Increased sugar consumption is increasingly considered to be a contributor to the worldwide epidemics of obesity and diabetes and their associated cardiometabolic risks. As a result of its unique metabolic properties, the fructose component of sugar may be particularly harmful. Diets high in fructose can rapidly produce all of the key features of the metabolic syndrome. Here we review the biology of fructose metabolism as well as potential mechanisms by which excessive fructose consumption may contribute to cardiometabolic disease.
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Introduction

Glucose is the predominant form of circulating sugar in animals, while sucrose, the disaccharide composed of equal portions of glucose and fructose, is the predominant circulating sugar in plants. As plants form the basis of the food chain, herbivores and omnivores are highly adapted to use sucrose for energetic and biosynthetic needs. Because fructose does not circulate at high levels in animals, ingested fructose may be uniquely positioned to convey signals related to sugar consumption. Therefore, understanding mechanisms by which fructose is sensed may be of consequence for understanding the adaptive physiology of sucrose metabolism as well as potential pathophysiological consequences of excessive sugar consumption.

Sugar in the form of sucrose or high-fructose corn syrup, both of which are composed of nearly equal amounts of glucose and fructose, is added to numerous manufactured food products. Sugar-sweetened beverages (SSBs) are a major source of added sugar in diets worldwide and include sodas, fruit-flavored drinks, and sport drinks. On average, SSBs contribute approximately 7% of daily calories (1) and nearly 50% of added sugars in the diet (2). Although trends in SSB consumption have declined in recent years, almost 66% of US youths still consume at least one SSB per day (3). Other major contributors to added sugar intake include candy and desserts, contributing approximately 4% to 9% of daily energy intake depending on age (2, 4).

Whether increased sugar consumption is a major contributor to the epidemics of obesity, type 2 diabetes, and nonalcoholic fatty liver disease remains controversial (5–7). While the relationships between some measures of dietary sugar exposure and cardiometabolic risk factors are inconsistent, greater SSB consumption consistently associates with indices of higher cardiometabolic risk (5). Several large meta-analyses associate increased SSB consumption with increased body weight, and much, though not all, of this increased weight is likely due to increased total energy consumption (5, 8). SSBs may increase cardiometabolic risk by increasing visceral adiposity, which accounts for much of the weight gain. A recent prospective study showed that daily SSB consumers had a 29% greater increase in visceral adipose tissue volume over 6 years compared with nonconsumers (9). A causal association is supported by evidence that intake of 1 liter of SSB daily for 6 months increased visceral and liver fat, but increases were not observed in those consuming isocaloric semiskim milk, noncaloric diet soda, or water (10). While increased visceral adiposity is a major cardiometabolic risk factor, SSBs may increase risk independently of adiposity. For instance, daily SSB consumption is associated with an unhealthy metabolic profile across BMI strata and with increased risk for type 2 diabetes independently of obesity (11, 12).

Hypertriglyceridemia is a major cardiovascular risk factor and is another mechanism by which SSBs might increase cardiovascular risk. Few large cross-sectional studies have examined the risk of dyslipidemia with SSB intake, and these studies show that dyslipidemia prevalence increases with higher SSB intake (13, 14). One prospective study reported that consuming more than 1 soft drink per day increased the odds of developing hypertriglyceridemia by 25% over 4 years compared with consuming less than 1 soft drink per day (15). Moreover, two recent prospective cohort studies showed that daily SSB consumption was associated with approximately 25% greater risk of developing coronary heart disease in both men and women compared with nonconsumers (13, 16).

SSB consumption also associates with hypertension, another major cardiovascular risk factor. A recent meta-analysis found a modest 12% increase in hypertension risk among the highest SSB consumers compared with the lowest (17). Thus, SSB intake may contribute to hypertension, but it may play a lesser role in this risk factor compared with other cardiometabolic risk factors.

