Fructose Malabsorption in Systemic Sclerosis

Isabelle Marie, MD, PhD, Anne-Marie Leroi, MD, PhD, Guillaume Gourcerol, MD, PhD, Hervé Levesque, MD, PhD, Jean-François Ménard, MD, PhD, and Philippe Ducrotte, MD, PhD

Abstract: The deleterious effect of fructose, which is increasingly incorporated in many beverages, dairy products, and processed foods, has been described; fructose malabsorption has thus been reported in up to 2.4% of healthy subjects, leading to digestive clinical symptoms (eg, pain, distension, diarrhea). Because digestive involvement is frequent in patients with systemic sclerosis (SSc), we hypothesized that fructose malabsorption could be responsible for intestinal manifestations in these patients.

The aims of this prospective study were to: determine the prevalence of fructose malabsorption, in SSc; predict which SSc patients are at risk of developing fructose malabsorption; and assess the outcome of digestive symptoms in SSc patients after initiation of standardized low-fructose diet.

Eighty consecutive patients with SSc underwent fructose breath test. All SSc patients also completed a questionnaire on digestive symptoms, and a global symptom score (GSS) was calculated.

The prevalence of fructose malabsorption was as high as 40% in SSc patients. We also observed a marked correlation between the presence of fructose malabsorption and: higher values of GSS score of digestive symptoms (P = 0.000004); and absence of delayed gastric emptying (P = 0.0007). Furthermore, in SSc patients with fructose malabsorption, the median value of GSS score of digestive symptoms was lower after initiation of standardized low-fructose diet (P = 0.0009).

Our study underscores that fructose malabsorption often occurs in SSc patients. Our findings are thus relevant for clinical practice, highlighting that fructose breath test is a helpful, noninvasive method by: demonstrating fructose intolerance; identifying the group of SSc patients with fructose malabsorption; and assessing which SSc patients are at risk of developing fructose malabsorption; and assessing the outcome of digestive symptoms in SSc patients after initiation of standardized low-fructose diet.

INTRODUCTION

Systemic sclerosis (SSc) is a systemic inflammatory disorder affecting the skin and other organs, particularly the gastrointestinal tract where lesions may lead to motor activity impairment. Gastrointestinal involvement occurs in 50 to 88% of patients with SSc. Gastric abnormalities are recognized to be associated with high morbidity, resulting in gastric hemorrhage related to bleeding from antral vascular ectasia (“watermelon stomach”) and delayed gastric emptying. Intestinal involvement may also be responsible for severe complications, including hemorrhage due to small intestinal ectasia, malabsorption syndrome, and intestinal pseudo-obstruction.

Recently, the deleterious effect of fermentable saccharides/polyols, especially fructose that is increasingly incorporated in many beverages, dairy products, and canned/baked or processed foods, has been described in non-SSc patients. Previous investigators have thus reported that 2.4% to 16% of healthy subjects exhibited fructose malabsorption. Other authors have further found that fructose malabsorption resulted in abdominal clinical manifestations (ie, pain, discomfort, bloating, distension, nausea, diarrhea) whereas decreased fructose intake led to significant improvement of gastrointestinal symptoms in these patients.

Currently, however, there are no studies in SSc assessing the relationship between food intake and gastrointestinal dysfunction; in particular, the link between fructose malabsorption and the occurrence of gastrointestinal impairment has not yet been documented in SSc. Indeed, the aims of the present prospective study were to: determine the prevalence of fructose malabsorption, using fructose breath test, in unselected patients with SSc; evaluate the correlation between fructose malabsorption and digestive symptoms, nutritional status, and gastrointestinal involvement; make prediction about which SSc patients exhibit fructose malabsorption; and assess the outcome of digestive symptoms in SSc patients after initiation of standardized low-fructose diet.

METHODS

Patients

From January 2011 to June 2014, 80 consecutive patients with a definite diagnosis of SSc, attending a tertiary care center, were included in the study. The criteria used for the diagnosis of
SSc were based on the American College of Rheumatology/European League against Rheumatism classification. Ethical approval was obtained from the local ethical committee (CERNI for the CCP de Haute Normandie), and informed consent was obtained from all patients.

The study cohort consisted of 14 men and 66 women with a median age of 52.5 years (range, 22–79 yr). The median duration of the disease, considered from the onset of the first non-Raynaud’s phenomenon clinical manifestations, was 4 years (range, 1–25 yr). Patients were grouped according to criteria of LeRoy et al.: 23 patients (28.8%) had diffuse years (range, 1–25 yr). Patients were grouped according to non-Raynaud’s phenomenon clinical manifestations, was 4 duration of the disease, considered from the onset of the first median age of 52.5 years (range, 22–79 yr). The median consent was obtained from all patients.

