Magnesium, Oxidative Stress, Inflammation, and Cardiovascular Disease

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Abstract: Hypomagnesemia is commonly observed in heart failure, diabetes mellitus, hypertension, and cardiovascular diseases. Low serum magnesium (Mg) is a predictor for cardiovascular and all-cause mortality and treating Mg deficiency may help prevent cardiovascular disease. In this review, we discuss the possible mechanisms by which Mg deficiency plays detrimental roles in cardiovascular diseases and review the results of clinical trials of Mg supplementation for heart failure, arrhythmias and other cardiovascular diseases.

Keywords: antioxidant; inflammation; mitochondria; arrhythmias; heart failure with preserved ejection fraction; heart failure with reduced ejection fraction

1. Introduction

Cardiovascular disease (CVD) refers to a number of conditions such as heart failure (HF), arrhythmia, atherosclerosis, and stroke. Two types of HF can occur when the heart fails to pump enough blood to support other organs in the body. One is systolic HF as a result of poor contractile function, also known as HF with reduced ejection fraction (HFrEF). The other is diastolic HF as a result of reduced heart filling in diastole, also known as HF with preserved ejection fraction (HFpEF). Recently, our group has reported that magnesium (Mg) supplementation demonstrates significant protective effects on cardiac diastolic function, providing a new possible therapy for HFpEF [1]. Therefore, review of the effects of Mg on CVD seems timely.

CVDs are the number one cause of death globally. About 650,000 Americans die from CVD each year. In the United States, about 6.5 million adults have HF, compared with ~5.7 million between 2009–2012 [2]. HF caused one in eight deaths in 2018 [3], in spite of the remarkable advances in therapeutic treatments and prevention. Certain medical conditions increase risk for HF, such as diabetes mellitus (DM), obesity, and hypertension. Primary prevention of HF starts with a healthy lifestyle by following the Life’s Simple 7 goals (diet, smoking, body mass index, physical activity, cholesterol, glucose, and blood pressure) [2]. Treatment of systolic HF can involve medicines, mechanical support, arrhythmia prevention, and transplantation. There are no specific therapies for diastolic HF.

Hypomagnesemia is commonly observed in HF [4–10], DM [1,11], CVD [6,12–14], hypertension [15,16], and stroke [17,18], and the presence of hypomagnesemia in these conditions has been nicely reviewed [6–8,19,20]. Dietary Mg intake is inversely correlated with the occurrence of metabolic diseases [21,22], such as DM (type 1, 2 and 3) [23–26] and hypertension [27,28]. Low serum Mg is a predictor for cardiovascular and all-cause mortality [13], and hypomagnesemia is associated with unstable cardiac electrical repolarization and contributes to sudden death in HF [29].

Increased consumption of processed food, filtered/deionized drinking water, and crops grown in Mg-deficient soil has led to a significant decline in Mg intake [30–32]. The majority of North Americans...
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consume 185–235 mg/day Mg, compared with 450–485 mg/day in ~1900 [33]. Studies have reported that ~50% of the US population, especially the elderly, consume less than the estimated average requirement for Mg [33–35] and ~23% of US adults have hypomagnesemia (serum Mg <0.7 mM or 1.7 mg/dL) [36,37]. Dietary Mg intake <2.3 mg/kg body weight is related to increased risk of HF hospitalization in black adults [38]. Moreover, chronic diseases and medications such as diuretics, thiazides, cytotoxic drugs, digoxin, aminoglycosides, and steroids can further increase Mg loss, decrease Mg absorption, and cause hypomagnesemia.

Mg supplementation has shown significant therapeutic effects in HF and CVD (see reviews [4,6,12,14,39,40]). It has also been shown to improve arrhythmias including torsades de pointes (TdP, polymorphic ventricular tachycardia with marked QT prolongation on the electrocardiogram) [41,42], atrial fibrillation (AF) [43–45], ventricular arrhythmias (VA) [46,47], and arrhythmias in acute myocardial infarction (MI) [48].

Mg is the fourth most abundant mineral and the second most abundant intracellular divalent cation. It acts as a cofactor for more than 300 metabolic reactions, such as ATP production [49–51], protein synthesis, DNA/RNA synthesis [52], and mitochondrial function maintenance [53,54]. As a natural calcium (Ca) antagonist, Mg also participates in the regulation of Ca homeostasis. Mg has been reported to play critical roles in heart rhythm [55–58], muscle contraction [59–61], blood pressure [15,16,62], insulin/glucose metabolism [63,64], and bone integrity [65–67]. In this review, we will focus on the effects of Mg on CVD. First, we will discuss the mechanisms of how Mg deficiency affects metabolism including Mg deficiency-induced oxidative stress, inflammation, and insulin resistance. Then, we will review clinical and animal studies on how Mg supplementation has been used to improve HF, arrhythmias, metabolic syndromes, and vascular diseases. In the end, we will discuss the methods of Mg treatment including formulations, routes of delivery, dosing, and duration in different diseases. The limitations of Mg as a treatment will also be discussed.

2. Mg Deficiency Induces Metabolic Derangements

Normal serum Mg concentration is between 0.7–1.0 mM (1.7–2.4 mg/dL) [68,69]. Hypomagnesemia is defined as serum Mg < 0.7 mM. Mild to moderate hypomagnesemia occurs when serum Mg is between 0.5–0.69 mM (1.20–1.88 mg/dL) and severe hypomagnesemia is a serum Mg < 0.5 mM. For cellular Mg, about half of the total Mg is bound to nucleotide triphosphates, primarily ATP (MgATP), and ~20% is in the cytoplasm and the lumen of organelles. In cardiomyocytes, free ionized intracellular Mg (Mg$_i$) concentration ([Mg]$_i$) is tightly maintained in the range of 0.8–1.0 mM [70,71].

