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Sugary beverages consumption and latent autoimmune diabetes in adults: systematic review and meta-analysis

ABSTRACT

Introduction. Sugary beverages consumption (SBC) has amplified globally. SBC is associated with and leads to obesity and chronic diseases, nonetheless the role of SBC in development of autoimmune disorders such as latent autoimmune diabetes in adults (LADA) has not been addressed adequately among the different ethnic groups. We conducted this meta-analysis to compare the random effect of SBC intake on the risk of development of LADA.

Methods. We scrutinized the MEDLINE database up until January 2019 for articles addressing the association between sugary beverages, coffee consumption and LADA. We found 6 studies all of them addressed the LADA. We have included them in the meta-analysis and compared the random effect of SBC from the uppermost to the lowermost quantiles parallel to the risk of LADA.

Results. According to the research conducted, and data extracted, which involved 15027 contributors and 1862 patients with LADA, the participants in the uppermost quantile of SBC intake (used 1–2 servings per day in most cases) were at risk of developing LADA more than those in the lowermost quantile (≤ 1 serving per month) (odds ratio [OR] 1.37 [95% CI 1.23–1.52]).

Conclusion. According to the meta-analysis results excessive SBC intake may increase the risk of development of latent autoimmune diabetes in adults. However, no definite conclusions could be drawn due to heterogeneous data from low quality researches and the analysis was based on observational and case-control studies only. (Clin Diabetol 2020; 9)

Key words: sugary beverages consumption, latent autoimmune diabetes in adults, systematic review, meta-analysis

Introduction and background

In almost 50 years sugary beverages consumption (SBC) has increased at an alarming rate worldwide. For instance, in the United States, from 1970 to 2006 SBC per each individual reised from 64.4 to 141.7 kcal/day, forming double or twice the increase [1]. Similar results have been revealed in Mexico, where presently more than 12% of total calorie intake was represented by SBC [2]. The rapid and dramatic increase of SBC in several developing republics where SBC has increased concurrently in relation with increasing rates of growth and urbanization. In the 2007 annual report, the Coca-Cola company shows that, the amount of SBC sold in India and China increased by 14% and 18% respectively in one year, indicative of the considerable upsurges in trade at the national level [3]. For clarification sugary beverages include carbonated sodas, energy drinks, sport drinks, juice drinks, sweetened tea, iced tea, fruit drinks, and vitamin drinks. Recent research has been
The most common biochemical marker in LADA is acid decarboxylase antibodies (GADA) with LADA. Of autoimmune indicators such as anti-glutamic acid decarboxylase antibodies (GADA) with LADA. On the other hand, a drink that is a 100% pure and natural fruit juice and not mixed with extra sugars is not considered a sugary sweetened beverage. Progressively, teams of researchers and institutes are calling for maximum reductions in SBC [5, 6]. Results from significant prospective researches in epidemiology have demonstrated a reliable positive correlation between SBC and obesity among children and adults [7]. Furthermore evolving proof also proposes that habitual SBC is correlated with a higher risk of developing diabetes and other metabolic disorders [8]. SBC has been proven to be the cause of obesity due to their more added sugar and imperfect recompense for total calorie intake [7].

Due to the high amount of fast absorbable sugars such as fructose corn syrup and sucrose, in combination with the large amounts consumed, SBC might increase the risk of diabetes through obesity and by raising the dietary glycemic index, and insulin resistance, which contributes to β-cell dysfunction [9]. An increase in metabolic impacts of SBC may also cause elevated blood pressure and the buildup of visceral fatty tissue and ectopic adiposities due to high liver de novo lipogenesis [10], which will in turn lead to the development of more triglycerides, LDL and decrease the level of HDL. The correlation between SBC and LADA is less clear [11, 12], but current research proposes that SBC may elevate the risk of diabetes in hereditarily predisposed subjects [11]. Conceivable mechanisms for SBC participation in autoimmune pathogenesis involve prompted beta cell apoptosis [13], perhaps due to prompted oxidative stress, high glucose levels [14, 15] or an overwhelmed beta cell, probably because it is more visible and exposed to the body’s immunity [16].

