Review Article
The Regulation and Characterization of Mitochondrial-Derived Methylmalonic Acid in Mitochondrial Dysfunction and Oxidative Stress: From Basic Research to Clinical Practice

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Methylmalonic acid (MMA) can act as a diagnosis of hereditary methylmalonic acidemia and assess the status of vitamin B12 deficiency [1]. And existing cross-sectional studies suggest that MMA is considered a potential risk factor for cardiovascular events. Although plasma levels of MMA exceed the universal reference interval with age, other factors may be involved in major organs. In epidemiological studies, MMA is related to the risk and poor prognosis of numerous chronic diseases. It is largely evidenced that MMA has a significant relationship to chronic diseases (e.g., aging, diabetes, obesity, and kidney dysfunction) [2, 3]. In particular, MMA is a mitochondrial toxin that can disrupt redox homeostasis by inhibiting electron transport complex II [4]. With the increase in concentrations of MMA, mitochondrial energy metabolism can be altered. Thus, disorders of mitochondrial energy metabolism can promote free radicals to be generated intracellularly, which damages mitochondrial DNA [5]. Furthermore, MMA is suggested to be a biomarker of oxidative stress conditions [6]. As a promising novel biomarker, MMA may optimize the risk stratification of diseases and provide a potential target for the study of disease mechanisms.

2. Circulating MMA in Clinical Diagnosis

2.1. MMA and Vitamin B12 Deficiency. The relationship between MMA and vitamin B12 (vitamin B12, Cb1) deficiency dates to the 1960s [7, 8]. Some medical institutions consider the MMA assay a more appropriate response to vitamin B12 activity than the conventional vitamin B12 test. Still, MMA testing has not been widely used, and there is no consensus on its clinical use. MMA is one of the indicators of Cb1 deficiency. Vitamin B12 deficiency refers to a
2.2. MMA and Hereditary Methylmalonic Acidemia. Methylmalonic acidemia is an autosomal recessive disorder. Abnormal metabolism of the MMA cofactor 5-deoxyadenosyl cobalamin or defective MCM in the mitochondria results in the accumulation of large amounts of MMA in major tissues due to impaired metabolism [18, 19]. During acute episodes of methylmalonic acidemia, MMA accumulates in the blood and cerebrospinal fluid of patients [20, 21].

The metabolic blockage causes considerable MMA to be accumulated mainly in a tissue. Overall, concentrations of MMA in blood and cerebrospinal fluid are approximately 2.5 mmol/L during acute metabolic attacks [20, 21]. The catabolic metabolites of MMA (e.g., propionic, methyl citric and hydroxy propionic acids, and propionyl glycine) can also be detected in the patient’s body fluids at the same time, but are not as easily detected as MMA due to its low concentration [22]. According to several available studies, MMA not only is a new risk factor for neurodegenerative diseases but also causes accumulation of organic acids and synergistic secondary toxic reactions [23], suggesting that acquired high levels of MMA can be a risk factor for adverse outcomes and not only an indicator of Cbl deficiency status.

Clinically, the major long-term complications consist of chronic renal failure [24] and neurological deficit [25], cardiomyopathy [26], pancreatitis [27], and recurrent infections attributed to leukopenia. Metabolic brain damage and
the progression of heart disease can be induced [28]. Congenital methylmalonic acidemia adversely affects multiple organs, which may suggest the acquired high levels of MMA as a risk factor for adverse outcomes.

2.3. MMA and Cardiovascular Disease. Despite the lack of large-scale epidemiological studies on circulating MMA and coronary artery disease, MMA has been described as a risk factor for CHD in the available cross-sectional studies. The most frequent complications in the MMA group with renal disease are dilated cardiomyopathy and arterial hypertension [29]. Moreover, in a cross-sectional clinical study enrolled in a total of 120 patients, MMA serum levels significantly increased with acute myocardial infarction (AMI) or acute heart failure (HF) compared with healthy controls. This result suggested that the differences were independent of demographic, medical, and other comorbidities [6]. There were also studies in which patients with chronic decompensated HF had higher MMA concentrations than patients with newly diagnosed HF in the control group. A retrospective study also demonstrated that MMA levels in patients suffering from oxidative stress (e.g., atrial fibrillation or arterial hypertension) were elevated [30], since high levels of MMA may be related to cobalamin deficiency and impaired mitochondrial function, in addition to long-lasting oxidative stress [31]. Accordingly, MMA critically impacts cardiac diseases, as the impairment of respiratory chain complex I have been identified in the cardiomyocytes, which inhibits carnitine uptake in the heart and causes cardiac insufficiency [32]. As suggested from a retrospective observational study in cblC-type methylmalonic aciduria, significant structural heart defects seem to be highly common probably due to MMA-influenced abnormal DNA methylation [33]. However, the mentioned studies have significant limitations, and convincing studies are rare on the association between MMA and the risk of cardiovascular events. Our recent cohort study in 23437 general adults demonstrated the significant relationship between circulating MMA and cardiovascular deaths [34]. Participants in MMA ≥ 250 nmol/L showed an increase in cardiovascular and all-cause mortality risk compared with those with MMA < 120 nmol/L. Thus, the accumulation of MMA is of critical importance for the evaluation of cardiovascular adverse events (risk and prognosis).