Several Mg transporters and channels have been identified in the heart and regulate Mg influx and efflux across the plasma membrane, including the transient receptor potential melastatin 7 channel (TRPM7, mainly for Mg influx), solute carrier family 41 A1 (SLC41A1, for Mg efflux), Mg transporter 1 (MagT1, for Mg influx), and the ancient conserved domain protein 2 (ACDP2, or cyclin and CBS domain divalent metal cation transport mediator 2: CNNM2, for Mg influx) [72–74]. Mitochondrial RNA splicing 2 protein (MRS2, for Mg influx into mitochondria) [75] and solute carrier family 41 A3 (SLC41A3) (for Mg efflux from mitochondria) [76] appear to transport Mg across the mitochondrial membranes. Dysregulation of these Mg transporters and channels is caused by and contributes to disturbed Mg homeostasis [72,77,78]. For example, gene upregulation of SLC41A1 and MagT1 have been reported with Mg deficiency [72,79,80]. Under hypomagnesemia, low serum (i.e., extracellular) Mg levels can activate Mg transporters such as TRPM7 and SLC41A1 to induce Mg efflux from cells to increase serum Mg levels. This would be expected to decrease [Mg]$_i$ and affect Mg/MgATP-dependent cellular signaling and functions. A decreased [Mg]$_i$ could trigger Mg stores such as mitochondria [51,81] to release Mg through SLC41A3 [76]. Decreased mitochondrial Mg levels ([Mg]$_{m}$) could affect further Mg/MgATP-associated mitochondrial signaling and functions, which may explain the mitochondrial reactive oxygen species (ROS) overproduction and decreased ATP levels observed in Mg deficient mice [1,82].
CVD patients often have hypomagnesemia, resulting from low Mg intake and increased Mg loss. In a study of congestive HF, 38% of patients have hypomagnesemia, and 72% of patients have excessive Mg loss [10]. This could be attributed to electrolyte disturbances induced by the excessive activation of the renin-angiotensin-aldosterone system (RAAS) [5,83–85] and by long-term use of medications such as diuretics and digoxin in chronic HF [9,86–88]. Hypomagnesemia may increase the risk of CVD by suppressing mitochondrial function with increased oxidative stress and decreased energy production [1,50,89,90], modulating cardiac ion channels to cause myocardium electrical remodeling [8,47,56], and disturbing Ca homeostasis and cardiac contraction [1,8,49,82,91]. Mg deficiency also induces sustained inflammation [35,92–95] and interrupts insulin signaling [26,64,96,97], both of which contribute to the development and progression of CVD. In this section, we will discuss possible mechanisms of Mg deficiency-associated CVD, including Mg deficiency-induced oxidative stress, inflammation, and insulin resistance.

2.1. Mg Deficiency and Oxidative Stress

Oxidative stress is common in CVD [98–101]. Mg deficiency has been associated with oxidative stress in HF, DM, obesity, and hypertension [1,4,5,9,86,102,103] (for reviews [77,104,105]). Obesity, which is a strong risk factor for CVD, is linked with decreased serum Mg and increased malondialdehyde, a marker for oxidative stress resulting from lipid peroxidation [106,107]. ROS in the heart are mainly produced by mitochondria, nicotinamide adenine dinucleotide phosphate (NADPH) oxidases, uncoupled nitric oxide synthase (NOS), and xanthine oxidase. Mitochondria occupy ~30% of the cardiomyocyte volume and are not only the major site for energy and ROS production in the heart but also are major Mg stores (with three- to five-fold higher concentrations of [Mg]$\text{_{m}}$ compared with [Mg]$\text{_{i}}$) [75,108]. Recently, our group has reported that Mg deficiency in diabetic mice increases mitochondrial oxidative stress and contributes to cardiac diastolic dysfunction [1]. Both mitoTEMPO (a mitochondria-targeted ROS scavenger, (2-(2,2,6,6-Tetramethylpiperidin-1-oxyl-4-ylamino)-2-oxoethyl)triphenylphosphonium chloride) treatment [109] and Mg supplementation [1] reverse diastolic function by reducing the oxidative stress and subsequent S-glutathionylation of cardiac myosin binding protein C. In a low-Mg diet-induced Mg deficient mouse model, mitochondrial oxidative stress is also detected and contributes to cardiac diastolic dysfunction [82]. Both mitoTEMPO treatment and Mg repletion can suppress mitochondrial ROS overproduction and reverse diastolic dysfunction [82]. Therefore, Mg can act as a mitochondrial antioxidant.

Presented in Figure 1 are the effects of Mg deficiency on mitochondria. Intracellular Mg deficiency has been shown to disrupt mitochondrial function by altering coupled respiration [50,89,90], increasing mitochondrial ROS production [1,82,110], suppressing the antioxidant defense system (such as superoxide dismutase (SOD), catalase, glutathione, and Parkinsonism associated deglycase PARK7/DJ-1) [105,111–114], inducing mitochondrial Ca overload via the mitochondrial Ca uniporter [1,54,115], attenuating pro-survival signaling [116–118], and promoting opening of mitochondrial ATP-sensitive potassium (K) channel [119], inner membrane anion channel [120] and mitochondrial permeability transition pore (PTP) [121]. The effects result in depolarization of the mitochondrial membrane potential ($\Delta$Ψ$_{m}$) [54]. Mg repletion, on the other hand, has been found to improve mitochondrial function by suppressing mitochondrial ROS overproduction [1,82], inhibiting mitochondrial PTP opening and cytochrome C release [122–124], preserving $\Delta$Ψ$_{m}$ [53,125], diminishing mitochondrial Ca accumulation [1,49,126,127], enhancing protein expression of the anti-apoptotic B-cell lymphoma 2 (Bcl-2) family and concomitantly decreasing pro-apoptotic protein expression such as Bcl-2-associated X protein (Bax) [118,125], decreasing apoptosis by suppressing activation of hypoxia-inducible factor 1α (HIF-1α) and p38 mitogen activated protein kinase/c-Jun N-terminal kinase (p38/JNK) signaling [125], and downregulating autophagy [127].
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Figure 1. Intracellular magnesium (Mg) deficiency (Mg↓) induces mitochondrial oxidative stress and dysfunction. Mg deficiency downregulates the electron transport chain (ETC) and increases reactive oxygen species (ROS) production. Mg deficiency also suppresses the antioxidant defense system with decreased protein levels such as manganese SOD (MnSOD) and catalase. Mg deficiency downregulates ATP synthase (F0F1) and decreases ATP production. Intracellular Mg deficiency inhibits Mg transport into mitochondria via mitochondrial RNA splicing 2 protein (MRS2) and promotes mitochondrial Mg (Mgm) efflux via SLC41A3. Mg deficiency promotes apoptosis by increasing cytochrome C release via Bax and voltage dependent anion channel (VDAC), decreasing anti-apoptotic proteins such as the Bcl-2 family, and increasing pro-apoptotic proteins such as HIF-1α and p38/JNK. Mg deficiency induces mitochondrial membrane (ΔΨm) depolarization by increasing the permeability of mitochondrial permeability transition pore (PTP), ATP-sensitive K channel (KATP), and inner membrane anion channel (IMAC). Mg deficiency increases mitochondrial Ca (Ca2+) via the mitochondrial Ca uniporter (MCU). Ca leak from mitochondria via VDAC is also increased with Mg deficiency.

