Variation in Nutrient Intakes and Required Number of Days for Assessing Usual Nutrient Intake among Different Populations

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Abstract

Estimating the usual intake of a population is essential in the process of establishing the protective effects of nutrients against the development of certain diseases and assessing nutrient intake adequacy [1]. Common assessment tools used in epidemiological studies include diet histories, dietary recalls (DR), food frequency questionnaires (FFQ), and food records (FR) [2]. A long term assessment of daily intakes is required to assess usual intakes, however, due to the cost and burden of this requirement, most studies employ shorter term assessments. Many limitations exist with dietary assessment tools which can alter the results and conclusions of intake data. This article was aimed to review on the variation of nutrient intakes and required number of days to assess usual nutrient intake status of diverse population groups based on previous publications.

Keywords: Usual nutrient intake; Misreporting; Within-person variation; Between-person variation; Antioxidants

Introduction

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of misunderstanding or error in dietary instruction on portion sizes for the participant or lack of cooperation in consistently measuring food intake. In addition, elimination of certain foods or amounts can be the result of the study participant intentionally or unintentionally selecting foods due to the bias of the study towards analysis of certain nutrients [20]. Regardless of classification of misreporting, the inclusion of individuals who provide inaccurate nutrient intakes can alter the results and conclusions significantly. A common method for identifying such individuals is the Goldberg's cut off equation [21]. This equation requires the average energy intake, average basal metabolic rate, and daily physical activity of the population to generate critical values for energy intake applied to the average energy intake of each individual participant. The population critical values are represented by energy intake: BMR estimated ratio (EI/BMR).

A review by Black provides a guide for the use of the Goldberg cut off in nutritional assessment research [4]. In the review, Black emphasizes the importance of selecting a physical activity level (PAL) for each population dependent on reported daily physical activity and classifications provided by the World Health Organization (WHO) [22]. For the average student population group, a 1.6-1.7 PAL is suggested for determining energy requirements [22]. In data reported by Black [4], a PAL of 1.7 was used for young adults aged 18-29 y who were predominately non-Hispanic White and participated in moderate leisure activities. However, a high PAL value can inaccurately identify individuals as low energy reporters especially in a study with a small sample size [4]. In order to increase the sensitivity and specificity of the Goldberg cut off in a study with a population size (n) < 100, the number of days of dietary intake assessment should be increased [23]. However with larger sample sizes, fewer numbers of dietary assessments may be used. Results from the National Health and Nutrition Examination Survey 1988-1994 (NHANES III) for misreporting using the Goldberg cut off include a critical value of 0.9 to 1.54 with a mean EI_BMR of 1.36 for all adults. In addition, 18% of males and 28% of females were classified as under-reporters of energy intake [6].

After the identification of misreporters in a population group, causality should be determined before the decision to include or exclude individuals from the results. In a recent review, Poslunsa et al. summarized the main causes of errors in 24-hr DR and FR most frequently reported in 38 nutritional studies [24]. Results indicated that the major determinants for misreporting included body mass index (BMI), age and sex, socioeconomic status and education, health related activities, psychological factors, and eating habits. While misreporting includes both underreporting and overreporting of nutrient intake, overreporting was identified less frequently in these studies. The most consistent factor reported in the review was that as BMI increased, a larger percentage of the population was classified as misreporters, specifically among females. In addition, more females than males tended to misreport their nutrient intake [24]. Similar gender results were found in a study with 53 non obese, weight stable adults. They reported 49% of the females and 14% of the males were identified as under-reporters from a 7-day DR [19].

It remains unknown whether males tend to underreport less than women do or if their higher energy requirements allow them to rarely fall below the cutoff limits when applied to an entire study population [23]; however, Asbeck et al. reported that the higher percentage of female underreporters was due to restrained eating practices evidenced by scores from an eating practice survey in a normal weight population [19]. Leibman et al. conducted a study with 324 college students analyzing the relationship between dieting practices, gender, and psychological variables such as self image and body perceptions [25]. The results showed that 38% of females and 13% of males had dieted to lose weight within the past year and more females reported patterns of disordered eating, such as fat avoidance or replacement, and body dissatisfaction [25]. Body weight dissatisfaction, frequent dieting, and societal pressures seem to be an area of concern in young adult and adult female populations; therefore the validity of dietary assessments from these population groups should be analyzed before average intake results are reported [26].