Approval was obtained from the local ethical committee our patients, median values for levels of anxiety and depression (GSS score), calculated as the sum of all symptom scores, was 4 assigned to each patient (maximum score, 33), as described and (60%) had severe esophageal motor impairment. According to Hurwitz’s criteria, 48 patients stage IV: both aperistalsis and decreased low esophageal sphincter pressure and/or CH4 levels above 20 p.p.m.\textsuperscript{18} SIBO when, at least one of the following criteria was present: H\textsubscript{2} and/or CH\textsubscript{4} increase \textgreater\textsubscript{10} p.p.m. on 2 consecutive measurements within the 2 first hours; and H\textsubscript{2} and/or CH\textsubscript{4} increase \textgreater\textsubscript{10} p.p.m. between minimal and maximal values after glucose ingestion.\textsuperscript{18,33} When the glucose breath test was positive, patients received an antibiotic treatment with quinolone or metronidazole for 10 days per month over 3 successive months. A second glucose breath test was then performed. The fructose breath test was only made in patients exhibiting negative control glucose breath test during the next following day.

Glucose Breath Test
SSc patients underwent glucose H2/CH4 breath test as previously described.\textsuperscript{10,18} The test was considered positive for SIBO when, at least one of the following criteria was present: H\textsubscript{2} and/or CH\textsubscript{4} increase \textgreater\textsubscript{20} p.p.m. above basal value; H\textsubscript{2} and/or CH\textsubscript{4} increase \textgreater\textsubscript{10} p.p.m. on 2 consecutive measurements within the 2 first hours; and H\textsubscript{2} and/or CH\textsubscript{4} increase \textgreater\textsubscript{10} p.p.m. between minimal and maximal values after glucose ingestion.\textsuperscript{18,33} When the glucose breath test was positive, patients received an antibiotic treatment with quinolone or metronidazole for 10 days per month over 3 successive months. A second glucose breath test was then performed. The fructose breath test was only made in patients exhibiting negative control glucose breath test during the next following day.

Fructose Breath Test
Patients ingested 25 g of fructose dissolved in 250 cc of sterile water; then, end alveolar breath samples were collected every 30 min for 5 h, as described previously.\textsuperscript{18} Both H\textsubscript{2} and CH\textsubscript{4} levels were calculated. The test was considered positive and defined a fructose malabsorption in case of a rise of H\textsubscript{2} and/or CH\textsubscript{4} levels above 20 p.p.m.\textsuperscript{18} Moreover, during the test, gastrointestinal symptoms were closely monitored and were collected by the laboratory technician every 30 min of the fructose breath test. The occurrence of gastrointestinal symptoms was then analyzed and defined fructose intolerance.\textsuperscript{18}

Comparison of Fructose Breath Test Findings and Other Manifestations of SSc
We compared various characteristics between SSc patients with and without abnormal fructose breath test.

First, digestive manifestations were compared between these groups of SSc patients. Digestive manifestations included: the values of BMI; the values of: GSS score of digestive symptoms; and the presence of cardiometabolic disease, Crohn disease, ulcerative colitis diseases, diabetes mellitus, gastric surgery, or vagotomy. Moreover, no patient received nonsteroidal anti-inflammatory drugs.
Fructose Malabsorption in Systemic Sclerosis

RESULTS

Prevalence of Fructose Malabsorption

In our cohort of 80 SSc patients, the fructose breath test was positive in 32 patients (40%). Furthermore, fructose intolerance, defined as the occurrence of gastrointestinal symptoms during fructose breath test, was found in 25 of 32 patients with SSc (78.1%).

Predictive Factors of Fructose Malabsorption

Digestive Symptoms in SSc Patients

The median value for the GSS score of digestive symptoms was significantly higher in SSc patients with fructose malabsorption than in those without (4 vs. 1; \( P = 0.000004 \)). There were no significant differences between SSc patients with and without fructose malabsorption for: BMI levels \( (P = 0.533) \), scales of anxiety \( (P = 0.267) \), and depression \( (P = 0.148) \) (Table 1).

Esophageal and Gastric Involvement in SSc Patients

As shown in Table 1, patients with normal gastric emptying on \(^{13}\)C octanoic acid breath test more often exhibited fructose malabsorption \( (P = 0.007) \). We failed to show any significant difference between SSc patients with and without fructose malabsorption for severe esophageal motor disorders (stage IV) on manometry \( (P = 0.489) \) (Table 1).

