Investigation of intestine function during acute viral hepatitis using combined sugar oral loads

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SUMMARY One fifth of all cases of A virus hepatitis (AVH) have symptoms of gastroenteritis at the onset. This study investigated the mediated intestinal absorption of D-xylose (D-xyl) and 3-o-methyl-D-glucose (3-omG) and the non-mediated permeation of lactulose (LacI, mol wt 342) and L-rhamnose (L-rh, mol wt 164) during acute and remission phases of AVH. Ten patients with AVH were given an oral load containing these sugars (5 g D-xyl: 2.5 g 3-omG, 1 g L-rh, 5 g lacI in 250 ml water) once during the acute phase and again during remission. The same load was given once to a group of 22 healthy controls. The mean concentration of D-xyl in urine and the ratio of D-xyl to 3-omG in plasma and urine were normal in both the AVH phases, ruling out intestinal malabsorption even in the acute phase. This study showed a significant increase in non-mediated permeation to LacI, but not to L-rh, during the acute phase. These data indicate that the barrier function of the intestine is compromised in AVH infection while the absorptive function is not. An abnormally low concentration of D-xyl and 3-omG in plasma at one hour was found in all patients during the acute phase. This finding cannot be explained by alterations in intestinal absorption, but could be accounted for by increased space distribution of the sugars because of increased diffusion into tissue cells and/or expansion of the extracellular space by fluid retention.

A-virus hepatitis (AVH) is associated with gastroenteritis symptoms in approximately one of five cases. Recent reports have suggested direct replication of the A hepatitis virus in intestinal cells, which could result in altered intestinal function. An acquired monosaccharide malabsorption is often observed in the course of viral gastroenteritis and can persist after disease regression. Early studies of intestinal absorption during viral hepatitis have given inconsistent data. In these studies urinary excretion of D-xylose after oral load was used as a measure of active absorption of monosaccharides. Modifications to this test have recently been proposed to correct interference by variables such as distribution space, gastric emptying, and renal clearance. One of these modifications is the determination of the D-xylose (D-xyl) plasma concentration peak at one hour corrected by body surface area (Haeney’s method). Another modification is the addition to D-xyl of a second absorption marker, 3-o-methyl-D-glucose (3-omG) which is completely excreted in the urine unmetabolised and whose absorption is much less hindered by intestinal disease than is that of D-xyl. This modification allows the correction of plasma and/or urine xylose concentrations by calculating the ratio of D-xyl to 3-omG (Menzies’ method). Because little is known about intestinal dysfunction during viral hepatitis, we designed a sugar absorption study using these new procedures, in patients with AVH.

The measurement of passive intestinal permeability has been recognised recently as a more sensitive tool to explore the functional integrity of the gut. To include this new evaluation we have used a multiple sugar oral load consisting of D-xyl, 3-omG, lactulose (LacI) and L-rhamnose (L-rh). This test allows simultaneous evaluation of mediated absorption (D-xyl and 3-omG), and non-mediated
permeation to small molecules (L-rh, mol wt 164), which are passively absorbed through cell aqueous pores as well as to disaccharide molecules (Lacl, mol wt 342), whose permeation is thought to be through larger channels related to the tight junctions and/or cell extrusion zones.

An increase in non-mediated intestinal permeation to lactulose was recognised in the acute phase of AVH, while the mediated absorption appeared to be unaffected.

**Methods**

**Patients and Subjects**

Twenty-two healthy subjects and 10 patients with AVH participated as volunteers after giving informed consent. Hepatitis was diagnosed by typical clinical features, raised transaminase values and the presence of antiviral A IgM. After overnight fast, each subject was given an oral load consisting of 5 g D-xyl, 2.5 g 3-omG, 1 g L-rh and 5 g Lacl in 250 ml water. Hepatitis patients received this load on two occasions: early the first morning after admission to hospital (within a week of hepatitis symptom onset) and another 30 days after (regression phase).

