Evaluation of the association between the TAS1R2 and TAS1R3 variants and food intake and nutritional status in children

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Abstract

Taste perception plays a key role in determining individual food preferences and dietary habits and may influence nutritional status. This study aimed to investigate the association of TAS1R2 (Ile191Val - rs35874116) and TAS1R3 (-1266 C/T -rs35744813) variants with food intake and nutritional status in children followed from birth until 7.7 years old. The nutritional status and food intake data of 312 children were collected at three developmental stages (1, 3.9 and 7.7 years old). DNA was extracted from blood samples and the polymorphisms were analyzed by real-time polymerase chain reactions (qPCR) using hydrolysis probes as the detection method. Food intake and nutritional status were compared among individuals with different single nucleotide polymorphism (SNP) genotypes. At 3.9 years old, children homozygous (Val/Val) for the TAS1R2 Ile191Val polymorphism ingested less sugar and sugar-dense foods than children who were *Ile carriers. This finding demonstrated that a genetic variant of the T1R2 taste receptor is associated with the intake of different amounts of high sugar-content foods in childhood. This association may provide new perspectives for studying dietary patterns and nutritional status in childhood.

Keywords: food intake, polymorphism, TAS1R2 gene, TAS1R3 gene, taste receptors.

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Introduction

The prevalence of childhood obesity throughout the world has increased rapidly in recent decades (Oliveira and Fisberg, 2003; Allison et al., 2008; WHO, 2010). The perception of flavors (sweet, bitter, salty, sour and umami) can play a key role in determining the preference for foods and dietary habits, thereby influencing nutritional status (Garcia-Bailo et al., 2009). Individual food preferences are based on physiological and nutritional factors, as well as economic and socio-cultural aspects. However, the sensory properties of food have a critical effect on dietary choices, and flavor can be a major determinant of food selection (Leterne et al., 2008; Garcia-Bailo et al., 2009). The T2R family of taste receptors is associated with the perception of bitter taste while the T1R family is associated with the perception of sweet and umami tastes (Nelson et al., 2001; Zhao et al., 2003). The T1R family consists of the T1R1, T1R2 and T1R3 receptors that are coupled to a heterodimeric G protein (Hoon et al., 1999; Zhao et al., 2003). The T1R receptors combine to form heterodimeric receptor complexes: T1R1-T1R3 is responsive to L-amino acids (umami taste perception) and T1R2-T1R3 is responsive to a variety of molecules, including natural sugars and artificial sweeteners (perception of sweetness) (Zhao et al., 2003). The receptor complex T1R2-T1R3 is encoded by the TAS1R2 and TAS1R3 genes, both of which are located on chromosome 1 (Liao and Schultz, 2003; Fushan et al., 2009). The TAS1R2 gene is characterized by a high natural diversity and the Ile191Val polymorphism (rs35874116) leads to the substitution of an isoleucine for a valine at position 191, which is a putative ligand-binding domain. Eny et al. (2010) reported an association between this polymorphism and sugar intake and differences in the response to dietary counseling. Fushan et al. (2009) described two polymorphisms within the TAS1R3 gene, -1572 C/T (rs307355) and -1266 C/T (rs35744813), that are in strong linkage disequilibrium and could be related to the perception of the taste of sucrose; these polymorphisms account for 16% of the variation in sensibility.
Genetic variation in taste receptors may be associated with greater sensitivity to particular flavors and result in a higher intake of some foods considered to be palatable but which are highly energetic and may be associated with an increased risk of overweight. The aim of this study was to examine the association of the *TAS1R2* gene and the *-1266 C/T* SNP in the *TAS1R3* gene with food intake and nutritional status at three developmental phases in children followed up until 7.7 years old.

**Subjects and Methods**

**Subjects**

The subject cohort was drawn from a randomized trial conducted in the first year of life to assess the impact of dietary counseling given to mothers during the first year of the infants’ lives and dealt with food consumption, nutritional status and lipid profile (Vitolo et al., 2010). The initial sample consisted of 500 children born from October 2001 to July 2002. Mothers were recruited from the maternity ward of the only publicly funded city hospital in São Leopoldo, a city in the southern Brazilian state of Rio Grande do Sul. HIV-positive mothers and infants with congenital malformations were excluded from the study. All eligible mothers were informed by field workers about the overall aims of the study and all research procedures. The intervention group received dietary advice during home visits in the first year of life, based on the ‘Ten Steps to Healthy Feeding’, a Brazilian national health policy for primary care, supported by the World Health Organization (Brasil Ministério da Saúde and Organização Pan-Americana da Saúde, 2002). The dietary advice encouraged exclusive breastfeeding during the first six months, after which the mothers were encouraged to continue breastfeeding and gradually introduce foods. Both groups (control and intervention) received visits at six and 12 months and routine follow-up by their pediatricians. This intervention was not the primary objective of the present research and participation in the intervention/control group was considered a confounding factor in subsequent statistical analyses.

