Vanillic Acid Restores Coenzyme Q Biosynthesis and ATP Production in Human Cells Lacking COQ6

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Received 3 February 2019; Revised 30 April 2019; Accepted 26 May 2019; Published 10 July 2019

Guest Editor: Livia Hool

Coenzyme Q (CoQ), a redox-active lipid, is comprised of a quinone group and a polyisoprenoid tail. It is an electron carrier in the mitochondrial respiratory chain, a cofactor of other mitochondrial dehydrogenases, and an antioxidant. CoQ requires a large set of enzymes for its biosynthesis; mutations in genes encoding these proteins cause primary CoQ deficiency, a clinically and genetically heterogeneous group of diseases. Patients with CoQ deficiency often respond to oral CoQ 10 supplementation. Treatment is however problematic because of the low bioavailability of CoQ 10 and the poor tissue delivery. In recent years, bypass therapy using analogues of the precursor of the aromatic ring of CoQ has been proposed as a promising alternative. We have previously shown using a yeast model that vanillic acid (VA) can bypass mutations of COQ6, a monooxygenase required for the hydroxylation of the C5 carbon of the ring. In this work, we have generated a human cell line lacking functional COQ6 using CRISPR/Cas9 technology. We show that these cells cannot synthesize CoQ and display severe ATP deficiency. Treatment with VA can recover CoQ biosynthesis and ATP production. Moreover, these cells display increased ROS production, which is only partially corrected by exogenous CoQ, while VA restores ROS to normal levels. Furthermore, we show that these cells accumulate 3-decaprenyl-1,4-benzoquinone, suggesting that in mammals, the decarboxylation and C1 hydroxylation reactions occur before or independently of the C5 hydroxylation. Finally, we show that COQ6 isoform c (transcript NM_182480) does not encode an active enzyme. VA can be produced in the liver by the oxidation of vanillin, a nontoxic compound commonly used as a food additive, and crosses the blood-brain barrier. These characteristics make it a promising compound for the treatment of patients with CoQ deficiency due to COQ6 mutations.

1. Introduction

Coenzyme Q (CoQ) is a key component of the mitochondrial respiratory chain (RC) where it shuttles electrons from complexes I and II to complex III. It is also a cofactor of other mitochondrial dehydrogenases and of uncoupling proteins, an antioxidant, and a modulator of the mitochondrial permeability transition pore [1]. CoQ is comprised of a
quinone group and of a polyisoprenoid tail, which varies in length in different species: ten isoprene units in humans (CoQ10), nine in mice (CoQ9), and six in yeast (CoQ6). The CoQ biosynthetic pathway is still incompletely characterized, especially in higher eukaryotes [2]. The biogenesis of the isoprene tail begins in the cytosol through the mevalonate pathway, sharing its initial steps with cholesterol biosynthesis. The individual isopentyl diphosphate units produced in these pathways are then joined together to form the all-trans polyproprenyl tail by COQ1 (in yeast) and PDSS1 and PDSS2 (in mammals) within mitochondria. The quinone group is synthesized from tyrosine through a poorly characterized set of reactions that produce 4-hydroxybenzoate (4-HB) [3, 4]. 4-HB is then joined to the polyisoprenoid tail by COQ2, an integral protein of the mitochondrial inner membrane [5]. Subsequent biosynthetic steps occur within the mitochondrial matrix and are catalyzed by a set of enzymes that modify the quinone ring. In eukaryotes, these enzymes form a multiprotein complex associated with the inner mitochondrial membrane [6]. This complex is still poorly characterized in mammals, and the precise sequence of reactions is not known [2]. Among the different genes coding the components of this complex, COQ6 encodes a FAD-dependent monoxygenase responsible for the addition of the hydroxyl group in position C5 of the quinone ring [7].