No subject showed intolerance to the load; the osmolarity of the test solution was kept below the level known to induce an alteration in intestinal permeability or motility (340 mOsm/l). Heparinised blood was obtained before and at 30, 60, 90, and 120 minutes after oral load, and plasma samples were stored at −20°C. A complete five hour urine collection was made after the oral load and an aliquot of each urine collection was preserved with merthiolate (10 mg/ml), and frozen. Monosaccharides in plasma and urine were estimated by quantitative thin layer chromatography according to Menzies. Sugar separation was achieved by multiple development on half plates (10 cm×20 cm) of F1500 plastic backed silica gel (Schleicher and Schull, Dassel FRG), using for sugars in the plasma (D-xyl, 3-omG) three consecutive ascending runs (8.5 cm each), one with solution A (Butan-l-o/ethanol-o/acetic acid/water; 60:50:10:10) and two with solution B (Butan-l-o/ethyl-acetate/pyridine/acetic-acid/water; 5:70:15:10:10). For sugars in the urine (D-xyl, 3-omG, L-rh) three consecutive ascending runs (8.5 cm each) were made with solution B. The precision of both methods was 7%, recovery above 90% and minimum level of detection below 0.1 mmol/l for the three sugars tested.

Lactulose was estimated in urine by gas liquid chromatography according to Laker’s method with modification in the gas-carrier and in its flow rate. The chromatograph was a Fractovap 4002 from Carlo Erba (Italy). The operative conditions were: oven 250°C, injector 275°C, detector (ionisation flame) 305°C, gas-carrier (N₂), flow 35 ml/min, attenuation input 1:100 and output 1:2, back-off 1:100. The column was a 3-metre OV 17 (8% phases of gas-chrom Q 80–100 mesh). The lactulose retention time relative to that of turanose (internal marker 10 mg/100 ml) was 0.73. The variation coefficient was above 8% and the minimum concentration value of lac in urine was 80 μg/ml. Values representing timed D-xyl and 3-omG concentrations in plasma for each test were processed (Sperry Univac personal computer) calculating second to tenth degree equations with their integrals. The fourth degree equation, according to correlation coefficients, most accurately expressed the kinetics of the two sugars in plasma. This was used to calculate the area under the 0–120 min plasma curves (PCTA) for each patient and control subject. The values for each group were statistically analysed according to the Student's t and Wilcoxon's tests. Patient biochemical data and sugar parameters were compared according to Pearson's correlation coefficient.

**Results**

Table 1 shows biochemical data of the hepatitis patients. All were icteric at the onset of the disease; bilirubin and transaminase values had returned to control levels at the end of the thirty day study period. Renal function was normal in patients and controls, as proven by creatininaemia and urine data.

All patients were untreated throughout the observation period. Figure 1 shows the mean plasma levels of D-xyl and 3-omG at different times after oral load.

Figure 2 reports plasma concentrations of D-xyl and 3-omG at one hour in each patient and control. The concentrations of both sugars were lower than normal in the acute phase of hepatitis and had returned to normal 30 days later.

Table 2 shows the mean one hour xyloxaemia corrected according to Haeney, the mean plasma concentration/time area (PCTA) of D-xyl and 3-omG in patients with type A hepatitis (AVH) in acute and remission phases (mean values±SD).

The measure units and normal range are reported in brackets.

<table>
<thead>
<tr>
<th>Biochemical Data</th>
<th>Acute phase</th>
<th>Remission phase</th>
</tr>
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<tbody>
<tr>
<td>AST (0–40 U/l)</td>
<td>1539±442</td>
<td>35:4±9:3</td>
</tr>
<tr>
<td>ALT (0–40 U/l)</td>
<td>2472±884</td>
<td>47:8±1:0</td>
</tr>
<tr>
<td>Bilirubin</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AVH (10)</td>
<td></td>
<td></td>
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<tr>
<td>AGE (17–23)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tot (0–2–1 mg/dl)</td>
<td>91±1·7</td>
<td>0·8±0·3</td>
</tr>
<tr>
<td>Glutamic (70–100%)</td>
<td>86±8·9</td>
<td></td>
</tr>
<tr>
<td>Glucose (60–120 mg/dl)</td>
<td>78±8·9</td>
<td></td>
</tr>
<tr>
<td>Creatininaemia</td>
<td>(0·7–1·5 mg/dl)</td>
<td>0·72±0·04</td>
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</table>
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![Graphs showing plasma concentrations of D-xylose and 3-O-methylglucose in AVH remission-phase and acute-phase, as well as normal values.]

