Ketone body metabolism and cardiovascular disease

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Cotter DG, Schugar RC, Crawford PA. Ketone body metabolism and cardiovascular disease. Am J Physiol Heart Circ Physiol 304: H1060–H1076, 2013. First published February 8, 2013; doi:10.1152/ajpheart.00646.2012.—Ketone bodies are metabolized through evolutionarily conserved pathways that support bioenergetic homeostasis, particularly in brain, heart, and skeletal muscle when carbohydrates are in short supply. The metabolism of ketone bodies interfaces with the tricarboxylic acid cycle, β-oxidation of fatty acids, de novo lipogenesis, sterol biosynthesis, glucose metabolism, the mitochondrial electron transport chain, hormonal signaling, intracellular signal transduction pathways, and the microbiome. Here we review the mechanisms through which ketone bodies are metabolized and how their signals are transmitted. We focus on the roles this metabolic pathway may play in cardiovascular disease states, the bioenergetic benefits of myocardial ketone body oxidation, and prospective interactions among ketone body metabolism, obesity, metabolic syndrome, and atherosclerosis. Ketone body metabolism is noninvasively quantifiable in humans and is responsive to nutritional interventions. Therefore, further investigation of this pathway in disease models and in humans may ultimately yield tailored diagnostic strategies and therapies for specific pathological states.

mitochondrial function; substrate competition; HMGCS2; CoA transferase/SCOT; lipogenesis

Introduction

Metabolism of ketone bodies is conserved among eukarya, bacteria, and archaea (8, 33, 113). Ketone bodies are synthesized in liver from acetyl-CoA derived primarily from fatty acid oxidation and are transported to extrahepatic tissues for terminal oxidation during physiological states characterized by limited carbohydrate and surplus fatty acid availability [reviewed in (137, 167); Fig. 1, A and B]. Ketone body oxidation becomes a significant contributor to overall energy metabolism within extrahepatic tissues in numerous physiological states, including the neonatal period, starvation, postexercise, and adherence to low-carbohydrate diets, when circulating ketone body concentrations increase from ~50 μM in the normal fed state to up to 7 mM. Circulating ketone body concentrations rise to ~1 mM after 16–20 h of fasting in healthy adult humans but can accumulate to as high as 20 mM in pathological states such as diabetic ketoacidosis (33, 100, 167). Ketone body metabolism is not solely rooted in energy metabolism, as ketone bodies also serve as lipogenic and sterol biosynthetic substrates in many tissues, including the developing brain, lactating mammary gland, and liver (51, 59, 144, 168) (Fig. 1C). Furthermore, hepatic ketogenesis interfaces with fatty acid β-oxidation, the tricarboxylic acid (TCA) cycle, and gluconeogenesis (Fig. 2). Derangements of ketone body metabolism occur in numerous disease states, including types 1 and 2 diabetes and heart failure, and ketone body metabolism changes over the course of normal aging (56, 77, 115, 124, 125, 145, 158, 186, 193). In this review we examine 1) the biochemistry of ketone body metabolism and its regulation (in Ketone Body Metabolism), 2) the roles of ketone body metabolism in normal and pathological states of the myocardium (in Bioenergetics of Myocardial Ketone Body Oxidation and Ketone Bodies and Myocardial Disease), 3) the concept, and prospective cardiovascular disease relevance of hepatic ketogenesis, which may channel ketone bodies into lipogenesis (in Reversibility of the CoA Transferase Reaction Extends Cardiovascular Disease Targets of Ketone Body Metabolism), and 4) intersections among ketone body metabolism, cellular signaling pathways, and the gut microbiota (in Signaling Roles for Ketone Bodies and Ketone Body Metabolism, the Microbiota, and Cardiovascular Biology and Disease). Analysis of the known and prospective cardiovascular disease targets of ketone body metabolism is also provided (in Diagnostic and Therapeutic Targets of Ketone Body Metabolism). The effects of ketone body metabolism are summarized in Table 1.

Glossary

For this article we have included a short glossary of terms and concepts for the convenience of the reader.
Fig. 1. Ketone body metabolism pathways. A: ketogenesis within hepatic mitochondria is the primary source of circulating ketone bodies. B: primary metabolic fate of ketone bodies is terminal oxidation within mitochondria of extrahepatic tissues via CoA transferase [succinyl-CoA:3-oxoacid-CoA transferase (SCOT)]. Substrate competition with pyruvate-derived and fatty acyl-CoA-derived (the corkscrew arrows represent the activities of the β-oxidation spiral) acetyl-CoA are shown. C: cytoplasmic de novo lipogenesis (DNL) and cholesterol synthesis are nonoxidative metabolic fates of ketone bodies. For simplicity of C, only acetooacetate (AcAc) is depicted, although β-hydroxybutyrate (βOHB) is also a substrate for lipogenesis after it has been oxidized to AcAc via mitochondrial βOHB dehydrogenase (BDH1). AACS, acetoacetyl-CoA synthetase; ACC, acetyl-CoA carboxylase; AcAc-CoA, acetoacetyl-CoA; ATP, adenosine triphosphate; CoA-SH, free coenzyme A; FAS, fatty acid synthase; HMG-CoA, 3-hydroxymethylglutaryl-CoA; HMGCL, HMG-CoA lyase; HMGCS1, cytoplasmic HMG-CoA synthase; HMGCS2, mitochondrial HMG-CoA synthase; HMGCR, HMG-CoA reductase; NAD+, nicotinamide adenine dinucleotide oxidized (reduced); PDH, pyruvate dehydrogenase; TCA, tricarboxylic acid; mThiolase, mitochondrial thiolase; cThiolase, cytoplasmic thiolase. Thiolase activity is encoded by at least 6 genes: ACAA1, ACAA2 (encoding an enzyme known as T1 or CT), ACAT1 (encoding T2), ACAT2, HADHA, and HADHB.

AACS, acetoacetyl-CoA synthetase; a cytoplasmic enzyme that catalyzes the ATP-dependent covalent activation of acetooacetate (AcAc) by CoA to generate AcAc-CoA.

Anaplerosis, essential replenishment of TCA cycle intermediates that is required to preserve terminal oxidation of acetyl-CoA. The primary anaplerotic reaction is catalyzed by pyruvate carboxylase, which generates oxaloacetate from pyruvate, but oxidation of odd-chain fatty acids and amino acid catabolism also yield anaplerotic substrates.

