Bioavailability of starch and postprandial changes in splanchnic glucose metabolism in pigs

LIONEL NOAH, MICHEL KREMPF, GÉRARD LECANNU, PASCALE MAUGÈRE, and MARTINE CHAMP

Institut National de la Recherche Agronomique, 44316 Nantes; and Centre de Recherches en Nutrition Humaine, Groupe Météabolisme, 44035 Nantes, France

Noah Lionel, Michel Krempf, Gérard Lecannu, Pascale Maugère, and Martine Champ. Bioavailability of starch and postprandial changes in splanchnic glucose metabolism in pigs. Am. J. Physiol. Endocrinol. Metab. 278: E181–E188, 2000.—Changes in splanchnic metabolism in pigs were assessed after meals containing slowly or rapidly digested starch. The pigs were fed a mixed meal containing a “slow” native (n = 5) or a “rapid” pregelatinized (n = 5) cornstarch naturally enriched with [13C]glucose. Absorption of [13C]glucose was monitored by the arteriovenous difference technique, and infusion of D-[6,6-2H2]glucose in the jugular vein was used to calculate the systemic appearance of [13C]glucose. Arteriovenous balance data obtained during a 12-h study period showed that the fraction of ingested glucose equivalent appearing as glucose in the portal vein was 49.7 ± 7.2% for the slow starch and 48.2 ± 7.5% for the rapid starch (P = 0.86). These values, corrected for the gut extraction of circulating [13C]glucose, became 66.4 ± 5.6 and 65.3 ± 5.6%, respectively (P = 0.35). Isotope dilution data indicated that systemic appearance of exogenous [13C]glucose represented 62.9 ± 7.6 and 67.4 ± 3.0% of the oral load for slow and rapid starch, respectively (P = 0.68). Arterial glucose utilization by the gut increased from 7.3 ± 0.9 µmol·kg⁻¹·min⁻¹ before the meal to 8.5 ± 1.6 µmol·kg⁻¹·min⁻¹ during absorption, independently of the nature of the starch. Thus splanchnic glucose metabolism was unaffected by the nature of starch ingested.

The rate of absorption is probably a key factor in splanchnic metabolism, because it regulates insulin secretion or action (20, 27). Although long-term consumption of slowly digested starches can prevent the development of insulin resistance (12, 20, 27) and obesity (15, 27) in rats, the postprandial effects of starch ingestion on splanchnic glucose metabolism are not clearly known. This is especially true when starch is ingested with other food products, which can affect its bioavailability. Most studies in animal models have been carried out in pigs. After the meal, which is rarely used to indicate this fact.

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first 12 h after a meal was, respectively, 0.65 and 0.73% for pregelatinized starch and 0.46 and 0.30% for native starch. Only 2.8% (pregelatinized starch) and 0.8% (native starch) of the oral load still remained within the gut lumen after the pigs were killed at 12 h. The amounts of available glucose equivalent supplied by the two products were 207.7 g (native starch) and 193.0 g (pregelatinized starch).

Although both starches were highly digestible, the susceptibility of starch to hydrolysis was greater for pregelatinized than for native starch, as measured by the method of Bornet et al. (3). After 30 and 180 min of incubation with α-amylase, the proportion of starch converted to alcohol-soluble maltoligosaccharides was, respectively, 65 ± 1 and 71 ± 3% for pregelatinized starch (n = 2) and only 8 ± 1 and 22 ± 1% for native starch (n = 2).

Experimental meals. Two experimental meals providing 5,720 kJ (31% from lipids, 16% from proteins, and 53% from carbohydrates) were given to the pigs (Table 1). The mixed meals contained either "slow" native or "rapid" pregelatinized starch.