**Fig. 1** Plasma values (mean ± SD) of D-xylose and 3-O-methylglucose in 22 normal subjects and in 10 patients with A viral hepatitis (AVH), in the acute phase of disease and 30 days after.

**Fig. 2** One hour plasma concentrations of D-xylose (D-xyl) and 3-O-methylglucose (3-omG) in acute phase (0) and 30 days after (30) in patients with acute viral hepatitis and in controls (C), with mean values.

3-omG, and the mean plasma ratio of D-xyl to 3-omG. Only the D-xyl/3-omG ratio shows no significant variation among acute and remission phases of AVH and controls.

Table 3 shows per cent recovery of the different sugars in urine, and the D-xyl/3-omG and Lac/l-rh ratios. The excretion of D-xyl and 3-omG was similar in patients during both AVH phases and in controls.

<table>
<thead>
<tr>
<th>Table 2</th>
<th>Plasma parameters of sugars after combined oral loads</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>AVH</td>
</tr>
<tr>
<td></td>
<td>Controls (C)</td>
</tr>
<tr>
<td>D-xyl</td>
<td>0.77±0.10</td>
</tr>
<tr>
<td>D-xyl/3omG</td>
<td>0.85±0.18</td>
</tr>
<tr>
<td>PCTA D-xyl</td>
<td>1119±219</td>
</tr>
<tr>
<td>PCTA 3omG</td>
<td>1037±203</td>
</tr>
</tbody>
</table>

*Haney's correction; †mmol/l; ¶Wilcoxon's test; $Student's t$ test; PCTA = plasmatic concentration/time curve (mg×min⁻¹).
and therefore the D-xyl/3-oMG ratio remained unmodified; L-rh excretion was also unmodified during both disease phases with respect to controls. LacI urinary excretion, however, underwent significant variations during hepatitis. In the acute phase it was three times greater, but had returned to near normal levels 30 days later. The LacI/L-rh ratio followed the same course. Figure 3 shows this pattern of lactulose excretion in each patient in the two phases studied, and the control data.

### Discussion

Our study revealed a functional gut derangement in acute AVH which was still evident during remission phase. We observed a marked increase in non-mediated intestinal permeation to LacI at the onset of hepatitis, while the transmucosal transfer of small molecules like L-rh remained unaffected. Two distinct pathways are hypothesised for unmediated mucosal permeation, one consisting of small pores in the enterocyte cell membrane through which L-rh passes, and the other of larger channels related to the tight junctions and/or enterocyte extrusion zones.15 As LacI is a probe molecule of the latter pathway our results indicate damage to these paracellular mechanisms. An analogous phenomenon has been described in allergic conditions such as atopic eczema,16,17 in rheumatoid arthritis,18 and during cytotoxic therapy.21 The mechanisms involved in these pathological conditions are at present unknown. In the acute phase of hepatitis it is possible that a transient increase in portal pressure with a secondary alteration in cell metabolism is capable of influencing the paracellular pathways. An intestinal inflammation is less likely to occur because in typical acute gastroenteritis by rotavirus or adenovirus absorption of LacI is increased and L-rh absorption is decreased,22 while our results showed a normal urinary recovery of L-rh. It is of interest that an abnormal increase in intestinal permeation to LacI is also present in coeliac disease.23 This disease, however, is characterised by marked malabsorption of monosaccharides such as D-xyl and L-rh, as a consequence of reduced absorptive gut surface as a result of mucosal atrophy.24 In intestinal biopses

