Evidence for an Effect of Insulin on the Peripheral Utilization of Ketone Bodies in Dogs

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**Abstract** The rates of transport and oxidation of acetoacetate have been measured in seven anesthetized, pancreatectomized, ketotic dogs using a constant infusion of acetoacetate-3-14C. Control experiments were performed in 14 normal dogs. In addition to the acetoacetate-14C, the latter were infused at a constant rate with varying amounts of unlabeled acetoacetate so as to obtain a range of ketone transport (26-65 μmole/min·kg) comparable with that observed in the diabetic dogs (21-41 μmole/min·kg). The specific activities of acetoacetate and β-hydroxybutyrate in blood became equal during the infusion of labeled acetoacetate, indicating that the net transport of acetoacetate represents that of total ketones. In each group, the concentration of ketones was an exponential function of the rate of transport, but for any value below 30 μmoles/min·kg, ketone concentration in the diabetic dogs was about 3 times that in normal dogs, indicating an impairment of mechanisms for utilizing ketones in insulin deficient animals. Maximal capacity to utilize ketones in diabetic dogs was slightly more than half that of normal ones. A similar fraction (32-63%) of the infused 14C appeared in respiratory CO2 in the two groups and was independent of the rate of transport. In seven of the normal dogs, administration of insulin and glucose increased removal of the infused ketones and increased the fraction of 14C appearing in respiratory CO2. These results demonstrate that utilization of ketones in extrahepatic tissues is influenced by insulin; impaired utilization contributes to diabetic ketosis and is probably essential to the production of severe ketoacidosis.

**Introduction**

It is well established that ketone bodies are manufactured primarily in the liver and transported through blood to peripheral tissues where they are readily oxidized. Under certain conditions, ketones produced in liver may account for a considerable fraction of caloric requirements (2, 3). The regulation of production of ketones by the liver has been extensively studied both in vivo and in vitro (4-7). In the intact organism, high rates of ketogenesis are observed essentially in situations characterized by high free fatty acid (FFA) levels associated with carbohydrate deprivation as in starvation or diabetes. Little is known about factors controlling the peripheral utilization of ketone bodies. The only well-established data are that ketone utilization is accelerated by increased concentrations of ketones in blood (8, 9) and by increase in the metabolic rate such as occurs in muscular exercise (10). A possible role of insulin in the control of ketone utilization has been investigated repeatedly during the last 40 yr. Conflicting reports have appeared. Earlier work in several mammalian species indicated that ketone body utilization is not impaired in diabetes (11-16), whereas more recent reports indicate that some impairment may exist in diabetic rats and that it can be corrected by insulin (17-20).

Recent work from this laboratory has shown that hyperketonemia in dogs with diabetic ketosis exceeds that observed in norepinephrine-infused, nondiabetic animals with comparable rates of hepatic ketogenesis (21). This observation has led us to reevaluate the problem of ketone utilization in this species with isotopic techniques which permit measurement of rates of transport and oxidation of ketones. For this purpose, diabetic dogs have been compared with normal animals made ketotic by infusing acetoacetate (AcAc).

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METHODS

Experimental animals and procedures

Two groups of male mongrel dogs weighing 13-28 kg were studied: (a) normal dogs maintained on a diet containing approximately 75% protein and 25% fat and fasted for 18 hr before (b) pancreactomized dogs fasted for 18 hr and deprived of insulin for 48-72 hr to allow the development of variable degrees of ketosis. The animals were depancreatized at least 1 month earlier and maintained on the same diet as that of normal animals but supplemented with crude pancreatin and given NPH insulin subcutaneously once daily to maintain urinary excretion of glucose below 15 g daily. Animals from both groups were in excellent nutritional condition.

On the day of the experiment, anesthesia was induced with sodium pentobarbital (25-30 mg/kg) and maintained by additional smaller doses to prevent shivering or other muscular activity. The animals were intubated and ventilated artificially with air at a constant rate (0.17-0.29 liter/min/kg). Catheters were introduced into the vena cava through a superficial leg vein for infusion, into a femoral artery through a side branch for blood sampling and into the bladder for urine collection. The diabetic animals were given a constant infusion of trace amounts of AcAc-3-14C diluted in saline (0.01-0.02 μCi/min/kg). The normal animals were infused with a mixture of AcAc-3,4-14C (0.001-0.02 μCi/min/kg) and unlabeled AcAc (26.3-77.5 μmoles/min/kg). In both diabetic and normal dogs, infusions were continued for 3-6 hr and were preceded by injection of a priming dose equivalent to 15 times the infusion rate per min. Samples of arterial blood, expired air, and urine were collected at intervals during the course of the experiment.

In a few experiments on normal dogs, blood was also sampled from the femoral vein and the portal vein through appropriate catheters (the portal catheters were implanted through a branch of the splenic vein at laparotomy about 6 days before the experiment [21]). At the end of two experiments on normal dogs infused with AcAc, samples of adipose tissue, skeletal muscle, liver, and kidney were taken for analysis. In one of these experiments (dog 58), the infused tracer was [1-14C]-β-hydroxybutyrate 3-14C rather than AcAc-3-14C. The influence of blood pH and of administration of insulin on the utilization of ketones were also studied. The experimental protocols are described together with the results.