BDH1, mitochondrial β-hydroxybutyrate (βOHB) dehydrogenase; an enzyme that catalyzes the NAD+/NADH coupled interconversion of AcAc and β-OHB. The $K_{eq}$ of this reaction favors β-OHB formation.

β-Oxidation, sequential derivation of two carbon units, in the form of acetyl-CoA, from fatty acyl chains that primarily occurs within the mitochondrial matrix. Some β-oxidation also occurs within peroxisomes.

DNL, de novo lipogenesis. The synthesis of new lipid from acetyl-CoA, dependent on the enzymes acetyl-CoA carboxylase and fatty acid synthase (FAS).

Fluxomics, analysis of metabolic fates of labeled substrates using sophisticated chemical profiling platforms including, nuclear magnetic resonance spectroscopy and mass spectrometry, coupled to statistical computing platforms.

Gnotobiotic, Greek roots: gnos, knowledge; bios, life; a technology that allows cultivation of experimental animal models in the presence of defined microbial communities.

GPCR, G protein-coupled receptor; a plasma membrane receptor with seven transmembrane domains that mediates an intracellular signaling cascade via an interaction with a G protein.

HMGCL, HMG-CoA lyase; a mitochondrial enzyme that cleaves HMG-CoA to liberate AcAc and acetyl-CoA during ketogenesis. This enzyme mediates ketone body production from β-oxidation-derived acetyl-CoA and leucine catabolism.

HMGCS2, HMG-CoA synthase 2; a mitochondrial enzyme that condenses β-oxidation-derived AcAc-CoA and acetyl-CoA to generate HMG-CoA and liberate a CoA moiety.

Ketogenesis, production of ketone bodies. AcAc and βOHB are the predominant circulating ketone bodies.

Ketolysis, breakdown of ketone bodies either within the mitochondria for terminal oxidation or in the cytoplasm for lipogenesis.

Ketone body oxidation, contribution of acetyl-CoA derived from ketone bodies to the TCA cycle.

Microbiota, ecosystem of 100 trillion prokaryotic cells that comprise a ~1-kg organ that harbors a massive aggregate of genomes (the microbiome).

NAFLD, nonalcoholic fatty liver disease.
P-to-O ratio, the ATP yielded per oxygen invested, which depends on many factors, including the inherent energy of combustion for a substrate, the initial energetic investment required for oxidation, and the effect of oxidizing the substrate on the electrochemical potential effected by the mitochondrial electron transport chain.

Pseudoketogenesis, quantifiable dilution of isotopically labeled ketone bodies, mediated by reversibility of thiolase and CoA transferase reactions in extrahepatic tissues.

SCOT, succinyl-CoA:3-oxoacid-CoA transferase, encoded by nuclear Oxct1. A mitochondrial enzyme that catalyzes the following near equilibrium reaction: succinyl-CoA + AcAc ↔ succinate + AcAc-CoA.

Sirtuin, silent mating type information regulation 2 homolog (SIRT); In mammals, these enzymes comprise a family of seven NAD+/H+ -dependent enzymes that coordinate an array of cellular processes via deacetylace, demalonylase, desuccinylase, and ADP-ribosylase enzymatic activities.

TCA cycle, tricarboxylic acid cycle. Also termed Krebs cycle, citric acid cycle.

Terminal oxidation, contribution of acetyl-CoA to the TCA cycle via citrate synthase enzymatic activity that results in

Fig. 2. Hepatic integration of ketogenesis. Integration of hepatic ketogenesis with hepatic TCA cycle, DNL, and pyruvate cycling/glucose metabolism. FAO, β-oxidation of fatty acids; GDH, glutamate dehydrogenase; OAA, oxaloacetate; ME, malic enzyme; PEPCK, phosphoenolpyruvate carboxykinase; PC, pyruvate carboxylase; PK, pyruvate kinase; PEP, phosphoenolpyruvate cytosolic; α-KG, α-ketoglutarate.

Fig. 3. Ketogenic flux through CoA transferase. Because CoA transferase catalyzes an equilibrium reaction, select spatiotemporal metabolic conditions may favor channeling of mitochondrial acetyl-CoA to AcAc, generating a monocarboxylate shuttle that complements the citrate-dependent tricarboxylate shuttle and permits mitochondrial efflux of β-oxidation-derived acetyl-CoA independent of the TCA cycle (TCAC). ACLY, ATP-citrate lyase; ETC, electron transport chain.
conversion of 1 mol of acetyl-CoA to 2 mol of CO₂ and 2 mol of NADH.

Thiolase, any enzyme that catalyzes the reversible condensation of acetyl-CoA to acetoacetyl-CoA. This enzymatic activity is redundant in mammals.

**Ketone Body Metabolism**

The fundamental biochemical and physiological roles of ketone body metabolism have been extensively studied and reviewed (68, 137, 167). In this section, we provide a brief overview of ketone body metabolism and then focus on recent studies revealing novel aspects of the pathway’s regulatory mechanisms.

**Hepatic ketogenesis.** In liver, 3-hydroxybutyryl-CoA synthase 2 (encoded by the nuclear gene *Hmgcs2*) catalyzes a rate committing ketogenic condensation: β-oxidation of acetoacetate-acetylated-CoA (AcAc-CoA) and acetyl-CoA to generate HMG-CoA, which is cleaved by HMGCL to generate AcAc. AcAc is reduced to D-OHB, in an NAD⁺/NADH-coupled equilibrium reaction catalyzed by phosphatidylcholine-dependent mitochondrial N-β-OHB dehydrogenase (BDH1), in which the *Kₐₙ* favors N-β-OHB formation (26, 119) (Fig. 1A). Because the BDH1 catalyzed reduction/oxidation of AcAc and N-β-OHB is common to the final reaction of ketogenesis and the first reaction of ketone body oxidation (Fig. 1A, and B), BDH1 modulates mitochondrial redox potential in liver and extrahepatic tissues, wherein the AcAc-to-β-OHB ratio is directly proportional to the mitochondrial NAD⁺-to-NADH ratio (230). A cytoplasmic N-β-OHB dehydrogenase (BDH2) with only 20% sequence identity to BDH1 has been identified (72). More work is needed to elucidate the mechanisms by which the BDH enzymes are regulated and function in vivo. AcAc is also nonenzymatically decarboxylated to acetone. Ketone bodies are released by the liver via solute carrier 16A (SLC16A) family members 1, 6, and 7 and circulate to extrahepatic tissues where they primarily undergo terminal oxidation (75, 76, 91).

