Evaluation of differential disaccharide excretion in urine for non-invasive investigation of altered intestinal disaccharidase activity caused by α-glucosidase inhibition, primary hypolactasia, and coeliac disease

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Abstract
Background/Aim—The reliability of a quantitative method for the non-invasive assessment of intestinal disaccharidase hydrolysis was assessed.

Methods—Differential excretion of intact disaccharide, expressed as ratios of lactulose to appropriate hydrolysable disaccharides in urine collected following combined ingestion, has been investigated in healthy volunteers with drug induced α-glucosidase inhibition, in subjects with primary hypolactasia, and patients with coeliac disease.

Results—Oral administration of the α-glucosidase inhibitor ‘Acarbose’ (BAY g 5421, 200 mg) together with sucrose and lactulose increased the urinary sucrose/lactulose excretion ratios (% dose/10 h) fivefold. The effect was quantitatively reproducible, a higher dose of ‘Acarbose’ (500 mg) increasing the excretion ratio to about 1.0 indicating complete inhibition of intestinal sucrase activity. The suitability of the method for measuring differences in dose/response and duration of action was assessed by comparing three different α-glucosidase inhibitors (BAY g 5421, BAY m 1099, and BAY o 1248) and found to be satisfactory. Subjects with primary adult hypolactasia had urine lactose/lactulose excretion ratios raised to values indicating reduced rather than complete absence of lactase activity whereas sucrose/lactulose ratios were not significantly affected. ‘Whole’ intestinal disaccharidase activity assessed by this method demonstrated impairment of lactase, sucrase, and isomaltase in eight, one, and seven, respectively, of 20 patients with coeliac disease. By contrast in vitro assay of jejunal biopsy tissue indicated pan-disaccharidase deficiency in all but five of these patients. This shows the importance of distinguishing between ‘local’ and ‘whole’ intestinal performance.

Conclusions—Differential urinary excretion of ingested disaccharides provides a reliable, quantitative, and non-invasive technique for assessing profiles of intestinal disaccharidase activity.

(Gut 1996; 39: 374–381)

Keywords: coeliac disease, intestinal function, intestinal disaccharidases, intestinal permeability.
following their combined ingestion, is based on the finding that recovery of intact disaccharide in urine after ingestion is inversely related to the rate of their intestinal hydrolysis. The test solution contains a non-hydrolysable ‘reference’ disaccharide (for example, lactulose) together with hydrolysable disaccharides (lactose, sucrose, palatinose). As renal excretion of these disaccharides after entering the circulation is known to be complete, without metabolic loss, the excretion ratios of hydrolysable/

non-hydrolysable disaccharide should relate inversely to the activity of intestinal disaccharides (lactase, sucrase and, in the case of palatinose, isomaltase). This technique has been used to demonstrate lactase deficiency associated with rotaviral infection and combined sucrase and isomaltase deficiency in asucrasis. The main purpose of this study was further evaluation of the reliability of the urine disaccharide ratio technique for both clinical and research purposes. This has been undertaken by assessing the ability of the differential disaccharide excretion technique to distinguish the differences in dose/response and duration of action of three orally administered α-glucosidase inhibitors and, secondly, to quantitate intestinal lactase activity in subjects with confirmed genetic hypolactasia (primary adult hypolactasia). Intestinal disaccharide hydrolysis assessed in vivo by the combined disaccharide ratio test was also compared with in vitro assay of disaccharidase activity in homogenised jejunal biopsy tissue in patients with coeliac disease to study the effect of a disorder that mainly involves the jejunum.

Methods
Differential Urinary Excretion of Ingested Disaccharides: Principles Involved
The principle of the differential urinary excretion of ingested sugar probes has been detailed elsewhere. Figure 1 briefly outlines the principle and how urinary excretion ratios of a non-hydrolysable disaccharide and hydrolysable disaccharides can give quantitative information of the rate of intestinal hydrolysis of some disaccharides.

Studies With α-Glucosidase Inhibitors
Fourteen healthy adult white volunteers were studied. A strict sucrose free diet was implemented 18 hours before and throughout the test period. After an overnight fast each subject voided urine directly before the test to confirm absence of sucrose. At 8 am the test solution containing 20 g sucrose (α-D-glucopyranosyl β-D-fructofuranoside) and 6.7 g lactulose (β-D-galactopyranosyl-fructose=10 ml ‘Duphalac’ lactulose syrup, Duphar Laboratories, Southampton) dissolved in 300 ml water (300 mmol/l) was ingested within a period of four minutes. Food and fluids were not permitted until 2-5 hours later. A complete urine collection was made from 8 am to 6 pm (10 hours), the volume recorded, and an aliquot preserved with methanol (10 mg/100 ml minimum) for analysis of sugars by quantitative paper or thin layer chromatography. Results are expressed as urinary sucrose/lactulose ratios of percentages of the oral doses excreted during 10 hours.