LADA is a form of diabetes combining the features of both of type 1 and type 2 diabetes. Besides the involvement of autoimmune indicators such as anti-glutamic acid decarboxylase antibodies (GADA) with LADA. The most common biochemical marker in LADA is a mild or moderate insulin resistance [17]. Therefore, it is likely that SBC may impact the risk of LADA by the pathogenesis related to autoimmune disorders or insulin resistance, but then again this is still unclear. 9% of all cases diagnosed as adult-onset diabetes was recognized as LADA [18], which is considered a mixture of different elements from different types of diabetes. After review of the available literature, we conducted a systematic review and a meta-analysis to observe the relationship between SBC and the risk of development of LADA.

Research methodology and design

Search of literature

Guidelines of the PRISMA 2009 Statement have been adopted — step by step as we conducted our meta-analysis [19]. Pertinent, applicable and multi-ethnic researches written in the English were recognized and acknowledged by an in-depth and meticulous probing of the following databases: MEDLINE electronic database; Cochrane Library; PsycINFO — American Psychological Association; Embase; CABI Abstracts; Web of Science by Clarivate Analytics (formerly known as ISI Web of Knowledge); CINAHL Database, BIOSIS (King Saud University Medical City Library of Medicine, Salah, MD) for studies from 1983 to January 2019, which included SBC and sugar-sweetened beverage consumption, such as: (soft drinks, soda, carbonated drinks, sweetened coffee, iced tea, alcoholic beverages, fruit drinks, squashes, sports drinks, soda-pop, cordials, energy drinks, punch, vitamin water drinks and sugary lemonade) and the risk of LADA. We searched for keywords including those mentioned above as well as those combined with “auto antibodies”, “autoimmune disorders of β-cells of the pancreas”, “latent diabetes”, “latent autoimmune diabetes in adults”, and “LADA.” We used this method as extensively as possible as our primary means of exploration and in the next successive medical subheading (MESH) terms examination. Every relevant article that was found, we followed its references, searching for any hints of another thread. We searched for references and cross-references that would possibly guide us to other references. We searched not only for articles published in journals, but also those in the press, books, magazines, newspapers, websites, documentary films, dissertations, congressional publications, international organizational reports, and even editorials by deploying the (Citation Machine™) as a means of accomplishing our purpose and achieving our objective.

Due to the high possibility for confusing and converse causation, we have excluded cross-sectional researches. We have also excluded short-term trials as they were unable to address the long-term relationship that we are exploring. However these short-term studies do provide significant intuition about the possible causal biological mechanisms and thus has helped further our understanding of the causality in some capacity.
Inclusion criteria and extracting data

In our meta-analysis we included population based observational epidemiological studies as inclusion criteria. Criteria for inclusion comprised the end points of LADA, associated measure of variance (standard error or confidence interval) and relative risk as well as measures of SBC and potential mediators’ adjustment. After we applied these criteria, our collected works selected eight identified articles out of 148 relevant references. Those 148 references were derived from 7534 citations (Figure 1). Each of the eight studies hit our target precisely [20–27]. Two of these studies have been excluded as one of them was review of literature [27] and the other was a master’s thesis which was published as a paper later on, and selected from our target group [24].

The remaining six studies [20–27], all of which were held in Sweden, and written in English and those were two main weak points and mentioned as limitations. Only two addressed the genetic susceptibility as an independent variable [20, 25]. Estimated adjusted odds ratio (OR) of diabetes were entailed in relation to SBC. Standard errors and coefficients of variation were attained from Rasouli et al. [21] and Löfvenborg et al. [22] through subsequent communications. Two of the team members independently extracted the data. No variances were noticed in the extracted data to provide estimation of the effect, comparing skewed or drastic quantiles of SBC. All studies have defined one serving as 200 ml however there are some notable variations in estimation of the serving size including Löfvenborg et al. [20] in which the maximum level of SBC was 200 ml servings per day and lowest amount of intake was 200 ml servings per week. With Rasouli et al. [21], the average intake of alcoholic beverages was 12 grams per day, while the maximum amount served was 25 grams per day. Löfvenborg et al. [24] classified the amount of servings of soft drinks and sodas as (< 1, 1–2 and > 2 servings per day). Finally the least amount of coffee served with Rasouli et al. [25] was < 1 cup per day and the highest was > 4 cups.
Patient and public involvement statement

Our current study design was conducted in the form of meta-analysis and systematic research on already published study articles, so no patient involvement was documented, and the used materials was only published data.

