Effect of Raw Milk on Lactose Intolerance: A Randomized Controlled Pilot Study

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ABSTRACT

PURPOSE This pilot study aimed to determine whether raw milk reduces lactose malabsorption and/or lactose intolerance symptoms relative to pasteurized milk.

METHODS We performed a crossover trial involving 16 adults with self-reported lactose intolerance and lactose malabsorption confirmed by hydrogen (H₂) breath testing. Participants underwent 3, 8-day milk phases (raw vs 2 controls: pasteurized, soy) in randomized order separated by 1-week washout periods. On days 1 and 8 of each phase, milk consumption was 473 mL (16 oz); on days 2 to 7, milk dosage increased daily by 118 mL (4 oz), beginning with 118 mL (4 oz) on day 2 and reaching 710 mL (24 oz) on day 7. Outcomes were area under the breath H₂ curve (AUC ∆H₂) and self-reported symptom severity (visual analog scales: flatulence/gas, audible bowel sounds, abdominal cramping, diarrhea).

RESULTS AUC ∆H₂ (mean ± standard error of the mean) was higher for raw vs pasteurized on day 1 (113 ± 21 vs 71 ± 12 ppm·min·10⁻², respectively, P = .01) but not day 8 (72 ± 14 vs 74 ± 15 ppm·min·10⁻², respectively, P = .9). Symptom severities were not different for raw vs pasteurized on day 7 with the highest dosage (P > .7). AUC ∆H₂ and symptom severities were higher for both dairy milks compared with soy milk.

CONCLUSIONS Raw milk failed to reduce lactose malabsorption or lactose intolerance symptoms compared with pasteurized milk among adults positive for lactose malabsorption. These results do not support widespread anecdotal claims that raw milk reduces the symptoms of lactose intolerance.

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INTRODUCTION

Lactose malabsorption is an extremely common condition worldwide, and its incidence increases with age and varies between ethnicities.¹,² Bacteria in the colon ferment undigested lactose into short-chain fatty acids, hydrogen, methane, and carbon dioxide, resulting in such symptoms as bloating, abdominal pain, and/or diarrhea after ingestion of lactose.¹ National Institutes of Health (NIH) consensus development panel defined lactose intolerance as a clinical syndrome characterized by the onset of gastrointestinal symptoms following a lactose challenge,³ whereas true lactose malabsorption is identified through a hydrogen breath test (HBT). Many people with lactose malabsorption do not report clinical lactose intolerance.⁴ Conversely, many individuals with perceived lactose intolerance do not experience malabsorption.⁶ Because of inconsistent definitions in previous studies, the true prevalence of lactose intolerance is unknown.⁷

Strategies to reduce unpleasant gastrointestinal symptoms associated with lactose intolerance include (1) choosing lactose-free or reduced-lactose dairy foods,⁸,¹¹ (2) choosing fermented dairy foods,¹¹,¹⁵ (3) combining lactose intake with a meal,¹⁴,¹⁶ (4) taking lactase enzyme tablets,¹⁰,¹¹,¹⁷ (5) consuming probiotics,¹¹,¹⁸-²⁰ (6) colonic adaptation,²¹-²² and (7) psychological and behavioral approaches.²³-²⁵ None of these strategies, however, have been shown to fully eliminate symptoms. A recent review by the
NIH highlighted the need for further studies evaluating the effectiveness of interventions addressing lactose intolerance.\textsuperscript{3,7}

Recently, unpasteurized raw milk consumption has increased in popularity and emerged into a nationwide movement despite the acknowledgement of risks associated with foodborne pathogens.\textsuperscript{26} Raw milk proponents and producers purport that consumption of raw milk is associated with reductions in atopic conditions,\textsuperscript{27-29} autism, inflammatory bowel disease, and lactose intolerance; these claims currently lack definitive scientific evidence. In the case of raw vs pasteurized yogurt, it has been shown repeatedly that compared with pasteurized yogurt, unpasteurized yogurt containing live bacterial cultures with lactase activity significantly reduces lactose malabsorption.\textsuperscript{30-34} These findings are apparently due to bacterial lactase activity provided in the small intestine and/or changes in the intestinal milieu.\textsuperscript{30,34} As in the case of raw yogurt, the omission of pasteurization in the raw milk production process results in a greater number of surviving microflora,\textsuperscript{26} including naturally occurring strains of lactobacilli.\textsuperscript{35} It has been hypothesized that these additional microflora may aid in the digestion of lactose, reducing lactose malabsorption for raw milk relative to pasteurized milk. The objective of this pilot study was to examine whether breath hydrogen (H\textsubscript{2}) (standard measure of lactose malabsorption, primary outcome) and/or symptoms of lactose intolerance (secondary outcomes) would be reduced after consuming raw milk vs pasteurized milk, using soy milk as a negative control.