2.4. MMA and Other Chronic Diseases. It is known that plasma serum MMA levels increase above commonly accepted reference intervals with age [35]. Though vitamin B12 deficiency cannot be ruled out as a possibility contributing to the high level of MMA in the elderly, it may be related to other factors within the main organs [36]. Considerable data indicate a significant association between MMA and chronic diseases (e.g., renal dysfunction, neurodegenerative disease, obesity, and cancer) [2, 3].

To be specific, chronic renal failure is well-recognized complication of methylmalonic acidemia [24]. Moreover, the renal function acts as a vitally important determinant of MMA concentrations that is independent of vitamin B12 status in the patients [35]. The mitochondrial enrichment of epithelial cells of kidney tubules accounts for the transport functions and integrity [37]. The accumulation of MMA could break mitochondrial homeostasis and drive various degrees of tubular dysfunction [38]. As demonstrated by Horster and Hoffmann, accumulation of MMA on the tubular cell of the kidney could cause a disturbance of energy metabolism [39]. As opposed to the mentioned, kidney function might be restored by modulating mitochondrial function in mouse models [40]. In addition, tubular mitochondrial dysfunction refers to the key pathogenic mechanism of MMA-associated renal disease, while antioxidants were reported to reduce renal disease in some MMAAs [41]. MMA can induce DNA damage in the kidney, which may explain kidney failure in patients of methylmalonic aciduria. Besides being toxic to the kidney, it is also known as a neurotoxin [42].

It is suggested that MMA, as an endogenous toxic metabolite, could impair energy metabolism [43, 44] and elicit oxidative stress in brain of rats [45]. It is demonstrated in a study that intrastriatal administration of MMA induces a change of behavior (e.g., rotational behavior and striatal lesions in rats) [46, 47]. MMA could induce neuron apoptosis in different cell culture systems and cause secondary excitotoxicity together with the complex II inhibitors malonate (MA) [48, 49]. As suggested from a longitudinal study in the United Kingdom, MMA is a significant predictor of cognitive impairment. The rate of decline in cognitive performance of subjects was upregulated by 50% with an increased concentration of MMA 0.25-0.50 μmol/L in a longitudinal study in the United Kingdom [50]. The mentioned result might also explain the neurological deficit symptoms of methylmalonic acidemia which may have nonspecific symptoms (e.g., epilepsy or motor dysfunction of various degrees) [51–53]. These lesions typically occur during the acute phase of metabolic disorders. The neuroimaging of this inherited disease has shown basal ganglia degeneration and especially severe necrosis revealed by histopathology in the globus pallidus [4, 54]. By inhibiting the tricarboxylic acid cycle (TCA cycle) and mitochondrial respiratory chain, MMA causes one symptom of bioenergetic stroke after toxic metabolites accumulate [55].

According to one recent report, MMA could predict severe obesity together with homocysteine [56]. It is well known that insulin resistance is one of the key pathophysiological processes involved in obesity and diabetes mellitus. Some potential reasons might explain the Cb1 deficiency, and MMA could influence insulin resistance under the folate presence [57]. As a coenzyme, Cb1 could impair the activation efficiency of MMA and induce the expression of SIRT1 and lipogenesis subsequently [58, 59]. Studies have demonstrated that elevated MMA levels in diabetes patients showed more severe peripheral neuropathy [60, 61]. Similarly, our recent prospective cohort study in type 2 diabetes patients also found that MMA accumulation was positively associated with increased mortality risk [62]. Obesity and diabetes may play a critical biological role in the metabolic process of MMA in the setting of diseases.