Limitations of the study

The first limitation and point of weakness is that we conducted this meta-analysis and there was heterogeneity of results. All studies had different populations, different designs and outcomes. However all of them addressed the same topic and the same research question. Available literature regarding this topic is very scarce, in addition to heterogeneous data from low quality studies depending on retrospective, observational or case-control designs only. This in turn reflected negatively on the level of evidence and conclusion. Moreover, there were no prospective or long term experimental, interventional or randomized studies, with sufficient follow-up period, to demonstrate the potential relationship between sugary beverage consumption and LADA, and most probably such researches will never be conducted due to ethical reasons. The included studies were restricted to those published in English which lead to exclusion of non-English studies with their evidence base. This may increase the likelihood of selection bias. Also, there was point of limitation and unavoidable weaknesses. All of the studies were in Sweden. And we could do nothing to overcome all these point. Wide range of the dates of publications of the included studies, almost five decades, increases the validity and significance of the results.

Analysis and investigation

A total of six studies with nine data points are comprised in this meta-analysis of LADA and sugary beverages consumption [20–25]. We used STATA (version 9.0; Stata Corp, College Station, TX, USA) to attain instantaneous relative risks employing random effects models as well as fixed effects models designed from the logarithm of the relative risks and matching 95% confidence intervals of the separate studies [20–27]. A random-effects model was used primarily because it integrates the constituents of variance within the study itself and also between the studies. Egger’s test was acknowledged to be employed in case of heterogeneity between studies and it’s also considered to be the more conventional method [28]. We assessed the heterogeneity significance of the results throughout our selected studies by the application of Cochrane Q test, in spite of the presence of lack of sensitivity.

We followed Cochrane Q test by an I² statistical analysis which embodies the proportion of whole disparity across studies because of inter-study heterogeneity [29]. Sensitivity analysis has been conducted to avoid heterogeneity, which might occur as a result of the total calorie intake modification which includes a follow up procedure and other potential mediators. We used these combined mediators as conjecturers and forecasters of effect in the meta-regression analyses. Those mediators were likely be able to influence the association between SBC and LADA, so we are therefore obliged to adjust all of these mediators to weaken and lessen the effect. We used a visual assessment of the Begg funnel plot and applied the Begg and Egger analysis to evaluate and appraise any possibility for publication bias [30, 31]. Generally case-control studies can study rare diseases which have multiple risk factors for one disease as they are relatively cheap, quick and easy to design due to retrospective recall because of the already existing data. However, this design could not study several diseases, rare exposures or even estimate the incubation period between risk factor of the disease in question, and disease itself. Neither could it measure the risk directly nor even the occurrence rates including the incidence and prevalence of the same. Relative risk could not be calculated but the Odds ratio could be. Thus given everything mentioned above, in terms of strength of association, case control studies showed the same strength as cohort because both were analytical studies [32].

