Is there a Need for Ubiquinone (CoQ10) Supplementation in Statin-Associated Myopathy?

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Abstract: Statins are currently the most effective drugs in reducing low-density lipoprotein cholesterol (LDL-C). With their mechanism of action, by inhibiting 3-hydroxy-3 methylglutharyl coenzyme A reductase, statins decrease cholesterol production. The same biosynthetic pathway is shared by ubiquinone or coenzyme Q10. Statins block production of dekaprenyl-4-benzoate, a precursor of coenzyme Q10 (CoQ10), which is an essential component in mitochondrial transport system. Ubiquinone deficiency may affect oxidative phosphorylation and adenosine triphosphate production, which can subsequently result in impairing of muscle energy metabolism and contribute to development of myopathy. Statin therapy also decreases antioxidant status in the body, resulting in to increase in free radical damage to cells in various parts of body: liver, nerves, muscles. Statins can decrease natural antioxidant protection present in our body and predispose toxicity.

Statin-associated myopathy is the most common side effect of statin treatment often lead to statin dose reduction, or therapy cessation, which can negatively affect cardiovascular risk management. The spectrum of statin-related myopathy ranges from common but clinically benign myalgia to rare but life-threatening rhabdomyolysis. Observational studies suggest that myalgia can occur in up to 10% of persons prescribed statins, whereas rhabdomyolysis continues to be rare.

Statins lower circulating levels of CoQ10 up to 54%, whereas several studies did not confirmed a lowering of CoQ10 levels in muscles during statin therapy and the evidence was given also on low dose of statin therapy, which did not appear to reduce intramuscular levels of CoQ10 in symptomatic patients with statin myopathy. The conflicting results have been published on the impaired mitochondrial function, which was found during the analysis of the myocyte cells in patients treated with statins. The supplementation of CoQ10 results in increasing of serum CoQ10, but it is not clear if it relieves myopathic symptoms in statin treated patients. Yet available intervention studies reported contrasting results and just two of them proved benefit of CoQ10 supplementation. Hence coenzyme Q10 supplementation is not currently recommended for routine use in the prophylaxis of statin toxicity.

Keywords: Statins, statin-associated myopathy, CoQ10, ubiquinone.

INTRODUCTION

Statins or 3-hydroxy-3 methylglutharyl coenzyme A reductase inhibitors competitively inhibit cholesterol production by reducing the synthesis of mevalonate, a critical intermediary in the cholesterol pathway. Statins are well tolerated by many patients, but muscle toxicity, also known as myopathy or myotoxicity, is a feared adverse effect of this class of drug. Myopathy is a general term referring to any disease of muscles and myalgia means muscle pain or weakness without increase in creatinine kinase. The most severe form of myotoxicity is rhabdomyolysis, which can occur in all statins, either in monotherapy or in combination therapy, especially with fibrates. Less common adverse effects of statin therapy are hepatotoxicity, peripheral neuropathy, impaired myocardial contractility and autoimmune diseases [1-3]. The mechanisms of development of statin-myopathy are unclear. There are several theories proposed to explain it. The first one says that blockage of cholesterol synthesis through statins causes the reduction of sacrolemmal cholesterol, making the skeletal muscle cells unstable [4]. The second theory is based on the assumption that inhibition of small guanosine triphosphate-bindings proteins formation, important to promote cell maintenance and growth and at the same time diminishes apoptosis [4]. Finally, the depletion of CoQ10 in muscle mitochondria during the statin therapy is to increase muscle cell fluidity and thus possibly cause myopathy [4, 5]. Poor coenzyme Q10 status and antioxidant status in the body are also predisposing factors for statin toxicity [6-10]. A number of observations suggest that statin toxicity may be due to the metabolic effects of lipid lowering in patients with minor muscle disorders [11-13]. These patients have a high frequency of mutations for metabolic muscle diseases and often have depleted mitochondrial enzymes. Their exercise physiology and biopsy findings indicate reduced oxidation of...
fats and mitochondrial dysfunction. These subjects are often intolerant of other lipid lowering therapies in addition to statins, which suggests that the myopathy is due to lipid-lowering itself more than a simple pharmacokinetic reaction to high statin levels. Altogether, these findings support the concept that statin myopathy is a metabolic muscle disease [14].

