influencing energy harvest from dietary substances (Fiaf) and affecting genes that regulate how energy is expended and stored.

CONCLUSION

Evidence that gut microbiota composition can differ between obese and lean humans has led to the speculation that gut microbiota can participate in the pathophysiology of obesity. Different mechanisms have been proposed to explain the link between gut flora and obesity. The first mechanism consists in the role of gut microbiota to increase energy extraction from indigestible dietary polysaccharides. The second, consists in the role of gut flora to modulate plasma LPS levels which triggers chronic low-grade inflammation leading to obesity and diabetes. A third mechanism proposes that gut microbiota may induce regulation of host genes that modulate how energy is expended and stored. However, many questions are still unanswered. First, it is not clear whether small changes in caloric extraction can actually lead to meaningful differences in body weight. Furthermore, it is essential to prove whether the differences observed in gut microbiota of obese people are the cause or consequence of obesity.

The human intestine is more densely populated with microorganisms than any other organ and manipulation of the gut microbiota may represent a new approach for treating obesity. However, the role of prebiotics, probiotics and antibiotics in the modulation of gut microbiota in obesity is unknown. Prebiotics are non-digestible oligosaccharides that are fermented by gut flora enhancing the growth of beneficial commensal microorganisms. In the future, prebiotics, probiotics and antibiotics may be used to modulate gut flora and perhaps they will have a role in the treatment of obesity and its complications.

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REFERENCES