CoQ biosynthesis is relevant for human diseases because mutations in genes involved in CoQ biosynthesis cause primary CoQ deficiency, a clinically heterogeneous group of disorders. In particular, patients with mutations in COQ6 present with steroid-resistant nephrotic syndrome (SRNS) associated with sensorineural deafness and a variable degree of encephalopathy [8]. CoQ-deficient patients respond to oral CoQ10 supplementation, making this one of the few treatments for SRNS [11, 12]. In the case of COQ6, the mutated sequence (G255R, and a negative control isoform COQ6Isw) was used to transiently transfect HeLa cells stably expressing mtRFP seeded on glass cover slips [16]. 48 hours after transfection, cells were examined using a Zeiss Axio Imager M2 fluorescence microscope.

In this work, we report the generation of a cell line lacking functional COQ6, which was used to test the effect of VA supplementation.

2. Materials and Methods

2.1. Cell Culture and Reagents. HEK 293 cells were cultured at 37°C using DMEM 4.5 g/L glucose, L-glutamine (6 mM) (Gybro™) supplemented with sodium pyruvate solution (1 mM) (Sigma), an antibiotic/antimycotic solution (Sigma), uridine (10 µM) (Sigma), and 10% fetal bovine serum (FBS) (Gybro™). When required, supplemental CoQ prediluted in ethanol 100% was added to the medium at a final concentration of 100 µM (coenzyme Q10 ≥98%, HPLC, Sigma) and supplemental vanillic acid was added at a final concentration of 500 µM (vanillic acid, ≥97%, HPLC, Sigma). For all biochemical assays, cells were incubated for 48 hours in DMEM containing 2 mM glucose and 5% FBS, to force mitochondrial respiration.

2.2. Generation of CRISPR Knockout Cell Lines. The CRISPR/Cas9 constructs were purchased from transOMIC technologies. We performed transfection by using Lipofectamine 2000 (Life Technologies), according to the manufacturer’s instructions. Two pCLIP-All-EFS vectors, which express Cas9 and the target gRNAs, together with the puromycin resistance gene (TEVH-1164665: 3′-TCTCTGTAGAGACCGTGCTACT-5′; TEVH-1097523: 3′-CTAGGGTAAATGAAACCCAA-5′), were cotransfected into HEK293 cells. Cells were incubated in medium with 1.5 µg/mL puromycin for selection. Cell clones were obtained by cell sorting into 96-well plates, cultured until confluence, and duplicated for genotyping PCR. Culture media were supplemented with CoQ10 and uridine as described [14]. Three positive clones were expanded and mixed (1:1:1) to obtain a homogeneous cell population.

2.3. Construction and Transfection of Lentiviral Vectors. The coding sequence of human COQ6 isoform a and isoform c, the mutated sequence COQ6 G255R, and a negative control expression sequence were transferred into the lentiviral expression plasmid (pLent6/V5-DEST™ Gateway™ Vector, Invitrogen). To generate lentiviral particles, vectors were cotransfected with packaging vectors (ViraPower Packaging Mix, Invitrogen) into HEK293-FT cells using Lipofectamine 2000 (Life Technologies). Culture supernatants were harvested on day 3 and used to transduce HEK293 COQ6KO cells. Selection was carried out as described [15].

2.4. GFP Fusion Vectors. A COQ6_Iso_c-GFP fusion gene was constructed using the pEGFP-N1 as previously reported [8], except that the forward primer employed was specific for COQ6 isoform c (5′-TCTAAAGCTTGTATCGGGGCG 5′). As the PCR template, we employed the plasmid generated for the lentiviral construct. The pEGFPN1-COOQ6_Iso_c was used to transiently transfect HeLa cells stably expressing mtrFP seeded on glass coverslips [16]. 48 hours after transfection, cells were examined using a Zeiss Axio Imager M2 fluorescence microscope.
2.5. Immunoblot Assay. Standard techniques were employed for SDS-PAGE and Western blotting to PVDF membranes using the antibody COQ6 12481-1-AP (ProteinTech). Peroxidase-conjugated anti-rabbit IgG was used as secondary antibodies (Santa Cruz). Visualization of antibody protein complexes was achieved by enhanced chemiluminescence (LiteAblot Turbo, EuroClone) and the ChemiDoc™ XRS+ System (Bio-Rad).