Production rate of ketones by the liver was measured directly in seven normal, fasted dogs fitted with catheters in an artery, the portal vein, and an hepatic vein (21). Hepatic blood flow was measured using a constant infusion of 3H-labeled rose bengal into a peripheral vein (22). Calculations of production rates of ketones were made assuming that 80% of the hepatic blood flow was derived from the portal vein (21).

Analyses

Blood samples were collected with glass syringes and placed in tubes containing dry heparin. Samples of 5 ml blood were immediately mixed with 5 ml of chilled perchloric acid (30% w/v). After centrifugation at 4°C, the supernatant fluid was collected, neutralized with 20% KOH, and centrifuged again, and the resulting protein-free filtrate was kept on ice until further use. Duplicate 0.05 ml blood samples were deproteinized with zinc sulfate and barium hydroxide solutions for measurement of glucose (23). The rest of the blood was centrifuged, and the plasma was analyzed for content of FFA (24) and immunoreactive insulin (IRI) (25).

AcAc and β-hydroxybutyrate (βOHB) were estimated in the perchlorate filtrates using a fluorimetric adaptation of the enzymatic method of Williamson and Krebs (21). Co2 in ketone bodies was measured by the method of Mayes and Felts (26). This method permits the determination of 14C activity separately in AcAc and βOHB; βOHB-14C is converted enzymatically to AcAc-14C; AcAc-14C is decarboxylated to acetone and CO2 which are trapped separately in a double-well flask and assayed for 14C in a liquid scintillation spectrometer. Since the AcAc used in these experiments was labeled in the carbonyl carbon, only the labeled acetone produced was measured. The recovery of AcAc-14C and βOHB-14C [ν(ν- or ν(-)] added to nonradioactive blood was determined with each set of determinations and found to be 88-107% for AcAc and 74-100% of βOHB [when ν-βOHB was used to check recovery, only half of the 14C was expected to be recovered since the assay is specific for the ν(-) isomer]. Values for ketone radioactivity in blood samples were corrected accordingly. The determinations of unlabeled and labeled AcAc were done on the day of each experiment owing to the instability of the compound.

Blood samples for analysis of gas tensions were taken in heparinized syringes and analyzed promptly (27). Samples of expired air were collected in plastic balloons and analyzed for content of CO2 and 14CO2 by the method of Frederickson and Ono (28). In two diabetic dogs receiving an infusion of AcAc-14C, expired air was analyzed for acetone-14C by using the technique described by Frederickson and Ono for expired 14CO2 except that a mixture of lactic acid and hydrazide hydrate (5:2, v/v) was used to trap acetone. Recovery of labeled acetone by this procedure was about 85%. Results were not corrected for incomplete recovery. No detectable amounts of 14CO2 were trapped by the lactic acid-hydrazide mixture.

The urine samples were collected in graduated cylinders kept on ice, immediately neutralized with 10 N NaOH to avoid degradation of AcAc, and filtered. Samples of urine were assayed for 14C. The 14CO2 content of urine was estimated using the procedure described elsewhere for blood (21). The 14C in urine not accounted for by bicarbonate was assumed to be in ketone bodies.

Lipids were extracted from tissues in ethanol-acetone (1:1) in a blender and reextracted in chloroform-methanol (2:1). An equal volume of 0.005 N sulfuric acid was added to the latter extract. Lipids in the chloroform phase were separated on columns of silicic acid (29). Separated fractions containing phospholipids and neutral glycerides were saponified, and the liberated fatty acids and glycerol were separated after acidification (30). Glucose was isolated from the aqueous methanol phase as the gluconate derivative (31). The washed precipitate of proteins from the ethanol-acetone extract was dissolved in hot 20% KOH. Glycogen was precipitated from a KOH digest of fresh tissue (32) and hydrolyzed in 2.4 N HCl.

All samples were assayed for 14C in a liquid scintillation spectrometer. Methanol was added when necessary to make a single-phase system in toluene containing phosphors. Quenching of radioactivity was determined with an internal standard. All analyses were performed in duplicate.

Preparation of materials for injection

Unlabeled AcAc. Sodium AcAc was prepared by thoroughly mixing 54 ml of ethyl AcAc (Eastman Organic Chemicals, Rochester, N. Y.) with 200 ml of 2 N NaOH.
After 15 hr, the neutrality of the solution was checked with a pH paper. The solution (pH 7.0–7.5) was washed three times with equal amounts of diethyl ether to remove excess ethyl AcAc. The ether, the ethanol generated by the hydrolysis, and any acetone produced by spontaneous decarboxylation of the AcAc were removed by blowing a stream of nitrogen over the solution for at least 45 min.

**Labeled ketones.** Chromatographically pure ethyl AcAc-3-14C (specific activity 0.046 mCi/mg) (New England Nuclear Corp., Boston, Mass.) was mixed with 100 µl of ethyl acetocetate as carrier, hydrolyzed to AcAc-3-14C, and purified as indicated for the unlabeled material. n(-)3-OHB-3-14C was prepared from AcAc-3-14C with βOHB dehydrogenase and NADH and purified by liquid/liquid extraction in a Soxhlet apparatus and paper chromatography (26). Solutions of labeled and unlabeled AcAc were used immediately after preparation and chilled with ice during the course of the infusion.