Dose/response studies
To evaluate the reliability of the method for the quantitative differentiation of intestinal

Figure 1: Principle of the differential urine excretion of orally administered test substances. The precise quantity of intact disaccharide passing by unmediated diffusion across the intestinal mucosa into the circulation is determined by the factors shown in the Figure. The inclusion of lactulose, which resists the action of intestinal hydrodases in a test solution containing hydrolysable disaccharides (lactose, sucrose or palatinose either individually or in combination), enables a correction for non-mucosal variables on urinary excretion to be obtained. These four disaccharides permeate the intestine as intact molecules by a common mucosal pathway in quantities that depend upon their individual concentrations within the intestine. In turn, these concentrations are determined by the efficiency of hydrolysis by appropriate disaccharidases. As lactulose will be affected in the same way as the ‘test’ disaccharides by all these factors except for the activity of mucosal disaccharidase, this disaccharide resists, the urinary excretion ratios of lactose, sucrose, or palatinose, or all three, to lactulose can be regarded as specific indices of the efficiency with which the concentrations of lactose, sucrose, and palatinose are reduced by hydrolysis within the small intestine. In the absence of hydrolysis of lactose the fraction of the ingested dose permeating the intestine will approach that of lactulose (which resists hydrolysis) so that the lactose/lactulose ratio of percentages of the oral dose excreted in urine would be 1.0. With normal active intestinal hydrodases of lactose this ratio is much lower, usually between 0.1-0.3, and intermediate values provide a quantitative measure of the degree of impaired hydrolysis. Nil=No effect, + = Plays a part in the overall urine excretion of the test substance but the effect on simultaneously ingested test substances is equal. ++ = Plays a part in the overall urine excretion of the test substance but the effect on simultaneously ingested test substances differs.
disaccharidase activity a study of the effect of different dose levels of a known α-glucosidase inhibitory drug (BAY g 5421, ‘Acarbose’) was undertaken. Subjects underwent a control test without α-glucosidase inhibition (n=14), which was then repeated (with intervals of at least one week) adding either BAY g 5421 (Acarbose: dose range 25–500 mg, n=4–8), BAY m 1099, (dose 100 mg, n=7) or BAY o 1248 (dose 10 mg, n=6) to the test solution. One subject underwent 13 tests with BAY g 5421 in the dose range 0–500 mg added to a test solution that contained lactose (β-1-4-D-galacto-pyranosyl-α-D-glucose) 20 g in addition to sucrose 20 g and lactulose 6.7 g dissolved in 300 ml drinking water, to assess response in relation to both sucrose and lactose hydrolysis.

Duration of action
To demonstrate the ability of the method to resolve differences in the duration of disaccharidase inhibition the action of the three α-glucosidase inhibitors on intestinal sucrose hydrolysis over a timed period was also assessed in a single individual as follows: (a) BAY g 5421 (‘Acarbose’ 500 mg) with, and then 30, 60, 120 minutes and 48 hours before oral administration of the test solution. (b) BAY m 1099 (200 mg) with, and then 30, 60, 120 minutes and 48 hours before the test solution. (c) BAY o 1248 (20 mg) with, and then 60, 120, 180 minutes and 48 hours before the test solution.