2.6. Lipid Extraction and HPLC Analysis. Harvested cells (1 mg protein) were resuspended in 0.3 mL of a 0.15 M KCl solution. Then, 200 μL glass beads, 10 μL of a 5 μM UQ8 standard solution, and 3 mL methanol were added. The tubes were vortexed for 1 min, 2 mL petroleum ether (40-60° boiling range) was added, and vortex was repeated for 1 min. The tubes were centrifuged at 700 rpm for 1 min, the upper phase was collected, and the methanol phase was extracted again with 2 mL petroleum ether. Both petroleum ether phases were combined and dried under a nitrogen flow, and the lipid extracts were resuspended in 200 μL ethanol. HPLC analysis was conducted essentially as described [17] with the following modifications. Samples corresponding to 0.2 mg protein were injected onto the C18 column, and separation was obtained at a flow rate of 1 mL/min with a mobile phase composed of 25% isopropanol, 45% methanol, 20% ethanol, and 10% of a solution composed of 90% (v/v) isopropanol, 10% (v/v) 1 M ammonium acetate, and 0.1% (v/v) formic acid. The precolumn electrode (5020 Guard Cell, Thermo) was set either at +650 mV (oxidizing mode) or at -650 mV (reducing mode). Mass spectrometry detection was conducted in positive mode with electrospray ionization, probe temperature of 400°C, and cone voltage of 80 V. 

HP10 was detected with single-ion monitoring at m/z 806.5 (M + NH4+) and its reduced form, 3-decaprenyl-1,4-benzoquinol, at m/z 808.5 (M + NH4+).

CoQ10 biosynthesis rates were measured as previously described [18], by evaluating incorporation of 14C-labelled 4-HB.

2.7. Respiratory Chain Activities and ATP Levels. Activities of mitochondrial respiratory chain complexes were measured as described [19]. ATP levels in cells were determined using the ATPlite Luminescence Assay System (PerkinElmer) according to the manufacturer’s instructions and using a Victor3 (PerkinElmer) multilabel plate reader.

2.8. ROS Measurement. Intracellular ROS was measured using two different assays. To measure the mitochondrial redox state, we used the mitochondrially targeted redox-sensitive GFP (ro-GFP) system [20]. A vector expressing ro-GFP was transfected into the different cell lines using Lipofectamine 2000 (Life Technologies). After 48 hours of growth and treatment, fluorescence was measured in a Victor3 (PerkinElmer) multilabel plate reader with an excitation of 405 nm and emission of 485 nm.

Total oxidant levels were measured using the oxidant-sensitive fluorescent probe CM-H2DCFDA (Invitrogen). ROS were detected using CM-H2DCF-DA, a chloromethyl derivative of H2DCFDA. Cells were seeded onto 24 mm diameter glass coverslips placed in 6-well plates and cultured in the appropriate medium as described above. 30 min before measurements, cells were loaded with CM-H2DCF-DA (2.5 μM) and then washed twice. All the steps were carried out at 37°C with 5% CO2. The chambered coverslips were transferred to a Leica (Wetzlar, Germany) DMi6000B microscope, equipped with a digital camera. Fluorescence was measured in 5-7 random fields per chamber. For each group, 4-6 chambers were analyzed. Fluorescence emission was monitored by using 560 ± 20 nm excitation and 645 ± 37 nm emission filter setting. Data were acquired and analyzed using MetaFluor software (Universal Imaging).

2.9. Measurement of Oxygen Consumption by the Seahorse XF24 Extracellular Flux Analyzer. The oxygen consumption rate (OCR) was determined using a Seahorse XF24 Extracellular Flux Analyzer following the manufacturer’s instructions. 24 h before seeding the Seahorse plate, cells were treated with vanillic acid or CoQ. 106 cells per well were seeded onto poly-D-lysine precoated (Sigma) Seahorse 24-well plates for 48 hrs. Then, medium was replaced for 2 mM glucose DMEM and cells were incubated with this medium for 24 h. Prior to the measurements, medium was replaced with Seahorse XF base medium supplemented with 2 mM glucose, 2 mM glutamine, and 1 mM sodium pyruvate and incubated for 1 hr at 37°C without CO2. OCR was measured under basal conditions and after the sequential addition of oligomycin (1 μM), FCCP (0.2 μM), rotenone (1 μM), and antimycin A (2.5 μM). To normalize respiration rates, cells were harvested and counted after the assay.