**Calculations**

Concentration and specific activity of AcAc in blood were stable 1–4 hr after starting the constant infusion. The infusion was continued for at least 60 min after a steady state was reached, and 3–10 determinations were obtained during that period. Transport of AcAc to tissues was calculated by the following equation:

\[
(A) \text{Transport of AcAc (µmoles/min) = \frac{\text{infusion rate of AcAc-14C (cpm/min)}}{\text{specific activity of arterial AcAc (cpm/µmole)}}}
\]

For normal dogs infused with AcAc, stable concentrations of ketones in blood were obtained 1–4 hr after starting infusions. Ketone transport to tissues was calculated by the following equation:

\[
(B) \text{Transport of AcAc (µmoles/min) = \frac{\text{infusion rate of AcAc (µmoles/min)}}{\text{specific activity of arterial AcAc (cpm/µmole)}} - \text{ketones excreted in urine (µmoles/min)}}
\]

This equation assumes that endogenous production of total ketones in these animals is negligible with respect to the amount infused. This assumption is supported by the following data: (a) in seven normal fasted dogs, we found (Table VI) that the production rate of total ketones averaged 2.46 µmoles/min/kg (1.96–3.55), whereas the infusion rates of AcAc varied between 28.0 and 77.5 µmoles/min/kg, and (b) in five of the normal dogs, the specific activity of blood AcAc averaged 93 ±4% of the specific activity of infused AcAc. These data indicate that the values for AcAc transport calculated from (B) are underestimated by less than 10%. The amount of unlabeled ketones excreted in urine was calculated assuming that the specific activity of ketones in urine was equal to that of ketones in blood.

The efflux rate of 14CO2 and the specific activity of CO2 in expired air became virtually constant 2–4 hr after starting the infusion. Calculations of oxidation were made as follows:

\[
(C) \text{Percentage of AcAc-14C oxidized promptly to 14CO2 = \frac{100 \times \text{efflux rate of 14CO2 in expired air (cpm/min)}}{\text{influx rate of AcAc-14C (cpm/min)} - \text{14C in ketones excreted in urine (cpm/min)}}}
\]

\[
(D) \text{Apparent percentage of exhaled CO2 derived from rapid oxidation of AcAc = \frac{100 \times \text{specific activity of exhaled CO2 (cpm/µmole)}}{\text{specific activity of blood AcAc (cpm/µatom C)}}}
\]

In normal animals, for reasons outlined above, the specific activity of the infused AcAc (cpm/µatom C) was used instead of that of blood AcAc in the latter equation.

**RESULTS**

Major data obtained from an experiment on a moderately ketogenic diabetic dog are presented in Fig. 1. In this experiment, a steady state was present after about 180 min. Individual results from 7 experiments on diabetic dogs and 14 experiments on normal animals are summarized in Tables I–IV.

**Concentration of metabolites in blood.** The diabetic dogs (Table I) showed high FFA (1.56–2.72 µmoles/ml) and glucose levels (217–524 mg/dl) and elevated ketone concentrations (1.14–18.21 µmoles/ml), with βOHB-AcAc ratios varying between 0.78 and 2.56 (mean = 1.74). The most ketogenic dogs were also acidotic. Infusion of AcAc into the normal dogs (Table III) increased their blood ketones to levels (0.49–10.58 µmoles/ml) in the diabetic range with no systematic change in the βOHB-AcAc ratio which averaged 1.21 (0.14–2.80) before and 0.86 (0.41–1.18) during the infusion. The infusion of ketones also produced a rise in blood pH and a fall in blood glucose but no systematic changes in FFA or IRI concentrations (Fig. 2).

**Specific activity of blood ketones.** In three model experiments where measurements were made, the ratio of the specific activity of βOHB to that of AcAc was close to unity (0.90; 1.00; 1.18).

**Excretion of ketones in urine and expired air.** The values for transport of AcAc (Tables II and IV) have been corrected for urinary losses of ketones. As shown in Fig. 3, excretion was roughly proportional to the blood ketone concentration without apparent difference between normal and diabetic animals. In most cases, less

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1 Nomenclature used here is that recommended by Task group on Tracer Kinetics of International Commission on Radiation Units (33).
than 10% of the infused label was excreted in urine as ketones (Table II). In two diabetic dogs, acetone in expired air amounted to only 0.15 and 0.18% of transport of AcAc.

**TABLE I**

*Arterial Concentration of Metabolites in Pancreatectomised Dogs*

<table>
<thead>
<tr>
<th>Dog no.</th>
<th>FFA*</th>
<th>Glucose*</th>
<th>pH*</th>
<th>AcAc†</th>
<th>3-OHB‡</th>
<th>AcAc</th>
</tr>
</thead>
<tbody>
<tr>
<td>34</td>
<td>1.87</td>
<td>320</td>
<td>7.29</td>
<td>3.00 ±0.29</td>
<td>6.30 ±0.35</td>
<td>2.10 ±0.30</td>
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<td>35</td>
<td>1.29</td>
<td>305</td>
<td>7.23</td>
<td>5.11 ±0.20</td>
<td>13.10 ±0.14</td>
<td>2.56 ±0.08</td>
</tr>
<tr>
<td>46</td>
<td>1.58</td>
<td>401</td>
<td>7.41</td>
<td>0.64 ±0.04</td>
<td>0.59 ±0.08</td>
<td>0.78 ±0.12</td>
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<tr>
<td>43</td>
<td>2.10</td>
<td>524</td>
<td>7.33</td>
<td>0.83 ±0.06</td>
<td>1.59 ±0.17</td>
<td>1.92 ±0.22</td>
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<tr>
<td>40</td>
<td>2.72</td>
<td>490</td>
<td>7.28</td>
<td>2.08 ±0.18</td>
<td>3.58 ±0.60</td>
<td>1.72 ±0.23</td>
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<tr>
<td>50</td>
<td>1.85</td>
<td>324</td>
<td>7.38</td>
<td>0.98 ±0.09</td>
<td>1.72 ±0.10</td>
<td>1.75 ±0.19</td>
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<td>2.08</td>
<td>325</td>
<td>7.56</td>
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<tr>
<td>52§</td>
<td>1.56</td>
<td>217</td>
<td>7.55</td>
<td>1.25 ±0.11</td>
<td>1.92 ±0.11</td>
<td>1.54 ±0.09</td>
</tr>
</tbody>
</table>