**KETOGENIC SUBSTRATES.** Ketogenesis predominantly occurs in liver mitochondria at rates proportional to β-oxidation when dietary carbohydrates are limiting and is highly integrated with the TCA cycle and gluconeogenesis (Fig. 2). Biochemical studies by pioneering investigators including Krebs, McGarry, and Foster demonstrated that hepatic metabolic fluxes of acetyl-CoA govern rates of ketogenesis [reviewed in (68, 137)]. Fatty acid β-oxidation-derived AcAc-CoA and acetyl-CoA are the primary ketogenic substrates. Glucose metabolism accounts for <1% of circulating ketone bodies in states of low-carbohydrate intake because pyruvate predominantly enters the hepatic TCA cycle via carboxylation to oxaloacetate or malate rather than decarboxylation (to acetyl-CoA) (99, 134, 142). In addition, amino acid catabolism accounts for a small percentage of circulating ketone bodies, with leucine catabolism generating up to 4% of circulating ketone bodies in the post-absorptive state (207).

**REGULATION OF KETOCENIC MEDIATORS.** Key regulatory steps in ketogenesis include lipolysis of fatty acids from triacylglycerols, transport to and across the hepatocyte plasma membrane, transport into mitochondria via allosterically regulated carnitine palmitoyltransferase 1, the β-oxidation spiral, and the hormonal regulators of these processes, predominantly glucagon and insulin. These classical mechanisms have been reviewed (55, 137, 138, 194). Hepatic ketogenesis is a spillover pathway for β-oxidation-derived acetyl-CoA generated in excess of the liver’s energetic needs (Fig. 2). Acetyl-CoA subserves several roles integral to hepatic intermediary metabolism beyond ATP generation via terminal oxidation. Acetyl-CoA allosterically activates 1) pyruvate carboxylase, thereby activating a metabolic control mechanism that augments anaplerotic entry of metabolites into the TCA cycle (154, 185) and 2) pyruvate dehydrogenase kinase, which phosphorylates and inhibits pyruvate dehydrogenase (39), thereby further enhancing flow of pyruvate into the TCA cycle via anaplerosis. Furthermore, cytoplasmic acetyl-CoA, whose pool is augmented by transport mechanisms that convert mitochondrial acetyl-CoA to transportable metabolites (see CoA transferase-dependent ketogenesis in extrahepatic tissues), inhibits fatty acid oxidation: acetyl-CoA carboxylase catalyzes the conversion of acetyl-CoA to malonyl-CoA, the lipogenic substrate and an allosteric inhibitor of mitochondrial carnitine palmitoyltransferase 1, decreasing delivery of acyl chains to the mitochondrial matrix for terminal oxidation [reviewed in (102, 137)]. Thus the mitochondrial acetyl-CoA pool regulates and is regulated by the spillover pathway of ketogenesis, which orchestrates key aspects of hepatic intermediary metabolism. The studies described below focus on recently discovered mechanisms of ketogenic regulation.

Following procession through β-oxidation, ketogenic fate is determined by HMGCS2, whose regulatory mechanisms are schematized in Fig. 4A. Because of its ability to support high rates of hepatic ketogenesis, divergence of this mitochondrial HMGCS ~500 Mya from the gene encoding cytoplasmic HMGCS1 may have supported the emergence of increasing brain weight-to-body weight ratios during vertebrate evolution (30, 43, 161). The *Hmgcs2* gene and encoded protein are regulatory targets during the transition to extrauterine life; in starvation, diabetes, and aging; and during adherence to low-carbohydrate/high-fat (ketogenic) diets, and in aging (17, 33, 68, 84, 179, 186). The *Hmgcs2* gene is dynamically regulated at the transcriptional level. Methylation of 5’ regulatory sequences within the *Hmgcs2* gene silences its transcription in fetal liver and in nonketogenic adult tissues (12). At birth,
In addition to cysteine palmitoylation, HMGCS2 enzymatic activity is responsive to other forms of post-translational modification. Mitochondrial activity of HMGCS2 is inhibited through allosteric regulation by succinyl-CoA (84, 128, 164) and through succinylation of lysine residues (9, 84, 162, 209). While the regulatory mechanisms remain to be determined, desuccinylation is stimulated by glucagon (162), and recent discoveries that identify sirtuin (SIRT5) as a mitochondrial NAD⁺-dependent lysine desuccinylase prompt the hypothesis that SIRT5 function dynamically regulates mitochondrial HMGCS2 activity (47, 243). HMGCS2 activity is also induced in the post-absorptive state by SIRT3, a mitochondrial NAD⁺-dependent deacetylase (188). Because SIRT3 regulates multiple enzymatic mediators of β-oxidation, sirtuins may be multilayered regulators of ketogenesis. Genetically obese mice were recently shown to exhibit increased hepatic HMGCS2 serine phosphorylation, a post-translational modification that enhances HMGCS2 activity in vitro and was correlated with elevated serum \( \Delta \)-BOHB levels in vivo (70).

Despite all of the described transcriptional and post-translational mechanisms that regulate HMGCS2, it is unknown whether regulation of Hmgs2 gene expression correlates with changes in protein abundance and whether these changes in gene expression directly regulate ketogenesis. Importantly, it remains undetermined whether HMGCS2 post-translational modification itself is a primary determinant of ketogenic flux. Nonetheless, HMGCS2 activity is required for the normal ketogenic response to states of diminished carbohydrate intake, as its absence in humans yields hypoketotic hypoglycemia and fatty liver in states of diminished carbohydrate intake (7, 28, 208).

Ketone body utilization. Ketone body catabolism generates acetyl-CoA that can be terminally oxidized within the TCA cycle or used for sterol biosynthesis and DNL. Ketone bodies are an alternative and glucose-sparing fuel source, avidly oxidized in heart and muscle. Moreover, neurons do not effectively generate high-energy phosphates from fatty acids and consequently oxidize ketone bodies during starvation and in the neonatal period [presented and reviewed in (33, 50, 167, 225, 235)]. Below we review the biochemical activities of the enzymes of ketone body utilization, focusing on recent studies that reveal the precise physiological roles of ketone body utilization in vivo.