Other Predictive Factors for Fructose Malabsorption

General Clinical Characteristics

We failed to show any statistically significant differences between SSc patients with and without fructose malabsorption regarding: median age \( (P = 0.992) \), sex \( (P = 0.381) \), SSc duration \( (P = 0.487) \), median modified Rodnan score \( (P = 0.710) \); and prevalence of digital ulcers/pitting scars \( (P = 0.650) \), joint involvement \( (P = 0.350) \), ILD \( (P = 0.806) \), pulmonary arterial hypertension \( (P = 1) \) and previous history of renal crisis \( (P = 0.157) \) (Table 2). Additionally, fructose malabsorption tended to be more common in patients with lcSSc than in those with dcSSc, although not significantly so \( (P = 0.076) \). Finally, immunosuppressive therapy did not differ between SSc patients with and without fructose malabsorption for low-dose steroids \( (15.6\% \text{ vs. } 12.5\%; \ P = 0.747) \), methotrexate \( (9.4\% \text{ vs. } 12.5\%; \ P = 0.734) \), azathioprine \( (3.1\% \text{ vs. } 6.3\%; \ P = 0.646) \), and mycophenolate mofetil \( (15.6\% \text{ vs. } 6.3\%; \ P = 0.256) \).

Laboratory Findings

As seen in Table 3, regarding biochemical tests, SSc patients with and without fructose malabsorption did not exhibit different median levels of ESR \( (P = 0.579) \), C-reactive protein levels of ferritin, plasma folic acid, vitamin B12, magnesium/zinc/selenium, and 25 (OH) vitamin D.

Fructose Restrictive Diet

The SSc patients with fructose intolerance were referred to an experienced dietician for a standardized diet adaptation, consisting of a 1-month low-fermentable oligo-saccharides, disaccharides, monosaccharides, and polyol (FODMAP) diet.\(^{16}\) These SSc patients were interviewed, using a standardized questionnaire regarding occurrence of gastrointestinal symptoms, using the GSS score.\(^{5,9,10,33}\)

Dietary compliance was checked by direct interview by the dietician and the physician, 5 weeks after the initiation of dietary changes; compliance was considered adequate if patients confirmed that they adhered to the dietary guidelines during at least 55% of the meals consumed.

Statistical Analysis

For group comparison involving binary data, we used either the \( \chi^2 \) test or Fisher exact test, depending on expected cell count \((\geq 5 \text{ or } < 5\), respectively\). Comparisons involving continuous data were performed using: Student \( t \) test when distribution of variables was normal; and Mann–Whitney \( U \) test in other cases. The results were regarded as significant when the \( P \) value was less than 0.05.

TABLE 1. Digestive Features of SSc\(^{*}\) Patients With Fructose Malabsorption Compared With Those Without

|                          | Fructose Malabsorption (n = 32) | Absence of Fructose Malabsorption (n = 48) | \( P \) |
|--------------------------|---------------------------------|-------------------------------------------|-------|
| GSS score of digestive symptoms | 4 [1–21]                        | 1 [0–9]                                    | 0.000004 |
| Scale of anxiety          | 8 [2–16]                        | 7.5 [2–15]                                 | 0.267 |
| Scale of depression       | 6 [1–16]                        | 2.5 [0–13]                                 | 0.148 |
| Body mass index           | 24.9 [15.8–32]                  | 24 [17.8–34.5]                             | 0.533 |
| Esophageal manometry findings |                              |                                           |       |
| Severe motor esophageal involvement | 56.3%                          | 64.6%                                      | 0.489 |
| \(^{13}\)C octanoic acid breath test findings | | | |
| Absence of delayed gastric emptying | 86.7%                          | 61.9%                                      | 0.007 |

\(^{*}\) GSS of digestive symptoms: global symptom score of digestive symptoms; SSc: systemic sclerosis. Except were indicated, values are median.
(P = 0.827), albumin (P = 0.276), prealbumin (P = 0.455), ferritin (P = 0.941), plasma folic acid (P = 0.444), vitamin B12 (P = 0.115), magnesium (P = 0.835), zinc (P = 0.677), selenium (P = 0.077), and 25 (OH) vitamin D (P = 0.155).

Additionally, autoantibody status did not differ between SSc patients and without fructose malabsorption for anti-centromere (P = 1), anti-Scl 70 (P = 0.447), and anti-RNA polymerase III (P = 0.156) antibodies (Table 3).

### Outcome of Gastrointestinal Symptoms After Low-FODMAP Diet

Complete outcomes were available in all 32 SSc patients with fructose malabsorption who received low-FODMAP diet. Furthermore, 91% of these patients reported adequate dietary compliance. In overall 32 SSc patients, who received low-FODMAP diet, other specific therapy of SSc was unchanged.