**EPIDEMIOLOGY AND RISK FACTORS OF STATIN MYOPATHY**

In randomized clinical trials (RCTs), statin myopathy incidence is about 1.5% to 5.0% [15, 16]. However, incidence of statin myopathy in clinical trials seems to be lower that in real world clinical practice. In clinical trials patients with risk factors of statin toxicity, history of statin related intolerance or developed biochemical abnormalities during the screening period before randomization are usually excluded. In addition, persons who have had previous statin intolerance would probably not enroll in clinical trials, whereas motivated enrolled patients might minimize reporting of mild statin-related myalgias [17]. Several observational studies have documented a 5% to 10% incidence of statin-associated myalgia [18, 19]. The large, observational PRIMO (Prediction of Muscular Risk in Observational Conditions) study [18] of 7924 patients exposed to high-dose statins found that 10.5% had muscle-related symptoms over 12 months. One observational study of 32 225 patients reported that 5.8% and 6.7% of diabetic and nondiabetic patients, respectively, had statin-related myalgia [19].

Postmarketing surveillance through the FDA Adverse Event Reporting System (AERS) has documented low reporting rates of statin-related myopathy, myositis, and rhabdomyolysis. From 1998 to 2000, reporting rates for all statins except cerivastatin were 0.38, 0.43, and 1.07 cases per 1 million prescriptions, respectively. From 2002 to 2004, these rates increased to 0.74, 0.57, and 3.56 cases per 1 million prescriptions, respectively, probably because of heightened awareness after the withdrawal of cerivastatin in 2001 [20]. From 2002 to 2004, the FDA AERS rates for myopathy were lowest for fluvastatin (0.43 cases per 1 million prescriptions) and highest for rosvastatin (2.23 cases); for myositis, rates were lowest for atorvastatin (0.27 cases) and highest for rosvastatin (2.37 cases); and for rhabdomyolysis, rates were lowest for pravastatin (1.63 cases) and highest for rosvastatin (13.54 cases) [20]. The high AERS rates for rosvastatin, which is the only statin launched after cerivastatin’s withdrawal, were attributed to a biased “new drug” reporting effect and to widespread lay media coverage in 2004 [20]. Of importance, the proportionate AERS rate for rosvastatin was about the same as, or was lower than, that for all statins [20]. Limitations of FDA-derived data on statin myopathy include reliance on voluntary reporting and diagnostic criteria for myopathy or rhabdomyolysis, which required much higher creatine kinase elevations than the ACC/AHA/NHLBI clinical advisory and perhaps caused underestimation of myopathy incidence [17].

The 2001 AERS rates of fatal rhabdomyolysis varied by agent (1 reported case per 5.2 million prescriptions for lovastatin, 23.4 million prescriptions for atorvastatin, 27.1 million prescriptions for pravastatin, and 8.3 million for simvastatin). These low rates starkly contrast with the rate of 1 reported case of fatal rhabdomyolysis per about 316 000 prescriptions for cerivastatin. No case of fatal rhabdomyolysis has been reported yet with fluvastatin [21]. Thus, although rates of myopathy are higher in clinical practice than in clinical trials and the AERS, the rates of rhabdomyolysis are still reassuringly low (about 0.1 to 0.2 case per 1000 person-years) and are similar to those reported in clinical trials [22].

Higher risk patients for statin myopathy are those with advanced age, females, with a small body frame and frailty, on higher statin doses, on other medications, or with other systemic diseases including hepatic or renal diseases, diabetes mellitus, or hypothyroidism [14]. For example, thin elderly women may represent a demographic category with increased risk for creatine kinase elevations greater than 10 times the upper limit of normal with statin therapy because 5% to 7% of women who received cerivastatin, 0.4 to 0.8 mg/d, had these creatine kinase elevations [23]. Also, only 6 of 22 professional athletes with familial hypercholesterolemia who received statins could tolerate 1 of several statins attempted, indicating that intense physical activity might be a risk factor [24]. During hospitalization for major surgery, the ACC/AHA/NHLBI advises short-term cessation of statin therapy to minimize myopathy risk during the perioperative period [13].