**Within- and between-person variation of nutrient intake**

Day-to-day variability in nutrient intakes can significantly alter the statistical outcomes and interpretations of dietary assessment data. This fluctuation is defined as within-person variation and can be attributed to environmental and cultural factors [27]. Micronutrients have a higher concentration in specific foods and tend to have greater variation due to seasonal variation or the wide array of food choices available in many developed countries when compared to macronutrients which remain more stable in the diet. However, seasonal variation has a greater impact in developing countries where all foods are not as easily accessible [27]. Day of the week sampled by a dietary assessment tool is another source of within-person variation. Energy and protein consumption are typically larger on the weekends compared to the weekdays and should be considered when using 24-hr DR [9]. Within-person variation can be estimated and must be adjusted for statistically due to its high correlation to the mean of the sample day. This is crucial in the interpretation when the study design only includes a small number of days of dietary intake [5]. However, increasing the number of days of diet recorded can decrease the within-person variation significantly [3]. Another important consideration is a large variation between individuals of a population because it may misconstrue the relationship between nutrient consumption and disease risks [4,28]. Between-person variation can be reduced by accounting for certain sociodemographic and lifestyle factors specific to the group of study [12,27,29]. The ratio of the within to between variation can be used to further describe the effect of the within-person variation [1]. The greater the variance ratio, the greater the within-person variation in daily intakes.

Several methods have been developed to assess usual dietary intake among populations [1,8-10]; however all methods require estimation of within- and between-person variation. Therefore, these values must be calculated from multiple numbers of diet records or values can be borrowed from an appropriate subset population [1]. Chang et al. analyzed the within- and between-person variation among Taiwan college students who completed a total of three 5-day DR [1]. They found that males had larger within to between ratios for fat, protein, polyunsaturated fatty acid, vitamin A, thiamin, and riboflavin than the females which they attributed to the irregular eating patterns and possible binging of male college students. Females had larger within-person variation in the intakes of carbohydrates which could be a result from the common practice of dieting or meal skipping in this population group [1]. In another study, Jahns et al. analyzed the effects of gender as well as age and culture on the estimation of within- and between-person variation in U.S. and Russian older children and adolescents [29]. Results were reported from nonconsecutive 24-hr DR from the Russia Longitudinal Monitoring Survey (RLMS) and the Continuing Survey of Food Intake by Individuals (CSFII). They analyzed energy intake and 10 additional macro- and micro-nutrients: protein, carbohydrate, fat, calcium, iron, magnesium, thiamin, riboflavin, niacin and the antioxidant nutrient as vitamin C. Among the U.S. population, they found that the girls had higher within-person variation than the boys for all nutrients excluding carbohydrates and the girls had higher...
between-person variation as well. Results pertaining to the differences in age groups reported that the older Russian girls had higher within-person variation for all nutrients except riboflavin, niacin, and vitamin C as well as higher between-person variation for all nutrients except magnesium and thiamin than the younger girls. No observable patterns were found among the U.S. age groups for within-person variation but the between-person variation was higher for the older girls for 9 out of the 11 nutrients including vitamin C [29].

In U.S. men and women, Neuhaus et al. analyzed the ratios of within-person variation to between-person variation in different age groups for energy, 3 macronutrients, and 9 micronutrients including vitamin C [30]. They found that as age increased, the variance ratio decreased meaning the within-person variation was greater than the between-person variation. These results were significant among men for most nutrients, however, a decreasing trend was not as apparent for women [30]. Overall, the results seem to indicate that younger adult populations may have larger day-to-day variability in nutrient intake, which has important implications with estimating usual nutrient intakes of a population. While these studies do include within- and between-person variation among adolescents and young adults, there is a gap in the literature pertaining to antioxidant intakes among this age group in the U.S.

Due to the numerous errors associated with dietary intake assessment, it is important to consider analysis through biomarkers as a complimentary methodology. For example, a review by Hudson et al. provided evidence to support the use of tissue and blood fatty acids as biomarkers for intake due to its correlation of precisely controlled nutrient intake [31]. In addition, multiple studies have identified misreporting by verifying intake using doubly labeled water to measure total energy expenditure (TEE) and 24 hr urinary nitrogen to measure protein intake. However, these methodologies are time consuming for the researchers and participants as well as expensive, specifically in reference to the doubly labeled water technique. The measurement of antioxidant intake has also been assessed through biomarkers. For example, carotenoids can be measured in the plasma, serum, or the skin through the use of high performance liquid chromatography (HPLC) [32]. In addition, vitamins C and E can be measured using HPLC. The lipid soluble antioxidants, carotenoids and vitamin E can be measured in adipose tissue; however, due to the procedure involved, participation from eligible subjects may be a challenge [32]. There are limitations associated with the methodologies utilized in measuring biomarkers, which include lack of reproducibility, specificity, variation within- and between- individuals, and lack of detection. Despite the limitations associated with any method of assessing nutrient intake, there is promising research that nutrient intake data used in conjunction with biomarkers can produce valid and reliable results.