KETONE BODY OXIDATION. Ketone bodies are extracted from the circulation by peripheral tissues via SLC16A1 and -7 (75, 76). Within mitochondria of peripheral organs, BDH1 catalyzes the oxidation of \( \Delta \)-BOHB to AcAc. AcAc is activated to AcAc-CoA by SCOT (CoA transferase), the only mammalian CoA transferase that catalyzes a near equilibrium reaction which exchanges coenzyme A between succinate and AcAc (Fig. 1B). The free energy released by hydrolysis of AcAc-CoA is greater than that of succinyl-CoA (198). Therefore, the equilibrium of this reaction thermodynamically favors the formation of AcAc ([229], also see Reversibility of the CoA Transferase Reaction Extends Cardiovascular Disease Targets of Ketone Body Metabolism). Thus ketone body oxidative flux occurs by mass action: an abundant supply of AcAc and rapid utilization of acetyl-CoA through citrate synthase favors AcAc-CoA formation and contribution of ketone bodies to the TCA cycle. A reversible AcAc-CoA thiolase reaction yields two molecules of acetyl-CoA that enter the TCA cycle (Fig. 1B) (61, 66, 167, 229). CoA transferase is necessary, but not

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**Fig. 4. Regulatory mechanisms for HMGCS2 and CoA transferase (SCOT).** HMGCS2 (A) and SCOT (B) both undergo transcriptional (red) and post-translational modes of regulation (blue). Some regulatory factors exhibit both classes of effects (violet). mTORC1, mammalian target of rapamycin complex 1; PPARα, peroxisome proliferator-activated receptor-α; PTM, post-translational modification; SIRT3, sirtuin (silent mating type information regulation 2 homolog) 3.
Ketone body oxidation becomes a primary contributor to bioenergetic homeostasis during ketotic states, during which ketone bodies are oxidized in proportion to their delivery in heart, brain, and muscle until saturation of either uptake or oxidation occurs (17, 33, 155, 166, 199, 225). Relative to ketogenesis, little is known about the regulation of mediators of ketone body oxidation. Perhaps paradoxically, expression of the gene encoding CoA transferase (Oxct1), CoA transferase protein abundance, and CoA transferase enzymatic activity are all diminished in rodent heart and muscle during states of sustained ketosis (53, 54, 71, 152, 213, 228). In pathological contexts such as diabetic ketoacidosis, diminution of CoA transferase abundance and activity limit ketone body disposal, whereas insulin deficiency (or severe impairment of its signaling) stimulates peripheral fatty acid mobilization and unabated hepatic ketogenesis. This mismatch of ketogenesis and peripheral disposal predisposes to the extreme hyperketonemia that can emerge in diabetic ketoacidosis.

In ketotic states, expression of Oxct1 may be negatively regulated through mechanisms involving PPARs that are abrogated by insulin [(228); Fig. 4B]. Myocardial Oxct1 expression may also be downregulated under select nonketotic metabolic states, as transgenic mice overexpressing the non-insulin-dependent glucose transporter (GLUT1/SLC2A1) in cardiomyocytes exhibit decreased Oxct1 expression and diminished ketone body oxidation (234). Future experiments are needed to determine whether regulation of Oxct1 expression occurs through transcriptional or post-transcriptional mechanisms (or both), which may help support insight into whether these events are adaptive or maladaptive. Post-translational modification of CoA transferase occurs through nonenzymatic tyrosine nitration, which has been observed in hearts of endotoxin-treated rats, streptozotocin-treated rats, and db/db mice (135, 213, 223). Tyrosine nitration correlates with impairments of enzymatic activity (Fig. 4B). Conversely, nitration of CoA transferase on tryptophan residues in aged rat hearts and kidneys appears to augment enzymatic activity (31, 163). Molecular mechanisms of residue-specific nitration or denitration designed to modulate CoA transferase activity may exist and require elucidation, as do additional prospective post-translational modifications of CoA transferase.

Although touted as energy-efficient substrates (180, 219), an energetic requirement for ketone body oxidation at the cellular level has never been demonstrated, even in states of diminished carbohydrate supply. Nonetheless, human inborn errors of ketone body oxidation do result in clinically significant disease, as ~30 CoA transferase-deficient patients have been identified. These patients present early in life with severe ketoacidosis that results in lethargy, vomiting, and coma, requiring aggressive intravenous hydration, bicarbonate, and glucose and insulin therapy (20, 62, 105, 143, 149, 181, 192, 210). Some CoA transferase-deficient patients also present with hypoglycemia, and CoA transferase deficiency may contribute to a subset of idiopathic ketotic hypoglycemia cases (20, 27, 52, 83, 92, 108). If maintained on carbohydrate-rich diets that are mildly reduced in protein content, CoA transferase-deficient patients seem to thrive, albeit with persistent hyperketonemia. To mechanistically dissect the homeostatic roles of ketosis, germline CoA transferase-knockout (SCOT-KO) mice were developed (40). These mice cannot oxidize ketone bodies and invariably die within 48 h of extrauterine life because of hyperketonemic hypoglycemia that is marked by hypolactatemia and an increase in the serum AcAc-to-β-HB ratio. The mechanisms and consequences of this unusually high ratio remain to be determined. Despite the critical nature of CoA transferase and ketone body disposal at an organismal level, recently published studies demonstrated that mice with selective loss of CoA transferase in either cardiomyocytes, neurons, or skeletal myocytes—the three greatest consumers of ketone bodies (64, 167)—survive the neonatal period and starvation as adults, with only subtle metabolic abnormalities in these two states (41). Future studies using these and related models will be important to determine the metabolic and bioenergetic adaptations to CoA transferase deficiency and the inability to derive energy from ketone bodies.

Nonoxidative metabolism of ketone bodies. Ketone bodies also contribute to lipogenesis and sterol biosynthesis in developing brain, lactating mammary gland, and liver following enzymatic and endergonic activation of AcAc to AcAc-CoA by cytoplasmic AACS [(51, 59, 144, 168) and Fig. 1C]. While AcAc-CoA serves as a direct substrate for cytoplasmic HMGS1, which catalyzes the fate-committing step of sterol biosynthesis, channeling of AcAc-CoA into DNL requires a thiolytic cleavage reaction (i.e., cytoplasmic ketolysis) to yield acetyl-CoA, which becomes the lipogenic substrate malonyl-CoA upon carboxylation (19, 49, 51, 67, 226). Recent studies support the physiological importance of ketone bodies as anabolic substrates. Genetic knockdown of AACS in mouse liver lowered total blood cholesterol in vivo, and in vitro AACS knockdown impaired differentiation of primary mouse embryonic neurons and inhibited adipocyte differentiation of 3T3-L1 cells [(79–81), also see Reversibility of the CoA Transferase Reaction Extends Cardiovascular Disease Targets of Ketone Body Metabolism]. While lipogenic fates of ketone bodies may support essential biological functions, these pathways do not serve as a primary pathway for disposal of ketone bodies when hepatic ketogenesis is stimulated, because CoA transferase deficiency in mice and humans, which abrogates terminal ketone body oxidation, causes severe hyperketonemia (20, 40, 60, 62, 105). Nonetheless, future studies in models of AACS disruption are merited to firmly establish the physiological roles for nonoxidative fates of ketone bodies.