* Mean of two to three values obtained during the period of steady blood ketone radioactivity.
† Mean ±SD of three to six values obtained during the last 60–220 min of AcAc-14C infusion (steady period of blood ketone radioactivity).
§ Dogs made alkaliotic by infusing sodium bicarbonate (see text).
TABLE II
Transport and Oxidation of Acetoacetate in Pancreatectomized Dogs

<table>
<thead>
<tr>
<th>Dog no.</th>
<th>% of infused ketones in urine</th>
<th>Specific activity of AcAc in blood*</th>
<th>Net transport of AcAc*</th>
<th>Efflux rate of CO2</th>
<th>Efflux rate of AcAc-14C Specific activity</th>
<th>% AcAc oxidised</th>
<th>% CO2 from AcAc</th>
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</thead>
<tbody>
<tr>
<td>34</td>
<td>14.5</td>
<td>56.8 ± 3.8</td>
<td>37.7</td>
<td>208</td>
<td>3000</td>
<td>14.4</td>
<td>35.1</td>
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<tr>
<td>35</td>
<td>29.0</td>
<td>46.7 ± 1.7</td>
<td>38.0</td>
<td>219</td>
<td>3870</td>
<td>17.7</td>
<td>54.2</td>
</tr>
<tr>
<td>46</td>
<td>4.0</td>
<td>110.6 ± 7.4</td>
<td>21.6</td>
<td>212</td>
<td>5370</td>
<td>25.3</td>
<td>56.0</td>
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<td>43</td>
<td>2.6</td>
<td>115.0 ± 10.8</td>
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<td>275</td>
<td>6100</td>
<td>22.2</td>
<td>62.6</td>
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<td>4.3</td>
<td>58.0 ± 3.0</td>
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<td>5.0</td>
<td>78.4 ± 5.3</td>
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<td>5020</td>
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<td>66.7</td>
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<td>92.7 ± 5.6</td>
<td>25.9</td>
<td>170</td>
<td>3620</td>
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<td>37.7</td>
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Data pertaining to radioactivity have been converted to influx rate of 10,000 cpm/min·kg.
* Mean ±SD of three to six determinations obtained during the last 60–220 min of AcAc-14C infusion (steady period of blood ketone radioactivity).
‡ Mean of two to six values obtained during the last 30–150 min of AcAc-14C infusion (steady-state period for efflux rate of 14CO2).
§ Dogs made alkalotic by infusing sodium bicarbonate.

of ketone concentration is shown at the lower part of the figure. Values corresponding to normal dogs in whom hepatic ketone production was directly measured by catheterization of the hepatic and portal veins are situated on this regression line. This suggests that the same linear relationship between transport and concentration holds for a very wide range of concentrations from the low levels encountered in normal fasted dogs up to the high levels obtained by infusing AcAc. Fig. 4 shows clearly that all the values for diabetic animals are situ-
Table IV
Transport and Oxidation of Acetooacetate in Normal Dogs Infused with Acetooacetate

<table>
<thead>
<tr>
<th>Dog no.</th>
<th>AcAc infused</th>
<th>Specific activity of AcAc in blood*</th>
<th>% of infused AcAc excreted in urine</th>
<th>Net transport of AcAc*</th>
<th>Efflux rate of CO2</th>
<th>Efflux rate of 14CO2</th>
<th>Specific activity</th>
<th>% AcAc oxidized</th>
<th>% CO2 from AcAc</th>
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<tr>
<td></td>
<td>μmoles/min·kg</td>
<td>cpm/μmol C</td>
<td></td>
<td></td>
<td>cpm/min·kg</td>
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<tr>
<td>36</td>
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Data pertaining to radioactivity have been converted to influx rates of 10,000 cpm/min·kg.
* Mean of three values obtained during the last hour of ketone infusion (steady-state period for efflux rate of 14CO2). For dogs 38 and 39, mean of two values recorded during the last 30 min of infusion.

Oxidation of ketones. The fraction of AcAc taken up by tissues and promptly oxidized to CO2 averaged 43.2% (31.7-54.7%) in normal animals, and 49.9% (35.1-62.6%) in diabetic dogs. The difference between groups was not significant. The apparent percentage of respiratory CO2 derived from AcAc was positively correlated with transport of AcAc as shown in Fig. 5. At high rates of transport (50-60 μmoles/min·kg), about 50% of expired CO2 could be accounted for by the oxidation of AcAc. As discussed below, excretion of 14CO2 underestimates the actual contribution of ketones to oxidative metabolism.