Experimental Design

Each pig received the two experimental meals in randomized order, with an interval of 1 wk between experiments. Two pigs got only one of the two experimental meals. A total of 10 experiments was performed for the entire protocol (5 for the slow starch meal and 5 for the rapid starch meal). One week before the study, the pigs were fed a meal without naturally occurring foods (Table 1). At 7 AM on the day of the experiment (t = −120 min), a bolus of tracer (22 mmol/kg body wt; δ[6,6-2H2]glucose, 99 mol% enrichment, CIL, Andover, MA) was injected into the jugular vein, and constant infusion was begun at a rate of 20 µmol·kg⁻¹·h⁻¹. This sterile, pyrogen-free saline solution was infused through an 0.2-mm Millipore filter. After 120 min (time 0), the pigs were fed the experimental meal. Portal vein blood flow was continuously recorded, and blood samples were taken from the portal vein and the carotid artery from −30 min before the meal until the end of the study (t = 720 min).

Blood samples (5 ml) were taken every 15 min during the first 4 h (t = −120 min to t = 120 min), every 30 min during the next 3 h (t = 120 min to t = 300 min), and then every hour until the end of the study (t = 300 min to t = 720 min). Blood was collected in heparinized tubes (Terumo, Leuven, Belgium) and centrifuged (9,000 g for 10 min at 4°C). Plasma was collected in heparinized tubes (Terumo, Leuven, Belgium) and centrifuged (9,000 g for 10 min at 4°C). Plasma was isolated and kept at −20°C until analyzed.

Table 1. Composition of experimental and preexperimental meals

<table>
<thead>
<tr>
<th>Preexperimental Meal</th>
<th>Experimental Meals</th>
<th>Diet N</th>
<th>Diet P</th>
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<tbody>
<tr>
<td></td>
<td>g/kg dry matter</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Native corn starch*</td>
<td>573</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pregelatinized corn starch*</td>
<td>563</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pregelatinized potato starch*</td>
<td>393</td>
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<td></td>
</tr>
<tr>
<td>Native wheat starch*</td>
<td>119</td>
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<tr>
<td>Meat meal</td>
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<td>307</td>
<td>314</td>
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<tr>
<td>Soybean oil</td>
<td>118</td>
<td>115</td>
<td>118</td>
</tr>
<tr>
<td>Wheat bran</td>
<td>81</td>
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</tr>
<tr>
<td>Pea hulls</td>
<td>5</td>
<td>5</td>
<td>5</td>
</tr>
</tbody>
</table>

| Diet N is a native starch meal, and diet P is a pregelatinized starch meal. Preexperimental meal consisted of 430 g. Experimental diet (326 g for diet N, and 318 g for diet P) provided −5,720 kJ. Meals were mixed with water at a ratio of 1 to 3 (wt/vol). * Starches were supplied by Roquette Industries, Lestrem, France.

Analytical Procedures

13C enrichment of starch. The 13C enrichment of glucose from starch was determined by the gas chromatography–combustion–isotope ratio mass spectrometry (GC-C-IRMS) method described by Tissot et al. (29). Native and pregelatinized starches were first hydrolyzed into glucose by use of a method modified by Thivend et al. (28). Five hundred milligrams of starch, mixed with 25 ml of distilled water, were heated at 100°C for 30 min. Then 2.5 ml of acetate buffer (pH = 4.6) were added, and the solution was completed with 45 ml of distilled water. Ten milligrams of amyloglucosidase (thermostable, Merck-Clévenot, Nogent-sur-Marne, France, cat. no. 1.01332, 2217 nKat/mg glucose equivalent) were added. Hydrolysis was stopped after 2 h of incubation by heating the solution at 100°C for 10 min. The hydrolysis products were then extracted by adding 188 ml of pure ethanol and mixing and centrifuging at 1,500 g (10 min). Supernatant was isolated and evaporated under vacuum. Glucose was then derivatized to glucose pentaacetate (21). The derivatized product was diluted in 1 ml of ethyl acetate (Sigma-Aldrich Chimie), and a sample (1 µl) was injected into a gas chromatograph (capillary column 30 m × 0.32 mm × 0.25 µm, type DB1, J W Scientific, Courtaboeuf, France). The gas chromatograph was coupled to a combustion furnace (CuO, NiO, Pt, 940°C), and an isotope ratio mass spectrometer (Delta S, Finnigan Mat, Bremen, Germany). Isotopometric ions of CO₂ were separated and recorded according to their mass-to-charge ratio (m/z) (44, 45, or 46). A standard CO₂ sample of known enrichment was injected intermittently to ensure the high precision of measurements.