**Bioenergetics of Myocardial Ketone Body Oxidation**

In the normal adult heart, mitochondrial oxidative phosphorylation provides more than 95% of the ATP generated for its mechanical, electrical, and homeostatic activities. Fatty acid oxidation provides up to 70% of the ATP produced by the heart, with metabolism of glucose, lactate, amino acids, and ketone bodies supplying the balance [reviewed in (127)]. Cardiomyocytes demonstrate considerable metabolic flexibility during dynamic alterations of nutrient state and hemodynamic stress, conferring an important adaptive property to the myocardium (22, 203, 204, 239). Myocardium is the highest ketone body consumer per unit mass (17, 33, 61). Cardiomyocytes oxidize ketone bodies in proportion to their delivery, at the expense of terminal fatty acid oxidation and glucose oxidation (22, 42, 65, 82, 98, 156, 206, 234). Competition between...
oxidation of ketone bodies and fatty acids is independent of changes in myocardial malonyl-CoA concentrations (196).

Ketone body oxidation is more energetically efficient than terminal fatty acid oxidation. While terminal fatty acid oxidation yields the highest theoretical payoff of ATP per C2 unit of all the myocardial substrates, the initial ATP investment required to activate long-chain fatty acids for oxidation (i.e., to generate a fatty acyl-CoA thioester) and fatty acid-induced expression of uncoupling proteins may actually confer greater energetic density to ketone bodies (219). Furthermore, compared with terminal fatty acid oxidation, proportionately more of the reducing equivalents generated by ketone body oxidation are delivered via NADH to complex I within the electron transport chain. Oxidation of βOHB also increases the redox span between complexes I and III by keeping mitochondrial ubiquinone oxidized. This may increase the potential energy harvested from oxidation of ketone bodies (relative to fatty acids), thereby yielding more energy available for P-to-O ratio, improving the energetic efficiency of oxidizing ketone bodies over fatty acids (104, 180, 219). Because unrestrained mitochondrial fatty acid oxidation may augment the generation of reactive oxygen species (ROS) to levels that exceed scavenging mechanisms, competitive contribution of ketone bodies to the TCA cycle in cardiomyocytes may be adaptive under conditions in which the ability to switch between fatty acids and glucose becomes impaired (104, 180, 219). Moreover, ketone bodies diminish oxidative stress by scavenging free radicals and by maintaining ubiquinone in the oxidized state (Fig. 5) (74, 130, 219).

CoA transferase-dependent activation of AcAc to AcAc-CoA sequesters free CoA-SH (90, 176). Rodent hearts perfused ex vivo with AcAc alone exhibit depleted free CoA-SH pools and require addition of either CoA precursors or anaplerotic substrates to maintain anaplerotic and TCA cycle flux and to prevent functional decline in cardiac performance (175–177). Although ketone bodies do not serve as the sole myocardial fuel under any in vivo circumstance, these studies underscore the important role of cardiac anaplerosis in maintaining myocardial bioenergetic homeostasis and cardiac function, a topic that was recently reviewed by Des Rosiers and colleagues (44).

Until recently, it was not known how the myocardium adapts to chronic ketosis, an important question relevant to the role that ketone body metabolism may serve in myopathic hearts. Mouse models of physiological ketotic nutritional states (24 h...
of fasting or 4 wk of a ketogenic diet) demonstrate that the myocardium engages a transcriptional program including, as described above, transcriptional suppression of Oxt1 and diminution of CoA transferase protein (228). Consistent with diminution of CoA transferase, nuclear magnetic resonance profiling demonstrated that maintenance on a ketogenic diet decreased myocardial $^{13}$C-enrichment of glutamate from $^{13}$C-labeled ketone bodies that were delivered in vivo or ex vivo by 25%, indicating diminished terminal oxidation of ketone bodies in the TCA cycle (101, 228). Furthermore, attenuation of ketone body oxidation correlated with failure of ketone bodies to inhibit contribution of fatty acids to the TCA cycle. Together, these results indicate that ketotrophic environments induce mechanisms that modulate myocardial utilization of ketone bodies. Because hemodynamic parameters were preserved in hearts of mice fed a ketogenic diet, these results suggest that modulation of ketone body metabolism may be adaptive in the setting of sustained ketosis.

**Ketone Bodies and Myocardial Disease**

Numerous experimental approaches have established that cardiomyopathy is associated with changes in cardiac substrate and energy metabolism and that altered energy metabolism can cause cardiomyopathy [reviewed in (1, 10, 29, 44, 89, 94, 97, 109, 118, 127)]. Both metabolic and bioenergetic lines of evidence indicate that ketone body metabolism could play a significant role in the myopathic heart. Hepatic ketogenesis is stimulated and ketone bodies circulate at increased concentrations in the setting of heart failure in a relationship directly proportional to filling pressure (115, 124, 125, 145, 158). As previously noted, the myocardium oxidizes ketone bodies at the expense of fatty acid oxidation (22, 82, 196, 218), and thus reductions of myocardial fatty acid oxidation that occur during the development of advanced cardiomyopathy may not be coupled to reductions of ketone body oxidation (1, 10, 127). A study in humans with advanced heart failure indicated that myocardial extraction of delivered ketone bodies is maintained in the failing heart but not skeletal muscle (96). Similarly, the contribution of ketone bodies to cardiac energy metabolism may be elevated in patients with dilated and hypertrophic cardiomyopathies (174). Therefore, while the data on ketone body metabolism in heart failure are currently very limited, diminished myocardial ketone body oxidation could promote pathological outcomes. Of the identified CoA transferase-deficient patients, two were reported to present with dilated cardiomyopathy (63, 126, 210). Future studies in humans that specifically measure CoA transferase function and ketone body oxidation in cardiomyopathic states, complemented by mechanistic studies using tissue-selective genetic rodent models, will be required to definitively determine how myocardial ketone body metabolism changes in pathophysiological states and the contexts in which myocardial ketone body utilization may be adaptive or maladaptive.