Results and findings

Characteristics of all the study population included within our meta-analyses are presented in Table 1. Each research study assessed the risk of SBC in development of LADA (nine data points) [20–25], comprising males and females of the Caucasian population from Sweden and all of whom were adults. Regarding all the selected case-control studies, each of which were compared via a retrospective recall of previous exposure to the risk factor, which was sugary beverages consumption, including coffee and alcohol. Cases were matched with controls in relation to number of participants and demographic characteristics and was conducted by interviews and structured questionnaires which included food frequency questionnaires (FFQs). There were 15027 participants involved with and 1862 patients with LADA. The research articles assessed the dietary intake [20–25] revealed effect estimations that were not adjusted for total calorie intake or measures of BMI. According to the data from those six articles, the shared odds ratio [OR] for LADA was 1.37 [95% CI 1.23–1.52]]. Overall P value 0.001, I² = 73.1%, for the
<table>
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<th>Ref.</th>
<th>Population</th>
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<th>Study design</th>
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<th>Conclusion</th>
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<tr>
<td>Löfvenborg et al. 2019 [20]</td>
<td>Sweden 386 LADA cases 1545 LADA controls 1253 type 2 diabetes mellitus cases</td>
<td>40–69</td>
<td>Observational analytical Case-control study</td>
<td>Diet history</td>
<td>High sweetened beverage intake</td>
<td>Age, sex, BMI, family history, smoking and education</td>
<td>OR (95% CI) between extreme quartiles of median SBC (0 vs. 143 g/day): 1.67 (0.98–2.87); P trend &lt; 0.01</td>
<td>High sweetened beverage intake encompasses autoimmune forms of diabetes</td>
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<td>Rasouli et al. 2014 [21]</td>
<td>Sweden 250 LADA cases 1012 LADA controls 764 type 2 diabetes mellitus cases</td>
<td>45–64</td>
<td>Observational analytical Case-control study</td>
<td>FFQs</td>
<td>Coffee intake</td>
<td>High-risk HLA genotypes, age, sex, BMI, family history, smoking and education</td>
<td>Men: OR (95% CI) between extreme quartiles of SBC (&gt; 1–8-oz serving/day vs. &lt; 2–8-oz servings/day): 1.09 (0.89–1.33); P trend &lt; 0.68 Women: OR (95% CI) between extreme quintiles of SBC: 1.17 (0.94–1.46); P trend &lt; 0.05</td>
<td>Coffee intake is positively associated with LADA among carriers of high-risk HLA genotypes</td>
</tr>
<tr>
<td>Löfvenborg et al. 2014 [22]</td>
<td>Sweden 245 LADA cases 990 LADA controls 759 type 2 diabetes mellitus cases</td>
<td>38–65</td>
<td>Observational analytical Case-control study</td>
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<td>Intakes of sweetened beverages</td>
<td>Genetic susceptibility conferred by genotypes of HLA, FTO or TCF7L2</td>
<td>OR (95% CI) between extreme quartiles of SBC (&lt; 1 serving/month vs. &lt; 1 serving/day): 1.83 (1.42, 2.36); P trend &lt; 0.001</td>
<td>High intakes of sweetened beverages increase the risk of both LADA and type 2 diabetes mellitus</td>
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<tr>
<td>Rasouli et al. 2012 [23]</td>
<td>Sweden 140 LADA cases 1841 type 2 diabetes mellitus cases</td>
<td>42–57</td>
<td>3 cross-sectional surveys</td>
<td>68 item FFQs</td>
<td>Alcohol consumption</td>
<td>Age, sex, race, education, center, total calories, smoking, physical activity, intake of meat, dairy, fruits and vegetables, whole grains, and refined grains</td>
<td>OR (95% CI) between extreme quintiles of SBC (0 vs. &gt; 1 serving/day): 0.15 (0.12–1.42); P trend &gt; 0.65</td>
<td>Alcohol consumption may improve insulin sensitivity and reduce the risk of type 2 diabetes LADA</td>
</tr>
<tr>
<td>Löfvenborg et al. 2016 [24]</td>
<td>Sweden 357 LADA cases 1136 LADA controls 1371 type 2 diabetes mellitus cases</td>
<td>39–63</td>
<td>Observational analytical Case-control study</td>
<td>FFQs</td>
<td>Coffee</td>
<td>BMI, physical activity, family history of diabetes, postmenopausal hormone use, alcohol use, smoking, and total energy intake physical activity, family history of diabetes, smoking, postmenopausal hormone use, oral contraceptive use, cereal fiber, magnesium, trans fat</td>
<td>OR (95% CI) between extreme quintiles of SBC: (&gt; 1–12-oz serving/month vs. 2–3 12-oz servings/day): 1.31 (0.99–1.74); P trend &lt; 0.001</td>
<td>Coffee may promote autoimmunity and possibly even increase the risk of autoimmune diabetes</td>
</tr>
<tr>
<td>Rasouli et al. 2018 [25]</td>
<td>Sweden 484 LADA cases 1609 LADA controls 885 type 2 diabetes mellitus cases</td>
<td>37–67</td>
<td>Observational analytical Case-control study</td>
<td>FFQs</td>
<td>Alcohol consumption</td>
<td></td>
<td>OR (95% CI) between extreme quintiles of SBC (&lt; 1 12-oz serving/month vs. &gt; 2–12-oz servings/day): 1.24 (1.06–1.45); P trend &lt; 0.002</td>
<td>Moderate alcohol consumption reduces risk of both type 2 diabetes and autoimmune diabetes</td>
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difference between extreme quantiles of SBC indicates an additional risk of 26% related with increased SBC.

Even though all researches apart from two [21, 23] exhibited positive associations and significant correlations between SBC and LADA, there was considerable heterogenic difference among them in the analysis, where the P value was calculated for the test of heterogenic difference for LADA: 12–66% (95% CI 31–83), however P value, was 0.003. Rasouli et al. [23] performed as a cross sectional study survey assessing LADA and expressed non-significant negative association [23]. If we exclude this research from our analysis, the heterogenic difference will be reduced slightly, 12–62% (95% CI 17–82), however the P value, was 0.01. On the other hand, the remaining studies showed clear significant positive association, except for one study [21] which also provided clear non-significant negative association between alcohol and LADA among men. However, it did conclude that alcohol could be a potential protective factor against LADA in women, due to the significant negative correlation (P < 0.68 for men [21] P < 0.05 for women [21] P < 0.65 [23]). In spite of this condition, findings from the meta-regression analysis did not find that the noted variation in either study [21, 23] to make a significant difference.