Plasma variables. Plasma glucose was measured by the glucose-oxidase method (Glucose Analyzer II, Beckman, Fullerton, CA). Insulin was measured by radioimmunoassay (ORIS, Gif sur Yvette, France). Plasma lactate was analyzed using an enzymatic test kit (Boehringer Mannheim, Meylan, France, cat. no. 139084).

Plasma [2H₂]glucose and [13C]glucose isotopic enrichment. Fifty microliters of plasma were deproteinized by addition of 300 µl acetone. After 10 min at 4°C, the tube was centrifuged for 10 min at 4,000 g, and the supernatant was isolated. The supernatant was then evaporated under nitrogen, and the sample was derivatized.

13C plasma glucose enrichment ([13C%]) was analyzed by the GC-C-IRMS method we have described.

Plasma δ[6,6-2H₂]glucose was analyzed by GC coupled to MS (Hewlett-Packard MSD 5971A, Les Ulis, France) by use of a previously described method (14).

Calculations

Net [13C]glucose and lactate balance across the gut. Net gut balance (NGB) was calculated according to the formula described by Rérat et al. (24).

\[
NGB = \sum (C_{pv} - C_{ca}) \cdot PVF \cdot dt
\]

where \(C_{pv}\) and \(C_{ca}\) are the [13C]glucose (exogenous glucose) or lactate concentrations in portal vein and carotid artery blood, respectively, and PVF is the portal vein blood flow during the studied time interval (dt). Plasma concentrations were converted into whole blood concentrations using the \((1 - 0.3 \times \text{hematocrit})\) correction factor proposed by Dill (6).

The fraction of vascular glucose extracted by the gut (%) was calculated as follows:

\[
\text{gut fractional extraction} = \left(\frac{[D_{ca} - D_{pv}]}{D_{ca}}\right)\cdot 100
\]
where $D_{pa}$ and $D_{pw}$ are the $\nu$-[6,6-$^3$H$_2$]glucose concentrations in carotid artery and portal vein plasma, respectively.

The extraction ratio of vascular glucose was then used to correct the net $[^{13}$C]glucose balance across the gut. To take into account the gut extraction of recirculating glucose, arterial blood flow of $[^{13}$C]glucose was multiplied by the gut extraction ratio. The obtained value was added to the net portal $[^{13}$C]glucose flux.

Rate of glucose appearance in the systemic pool. The isotopic abundance of the samples was expressed in $^{13}$C atom % (AP). The AP of ingested and plasma glucose was transformed into atom % excess (APE) by the following formula:

$$\text{APE} = \text{AP} - \text{AP}_b$$

where $\text{AP}$ is the AP of the plasma sample, and $\text{AP}_b$ is the AP of plasma glucose before the meal.

The rate of glucose appearance (RaT, mol·kg$^{-1}$·min$^{-1}$) was calculated from the $^3$H enrichment of plasma glucose according to the equation of Steele (26), as modified by De Bodo et al. (5).

$$\text{RaT} = \frac{F - V \cdot p \cdot [(G_1 + G_2)/2] \cdot \{[\text{MPE}_2 - \text{MPE}_1]/(t_2 - t_1)\}}{\text{MPE}_2 + \text{MPE}_1}/2$$

where $F$ is the infusion rate of tracer $\nu$-[6,6-$^3$H$_2$]glucose (mol·kg$^{-1}$·min$^{-1}$), $\text{MPE}_1$ and $\text{MPE}_2$ are the mol % excess values of $[^{3}$H$_2$]glucose of plasma glucose, respectively, at time $t_1$ and $t_2$ (min), and $G_1$ and $G_2$ (mol) are the plasma glucose concentrations, respectively, at times $t_1$ and $t_2$. The distribution volume ($V$) was assumed to be 0.2 l/kg, and a pool fraction value ($p$) of 0.65 was chosen for the study (19).