Influence of blood pH on the concentration and the rate of utilization of blood ketones. The influence of an increase in blood pH on the rate of ketone uptake was tested in two diabetic dogs. Dog 50 (Tables I and II) was studied during two successive stages of the same experiment. During the first stage (0-270 min), blood pH was 7.38, the concentration of ketones was 2.72 μmoles/ml, and transport of AcAc was 30.2 μmoles/min·kg. During the second stage (270-330 min), the constant infusion of labeled AcAc was continued, whereas the blood pH was increased to 7.56 by infusing large amounts of sodium bicarbonate. During the last 60 min of that period, the ketone concentration was 3.73 μmoles/ml (+38%), and transport was 33.9 μmoles/min·kg (+12%). This increase in transport and in the concentration of ketones was most probably independent of the infusion of bicarbonate and corresponds to the expected aggravation of ketosis with time in an animal deprived of insulin. By referring to Fig. 4 where values from both stages of the experiment have been plotted, it can be seen that alkalization did not alter the abnormal relationship between transport and concentration of ketones. The same conclusion can be drawn from experiment on dog 52 (Tables I and II and Fig. 4) where alkalosis (pH 7.55) was produced during the entire study.

Oxidation of ketones. The fraction of AcAc taken up by tissues and promptly oxidized to CO2 averaged 43.2% (31.7-54.7%) in normal animals, and 49.9% (35.1-62.6%) in diabetic dogs. The difference between groups was not significant. The apparent percentage of respiratory CO2 derived from AcAc was positively correlated with transport of AcAc as shown in Fig. 5. At high rates of transport (50-60 μmoles/min·kg), about 50% of expired CO2 could be accounted for by the oxidation of AcAc. As discussed below, excretion of 14CO2 underestimates the actual contribution of ketones to oxidative metabolism.

Influence of glucose and insulin on peripheral metabolism of ketones. The influence of the magnitude of glucose assimilation on the rate of uptake and oxidation of ketones was tested in seven normal dogs (Fig. 6). The animals were infused at a constant rate with labeled and unlabeled AcAc (33.2-65.2 μmoles/min·kg) for about 300 min. In all animals except one, a steady concentration of ketones (1.1-5.9 μmoles/ml) was achieved in blood. Simultaneously, a constant efflux rate of expired 14CO2 was observed, corresponding to 35-55% of the infused 14C. After about 180 min, a constant infusion of large amounts of glucose (16-48 mg/min·kg as a 50% solution) and insulin (4.8-6.7 mU/min·kg) was started, preceded by a priming dose corresponding to the volume infused in 15 min. The following changes were observed during the glucose-insulin infusion: (a) reduction in blood ketone levels by an average of 35% (21-49) with
a fall in βOHB-AcAc ratio from 0.86 to 0.68 (no change in rate of ketone excretion by the kidney was observed which indicates that the fall in ketone concentration resulted from an increased uptake by tissues), (b) a 50% (33–67) decrease in concentration of plasma FFA, and (c) a 40% (29–51) increase in the efflux rate of expired 14CO2.

**Pattern of ketone disappearance from blood after stopping a constant infusion.** A steady state of hyperketonemia was obtained in a normal dog by infusion of AcAc for 220 min. A tracer amount of AcAc-14C was simultaneously infused for the first 180 min and then stopped. The disappearance of labeled ketones from the blood compartment was not a single exponential function of time (Fig. 7). The rate of decay was at first rapid (with a half-life of a few minutes) and then decreased progressively. The same pattern was observed for βOHB-14C but with slower rates. In other experiments, similar patterns were observed for ketone concentration after stopping an infusion of unlabeled AcAc.

**Arteriovenous differences in ketone bodies.** The arteriovenous (A-V) differences of AcAc and βOHB across the extrahepatic splanchnic bed or across the leg were measured in normal dogs at normal fasting levels and at high concentrations observed during infusion of AcAc (Table V). At low concentrations (0.040 μmoles/ml), extraction ratio of AcAc in extra-hepatic splanchnic tissues averaged 55%, whereas it amounted to only 23% at higher concentrations (1.3 μmoles/ml). Similar values were observed for the leg. In each experiment, extraction ratio of AcAc was higher than that of βOHB.

**Hepatic production of AcAc and βOHB in normal dogs.** Blood concentrations and hepatic production rates of AcAc and βOHB were measured directly in normal animals (Table VI). AcAc accounted for 75% of the total ketone production but for only about 50% of total ketones in arterial blood.

**14C in tissues.** In dog 58, 0.5 and 0.2% of infused βOHB-14C were recovered in lipids of liver and kidneys, respectively, at the end of the experiment. Estimated recovery in muscle (estimated from 14C in lipids of thigh muscle with the assumption that muscle mass was 40% of body weight) was 1.1%, and in adipose tissue (estimated from 14C in lipids of mesenteric fat with the assumption that adipose tissue mass was 5% of body weight) was 5.5%. In another normal dog infused with AcAc and AcAc-314C for 4 hr (not reported in Tables III and IV), corresponding values were 2.0% for liver, 0.2% for kidney, and 1.1% for muscle. In liver and kidney, radioactivity was equally divided between neutral glycerides and phospholipids, and in both tissues 70–80% was in the fatty acyl moieties of these lipids. Small amounts of 14C were found in glycogen (0.1% in liver, 0.1% in kidneys, and 0.5% in muscle) and in tissue glucose (0.2% in liver and 0.1% in kidneys). Substantial quantities were, however, recovered in proteins (2.0%...
in liver, 0.4% in kidneys, and 9.4% in muscle). Thus, about 20% of the infused \(^{14}\)C was recovered in lipids and proteins of the four tissues studied.