As shown in Table 4, after low-FODMAP diet initiation, SSc patients with fructose malabsorption less commonly exhibited: nausea (P = 7.09 × 10⁻⁷), vomiting (P = 0.024), abdominal pain (P = 6.62 × 10⁻⁴), bloating (P = 0.0009), diarrhea (P = 0.022), and abdominal tenderness (P = 0.008). In addition, median value of GSS score of digestive symptoms was significantly lower (4 before vs. 1 after; P = 0.00009) after initiation of low-FODMAP diet in SSc patients.

### DISCUSSION

Fructose is a 6-carbon monosaccharide molecule that is found in fruits and vegetables; furthermore, it is increasingly incorporated in processed foods. Fructose malabsorption has been shown to be responsible for gastrointestinal symptoms. In clinical practice, fructose breath test has been reported as a helpful and reliable method for

---

**TABLE 2. Clinical Characteristics of SSc Patients With Fructose Malabsorption Compared With Those Without Fructose Malabsorption**

| Fructose Malabsorption (n = 32) | Absence of Fructose Malabsorption (n = 48) |
|---------------------------------|------------------------------------------|
| **Clinical parameters**         |                                          |
| Age (yr)                        | 54 [range, 27–79]                        | 53 [range, 22–79] |
| Sex: male/female                | 12.5% males/87.5% females                | 29.7% males/70.3% females |
| SSc duration (yr)               | 5 [range: 1–17]                         | 4 [range: 1–25] |
| SSc subset                      | 78.1% of lcSSc/21.9% of dcSSc            | 66.7% of lcSSc/33.3% of dcSSc |
| Modified Rodnan score           | 4 [range: 15.8–32]                      | 4.5 [range: 17.8–34.5] |
| Digital ulcers/pitting scars    | 50%                                      | 56.3% |
| Joint involvement               | 31.3%                                    | 43.8% |
| Intestinal lung disease         | 28.1%                                    | 33.3% |
| Pulmonary arterial hypertension | 6.3%                                     | 4.2% |
| Previous history of renal crisis| 6.3%                                     | 0% |

* dcSSc: diffuse cutaneous SSc; lcSSc: limited cutaneous SSc; SSc: systemic sclerosis. Except where indicated, values are median.

---

**TABLE 3. Biochemical Characteristics of SSc Patients With Fructose Malabsorption Compared With Those Without Fructose Malabsorption**

| Fructose Malabsorption (n = 32) | Absence of Fructose Malabsorption (n = 48) |
|---------------------------------|------------------------------------------|
| **Biochemical parameters**      |                                          |
| ESR (mm/hour)                   | 10 [range: 1–32]                        | 8.5 [range: 1–89] |
| C-reactive protein (mg/L)       | 3 [range: 1–54]                         | 3 [range: 1–19] |
| Albumin (g/L)                   | 41 [range: 28–46]                       | 41.25 [range: 28–50] |
| Prealbumin (g/L)                | 0.28 [range: 0.17–0.41]                 | 0.27 [range: 0.13–0.42] |
| Ferritin (µmol/L)               | 82 [range: 8–334]                       | 735 [range: 6–409] |
| Plasma folic acid (mmol/L)      | 17.2 [range: 3–45]                      | 18.6 [range: 6.5–49] |
| Vitamin B12 (pmol/L)            | 357.5 [range: 126–806]                  | 305.5 [range: 40–861] |
| Magnesium (mmol/L)              | 0.79 [range: 0.67–0.97]                 | 0.81 [range: 0.38–0.95] |
| Zinc (µmol/L)                   | 11.5 [range: 9.4–15.1]                  | 11.9 [range: 7.8–14.4] |
| Selenium (µmol/L)               | 0.83 [range: 0.53–1.46]                 | 0.86 [range: 0.55–1.3567] |
| Vitamin C (µmol/L)              | 27.9 [range: 2–83]                      | 443 [range: 3–92] |
| 25 OH vitamin D (µmol/L)        | 25 [range: 14–79]                       | 43 [range: 12–104] |
| Anti-centromere antibody        | 43.8%                                    | 41.7% |
| Anti-RNA polymerase III antibody| 6.3%                                     | 0% |

* ESR: erythrocyte sedimentation rate; SSc: systemic sclerosis. Except where indicated, values are median.
assessment of fructose malabsorption. However, to date, there have been no large studies using fructose breath test in SSc and consequently, the prevalence of fructose malabsorption has not yet been determined in these patients.

The present prospective study is, to the best of our knowledge, the first to evaluate fructose malabsorption in SSc. In this instance, we found a high frequency (40%) of fructose malabsorption in patients with SSc. Since we investigated 80 SSc patients without any prior selection based on clinical and digestive presentation, our sample appeared to be representative of the entire SSc population. Our findings therefore underscore that fructose malabsorption is prevalent in the whole population of SSc patients. In the current study, fructose breath test was not performed in healthy controls, since previous investigators reported that these latter subjects exhibited fructose malabsorption in only 2.4 to 16% of cases. Other authors have also observed fructose malabsorption, in 14.4 to 39% of patients with irritable bowel syndrome (IBS).