2.10. In Silico and Statistical Analyses. The structure of human COQ6 was modelled on that of P. fluorescens para-hydroxybenzoate hydroxylase (Protein Data Bank code 1PBE) as reported [13]. Statistical analyses were performed as described [21].

3. Results

3.1. COQ6 Is Essential for CoQ10 Biosynthesis in Mammalian Cells. Wild-type HEK293 cells were transfected with the CRISPR-Cas9 constructs, and after selection, individual clones were genotyped. All analyzed clones harbored a deletion of 75 nucleotides (Figure 1(a)), which caused a deletion of 25 amino acids (Figure 1(b)) in a region of the protein which participates to the FAD-binding site (Figure 1(c)) and is contained in all three isoforms of human COQ6.

Western blot analysis of the mixed clones did not detect residual COQ6 protein (Figure 1(d)), and when the mutant cDNA was expressed in delta coq6 yeast, VA could not rescue growth in nonfermentable media (not shown). These results indicate that the deletion results in an inactive, unstable protein. From now, we will refer to these cells as COQ6Δ25.

CoQ10 levels were markedly reduced in COQ6Δ25 cells compared to wild-type cells (Figure 1(e)). Incorporation of 14C-labelled 4-HB was virtually undetectable (Figure 1(f)), indicating that, in analogy with what we observed in COQ4KO cells [22], the residual CoQ10 detected in these cells is not produced endogenously but it is derived from the
The chromatogram showed an additional radioactive peak present only in COQ6Δ25 cells, which eluted faster than CoQ10 (Figure 1(f)). Activities of individual respiratory chain enzymes were normal; however, combined activity of complexes II and III was markedly reduced in these cells, consistent with severe CoQ10 deficiency (Figure 1(g)). ATP levels were also markedly reduced (see below). Complex I + III activity was not assayed as results are unreliable in cultured (even when mitochondria-enriched preparations are used), due to the residual high levels of rotenone-insensitive activity [19].

3.2. COQ6Δ25 Cells Accumulate 3-Decaprenyl-1,4-benzoquinone. To characterize the additional product detected in COQ6Δ25, we analyzed lipid extracts from WT and COQ6Δ25 cells by HPLC-MS. Electrochemical detection confirmed a marked reduction of CoQ10 in COQ6Δ25 cells and the presence of an electroactive compound that eluted
mass (M+N H 4) with single-ion monitoring for motion of 4-HP10. Overall, these data show that the impairment at 806.5 around 10.4 min (Figure 2(c)), consistent with the two-electron reduction of isoform (red). The chemical structures of (b) 3-decaprenyl-1,4-benzoquinone and (c) 3-decaprenyl-1,4-benzoquinol are shown.

Figure 2: (a) HPLC electrochemical detection analyses (with precolumn electrode in oxidizing mode) of lipid extracts from wild-type (WT) and COQ6Δ25 cells (0.2 mg protein) with UQα used as standard. UQ10 and 4-HP10 have different retention times as shown by the dotted lines. (b, c) Single-ion monitoring ((b) m/z 806.5, (c) m/z 808.5) in HPLC mass spectrometry analyses of lipid extracts from WT and COQ6Δ25 cells (0.2 µg protein) with the precolumn electrode set in oxidizing mode (ox) or reducing mode (red). The chemical structures of (b) 3-decaprenyl-1,4-benzoquinone and (c) 3-decaprenyl-1,4-benzoquinol are shown.

At 10.4 min, right after CoQ10 (Figure 2(a)). This compound was characterized by HPLC-coupled mass spectrometry, and mass scanning (m/z 600-900) showed a prominent ion at 806.5 around 10.4 min (data not shown). A signal at 10.4 min was indeed specifically obtained in COQ6Δ25 cells with single-ion monitoring for m/z 806.5 (Figure 2(b)), a mass (M + NH₄⁺) compatible with that of 3-decaprenyl-1,4-benzoquinone (4-HP10). Upon shifting the precolumn electrode to a reducing mode, the signal at m/z 806.5 disappeared (Figure 2(b)) and a signal at m/z 808.5 appeared at 5.8 min (Figure 2(c)), consistent with the two-electron reduction of 4-HP10. Overall, these data show that the impairment of CoQ10 biosynthesis in COQ6Δ25 cells leads to the accumulation of 4-HP10, in agreement with the accumulation of 4-HP6 previously reported in yeast Δcoq6 cells [7].

