Improved specificity of the PABA test with p-aminosalicylic acid (PAS)

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SUMMARY  Until now use of the PABA test together with ["C"] PABA to calculate the PABA excretion index has probably been the best adaptation suggested to enhance the specificity of this non-invasive pancreatic function test. Drawbacks of the method are the application of radioactivity, the fact that children, pregnant women, and patients with renal insufficiency have to be excluded from the test, and the possible interference of drugs and isotopes. We propose simultaneous administration of p-aminosalicylic acid (PAS) in the PABA test and quantification of the urinary PABA and PAS excretion with liquid chromatography. Urinary PABA and PAS excretion in six hours are comparable (69.5±8.4% and 65.6±18.4% respectively in five healthy volunteers). Application of the PABA/PAS ratio was compared with the urinary PABA excretion in 21 normal controls, 38 patients with pancreatic disease, and 42 patients without pancreatic pathology. The PABA/PAS ratio and the per cent PABA excretion correlated very well in pancreatic patients: (PABA/PAS ratio)=0.0149(% PABA)+0.052 (r=0.902). Use of the PABA/PAS ratio enhanced the specificity of the test from 76 to 89%.

Since its introduction, the PABA test has gained an established position in the battery of tests available for the diagnosis of exocrine pancreatic insufficiency.1 Advantages of the test are the fact that no invasive procedures are necessary and that simple colorimetric procedures are available for the quantification of p-aminobenzoic acid (PABA) in urine.2±

It shares, however, a lack of specificity with other non-invasive and indirect pancreatic function tests.1 False low excretion rates can be obtained by defects in gastric emptying, in endogenous stimulation of the pancreas, failure in raising the intestinal pH, abnormalities in intestinal PABA resorption, in liver function and in renal excretion.

For a screening test to be used in a patient population with a low prevalence of the disease, a high specificity is of the utmost importance to obtain an acceptable predictive value of a positive result. Administration of an equimolar dose of free PABA on a separate occasion and calculation of a PABA excretion index has been recommended to compensate for abnormalities in PABA resorption and metabolism.1 With ["C"] PABA the duration of the procedure can again be brought back to six hours,1 but the radioactivity involved excludes use of the test in children, pregnant women, and patients with renal insufficiency and prevents its widespread use. No single report is available in the literature of use of this test modification outside the UK. Besides, drug or isotopic interference inhibits test interpretation in about 10% of the cases.4

Determination of PABA with high performance liquid chromatography (HPLC) is a simple and accurate procedure5,6 with the advantage of high analytical specificity. It permits the incorporation...
into the PABA test of a PABA-like marker substance. Calculation of the ratio of PABA and marker excretion should then eliminate influences of resorption, metabolism and excretion of PABA on the PABA test in the same way as the PABA excretion index does. A one day tubeless pancreatic function test is then obtained without radioactivity.

We propose p-aminosalicylic acid (PAS) as the marker substance, because it has a molecular structure very similar to PABA, consequently similar pharmacokinetic properties and it is readily separated from PABA in our HPLC assay. In this study data are presented of the PABA/PAS ratio, obtained in healthy volunteers and patients and a comparison is made with the results of the PABA test, expressed in the usual way.

Methods

Patients

The PABA test was performed with 1 g N-benzoyl-L-tyrosyl-PABA (BTPABA; bentiromide; Hoffman-La Roche, Basle Switzerland). The substrate and 360 mg PAS (500 mg PAS Na₂H₂O) were administered together with a Lundh test meal (150 ml). Urine was collected during six hours. In approximately one half of the tests the urine was collected in separate one hour portions. In the other patients and controls the urine was collected in one 6 hour portion. Always a urine sample from before the start of the test was analysed also to ensure the absence of substances which could interfere with the analysis.

After alkaline hydrolysis of the PABA and PAS metabolites in the urine, analysis was undertaken by high performance liquid chromatography (HPLC) on a reversed phase column (10 μm Lichrosorb RP 18) with a 3/1 mixture (by vol) of 0.01 M Na-acetate buffer pH 4.0 and methanol as the mobile phase. Details of the procedure are described elsewhere.

The amount of PABA recovered in the urine in six hours was expressed as a percentage of the PABA equivalents (340 mg) administered orally as BTPABA. The lower limit of normal PABA excretion was 50%. The amount of PAS, which was recovered in the urine, was also expressed as a percentage of the 360 mg PAS administered orally. The PABA/PAS ratio, the ratio of these percentages of recovered PABA and PAS was calculated.