The study sample consisted of 312 children who completed the initial phase of the study. The anthropometric measures and dietary data for each child were evaluated at three time points (1, 3.9 and 7.7 years old) at the child’s home. The main reason for the loss of subjects was the inability to locate the child’s home, usually because the family had moved to another city. Other reasons included refusal to continue in the study and children/maternal death. Race or ethnicity was classified by the interviewer based on skin color. Additional details of the traits studied are described in Vitolo et al. (2008).

The study protocol was approved by the ethics committee of the Universidade Federal de Ciências da Saúde de Porto Alegre and was in accordance with The Code of Ethics of the World Medical Association (Declaration of Helsinki) for experiments involving humans. All parents/guardians of the participants provided written informed consent before commencing the study.

**Nutritional status and dietary data**

Undergraduate nutrition students were trained to collect data related to the children’s anthropometric, dietary, sociodemographic and health status variables. Food intake was assessed through 24-h dietary recalls. At 1 year old, one 24 h dietary recall was recorded for each child. At 3.9 and 7.7 years old, two 24 h dietary recalls were recorded on two nonconsecutive days and the mean values were used in the analyses. The first dietary recall was done at the subjects home and the second was done in the municipal health unit. Portion sizes were confirmed with the aid of an album that was specifically designed for this study and contained photographs of utensils and foods, as well as domestic measurements, such as cups, tablespoons and teaspoons. The Nutrition Support Software (Nutwin 1.5, Federal University of São Paulo, São Paulo, Brazil) was used to convert the average sizes into grams for different measurements and to estimate the energy intake from the selected foods (Philippi, 2002; NEPA, 2006).

The energy intake from foods with a high sugar content (sugar-dense foods – SDF) was calculated from the two dietary recalls. The foods were classified as SDF when the sugar content was ≥50% and included items such as soda, Jell-O, candies and artificial juice. The foods were chosen based on the results of a study conducted in Brazil that indicated which foods were most commonly present in children diets. The average food intake from the two dietary recalls was used for the analysis.

At 1 year old, each child’s height and weight were evaluated. At ages 3.9 and 7.7 years, the height, weight, tricipital and subscapular skinfold thickness and waist circumference were measured. The body mass index (BMI) was calculated as weight (kg)/height$^2$ (m$^2$) and the values were transformed into Z-score (BMIzs). Details of the measurement procedure have been described by Vitolo et al. (2010). The percent variation in BMI for each child was computed with the formula \([\text{BMI} \_a - \text{BMI} \_b]/\text{BMI} \_b \times 100\), where \(a = \) final BMI and \(b = \) initial BMI.

**DNA analyses**

At the age of 3.9 years, blood samples were collected for hematological analyses (data not used in this study) and for DNA extraction. The blood samples were collected and transported to the Molecular Biology Laboratory of the Universidade Federal de Ciências da Saúde de Porto Alegre. Genomic DNA was extracted from peripheral blood leukocytes using a standard salting-out procedure (Lahiri and Nurnberger, 1991). The DNA samples are currently stored in this same laboratory. The *Ile191Val* polymorphism of the *TAS1R2* gene and the *-1266 C/T* polymor-
The number of children analyzed differed among the polymorphisms in each phase of the study because some biological samples were not analyzed for all polymorphisms. In samples collected from children at 3.9 years of age (n = 312), the average age was 3.98 ± 1.07 years old (mean ± SD), while for children at 7.7 years old (n = 274), the average age was 7.71 ± 0.72 years. At the age of 7.7 years, 49.3% of the children were boys and 43.2% were defined as white.

The allele and genotype frequencies for the TASIR2 Ile191Val and TASIR3 -1266 C/T polymorphisms are shown in Table 1. The genotype frequency distributions for the variants studied did not differ from those expected for populations under Hardy-Weinberg equilibrium (TASIR2 Ile191Val, $\chi^2_{G L=2} = 0.024$, $p = 0.988$; and TASIR3 -1266 C/T, $\chi^2_{G L=2} = 0.558$, $p = 0.757$). The most frequent allele for TASIR2 Ile191Val was *Ile (72%) and the most frequent allele for TASIR3 -1266 C/T was *C (87%). There were no significant differences in the allele and genotype frequencies between white and non-white children (data not shown).