Studies in rodents demonstrate a possible role for ketone body metabolism in myocardial adaptation to ischemia-reperfusion injury. In two studies using ex vivo perfusion approaches in rat hearts, maintenance on low-carbohydrate diets before ischemia-reperfusion protocols gave conflicting results with regard to infarct size and hemodynamic performance (6, 221). Prospective cardioprotective effects of a low-carbohydrate diet may be attributable to an increase in the number of myocardial mitochondria or transcriptional upregulation of key mediators of oxidative phosphorylation (6, 112). Cardioprotective effects have been observed using in vivo ischemia-reperfusion approaches in rats subjected to starvation-induced ketosis, initiated through prolonged fasting, and also via intravenous injection of DL-BOHB immediately before ischemic injury, which conferred a significant decrease in both infarct size and myocardial cell death (191, 244). These studies should stimulate further investigation using genetic and pharmacological approaches to determine the validity and mechanisms of these preliminary studies and whether the observations actually relate to myocardial metabolism of ketone bodies.

Relationships among ketone body metabolism, membrane excitability, and arrhythmogenesis have also been reported. Metabolomic and proteomic analysis of atrial tissue harvested during cardiac surgeries demonstrated an increase in both myocardial βOHB content and CoA transferase abundance in patients with atrial fibrillation compared with control patients (136). Although myocardial ketone body utilization in the setting of atrial fibrillation has not been investigated, mechanisms that directly link ketone bodies to membrane excitability and arrhythmogenesis have been preliminarily assessed. βOHB blocks the transient outward K$^{+}$ current in murine ventricular myocytes, causing action potential prolongation (46). However, only L-βOHB inhibits the transient outward K$^{+}$ current. L-βOHB is measurable in myocardial extracts, but does not circulate and likely results from hydrolysis of the β-oxidation intermediate L-βOHB-CoA (95, 122, 165, 184). Hepatic ketogenesis only produces D-βOHB, which is the only form that is a BDH substrate, and thus a substrate for oxidation. SLC16A transporters in rat myocytes demonstrate no stereoselectivity for βOHB (222). Prospective pathophysiological significance of the L-βOHB stereoisomer requires investigation.

**Reversibility of the CoA Transferase Reaction Extends Cardiovascular Disease Targets of Ketone Body Metabolism**

Each enzyme involved in ketone body oxidation catalyzes a reversible reaction. Thus each tissue that oxidizes ketone bodies has the enzymatic potential to synthesize them.

**Pseudoketogenesis.** Brunengraber and colleagues (57) quantified exchange between isotopically labeled fatty acids and unlabeled ketone bodies in rat hearts perfused ex vivo. Des Rosiers et al. (45) subsequently quantified dilution of labeled ketone bodies in hepatectomized dogs in vivo. This isotopic dilution by extrahepatic tissues was termed pseudoketogenesis, which occurs because CoA transferase and AcAc-CoA thiolase catalyze reversible reactions. While a physiological role for pseudoketogenesis has yet to be established, the process leads to overestimation of ketone body turnover in whole body tracer dilution studies, because its rate is up to one-third of the rate of ketone body uptake (57).

CoA transferase-dependent ketogenesis in extrahepatic tissues. CoA transferase is expressed in all mammalian cells that harbor mitochondria, except hepatocytes, but not all cell types oxidize ketone bodies, suggesting that CoA transferase may mediate nonoxidative metabolic functions in certain tissues or physiological states (11, 64, 153, 229). Because AcAc exits mitochondria via monocarboxylate transporters, ketogenic flux
through thiolase and CoA transferase may permit carbon efflux from mitochondria independent of citrate synthase, thus generating a monocarboxylate transport system that complements both citrate-dependent tricarboxylate transport (Fig. 3) and acetylcarbarnine transport (172). Such a role for CoA transferase reflects its dual evolutionary history as an enzyme of ketone body production and utilization (157, 240). Ketogenic flux through mitochondrial thiolase and CoA transferase may be poised to regulate discrete cellular functions directly relevant to metabolic syndrome, diabetes, and atherosclerosis, because ketone bodies are substrates for cytoplasmic DNL and sterol biosynthesis [Fig. 3 and (19, 49, 51, 67)]. A series of studies have implicated ketogenic flux through CoA transferase as a putative mechanism through which ketone bodies act as insulin secretagogues in pancreatic β-cells (78, 131–133). The authors of these reports demonstrated that ketone bodies and ketone body precursors potentiate insulin release, possibly by contributing to formation of cytoplasmic short-chain acyl-CoAs (131–133) and that diminished CoA transferase expression correlated with impaired glucose stimulated insulin secretion in a rat insulinoma cell line (78). Future experiments will be required to determine the biological roles of ketogenic flux in cell types not classically associated with lipogenesis, including β-cells of the pancreas and cardiomyocytes.

An additional nonoxidative role for CoA transferase may emerge in macrophages, which exhibit robust CoA transferase enzymatic activity but do not terminally oxidize ketone bodies (146, 147). Recent discoveries have highlighted the importance of macrophage metabolism in cardiovascular disease [reviewed in (151)]. As macrophage-specific deficiency of the enzyme required for DNL, FAS, ameliorates diet-induced atherosclerosis in mice (182), it is intriguing to consider that anabolic procession of metabolites through mitochondrial thiolase and CoA transferase acts upstream of FAS to contribute to macrophage DNL flux. CoA transferase is also abundantly expressed in adipose tissue, in which the enzyme that catalyzes the reaction downstream from CoA transferase, AACS (Fig. 3), is dynamically regulated by PPARγ, a key mediator of adipogenesis and adipocyte function (2, 81, 233). Adipocyte-specific FAS deficiency increases baseline energy expenditure and improves diet-induced obesity in mice (123). Because ketone bodies have been proposed as quantitatively significant DNL substrates in multiple cell types (51, 238), these recent findings suggest that use of genetic models is warranted to draw implications for ketone body metabolism in a multitude of disease states, which will benefit from the development and application of both pharmacological and genetic tools.

**Signaling Roles for Ketone Bodies**

Because serum ketone body concentrations vary over a large dynamic range, they may act as physiologically relevant signals for cell-surface and intracellular receptors. In fact, d-β-OHB is an endogenous ligand for a niacin receptor, GPCR 109A (GPR109A), with an EC₅₀ of 770 μM (3). Like niacin, d-β-OHB can inhibit adipose tissue lipolysis, which has been proposed to create a negative feedback loop in which ketosis curtails ketogenesis by limiting delivery of nonesterified fatty acids to the liver (205, 212). GPR109A signaling also promotes reverse cholesterol transport in macrophages (129). It is unknown whether d-β-OHB plays a role in this cascade.