The 12 healthy volunteers, participating in this study, were members of the hospital staff. Patient controls (n=9) were without organic disease which could possibly interfere with the PABA test. For the pancreatic disease group, 38 patients were analysed with established disease. Pancreatic disease was confirmed by ultrasound, computed tomography, ERCP, endosonography, abdominal radiograph or histology. Diagnosis was chronic pancreaticitis in 29 patients, pancreatic carcinoma in five patients, acute pancreatitis in three patients, and in one patient cystic fibrosis with pancreatic insufficiency. Pancreatic insufficiency was not always present. Faecal fat excretion was assessed in 21 patients and was abnormal in 18. The majority of the 42 patients in the group with gastrointestinal disease was investigated because of upper abdominal complaints and/or diarrhoea. In these patients no independent proof of pancreatic involvement was obtained. In addition some patients with established malabsorption syndrome (n=7), impaired gastric motility (n=3), or severe liver cirrhosis (n=3) were investigated.

Results

For use of PAS instead of PABA as internal reference of resorption, metabolism, and excretion, a comparison of their respective pharmacokinetics is essential. Therefore 340 mg PABA and 360 mg PAS (500 mg PAS Na₂H₂O) were administered together to five healthy volunteers with 150 ml Lundh test meal. Urine was collected in one hour portions and PABA and PAS measured therein. Urinary excretion in six hours amounted to 69.5 ± 8.4% (X ± 2 SD) for PABA and to 65.6 ± 18.4% (X ± 2 SD) for PAS. Over the six hour period PABA and PAS excretion proved to be very comparable (Fig. 1). The PABA excretion was not different from data obtained in earlier experiments in five healthy volunteers after administration of 340 mg PABA alone with 150 ml Lundh test meal, measured with the colorimetric method. A different urinary excretion curve was

![Graph](http://gut.bmj.com) Fig. 1 Comparison of the time curves of urinary excretion of PABA and PAS. Mean ± SD are given after oral administration of 340 mg PABA (○), 500 mg PAS Na₂H₂O (□) and 1 g BTPABA (●) together with a test meal to 5 healthy volunteers.
found for PABA, when it was administered as BTPABA with the test meal (Fig. 1), although the total excretion in six hours was equal.

The PABA/PAS ratio was evaluated in 12 healthy volunteers and nine patient controls, in 38 patients with evidence of disease of the exocrine pancreas and in 42 patients with gastrointestinal or liver disease without evidence of exocrine pancreatic involvement. After administration of 1 g BTPABA and 360 mg PAS together with the test meal, urinary PABA excretion in the 12 healthy volunteers was 65.3 ± 20.0% (× ± 2 SD). With the exception of one abnormal test result (43.4%), this PABA excretion was in good agreement with the value reported earlier for 20 healthy volunteers, 74.0 ± 18.8% (× ± 2 SD) measured with the colorimetric method.\textsuperscript{11} PAS excretion in these healthy volunteers amounted to 74.2 ± 18.3% (× ± 2 SD) of the administered dose. The PABA/PAS ratio for the 12 healthy volunteers was 0.89 ± 0.28 (× ± 2 SD) with values ranging from 0.62 to 1.15.

In five healthy volunteers PAS excretion was measured once together with BTPABA and once with free PABA. Excretions measured were 74.7 and 73.3%, 65.7 and 52.7%, 54.5 and 59.0%, 79.7 and 73.0%, 70.1 and 69.8% respectively. Calculation of the PABA excretion index was also possible for these persons. Results for the PABA/PAS ratio and the PABA/PAS ratio for these five volunteers are given in the Table.

Based on the results in the 21 normal controls 0.75 was arbitrarily accepted as lower limit of normal for the PABA/PAS ratio. The lower limit of normal for PABA excretion was 50%. The results for the normal controls and the patients are given in Figure 2, expressed as per cent PABA recovered and as PABA/PAS ratio. In 25 of the 38 patients with pancreatic disease the PABA/PAS ratio was abnormal, giving a sensitivity of 66%, while PABA excretion was <50% in 28 patients (74%). In the group of patients without pancreatic disease and the controls the PABA/PAS ratio was abnormal in seven of 63 cases, resulting in a specificity of 89%, compared with an abnormal PABA excretion in 15 of the 63 cases (specificity 76%). Results of the PABA/PAS ratio and the six hour PABA excretion are compared in Figure 3.

An excellent correlation between PABA/PAS ratio and PABA excretion was found for the patients with pancreatic disease: (PABA/PAS ratio)=0.0149 (% PABA)+0.052 (r=0.902; n=38). The correlation was much weaker for the patients without pancreatic disease: (PABA/PAS ratio)=0.0056 (% PABA)+0.55 (r=0.510; n=42).

When a prevalence of pancreatic disease of 15% was assumed,\textsuperscript{4} a positive predictive rate was calculated of 35% for the PABA test and of 50% for the PABA/PAS ratio. The predictive value of a negative result for absence of pancreatic disease was 94% for both tests.