There are two identified mitochondrial Mg transporters, MRS2 and SLC41A3 (Figures 1 and 2). MRS2 mainly transports Mg into mitochondria, although Mg efflux via MRS2 is also observed when ΔΨm is depolarized [75]. MRS2 knockdown has been reported to reduce Mg uptake capacity, decrease [Mg]m, induce loss of complex-I of the electron transport chain, reduce cellular and nuclear ATP levels, depolarize ΔΨm, make cells vulnerable to oxidative stress and apoptotic stimulation, and increase cell death (Figure 2) [51,128]. MRS2 upregulation, on the other hand, exhibits protective effects against apoptosis. Overexpression of MRS2 leads to increased total cellular Mg levels and enhanced resistance to apoptotic inducing drugs [129], probably by inhibiting Bax-induced cytochrome C release from mitochondria [118]. These results suggest the vital roles of MRS2 to maintain [Mg]m and mitochondrial function.

SLC41A3 is highly expressed in the heart, central nervous system, small intestine, and lung and is responsible for Mg efflux from mitochondria [76,130]. Its exact location on either inner or outer membrane of mitochondria has not been determined yet. In Figure 2, SLC41A3 is depicted on the outer mitochondrial membrane for convenience. Under Mg deficiency, the mRNA and protein levels of SLC41A3 are increased, probably as a compensation so that the mitochondria can release Mg to cope with Mg deficiency [131]. SLC41A3 overexpression leads to significantly increased Mg efflux from mitochondrial in a sodium (Na) dependent and temperature sensitive manner [76]. Cells with SLC41A3 overexpression demonstrate lower cellular ATP levels, suggesting that a decreased [Mg]m suppresses mitochondrial ATP production [76]. SLC41A3 knockout in mice results in hypomagnesemia under a normal diet, indicating that mitochondria play important roles in Mg homeostasis. Mechanisms for hypomagnesemia include increased intestinal mRNA expression of TRPM6, TRPM7, and SLC41A1.
and therefore increased Mg extrusion [130]. Under Mg deficiency, the electro-chemical balance of Mg across the mitochondrial membranes will be altered. MRS2 and SLC41A3 can respond to this change by decreasing mitochondrial Mg influx and increasing mitochondrial Mg efflux, respectively. This would increase \([\text{Mg}]_i\) and decrease \([\text{Mg}]_m\). Fluctuation of \([\text{Mg}]_m\) could lead to changes in Mg-dependent signaling cascades to modulate the mitochondrial electron transport chain, mitochondrial ATP and ROS production, and other mitochondrial functions.

Figure 2. Mitochondrial Mg transporters MRS2 and SLC41A3 mediate mitochondrial and cellular changes by modulating mitochondrial Mg homeostasis. MRS2 and SLC41A3 carry out mitochondrial Mg (\(\text{Mg}_m\)) influx and efflux, respectively. Changes in MRS2 have shown to alter \(\text{Mg}_m\) and affect complex I, ATP production, mitochondrial membrane depolarization (\(\Delta \Psi_m\)), intracellular Mg (\(\text{Mg}_i\)), and cellular oxidative stress and apoptosis. Changes in SLC41A3 has been reported under hypomagnesemia and TRPM7 depletion. SLC41A3 impacts ATP production, expression of other Mg transporters such SLC41A1. The double-headed black arrows indicate that changes in Mg transporters can result in and from disturbed Mg homeostasis.

Aside from the association between Mg deficiency and mitochondrial oxidative stress, studies have reported correlations between Mg deficiency and other ROS sources. Mg deficiency is found to interfere with nitric oxide (NO) release from coronary endothelium [132]. Mg deficiency results in activation of endothelial and neuronal nitric oxide synthase as well as reduction of serum Mg and tissue glutathione in all chambers of the heart [111]. Mg deficiency also causes an increased activity in NADPH oxidase induced by decreased ATP [133].

2.2. Mg Deficiency and Inflammation

Mg deficiency has been linked to inflammation for decades [66,94,134–137], and this association has been reviewed many times [35,81,92,138,139]. Reduced Mg levels result in activation of inflammation, while elevated Mg levels suppress the inflammatory response. Table 1 lists some inflammatory biomarkers that have been reported to be elevated under Mg deficiency. For example, Mg levels in HF are inversely related to inflammatory markers C-reactive protein (CRP) and tumor necrosis factor-\(\alpha\) (TNF-\(\alpha\)) [81,140]. Decreased plasma Mg is associated with increased monocyte chemoattractant protein-1 (MCP-1) [141]. In animal studies, low-Mg diet increases interleukin-1 (IL-1), IL-6, TNF-\(\alpha\), IL-8, and MCP-1 [66,134,136,142–144]. Moderate Mg deficiency (~50% of requirement) is associated with increased TNF-\(\alpha\), oxidative stress, and increased risk to develop CVD [142,143]. Protein kinase C (PKC) activation induced by Mg deficiency seems to contribute to the synthesis and release of cytokines [145].
Table 1. Mg deficiency activates inflammation with increased inflammatory markers reported in clinical and animal studies.

| Increased Inflammatory Markers | Tested Targets                        | Reference |
|--------------------------------|---------------------------------------|-----------|
| CRP                            | Human HF                              | [140]     |
|                                | Postmenopausal women                  | [146]     |
| MCP-1                          | Human carotid artery stenosis         | [141]     |
|                                | Human dermal microvascular cells      | [144]     |
|                                | Mouse hearts                          | Our unpublished work |
| substance P                    | Rats                                  | [147,148] |
|                                | Rats with cardiomyopathic lesions     | [149]     |
|                                | Mouse megakaryocytes and lymphocytes  | [66]      |
| IL-1                           | Human umbilical vein endothelial cells| [136]     |
|                                | Mouse osteoclasts                     | [66]      |
|                                | Hamsters and rats with cardiac lesions| [134]     |
|                                | Rat bone cells                        | [148]     |
| IL-6                           | Postmenopausal women                  | [146]     |
|                                | Aging rats with CVD                   | [142,143] |
|                                | Hamsters and rats with cardiac lesions| [134]     |
| IL-8                           | Human dermal microvascular cells      | [144]     |
|                                | Hamsters and rats with cardiac lesions| [134]     |
| TNF-α                          | Type 2 diabetes patients              | [150]     |
|                                | Postmenopausal women                  | [146]     |
|                                | Aging rats with CVD                   | [142,143] |
|                                | Hamsters and rats with cardiac lesions| [134]     |
|                                | Rat bone cells                        | [148]     |
|                                | Mouse osteoclasts                     | [66]      |