**METHODS**

**Study Participants**
Participants were recruited from the local community near Stanford University through email and radio advertisements and letters to previous study participants. Participants were screened first using an online survey. The initial eligibility criteria were self-reported lactose intolerance symptoms of "moderate" to "severe" severity. Exclusion criteria included self-reported symptoms of "mild" or "extremely severe" severity, recent or planned antibiotic consumption, a history of diarrheal illness within the past month, and a history of any gastrointestinal conditions other than lactose malabsorption. Remaining eligible participants were screened using the standardized HBT. Participants whose peak hydrogen concentrations rose to 25 ppm or higher above baseline and who simultaneously experienced symptoms of lactose intolerance were included in the study. Participants received $150 for completing all 3 milk phases. All participants provided written informed consent, and the study was approved by the Stanford University Human Subjects Committee.

**Study Design**
We used a randomized, double-blind, 3-way crossover design. The protocol required each participant to consume 3 types of milk for 8 days each: organic whole raw milk (R), organic whole pasteurized milk (P), and plain (unflavored) soy milk (S). Each 8-day treatment phase was separated by a washout phase of 1 week. Participants were instructed to avoid consuming all dairy and lactose-containing products (other than study products) throughout the treatment and washout phases, as well as during a 1-week run-in period that preceded the first treatment week.

Milk dosage for each 8-day phase is displayed in Figure 1. Eight-day milk phases were selected to explore potential adaptation to each of the milk products resulting in possible changes in the intestinal milieu.\textsuperscript{30,34} Each 8-day phase was anchored on days 1...
and 8 by a dosage of 473 mL (16 oz) to allow for examination of potential adaptation. Each phase included an incrementally increasing dosage on days 2 to 7 to explore participant tolerance to doses ranging from small to substantial. Participants had the option of consuming less than the assigned dose at any point if their symptoms were unbearable. The amount of milk consumed was confirmed verbally twice during each phase, and participants were asked to return leftover milk, which was measured by study staff to corroborate verbal accounts.

The 8-day treatment duration, escalating dosage (4 to 24 oz), and compensation amount ($150) were selected in the design phase of the project and informed by 2 focus groups conducted among lactose intolerant individuals. Dosage and duration were selected to lead to symptoms sufficiently severe to allow for a detectable difference in treatment arms but sufficiently bearable to allow for successful recruitment of typical lactose intolerant individuals, thereby allowing for a reasonable level of generalizability of the findings. Participants were randomly assigned to 1 of 6 possible sequences of the 3 milks: RPS, RSP, PRS, PSR, SRP, or SPR. Randomization was done in blocks of 12 (ie, 2 of each of the 6 possible sequences) and was performed by a researcher (S.M.) who was blinded to the study assignments and who selected pieces of paper with order assignments from an envelope.

**Milk Products**

The raw milk was an organic, grade A, whole milk produced by Organic Pastures, which follows state mandated testing for human pathogens. The pasteurized milk was an organic, grade A, whole milk produced by Horizon Organic. Among 12 different types of soy milk taste tested by a panel of study staff, the soy milk that most resembled the taste and appearance of the cow’s milk was selected: an organic, Soy Dream Original Classic brand produced by The Hain Celestial Group. Energy content and macronutrient composition of the milk products were obtained from label information (Table 1).

**Blinding**

Preparation and distribution of each of the milk types was made as comparable as possible so study participants and researchers were blinded to the type of milk they received. To maximize palatability and masking of milk taste, sugar-free vanilla syrup flavoring (Torani brand) was added to all milks in a flavoring to milk ratio of 1:31. To maintain staff blinding, milk containers were prepared and coded by a researcher (S.M.) who did not have participant contact, while separate researchers (Q.V. and J.H.) distributed milks to participants and remained blinded to assignment phase for the duration of the study. All milks were provided in standardized, unlabeled containers. Participants were asked at the end of each treatment phase to try to identify their milk assignment.

**Data Collection**

**Hydrogen Breath Test**

The primary study outcome was breath H$_2$ excretion as measured by HBT. The increase in H$_2$ production after the consumption of lactose corresponds to the degree of lactose malabsorption. Those determined eligible after the initial online survey screening were invited to complete a 3-hour HBT: after an overnight fast, end-alveolar air samples containing 20 mL or more were collected before and at 9 consecutive 20-minute intervals after an oral load of 25 g of lactose dissolved in water, with continued fasting throughout the 3-hour test. Expired H$_2$ concentrations were stored in plastic syringes with stopcocks and measured within 12 hours, in parts per million, using gaseous chromatography (Breath Tracker Digital Microlyzer, model SC, Quintron Instruments). Participants whose H$_2$ levels rose 25 ppm or more above baseline and who experienced any symptom(s) of lactose intolerance during the test were included in the study.