The rate of appearance of exogenous glucose in the systemic pool (RaE, mol·kg$^{-1}$·min$^{-1}$) was calculated from the enrichment of derivatized plasma $[^{13}$C]glucose, according to the formula suggested by Proietto et al. (22).

$$\text{RaE} = \frac{[\text{RaT} \cdot (\text{APE}_1 + \text{APE}_2)/2] + [V \cdot p \cdot (G_1 + G_2)/2 \cdot (\text{APE}_2 - \text{APE}_1)/(t_2 - t_1)]}{\text{APE}_{glu.ing}}$$

where $\text{APE}_1$ and $\text{APE}_2$ are the $^{13}$C enrichments of plasma glucose, respectively, at time $t_1$ and $t_2$ (min), and $\text{APE}_{glu.ing}$ is the AP excess of ingested glucose.

Endogenous glucose production (Rae) was calculated by subtracting RaE from RaT. The suppression of Rae caused by gut extraction was calculated from the formula suggested by Proietto et al. (22)

$$[\text{RaT} \cdot (\text{APE}_1 + \text{APE}_2)/2] + [V \cdot p \cdot (G_1 + G_2)/2 \cdot (\text{APE}_2 - \text{APE}_1)/(t_2 - t_1)]$$

where $\text{APE}_1$ and $\text{APE}_2$ are the $^{13}$C enrichments of plasma glucose, respectively, at time $t_1$ and $t_2$ (min), and $\text{APE}_{glu.ing}$ is the AP excess of ingested glucose.

Endogenous glucose production (Rae) was calculated by subtracting RaE from RaT. The suppression of Rae caused by gut extraction was calculated from the formula suggested by Proietto et al. (22)

$$\text{Rae} = \frac{(\text{basal production} - \text{postprandial endogenous production})}{\text{basal production}} \times 100$$

where basal production is the mean Rae before the meal.

**Statistical Analysis**

Statistical analyses involved standard error of the mean (SE), the Mann-Whitney test to compare the mean enrichments of native and pregelatinized cornstarch, and two-way analysis of variance to determine the effects of animals and test meals on metabolic variables. Analyses were performed using the Statgraphics 3.0 software package (STSC, Rockville, MD).

**RESULTS**

$^{13}$C Enrichment of Starch

After the starches were enzymatically hydrolyzed, no difference was observed between the $^{13}$C enrichments of derivatized glucose from slow or rapid starch ($P =$ 0.87). The mean $^{13}$C enrichment of derivatized glucose was $1.0753 \pm 0.0001$ (n = 12) atom % $^{13}$C (AP).

Absorption and Gut Metabolism (Arteriovenous Difference Technique)

Portal blood flow. Basal portal blood flows were $21.25 \pm 1.05$ ml·kg$^{-1}$·min$^{-1}$ (n = 5) and $21.07 \pm 0.65$ ml·kg$^{-1}$·min$^{-1}$ (n = 5) for slow starch and rapid starch meals ($P =$ 0.36). Portal flows increased to a mean of $31.43 \pm 2.01$ ml·kg$^{-1}$·min$^{-1}$ at 45 min (slow starch) and $30.57 \pm 2.18$ ml·kg$^{-1}$·min$^{-1}$ at 90 min (rapid starch) during the postprandial period, and they returned to the basal level after 420 min. There was no difference between average portal flows measured after meals ($25.21 \pm 1.59$ and $25.50 \pm 1.17$ ml·kg$^{-1}$·min$^{-1}$) for slow and rapid starch, respectively ($P =$ 0.55).