Furthermore, MMA is a mediator of tumor progression by inducing aggressive features in cancer cells. As reported in a recent study, MMA is sufficient to stimulate the
progression and aggressiveness of the tumor by inducing SOX4 by activating autocrine TGFβ signaling. The accumulation of MMA in the aged host could represent antagonistic properties of cancer cells [36]. In brief, the association between MMA, an underestimated biomarker, and disease requires in-depth study and verification. As a systematically increased aging-induced metabolite [36], MMA could induce aggressive effects of aging in cancer cells and contribute to tumor progression and aggressiveness [63]. The pathophysiological mechanism related to MMA in the disease of cancer is not only mitochondrial dysfunction, impairment of tricarboxylic acid cycle, and oxidative stress mentioned before but also the effects of oncometabolites [36]. Several recent studies demonstrated that MMA could induce a proaggressive EMT-like phenotype with a decline in E-cadherin and a concurrent increase in fibronectin and vimentin. MMA is sufficient to endow cancer cells with migratory and invasive capacity by inducing SOX4 expression by activating autocrine TGFβ signaling [35]. Therefore, MMA is indicated to be a potential therapeutic target in carcinoma treatment. In brief, the association between MMA, an underestimated biomarker, and disease requires in-depth study and verification.

### 3. Pathophysiology of MMA

#### 3.1. Mitochondrial Homeostasis

Mitochondria are organelles with the energy of nutrients converted into adenosine triphosphate (ATP), which supports cellular metabolism and functions [64]. Thus, mitochondrial dysfunctions and homeostasis imbalances could result in crushing destruction to many cells, leading to a wide spectrum of diseases [65]. Accumulation of MMA due to mitochondrial dysfunction is recognized as the most common form of methylmalonic acidemia attributed to the mitochondrial enzyme MUT mutation and the lack of synthesis or transport of mitochondrial proteins [66, 67]. As a matter of fact, the metabolic pathway by which MMA enters the TCA cycle via propionyl-CoA and methylmalonyl-CoA produces large amounts of total ATP production under normal conditions [68]. In tissues and biological fluids, high levels of MMA might impair the energy metabolism of mitochondria. Disorders of mitochondrial energy metabolism consequently induce the elevation of intracellular free radical generation [5], leading to damage to mitochondrial DNA [69]. MMA is considered a mitochondrial toxin that interferes with redox equilibrium by inhibiting the electron transport complex II [4] and multiprotein in the TCA cycle, due to the exhibiting structural similarity with the respiratory chain complex II inhibitor MA [67]. Antagonists of ionotropic glutamate receptors and antioxidants prevented striatal lesions and cell injury induced by methyl methacrylate in rats [66]. Furthermore, one study suggests that MMA plays a role in inhibiting the transport of mitochondrial malate and succinate by a dicarboxylate carrier. Accordingly, one experiment demonstrated that MMA does not inhibit the consumption of succinate-supported oxygen in isolated mitochondria. Inhibition of the mitochondrial dicarboxylate carrier induced by methylmalonyl-CoA probably results in decreased gluconeogenesis and increased glycine in the kidneys and liver, and these could explain the disorders of failure to thrive and the delayed development in later childhood of methylmalonic acidurias [68, 70]. In the brain, the mitochondrial dicarboxylate carriers that could transport through the mitochondrial inner membrane are probably associated with de novo glutamate synthesis intermediates. Since pyruvate carboxylation is considered to occur only in astrocytes rather than neurons, high levels of MMA can compromise neuronal energy metabolism and glutamatergic neurotransmission [71]. In addition, MMA also inhibits the transport of glutathione in mitochondria via a dicarboxylate carrier at the same time inhibiting the mitochondrial dicarboxylate carrier, resulting in redox imbalance and mitochondrial antioxidant defense exhaustion [72]. As suggested by a number of studies, MMA leads to lipid peroxidation in cerebral tissues, which might explain the symptoms of neurologic deficit in methylmalonic acidemia [73–75].