INVESTIGATION OF SUBJECTS WITH PRIMARY (GENETIC) ADULT HYPOLACTASIA
Twenty three adult white volunteers acted as normolactasic control subjects, 16 Asian or Afro-Caribbean together with two white subjects comprised the hypolactasic group. Hypolactasia was independently established by demonstrating a positive conventional lactose tolerance test (blood glucose rise <20 mg/100 ml and a diarrheal response after ingestion of 50 g lactose) and absence of intestinal disease apart from milk intolerance. A lactose and sucrose free diet was implemented 18 hours before, and throughout, each test. After an overnight fast each subject collected a baseline urine sample, which ensures that errors of interpretation due to inadequate dietary restriction, unsuspected lactulose medication, endogenous lactosuria of pregnancy, etc, are avoided. Patients then ingested a test solution containing lactose (20 g) sucrose (20 g) and lactulose (6.7 g) dissolved in 300 ml water. A complete urine collection was then made in two consecutive five hour periods (that is, 0–5 hours and 5–10 hours). The urine volumes were recorded and aliquots preserved (10 mg merthiolate/100 ml minimum) for quantitative analysis of lactose, sucrose, lactulose, and galactose by quantitative paper chromatography or an adaptation of this method to thin layer chromatography. Results were expressed as percentage of the dose of each sugar excreted in urine during 0–5 or 0–10 hour period and as lactose-sucrose/lactulose ratios (of percentages excreted). Lactose/galactose urine excretion ratios were also calculated to assess whether this ratio gave a better discrimination of hypolactasic from normolactasic subjects than lactose or galactose alone as has previously been suggested. 7 8

INVESTIGATION OF PATIENTS WITH COELIAC DISEASE
Nineteen patients undergoing jejunal biopsy as a part of routine gastrointestinal investigation, in whom jejunal histology was normal and significant intestinal pathology not discovered, served as a control group. The mean age of this group was 41 years (range 21–67 years).

Eleven patients with symptomatic, untreated, and nine with treated coeliac diseases were studied, the latter having achieved a full clinical recovery with a gluten free diet of mean duration six years (range nine months–12 years). The mean age of the whole group was 54 years (range 28–74). Each patient was admitted to the metabolic research ward for investigation, in vivo studies and jejunal biopsies being performed within a week of each other.

Assessment of intestinal disaccharide hydrolysis by urinary excretion ratio
Dietary sources of sucrose, lactose, and lactulose were excluded for 18 hours before and throughout the test period. Each subject fasted overnight and voided urine directly before the test, a ‘baseline’ urine being retained to exclude the presence of disacchariduria. At 8 am each subject ingested a 300 ml test solution containing: Lactose: 10 g (to assess lactase activity), Sucrose: 10 g (to assess sucrase activity), Palatinase: 10 g (to assess palatinase (isomaltase) activity (α-1-6 D glucopyranosyl-D-fructofuranose, isomaltulose), Lactulose: 6.7 g (as non-hydrolysable reference for disaccharide permeation, also to assess ‘large pore’ permeation), L-rhamnose: 1 g (to assess ‘small pore’ permeability).

Incorporation of L-rhamnose permits simultaneous assessment of intestinal permeability by measurement of lactulose/L-rhamnose urine excretion ratios, but as the data from this study did not expand on the already extensive published literature these results are not presented. 13

Foods and fluids were permitted 2.5 hours after ingestion of the test solution.

Urine was collected for 0–5 and 5–10 hours, with merthiolate (10 mg/100 ml) as a preservative, for marker analysis as described. 17 18

Assessment of intestinal disaccharide activity by in vitro assay
Intestinal biopsy specimens were obtained by intubation technique with an adult Watson biopsy capsule, under radiological guidance, just distal to the ligament of Trietz. A portion of the biopsy specimen was allocated for histological examination, the remainder being
Differential disaccharide excretion and intestinal disaccharidase activity

Figure 2: Use of the disaccharide ratio method to compare the dose/response relation of three different α-glucosidase inhibitors on intestinal sucrose hydrolysis. This is assessed by measuring the excretion ratio of sucrose/lactulose (% dose in a 10 hour urine collection) following ingestion of the disaccharide test solution together with increasing dose of BAY g 5421 (Acarbose), BAY m 1099, and BAY o 1248. Equivalent effects were produced by 0·2 g, 0·1 g, and 10 mg doses of these inhibitors, respectively.

‘flash’ frozen in liquid nitrogen and stored at −70°C. No more than two minutes lapsed between firing the capsule and freezing the samples. Lactase and sucrose activities in homogenised biopsy tissue were measured by a micro-assay as described previously.19 The same method was used for measurement of isomaltase activities using palatinose as the substrate, dissolving 0·056 mol/litre in 0·1 M sodium maleate buffer, pH 6·0. Enzyme units were expressed as units/mg protein. All subjects gave informed consent and the studies were approved by the Harrow Health Authority and St Thomas’s Hospital Ethical Committee.

RESULTS

STUDIES WITH α-GLUCOSIDASE INHIBITORS

Figure 2 shows the 10 hour urine sucrose/lactulose excretion ratios obtained when the test solution had been ingested without, and then together with progressively increasing doses of the three α-glucosidase inhibitors. The normal control 10 hour sucrose/lactulose excretion ratio (mean SD) was 0·12 (0·10), range 0·05–0·30.