Hypomagnesemia activates systemic inflammation in at least seven ways (Figure 3): (i) induction of cellular oxidative stress that promotes inflammation [92,151,152]; (ii) activation of the RAAS that leads to inflammation [84,92,153]; (iii) relief of Mg inhibition on Ca channel opening that induces abnormal Ca handling [151,154,155]; (iv) activation of phagocytic cells [134,154,156–158]; (v) relief of Mg antagonism on the N-methyl-D-aspartate (NMDA) that leads to over-activated NMDA receptors and release of substance P [95,138,151]; (vi) activation of nuclear factor-κB (NF-κB) signaling and upregulation of transcription of cytokines and pro-inflammatory genes such as IL-1α [136,159–161]; and (vii) reduction of anti-inflammatory mediators such as NO, lipoxins, resolvins, and protectins [103,162]. For instance, low-Mg diet significantly increases macrophage-derived cytokines (IL-1, IL-6, and TNF-α) in hamsters and rats [134]. A few days of experimental Mg deficiency in rats induces activation of leukocytes and macrophages and increases release of inflammatory cytokines [93]. Low Mg increases the oxidative activity of neutrophils, while high Mg significantly decreases oxyradical production in rats and human polymorphonuclear cells [163,164]. Phagocyte priming has been proposed as a mechanism of Mg deficiency-induced immuno-inflammatory processes [156,158].

Mg transporters TRPM7 and MagT1 have been shown to affect and be affected by inflammation. TRPM7 is highly expressed in leukocytes and many organs such as the heart and kidney. It is a rare channel-enzyme with a channel domain to transport multiple cations (including Mg and Ca) and a kinase domain to phosphorylate serine/threonine [165,166]. TRPM7 acts as a key transporter to regulate Mg homeostasis by mediating Mg influx [166–168]. Its channel function is inhibited by physiological [Mg]i and [Ca]i, but activated under Mg deficiency [166]. Its kinase function is elevated by Mg [169,170]. Transgenic mice with heterozygous deletion of TRPM7 kinase domain (TRPM7+/Δkinase) have cardiac hypertrophy, inflammation, fibrosis, decreased cardiac [Mg]i, elevated expression of cytokines (IL-6,
-10, -12, and TNF-α), increased cardiac collagen and fibronectin content, and increased infiltration of macrophages [171]. Inhibition of TRPM7 can abolish IL-4-induced macrophage proliferation and prevent macrophage polarization towards the anti-inflammatory M2 phenotype [172].

**Figure 3.** Mg deficiency induces inflammation through several signaling pathways. Abbreviation: NMDA, N-methyl-D-aspartate; RAAS, the renin-angiotensin-aldosterone system.

MagT1 conducts Mg influx and plays a key role in T cell-activated immune response [155,173]. MagT1 deficiency leads to the absence of a T cell antigen receptor-stimulated Mg flux and attenuation of T cell activation [155,173]. Matsuda-Lennikov et al. have reported that MagT1 mediates N-linked glycosylation and expression of immune-response genes in T cells [174].

### 2.3. Mg Deficiency and Insulin Resistance

Mg deficiency, insulin resistance, and oxidative stress often co-exist in patients with diabetes, obesity, and hypertension [64,175]. Mg is a second messenger for insulin signaling [20,155,176]. It affects glucose transportation across the membrane [177,178] and insulin secretion and release [179,180]. Mg can also affect insulin signaling through its regulation of redox balance and enzyme/kinase activities [1,96,181–183].

Oxidative stress induced by Mg deficiency is a major cause of disrupted insulin signaling [181]. Animal studies show that rats and mice fed with low Mg diet exhibit increased plasma glucose levels, insulin resistance, and oxidative stress [96] (and our unpublished results). Decreased serum Mg and increased oxidative stress are observed in diabetic mice, while Mg supplementation decreases blood glucose levels and improves glucose tolerance and mitochondrial oxidative stress [1]. Vitamin E, an antioxidant, has been shown to increase [Mg], levels and insulin sensitivity [184]. On the other hand, Mg, being a cofactor for adenine nucleotides, is critical for the activity of ATP-consuming enzymes and kinases (such as hexokinase, phosphofructokinase, aldolase, phosphoglycerate kinase, pyruvate kinases, protein kinase B also known as Akt2, and insulin-related substrate-1) that regulate glycolysis [64,185–187] and insulin action [182,183] (Figure 4). For instance, Mg supplementation upregulates Akt2 and insulin-related substrate-1 to decrease blood glucose levels and improve glucose tolerance [183]. In summary, Mg deficiency can cause downregulation of insulin signaling and elevated insulin resistance.
Figure 4. A possible signaling pathway linking intracellular Mg deficiency and insulin resistance in a positive feedback loop.

For decades, insulin has been reported to regulate Mg levels [63,176,188]. SLC41A1 transports Mg out of the cells. Insulin signaling has been shown to inhibit SLC41A1 expression and function [176,188]. Under insulin resistance, there is less inhibition of SLC41A1, which leads to elevated cytoplasmic Mg efflux, causing intracellular Mg deficiency [189]. Increased SLC41A1 has been observed in Mg deficiency [72,79]. Mg deficiency can in turn exacerbate oxidative stress and downregulation of enzymes and kinases involved in glycolysis and insulin signaling, forming a positive feedback. This possible signaling pathway is summarized in Figure 4. These signaling changes may explain why Mg deficiency is often observed in patients with DM [23–26]. Mg supplementation and insulin both have protective effects for [Mg], levels by inhibition of SLC41A1 expression [188,189].

3. Mg Supplementation as a Therapeutic Treatment for Cardiomyopathy

Chronic Mg deficiency contributes to the development and progression of HF, hypertension, atherosclerotic vascular disease, and metabolic diseases, while acute Mg deficiency has been shown to be associated with cardiac arrhythmia and neuromuscular hyper-excitability (such as pre-eclampsia and epilepsy) [190]. Recently, our group found that Mg deficiency independently induced systolic and diastolic dysfunction in mice with low Mg diet [82]. The underlined mechanisms include Mg deficiency-induced mitochondrial dysfunction, oxidative stress, oxidation of cardiac myosin binding protein C, dysregulation of Ca handling proteins such as the sarcoplasmic reticulum (SR) Ca-ATPase (SERCA) and the SR Ca release channel (i.e., the ryanodine receptor 2 (RyR2)), and suppression of Ca transient. Mg repletion reversed all the detrimental changes and restored heart function.