On the basis of short-term overfeeding studies conducted predominantly in animals, the fructose component of SSBs and added sugar appears to be particularly harmful. Feeding animals large amounts of fructose can rapidly produce multiple features of the metabolic syndrome, including obesity, dyslipidemia, fatty liver, hypertension, insulin resistance, and diabetes (18, 19). Some, but not all, short-term dietary intervention studies in humans also demonstrate that overfeeding fructose, but not glucose, can increase visceral adiposity, postprandial hypertriglyceridemia, and insulin resistance, and effects on specific traits may be impacted by gender (20, 21). One concern with such studies is that the amount
of fructose consumed often exceeds that commonly found in ad libitum diets. The average consumption of fructose in US populations accounts for approximately 9% of total energy intake, while consumers in the 95th percentile average approximately 15% of total energy from fructose (22). In contrast, many interventional studies are short in duration (less than 4 weeks) and include dietary intakes closer to 25% of total energy intake from fructose (23, 24).

Large randomized controlled dietary intervention studies assessing the effects of added sugars on cardiometabolic risk factors over long periods of time are lacking. Complexity, cost, compliance, and potential ethical issues likely prohibit the conducting of such studies. Nevertheless, some short-term interventional studies, even those within the range of “normal” fructose consumption, show that fructose can rapidly impair intermediate physiological endpoints like circulating lipids and insulin sensitivity in humans (25). Several recent reviews comprehensively discuss the physiological effects of added fructose or sugar on pathophysiological endpoints in human subjects (26, 27).

Understanding the mechanisms by which the isolated monosaccharide fructose might contribute to the development of metabolic disease may provide fundamental insights into pathogenic mechanisms that can be used to develop new diagnostic, preventative, and therapeutic strategies. Here we will review the biochemistry and molecular genetics of fructose metabolism as well as potential mechanisms by which excessive fructose consumption contributes to cardiometabolic disease. We hope that lessons learned from improved understanding of fructose metabolism and fructose-induced cardiometabolic risk may also apply to other forms of diet-induced and genetically induced metabolic disease.

Fructose absorption
Ingested fructose is predominantly absorbed passively from the intestinal lumen via the hexose transporter SLC2A5, also known as GLUT5, which has high affinity for fructose ($K_m = 6$ mM). GLUT5 is highly expressed on enterocytes’ luminal membrane and is also expressed basolaterally (28). Deletion of Glut5 in mice reduces fructose absorption by 75% and causes cecum and colon dilatation as well as gas accumulation (29). These features are suggestive of fructose malabsorption, frequently cited as a cause of gastrointestinal symptoms in humans (30, 31). The intestine’s capacity to absorb fructose is saturable (32), and a healthy adult’s ability to absorb free fructose ranges from less than 5 g to more than 50 g (33). Unabsorbed fructose can impose an osmotic load on the distal small intestine and the colon, which may contribute to gastrointestinal symptoms (32). Moreover, fructose can serve as a substrate for bacterial fermentation, leading to formation of gas and other bacterial metabolites, which can affect intestinal motility and cause various symptoms such as abdominal pain and bloating (34).

Intestinal GLUT5 mRNA levels and fructose transport rates are very low prenatally and rapidly increase with weaning independently of diet, but they can be further induced following weaning to diets containing fructose (35). Recent data showed that high-fructose feeding induces intestinal thioredoxin-interacting protein (TXNIP), which binds and regulates GLUT5-mediated intestinal fructose transport (36). Consistent with this, we recently showed that carbohydrate-responsive element-binding protein (ChREBP), a transcription factor that responds to intracellular carbohydrate nutrients and a known transcriptional regulator of TXNIP (37), also regulates intestinal GLUT5 expression and is required for systemic fructose tolerance (38). In the future, it will be interesting to determine whether variability in the expression or function of GLUT5 or its regulatory factors contributes to the variability in fructose absorption in humans.

Intermediary fructose metabolism
Fructose concentrations in peripheral plasma are typically about 0.04 mM, can acutely increase 10-fold after fructose consumption, and return to fasting levels within 2 hours (39–41). This rapid clearance is mediated in large part by efficient extraction by the liver. Whereas the liver extracts only 15% to 30% of an oral glucose load, it is capable of extracting 70% of an oral fructose load (42, 43). Following fructose ingestion, plasma fructose can achieve low millimolar concentrations in the portal vein accompanied by peripheral circulation levels of approximately 0.2 mM, indicating that peripheral fructose concentrations rarely exceed the high micromolar range (44).