Once enrolled, on days 1 and 8 of each phase, participants completed a similar 4-hour HBT after consumption of the assigned milk. Samples were collected for 4 rather than 3 hours during the tests conducted with study milks to account for the longer digestion time of milk relative to the lactose solution used for the screening HBT. H$_2$ concentrations were expressed as area under the H$_2$ curve above baseline (AUC ∆H$_2$) in parts per million per minute per $10^{-2}$ (ppm · min · 10$^{-2}$), calculated according to the linear trapezoidal rule ignoring any area below the baseline, and as maximal increase over baseline concentration (peak ∆H$_2$) in parts per million.

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**Table 1. Composition of Milk Products per 237 mL (8 oz)**

| Component     | Raw Milk | Pasteurized Milk | Soy (Nondairy) Milk |
|---------------|----------|------------------|--------------------|
| Energy (kcal) | 145      | 145              | 126                |
| Macronutrients (g) |          |                  |                    |
| Fat           | 7.8      | 7.8              | 3.9                |
| Protein       | 7.8      | 7.8              | 6.8                |
| Carbohydrate  | 11.6     | 11.6             | 15.5               |
| Sugars        | 11.6     | 11.6             | 8.7                |
| Lactose       | 11.6     | 11.6             | 0                  |

Note: All milk products included sugar-free vanilla syrup flavoring in a flavoring to milk ratio of 1:31.
Symptoms of Lactose Intolerance
To assess the incidence and severity of symptoms, a validated gastrointestinal symptom log was used asking participants to mark on a 10-cm visual analog scale of 0 to 10 the severity of 4 symptoms: flatulence/gas, diarrhea, audible bowel sounds, and abdominal cramping. Symptom logs were completed at 52 time points: 4 times during the screening HBT (at baseline and at 3 consecutive 1-hour intervals); 5 times during the HBT on days 1 and 8 of each phase (at baseline and at 4 consecutive 1-hour intervals); and once per day on days 2 through 7 of each phase.

Statistical Methods
Sample size was determined based on the selection of a 25% decrease in AUC ∆H2 as the minimal difference that would be clinically significant—a projected effect size of 1.0. With a crossover design, it was determined that 15 subjects would yield 95% power using $\alpha = .05$ to detect a 25% decrease. Differences in AUC ∆H2 and peak ∆H2 concentrations (primary outcomes) among milk phases were examined by repeated-measures analysis of variance (ANOVA). Repeated-measures ANOVA was also used to test for differences within each of the symptom categories (secondary outcomes) on day 7. For both the H2 and symptom levels, when statistical significance was reached by ANOVA, pairwise differences were examined using matched pairs $t$ tests. Matched pairs $t$ tests were also used to test for differences in AUC ∆H2 and peak ∆H2 concentrations between days 1 and 8 of each phase. In each analysis, only those study participants with complete data were included in the statistical testing. All statistical tests were 2-tailed using $\alpha < .05$.

RESULTS
Study Population
Participant enrollment began in May 2010, and the study ended in September 2010. Of 63 potential participants screened using the HBT, 27 (43%) tested positive for lactose malabsorption (Figure 2), all of whom reported symptoms of lactose intolerance during the HBT. Among these 27 individuals, 11 chose not to continue. Among the 16 participants who were randomized, 2 to 4 were assigned to each of the 6 possible orders of study milks. Randomized participants were aged a mean of 40 years (SD = 14 years), had a mean of 16 years (SD = 3 years) of education, and had a mean body mass index of 24 kg/m² (SD = 2 kg/m²).

Retention and Adherence
All 16 participants completed all 3 milk phases. There was 100% adherence on day 1 of all 3 milk phases. Four participants failed to adhere to the full 8-day protocol during 1 of the 3 milk phases: 3 participants opted to consume less than the assigned dosage during 1 of the phases due to unbearable symptoms, and 1 participant accidentally did not consume milk on day 6 of the R phase.

There was no significant difference in adherence to the milk consumption protocol among the 3 milk phases ($P = .3$), and 90% or more of the total assigned milk intake was consumed by 88%, 94%, and 100% of the participants during the R, P, and S phases, respectively.
Blinding
Of the 16 participants, 12 (80%) correctly guessed assignment to the soy milk phase, whereas 6 (40%) correctly identified both dairy milks. Successful identification of milk assignment increased with each successive milk phase from 43% to 69% to 88% during the first, second, and third phases, respectively.