Notably, MMA also shows a relationship to mitochondrial function and homeostasis. As discovered recently by Luciani et al., the MUT deficiency in methylmalonic acidemia might inhibit the autophagy of damaged mitochondria under stress-induced conditions by influencing Parkin’s E3 ubiquitin ligase. Thus, damaged mitochondria may accumulate and generate epithelial stress and tissue injury [76]. MMA also induces mitochondrial damage via other mechanisms. In one study, the MMA metabolite was reported to suppress the activity of the citrate cycle rate-limiting enzyme, α-ketovalerate dehydrogenase complex (αKGDH), by competing with α-ketovalerate. The αKGDH activity was measured to decrease by 73% and 60% at high concentrations of MMA (10 mm) with 0.1 and 0.25 mm α-ketopentanoic acid, respectively. The mentioned effects will lead to a decrease in the number of intermediates in the TCA cycle, which can apparently damage the organs enriched with mitochondria by suppressing oxidative metabolism [28]. It has also been argued that MMA does not directly impact the mitochondrial respiratory chain. In addition, MMA-induced damage may be driven by metabolites derived from propionyl-CoA and its substitutes, including propionic, malonic, 2-methylcitric, and 3-hydroxypropionic acids, causing a synergistic inhibition of the TCA cycle and respiratory chain. As demonstrated by one study, the neuronal damage induced by MMA involves the intracellular accumulation of MA and 2-methylcitrate metabolite in rat embryos, thereby inhibiting the TAC in multiple pathways [55].

Mitochondria-rich organs in the body (e.g., heart, kidney, and brain) may also be affected by MMA to affect various functions [23]. It has been demonstrated in many studies that impaired mitochondrial function is closely related to cardiac ischemia-reperfusion injury [77, 78] and that improved mitochondrial quality control can reduce cardiac microvascular ischemia-reperfusion injury [79, 80]. As suggested by Wang et al. [81], MMA accumulation will result in significant trapping of mitochondrial coenzyme A in cardiomyocytes, promoting increased myocardial oxygen consumption and impeding normal cardiac productivity. Thus, circulating MMA is associated with heart failure and cardiac hypertrophy. On the other hand, MMA-induced
mitochondrial damage also plays a pathogenic role in ischemic acute kidney injury (AKI) by disrupting mitochondrial mass and activating mitochondrial apoptosis [82, 83]. However, few relevant in vivo studies have been conducted on the biological function of methyl methacrylate in cardiovascular disorders, and the potential clinical significance of MMA remains unclear.

3.2. Oxidative Stress. MMA, as a promising novel biomarker, may represent the condition of oxidative stress [6]. Oxidative stress is identified as an important contributor to physiopathology of several diseases including cancer [84] and metabolic diseases [85]. There have been numerous studies based on animal models, and patients report the existence of oxidative stress within methylmalonic acidemia [86–88]. One experiment demonstrated that primary neuronal cultures and mouse models have suffered by significant toxic effects by consecutive intrastratal administration [66]. And another laboratory reported that injecting MMA into the encephalocoele in animal models could interfere with redox homeostasis [88]. The source of oxidative stress is usually multifactorial and seldom attributable to a single mechanism [89]. These factors include the accumulation of toxic metabolites in metabolic diseases and the production of reactive oxygen species (ROS) as well as reactive nitrogen species (RNS) related to pathogenesis of other diseases. In methylmalonic acidemia, the primary reason for oxidative stress is considered to be the generation of mitochondrial ROS induced by electron transport chain (ETC) dysfunction [73]. The results of some studies have suggested that inhibiting the enzymatic activity of ETC due to the accumulation of malonate and methylcitrate rather than MMA itself [55]. However, the results from other studies indicate that MMA tends to inhibit ETC activity [90–92]. Accordingly, synergistic inhibition by MMA, methylcitrate, and malonate might be responsible for the ETC dysfunction associated with methylmalonic acidemia [21].

Oxidative stress inevitably results in oxidative damage of proteins and lipids as well as a decrease in the antioxidant defense in vivo, while succinic acid, antagonists of ionotropic glutamate receptors, and antioxidants were prone to effectively prevent injury induced by MMA owing to their antioxidant [66]. There is extensive evidence that the antioxidant status of cells has an impact on methylmalonic acidemia. However, few studies have suggested antioxidant as an adjunct therapy to the methylmalonic acidemia treatment regimen. According to an animal model of methylmalonic acidemia, the combination of CoQ10 and vitamin E improves the glomerular filtration rate [41]. Furthermore, Salmi et al. showed that patients with methylmalonic acidemia have a deficiency of glutathione and have an overproduction of Oxic-nitrile (NO) [93]. There is also evidence that protein damage is mediated by NO, since seizures induced by MMA decline significantly in endothelial nitric oxide synthase knockout mice compared to normal controls [68, 94]. Thus, the increasing dose of MMA is also suggested to be able to reduce immune defenses. These findings may have the potential to offer a novel biomarker for disease prevention and treatment.