Glucose. Basal arterial plasma glucose concentration was not different between the slow starch meals ($4.71 \pm 0.14$ mmol/l; n = 5) and the rapid starch meals ($4.55 \pm 0.19$ mmol/l; n = 5) ($P =$ 0.30). Mean gut fractional extraction of glucose during the study was $7.6 \pm 1.4$ and $7.7 \pm 2.5\%$, respectively, for slow starch and rapid starch meals ($P =$ 0.16). The fraction of arterial glucose extracted by the gut decreased after the meal and was minimal at 30 min for slow starch and at 90 min for rapid starch (Fig. 1). In basal state, the mean rate of arterial glucose utilization by the gut was $6.6 \pm 1.3$ µmol·kg$^{-1}$·min$^{-1}$ for slow starch and $7.5 \pm 1.4$ µmol·kg$^{-1}$·min$^{-1}$ for rapid starch ($P =$ 0.91) (mean for both starches was $7.3 \pm 0.8$ µmol·kg$^{-1}$·min$^{-1}$). Under the meal fed condition, this rate was $8.8 \pm 1.3$ and $8.3 \pm 3.2$ µmol·kg$^{-1}$·min$^{-1}$, respectively, for slow and rapid starch ($P =$ 0.20) (mean for both starches was $8.5 \pm 1.6$ µmol·kg$^{-1}$·min$^{-1}$). During absorption, the total amount of arterial glucose extracted was $57.4 \pm 11.3$ g for slow starch and $51.3 \pm 20.0$ g for rapid starch ($P =$ 0.20).
plateau at ~75 min both for slow starch (≈0.0135 APE) and for rapid starch (≈0.0150 APE) (P = 0.23) (Fig. 2). For both meals, glucose $^{13}$C enrichment remained relatively high at the end of the study (≈0.0100 APE). At that time, enrichments were not significantly different between portal vein and carotid artery, either for slow starch (P = 0.06) or for rapid starch (P = 0.11).

As shown in Fig. 3, net absorption of $^{13}$C glucose lasted until 720 min after the meals. The cumulated amount of “absorbed” $^{13}$C glucose tended to be lower (P = 0.05) for the slow starch than for the rapid starch meal during the first 90 min (respectively, 30.9 ± 4.6 and 37.8 ± 3.5 g) and 105 min (respectively, 36.1 ± 5.3 and 44.3 ± 4.7 g). The amounts of $^{13}$C glucose appearing in the portal vein during the 720 min of the experiment were 103.2 ± 15.0 g (49.7 ± 7.2% of the ingested amount) and 95.5 ± 14.9 g (48.2 ± 7.5% of the ingested amount) for the slow starch meal (n = 5) and the rapid starch meal (n = 5), respectively (difference not statistically significant, P = 0.86). When net balance of $^{13}$C glucose across the gut was corrected for the extraction of circulating $^{13}$C glucose, the rates of $^{13}$C glucose absorption were still found to be 6.5 ± 4.6 and 6.0 ± 1.9 µmol·kg$^{-1}$·min$^{-1}$ for the slow starch meal (n = 5) and the rapid starch meal (n = 5), respectively, at the end of the study (Fig. 3). Corrected for the extraction of circulating $^{13}$C glucose, the total amounts of absorbed $^{13}$C glucose during the 720 min of the experiment were 137.8 ± 11.6 g for the slow starch (66.4 ± 5.6% of the ingested amount) and 129.3 ± 11.0 g for the rapid starch (65.3 ± 5.6% of the ingested amount) (P = 0.35).

Insulin. Arterial plasma insulin concentration was 34.3 ± 7.3 pmol/l (n = 5) and 37.1 ± 13.8 pmol/l (n = 5) before the slow starch and rapid starch meals (P = 0.99). Maximal increment over basal state, obtained 30...
min after the beginning of the meal (Fig. 4), was 216.4 ± 46.8 and 247.5 ± 41.7 pmol/l, respectively (P = 0.77). No difference was observed between the two meals for the area under the postprandial curve of insulin (P = 0.88).