The current study interestingly underscores that fructose breath test is a reliable method for noninvasively disclosing fructose malabsorption in SSc. Our experimental protocol was carefully designed to avoid any methodological bias that could interfere with the interpretation of the test.

First, SIBO can reasonably be ruled out as a cause of increased proportion of fructose malabsorption in our SSc patients; indeed, all these patients had a negative glucose breath test making SIBO very unlikely. Indeed, because colonic bacteria population migrates proximally into the small intestine in SIBO (gaining access to sugars), this shift in fermentation has been reported to lead to abnormal fructose breath test. However, 1 potential limitation of our study was the use of a glucose breath test for SIBO diagnosis. In fact, the gold standard for the diagnosis of SIBO remains the culture of small bowel aspirates, even such aspirates cannot detect bacterial overgrowth in the distal jejunum and proximal ileum.

Second, we chose 25 g as the dose of fructose load for testing SSc patients, because it is usually accepted as the appropriate dose for clinical use of fructose breath testing. Although Goebel-Stengel et al used 50 g as the dose of fructose load, other authors have shown that healthy subjects can only absorb up to 25 g of fructose per day; indeed, while it is possible that some healthy volunteers may consume up to 50 g of fructose per day, this amount is usually not ingested all at once. A series of 17 healthy controls, increasing fructose consumption from 15–25 g to 50 g resulted in increased prevalence of fructose malabsorption from 5.9% to 52.9%. Another series, which compared 3 doses of fructose load (15, 25, and 50 g), also demonstrated that 100% of healthy volunteers could absorb 15 g of fructose, 90% could absorb 25 g of fructose, but only 20 to 30% could absorb 50 g. Altogether these findings suggest that the physiologic dose of 25 g fructose appears to be relevant to characterize clinically significant fructose malabsorption, higher doses leading to false-positive diagnosis of fructose malabsorption.

Furthermore, we used a 5-h fructose breath test duration, because previous authors have found that 10 to 16% of fructose malabsorption would be missed with a 3-h fructose breath test duration. In this instance, the collection of breath samples was also performed according to previous methodological recommendations to ensure end-alveolar breath samples; we measured both H2 and CH4 levels, as it has been shown that false-negative findings were observed in 14% of cases when CH4 is not measured.

Another main finding in the present series was that fructose malabsorption was strongly associated with a greater prevalence and intensity of gastrointestinal clinical manifestations in SSc, as shown by the higher median value of the GSS score of digestive symptoms in SSc patients exhibiting fructose malabsorption. Previous investigators have indeed reported that unabsorbed fructose will reach the colon in patients, where colonic fermentation leads to gas production (including hydrogen, carbon dioxide, hydrogen sulphide, methan, and short-chain fatty acids), resulting in gastrointestinal symptoms such as abdominal pain and discomfort, bloating, distension, nausea, and diarrhea. In our experience, SSc patients with fructose malabsorption did not show a more altered psychological profile than SSc patients without fructose malabsorption, as shown by similar scales of anxiety and depression.

Taken together, our data suggest that fructose breath test should be performed in SSc patients without SIBO exhibiting unexplained clinical gastrointestinal symptoms to detect underlying fructose malabsorption.

In the current study, neither the patients’ age nor SSc duration could be considered to be predictive of fructose malabsorption. Furthermore, our study shows that the prevalence of fructose malabsorption was similar in patients with dcSSc and lcSSc. In addition, we found that fructose malabsorption was not correlated with other extradigestive manifestations of SSc, including digital ulcers/pitting scars, joint tying.

### Table 4. Gastrointestinal Features of the 32 SSc Patients With Fructose Malabsorption Before and After FODMAP Diet Initiation

| Clinical parameters | Before FODMAP Diet | After FODMAP Diet | P   |
|---------------------|-------------------|------------------|-----|
| Nausea              | 46.9%             | 0%               | 7.09×10^-6 |
| Vomiting            | 18.8%             | 0%               | 0.024  |
| Abdominal pain      | 81.3%             | 9.4%             | 6.62×10^-9 |
| Bloating            | 71.9%             | 28.1%            | 0.009  |
| Diarrhea            | 31.3%             | 6.3%             | 0.022  |
| Constipation        | 12.5%             | 6.3%             | 0.672  |
| Abdominal tenderness| 53.1%             | 18.8%            | 0.008  |
| GSS score of digestive symptoms | 4 (1–21) | 1 (0–13) | 0.00009 |