Poor coenzyme Q10 status and antioxidant status in the body are also predisposing factors for statin toxicity [6-10]. One of the most important factors for predicting risk of myopathy is also way of metabolization of statins. Most of them are metabolized by cytochrome P 450 family with the exception of pravastatin [25-27]. Because lovastatin, simvastatin and atorvastatin are metabolized by cytochrome P4503A4 (CYP4503A4), inhibitors of CYP3A4 like cyclosporine, nifedipine, felodipine, amiodarone, fibrates could theoretically increase serum statin levels and exposure to susceptible tissues [28]. Protease inhibitors are potent CYP3A4 inhibitors and thus can increase up to 30 times plasma concentration of statins [29, 30] Cyclosporine is a potent inhibitor of not only CYP4503A4 but also several membrane transporters and it was found to cause many reported cases of rhabdomyolysis [31]. Because pravastatin undergoes renal metabolism, fluvastatin and rosvastatin are primarily metabolized by CYP2C9, there 3 statins may have a lower myopathy risk, especially in the context of polypharmacy.

Genetic predisposition to statin myopathy is a rapidly expanding area of investigation. Genes of interest include those involved in the pharmacokinetics of the statin response, muscle atrophy, exercise intolerance, pain perception, and mitochondrial energy metabolism [14]. Common DNA polymorphisms in genes encoding cytochrome P450 enzymes, intestinal P-glycoproteins, and organic anion-transporting polypeptide are inconsistently associated with statin myopathy [32-37]. DNA polymorphisms of genes involved in metabolism of coenzyme Q10 and serotonin pain receptors were also inconsistently associated with statin myopathy [38, 39]. Recently, a common DNA polymorphism in the SLC01B1 gene encoding organic anion-transporting polypeptide was strongly associated with simvastatin-associated myopathy [32], but this association was not seen in patients with atorvastatin associated myopathy [35]. Finally, among 110 patients with statin myopathy, about 10% had heterozygous mutations in 1 of several genes that
normally cause rare myopathy syndromes [40], suggesting that genetic susceptibility to statin myopathy may comprise a complex mixture of rare DNA variants and common DNA polymorphisms.

**COQ10**

CoQ10 is naturally occurring, oil-soluble substance, localized in the cell membranes. Around half of ubiquinone is obtained though dietary fat ingestion, while remaining half though endogenous synthesis [41]. Synthesis of CoQ10 in human body decreases with age. CoQ10 is an antioxidant, participating in mitochondrial electron transport during oxidative phosphorylation in mitochondria, transferring electrons from complex I and complex II to complex III of respiratory chain. It forms a very effective redox system, composed of ubiquinone (reduced form), semiquinone radical and ubiquinone (oxidized form). Its reduced form is important scavenger of free oxygen radicals, prevents low-density lipoprotein cholesterol (LDL-C) oxidation and protects membrane lipids and proteins and deoxyribonucleic acid (DNA) against oxidative damage [42]. It also regenerates active forms of the antioxidants ascorbic acid and tocopherol (vitamin E) [43, 44]. In 1980s, it appeared a theory that blocking of farnesyl pyrophosphate resulting in CoQ10 depletion is involved in pathogenesis of statin-associated myopathy [45, 46]. Statins, by reductions of CoQ10 synthesis, can decrease natural antioxidant protection present in our body and predispose toxicity [1-3, 47-51]. The antioxidant status of our body depends on the presence of coenzyme Q10, vitamin A, E, and C and beta-carotene, superoxide dismutase, catalase and ceruloplasmin. Apart from these antioxidants, flavonoids, anthocyanins, minerals; selenium, chromium and copper and ω-3 fatty acids may also provide protection against oxidative stress predisposed by statins [6-8]. It also has been shown that increased levels of CoQ10 may protect cells from chemotherapy-induced oxidative stress [52]. It is suggested that prior or simultaneous administration of antioxidants may be protective against P450 induced cell damage by the statins, resulting in decrease of statin toxicity [14].