The SLC2A2 glucose transporter, also known as GLUT2, has lower affinity for fructose ($K_m = 11$ mM) than GLUT5 (45). GLUT2 is a minor contributor to intestinal fructose transport (45), whereas it is likely a major contributor to hepatic fructose uptake, since GLUT5 is not robustly expressed in the liver (46, 47). SLC2A8, also known as GLUT8, may also contribute to hepatocellular fructose transport (48). Fructose is a poor substrate for the hepatic hexokinase glucokinase (GCK). Instead, ketohexokinase (HKH, also known as fructokinase) rapidly phosphorylates fructose to generate fructose-1-phosphate (F1P). HKH’s high activity and insensitivity to cellular energy status account for the liver’s ability to efficiently extract fructose. F1P is metabolized to dihydroxyacetone phosphate (DHAP) and glyceraldehyde 3-phosphate (G3P), which enter the glycolytic/gluconeogenic metabolite pools (Figure 1).

Cellular metabolic status and energy status tightly regulate the phosphofructokinase (PFK) step in glycolysis, which limits hepatic glycolytic flux (49). In contrast, fructose-derived metabolites enter the triose-phosphate pool distal to PFK and therefore bypass this restriction. As hepatic fructolysis is unrestricted, fructose loads can lead to large, rapid expansions in the hexose- and triose-phosphate pools, potentially providing increased substrate for all central carbon metabolic pathways, including glycolysis, glycogenesis, gluconeogenesis, lipogenesis, and oxidative phosphorylation.

The disposition of fructose-derived carbon among the major metabolic pathways depends on the overall nutritional and endocrine status of the animal as well as the status of key regulatory checkpoints in intermediary metabolism. For instance, in starved animals, low levels of fructose-2,6-biphosphate inhibit PFK activity and glycolysis and activate fructose-1,6-biphosphatase and glucose production (50). Thus, in a starved animal, fructose-derived triose-phosphates are preferentially routed through the gluconeogenic path (51, 52). The fate of ingested fructose may also depend on coinherited nutrients. For instance, infusing physiological concentrations of fructose to fed rats and humans increases serum glucose and lactate levels without affecting hepatic glycogen accumulation (53, 54). However, when fructose is infused with glucose, which stimulates insulin secretion, marked glycogen accumulation occurs (55). Chronic fructose consumption can affect metabolic
glucose uptake and phosphorylation, leading to rapid glycogen accumulation (66). F1P may also enhance glycogen synthesis by allosterically inhibiting glycogen phosphorylase (67, 68). Lastly, F1P also allosterically activates pyruvate kinase, the terminal step in glycolysis, contributing to increased circulating lactate levels following fructose ingestion (69). In rodent liver, hepatic F1P levels increase 10-fold to approximately 1 mM within 10 minutes after fructose ingestion and remain elevated for several hours (70). F1P concentrations of only approximately 200 μM are sufficient to allosterically inhibit glycogen phosphorylase (GCK) (71). Thus, fructose ingestion is likely to have rapid, robust, and sustained effects on hepatic glucose uptake and intermediary metabolism.

While the efficiency and rapidity with which the liver can extract and phosphorylate ingested fructose are likely important for its role in integrating nutritional and systemic fuel metabolism, this robust metabolism may also have deleterious consequences. For instance, decreases in intracellular free phosphate due to rapid hepatic fructose phosphorylation can increase uric acid production through activation of AMP deaminase, which leads to catabolism of AMP to uric acid (72, 73). Fructose feeding may also