Lactate. Mean arterial lactate concentration was 0.62 ± 0.08 mmol/l (n = 5) and 0.65 ± 0.09 mmol/l (n = 5), respectively, before the slow starch and rapid starch meals (P = 0.83). Net production of lactate by the gut occurred during the absorption period, and the rate of production at maximum level (30 min) was not significantly different (P = 0.10) between the two meals (13.0 ± 1.3 µmol·kg⁻¹·min⁻¹ for slow starch and 13.0 ± 1.2 µmol·kg⁻¹·min⁻¹ for rapid starch; n = 5). The total production of lactate by the gut amounted to 7.3 ± 1.4 g (3.5 ± 0.7% of the glucose equivalent ingested; n = 5) and 6.7 ± 0.7 g (3.3 ± 0.3% of the glucose equivalent ingested; n = 5) for slow starch and rapid starch meals, respectively (difference not statistically significant, P = 0.35).

Isotope Dilution Data

Rate of systemic appearance of glucose. The rate of total glucose appearance (RaT) was 17.23 ± 1.18 and 13.92 ± 1.11 µmol·kg⁻¹·min⁻¹, respectively, before the slow starch meal (n = 5) and the rapid starch meal (n = 5) (P = 0.10). Glucose production increased to maximal values at 90 min for slow starch (70.25 ± 9.08 µmol·kg⁻¹·min⁻¹) and at 75 min for rapid starch (61.35 ± 3.49 µmol·kg⁻¹·min⁻¹) (P = 0.45).

Systemic appearance of endogenous and exogenous glucose. During absorption, endogenous glucose production was suppressed an average of 59.9 ± 8.3% (n = 5) after the slow starch meal and 69.7 ± 5.7% (n = 5) after the rapid starch meal (Table 2), corresponding to equivalent amounts of glucose retained in the splanchnic bed of 59.0 ± 10.2 g and 54.8 ± 6.5 g, respectively (P = 0.55) (Fig. 5).

The rate of systemic appearance of exogenous [¹³C]glucose (RaE) increased to maximal values of 58.28 ± 9.38 µmol·kg⁻¹·min⁻¹ at 90 min and 61.63 ± 7.43 µmol·kg⁻¹·min⁻¹ at 75 min, respectively, after slow starch and rapid starch meals (P = 0.80) (Fig. 5). As calculated by the area under the curve, the cumulated amount of

<table>
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<tr>
<th>Ingested [¹³C]glucose equivalent, g</th>
<th>Native Starch</th>
<th>Pregelatinized Starch</th>
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<tr>
<td>207.7</td>
<td>198.0</td>
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<th>Systemic appearance of exogenous [¹³C]glucose, g</th>
<th>Native Starch</th>
<th>Pregelatinized Starch</th>
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<tr>
<td>130.7 ± 15.8</td>
<td>133.4 ± 5.9</td>
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<tr>
<th>Exogenous [¹³C]glucose retained in the splanchnic bed, g</th>
<th>Native Starch</th>
<th>Pregelatinized Starch</th>
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<tr>
<td>77.0 ± 15.8</td>
<td>64.6 ± 5.9</td>
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<th>Suppression of basal hepatic glucose production, %</th>
<th>Native Starch</th>
<th>Pregelatinized Starch</th>
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<tbody>
<tr>
<td>59.9 ± 8.3</td>
<td>69.7 ± 5.7</td>
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</table>

Values are means ± SE (n = 5). Native (“slow”) and pregelatinized (“rapid”) cornstarches were supplied by Roquette Industries, Les-trem, France.

Fig. 4. Incremental insulin concentration in arterial plasma after slow starch meal (●) and rapid starch meal (○). Values are means ± SE; n = 5. Time 0 is beginning of experimental meal.

Fig. 5. Rates of systemic appearance of endogenous (●, ○ in A) and exogenous (▲, △ in B) glucose for native starch meal (A) and pregelatinized starch meal (B). Values are means ± SE; n = 5. Time 0 is beginning of experimental meal.