**β-OHB signaling may also influence cardiovascular biology through additional GPCRs. A recent study in mice revealed that β-OHB decreases sympathetic outflow and reduces heart rate and total energy expenditure by antagonizing (through an unknown mechanism) GPR41, a Gᵢₒ-coupled receptor for short-chain fatty acids (SCFAs, e.g., acetate, propionate, and butyrate) that is abundantly expressed in sympathetic ganglia (106). Thus evidence implicating ketone bodies as signaling molecules further supports roles for ketone bodies that transcend energy metabolism, indicating that additional experimentation is required to elucidate the receptors and mechanisms through which ketone bodies serve as extracellular signals.

Recently published findings also demonstrated that n-β-OHB inhibits class I histone deacetylases (HDACs), resulting in increased histone acetylation and thus increased expression of genes encoding mediators of resistance to oxidative stress (189). The molecular effects of n-β-OHB were observed at high-IC₅₀ concentrations (2.4–5.3 mM, depending on the HDAC isoform) and were recapitulated by caloric restriction, fasting, or AcAc. To determine if these effects were dependent on β-OHB metabolism, small interfering RNAs against BDH1/2 were transfected into cells, which were then treated with n-β-OHB. The authors observed that with up to 3 mM n-β-OHB, the effects on histone acetylation were preserved, leading to the conclusion that most of the influence of n-β-OHB on HDAC function was independent of n-β-OHB metabolism. However, residual BDH persisted in the small interfering RNAs-transfected cells, and the control experiment confirming diminished n-β-OHB oxidation in BDH knockdown cells was not performed. Thus it remains plausible that oxidation of n-β-OHB to AcAc by BDH1 could alter mitochondrial redox potential and that CoA transferase-dependent contribution of AcAc to the TCA cycle for terminal oxidation could alter cellular energy metabolism (see KETONE BODY OXIDATION and Fig. 5), each of which may contribute to the observed effects on histone acetylation. Nonetheless, these observations reveal exciting roles for ketone body metabolism in a multitude of disease states, which will benefit from the development and application of both pharmacological and genetic tools.

**Ketone Body Metabolism, the Microbiota, and Cardiovascular Biology and Disease**

A therapeutically tractable interface between ketone body metabolism and cardiovascular disease is offered through the dynamic community of microorganisms living on our cutaneous and mucosal surfaces, the microbiota. This ecosystem of ~100 trillion cells, 10 times the total number of our own cells, comprises a ~1 kg organ that harbors a massive aggregate of genomes (the microbiome) encoding millions of genes, 100 times that of the human genome, that influence the human metabolome and coordinate immune function [recently reviewed in (13, 25, 69, 85, 87, 92a, 92b, 148, 211)]. The data supporting causal links between the microbiota and host cardiovascular disease in both rodent models and humans are compelling. Gut microbial ecology undergoes marked transformation in obese and type-2 diabetic humans and rodents, and the dynamic energy-harvesting capacity of the gut microbial community plays a significant contributing role to energy homeostasis in the host (14, 15, 34–36, 120, 121, 141, 160,
214–216, 232, 241). Acting in synergy, the host’s genome and microbiome coordinate supraoorganismal metabolic processes, creating a suite of metabolites that influence a panopoly of pathophysiological processes including NAFLD, hypertension, insulin resistance, and atherosclerosis (38, 48, 86, 224). Furthermore, the ability of indigenous microbial communities to coordinate the functions of the host’s immune system is likely a key influence over the development of cardiovascular disease [reviewed in (87)], including the development of type-1 diabetes (227). Together, comparative genomics, metabolomics, and gnotobiotic studies (Greek roots: gnosis, knowledge; bios, life; a technology that allows cultivation of experimental animal models in the presence of defined microbial communities) may usher an era that ultimately allows caregivers to administer individualized forms of therapy that configure the microbiota using probiotics and prebiotics to ameliorate cardiovascular disease.

Studies of gnotobiotic animals have revealed intriguing relationships among the microbiota, ketone bodies, and cardiovascular biology. Normalized to tibial length or body weight, hearts of germ-free mice, which are born and raised in sterile gnotobiotic isolators, are ~15% smaller than those of normally colonized mice, but myocardial mass increases within 2 wk of microbial colonization (42). In vivo and ex vivo functional parameters of hearts of germ-free mice are normal, but myocardial glucose oxidation rates are higher. A systematic assessment of substrate delivery and metabolism in germ-free mice revealed that while circulating ketone body concentrations were not significantly different under fed conditions, ketosis was blunted during nutrient deprivation in germ-free mice because of diminished hepatic ketogenesis (42). While reduced adiposity of germ-free mice (14) is a likely contributor to blunted ketogenesis during fasting, hepatic ketogenic machinery, including expression of Hmgs2, was curtailed in fasting germ-free mice. The reduction in cardiac size and alterations of systemic and myocardial metabolism were abrogated when germ-free mice were maintained on a ketogenic diet (42). It is also intriguing to consider the aforementioned cross talk between SCFAs and ketone bodies [(106); see Signaling Roles for Ketone Bodies, SCFAs are produced through the fermentative actions of glycan-digesting enzymes uniquely encoded by the microbiome, contributing to the host’s energy harvest and to signaling through GPR41 (88, 106). Thus the microbiome may supervise an expansive and integrated network in which ketone bodies influence cardiac metabolism, size, and physiology. Future studies using genetically modified gnotobiotic animals will permit further construction of the relationship between ketone metabolism and the microbiome in diseases of the heart and vasculature.

Diagnostic and Therapeutic Targets of Ketone Body Metabolism

The diversity of metabolic and signaling processes in which ketone body metabolism participates provides a range of diagnostic and therapeutic applications, only a subset of which is under active investigation.