Figure 1. Fructose biochemistry. Upon entering hepatocytes, fructose is phosphorylated by KHK to F1P. F1P is cleaved to DHAP and glyceraldehyde by ALDOB. Glyceraldehyde is phosphorylated by triose-kinase (TKFC, also known as dihydroxyacetone kinase 2 or DAK) to form the glycolytic intermediate glyceraldehyde 3-phosphate (GA3P). Both fructose-derived DHAP and GA3P enter the glycolytic/gluconeogenic metabolite pool at the triose-phosphate level, and these metabolites have numerous metabolic fates. F1P also allosterically regulates metabolic enzymes (red and green lines) to regulate the disposition of fructose-derived substrate and other metabolic products like uric acid. AMPD3, adenosine deaminase; GA, glyceraldehyde; IMP, inosine monophosphate; MTTP, microsomal triglyceride transfer protein; PYGL, glycogen phosphorylase L; GYS2, glycogen synthase 2; PKLR, pyruvate kinase, liver and red blood cell; PEP, phosphoenolpyruvate; TAG, triacylglycerol.
stimulate purine synthesis, contributing to uric acid production (74). Increased circulating uric acid levels increase the risk of gout, a condition characterized by painful inflammation due to deposition of uric acid crystals in joints. Indeed, a growing body of evidence implicates sugar intake as a risk factor for gout (75). Moreover, elevated serum uric acid levels and gout are associated with other cardiometabolic risk factors in diverse populations (76–78). A substantial body of work suggests that increased uric acid levels may independently regulate important aspects of metabolism and contribute to cardiometabolic risk (79–83). However, Mendelian randomization studies do not strongly support a causal role for nutrient consumption and integrating peripheral nutrient status to regulate systemic fuel storage versus provisioning. While hormones like insulin and glucagon help inform the liver of systemic fuel status, the liver is also well configured to integrate signals from insulin and glucagon; these metabolic pathways contribute to steatosis, VLDL packaging and secretion, as well as glucose production and the generation of lipid intermediates that may affect hepatic insulin sensitivity and other biological processes. ACACA, acetyl-CoA carboxylase α; FASN, fatty acid synthase; GPAT, glycerol-3-phosphate acyltransferases; AGPAT, acylglycerol-3-phosphate acyltransferase; DGAT, diacylglycerol acyltransferase; DAG, diacylglycerol.

Genetic lessons about fructose metabolism

KHK exists as two alternatively spliced isoforms produced by mutual exclusion of the adjacent exons 3C and 3A within the KHK gene (87, 88). The “A” isoform is ubiquitously expressed but has low activity due to relatively low affinity for its substrate (Kₘ = 8 mM) (89). Expression of the “C” isoform is primarily restricted to metabolic tissues including the liver, kidney, and intestine, and this isoform has much higher affinity for fructose (Kₘ = 0.8 mM) (89, 90). Mice deficient in both isoforms were fully protected from fructose-induced metabolic disease even though blood and urinary fructose levels were markedly increased (91). Thus, elevated blood fructose itself is not deleterious; rather, fructose metabolism is essential for fructose-induced metabolic disease. Loss-of-function mutations in KHK cause the benign human disorder essential fructosuria, characterized by impaired hepatic fructose metabolism leading to high blood and urine fructose levels after sucrose or fructose consumption (92). Consistent with observations in mice, there are no documented adverse health effects observed in people with this condition. Altogether, these results suggest that inhibiting KHK may be a safe therapeutic strategy to prevent fructose-induced metabolic disease.

In contrast with global KHK deletion, selective deletion of the A isoform exacerbates the adverse metabolic effects of fructose feeding (91). These results suggest two important hypotheses: (a) fructose metabolism outside of tissues that express the C isoform is non-negligible and contributes to whole-body fructose clearance, and (b) fructose metabolism within the tissues expressing KHK-C is critical for fructose-induced metabolic disease. This is supported by recent data showing that selective knockdown of KHK in mouse liver protects against fructose-induced steatosis (93). Recent data also indicate that altered splicing between KHK-A and KHK-C isoforms may contribute to the development of distinct diseases like hepatocellular carcinoma and heart failure (94, 95).

Hereditary fructose intolerance (HFI) is a rare autosomal recessive disease caused by a deficiency of aldolase B (ALDOB),
which is highly expressed in the liver, kidney, and small intestine (96). People with HFI develop abdominal pain, vomiting, diarrhea, symptomatic hypoglycemia, hyperuricemia, and potentially liver failure and death following ingestion of foods containing fructose, sucrase, or sorbitol (97). The precise mechanisms by which ALDOB deficiency causes symptoms are not entirely clear. An Aldob-deficient mouse model mimics the human HFI condition (98). These mice fail to thrive and die when exposed to high-fructose diets. Interestingly, even on a fructose-free diet, Aldob-deficient mice develop steatosis (98), possibly due to impaired metabolism of endogenously synthesized fructose (99).