With increasing amounts of simultaneously ingested BAY g 5421 (Acarbose 25–500 mg) there was stepwise and significant (p<0·01) increase in urine excretion ratio of sucrose/lactulose with a mean of 0·61 (0·12) and 0·91 (0·09) for 200 and 500 mg BAY g 5421, respectively. Ingestion of the two other α-glucosidase inhibitors also produced a significant (p<0·01) rise in urinary sucrose/lactulose excretion ratios. The relation between dose and response was very different, approximately similar effects being produced by 200 mg BAY g 5421, 100 mg BAY m 1099, and 10 mg BAY o 1248, giving mean sucrose/lactulose ratios of 0·61 (0·12), 0·58 (0·13), and 0·55 (0·08), respectively (see Fig 2).

Figure 3 shows sucrose/lactulose and lactose/lactulose responses obtained from a single subject receiving nine dose levels of BAY g 5421 (range 10–500 mg). The progressive rise in the sucrose/lactulose excretion ratio shows satisfactory resolution between different levels of α-glucosidase inhibition while absence of a significant change in lactose/lactulose excretion ratio provides evidence of specificity.

Figure 4 shows the rate at which the α-glucosidase inhibitory action of the three drugs declines. Whereas the sucrose/lactulose excretion ratio after ingestion of BAY g 5421 and BAY m 1099 fall off rapidly, reaching 30% of the initial value when the test dose is delayed by 30 minutes, the effect of BAY o 1248 is more prolonged, taking three hours to reach the 30% level. The resolution obtained seems to be satisfactory for investigating the duration of action of such drugs.
lactase activity. Sucrose excretion did not differ significantly between the groups and, although lactulose excretion was less in hypolactasic subjects, this did not reach a statistical significance. The 10 hour urinary excretion ratio of lactose/lactulose differed significantly between the groups and none of the 18 subjects defined as hypolactasic by other criteria had a ratio that was below the mean (2 SD) (0.24) for the normolactasic group. Alternatively, although the discrimination of hypolactasic subjects by excretion of lactose or galactose is incomplete in 10 hour urine samples, combination of these measurements, which respond to reduced lactase activity in opposite directions, as lactose/galactose excretion ratios provides almost complete discrimination. Sucrose/lactulose ratios in Figure 5 show that eight of the hypolactasic subjects have higher values than any of the normolactasic subjects, which could suggest a coexistent impairment of sucrose hydrolysis. However, this seems to be the outcome of a decrease in lactulose rather than an increase in sucrose excretion, and more likely to be the consequence of osmotic accumulation of fluid due to retention of unabsorbed lactose (due to hypolactasia) in the intestine, lactulose permeation being reduced by the resulting dilution and hurry. 20 Sucrose uptake would not be affected to the same extent because, unlike lactulose, it is largely hydrolysed before reaching the lower part of the small intestine where these osmotically induced effects are most pronounced.

**DISACCHARIDE HYDROLYSIS IN PATIENTS WITH COELIAC DISEASE**

*Non-invasive in vivo assessment*

Figure 6 shows the 10 hour urinary excretion of the four disaccharides and the calculated lactose/galactose excretion ratio. The results are expressed as the mean (SD) of all subjects. Median lactulose excretion was significantly higher than lactose and galactose in both the normolactasic and hypolactasic groups. Lactose excretion was also higher than galactose in both groups. The lactose/sucrose/lactulose excretion ratio was significantly greater in the normolactasic group than in the hypolactasic group. Median lactulose excretion was significantly higher in the normolactasic group than in the hypolactasic group. Median lactose excretion was significantly higher than galactose in both groups. The lactose/sucrose/lactulose excretion ratio was significantly greater in the normolactasic group than in the hypolactasic group. Median lactulose excretion was significantly higher in the normolactasic group than in the hypolactasic group. Median lactose excretion was significantly higher than galactose in both groups. The lactose/sucrose/lactulose excretion ratio was significantly greater in the normolactasic group than in the hypolactasic group.
lactulose, sucrose/lactulose, and palatinose/lactulose excretion ratios from patients without gastrointestinal disease and patients with untreated and treated coeliac disease. Only the differential urinary excretion of lactose/lactulose from patients with untreated coeliac disease differed significantly (p=0.047) from controls. Eight, one and seven of the coeliac group were found to have varying degrees of defective lactose, sucrose, and palatinose hydrolysis, respectively. The five hour urinary data showed less discrimination (data not shown) between controls and patients with coeliac disease.