H₂ Outcomes
Mean H₂ production (± standard error of the mean [SEM]) during the course of the HBT on day 1 is displayed in Figure 3a and on day 8 is displayed in Figure 3b. AUC ΔH₂ and peak ΔH₂ concentrations for both days 1 and 8 are shown in Table 2. Contrary to what was hypothesized, on day 1, both AUC ΔH₂ and peak ΔH₂ concentrations were significantly higher for R relative to P (P = .01). In contrast, on day 8, no significant difference was observed between R and P (P = .9). On both days 1 and 8, AUC ΔH₂ and peak ΔH₂ concentrations were significantly higher for the 2 dairy milks (R and P) relative to the soy milk (P < .001). AUC ΔH₂ and peak ΔH₂ concentrations between days 1 and 8 indicated a borderline significant reduction in H₂ production during the course of the R phase (P = .05, and .06, respectively) (Table 2). No significant change was observed for the P phase (P > .6) or S phase (P = .7).

Symptom Outcomes
The highest milk dose, 710 mL (24 oz), occurred on day 7 of each milk phase and produced the most severe symptoms relative to all other days, as would be expected. All but 3 participants were able to complete this dose for all 3 milk phases. Mean self-reported symptom severity levels (± SEM) are displayed in Figure 4. There was no significant difference in severity levels between R and P for any of the 4 categories (P > .7).

DISCUSSION
Our trial was designed to determine whether lactose malabsorption and/or lactose intolerance symptoms experienced by adults positive for lactose malabsorption...
would be reduced with raw milk vs pasteurized milk. The hypothesis was not supported. To the contrary, \( H_2 \) results showed higher lactose malabsorption for raw vs pasteurized milk on day 1, and \( H_2 \) results showed comparable degrees of lactose malabsorption for both milks on day 8. Day 7 symptom severities were similar for raw and pasteurized milk. Inclusion of soy milk as a negative control showed that in all cases both dairy milks induced significantly greater degrees of lactose malabsorption and intolerance symptoms. Overall, this 3-arm crossover trial provided no evidence that raw milk is better tolerated by adults positive for lactose malabsorption, either objectively or subjectively.

Previous studies have shown that in the case of yogurt, the additional microflora in unpasteurized yogurt reduce lactose malabsorption relative to pasteurized yogurt.\(^{30-34}\) One possible reason that a similar effect was not observed in the current study stems from the greater viscosity of yogurt, which prolongs digestion time, potentially allowing more time for the microflora to hydrolyze lactose in the small intestine.\(^{30}\) The current findings for raw milk parallel those of sweet acidophilus milk, which also contains live bacteria but has similarly been shown not to reduce lactose malabsorption.\(^{32,40,41}\)

Interestingly, raw milk induced significantly greater \( H_2 \) production than pasteurized milk on day 1 but not on day 8, despite containing identical amounts of lactose. It is not clear why this was the case. Although the reduced \( H_2 \) production observed for raw milk on day 8 vs day 1 suggests a degree of adaptation to raw milk, this apparent adaptation led only to levels of lactose malabsorption comparable to those of pasteurized milk. In contrast to raw milk, no adaptation was observed for pasteurized milk. This finding does not support the colonic adaptation hypothesis for conventional milk and parallels findings from other randomized controlled trials.\(^{7,21,22}\)

Notably, 57% (36 of 63) of screened individuals in the current study tested negative for lactose malabsorption despite

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**Table 2. Breath H\(_2\) Results for Each 8-Day Crossover Phase, n = 16 (mean ± SEM)**

| Phase       | Milk Consumption Phases | \( P \) Value* |
|-------------|-------------------------|----------------|
|             | Raw (R) | Pasteurized (P) | Soy (S) | R vs P | R vs S | P vs S |
| AUC \( \Delta H_2 \),\(^{b}\) |          |                |        |        |        |        |
| Day 1       | 113 ± 21 | 71 ± 12 | 5 ± 2 | 0.01 | <.001 | <.001 |
| Day 8       | 72 ± 14\(^{c}\) | 74 ± 15 | 5 ± 2 | 0.9 | <.001 | <.001 |
| Peak \( \Delta H_2 \) |          |                |        |        |        |        |
| Day 1       | 117 ± 20 | 75 ± 11 | 11 ± 4 | 0.01 | <.001 | <.001 |
| Day 8       | 79 ± 14\(^{d}\) | 80 ± 13 | 12 ± 3 | 0.9 | <.001 | <.001 |

ANOVA = analysis of variance; AUC \( \Delta H_2 \) = area under the \( H_2 \) curve above baseline; Peak \( \Delta H_2 \) = highest \( H_2 \) concentration after milk ingestion, minus baseline concentration (ppm); \( H_2 \) = hydrogen; HBT = hydrogen breath test.