Endogenous fructose production
While the vast majority of metabolized fructose is derived from dietary sources of sugar, animals including humans are capable of synthesizing fructose endogenously. The sorbitol (polyol) pathway, which is active in a wide range of tissues, is responsible for endogenous fructose formation from glucose (100, 101). In this pathway, glucose is first reduced to sorbitol by aldose reductase (102). Sorbitol is then oxidized to fructose by sorbitol dehydrogenase (103). Physiologically, endogenously synthesized fructose is the primary energy source for sperm and may be important for fertility (104–106). The placenta may also synthesize sorbitol that the developing fetus may use to synthesize fructose, suggesting a broader role for endogenous fructose in reproductive and developmental biology (107).

Sorbitol pathway activity increases during diabetic hyperglycemia (108). Endogenous fructose synthesis and polyol metabolites are considered key players in the development of diabetic microvascular complications (109). Interestingly, semen fructose concentrations are increased in type 1 diabetes and in obesity, in which it is associated with impaired sperm parameters (105, 110). Whether endogenous fructose synthesis might occur at sufficient rates to contribute to other aspects of fructose-induced cardiometabolic risk has only recently been addressed. Glucose dose-dependently induces aldolase reductase in human tissues, and chronic exposure to a high-glucose diet induces polyol pathway activation in mice (99, 111). This may be a mechanism by which severe hyperglycemia may exacerbate cardiometabolic risks. Additionally, Lanasa et al. report that endogenous fructose production and KHK activation within the kidney contribute to the development of diabetic nephropathy (112). Although sorbitol dehydrogenase is expressed at high levels in human liver (113), whether this pathway is sufficiently active in humans to play an adverse metabolic role will require further investigation.

Fructose effects on lipid homeostasis
As noted above, excessive fructose consumption may have significant effects on lipid metabolism, contributing both to steatosis and to increased circulating triglyceride levels in the form of very low-density lipoprotein (VLDL). Hepatic lipid accumulation results from a combination of increased hepatic de novo lipogenesis (DNL), esterification of preformed fatty acids derived from the diet or adipose stores, decreased VLDL secretion, and decreased hepatic fatty acid oxidation. Activation of the lipogenic program is observed immediately after a single load of fructose and contributes to increased VLDL triglyceride secretion (114, 115). Fructose also acutely suppresses hepatic fatty acid oxidation (116). Thus, fructose contributes to hepatic triglyceride production both by providing substrate for fatty acid and triglyceride synthesis and by activating signaling systems to enhance lipid production (Figure 2).

The liver is the primary site of DNL, the process by which fatty acids are synthesized from dietary precursors, predominantly carbohydrates (117). Due to the differences in hepatic glucose and fructose metabolism, a larger fraction of diet-derived fructose than glucose metabolites are available for conversion to fat in the liver via DNL in animals and humans (20, 118–120). Additionally, fructose metabolites entering the triose-phosphate pool are in equilibrium with glycerol 3-phosphate, which is used to synthesize the glycerol backbone in triglyceride. Moreover, the metabolite malonyl-CoA generated via DNL limits fatty acid oxidation by inhibiting carnitine palmitoyltransferase 1A (CPT1A), the enzyme required for translocation of fatty acids into the mitochondria (121). CPT1A inhibition further increases the availability of fatty acids for triglyceride production. Triglyceride can be incorporated into lipid droplets, leading to steatosis, or can be incorporated into VLDL and secreted from the liver.

In addition to providing substrate for lipogenesis, chronic fructose consumption increases transcriptional regulation of DNL by activating key transcription factors, including sterol regulatory element–binding protein 1c (SREBP1c) and carbohydrate-responsive element–binding protein (ChREBP) (122). SREBP1c promotes lipid synthesis and is regulated at the transcriptional and posttranslational levels by nutrients and hormones. Insulin is a major hormonal activator of hepatic SREBP1c (123, 124). Although acute fructose feeding does not directly stimulate insulin secretion, chronic fructose ingestion can lead to hyperinsulinemia, which may increase hepatic SREBP1c expression and activation (125, 126). Fructose may also activate SREBP1c independently of insulin, since SREBP1c responds to high-fructose feeding in liver-specific insulin receptor–knockout (LIRKO) mice (125). Fructose consumption may also promote ER stress, which may induce proteolytic cleavage of SREBP1c and the lipogenic program (127, 128). Fructose-induced ER stress may also enhance lipogenesis via activation of the transcription factor x box-binding protein 1 independently of other lipogenic transcription factors (129).