Abnormalities of ketone body oxidation. CoA transferase-deficient humans typically present early in life with severe ketoacidosis that can result in vomiting, coma, and death if intravenous glucose and insulin therapy is not rapidly instituted (20, 150, 210). Neonatal mice with complete loss of terminal ketone body oxidation (SCOT-KO mice) invariably die within the first 48 h of life in a manner that mimics human sudden infant death syndrome (SIDS) (40). Furthermore, a recent observational study that performed metabolic autopsies on 255 SIDS patients found that three exhibited underlying disorders of ketone body metabolism (159). Therefore, a small subset of cases that ultimately receive a diagnosis of SIDS may actually be attributable to latent abnormalities of CoA transferase function. However, statewide neonatal screening protocols in the United States do not currently detect newborns with isolated disorders of ketone body oxidation, who exhibit hyperketonemia, but generally normal organic acid and acylcarnitine profiles (143). Improved screening methods, measuring both AcAc and β-OHB, may detect patients with latent deficiency of ketone body oxidation, further reduce SIDS incidence, and predict metabolic complications which may manifest later in life.

Dietary therapies: ketogenic diets, ketone esters, and odd-chain fatty acids. Ketogenic diets are actively used for weight loss and anticonvulsant therapy and are intensively studied as potential adjunctive therapy for brain cancers and neurodegenerative diseases including Parkinson’s and Alzheimer’s diseases (110, 139, 187, 236, 237). Additionally, limited studies raise the possibility that humans with NAFLD could benefit from low-carbohydrate diet therapy (32, 58). An experimental ketogenic diet has been used to mitigate a mitochondrial cardiomyopathy in mice (112). The full scope of cardiovascular diseases responsive to nutritional and/or pharmacological manipulation of ketone body metabolism and the associated metabolic mechanisms remain underexplored.

Ketogenic diets are often unpalatable, which leads to poor patient compliance. Additionally, such diets raise blood cholesterol and free fatty acids, increase the risk of nephrolithiasis, and cause constipation (23, 116, 201). Therefore, Veech and Clarke developed and tested ingestible ketone ester compounds, which can rapidly generate ketones exceeding 5 mM in rats and humans (37, 103, 195), while sparing the adverse consequences of high-fat diets. A small preliminary study indicates that oral ketone esters are safe in humans (37). Rodents fed these compounds acutely decrease food intake. Although these animals do not exhibit changes in body weight over an extended time period, they do become more insulin sensitive, possibly because of increased expression of uncoupling proteins in brain and brown adipose tissue (103, 195). Further studies will evaluate the efficacy of these compounds in the mitigation and prevention of metabolic, myocardial, and neurological diseases in rodent and human subjects.

In the last decade, studies analyzing the beneficial roles of the anaplerotic five-carbon (C5) ketone bodies and their precursor odd-chain fatty acids for the treatment of long-chain fatty acid oxidation (LCFAO) disorders have emerged (170, 171). Conventional management of LCFAO disorders consists of dietary therapy with the medium-chain fatty acid octanoate. Substitution of octanoate for the odd-chain fatty acid heptanoate (given as the triglyceride triheptanoin) further improves clinical outcomes, specifically by reducing the incidence of rhabdomyolysis and cardiomyopathy (170, 171). While only trace levels of C5 ketone bodies are normally found in human body fluids (114, 202), ingestion of odd-chain fatty acids promotes hepatic C5-ketogenesis: β-oxidation of odd-chain fatty acids yields propionyl-CoA, an anaplerotic substrate,
which, in the liver, can also be packaged into C5 ketone bodies (107). Oxidation of C5-ketone bodies in peripheral tissues occurs through CoA transferase (197), regenerating propionyl-CoA. However, as oxidation of heptanoate within myocytes locally generates anaplerotic propionyl-CoA, direct evidence linking C5-ketone body metabolism to improved clinical outcome in LCAO disorders currently remains lacking. Thus it may be useful to cross experimental models of tissue-specific CoA transferase deficiency to existing models of LCAO defects (41, 183) to establish the metabolic roles of C5-ketone body metabolism in myocyte oxidation. Future studies of the role of these three approaches—low-carbohydrate ketogenic diets, ketone ester-containing diets, or anaplerotic odd-chain fatty acid-containing diets in cardiomyopathy—remain to be systematically performed but are warranted (44, 117).

Modifying ketone body enzymatic pathways: targeting Hmgcs2 and CoA transferase. As both nutrients and hormonal factors affect ketogenic regulation, it is not surprising that ketone body metabolism is dynamically regulated in mouse models of diet-induced obesity and in humans with metabolic syndrome. Hepatic ketogenesis becomes suppressed at later stages of the evolution of hyperinsulinemic obesity (18, 21, 179, 190, 193, 200). Because transgene-mediated Hmgcs2 overexpression within hepatocytes enhances ketogenesis and simultaneously diminishes circulating free fatty acid concentrations in vivo (217), regulation of HMGCS2 activity (See regulation of ketogenic mediators) becomes a prospective therapeutic target. Diversion of hepatic fatty acyl chains into ketone bodies could diminish carbon that would otherwise require terminal oxidation, storage, or packaging into lipoproteins for secretion, prospectively mitigating hepatic steatosis and insulin resistance.

Whole body ketone body turnover is noninvasively quantifiable in humans, as is tissue-level oxidative flux of ketone bodies via positron emission tomography using 11C-labeled ketone body tracers, which have been used for studies in brain, but not yet in other organs (24, 173). Additionally, ketone body metabolism and the mitochondrial enzyme CoA transferase are amenable to a number of nutritional and, perhaps ultimately, pharmacological therapies. Importantly, as described above, myocardial CoA transferase activity plays an important role in the regulation of substrate selection, mitochondrial ROS generation, and cardiac work and bioenergetic efficiency (Fig. 5). Therefore, analysis of animal models with sophisticated genetic manipulations of ketone body metabolism, coupled with pharmacological targeting strategies, may provide rationale for pharmacological therapies. Importantly, as described above, myocardial CoA transferase activity plays an important role in the regulation of substrate selection, mitochondrial ROS generation, and cardiac work and bioenergetic efficiency (Fig. 5). Therefore, analysis of animal models with sophisticated genetic manipulations of ketone body metabolism, coupled with pharmacological targeting strategies, may provide rationale for pharmacological therapies. Importantly, as described above, myocardial CoA transferase activity plays an important role in the regulation of substrate selection, mitochondrial ROS generation, and cardiac work and bioenergetic efficiency (Fig. 5).

Conclusions

Ketone body metabolism maintains bioenergetic homeostasis when dietary carbohydrates are limiting. Defects in either the synthetic or oxidative arms of ketone body metabolism result in disease pathogenesis in humans, and ketone body oxidation is required for maintenance of glycemia and survival in neonatal mice. Ketone bodies regulate mitochondrial metab-


METABOLISM OF KETONE BODIES IN HEALTH AND DISEASE