CoQ10 deficiency has been observed in patients with Parkinsons disease, Huntington’s disease, tuberous sclerosis, motor neuron disease, and cerebellar ataxia, CoQ10 supplementation may be useful in these conditions [53-56]. Low levels of CoQ10 have been described also in myocardial biopsies from patients with various cardiovascular diseases [57, 58]. Lower than normal levels of CoQ10 have been also described in the blood from patients with cardiovascular diseases as compared with levels with healthy human subjects [59, 60].

**STATINS AND COQ10 IN SERUM, MUSCLES AND MITOCHONDRIA**

The meta-analysis of several clinical trials demonstrates that statin treatment decreases serum CoQ10 levels [61] and in some of the reports the decrease was above 50 % [10, 41, 45]. Only a few small studies did not confirm this evidence, which is to be explained by the small number of participants and low statin dose [62, 63]. In the plasma, CoQ10 is transported with the very low density lipoprotein (VLDL) fraction hence, its levels are closely related to the plasma VLDL levels which can be affected by both the disease processes and by dietary factors. Moreover, whereas the cellular level of CoQ10 is dependent mostly on endogenous biosynthesis [64], the plasma level has been suggested as representing the equilibrium between CoQ10 absorption and synthesis [65]. There are also some indications, considered contradictory that statin treatment may affect muscle CoQ10 levels. Resulting from the recent human studies the effect of statins may be drug and dose dependent [61] as the significant decrease of muscle CoQ10 levels was found only in the patients treated with simvastatin 80 mg daily [66]. Taken together, it appears that the plasma level of CoQ10 cannot always be regarded as a true indicator of cellular levels. Therefore, in studies in which the cellular levels of CoQ10 and its changes under pathological conditions are investigated, tissues other than serum or plasma are more desirable [67], but only a few works have examined the intramuscular levels of CoQ10 in patients, who suffered from statin myopathy. In patients with statin associated muscle symptoms or high serum creatinine kinase or both, the muscle biopsies were obtained from 18 patients. No evidence of myocyte apoptosis was found, but only 11 biopsies provided enough tissue to perform the test. The results on muscle CoQ10 levels were controversial, not showing significant CoQ10 level decrease [68].

There should be also an evidence of impaired mitochondrial function, if CoQ10 depletion mediates the statin related myopathy. The data from some animal studies showed structural and functional mitochondria damage in statin treated animals [69-72], while few studies have directly (muscle biopsy) or indirectly (measurement of lactate:pyruvate ratio) addressed this issue in humans. In some studies an increased lactate/pyruvate ratio was found after treatment with statins [5, 73-75], which is used to show a shift toward anaerobic metabolism and could be a possible marker of mitochondrial dysfunction, but the results gained from the other studies did not confirm this observation. In the recent case study, lactic acidosis occurred and was considered as side effect of simvastatin treatment and argues that mitochondrial dysfunction is a cause [76].

**COQ10 SUPPLEMENTATION IN STATIN MYOPATHY**

To prevent side effects of statins including myopathy, it should be useful to combine statins with coenzyme Q10; however, greater clinical studies are needed. In a randomized, controlled intervention trial, among 71 patients (Intervention group) and 73 patients (Control group) of acute myocardial infarction (AMI), who were administered coenzyme Q10 (120mg/day) or B vitamins for one year, half of the patients (n=36 vs 31) in both groups, received lovastatin (10-20mg/day) [56]. Adverse effects, such as nausea (30.1 vs 9.8%), vomiting (13.7 vs 11.2%), were more common in the coenzyme Q group whereas fatigue (6.8 vs 40.8%, p<0.01) was more common in the control group [56]. Fatigue is an early manifestation of muscle damage, indicating that coenzyme Q may have prevented this adverse effect of lovastatin in the intervention group. Initial attempts in administering CoQ10 to statin treated patients with myopathic symptoms had a positive outcome; pain severity significantly decreased after 30 days of CoQ10 supplementation therapy [77, 78].