In this section, we will review the published clinical and experimental studies on how Mg supplementation improves HF, arrhythmias, metabolic syndromes, and vascular diseases (Figure 5).

3.1. Mg Supplementation Improves HF

In HF patients, hypomagnesemia is frequently observed (with a reported 7%–38% range), together with other electrolyte abnormalities such as hypocalcemia [5,9]. Maintaining a normal Mg level with Mg supplementation plays a protective role on HF survival and on all-cause mortality [191]. Gottlieb
et al. reported that HF patients with low Mg levels have a two-year survival rate of 42% vs. 61% for patients with normal Mg levels [39]. Mg deficiency-induced oxidative stress and inflammation contribute to the development and progression of HF [5,81,103,142], while Mg supplementation suppresses oxidative stress and inflammation and plays protective roles in HF [1,82,152].

3.2. Mg Supplementation Shows Protective Effects against Arrhythmias

Mg has been used to treat different types of arrhythmias for decades, such as AF [43–45,198,199], TdP [41,42,200], and VA [46,47,199] (for reviews [6,201–204]). Mg is also frequently used in patients undergoing cardiac and pulmonary surgery when lethal arrhythmias often occur [44,199,204–206]. For example, a meta-analysis of randomized controlled trials on 2069 patients demonstrates that prophylactic Mg administration (IV MgSO4 or MgCl2) reduces the risk of supraventricular tachycardia and VA after cardiac surgery [205].
Long-term mild and moderate hypomagnesemia is associated with AF [45]. A pilot trial supports oral Mg oxide supplementation for AF prevention [47]. IV MgSO₄ decreased the heart rate and converted to sinus rhythm patients with rapid AF [207]. Higher incidence of postoperative AF is observed in patients with lower plasma Mg levels [208]. IV Mg prevents AF after cardiac and thoracic surgery [199,204,205,209] and non-cardiac thoracic surgery [198]. IV Mg also helps control the ventricular rate in patients with AF as an adjunct to digoxin [203].

TdP is most commonly caused by medications such as QT prolonging drugs [42]. Other causes for TdP include congenital long QT syndrome, electrolyte imbalance (hypomagnesemia and/or hypokalemia), bradycardia, hypothyroidism, and cardiac diseases [210]. IV Mg has been effective in treating TdP patients and is recommended as the initial therapy of choice to treat TdP, regardless of serum Mg levels [41,201,210,211]. In a study of six patients with TdP but normal Mg levels [41], IV MgSO₄ infusion (50 mg/min for 2 h) eliminated the arrhythmia within 20–30 min in all cases. A single bolus of IV MgSO₄ (2 g) has been shown to abolish TdP within 1–5 min in nine of 12 TdP patients [200]. In studies of canine TdP models, Mg rapidly prevents triggered ventricular tachycardia (VT) and eliminates TdP, probably by inhibiting early afterdepolarizations, a cellular arrhythmic mechanism [212,213].

Mg has been shown to alleviate other VAs in many studies [46,47]. Oral MgCl₂ intake decreased the occurrence of ventricular premature complexes, couplets and non-sustained VT in HF patients [195]. Oral Mg intake together with potassium exhibits antiarrhythmic effects on patients with frequent VA [46]. IV MgSO₄ administration reduced the numbers and lasting time of ventricular ectopic beats, couplets, and episodes of nonsustained VT [10,196]. A meta-analysis on 22 studies with over 6000 patients shows that the rates of VA (VT and ventricular fibrillation (VF)) and the incidence of supraventricular tachycardia are significantly lower in patients receiving IV MgSO₄ (11.9% and 10.4%, respectively) than placebo (24.2% and 23.9%, respectively) [91]. Intraoperative Mg treatment is associated with reduced occurrence of postoperative arrhythmias including VT, junctional ectopic tachycardia, and atrioventricular block [199].

Mg shows protective effects on arrhythmias in MI. Low serum Mg is associated with increased risk and mortality of acute MI [191,214], and Mg deficiency, in turn, aggravates MI by inducing mitochondrial dysfunction and increasing oxidative stress-induced ischemic injury [96,110,215,216]. In Mg-treated patients, there is ~20% reduction in infarct size [217], ~24%–50% decreased mortality [20,48,192,214,218,219], decreased rates of arrhythmias after infarction [48,217–220], increased ejection fraction [221], and improved myocardial contractile function [193]. For example, in a meta-analysis with eight clinical trials, IV Mg treatment of 930 acute MI patients showed a 49% reduction in VT/VF, a 58% reduction of incidence of cardiac arrest, and a 54% reduction in mortality [48]. The mechanisms underlying the protective effects of Mg have been studied in animal MI models. Mg reduces Ca overload [49,53,216,222,223], improves cellular ATP production [49], reduces myocardial oxygen consumption [49,53], attenuates catecholamine-induced elevated oxygen demand and myocardial necrosis [216,223], and protects the post-ischemic myocardium from oxidative damage [53,224]. Mg has been proposed as an adjunct therapy option in selected cases of high-risk acute MI patients, such as elderly patients, those with left ventricular dysfunction and chronic HF, and patients not suitable for reperfusion therapy [12].

The detrimental effects of Mg deficiency in arrhythmias and the protective roles of Mg supplementation have been thought traditionally to arise from Mg regulation of multiple cardiac ion channels, transporters and ionic exchangers that are responsible for the cardiac action potential, including the cardiac sodium channel [225], L-type Ca channel [58,226–228], T-type Ca channel [229], Na/Ca exchanger (NCX) [230,231], and K channels [232–237], such as the human ether-à-go-go-related K channel, the slowly inactivating K channel, and inward-rectifier K channels (for reviews [6,56,91,238]). Several mechanisms have been proposed for how Mg plays its roles, such as acting as a channel pore blocker, binding directly on channels, altering membrane surface potential, modulating ATP-consuming kinases and enzymes, and mediating allosteric effects on channels (see reviews [6,56,201,238]). For the cardiac sodium channel, Mg deficiency leads to a downregulation of channel function [225], which can
lead to increased risks of arrhythmias by virtue of decreased conduction velocity. Mg has been proposed as a Ca antagonist for Ca channels including the L-type Ca channel, the T-type Ca channel, the NCX, and the SR RyR2. Mg has at least two effects on L type Ca currents: inhibition of channel open probability and on channel activity under protein kinase A phosphorylation [58,226–228]. Under low [Mg]i, the NCX current is significantly increased, compared to under physiological concentration [230,231], which leads increased incidence of triggered arrhythmia [231]. Physiological concentrations of [Mg]i mainly inhibit Ca-induced Ca release from RyR2 [239] by competing with Ca [240,241]. Under Mg deficiency, increased Ca leak from RyR2 disturbs Ca homeostasis and increases the occurrence of early and delayed afterdepolarizations. Mg, at physiological concentrations mainly plays an inhibitory role on cardiac K channels by interfering with the passage of K ions and reducing the channel open probability [242]. Under Mg deficiency, K currents are increased [232–237], which can lead to arrhythmias [243]. Under Mg deficiency, the comprehensive effects of Mg on multiple cardiac ion channels, transporters, and ion exchangers lead to prolongation of the action potential duration, increasing the risk of triggered electrical activity, while Mg supplementation reverses these effects and alleviates arrhythmias.