Notes: End-alveolar air samples collected before and at 12 consecutive 20-minute intervals after ingestion of 473 mL (16 oz) of milk. All data for 2 participants who did not undergo the day-8 HBT during one of the milk phases were omitted from the day-8 analysis (\( n = 14 \)). In the analysis comparing days 1 and 8, all data were omitted for 1 participant in the R phase and 1 participant in the P phase analyses who did not complete the day-8 HBT in each respective phase (\( n = 15 \)).

\(^{a}\) Group differences were tested by repeated-measures ANOVA, showing significant differences among the 3 milks in all instances (\( P < .001 \)). Pairwise differences were subsequently calculated using matched pairs t tests.

\(^{b}\) AUC \( \Delta H_2 \) calculated by the linear trapezoidal method (ppm · min · 10\(^{-2}\)).

\(^{c}\) Matched pair t test for day 1 vs day 8 for raw milk (\( n = 15 \)): \( P = .05 \).

\(^{d}\) Matched pair t test for day 1 vs day 8 for raw milk (\( n = 15 \)): \( P = .06 \).
their personal belief they were lactose intolerant. This finding confirms conclusions drawn by a recent NIH review indicating that many who believe they are lactose intolerant are not true lactose malabsorbers.1

There were several strengths in the design of the current study, including each participant serving as his or her own control in the crossover design, the use of both objective (breath H₂) and subjective (symptom report) outcome measures, and the use of soy milk as a negative control. Strengths in conduct of the study included 100% retention across all 3 milk phases and high levels of adherence to the incrementally increasing dosage schedule. Double-blind study efforts proved highly effective for study staff and partially effective for participants. The current study also had strong ecological validity as one of the few studies on lactose intolerance that utilized commercially available milk products rather than isolated lactose. We believe it is also the first randomized controlled trial examining the effect of raw milk on symptoms of lactose intolerance. Overall, these strengths in design and conduct address many of the limitations identified in the recent NIH consensus report.1

The study also included several limitations. The sample size was small, the impact of which was mitigated to some degree by using a crossover design. The sample size was too small to examine potential ethnic/racial differences. The 8-day study phases precluded examining possible responses or adaptation to raw milk that might have occurred after longer periods of time. A requirement to elicit and sustain symptoms of lactose intolerance for longer than 8 days, however, would likely make recruitment substantially more challenging, resulting in a lower percentage of eligible participants enrolling and thus a decrease in generalizability. The study also did not address any questions regarding consumer behavior associated with the higher cost (2- to 3-fold) of raw milk relative to conventional milk.

Replication of the findings presented here would strengthen the conclusions. To that end, the design and conduct of this pilot study should be useful in informing future studies of a larger scale. In particular, feedback from study participants suggested that escalating the milk dosage over time, with the option to discontinue the escalation at any time because of unbearable symptoms, was an important strategy for addressing concerns about the discomfort level that would arise from participating. Perhaps equally informative, and unexpected by our research team, more than 80% of our participants were willing to complete the full 1-week regimen and consume the highest 24-oz dose in 1 sitting for all 3 milk types. Given that these participants were confirmed lactose intolerant individuals (subjective) who were also determined to suffer from lactose malabsorption (objective), we would not have anticipated such a high rate of adherence to the highest dosage level. These observations and others reported here should prove useful to other investigators designing future trials to test the ability of other products that might diminish the symptoms of lactose intolerance.

Among those who report intolerance symptoms because of lactose malabsorption, these data do not support the widespread claim that raw milk confers benefits in reducing the discomfort of lactose intolerance. Primary care physicians and gastroenterologists should be aware that the evidence supporting raw milk consumption remains anecdotal. Although other health benefit claims for raw milk are plausible, such claims remain similarly anecdotal and unsubstantiated and should be subjected to appropriately designed controlled trials.

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Author contributions: The authors’ responsibilities were as follows: Mummah and Gardner had full access to all of the data in the study and take responsibility for the integrity of the data and the accuracy of the data analysis. Mummah, Oelrich, Hope, and Gardner designed research. Mummah, Hope, and Vu conducted research. Mummah analyzed data. Mummah, Oelrich, and Gardner wrote the paper. Mummah and Gardner had primary responsibility for final content. Gardner provided guidance and consultation throughout. All authors read and approved the final manuscript.

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RAW MILK AND LACTOSE INTOLERANCE

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