The case control study by Löfvenborg et al. [22], which displays a marginal significant positive correlation, has the smallest number of contributors and significantly lower amounts of SBC intake comparative to the other studies (median SBC is 143 g/day in uppermost quartile of intake, where intake of one 12-oz serving equal to 336 g). Exclusion of this research from the analysis did not lessen heterogeneity, as was expected, given its low statistical significance and small proportion weight (P value, test for heterogenic difference 0.002). However the research done by Rasouli et al. [25], which had the biggest significance and which used frequent measures of SBC, described the robust estimation. Exclusion of this research from the mutual analysis has decreased heterogenic difference to a marginal significance (P value, test for heterogenic difference 0.05; I2 51% [95% CI 0–78%]).

Assessments for publication bias usually depend on the supposition that a few studies with big variabilities may be more susceptible to publication bias, in comparison to large research studies. A visual review of the Begg funnel plot (accompanying Fig. 2), where by the standard error of log the relative risk (putting in consideration the study size) from each research was strategized against the log relative risk (effect of treatment), exhibited balance about the plot, suggestive of an impossible bias of publication, even though values for LADA may not be mostly helpful and informative because of the small number of studies comprised within the analysis. Studies with a big standard of error and great effect may recommend the existence of what is called “a small-study effect”. In other words, the propensity of small research studies in the meta-analysis to show the big treatment effects (P value for LADA was 0.75 in the studies of both Begg and Egger [28, 30, 31], Fig. 2).

Findings from our analysis of sensitivity in which both calorie, and BMI, adjusted coefficients were omitted [20, 22] revealed a slight escalation in risk of LADA with a pooled relative risk of 1.28 and 95% CI (1.13–1.45). This is with regards to the random-effects-model and on the other hand, relative risk of 1.250 (1.18–1.34) regarding the fixed-effects-model. There was a larger increase which was distinguishable in the dose-response-meta-analysis and when we excluded those studies [20, 22]: relative risk was 1.350 (1.14–1.59) and this regarding the random-effects-model. On the other hand, the relative risk of 1.180 (1.12–1.24) with regards to the fixed-effects-model. However findings from the meta-regression did not adjust for calorie intake as it was not considered to be an important mediator of outcome (P = 0.380). Further analysis of sensitivity was not conceivable for studies of LADA because they are too scarce and yet, both studies that did adjust for those mediators of outcome had borderline insignificant associations [20, 22], whilst the research that had shown unadjusted estimations also showed a significant positive correlation [21].

**Discussion and conclusion**

As can be deduced from the presented meta-analysis there is a clear association between SBC and risk of LADA. This is based upon the coefficients from five case-control studies and one cross sectional survey, which involved 15027 contributors and 1862 patients with LADA. Contributors in the uppermost group of SBC
intake had a 20% more risk of developing LADA than those in the lowermost group of SBC intake.

Since we matched the extreme quantiles of SBC, mostly zero or one serving per month against one or two servings per day, groups of intake between the studies were not consistent or homogenous. Consequently, it is likely that a random biased classification fairly weakened the mutual estimation; though, findings were analogous to the dose-response-analysis, which used data from all groups. For those studies that did not outline a size of serving, an average serving of (12 Oz) was presumed, which may overestimate or undervalue the experiential SBC levels but ought not to substantially disturb our results. Certainly there is considerable difference in study designs and assessment of exposure, through the studies, which may elucidate the observable notch between heterogenic differences in studies we perceived. Meta-analysis is integrally not as strong as the distinct prospective cohort research but it is still beneficial in providing a holistic view about the effect size. Moreover, they also provide larger investigations and studies with less random disparities and more weight than the smaller studies. Publication bias is always seen as a prospective apprehension especially with meta-analyses. Nevertheless standard assessments and visual scrutiny of the funnel plots garnered no proof of any publication bias in our analysis.