Table 2. Splanchnic glucose metabolism during absorption
exogenous glucose appearing at the peripheral level was 130.7 ± 15.8 g (n = 5) for slow starch and 133.4 ± 5.9 g (n = 5) for rapid starch during the 720 min of the experiment. Thus, respectively, 77.0 ± 15.8 g (37.1 ± 7.6% of the ingested load) and 64.6 ± 5.9 g (32.6 ± 3.0% of the ingested load) were retained in the splanchnic area (P = 0.68) (Table 2). Actually, in the present study, splanchnic glucose retention was found to be equivalent to gut glucose retention.

Comparison of arteriovenous difference and isotope dilution data. The cumulated amounts of [13C]glucose appearing in the portal vein (calculated using the net balance across the gut, or the net balance corrected for the extraction of circulating [13C]glucose) and the cumulated amounts of [13C]glucose appearing in systemic blood (calculated using the isotope dilution method) were determined for slow and rapid starches (Fig. 6). With either test meal, the values calculated by the different methods were similar during the first 240 min of the study. Then, values obtained by the arteriovenous difference method, uncorrected for the extraction of circulating [13C]glucose, became lower than the values calculated by the isotope dilution method. For both meals, the cumulated amounts of glucose determined by the isotope dilution technique at 720 min were ≈ 25% higher than those obtained with the arteriovenous difference method. Values were significantly different between the two methods at 600 min, 660 min, and 720 min only for the rapid starch meal (P < 0.05). When corrected for the extraction of circulating [13C]glucose, the values calculated by the arteriovenous difference method were similar to those determined by isotope dilution.

DISCUSSION

Splanchnic metabolism of glucose from a slowly or rapidly digested starch was considered in this study. The pig, because of its physiological similarities with humans, was used as the animal model (18). Splanchnic glucose metabolism was assessed simultaneously with the arteriovenous difference technique (to measure net glucose absorption) and the isotope dilution (dual tracer) technique, after the pigs had received isocaloric (5,720 kJ) meals containing ≈ 53% of their energy intake as starch.

A low average blood flow rate was found after the test meals (≈ 25.5 ml·kg⁻¹·min⁻¹) compared with the range of values reported for pigs (24–40 ml·kg⁻¹·min⁻¹) (2, 8, 24). However, because the transit time ultrasound method has been validated in vivo (7), measurements may depend on the amount and type of diet rather than on the technique used. Moreover, no development of collateral circulation in the portal system was observed when pigs were autopsied at the end of each experiment.

In basal state, the rate of glucose utilization by the gut was ≈ 7.3 µmol·kg⁻¹·min⁻¹. During absorption, the gut switched from net glucose utilization to net glucose release. However, it is remarkable that glucose utilization by the gut increased substantially in the meal fed condition. The gut used both luminal glucose from starch digestion, and vascular glucose, whose rate of utilization increased during absorption. The extraction ratio of vascular glucose decreased between 30 and 90 min after the meals, but arterial glucose was still partially extracted by the gut during absorption, mainly in the latest phase of this absorption (after the first 240 min of the study). For both starches, the rate of utilization increased during absorption. Abumrad et al. (1) showed that glucose utilization by the gut is increased during absorption. Although Vaugelade et al. (32) observed a decreased glucose oxidation capacity of enterocytes in postprandial conditions (in the presence of glutamine), they reported a high glycolytic capacity for the intestinal muscular layer, which may have contributed to the metabolism of both endogenous glucose and [13C]glucose (from starch digestion) in our study.
Absorption of $[^{13}C]$glucose tended to be higher during the first 105 min after the rapid starch meal. The pig has high digestive capacities (18), which may explain why the difference in the rates of glucose absorption for the two starches was not as large as expected from in vitro results. The proportion of ingested $[^{13}C]$glucose equivalent that appeared in the portal vein was low ($\approx 50\%$) for both test meals. However, when the values were corrected for recycled $[^{13}C]$glucose uptake by the intestine, the proportion of ingested $[^{13}C]$glucose equivalent entering the portal vein ($\approx 65\%$ for both meals) was more in accordance with the 70–80% generally measured after a pure glucose load (1, 2). On the basis of these last data, the gut would have retained $\approx 35\%$ of the dietary glucose load ($\approx 70$ g) during the postprandial study period. However, this proportion is overestimated, because absorption was not fully ended 12 h after the meals. Indeed, the rates of $[^{13}C]$glucose portal flux were still $6.5 \pm 4.6 \mu\text{mol} \cdot \text{kg}^{-1} \cdot \text{min}^{-1}$ (slow starch) and $6.0 \pm 1.9 \mu\text{mol} \cdot \text{kg}^{-1} \cdot \text{min}^{-1}$ (rapid starch) at the end of our experimental period.