3.3. COQ6 Isoform c Does Not Rescue CoQ10 Biosynthesis. At least three different COQ6 isoforms exist in human cells: the most represented transcript (with GenBank accession NM_182476) encodes isoform a, but there are two additional transcripts (isoforms b and c), which are present at lower levels in cells and encode different proteins [8]. We have previously shown that isoform b, which contains an alternative first exon (exon 1b) and lacks exon 3, is not active [13]. Isoform c (accession NM_182480) differs from isoform a only for the first exon and is still predicted to contain a mitochondrial-targeting sequence, but it is not clear if it encodes an active enzyme (Figure 3(a)). We employed our model to investigate the role of isoform c. Using lentiviral vectors, we expressed either isoform a or isoform c in COQ6KO cells. After selection, cells were incubated in DMEM containing 2 mM glucose for 48 hrs. Only isoform a rescued complex II and III activity (and thus CoQ production), whereas isoform c had no effect (Figure 3(b)). We also checked the subcellular localization of isoform c using a C-terminal GFP fusion construct, analogous to what we had previously employed for isoform a. As seen in Figure 3(c), the fusion protein colocalizes with mitochondria but it could also be an artifact due to overexpression or to the presence of GFP.

3.4. Vanillic Acid Restores CoQ10 Biosynthesis, Cellular Respiration, and ATP Production in COQ6Δ25 Cells. VA differs from 4-HB for the presence of a methoxyl group in position C5 (Figure 4(a)). To test the possibility of bypassing the COQ6 defect in mammalian cells, we incubated different cell types with either CoQ10 or VA. We found that VA restored II + III activity in COQ6Δ25 cells transduced with the G255R point mutant, but also in cells transformed with the empty vector, indicating that contrary to what happens in yeast, rescue is not restricted to inactive but structurally stable alleles but it occurs also with null mutants. CoQ10 supplementation had a similar effect on II + III activity (Figure 4(b)). Direct measurement of CoQ10 levels confirmed this finding (Figure 4(c)). We noted that the G255R mutant displays relatively high
basal II + III activity but this is consistent with the fact that it is a hypomorphic allele [13] and that the lentiviral vector that we are using is probably overexpressing the transgene since it uses the strong CMV promoter. ATP levels were markedly reduced in COQ6Δ25 cells, but after VA supplementation, they were virtually normal (Figure 4(d)). VA treatment could also restore coupled respiration in COQ6Δ25 cells (Figures 4(e) and 4(f)). In untreated cell, this value was about 10% of the wild type, in line with what we reported for COQ4KO cells. The residual respiration is probably due to the presence of a small amount of exogenous CoQ collected from the serum in cell culture media [22].

3.5. VA but Not CoQ10 Normalizes ROS Production in COQ6Δ25 Cells. Because of the role of CoQ as an antioxidant, we studied the effect of VA supplementation on ROS production in COQ6Δ25 cells. In accordance with previously reported data [23], we did not detect a significant increase of mitochondrial ROS production using mitochondrially targeted ro-GFP (Figure 5(a)). Conversely, when we employed a different system, based on the CM-H2DCFDA probe,
which measures total cellular ROS [24], we found increased levels in the COQ6Δ25 cells. VA treatment was able to decrease ROS to basal levels, while CoQ supplementation was only partially effective (Figures 5(b) and 5(c)).