ChREBP couples carbohydrate metabolites to lipid synthesis by inducing enzymes required for DNL (130). ChREBP may also suppress fatty acid oxidation by downregulating enzymes like CPT1A, in part by antagonizing peroxisome proliferator–activated receptorα (PPARα), a key transcriptional regulator of the fatty acid oxidation gene program (131, 132). ChREBP is highly expressed in key metabolic tissues, including liver, adipose tissue, small intestine, pancreatic islets, and kidney, where it regulates carbohydrate metabolism in an insulin–independent manner (37, 125, 130). The observation that ChREBP-deficient mice are intolerant to diets containing fructose but not to diets containing dextrose suggests a specific role for ChREBP in regulating fructose metabolism (37, 133). Moreover, ChREBP activity was markedly higher in rats fed high-fructose compared with isocaloric high-glucose diets (126). We recently demonstrated that ingesting fructose, but not glucose, acutely and robustly induces hepatic expression of the potent ChREBP-β isoform along with its lipogenic, fructolytic, and glycolytic targets (133, 134). The mechanism by which sugar metabolism.
metabolites activate ChREBP remains controversial but involves allostERIC activation by glucose-6-phosphate as well as modulation by other carbohydrate metabolites and postranslational modifications (135–137). ChREBP knockdown using antisense oligonucleotides (ASOs) in fructose-fed rats reduced circulating triglyceride levels and confirmed a role for ChREBP in fructose-mediated dyslipidemia, although steatosis was unaffected (138). Consistent with this, GWAS have identified multiple common SNPs within the ChREBP locus associated with increased serum triglyceride and low HDL cholesterol levels (139, 140).

ChREBP knockdown’s selective effect on circulating triglycerides but not steatosis in the experiment described above highlights the fact that fat accretion in lipid droplets and VLDL secretion are distinct processes. ChREBP potently regulates DNL, and fructose-induced DNL strongly correlates with fructose-induced hypertriglyceridemia (141). However, in steatotic human subjects, DNL-derived fatty acids contribute a minor fraction of fatty acids to VLDL (142), and the mechanistic connection between DNL and VLDL secretion remains uncertain. Moreover, ChREBP may have effects to increase circulating triglycerides independently of increasing VLDL secretion. ChREBP may transactivate expression of the apolipoprotein APOC3 as well as angiopoietin-like 8 (ANGPTL8), both of which may inhibit lipoprotein lipase and limit VLDL clearance (refs. 143, 144, and Figure 3). Thus, it is possible that high-fructose feeding may increase circulating VLDL both by enhancing VLDL production and secretion and by reducing VLDL clearance, but the precise mechanisms remain to be determined.

PPARγ coactivator 1β (PGC1β) is a transcriptional coactivator that increases the activity of multiple key transcription factors, such as PPARγ, PPARα, estrogen-related receptors (ERRs), and liver X receptor (LXR) (145, 146). PGC1β can also bind SREBP1 and ChREBP and enhance their transcriptional activity (147, 148). ASOs targeting PGC1β prevented SREBP1c expression and lipogenesis, which in turn decreased lipid accumulation in fructose-fed rat livers. PGC1β-targeting ASOs also prevented increases in adiposity, glycemia, and plasma insulin and triglycerides in fructose-fed rats. Thus, PGC1β is uniquely positioned to coordinately regulate both ChREBP and SREBP1c activities in the context of high-fructose feeding.

Increased sugar and fructose consumption is implicated in both simple steatosis and the progression toward more advanced forms of nonalcoholic fatty liver disease (NAFLD), including nonalcoholic steatohepatitis, fibrosis, and hepatocellular carcinoma (149). Important steps in DNL and VLDL synthesis occur at the ER membrane, and fructose-induced lipogenesis may elicit ER stress and the ER stress response (150). Moreover, signaling elements in the ER stress response may contribute to NAFLD pathogenesis and progression (151). Recent work from Zhang et al. suggests that ChREBP may protect the liver against fructose-induced ER stress and hepatic inflammation (152). However, we have recently observed that liver-specific ChREBP-knockout mice do not develop ER stress or hepatic inflammation when challenged with high-fructose diets (38). Mechanisms by which fructose may contribute to progression of NAFLD will require further investigation.