The Number of Days Required to Accurately Assess Nutrient Intake

It is important, when developing a study design, to know how many days of dietary assessment are required to produce accurate and reliable intake results for a population group [11]. To assess usual nutrient intake levels among a population, within- and between-person variation should be estimated and included in a calculation to determine sufficient number of diet records necessary to produce accurate results [11,12].

The calculation of days (D) of nutrient intake includes the ratio of within-person variation \((S_b)\) to between-person variation \((S_w)\) [33]. The variability in daily nutrient intake among adults has been shown to be greater than the variability between individuals in a study population [11], and the smaller the ratio, the fewer number of days is required to estimate the nutrient intake within a specified level of accuracy \((r)\) between true intake and observed intake [11]. Nelson et al. analyzed data from 18 studies that reported mean nutrient intake, values for within- and between person variation, and the number of days required to estimate true intake within a given accuracy. They included studies with populations aging from infancy to older adults and reported a total of 29 nutrients including energy. Values presented for D were based on \(r = 0.9\). Most nutrients required more than 7 days of DR to estimate true intake in all age and gender groups. Among the adult populations, energy, protein, carbohydrate, and fat required 4-8 days depending on gender. Females required more days than males for all macronutrients. Vitamin A and carotene were required to require three weeks or more to estimate true intakes with adult females requiring over a month to estimate carotene. Results pertaining to vitamin C included 12 days for males and 7 days for females while vitamin E required 8 days for males and 16 days for females. In general, this study found that the population group that required the most days to estimate true intake was 5-17y with adults requiring an intermediate amount [11].

Mennen et al. reported analysis of the number of 24-hr DR required among French adults participating in the Supplémentation en Vitamines et Minéraux Antioxydants (SU.VI.MAX) Study which investigated the effects of antioxidant supplementation on cancer and heart disease [34]. Participants included in the additional analysis completed six 24-hr DRs over a year and the study was separated into two phases consisting of 2 years each. Nutrients included energy and macronutrients with vitamin C, vitamin E, and β-carotene as the antioxidant micronutrients. Results from the first phase included the highest variance ratio for β-carotene and the lowest for carbohydrate. Carbohydrate required 5 days of DR while β-carotene required 16 recalls. For protein, total fat, and vitamin C, results showed 8 DR would be needed while vitamin E required 10 recalls for this French adult population. In general, the women required the same or more DR to estimate true intake for the macro- and micro-nutrients included [34].

A study was conducted in preschool age children reporting the variation in macronutrients and 11 micronutrient intakes stratified by age groups and gender [12]. Huybrechts et al. concluded that as the age of the children increased, the larger the variance ratios became and more days of DR were required for all nutrients. A 7-day DR would be sufficient to estimate energy and macronutrients when analyzing gender; however, results from the age groups indicate than more than 7 days would be required for the older children. Vitamin C could be estimated in 5 days among all age groups and genders [12]. In an older adult population in Korea, the number of days to estimate energy, protein, fat, and carbohydrates among males was over 2 weeks; however, vitamin C required 54 days to estimate true intake [27]. Females required 8-23 days to estimate their macronutrient intake while vitamin C required 16 days. Oh et al. concluded these results were attributed to the large within-person variability and low between-person variability in this population group [27]. Due to the population demographic, many of these studies can only serve as implications for study design. There is limited data on nutrient variability and number of days needed to assess nutrient intake, including antioxidants, among populations of interest.

Conclusion

Estimation of usual nutrient intake from dietary sources among a target population is a vital part in assessing dietary quality and the
risk of developing diseases. In order to do so, nutrient intake data must be validated by identifying misreporters and determining the variation among this population. It should be noted that an important application of these methods of validation and estimation of variation components of nutrient intake among a population of study is the evaluation of nutrient intake adequacy according to the Dietary Reference Intakes (DRIs) and the Dietary Guidelines for Americans. While research is often devoted to assessing diet quality and disease risk factors in adults, there is limited data in comparison focusing on this significant population group. With the steady increase of obesity and other diet related illnesses, it is critical that research professionals report accurate nutrient intake in order to link diet and disease risk.

References