**In vitro assay of jejunal biopsy tissue**

Figure 7 shows the in vitro lactase, sucrase, and palatinase (isomaltase) activities. The control levels were 76 (8), 107 (9), and 34 (2) units/mg protein, respectively. Corresponding values for patients with coeliac disease were, 11 (3), 36 (5), and 11 (2), all of which differed significantly (p<0.001) from controls. Two treated patients had normal activity for all three disaccharidases and three patients each had normal activity of a single disaccharidase.

There was no significant (p>0.1) correlation between the lactose/lactulose urine excretion ratio (0–10 hours) and lactase activities by in vitro assay within the control group (r=0.40) or collectively for all the subjects studies (r=0.28). Similarly, there was no significant (p>0.5) correlation between sucrose/lactulose and palatinose/lactulose urine excretion values and in vitro sucrase (r=0.00) and palatinase (r=0.08) activities in the whole group of patients.

![Figure 6: Ten hour urine excretion of disaccharides and ratios of lactose-sucrose-palatinose/lactulose from controls (open circles), untreated (filled boxes), and treated (filled circles) patients with coeliac disease.](image)

![Figure 7: In vitro lactase, sucrase, and palatinase activity in jejunal biopsy samples from controls (open circles) and patients with untreated (filled boxes) and treated (filled circles) coeliac disease.](image)
Despite full clinical recovery with gluten restriction, jejunal biopsy specimens from the treated patients were histologically abnormal showing partial villous atrophy, in keeping with previous studies.21

Discussion

Construction of non-invasive systems for assessing aspects of intestinal function by non-invasive methods such as recovery of orally administered probes in urine calls for attention to the detailed conduct of the test, especially selection of test substances with suitable properties.13 An important problem affecting such indirect procedures concerns the distortion of results by adventitious non-mucosal factors such as the rate of gastric emptying and intestinal transit, systemic distribution and metabolism, and renal clearance. Use of the 'differential absorption' principle to overcome such irrelevant variables is intrinsic to the strategy, illustrated in Figure 1, for simultaneous estimation of intestinal lactase, sucrase, isomaltase, and permeability based on the measurement of lactose, sucrose, palatinose, lactulose, and L-rhamnose excreted in urine following oral administration. Such test systems should be based on sound theory, but also shown to be reliable and convenient for clinical and research purposes.

Evaluation of an original method is usually undertaken by demonstrating ability to quantitate a specific function, in this study the activity of intestinal disaccharidases, with respect to the detection of a clinically important defect. This is usually achieved by comparison with previously established techniques. A familiar problem, however, concerns the reliability of test procedures available for reference purposes. Existing methods that depend upon the rise in blood monosaccharide concentration or generation of breath 14C-carbon dioxide or hydrogen after oral administration of disaccharide are not only affected by the rate of hydrolysis, but also by the state of intestinal absorption and systemic metabolism of the monosaccharide products. Newcomer and McGill25 have shown that as many as 25% of subjects without lactase deficiency may have misleadingly 'flat' blood glucose responses to lactose tolerance tests (a rise of less than 20 mg/100 ml following 50 g and 100 g oral lactose). The lactose-hydrogen breath test may become unreliable in the presence of monosaccharide malabsorption, bacterial overgrowth in the upper intestine, or of colonic bacteria incapable of generating hydrogen, and these possibilities require exclusion.23 24 In vitro assays performed on a homogenate of intestinal biopsy tissue are generally considered reliable evidence of disaccharidase status, but the measurements obtained relate to the region from which the biopsy tissue was taken and do not necessarily parallel the hydrolytic performance of the whole intestine.

During this study performance of the disaccharide ratio system was assessed by comparing responses obtained from healthy 'normolactasic' volunteers, lactase status being verified by the conventional lactose tolerance test, and also in context with temporary impairment of sucrase activity in healthy volunteers by drug induced α-glucosidase blockade where the test distinguishes between the potency of the α-glucosidase inhibitors and the duration of drug action. The three α-glucosidase inhibitors used for the latter purposes are complex oligosaccharides or, more precisely, pseudo-tetrasaccharides, which have been shown to inhibit intestinal sucrose in vitro and, by indirect methods, in vivo.25 26 The use and reliability of the disaccharide ratio method combining the use of lactose, sucrose, and palatinose with lactulose reference for the investigation of patients with primary intestinal sucrase-isomaltase, has already been investigated and found to be satisfactory.11 12