After adjustment for confounding variables, 3.9 years old children homozygous (Val/Val) for the TASIR2 Ile191Val polymorphism ingested less energy from sugar-dense foods ($p = 0.027$) and sugars ($p = 0.025$) than Ile/Ile and Ile/Val children (Table 2). The percentage variation in BMI from 1 to 3.9 years was not significant among genotypes with the TASIR2 Ile191Val polymorphism, but there was a trend towards a smaller reduction in BMI in Val/Val homozygotes (-6.14 ± 9.13%) than in *Ile allele carriers (-9.13% ± 8.97%; $p = 0.052$) in this period (Table 2). For 1 and 7.7 years old children, no association was detected between food intake and nutritional status for this variant, but at 7.7 years there was a tendency for Val/Val homozygotes to have a greater BMI z-score; there was also greater variation in BMI in 3.9 to 7.7 years old children than in the corresponding *Ile allele carriers. The polymorphism TASIR3 -1266 C/T was not associated with food intake and nutritional status at the three evaluations done in this study.

**Table 1 - Allele and genotype frequencies for the SNPs TASIR2 Ile191Val and TASIR3 -1266 C/T.**

| SNP                  | % (n)    |
|----------------------|----------|
| TASIR2 Ile191Val     |          |
| Genotypes            |          |
| Ile/Ile              | 51.4 (160) |
| Ile/Ile              | 40.6 (126) |
| Val/Val              | 8.0 (25)  |
| Alleles %            |          |
| Ile                  | 72       |
| Val                  | 28       |
| TASIR3 -1266 C/T     |          |
| Genotypes            |          |
| C/C                  | 74.3 (232) |
| C/T                  | 24.7 (77)  |
| T/T                  | 1.0 (3)   |
| Alleles %            |          |
| C                    | 87       |
| T                    | 13       |

n = number of carriers of each genotype

$\chi^2 = 0.024$, $p = 0.988$; and $\chi^2 = 0.558$, $p = 0.757$.

Discussion

This study examined the association between two genetic variants of T1R taste receptors and food intake and anthropometric parameters during childhood. The Ile191Val polymorphism of the TASIR2 gene was found to

The statistical analysis

Allele frequencies were estimated by gene counting. A chi-squared goodness-of-fit test was used to determine whether the distribution of observed genotype frequencies agreed with those expected under Hardy-Weinberg equilibrium. The normality of the variables distribution was analyzed with the Kolmogorov-Smirnov test and by examining the histogram. BMI variation from 3.9 to 7.7 years old, SDF and sugar intake variables were ln transformed prior to analysis because of their skewed distribution, but the untransformed values are shown. The data for dietary intakes and parameters of adiposity were compared between genotype groups by using a General Linear Model with type III sum of squares. To construct the models, the daily energy intake, SDF and sugar intake, BMI Z-score and BMI variation between each child’s evaluations were included as covariates. The models for examining the association of polymorphisms with average daily energy intake, SDF and sugar intake were adjusted for intervention or control group status, sex and exact age, and exact age difference between each child’s evaluations were included as covariates. The most frequent allele for TASIR2 Ile191Val was *Ile (72%) and the most frequent allele for TASIR3 -1266 C/T was *C (87%). There were no significant differences in the allele and genotype frequencies between white and non-white children (data not shown).

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Discussion

This study examined the association between two genetic variants of T1R taste receptors and food intake and anthropometric parameters during childhood. The Ile191Val polymorphism of the TASIR2 gene was found to
Table 2 - Food intake and nutritional status in relation to the TAS1R2 and TAS1R3 polymorphisms in Brazilian children.