In 32 hypercholesterolemic patients with statin related myalgia, treatment with CoQ10 (100 mg daily) after 30 days, decreased the intensity of pain by 40 % in the group using
coenzyme Q10 supplements. In contrast, no change in pain intensity was observed in the group using vitamin E supplements (400 IU/daily) at the end of the trial. Additionally, a decrease of 38% in pain interference with daily activities was found in CoQ10 group in contrast with no difference in pain interference in the vitamin E group [78]. The pain was assessed using Brief Pain Inventory, which provided measures of pain severity and interference with daily activities [79]. In the randomized, double-blind, placebo-controlled study of Mabuchi et al. [80], low dosage of atorvastatin (10 mg/day) has never produced definite liver or muscle damages, and supplementation of CoQ10 (100 mg daily over 16 weeks) has never produced positive results compared with placebo. A more recent trial randomized 44 patients to CoQ10 200 mg daily or placebo for 12 weeks in combination with simvastatin [81]. All patients discontinued lipid-lowering therapy (except ezetimib) and instead started simvastatin dose 10 mg daily, which was uptitrated every 4 weeks towards maximum tolerated dose of 40 mg, if was possible. Plasma levels increased in CoQ10 group, but there was no difference in myalgia score or in statin tolerance between both groups. Pain was assessed by using visual analogue scale [82]. The benefit of CoQ10 supplementation was not shown in this trial due to a several limitations: patients did not experience sufficiently severe myalgias, the absorption of CoQ10 likely to be decreased as the patients were initiated CoQ10 at the same time as the statin therapy. More recently, benefit of CoQ10 supplementation was shown in another randomized, placebo controlled trial (currently only available in abstract), using CoQ10 200 mg daily during 3 months in patients with statin associated myopathy defined as myalgia, fatigue, muscle cramps and muscle weaknes [83]. The present evidence does not support routine coenzyme Q10 supplementation in statin-associated myopathy. However, guidelines of some associations recommend using CoQ10 supplements in patients with the risk factors of statin-associated myopathy [84]. Possible risks of widely prescribed statin therapy should be decreased by administering of lower doses, combination with coenzyme Q10 and avoidance of risky combinations of drugs with interfering metabolism via cytochrome P 450. The role of lipid lowering diet plus exercise is indisputable and may support lower doses of lipid lowering drugs or even lower use of these drugs [85] and may prevent prooxidative stage that is important for statin intolerance. Use of coenzyme Q10 even in smaller doses of 30 mg/day may be rewarding in the prevention of adverse effects of statins [56, 86].

CONCLUSIONS

Myalgia is the most common adverse event of statins leading to nonadherence or even stopping of statin therapy. It affects up to 10% of patients receiving statin treatment, statin-induced fatal rhabdomyolysis is extremely rare. It is plausible that statin block the production of mevalonate, which is the precursor of CoQ10. Statin interference with CoQ10 production prompted the hypothesis that pathogenesis of statin myopathy may be related in part to statin inhibition of the endogenous synthesis of coenzyme Q10. However, coenzyme Q10 supplementation is not currently recommended for routine use. There is at least a need to consider coenzyme Q10 administration in conjunction with statin when they are given >20mg/day, as it might also prevent deaths due to myopathy and other toxic effects because statin toxicity is considered a metabolic myopathy [27, 87].

There are not known risks for CoQ10 supplementation in mild to moderate dosage (30-300 mg daily), which may support its usage in common medical practice in patients with statin related myopathy. It is postulated, that effect of CoQ10 supplementation is dose and form dependent; the therapeutic response likely to correlate with CoQ10 plasma levels achieved through supplementation therapy (target treatment levels 2.5-3.5 µmol/L). Therefore, when CoQ10 supplementation is occurring, it is advisable to monitor the CoQ10 levels to ensure the efficacy, as there a variable bioavailability of commercial formulations and inter-individual variation in CoQ10 absorption exist. CoQ10 is available in several forms, including powder, suspension, oil solution, solubilized forms (All Q and Q-gel), all with different bioavailability. Using a lipid formulations and taking CoQ10 with food improves absorption [88].

Coenzyme Q10 supplementation may offer an alternative to stopping treatment with these drugs, if adverse effects appear. This may be of great importance, as the statins represent the basic pillars for primary and secondary prevention of atherosclerosis and their prescription rate is increasing dramatically.

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