Subjects with primary hypolactasia had significantly higher urinary lactose/lactulose excretion ratios than those with normal intestinal lactase activity and the range of values (range 3.0–15.0) simulated that most had partial rather than complete lactase deficiency, which would be suggested had the ratio approached 1.0. At the same time the opportunity was taken to compare urinary excretion of galactose with that of lactose in healthy normolactasic and hypolactasic subjects. These results also produced satisfactory discrimination provided they were expressed as galactose/lactose excretion ratios. However well urinary or blood galactose estimations7 may discriminate, it must nevertheless be emphasised that galactose intolerance due, for instance, to ethanol intake,27 galactose pathway defects or hepatic disease, in addition to impairment of galactose absorption, will render interpretation unreliable while the excretion ratios of lactose/lactulose would remain unaffected.13 16 Furthermore, abnormal intestinal permeability due, for instance, to villous atrophy or to the effect of non-steroidal anti-inflammatory drugs28 will interfere with interpretation by producing a false rise in the galactose/lactose excretion ratio.

The suitability of the disaccharide ratio systems for assessing temporary changes in intestinal disaccharidase activity is well illustrated by the results of the study of the three α-glucosidase inhibitors, BAY g 5421, BAY m 1099, and BAY o 1248, reported here. These are shown in Figures 5 and 6 and indicate that the sensitivity and precision of the method is adequate to resolve the quantitative differences in dose response and also differences in the duration of disaccharidase inhibition over a short time interval.

Lactosuria and sucrlosuria have long been known to occur in patients with coeliac disease29 but the extent to which this was due to increased intestinal (paracellular) permeability or reduced disaccharide hydrolysis has not been clear.13 30 Blood glucose responses following the conventional lactose tolerance test generally suggest a high prevalence of lactase deficiency in untreated patients,31 but the outcome of this test depends as much upon intestinal monosaccharide transport as upon disaccharidase activity, both of which are often impaired in coeliac disease.31 Changes in
Differential disaccharide excretion and intestinal monosaccharide absorption may, similarly, interfere with tests that depend upon the rise in blood level or recovery in urine of galactose, of breath $^{14}$CO$_2$ derived from ingested $^{14}$C-lactulose, and also breath hydrogen, which is generated when either unabsorbed monosaccharide or disaccharide enters the colon. 7 This problem has usually been solved by repeating the test using an equivalent dose of monosaccharide. In vitro assay of intestinal biopsy tissue usually demonstrates the presence of lactase deficiency in coeliac disease regardless of treatment. The contract between estimates of intestinal disaccharidase activity in coeliac disease by in vitro assay of jejunal tissue and disaccharide hydrolysis assessed in vivo by the disaccharide ratio test is therefore a matter of interest and importance. Ratios of urinary disaccharide excretion, which are not affected by overall variations in monosaccharide transport capacity or permeability to disaccharide, suggest that impairment of intestinal disaccharide hydrolysis may be less prevalent in coeliac disease than suggested by the jejunal disaccharidase assays.

It is of note that although palatinoselactulose excretion ratios suggest that the in vivo intestinal hydrolysis of palatinose is as efficient as that of sucrose, in vitro assays show that jejunal palatinase activity is much lower than that of sucrose (34 (2) compared with 107 (9) mU/mg protein, respectively). Evidently palatinose is a satisfactory substrate for in vivo assessment of intestinal isomaltase activity but, unlike isomaltose, it seems to exert severe substrate inhibition under the conditions of in vivo assay. 83

In summary, the evidence presented suggests that measurement of urinary hydrolysable/non-hydrolysable disaccharide excretion ratios following oral administration of appropriate disaccharidases is a reliable non-invasive technique capable of providing simultaneous quantitative assessment of intestinal lactase, sucrase, and isomaltase activity. We have found the method convenient for studying the dose/response and temporary duration of action of $\alpha$-glucosidase inhibitors on sucrose hydrolysis, and for the investigation of lactose hydrolysis in subjects with primary (genetic) adult hypolactasia. A study of patients with coeliac disease shows that in most cases impairment of in vivo disaccharide hydrolysis is only moderate.

We thank Bayer UK for the gift of the $\alpha$-glucosidase inhibitors and Tate and Lyle Industries Ltd, Reading, Berks, UK, for the gift of palatinose used in these studies.