Splanchnic metabolism of propylene glycol infused into the jugular vein of steers under washed rumen conditions*

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ABSTRACT

The splanchnic metabolism and blood to rumen flux of propylene glycol (PG) infused into the jugular vein of steers, implanted with permanent indwelling catheters in major splanchnic blood vessels as well as a rumen cannula, was investigated under washed rumen conditions. Up to 95% of the infused PG was taken up in splanchnic tissues. The hepatic uptake of PG accounted for up to 70% of the PG infused though the hepatic extraction ratio was only 8%. Up to 9% of the infused PG was transferred to the buffer incubated in the washed rumen. Increased hepatic balance of L-lactate could account half of the hepatic PG uptake. In conclusion the liver is the quantitatively most important organ for PG uptake from the blood, however, up to about one third of blood PG could be transferred back to the lumen of the gastrointestinal tract. L-lactate is a quantitatively important product of PG metabolism in the ruminant liver.

KEY WORDS: propylene glycol, splanchnic metabolism, steer

INTRODUCTION

In a previous study it was observed that PG accumulated in the blood of cows when absorbed from the washed rumen and no response to PG was observed for the whole body irreversible flux of glucose (Kristensen et al., 2002). The aim of the present study was to investigate the splanchnic metabolism of PG infused into the jugular vein of steers and test the hypothesis that blood PG is partly transferred to the lumen of the gastrointestinal tract and therefore have a higher availability for microbial metabolism than apparent from the relative fermentation and absorption rates of PG in the rumen.

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MATERIAL AND METHODS

Three Holstein steers fitted with permanent indwelling catheters in the mesenteric artery, mesenteric vein, hepatic portal vein, hepatic vein, and in the right ruminal vein as well as a ruminal cannula (#2C rumen cannula, Bar Diamond Inc., Parma, ID) were sampled 70±21 days after surgery, with an average body weight of 302±20 kg. One steer was sampled per sampling day. Continuous infusion of para-aminohippuric acid (pAH, 17.3±0.7 mmol/h) into the mesenteric vein and propylene glycol (PG, 119±2 mmol/h) into the jugular vein was initiated. The rumen was emptied, washed, and incubation of VFA buffers, as well as continuous intraruminal VFA infusion to maintain VFA absorption, was initiated as previously described (Kristensen and Harmon, 2004). The steers were subjected to 3 buffer incubations each of 125 min. The rumen was emptied and washed between each of the buffer incubations. Blood samples were simultaneously drawn from the arterial, portal, and hepatic catheters 60, 90 and 120 min after initiation of each buffer incubation. Plasma concentrations of PG were determined by the method described by Needham et al. (1982) with the following modifications: plasma (450 µL) was added 100 µL 5 mM 1,2 butanediol as internal standard and deproteinized with 4 mL of acetonitrile. The supernatant was dried under a stream of nitrogen at 70°C. One milliliter of 1% p-bromophenylboric acid in ethyl acetate was added and incubated for 30 min. The sample was dried and resuspended in 100 µL of cyclohexane. Samples were analysed by gas chromatography (Varian 3400, Palo Alto, USA), using a HP Ultra-1 column (25 m, 0.3 mm ID, 0.17 µm film) at a head pressure of 8 psi and an initial temperature of 60°C increased at 10°C/min to 200°C. Plasma concentrations of propionate, L-lactate, D-3-hydroxybutyrate, glucose, L-glutamate, and L-glutamine were determined as described previously (Kristensen and Harmon, 2004). Net portal fluxes and net hepatic fluxes were calculated as previously described (Kristensen and Harmon, 2004). Means within animal and buffer incubation were analysed by the repeated measures ANOVA of the GLM procedure of SAS (SAS Inst., Inc., Cary, NC). Net portal flux and net hepatic fluxes of lactate, glucose, 3-hydroxybutyrate, propionate, glutamate, and glutamine observed in the present study were compared to observations on the same three animals under identical experimental conditions except for the PG infusion by a paired t-test.

RESULTS AND DISCUSSION

The arterial concentration of PG increased (P=0.001) with time of infusion reaching 1.79 mmol/L blood in the third period (Table 1). In agreement with a lower infusion rate of PG, relative to the body weight in the present study, the arterial concentration of PG remained below the concentrations found in a
previous study with dairy cows (Kristensen et al., 2002). The net portal flux of PG was negative, indicating uptake of arterial PG across the PDV (portal drained viscera), however, the net portal flux was not affected by period (P=0.75). The net hepatic flux of PG was negative and about three times larger than the net portal flux. The PG uptake in the liver increased with period (P=0.01), the largest numerical increase was from period 1 to 2. The hepatic extraction ratio was 8%, but despite of a relatively low hepatic extraction ratio the liver removed more than 70% of the PG infused during period 2 and 3. The splanchnic tissues removed 95% of the infused PG during period 3. The generally larger differences observed between period 1 and 2 compared with the differences between 2 and 3 could indicate that infusion and utilization of PG was approaching a steady and that that PG during the first hours of infusion was accumulating in body fluids.

Table 1. Arterial concentration, net portal flux, net hepatic flux, and hepatic extraction ratio of propylene glycol (PG) as well as the proportion of propylene glycol infused into the jugular vein, taken up in splanchnic tissues in steers under washed rumen conditions

<table>
<thead>
<tr>
<th>Item</th>
<th>Period</th>
<th>SEM</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arterial concentration, mmol/L</td>
<td>0.89</td>
<td>1.45</td>
<td>1.79</td>
</tr>
<tr>
<td>Net portal flux, mmol/h</td>
<td>-18</td>
<td>-27</td>
<td>-28</td>
</tr>
<tr>
<td>Net hepatic flux, mmol/h</td>
<td>-56</td>
<td>-84</td>
<td>-85</td>
</tr>
<tr>
<td>Hepatic extraction ratio</td>
<td>0.08</td>
<td>0.08</td>
<td>0.07</td>
</tr>
<tr>
<td>Proportion of infused PG taken up in splanchnic tissues</td>
<td>0.61</td>
<td>0.93</td>
<td>0.95</td>
</tr>
</tbody>
</table>

The ruminal concentration of PG at the end of ruminal incubations, the ruminal accumulation rate of PG and the fraction of the hourly infused PG recovered in the ruminal buffer increased with period (P<0.05; Table 2). In average 30% of the PG taken up in the PDV was recovered in the ruminal buffer. To what extent the PDV tissues metabolized PG or all PDV uptake was simply transfer to the lumen of the gastrointestinal tract is not known. However, due to the fact that 20% of the infused PG was taken up across the PDV and 30% of that was found in the rumen, PG seems to have a higher availability to microbes in the gastrointestinal tract than inferred from it relative high absorption rate (Clapperton and Czerkawski, 1972).
There was no effect of PG infusion on the net portal fluxes of propionate, lactate, glucose, 3-hydroxybutyrate, glutamate, and glutamine or on the net hepatic fluxes of propionate, glucose, 3-hydroxybutyrate, glutamate, and glutamine (P>0.10; data not shown). However, the net hepatic flux of lactate changed (P<0.01) from minus 19 to plus 21±11 mmol/h. The increased lactate balance across the liver accounted for 56±12% of the PG uptake in the liver. This could indicate that only the (S)-1,2-propanediol is metabolized into L-(S)-lactate.

CONCLUSIONS

Up to 95% of intravenously infused propylene glycol (PG) was taken up by the splanchnic tissues. The hepatic extraction ratio was relatively low (8%) for PG, but the liver was the quantitatively most important organ for PG metabolism and released L-lactate to peripheral tissues, equivalent to half of the PG uptake.

REFERENCES