4. Discussion

Bypass therapy is a promising alternative to conventional CoQ supplementation for patients with primary CoQ
deficiency and has been successfully tested in mouse models of \textit{COQ7} and \textit{COQ9} defects \cite{25, 26} and in cells of patients with \textit{COQ7} mutations \cite{27, 28}. In particular, mice with a conditional ablation of the \textit{Coq7} gene (which encodes the C6-hydroxylase catalyzing the penultimate step of CoQ biosynthesis) at 2 months of age developed severe, multiorgan CoQ deficiency, which lead to death after 9 months. If these animals were treated immediately before death with 2,4-dihydroxybenzoate (2,4-DHB), a compound similar to VA which can bypass the Coq7 defect, CoQ biosynthesis was restored and the animals recovered virtually all symptoms and displayed a normal lifespan \cite{26}.

In this work, we have generated a cell line lacking functional \textit{COQ6} using a genome-editing approach. We are aware that this model has some limitations. We obtained an in-frame deletion, which could still produce some folded protein (below the threshold of detectability of our assays), and we could not rule out off-target effects, even though reexpession of the wild-type cDNA corrected the biochemical phenotype of these cells.

Using this model, we showed that VA can rescue CoQ biosynthesis when expressed in \textit{COQ6}\textsubscript{Δ25} cells, suggesting that it does not encode an active protein. This finding is critical for the correct interpretation of genomic analyses in patients. In fact, \textit{COQ6} exon 1b is targeted by most exome analysis kits and the gnomAD database (http://gnomad.broadinstitute.org/gene/ENSG00000119723) lists several possible loss-of-function mutations within this
Decaprenyl-PP can occur even in the absence of COQ6 because carbon C5 of the ring is already bound to a methoxyl group in VA.

The exact order of reactions that carry out the modifications of the aromatic ring of CoQ is still unclear in eukaryotes. Generally accepted models indicate that the C5 hydroxylation catalyzed by COQ6 is the first reaction to occur after the condensation of 4-HB with the polyisoprene tail [36]. However, the fact that COQ6 knockout cells (both human and yeast) accumulate 4-HP, a compound that is decarboxylated and hydroxylated in position C1 of the ring, indicates that these reactions (which are carried out by still unidentified enzymes) occur before or independently on C5 hydroxylation. Furthermore, this result also confirms the notion that C1 hydroxylation is not catalyzed by COQ6 [7]. The identification of the enzymes that catalyze these biosynthetic steps will be critical to validate this hypothesis. Figure 6 depicts a possible model of CoQ biosynthesis in mammals according to our data.

VA was able to restore biosynthesis even in the presence of a null mutant, implying that, contrary to yeast, mammalian cells can assemble the biosynthetic complex even in the absence of COQ6. Therefore, data obtained in yeast about complex assembly and stability should be extrapolated with caution to mammalian cells.

An open issue is whether all COQ6 mutations could be responsive to VA. In the case of COQ7, the response to bypass therapy with 2,4-DHB was more evident when CoQ biosynthesis was severely impaired, rather than when the defect was relatively mild [28]. In the case of COQ6, VA was effective in both situations (CoQ biosynthesis was virtually absent in COQ6Δ25 as in COQ7KO cells, while the biosynthetic defect in cells expressing the G255R mutant was similar to that found in cells with the mild COQ7 [M103T+L111P] allele). This peculiar behavior of COQ7 mutants was attributed to the fact that 2,4-DHB has also an inhibitory action on CoQ biosynthesis [28]. Conversely, VA apparently has no inhibitory effects on the pathway and it is reported to stimulate biosynthesis of other COQ proteins [11]. We have tested the majority of known human mutations in a yeast model, and they all appeared to be responsive to VA [13]. Nevertheless, before attempting to treat patients with VA, one should consider to assay the efficacy of the compound in cell lines (primary fibroblasts or lymphoblastoid cells) obtained directly from the patient.

**Data Availability**

The data used to support the findings of this study are available from the corresponding author upon request.

**Conflicts of Interest**

None of the authors have conflicts of interest to disclose.

**Authors’ Contributions**

Gloria Brea-Calvo and Leonardo Salviati are authors that share senior authorship.

**Acknowledgments**

This work was funded by Fondazione Telethon Grant 14187c (to L.S.), by grants from Fondazione IRP Città della Speranza (to L.S., E.T., and M.C.), by Fondation pour la Recherche...
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