### Table 3  Urine excretion after combined sugar loads

<table>
<thead>
<tr>
<th></th>
<th>5 h excretion (% oral dose)</th>
<th>AVH Controls (C)</th>
<th>AVH Basal (A)</th>
<th>AVH 0-030</th>
<th>P</th>
<th>AVH CV A*</th>
<th>AVH CV B*</th>
<th>AVH A B*</th>
</tr>
</thead>
<tbody>
<tr>
<td>D-xyl</td>
<td>34.0±6-3</td>
<td>34.0±6-3</td>
<td>29.7±3-9</td>
<td>ns</td>
<td>ns</td>
<td>ns</td>
<td>ns</td>
<td>ns</td>
</tr>
<tr>
<td>3-oMG</td>
<td>56.2±6-5</td>
<td>55.7±6-4</td>
<td>56.0±5-4</td>
<td>ns</td>
<td>ns</td>
<td>ns</td>
<td>ns</td>
<td>ns</td>
</tr>
<tr>
<td>D-xyl/3-oMG</td>
<td>0.61±0.09</td>
<td>0.55±0.05</td>
<td>0.52±0.02</td>
<td>ns</td>
<td>ns</td>
<td>ns</td>
<td>ns</td>
<td>ns</td>
</tr>
<tr>
<td>L-Rh</td>
<td>10.2±4-3</td>
<td>10.7±2.25</td>
<td>9.30±0.3</td>
<td>&lt;0.001</td>
<td>&lt;0.05</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Lacl</td>
<td>0.24±0.1</td>
<td>0.76±0.29</td>
<td>0.39±0.16</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Lacl/L-Rh</td>
<td>0.023±0.01</td>
<td>0.076±0.036</td>
<td>0.04±0.01</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
<td>&lt;0.01</td>
</tr>
</tbody>
</table>

*Wilcoxon’s test; †Student’s t test.
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taken during AVH, only inconstant and non-specific minimal lesions have been observed, and there is no decrease in cell mass of the small intestine mucosa. The normal D-xyl and L-rh excretion in urine we observed in both acute and remission phases of hepatitis are in agreement with these findings; thus sugar malabsorption can be excluded. Approximately 50% of iv administered D-xyl is metabolised in man, mainly by the liver. The tendency for urinary xylose to increase in the presence of reduced hepatic metabolism of this sugar could obscure a possible decrease caused by coincident malabsorption. The unaltered excretion during the acute phase of hepatitis of L-rh, however, which is much less efficiently metabolised by the liver than is D-xyl, indicates that absorptive capacity remained normal.

The ratio of D-xyl to 3-omG in plasma allows a correct evaluation, according to the principle of differential absorption of the intestine’s absorptive capacity, and is as reliable a test as xylose urinary excretion. In our study the D-xyl/3-omG ratios in plasma and urine of hepatic subjects did not differ from those of healthy controls, thus confirming the absence of malabsorption even in the acute phase of hepatitis.

The plasma curves of D-xyl and 3-omG, however, were abnormally low in the acute phase of AVH, as were the plasma concentration/time area (PCTA) and the one hour sugar concentration. These findings bear a statistical correlation with hepatic cytonecrosis (Alanine aminotransaminase, ALT, v D-xyl one hour r = -0.76, p = 0.01; ALT v 3-omG one hour r = -0.71, p = 0.022), and are the inverse of what one would expect in the case of reduced hepatic metabolism (which, furthermore, is possible only for D-xyl). These data cannot be related to alterations in intestinal absorption, but may possibly be related to increased space distribution of the test sugars within the body because of the expansion of the extra-cellular space by fluid retention and/or increased diffusion into tissue cells.

An altered water sodium homeostasis with minor abnormalities in water excretion has been reported in acute hepatic disease not complicated by renal failure as shown by altered response to water loading. On the other hand, increased permeability of blood capillaries, perhaps mediated by bile salt retention, has been observed in acute phase hepatitis by experiments with Evans blue diffusion.

In conclusion, our results rule out a monosaccharide malabsorption during acute AVH but disclose a marked increase in non mediated intestinal permeation to larger molecules like disaccharides. These data suggest a temporary alteration in large paracellular channels of the small intestine and appear difficult to explain by virus replication within intestinal cells. The low D-xyl and 3-omG concentrations in plasma could be interpreted by haemodilution and/or increased diffusion into tissue. The evaluation of the ratios of D-xyl to 3-omG in plasma and urine seem to express more satisfactorily, than other methods the small intestine’s absorption capacity to monosaccharides.

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References

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