The protective effects of Mg supplementation may also result from the suppression of Mg on oxidative stress and inflammation, which are common in patients with CVD such as HF [244,245] and AF [246–250]. Oxidative stress can induce fibrosis and electrical remodeling [246–248,251]. Our group has reported that Mg deficiency induces oxidative stress [1,82], which promotes arrhythmias [252] and downregulations on multiple cardiac ion channels and transporters including cardiac sodium channel [251,253–255], Ca channels [255–258], NCX [259], RyR2 [260], and K channels [255,256,261–263]. Mg supplementation can inhibit ROS overproduction [1,82] and reverse oxidative stress-induced channel changes. Mg deficiency has also shown to activate PKC [145], which can downregulate the cardiac sodium channel [251,253]. Inflammation that can be induced by Mg deficiency [81] is also associated with increased arrhythmic risks [264,265] and the suppression of Mg supplementation on inflammation [146,161] should have protective effects.

3.3. Mg Treatment of Other Cardiovascular Diseases

Mg regulation of insulin signaling speaks to its importance in metabolic diseases such as diabetes and obesity. Mg deficiency is often reported in DM [1,11,23,24,26,266] and obesity [105,267,268], both of which are high risk factors of developing CVD. Strong associations are reported between metabolic diseases and hypomagnesemia, inflammation, and oxidative stress [152]. Type 1 DM patients show Mg deficiency, and Mg hydroxide (Mg(OH)2, 500 mg/day for 21 weeks) improves insulin resistance [23] in part because of Mg upregulation of Akt2 and insulin receptor substrate 1 [183]. A meta-analysis study of randomized, double-blind controlled trials of 370 patients with type 2 DM shows that oral Mg supplementation at a median dose of 360 mg/day for a median duration of 12 weeks (4–16 weeks) is effective in reducing fasting plasma glucose levels and increasing high-density lipoprotein (HDL) cholesterol levels [266]. Long-term Mg supplementation significantly improves homeostatic model assessment of insulin resistance (HOMA-IR) index and fasting glucose in both diabetic and non-diabetic patients [269]. For example, oral MgCl2 (26 mmol/day for three months) shows significant improvement of insulin resistance with reduced HOMA-IR index, compared with placebo [97]. Oral MgCl2 supplementation of pre-diabetic patients with hypomagnesemia improves glycemic control with lower fasting and post-load glucose, and higher HDL cholesterol and serum Mg after four months of treatment [270]. A possible mechanism of Mg regulation of insulin signaling has been proposed in Figure 4.

Mg supplementation has shown important therapeutic effects in hypertension [15,16,271] and stroke [17,18]. Mg participates in the metabolism of L-arginine-NO system, essential fatty acids and eicosanoids. With Mg deficiency, the beneficial products such NO and resolvins are suppressed, while pro-inflammatory cytokines are elevated [103]. This can result in inflammation and vasoconstriction, cause high blood pressure and platelet aggregation [272], and induce
hypertriglyceridemia and pro-atherogenic changes in lipoprotein metabolism [273,274], leading to the development of atherosclerosis and stroke. Mg supplementation has shown efficacy to treat vascular disease. For instance, oral Mg pidolate (600 mg/daily for 12 weeks) significantly reduced ambulatory blood pressure in patients with mild hypertension, accompanied by increased serum and intracellular Mg levels [271]. Dickinson et al. reviewed 12 randomized controlled trials and reported that diastolic but not systolic blood pressure is significantly decreased by Mg supplementation [15]. A recent meta-analysis study of 22 trials shows that Mg supplementation (120–973 mg/day) slightly but significantly decreases both systolic and diastolic blood pressure in a dose-dependent manner [7]. Mg can inhibit tissue transglutaminase and lysyl oxidase, both of which are associated with hypertension and atherosclerosis [275]. Mg can be used to reduce thrombotic complications. The Mg transporter TRPM7 has been found to be a key modulator of phospholipase C and the platelet Ca transient. High serum Mg inhibits TRPM7 activity. Blockade of TRPM7 kinase activity causes a significant defect in platelet aggregation and exhibits protective effects from ischemic stroke [276]. Mg inhibition of mitochondrial ROS production plays a protective role in carotid artery stenosis [141].

Mg deficiency has been reported to contribute to other vascular disorders such as pre-eclampsia and eclampsia [277–281]. IV MgSO₄ became a standard treatment for pre-eclampsia and eclampsia seven decades ago [81,277–279] and the pharmacokinetic properties have been studied [282,283]. MgSO₄ decreases the risk of eclampsia to half and reduces the risk of maternal death [280]. Mg supplementation helps decrease oxidative stress, suppress inflammation, reduce clotting factors [284], and increase the expression of calcitonin gene-related peptides and substance P that have vasodilatory effects and improve pre-eclampsia in women [85,285,286].

3.4. Gender Differences in Mg Therapy

Clinical studies have shown gender differences in Mg deficiency and associated diseases. For example, lower serum Mg is associated with DM and obesity in women but not in men [267,268]. Lower serum Mg levels are associated with a higher risk of cardiovascular mortality and all-cause mortality in female patients but not in the male subgroup [287–289], and the protective effect of Mg intake against total cardiovascular disease mortality risk is also observed most strongly in women [290]. IV MgCl₂ shortens the atrial effective refractory period in women but prolongs the interval in men [291]. Mg intake is associated with the reduction in the systematic inflammation markers plasma CRP and E-selectin in women [137]. Generally, females seem more vulnerable to Mg deficiency. The mechanism of gender differences is unclear.