Part of the ingested starch was metabolized by the gut into lactate. After both test meals, lactate production by the gut ($\approx 7$ g) accounted for 3.5% of the ingested glucose equivalent, which is in agreement with values found in different species (1, 24, 30). Only minor gut production of alanine (3–5% of the glucose equivalent ingested) has been reported after an oral glucose load (1) or a mixed meal (17). Therefore, the fraction of the ingested glucose equivalent lacking in the balance ($\approx 30\%$) must be related either to the conversion of glucose into other metabolites (such as CO$_2$, for example) or to the uncompleted absorption at the end of the study. It is unlikely that part of the starch was fermented in the gut, because short-chain fatty acid concentration in ileal residues did not increase during the experiment (studies in one pig; results not shown) and little carbohydrate entered the colon in either animal (see Results).

Basal glucose production ($14–17 \mu\text{mol} \cdot \text{kg}^{-1} \cdot \text{min}^{-1}$) was similar to that reported for pigs (10, 19) and slightly higher than the values of $8–13 \mu\text{mol} \cdot \text{kg}^{-1} \cdot \text{min}^{-1}$ observed for humans (9, 13, 16, 23, 29). In the present article, the systemic appearance of glucose was calculated according to the single-compartment model described by Steele (26), which has mainly been used for glucose metabolism studies (22, 25, 29). This model does not really describe the complex physiology of glucose metabolism but is reliable at slow turnover rates (4). Moreover, although an error due to the use of too low a glucose distribution volume ($V = 0.2 \text{l/kg}$) has been found, it was shown to be partly counterbalanced by glucose recycling (16). The systemic appearance of exogenous glucose was not different for the two experimental meals. The similar cumulated amounts of $[^{13}C]$glucose calculated by the isotope dilution method were higher ($\approx 25\%$ higher at 720 min) than those calculated by the arteriovenous difference technique. Glucose recycling, via gluconeogenesis (11), might account for part of the discrepancy observed between stable isotope and arteriovenous difference data. However, it has been shown after a pure glucose load that the rate of glucose recycling is low during 240–360 min after the oral load (13, 16). It was also suggested that exogenous glucose (newly absorbed) may be taken up first by the liver and incorporated into the glycogen pool before being released into the systemic circulation at the end of the study period. Indeed, Moore et al. (17) reported net hepatic release of absorbed $[^{13}C]$glucose tracer in dogs 300 min after the ingestion of a mixed meal. Our findings do not support this hypothesis, because at no period of the study were the arterial glucose $[^{13}C]$enrichments higher than the portal glucose $[^{13}C]$enrichments. Moreover, when the cumulated amounts of $[^{13}C]$glucose obtained by arteriovenous difference were corrected for recycled $[^{13}C]$glucose uptake by the intestine, they were similar to those calculated for isotope dilution. This suggests that the difference between the two methods might only be related to the extraction of recirculating exogenous glucose by the gut. Our findings are consistent with an appropriate use of the isotope dilution method as an alternative approach to measure the rate of $[^{13}C]$glucose absorption.

In conclusion, the concomitant use of arteriovenous balance across the gut and isotope tracers showed that glucose utilization by the gut increased substantially during absorption and that this was unaffected by the nature of the starch ingested.

Address for correspondence and reprint requests: M. Champ, INRA, rue de la Géraudière, BP 71627, 44316 Nantes, Cedex 03, France (E-mail: champ@nantes.inra.fr).

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