Fructose effects on glucose homeostasis
Fructose does not directly stimulate pancreatic β cell insulin secretion (153, 154). However, high-fructose feeding readily induces hyperinsulinemia in animal models. Moreover, hyperinsulinemia is more pronounced in rodent models with high-fructose compared with high-dextrose feeding despite similar increases in body weight and adiposity (155, 156). Similarly, hypercaloric fructose feeding increases circulating insulin in human subjects (157). Fructose-induced hyperinsulinemia, often considered a proxy for insulin resistance, might be the result of insulin resistance in some combination of liver, muscle, and/or adipose tissue.

The mechanisms by which high-fructose feeding causes hyperinsulinemia and insulin resistance remain uncertain.
Fructose-induced steatosis may contribute to hepatic insulin resistance through increased hepatic diacylglycerol accumulation, PKC activation, and impairment of insulin-mediated Akt2 activation (158–160). However, whether steatosis itself can cause hepatic insulin resistance remains controversial (131, 161). In addition to ChREBP’s role in fructose-induced dyslipidemia, Erion et al. demonstrated that ChREBP knockdown enhanced peripheral insulin sensitivity in high-fructose-fed rats (138). Whether the improvement in peripheral insulin sensitivity was directly related to the improvement in circulating lipid levels or adiposity is uncertain. We recently demonstrated that while hepatic ChREBP is essential for fructose-mediated upregulation of fructolytic, glycolytic, and lipogenic enzymes, ChREBP also mediated upregulation of G6PC, the terminal enzyme in glucose production (133). We showed that a fructose-induced, ChREBP-mediated increase in G6PC activity is a major determinant of endogenous glucose production. Moreover, fructose activated ChREBP and induced G6PC in the absence of FOXO1a, indicating that substrate-driven activation of ChREBP and G6PC to enhance glucose production dominates over the suppressive effects of insulin (133).

This ChREBP/G6PC signaling axis is also conserved in humans. These results are consistent with dietary intervention studies in humans indicating that either eucaloric substitution or hypercaloric addition of fructose may have more significant effects on hepatic insulin resistance than peripheral insulin resistance (157). However, as hyperinsulinemia itself can induce peripheral insulin resistance (162, 163), we speculate that chronic hyperinsulinemia that compensates for fructose-induced glucose production may subsequently lead to peripheral insulin resistance. This hypothesis remains to be tested experimentally.

Effects of fructose on hypertension
The mechanisms by which fructose contributes to the development of hypertension are less well characterized than its effects on glucose and lipid homeostasis. High-fructose feeding in rodents can increase intestinal salt absorption in part through induction of an intestinal anion exchanger, Slc26a6 (188). Moreover, this induction and associated hypertension are prevented by the fact that GLUT5-knockout mice suffer generalized malabsorption and become ill when challenged with fructose. Johnson and colleagues have hypothesized that fructose-induced hyperuricemia may impair kidney function, contributing to hypertension (189). However, as discussed above, genetic data do not strongly support a major role for hyperuricemia in cardiometabolic disease. As fructose is robustly metabolized in the kidney, fructose-mediated changes in renal salt handling may also be important. However, this has yet to be rigorously studied and is an area ripe for further investigation.

Conclusions and future directions
The combination of mechanistic data supporting a role for excessive fructose ingestion and epidemiological data supporting a role for SSBs in the development of cardiometabolic disease supports recent dietary recommendations to limit sugar consumption published by several public health agencies, including the American Heart Association, the World Health Organization, and the Dietary Guidelines Advisory Committee (2, 190, 191).
Safe thresholds for sugar consumption and concrete recommendations for targets to reduce cardiometabolic risk remain in dispute. Moreover, implementing effective programs to alter dietary habits remains challenging. However, initial reports indicate that “sugar taxes” may be effective in reducing SSB consumption (192, 193). Time will tell whether such approaches can improve health outcomes. Hopefully, by improving our understanding of the underlying mechanisms by which sugar and fructose can cause disease, we will be able to bring informed, comprehensive approaches to bear on our current metabolic epidemics.

Acknowledgments
This work is supported by American Heart Association 16CSA28590003 (to MAH and NMM), NIH R01DK100425 (to MAH), NIH ST32HL069772-15 (to DEH), and US Department of Agriculture Agricultural Research Service agreement 58-1950-4-003 (to NMM).

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