**DISCUSSION**

Our data show that rapid uptake and oxidation of ketones takes place in tissues. The extrahepatic splanchnic area and the leg tissues, for example, extract ketones from blood very efficiently. Rapid interconversion of \(\text{AcAc}\) and \(\beta\text{OH}B\) must also take place in tissues since the specific activity of the two compounds became equal during the constant infusion of \(\text{AcAc}\)\(^{14}\)C. This indicates that the rate of transport of \(\text{AcAc}\) represents that of total ketone bodies. Bergman, Kon, and Katz have made the same observation in sheep (3). The data presented in Table V and Fig. 7 suggest that \(\text{AcAc}\) is utilized more rapidly than \(\beta\text{OH}B\). However, this is difficult to ascertain because of the interconvertibility of the two compounds. Williamson and Krebs observed no difference in rates of uptake of \(\text{AcAc}\) and \(\beta\text{OH}B\) (corrected for conversion into the other compound) in a perfused heart preparation in vitro (34), and Hagenfeldt and Wahren found similar extraction ratios for \(\text{AcAc}\) and \(\beta\text{OH}B\) across the region drained by the deep veins of the human forearm at rest (35). In contrast, values similar to those observed in this study have been obtained for the region drained by the femoral vein in resting and exercising young men (36). This is consistent with the observation that the ratio of arterial concentration to splanchnic production rate of \(\text{AcAc}\) is lower than that of \(\beta\text{OH}B\) in dogs (Table VI) and humans (37).

The main observations of this study compare the peripheral utilization of ketone bodies in normal and diabetic dogs. To make this comparison possible, the control dogs were made ketotic by infusions of sodium \(\text{AcAc}\). Ideally, the control dogs should have been infused with a mixture of \(\text{AcAc}\) and \(\beta\text{OH}B\) identical with that secreted into the hepatic vein in diabetic animals. Unfortunately, this is difficult to accomplish since \(\beta\text{OH}B\) (the physiological isomer) is not available commercially and is cumbersome to prepare in large quantities. Use of \(\text{AcAc}\) instead of \(\beta\text{OH}B\) would have been inappropriate since it is known that the \(\text{L}\) (+) isomer is oxidized by a different enzymatic pathway (38, 39) and at a different rate (35) than the \(\text{D}\) (−) form. Infusion of \(\text{AcAc}\) into the normal animals did not alter their \(\beta\text{OH}B\)-\(\text{AcAc}\) ratio which was on the average about half that of the diabetic animals. This difference probably reflects the more reduced redox state of mitochondria in diabetic tissues. Another consequence of the infusion of sodium \(\text{AcAc}\) in the control dogs was a rise in blood pH. Since alkalization of the blood of the diabetic animals did not change significantly the rate of uptake of ketones, it is unlikely that this influenced our results. Bergman and Kon, using techniques similar to ours, compared nondiabetic ketogenic sheep with...
normal sheep infused with AcAc and were unable to detect any difference in rate of uptake of ketones between the two groups (40). The evidence available, therefore, suggests that our dogs infused with AcAc are appropriate controls for the diabetic animals.

In both groups studied, the concentration of ketones was an exponential function of transport, but for any value less than 30 μmoles/min·kg, the total ketone concentration in the diabetic dogs was about 3 times that in normal dogs. Moreover, the maximal capacity to utilize ketones did not exceed 40 μmoles/min·kg in the pancreatectomized animals, whereas it amounted to about 70 μmoles/min·kg in the normal dogs. It is thus clear that a definite impairment of mechanisms for utilizing ketones exists in diabetic dogs and contributes to ketosis. At the highest rates of transport observed in diabetic dogs with uncontrolled hyperglycemia and high rates of transport of FFA, total ketone concentration would have been only about 2 mmoles/liter if capacity to remove ketones had remained normal (Fig. 4). Actual values were

![Figure 6](image-url)

**Figure 6** Effect of glucose and insulin on concentration and oxidation of ketone bodies in normal dogs infused with AcAc. The thick lines represent mean values.
It is reasonable to conclude, therefore, that diabetic ketoacidosis would not occur in the absence of the observed defect in utilization.

On an average, about 50% of the infused AcAc-\(^{14}\)C appeared as \(^{14}\)CO\(_2\) in the expired air irrespective of the quantity of AcAc utilized and without difference between the two groups. Similar values were obtained in normal and ketotic sheep by Bergman et al. (3). It must be assumed that the radioactivity which did not appear in expired CO\(_2\) was incorporated into other compounds. At the end of sample experiments, about 11% of the infused radioactivity was estimated to be in proteins (mainly in muscle), about 8% in lipids (mainly in adipose tissue but some in liver, muscle, and kidney) and a small amount in tissue glucose and glycogen. Since there is no net conversion of acetyl CoA–carbon to amino acids or trioses, the \(^{14}\)C in proteins and lipid-glycerol must have resulted from exchange processes. For protein, this could occur at several steps of the citric acid cycle such as the reversible conversion of \(\alpha\)-ketoglutarate to glutamate and of oxaloacetate to aspartate. Exchange reactions can also account for appearance of \(^{14}\)C in glucose and glycogen. Our estimates of oxidation to CO\(_2\) are artifactually low to the extent of these isotopic exchanges, which appear to represent at least 10–15% of the infused \(^{14}\)C. Since our isotopic data show that at high rates of transport of ketones (40–60 \(\mu\)moles/min·kg), the percentage of expired CO\(_2\) which is derived from oxidation of AcAc approaches 50%, the actual value probably exceeds 60% of the total CO\(_2\) production. That the fate of \(^{14}\)C in compounds metabolized in gluconeogenic tissues can differ from net fate of the substance has recently been reemphasized (37, 41). The present results indicate that similar phenomena occur in other tissues. The conversion of AcAc-\(^{14}\)C to lipids presumably represents net synthesis of fatty acids from acetyl CoA and appeared to be a substantial pathway in normal animals under the conditions of the experiment. Unfortunately, no such information was obtained in the diabetics.