FODMAP: low fermentable oligosaccharides, disaccharides, monosaccharides and polyol; GSS of digestive symptoms: global symptom score of digestive symptoms; SSc: systemic sclerosis. Except were indicated, values are median.
of fructose are ingested. Other fructose transporters have been described, including GLUT7, GLUT8, and GLUT12, allowing adaptation to high free fructose intake. In a GLUT5 knockout murine model, fructose absorption was decreased by 75% in the jejunum compared with wild-type GLUT5 knockout murine model, fructose absorption was decreased by 75% in the jejunum compared with wild-type GLUT5/Glut2, which is located in the basolateral membrane of intestinal epithelium, becoming relevant when high doses of fructose are ingested. Other fructose transporters have been described, including GLUT7, GLUT8, and GLUT12, allowing adaptation to high free fructose intake. In a GLUT5 knockout murine model, fructose absorption was decreased by 75% in the jejunum compared with wild-type mice; these knockout mice, fed with high-fructose diet, developed intestinal dilation and fluid retention reminiscent of fructose malabsorption in humans. Interestingly, higher prevalence of fructose malabsorption has been reported in patients exhibiting decreased expression of GLUT5; furthermore, in patients with noninsulin-dependent diabetes mellitus, increased expression of GLUT5 has been shown to be correlated with increased absorption of fructose. By contrast, fructose malabsorption is also associated with GLUT2 mutation. Recently, Wilder-Smith et al have shown that expression of the GLUT5/GLUT2 transporters may not be uniform along the small intestine. In SSc patients, we suggest that fructose malabsorption may also be, in part, explained by reduced fructose absorption by the enterocytes related to decreased expression of GLUT5/GLUT2 transporters. In essence, inflammation has been shown to decrease intestinal GLUT5 activity and mRNA expression in murine models; similarly, in SSc patients, histological analyses of biopsy specimens have shown mononuclear inflammatory cells within intestinal wall. However, a limitation of the current study is that we did not assess the expression of GLUT5/GLUT2 transporter proteins in our SSc patients; thus, other investigations are warranted to confirm these data.

Another main finding in the present series is that fructose malabsorption was more frequent in our SSc patients with normal gastric emptying. Indeed, in healthy subjects, faster is gastric emptying, higher is the fructose load in the intestine and subsequently, higher is the osmotic load related to fructose, leading to fructose intolerance; it has been postulated that in patients with delayed gastric emptying, the postprandial small intestinal load of fructose is lower, reducing the osmotic effect of fructose, reducing the osmotic effect of fructose subsequently. Our findings are in accordance with this hypothesis, as SSc patients with normal gastric emptying more often exhibited fructose malabsorption. Additionally, in SSc patients, we also suggest that fructose malabsorption may be related to collagenous fibrosis of small intestinal wall, resulting in decreased intestinal permeability and fructose malabsorption, subsequently; and impairment of enteric microbiome. Previous investigators have reported that fructose increases intestinal translocation of bacterial endotoxin and subsequent activation of Kupffer cells through TLR-dependant mechanisms in mice. However, no definite conclusion can be drawn from our findings, and further investigations are warranted to confirm these data.

Another finding in the present study was that SSc patients with fructose malabsorption more often exhibited fructose intolerance (78.1% of cases), determined by the onset of gastrointestinal clinical manifestations during the fructose breath test. We thus suggest that fructose breath test may be useful in identifying the group of SSc patients with fructose intolerance who may benefit from low-fructose diet.

In fact, fructose coexists in food with other poorly absorbed carbohydrates that are FODMAPS. The low-FODMAP diet is defined as a diet that excludes foods rich in short-chain carbohydrates, inducing luminal distension by promoting osmosis and colonic gas production. Previous authors have reported that the low-FODMAP diet is helpful in IBS patients with symptomatic fructose intolerance, resulting in rapid improvement (within 1 wk) of gastrointestinal manifestations, and patients with nonceliac gluten sensitivity, leading to improvement of gastrointestinal symptoms. The hypothetized mechanism of action of low-FODMAP diet is related to decreased microbial fermentation of dietary FODMAPs, resulting in lower luminal gas generation and distension. Until now, no studies have directly compared the efficacy of restricting all FODMAPs with that of restricting fructose/fructans alone in patients exhibiting fructose malabsorption. In a study, low-fructose/fructan diet was found to reduce digestive symptoms in patients with fructose malabsorption. Altogether, to date, low-FODMAP diet has been mainly recommended in patients with symptomatic fructose malabsorption.