All the research that was involved in our meta-analysis included well-thought-out adjustments for possible confusion by several factors such as diet and lifestyle, and mostly due to the persistence of positive association, signifying an autonomous effect of SBC. High levels of SBC could be indicator of a generally unhealthy diet as they lean towards the inclusion of other factors such as, ingestion of high saturated and trans-fatty acids and a low fiber intake [12]. So, an imperfect adjustment for several diet and lifestyle factors could possibly overstate the strength of the positive correlation between SBC and risk of LADA. However, the consistency of results from these different studies decreases the probability that an enduring variable is responsible for the results. Longitudinal studies assessing diet and the risk of chronic disease may similarly be exposed to inverse causality, i.e., persons alter their diet due to subclinical disease symptoms or associated obesity, which may result in false associations [26]. Though it is not imaginable to totally remove these factors, studies with long periods of follow up and frequent measurements of nutritional intake have a tendency to be less susceptible to this process.

In a few studies, LADA was evaluated by self-assessment; yet, it has been demonstrated in confirmation studies that self-reporting of LADA is highly precise according to the review of medical records [26]. The bulk of research studies have used validated Food Frequency Questionnaire to assess SBC, which is the strongest technique for assessing a personal average dietary consumption associated with other valuation methods such as the 24 hour dietary recall [27]. However, errors of measurement in dietary assessment are always unavoidable, but because the studies we deliberated are case-control in design, faulty classifica-
tion of SBC perhaps does not vary by case status. This non-differential faulty classification of exposure may undervalue the real association between SBC and risk of such consequences.

SBC are thought to cause obesity due to their high supplementary sugar content, low compensatory water intake, reduced satiety and inadequate compensatory reduction in calorie intake during meals which causes a positive energy balance and thus the body stores the extra food as fats [7, 8]. Even though, SBC increases the risk of LADA, partially due to their participation in weight gain, an autonomous effect may also come from the increased amounts of fast absorbable carbohydrates as extra sugars, used as flavors in beverages. The results by Löfvenborg et al. [20] suggested that nearly half of the consequences of SBC on LADA were arbitrated through obesity. In a recent longitudinal research which followed 88,000 females for 24 years, who were consuming 2 servings per day and had a 34% more risk of coronary insufficiency in comparison to occasional consumers after adjustment for other potential mediators (relative risk 1.35, and 95% CI of 1.1–1.7, where P-value < 0.01) [33]. Further adjustment of potential mediators like BMI and total calorie intake, weakened the associations, however they were still statistically significant, indicating that the effect of SBC is not fully mediated by those factors. SBC has been proven to increase blood sugar and insulin levels quickly and intensely [34] and if frequently used in big quantities, will undoubtedly lead to a high dietary glycemic load. High glycemic load (GL) nutrition will increase the risk of development of diabetes [5]. Interim experimental researches recommend that fructose, which is an essential component of fructose/sucrose corn syrup, can lead to predominantly metabolic adverse effects when compared with glucose. This is because fructose is otherwise metabolized to lipids inside the liver, causing increased biochemical processes of creating fatty acids from acetyl-CoA that are formed from a number of different mechanisms within the hepatic cell, leading to high levels of triglycerides, decreased high density lipoproteins, development of dyslipidemia, and also insulin resistance [38]. In brief, this meta-analysis has not shown that excessive SBC is associated with the risk of development of LADA. It offers moderate evidence to support the restricted intake of these drinks and the use of healthy substitutes instead like water to which will decrease the risk of chronic diseases. Nevertheless, there were no long term randomized studies, with sufficient follow-up period, to show potential relationship between sugary beverages and LADA, and probably such studies will never be conducted due to ethical reasons. So, no definite conclusions could be drawn due to heterogeneous data from low quality researches and the analysis was based on observational and case-control studies only. So the authors were urged to elaborate on the fact that no definite conclusions could be drawn.

Summary

Only six research papers worldwide addressed the relation between consumption of sugary beverages and the development of latent autoimmune diabetes in adults. These articles had contradictory findings regarding this risk factor. We conducted a systematic review and meta-analysis to establish statistical significance across studies that might otherwise seem to have conflicting results. This will increase the validity and reliability of information and any observed differences. This is the first systematic review and meta-analysis comparing this correlation, to find a clear significant association between sugary beverages consumption and latent autoimmune diabetes in adults.

Ethical approval and consent to participate

The authors certify that the guidelines of the PRISMA 2009 statement have been adopted.

Competing interests and conflict-of-interest statement

The authors certify that they have no conflicts of interest to declare including but not limited to financial, consultancy, advisory, institutional, and other relationships that might lead to a possible bias or misconstrue the results and/or conclusions of this research.

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