We did not test whether exogenous insulin could correct the defect in maximal capacity to utilize ketones in diabetic dogs because its administration would inhibit production of ketones by the liver. This would reduce the ketone concentration and make it very difficult to

**Table V**

| Group                  | No. of experiments | Metabolite | Artery Cpm/ml | Femoral vein Cpm/ml | Portal vein Cpm/ml | A-V | A-V/A
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<tbody>
<tr>
<td>Normal</td>
<td>8</td>
<td>AcAc</td>
<td>0.040 ± 0.006</td>
<td>0.019 ± 0.006</td>
<td>0.021 ± 0.004</td>
<td>0.55 ± 0.10</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td>(\beta)OHB</td>
<td>0.033 ± 0.004</td>
<td>0.028 ± 0.003</td>
<td>0.006 ± 0.001</td>
<td>0.16 ± 0.04</td>
<td></td>
</tr>
<tr>
<td>Normal infused</td>
<td>4</td>
<td>AcAc</td>
<td>1.30 ± 0.35</td>
<td>1.03 ± 0.30</td>
<td>0.27 ± 0.09</td>
<td>0.23 ± 0.07</td>
<td></td>
</tr>
<tr>
<td>with AcAc</td>
<td></td>
<td>(\beta)OHB</td>
<td>0.67 ± 0.13</td>
<td>0.68 ± 0.14</td>
<td>-0.01 ± 0.01</td>
<td>0.67 ± 0.14</td>
<td></td>
</tr>
<tr>
<td>Normal infused</td>
<td>3</td>
<td>AcAc</td>
<td>1.65 ± 0.43</td>
<td>1.32 ± 0.41</td>
<td>0.33 ± 0.15</td>
<td>0.20 ± 0.08</td>
<td></td>
</tr>
<tr>
<td>with AcAc</td>
<td></td>
<td>(\beta)OHB</td>
<td>0.82 ± 0.18</td>
<td>0.80 ± 0.18</td>
<td>0.02 ± 0.01</td>
<td>0.20 ± 0.08</td>
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In each experiment, three to seven determinations obtained over a 60–120 min period were averaged. Values in the table are the mean ± SEM of all experiments in each group.

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**Figure 7** Pattern of disappearance of labeled ketone bodies from blood at the end of a constant infusion of AcAc-\(^{14}\) in a normal dog infused with AcAc.
detect an effect of insulin on peripheral uptake. Therefore, we tested the action of insulin on ketone uptake using animals in which the inflow of ketones into the blood compartment could be maintained. Normal animals infused at a constant rate with AcAc in large amounts fulfilled these conditions. Indeed, their endogenous production of ketones was negligible compared with the amount infused. Under these experimental conditions, large amounts of insulin and glucose produced a 35% reduction in blood ketone concentration, indicating an equivalent increase in turnover rate (Fig. 6). Moreover, insulin and glucose increased significantly the fraction of AcAc-14C which appeared in CO2. These data are in apparent conflict with the observation that the fractional oxidation of ketones in diabetic dogs was not significantly different from that in normal animals. It should be pointed out, however, that there was substantial variation in the percentage of ketones converted to CO2 in both groups so that the influence of diabetes might be difficult to detect. The apparent increase in oxidation of AcAc may represent reduction in exchange reactions leading to incorporation of 14C. This interpretation is supported by the report of Söling, Zahlten, Reinold, and Willms that glucose decreased incorporation of AcAc-14C into glycogen and glyceride-glycerol in rat epididymal adipose tissue in vitro (42).

Many studies have been devoted to the question of the role of insulin in regulating ketone uptake by tissues. As early as in 1928, it was postulated by Shaffer (quoted by Campbell and Best [4]) that ketosis could be caused by decreased rate of utilization of ketones in peripheral tissues with or without change in rate of production by the liver. However, numerous studies which appeared in the 1930's concluded almost unanimously that neither diabetes nor insulin had any influence on peripheral utilization (4). More recent literature, however, contains data that lead to opposite conclusions. Scow and Chernick observed that the clearance of ketones from blood after single intravenous injection of D(-)-OHB or AcAc was impaired in pancreatectomized rats and was restored to normal by administration of insulin (20). Söling, Garlepp, and Creutzfeldt confirmed the existence of a relationship between glucose assimilation and ketone uptake in eviscerated, nephrectomized rats infused with glucose and AcAc at constant rates. They showed in fasted, starved, or alloxan-diabetic rats that increase in the rate of glucose infusion (with or without insulin) enhanced the uptake of AcAc by tissues (43).