4. Mg Treatment: Routes, Chemical Formulations, Doses, and Duration

Mg supplementation is normally administrated via mouth, IV or intramuscular injection. Table 2 lists Mg supplementations that are effective in some clinical studies on treatment of HF, TdP, AF, VA, acute MI and hypertension in different salt formulations, routes of delivery, doses and treatment duration. The serum Mg level was often elevated after Mg treatment, no matter whether the patients had normal Mg levels or hypomagnesemia. Oral administration normally lasts longer (weeks to months, even a year in [197]) and is suitable for patients with low serum Mg levels and chronic diseases that are associated with Mg deficiency. Daily Mg supplementation can improve Mg deficiency-induced oxidative stress and inflammation with little side effects. Common oral Mg formulations include Mg L-lactate dehydrate, Mg gluconate, MgCl₂, and Mg L-aspartate hydrochloride [292]. Organic formulations of Mg supplementation such as gluconate, lactate, and aspartate have been reported to have better [293] or equivalent [294] bioavailability compared with inorganic salts such as MgCl₂ and MgSO₄. Mg oxide and carbonate show extremely low bioavailability, probably because of their low solubility [292]. The oral dose applied in clinic studies listed in Table 2 ranges from 1.0 to 16 mmol Mg/day with a duration from three weeks to one year. Oral Mg therapy has some limitations, however, because of slow absorption and a simultaneous increase in urinary clearance of Mg. For example, oral Mg L-lactate does not elevate serum or intracellular Mg in >80% of patients who have intracellular Mg deficiency [295].
**Table 2.** Mg treatments in different salt forms, routes, doses, and treatment durations that present protective effects in clinical studies on heart diseases. Serum Mg levels before and after Mg treatment are listed if tested in the studies. Serum Mg values are mean ± SD unless otherwise stated.

| Salts     | Route | Dose              | Treatment Duration | Treating Diseases                  | Serum Mg (mmol/L) | Reference |
|-----------|-------|-------------------|--------------------|------------------------------------|-------------------|-----------|
| MgSO₄     | Oral  | 5.9 mmol/day MgCO₃| 8 days             | NYHA II-IV HF-VA                   | 0.66 ± 0.14       | [10]      |
|           | IV    | 0.1 mmol/h        |                    | NYHA II-IV HF-VA                   | 0.74 ± 0.0        | [196]     |
|           | IV    | 1.67 mmol/h       | 24 h               | NYHA II-III HF-QT variability, ischemic HF | 0.85 ± 0.02       | [194]     |
|           | IV    | 1st bolus 30 mmol/h| 20 min             | AF                                | 0.85 ± 0.10       | NA        |
|           | IV    | 2nd bolus 5 mmol/h | 2 h                |                                    |                    |           |
|           | IV    | 1st bolus 66 mmol/h| 15 min             | New-onset AF                       | 1.1 ± 0.4         | NA        |
|           | IV    | 2nd bolus 11 mmol/h| 6 h                |                                    |                    |           |
|           | IV    | 0.15 mmol/kg BW    | 10 min             | Chronic AF                         | 0.90 ± 0.09       | [297]     |
|           | IV    | 0.10 mmol/kg BW    | 10 h               |                                    |                    |           |
|           | IV    | 12–40 mmol total   | 20 min–24 h        | Acute onset AF                     | normal            | NA        |
|           | IV    | 24 mmol/h          | 20 min, twice in   | Postoperative AT                   | 0.85 ± 0.08       | [198]     |
|           | IV    |                    | 6 h for 3 days     |                                    | 0.95 ± 0.09       |           |
|           | IV    | 2.7 mmol/h         | 24 h               | AMI-VA and mortality               | 0.80 ± 0.01       | [219]     |
|           | IV    | 2.8 mmol/h         | 24 h               | AMI-VT                            | 0.8 ± 0.2         | [298]     |
|           | IV    | 3.8 mmol/h         | 48 h               | AMI-HF and mortality               | NA                | NA        |
|           | IV    | 60 mmol/h          | 20 min             | AMI-VA                            | 0.92 ± 0.02       | [220]     |
|           | IV    | 1st bolus 48 mmol/h| 10 min             | AMI-VA                            | 0.78 (0.61–0.93)  | [224]     |
|           | IV    | 2nd bolus 2.6 mmol/h| 24 h              |                                    | 1.30 (1.1–1.74)   |           |
|           | IV    | 1st bolus 96 mmol/h| 5 min              | AMI-HF and mortality               | 0.80 ± 0.10       | [192]     |
|           | IV    | 2nd bolus 2.7 mmol/h| 24 h              |                                    | 1.55 ± 0.10       |           |
|           | IV    | 1.5–10 mmol/h      | 1.7–11.1 h         | TdP                               | 0.94 ± 0.08       | [200]     |
| MgCl₂     | Oral  | 25 mmol/h          | 1.5 h, twice a     | TdP                               | 0.98 ± 0.11       | [41]      |
|           |       |                    | day for 3–4 days   |                                    | 1.08 ± 0.08       |           |
|           | IV    | 1st day 20 mmol/h  | ~50 min, 50 min,   | TdP                               | NA                | NA        |
|           |       | then 5 mmol/h      | once a day for 3–4 |                                    | normal            | [299]     |
|           | IV    | 33 mmol/h          | 30 min             | VT                                | NA                | NA        |
|           | IV    | 1.5 mmol/h         | 24 h               |                                    | NA                | NA        |
|           | IV    | 0.21–0.42 mmol/kg BW| intraoperation      | Postoperative arrhythmias         | NA                | NA        |
| Mg oxide  | Oral  | 9.9 mmol/day       | 12 w               | NYHA II-IV HF-VA                   | 0.87 ± 0.05       | [195]     |
| Mg citrate| Oral  | 1.4 mmol/day       | 5 w                | AMI-VA and mortality               | 0.75 (0.56–0.96)  | [218]     |
| Mg gluconate | Oral  | 7.2 mmol/day      | ~10 w              | SV A                             | 0.78 ± 0.03       | [301]     |
| Mg glutamate | Oral  | 2.6 mmol          | twice              | TdP                               | 0.79 ± 0.10       | [305]     |
| Mg-DL-     | Oral  | 6 mmol/day         | 3 w                | VA                               | 0.85 ± 0.03       | [46]      |
| hydrogen-  |       |                    |                    |                                   | 0.88 ± 0.04       |           |
| aspartate | Oral  | 21 mmol/day        | 12–24 h            | Hypertension of ICD patients       | 0.91 ± 0.08       | [295]     |
| Mg eoritate| Oral  | 1.00 ± 0.04 mmol/day| 1 year            | NYHA IV HF-survival and symptoms  | NA                | NA        |