The present series also suggests that low-FODMAP diet is useful in SSc patients with symptomatic fructose malabsorption, resulting in marked improvement of clinical manifestations (ie, nausea, vomiting, abdominal pain, bloating, diarrhea, and abdominal tenderness). Our findings thus underscore that fructose malabsorption may play a significant role in the onset of gastrointestinal symptoms in SSc patients without SIBO. Interestingly, we have also observed that the strategy of dietary advice, being delivered by trained dietician following fructose breath test, provided an appropriate base for SSc patients to adhere to low-FODMAP diet which is crucial for its success. Although there are limitations in our study, because this is a nonrandomized controlled study with the potential for a placebo effect from the intervention; further randomized series are therefore warranted to confirm our findings.

In conclusion, we report a high prevalence (40%) of fructose malabsorption in SSc. Our findings are thus relevant for clinical practice, highlighting that fructose breath test is a helpful, noninvasive method by: demonstrating fructose intolerance in patients with SSc; and identifying the group of SSc patients with fructose intolerance who may benefit from fructose restriction diet. Interestingly, because the present series also shows that low-FODMAP diet resulted in a marked decrease of gastrointestinal clinical manifestations in SSc patients with fructose malabsorption, our findings underscore that fructose malabsorption may play a significant role in the onset of gastrointestinal symptoms in these patients. Finally, we suggest that fructose malabsorption may be related to impairment of fructose absorption by the enterocytes and/or enteric microbiome and decrease of intestinal permeability in SSc.
41. Berg LK, Fagerli E, Martinussen M, et al. Effect of fructose-reduced diet in patients with irritable bowel syndrome, and its correlation to a standard fructose breath test. Scand J Gastroenterol. 2013;48:936–943.

42. Erdogan A, Adame EC, Yu S, et al. Optimal testing for diagnosis of fructose intolerance: over-dosage leads to false positive intolerance test. J Neurogastroenterol Motil. 2014;20:560.

43. Frielings T, Kuhlbusch-Zicklam R, Kalde S, et al. Fructose malabsorption: how much fructose can a healthy subject tolerate? Digestion. 2011;84:269–272.

44. Gibson PR, Newnham E, Barrett JS, et al. Review article: fructose malabsorption and the bigger picture. Aliment Pharmacol Ther. 2007;25:349–365.

45. Ravich WJ, Bayless TM, Thomas M. Fructose: incomplete intestinal absorption in humans. Gastroenterology. 1983;84:26–29.

46. Rumessen JJ, Gudmund-Hoyer E. Absorption capacity of fructose in healthy adults. Comparison with sucrose and its constituent mono-saccharides. Gut. 1986;27:1161–1168.

47. Sung HY, Kim YS. Fructose malabsorption in patients with irritable bowel syndrome-like symptoms: What is the role in the pathogenesis and clinical implication? J Neurogastroenterol Motil. 2014;20:135–137.

48. Goebel-Stengel M, Stengel A, Schmidtmann M, et al. Unclear abdominal discomfort: pivotal role of carbohydrate malabsorption. J Neurogastroenterol Motil. 2014;20:228–235.

49. Goebel-Stengel M, Monnikes H. Optimal testing for diagnosis of fructose malabsorption: under-dosage leads to false negative intolerance test. J Neurogastroenterol Motil. 2015;21:296–297.

50. Kyaw MH, Mayberry JF. Fructose malabsorption: true condition or a variance from normality. J Clin Gastroenterol. 2011;45:16–21.

51. Montonen J, Järvinen R, Knekt P, et al. Consumption of sweetened beverages and intakes of fructose and glucose predict type 2 diabetes occurrence. J Nutr. 2007;137:1447–1454.

52. Pimentel M, Chow EJ, Lin HC. Eradication of small intestinal bacterial overgrowth reduces symptoms of irritable bowel syndrome. Am J Gastroenterol. 2000;95:3503–3506.

53. Knudsen CD, Di Palma JA. Do you need to measure methane? South Med J. 2012;105:251–253.

54. de Lacy Costello BP, Ledochowski M, Ratcliffe NM. The importance of methane breath testing: a review. J Breath Res. 2013;7:024001.38.

55. Latulippe ME, Skoog SM. Fructose malabsorption and intolerance: effects of fructose with and without simultaneous glucose ingestion. Crit Rev Food Sci Nutr. 2011;51:583–592.

56. Ong DK, Mitchell SB, Barrett JS, et al. Manipulation of dietary short chain carbohydrates alters the pattern of gas production and genesis of symptoms in irritable bowel syndrome. J Gastroenterol Hepatol. 2010;25:1366–1373.

57. Simrén M, Stotzer PO. Use and abuse of hydrogen breath tests. Gut. 2006;55:297–303.

58. Biesiekierski JR. Fructose-induced symptoms beyond malabsorption in FGID. United European Gastroenterol J. 2014;2:10–13.