MMA along with Hcy serum levels are significantly elevated at the beginning of vitamin B12 deficiency, so both can be adopted to detect the vitamin B12 deficiency condition. Under normal conditions, MMA participates in the TCA cycle that requires Cbl in the formation of adenosylcobalamin (AdoCbl) to be a cofactor of this chemical reaction [95]. Additionally, Cbl is involved in converting Hcy into methionine and leads to methyl group generation through the tetrahydrofolate cycle [96]. Hcy has been extensively reported as a critical risk factor for the cardiovascular system and neurodegenerative diseases [97, 98]. Since existing studies suggested a positive relationship between Hcy and the severity of coronary heart disease, it plays a predictive role for guiding the disease assessment of coronary heart disease [16, 99]. Moreover, several studies reported that Hcy can activate nicotinamide adenine dinucleotide phosphate (NADPH) oxidase and generate massive production of ROS that causes damage to vascular endothelial cells and promotes lipid peroxidation and oxidation of low-density lipoproteins. Hcy could enhance oxidative activity and aggravate cellular oxidative stress injury [100, 101], thereby causing vascular damage and progression of atherosclerotic plaque to be aggravated. Though cell damage for B12 deficiency is partially attributed to accumulation of Hcy, the concentration of circulating MMA has more sensitivity and specificity to indicate the state of Cbl condition than for Hcy [8]. The site of Hcy metabolism is extensively cytoplasmic. However, MMA metabolism is directly determined by Cbl and mitochondrial MCM activity; it is not dependent of folic acid and vitamin B6 [17]. Figure 1 lists the relationship between MMA, Hcy, and Cbl. Thus, MMA may reflect oxidative stress-related pathological impairment than high homocysteine as a biomarker.

3.3. Neuronal Cell Apoptosis. Neuronal cell apoptosis may be one of the mechanisms related to the progression of MMA-induced disease. In a number of studies, MMA causes a decreased intracellular ATP/ADP ratio and depolarization of the plasma membrane in cultured neurons, eventually resulting in necrotic and apoptotic cell death [102]. Besides, high levels of MMA probably cause the interference of MAPK and p53 signaling pathways to decrease neuron viability and increase cell apoptosis [103]. Ionotropic glutamate antagonists’ receptors could be used to prevent MMA-induced cell death, which suggests that excitotoxic mechanisms are involved in this pathology of cellular damage [66]. Okun et al. reported that MMA loading in rats led to intracellular accumulation of 2-methylcitric acid (2-MCA) and MA induce cellular damage [66]. For this reason, the induction of MMA may be indirect, which involves MMA and MA acting synergistically to mediate the impairment of cellular energy metabolism [55]. The release of glutamate, which is mainly expressed by the overactivation of N-methyl-D-aspartate (NMDA) receptors and induces an influx of Ca2+ and Na+ subsequently, could relate to the process of central nervous system [104]. Under excitotoxic conditions, high doses of stressors, such as the mitochondrial matrix Ca2+ and ROS, may cause rupture of the outer mitochondrial membrane and mitochondrial dysfunction.
through mitochondrial permeability transition (MPT) [105]. As suggested by Kowaltowski et al., MPT was involved in MMA-mediated neurotoxicity, since cyclosporin A (an MPT inhibitor) has been found to prevent MMA-induced cell death [102]. Furthermore, increasing evidences have suggested that the pathogenesis of MMA is related to the dysfunction of intracellular trafficking [106]. These can explain the symptoms of neurologic deficiency in methylmalonic acidemia.

4. Summary

It is acknowledged that MMA is capable of monitoring early vitamin B12 deficiency and diagnosing genetic metabolic disorders clinically. Moreover, the metabolism of MMA depends on mitochondrial status, and MMA also affects the energy homeostasis of mitochondria. MMA metabolism also affects the deterioration of diseases characterized by mitochondrial impairment, including cardiovascular disease, chronic renal failure, neurodegenerative disease, overweight, and cancer. Accordingly, MMA might show a substantial potential to improve disease management and risk stratification by complying with its distinctive biological role. There has been insufficient evidence on the relationship between MMA and a series of diseases and its prognostic value. A considerable number of studies face obvious limitations (e.g., small sample size, insufficient sensitivity of detection methods, and inconsistent diagnostic criteria). MMA in the pathogenesis and clinical risk stratification in mitochondrial-related disease warrants further elucidation.

Conflicts of Interest

There are no conflicts of interest to declare.

Authors’ Contributions

YL and SW drafted the manuscript. XZ, HC, and JL were involved in revising it critically for important intellectual content. BY and SF critically reviewed the manuscript drafts. All authors read and approved the final manuscript. Yige Liu and Shanjie Wang contributed equally to this work.

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