1 Value = mean±SEM; 2 median (range). Abbreviations: min, minute/minutes; h, hour/hours; w, week/weeks; AF, atrial fibrillation; AT, atrial tachycardia; AMI, acute myocardial infarction; BW, body weight; HF, heart failure; NA, not available; SVA, supraventricular arrhythmia; TdP, torsades de pointes; VA, ventricular arrhythmias; VT, ventricular tachycardia; IV, intravenous.
The IV route of Mg supplementation mainly uses the inorganic salts MgSO$_4$ and MgCl$_2$ to acutely treat patients with arrhythmias. The dose ranges from 0.1–100 mmol/h. A high dose but short treatment for only a few minutes is often given to patients as the first bolus to boost Mg levels, followed by a significantly lower dose but lasting for hours as a prophylactic treatment. For example, in the LIMIT-2 trial, IV MgSO$_4$ infusion was administrated at 96 mmol/h for 5 min followed by 2.7 mmol/h for 24 h to treat acute MI patients [192]. The patient average serum Mg level increased from 0.80 ± 0.10 to 1.55 ± 0.10 mmol/L after the first dose and was maintained at 1.80 ± 0.15 mmol/L after the second dose. The treatment significantly decreased left ventricular failure and mortality [192].

Mg supplementation has beneficial effects when combined with other treatments. For example, oral Mg supplementation has been used concomitantly with antiarrhythmic drugs [201]. IV Mg administration improves the efficacy of ibutilide in converting AF or flutter [306,307]. IV Mg is considered as a safe and effective adjunct to digoxin in controlling the ventricular response in AF in a meta-analysis of 515 patients with AF [203]. Early Mg therapy, when conjoined with reperfusion therapy, is reported to decrease the mortality after MI and the occurrence of HF [192,308]. Mg orotate used as adjuvant therapy in severe NYHA IV HF patients shows increased survival rate and improves clinical symptoms when compared with the placebo [197]. For HF patients using diuretics, hypomagnesemia also predisposes to hypokalemia. It is therefore critical to maintain both Mg and K levels by using either Mg/K supplementation or K- and Mg-sparing diuretics that inhibit renal K and Mg excretion (such as spironolactone, triamterene and amiloride) [309–311].

Adverse side effects such as gastrointestinal distress, diarrhea, and nausea have been reported with Mg supplementation for both oral and IV administration [203,266]. For IV infusion, this could be eliminated by slower infusion. For example, slow infusion of IV MgSO$_4$ (2.5 mM/h for 2 h and 1.5 mM/h for 1.5 h) was used to treat patients with TdP, and patients showed no side effects [41]. For oral administration, smaller doses with multiple times per day may lead to less side effects.

Another way to maintain a normal Mg level is to increase dietary Mg intake. Recommended dietary allowances of Mg for adults are 400–420 mg for men and 310–320 mg for women [312]. In a study of young American adults (4,637 Americans of 18–30 years old), Mg intake is reported to be inversely associated with the incidence of metabolic syndrome regardless of gender and race [21]. People who had the highest daily Mg intake (~191 mg Mg/1000 kcal) had significantly higher HDL cholesterol, lower blood glucose, lower fasting insulin levels, and lower blood pressure, compared with people who had the lowest daily Mg intake (~96 mg/1000 kcal) [21]. In a meta-analysis of 40 prospective cohort studies with more than one million patients, Fang et al. reported that increasing dietary Mg intake (per 100 mg/day increment) is linked to a 22% reduction in the risk of HF, as well as reduced risks of all-cause mortality, stroke, and type 2 DM [313]. Mg-rich food includes dark leafy greens (spinach, Swiss chard, kale, collard greens, turnip greens), vegetables (acorn squash, artichokes, okra, sweet corn, potato), nuts (almond, cashew, peanuts, and Brazil nuts), seeds (flax, pumpkin, sesame seeds, chia seeds), legumes (black beans, kidney beans, soybeans, lima beans, lentils, chickpeas, and green peas), fatty fish (salmon, tuna, mackerel, and halibut), whole grains, fruits (bananas, dried figs, guavas, avocados, kiwi, papaya, berries, cantaloupe, and grapefruit), dark chocolate, and yogurt. Most of the popular diets such as DASH (dietary approaches to stop hypertension) diet, Mediterranean diet, MIND diet (combination of the Mediterranean and DASH diet), Mayo Clinic diet, and vegetarian diet emphasize eating more vegetables, whole grains, and fruits that are all Mg-rich food [314]. Some diets such as the Mediterranean diet also encourage nuts, seeds, legumes, and fatty fish that contain higher amounts of Mg. For example, daily Mg intake in the DASH diet is ~500 mg [315], which meets the recommended dietary allowance. These diets also discourage processed food with high saturated fat and sugar but low Mg.

5. Limitations and Controversial Reports of Mg Supplementation

Not all reports of Mg use are positive and not all preparations of Mg have the same effect. Despite efficacy on some arrhythmias, Mg showed no statistically significant effect on monomorphic VT,
shock-resistant VF, and postoperative AF incidence [316–318]. Mg showed little effect on implantable cardioverter defibrillator (ICD) therapy for ischemic cardiomyopathy. For instance, oral Mg L-lactate treatment of patients with ICDs who had Mg deficiency did not reduce the occurrence of arrhythmias in ICD patients and had little impact on the health-related quality of life [319]. While several studies show that IV Mg supplementation (MgSO₄ and MgCl₂) improved mortality in acute MI patients [192,218,219,221], Feldstedt et al. observed no improvement with IV MgCl₂ on either the in-hospital or the follow-up mortality after acute myocardial infarction [320]. Moreover, Mg infusion was accompanied by a significantly increased incidence of atrioventricular conduction disturbances. Eichhorn et al. reported no correlation between serum Mg levels and survival of HF patients [321]. These controversies could result from variances in Mg treatments (different salts, routes, doses, treatment timing and durations) or severity of disease. Nevertheless, it appears that Mg has limited efficacy to treat acute reentrant rhythms such as ventricular tachycardia or atrial fibrillation [318,322,323] and that Mg may not be as efficacious in ischemic heart disease [319,320], where the arrhythmic substrate may be based more on structural issues and reentrant mechanisms.

6. Conclusions

Mg deficiency is common in CVD, and Mg supplementation has shown antioxidant and anti-inflammatory properties in patients with and without Mg deficiency. Mg supplementation is well tolerated with few side effects. Therefore, it may represent a reasonable additional therapy for many CVDs.

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