The in vitro approach to the problem with various tissues has not given uniform results. Beatty and co-workers (17-19) found that both diaphragm and skeletal muscle fibers from diabetic rats took up and oxidized less AcAc than diaphragm and fibers from control rats. Addition of insulin alone or glucose and insulin enhanced the uptake of AcAc by muscle from both control and diabetic rats, but glucose alone was without effect. Insulin increased the production of CO2 from labeled AcAc but had no effect on the percentage of incorporated AcAc appearing as CO2. Neptune, Sudduth, Fash, and Reish showed that glucose stimulates to a small extent the oxidation of OHB by rat diaphragm but has no effect on the oxidation of AcAc (44). In contrast, Williamson and Krebs observed that insulin with or without glucose inhibited the removal and the oxidation of AcAc by the perfused rat heart (34). Recently, Söling et al. (42) showed that glucose enhanced the uptake of AcAc and OHB by rat adipose tissue, stimulated the conversion of ketones into fatty acids and inhibited ketone oxidation. These effects were increased by insulin. Finally, several authors have observed that glucose stimulates oxidation of ketones in rat brain (45-47).

The available literature contains very little quantitative information on the utilization of ketones in intact animals. Bates, Krebs, and Williamson (48) measured the half-life of ketones in rats from the decline in specific activity of OHB in blood after a single intravenous injection of D(-)-OHB-14C. Their data show that the average half-time is 3.4 min in normal fed rats,

<table>
<thead>
<tr>
<th>TABLE VI</th>
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<tr>
<td><strong>Arterial Concentration and Rates of Hepatic Production of Ketone Bodies</strong></td>
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<tr>
<td><strong>in Normal Postabsorptive Dogs</strong></td>
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<td></td>
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<tr>
<td>Arterial concentration (μmoles/ml)</td>
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<tr>
<td>Hepatic production rate (μmoles/min·kg)</td>
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In each experiment, three to six determinations obtained over a 60-120 min period were averaged. Figures in the table are mean ± SEM for seven experiments.

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2 If a steady state could be obtained, the effect of small continuous infusions of insulin into diabetic dogs might provide quantitative information on the ability of insulin to correct the defect in utilization.
11.7 min in rats starved 48 hr, and 18.6 min in alloxan-diabetic rats. The average ketone concentration was, respectively, 0.18, 1.96, and 9.4 μmoles/ml. These data do not permit comparison of removal capacities of peripheral tissues in normal and diabetic states because of the great difference in ketone concentrations among the three groups. Indeed, our data in dogs, those of Nelson, Grayman, and Mirsky in rats (8) and those of Wick and Drury in rabbits (9) indicate that the turnover rate of ketones decreases (half-time increases) as the concentration in blood rises. The work of Bates et al. (48) raises another comment. It is probably hazardous to calculate half-lifetimes from an apparent exponential rate of removal of labeled ketones in blood after pulse injection. When disappearance of labeled ketones from blood was measured at short intervals after stopping a constant infusion of AcAc-14C, we found that removal was not a single exponential function of time, thus preventing simple calculation of turnover rate (Fig. 7). Yet, our technique has the advantage of providing for complete equilibration of the label into its space of distribution at the start which is not the case with single injection techniques. The type of curve we obtained on semilogarithmic coordinates suggests recycling of 14C, but this cannot be the sole explanation since similar curves were obtained with unlabeled ketones. The results could be explained by variable rates of equilibration of ketones in different compartments or tissues with ketones in blood.

Uncontrolled diabetes mellitus is not the only condition in which decreased utilization of ketones contributes to ketosis. Recent studies from this laboratory have shown that ketosis in adults with glycogenosis type I (glucose-6-phosphatase deficiency) can be ascribed almost exclusively to defective peripheral utilization (49). This disease resembles uncontrolled diabetes in several features: reduced utilization of glucose, low insulin levels, and high concentrations of FFA. In contrast, utilization of ketones is unimpaired during the hypoglycemic ketosis of pregnant sheep (40).

Our data provide no information on how insulin (or the changes in glucose utilization it induces) regulates the utilization of ketones in peripheral tissues. One likely possibility is that insulin acts by decreasing the concentration and the rate of oxidation of FFA which might compete with ketones as a fuel. If this were true, then the high concentration of FFA in diabetes would be the cause of both the overproduction and the underutilization of ketones. Another possibility has been suggested by Hatfi and Fakouhi (50) who showed that the oxidation of AcAc by heart mitochondria is controlled by the concentration of inorganic phosphate, ADP and ATP. Through reactions at the level of substrate phosphorylation, inorganic phosphate and ADP reduce the mitochondrial concentration of succinyl CoA which is required for activation and subsequent oxidation of AcAc. They have suggested that the inhibition of AcAc oxidation by inorganic phosphate might play a role in the underutilization of ketones by diabetic tissues since serum inorganic phosphate levels are increased in diabetes and revert to normal when insulin is given.

Recent studies of Bieberdorf, Chernick, and Scow (51) indicate that suppression of ketogenesis by insulin is not entirely dependent upon inhibition of lipolysis in adipose tissue. Thus, it appears that insulin rapidly inhibits ketosis at multiple sites involving rates of fat mobilization from adipose tissue and the hepatic production and peripheral utilization of ketone bodies.

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