**Discussion**

It is well known from the literature that the limited specificity of the PABA test for exocrine pancreatic insufficiency can be improved by use of a PABA excretion index, to compensate for the influence of non-pancreatic factors on PABA excretion. Originally, this compensation was achieved by administration of an equimolar dose of free PABA on a different day and determination of its excretion as a reference.\textsuperscript{5} Later the use of a small dose of [\textsuperscript{13}C] PABA was described.\textsuperscript{6,7} This limited the procedure to six hours and eliminated the day to day variations in PABA excretion observed by Kay et al\textsuperscript{8} and ascribed by them to variations in gastric emptying. Radioactive compounds are, however, used only reluctantly and they prohibit use of the test in children, pregnant women, and patients with renal insufficiency.

The excretion index described by us here also has the advantage that it takes only a single six hour urine collection, but in addition it puts no restrictions on the patients to be investigated. For a reliable calculation of a PABA excretion index it is essential that the pharmacokinetic properties of the marker resemble the kinetics of PABA metabolism and excretion as closely as possible. Therefore we started to look for compounds with a molecular structure strongly related to PABA. PAS was chosen for the following reasons: \(1\) PAS differs from PABA only by one hydroxyl group. \(2\) In the literature data were available to prove its comparable metabolism\textsuperscript{11} and its high urinary excretion rate,\textsuperscript{12} supported later by our data reported here (Fig. 1). \(3\) PAS has been in use as a tuberculostatic drug for a long time in the recent past and much higher doses than used in this study have been administered safely to patients. Therefore we could be sure that no side effects were to be expected with the dose used in this study. This obviated the necessity of prior toxicologic studies. \(4\)

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**Table** Comparison of PABA/PAS ratio and PABA excretion index (PEI) in five healthy volunteers

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In our HPLC assay for PABA, PAS was detected also with a retention time suitable for accurate quantification of both compounds.

As reported here for five healthy volunteers day to day variations in PAS excretion were not always negligible, confirming the day to day variations in PABA excretion reported by Kay et al. As a consequence, in one case a difference of 0.22 was found between PABA/PAS ratio and PABA excretion index (Table), although in the other case both parameters corresponded very well.

It may be pointed out here that small differences in excretion rate between the chosen marker and released PABA will not decrease the value of the excretion index. Also free PABA resorption and excretion are not completely parallel to the resorption and excretion of PABA, that has to be split off first from BTPABA in the duodenum. In healthy controls it was shown that with free PABA there was a smaller lag between oral ingestion and urinary excretion, even when taken with a meal (Fig. 1). The difference may probably be explained by differences in gastric emptying of on the one hand free PABA, which is readily soluble at the low pH of the gastric contents, and on the other hand BTPABA, which is virtually insoluble at pH<5. For PAS, also acid soluble, only a slightly lower excretion rate was observed than for free PABA (Fig. 1) and PAS excretion differed therefore to the same extent as free PABA did from the excretion of PABA produced by BTPABA proteolysis in vivo.

The optimal dose of BTPABA is probably determined by the limited capacity of the body for handling PABA. This limited PABA excretion

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Fig. 2  Urinary PABA excretion and PABA/PAS ratio for normal controls (○), patients with pancreatic disease (●) and patients with gastrointestinal or liver disease (□). The dotted lines indicate the lower limits of the normal range.
Fig. 3 Relationship between PABA/PAS excretion ratio and urinary PABA recovery. Results for 21 normal controls (○), 38 patients with pancreatic disease (●) and 42 patients with gastrointestinal or liver disease (□). The regression line for patients with pancreatic disease is: \( (\text{PABA}/\text{PAS}) = 0.0149 \times (\% \text{PABA}) + 0.052; r = 0.902 \). The dotted lines indicate the lower limits of the normal range.

capability has been suggested as an explanation for some false positive tests observed when a 2 g BTPABA dose was used. The urinary excretion curves for high doses of PAS reported in the literature do not suggest a comparably limited capacity for PAS excretion. In the results obtained by us with combined administration of PAS and PABA, or PAS and BTPABA, no indications could be found for restricted PABA excretion. Still, the analytical precision of the HPLC procedure is such, that BTPABA and PAS doses may probably be halved without damage to the diagnostic accuracy of the test.

The data obtained in the patients and controls (Figs 2 and 3) support our opinion that PAS can be used instead of \(^{14}\text{C}\) PABA to obtain a reliable PABA excretion index. These encouraging results will have to be confirmed in more extensive patient studies.

References

5 Mitchell CJ, Humphrey CS, Bullen AW, Kelleher J, Losowsky MS. Improved diagnostic accuracy of a
Improved specificity of the PABA test with p-aminosalicylic acid (PAS)