| Food intake and nutritional status | TAS1R2 Ile191Val | TAS1R3_1266 C/T |
|-----------------------------------|------------------|-----------------|
| 1 year old | Ile/Ile (160) | Ile/Val (126) | Val/Val (25) | P | P<sub>b</sub> | C/C (232) | C/T plus T/T (80) | P |
| Average daily energy intake (kcal/day) | 966.5 ± 395.10 | 904.4 ± 383.61 | 867.32 ± 386.17 | 0.277 | 0.242 | 924.8 ± 380.31 | 970.2 ± 421.0 | 0.422 |
| BMI Z-score | 0.65 ± 1.10 | 0.62 ± 1.05 | 0.28 ± 1.18 | 0.288 | 0.122 | 0.59 ± 1.10 | 0.69 ± 1.04 | 0.440 |
| 3.9 years old | Ile/Ile (160) | Ile/Val (126) | Val/Val (25) | C/C (232) | C/T plus T/T (80) | P |
| Average daily energy intake (kcal/day) | 1503.5 ± 386.1/77 | 1510.8 ± 335.47 | 1528.3 ± 400.2/77 | 0.232 | 0.233 | 1583.8 ± 398.03 | 1561.0 ± 319.73 | 0.596 |
| SDF (kcal/day)<sup>c</sup> | 80.5 ± 66.67 | 86.5 ± 86.74 | 86.5 ± 71.44 | 0.634 | 0.510 | 82.3 ± 75.48 | 88.35 ± 75.84 | 0.472 |
| Sugar intake (kcal/day)<sup>c</sup> | 3.9 ± 12.37 | 4.26 ± 9.62 | 10.16 ± 13.14 | 0.116 | 0.065 | 5.83 ± 11.83 | 4.96 ± 10.86 | 0.759 |

Values are the mean ± standard deviation; SDF – sugar-dense food.

<sup>a</sup>General Linear Model: for average daily energy intake, SDF and sugar intake, the intervention or control group status, sex and exact age were used as covariates. For the BMI Z-score, only the variable intervention or control group status was included in the model. For BMI variations, the exact age difference between the child’s evaluations, intervention or control group status and sex were included as covariates.

<sup>b</sup>For statistical analyses, the variables were natural logarithmic transformed but the non-transformed values are shown in the table.

<sup>c</sup>For statistical analyses, the variables were natural logarithmic transformed but the non-transformed values are shown in the table.

*TAS1R2* gene polymorphism was not significantly associated with food intake or SDF in children and not the sensitivity to sweetness. We hypothesize that the TAS1R2 gene polymorphism may not be a causal polymorphism but may instead be in linkage disequilibrium with a causal polymorphism. These findings are consistent with the hypothesis that the TAS1R2 gene polymorphism is not yet known and there is currently no information on this matter in the PolyPhen database. This variation is located in the ligand-binding extracellular domain that is specific for the *Ile191Val* polymorphism. The functional significance of the *Ile191Val* polymorphism is not yet known and there is current no information on the *Ile191Val* polymorphism. The decrease in daily energy intake and SDF was not significantly associated with anthropometric parameters or with variation in weight gain in this cohort. Additionally, contrary to our initial hypothesis, there was a trend towards a higher BMI Z-score at 3.9 and 7.7 years old and the TAS1R3_1266 C/T polymorphism was not significantly associated with anthropometric parameters or with variation in weight gain in this cohort. The mean daily energy intake and SDF was not significantly associated with anthropometric parameters or with variation in weight gain in this cohort.
size that to detect an association between a genetic variant of these taste receptors and the habitual consumption of sugar, this variation needs to have a strong effect on protein functionality or expression. In this context, the approach of Fushan et al. (2009) is more sensitive for detecting small genetic effect.

Our findings suggest that genetic variation in one component of the T1R2-T1R3 heterodimer may account for interindividual differences in the consumption of sweet foods, although this association was not reflected in modifications in total energy intake or in anthropometric parameters. Our findings contribute to knowledge in the area of taste genetics and may help to explain the different ability of individuals to alter their dietary intake in response to dietary advice. However, this study has some limitations. Only one polymorphism was evaluated for each gene and these polymorphisms were selected based on previous studies describing their possible association with food intake and sweet sensitivity. Another relevant feature was the age of the children. In view of their young age, the children may not have had the freedom to choose their foods and they may instead have been supplied with foods that were palatable to adults. In addition, the participants may have consumed all the available foods equally, regardless of whether they presented a greater sensitivity or not, and our data on food intake do not necessarily reflect the sweet sensitivity and food preferences. Furthermore, the food categories analyzed (SDF and sugar intake) may not accurately reflect the issue of palatability and sweet taste because some foods, such as soda, may have a high concentration of sugar but be perceived as less sweet. However, the association detected between genetic variants of the sweet taste receptor gene and SDF and sugar intake makes biological sense. To the best of our knowledge, no other studies have evaluated the association between genetic variations in the taste receptor genes and food intake and nutritional status at different stages of childhood. The results described here may open new perspectives for researchers and contribute to the study of childhood dietary patterns and nutritional status.

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