59. Burant CF, Takeda J, Brot-Laroche E, et al. Fructose transporter in human spermatozoa and small intestine is GLUT5. J Biol Chem. 1992;267:14523–14526.

60. Davidson NO, Hausman AM, IKhovits CA, et al. Human intestinal glucose transporter expression and localization of GLUT5. Am J Physiol. 1992;262:C795–C800.

61. Gouyon F, Onesto C, Dalet V, et al. Fructose modulates GLUT5 mRNA stability in differentiated Caco-2 cells: role of CAMP-signalling pathway and PABP (polyadenylated-binding protein)-interacting protein (Pai1). 2. Biochem J. 2003;375:167–174.

62. DeBosch BJ, Chi M, Moley KH. Glucose transporter 8 (GLUT8) regulates enterocyte fructose transport and global mammalian fructose utilization. Endocrinology. 2012;153:4181–4191.

63. Douard V, Ferraris RP. Regulation of the fructose transporter GLUT5 in health and disease. Am J Physiol Endocrinol Metab. 2008;295:E227–237.

64. Jones HF, Butler RN, Brooks DA. Intestinal fructose transport and malabsorption in humans. Am J Physiol Gastrointest Liver Physiol. 2011;300:G202–G206.

65. Vos MB. Nutrition, nonalcoholic fatty liver disease and the microbiome: recent progress in the field. Curr Opin Lipidol. 2014;25:61–66.

66. Wasserman D, Hoekstra JH, Tolia V, et al. Molecular analysis of the fructose transporter gene (GLUT5) in isolated fructose malabsorption. J Clin Invest. 1996;98:2398–2402.

67. Douard V, Ferraris RP. The role of fructose transporters in diseases linked to excessive fructose intake. J Physiol. 2013;591:401–414.

68. Jones HF, Butler RN, Moore DJ, et al. Developmental changes and fructose absorption in children: effect on malabsorption testing and dietary management. Nutr Rev. 2013;71:300–309.

69. Dyer J, Wood IS, Palejwala A, et al. Expression of monosaccharide transporters in intestine of diabetic humans. Am J Physiol Gastrointest Liver Physiol. 2002;282:G241–G248.

70. Southgate DA. Digestion and metabolism of sugars. Am J Clin Nutr. 1995;62:203S–210S.

71. Horowitz M, Cunningham KM, Wishart JM, et al. The effect of short-term dietary supplementation with glucose on gastric emptying of glucose and fructose and oral glucose tolerance in normal subjects. Diabetologia. 1996;39:481–486.

72. Moukarzel AA, Sabri MT. Gastric physiology and function: effects of fruit juices. J Am Coll Nutr. 1996;15(5 Suppl):185–258.

73. Barrett JS, Gibson PR. Fermentable oligosaccharides, disaccharides, monosaccharides and polyols (FODMAPs) and nonallergic food intolerance: FODMAPs or food chemicals? Therap Adv Gastroenterol. 2012;5:261–268.

74. Gibson PR, Varney J, Malakar S, et al. Food components and irritable bowel syndrome. Gastroenterology. 2015;148:1158–1174.

75. Pedersen N, Vegh Z, Burisch J, et al. Ehealh monitoring in irritable bowel syndrome patients treated with low fermentable oligo-, di-, mono-saccharides and polyols diet. World J Gastroenterol. 2014;20:6680–6684.

76. Rao SS, Yu S, Fedewa A. Systematic review: dietary fibre and FODMAP-restricted diet in the management of constipation and irritable bowel syndrome. Aliment Pharmacol Ther. 2015;41:1256–1270.

77. Staudacher HM, Irving PM, Lomer MC, et al. Mechanisms and efficacy of dietary FODMAP restriction in IBS. Nat Rev Gastroenterol Hepatol. 2014;11:256–266.

78. Biesiekierski JR, Peters SL, Newnham ED, et al. No effects of gluten in patients with self-reported non-celiac gluten sensitivity after dietary reduction of fermentable, poorly absorbed, short-chain carbohydrates. Gastroenterology. 2013;145:320–e1–320–e3.

79. Murray K, Wilkinson-Smith V, Hoad C, et al. Differential effects of FODMAPs (fermentable oligo-, di-, mono-saccharides and polyols) on small and large intestinal contents in healthy subjects shown by MRI. Am J Gastroenterol. 2014;109:110–119.

80. Shepherd SJ, Gibson PR. Fructose malabsorption and symptoms of irritable bowel syndrome: guidelines for effective dietary management. J Am Diet Assoc. 2006;106:1631–1639.

81. Escobar MA Jr, Lustig D, Pfliugese BM, et al. Fructose intolerance/malabsorption and recurrent abdominal pain in children. J Pediatr Gastroenterol Nutr. 2014;58:498–501.