Identification of Coq11, a New Coenzyme Q Biosynthetic Protein in the CoQ-Synthome in *Saccharomyces cerevisiae**^S

Received for publication, December 11, 2014, and in revised form, January 26, 2015 Published, JBC Papers in Press, January 28, 2015, DOI 10.1074/jbc.M114.633131

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Background: Yeast synthesizes coenzyme Q via a macromolecular protein complex.

Results: The Q biosynthetic complex includes Coq8 and the uncharacterized protein YLR290C; the *ylr290c* Δ mutant exhibits impaired Q synthesis.

Conclusion: YLR290C (Coq11) is a novel protein required for efficient yeast Q biosynthesis.

Significance: Discovery and characterization of yeast Coq biosynthetic proteins leads to an improved understanding of coenzyme Q biosynthesis and regulation.

Coenzyme Q (Q or ubiquinone) is a redox active lipid composed of a fully substituted benzoquinone ring and a polyisoprenoid tail and is required for mitochondrial electron transport. In the yeast Saccharomyces cerevisiae, Q is synthesized by the products of 11 known genes, COQ1-COQ9, YAH1, and ARH1. The function of some of the Coq proteins remains unknown, and several steps in the Q biosynthetic pathway are not fully characterized. Several of the Coq proteins are associated in a macromolecular complex on the matrix face of the inner mitochondrial membrane, and this complex is required for efficient Q synthesis. Here, we further characterize this complex via immunoblotting and proteomic analysis of tandem affinity-purified consecutive nondenaturing affinity purificationtagged Coq proteins. We show that Coq8, a putative kinase required for the stability of the Q biosynthetic complex, is associated with a Coq6-containing complex. Additionally Q₆ and late stage Q biosynthetic intermediates were also found to co-purify with the complex. A mitochondrial protein of unknown function, encoded by the YLR290C open reading frame, is also identified as a constituent of the complex and is shown to be required for efficient de novo Q biosynthesis. Given its effect on Q synthesis and its association with the biosynthetic complex, we propose that the open

reading frame YLR290C be designated COQ11.

Fn2

ZSI

Coenzyme Q (ubiquinone or Q)² is a small lipophilic molecule consisting of a hydrophobic polyisoprenoid chain and a fully substituted benzoquinone ring. The length of the polyisoprenoid group is species-specific; contains 6, 8, 9, and 10 isoprenyl units in Saccharomyces cerevisiae, Escherichia coli, mice, and humans, respectively (1); and localizes Q to biological membranes. The primary function of Q is to serve as an electron carrier in respiratory electron transport, where it sequentially accepts up to two electrons from NADH at complex I or succinate at complex II and donates them to cytochrome c at complex III (2). S. cerevisiae lacks complex I and instead uses internal or external NADH:Q oxidoreductases on the inner mitochondrial membrane, which unlike complex I, do not facilitate proton transport across the inner membrane (3). Q also functions as an electron carrier in other aspects of metabolism including dihydroorotate dehydrogenase (pyrimidine synthesis), fatty acyl-CoA dehydrogenase (fatty acid β -oxidation), and other dehydrogenases that oxidize various substrates including choline, sarcosine, sulfide, and glycerol-3-phosphate (4-6). QH₂ has also been demonstrated to function as a lipid-soluble antioxidant capable of mitigating lipid peroxidation and regenerating α -tocopherol (vitamin E) (7, 8). Primary deficiencies in Q biosynthesis have been identified in several human patients and manifest a variety of symptoms including encephalomyopathy, ataxia, cerebellar atrophy, myopathy, and steroid-resistant nephrotic syndrome (9).

In S. cerevisiae Q synthesis is dependent on the product of 11 F1 known genes: COQ1-COQ9, YAH1, and ARH1 (Fig. 1). COQ1-COQ9 were identified as complementation groups of Q-deficient yeast mutants (10, 11), whereas YAH1 and ARH1 were identified through lipid analysis of Yah1- and Arh1-depleted strains under control of galactose-inducible promoters (12). Deletion of any of these COQ genes results in a loss of Q synthesis and consequently a failure to respire and grow on a nonfermentable carbon source (12, 13). Deletion of either YAH1 or ARH1 results in a loss in viability because the biosynthesis of iron-sulfur clusters and heme A requires these gene products



^{*} This work was supported, in whole or in part, by National Institutes of Health Grants S10RR024605 and R01GM103479 (to J. A. L.). This work was also supported by National Science Foundation Grant MCB-1330803 (to C. F. C.), Kirschstein National Research Service Award NIH GM100753 (to C. E. B.-H.), and Grant DE-FC02-02ER63421 from the Office of Science (Biological and Environmental Research), U.S. Department of Energy.

^S This article contains supplemental Data Set 1.

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AO: F ² The abbreviations used are: Q, coenzyme Q; CNAP, consecutive nondenaturing affinity purification; DMQ₆, demethoxy-Q₆; HAB, 3-hexaprenyl-4-aminobenzoic acid; 4HB, 4-hydroxybenzoic acid; HHB, 3-hexaprenyl-4-hydroxybenzoic acid; IDMQ₆, imino-demethoxy-Q₆; MRM, multiple reaction

monitoring; pABA, para-aminobenzoic acid; PC, protein C; DOD, drop out dextrose medium; Ni-NTA, nickel-nitrilotriacetic acid; SDR, short chain dehydrogenase/reductase.



FIGURE 1. **Proposed coenzyme Q biosynthetic pathway in S. cerevisiae.** Yeast utilize either 4HB or pABA to synthesize the first lipid intermediates of the pathway, HHB or HAB, respectively, through the actions of Coq1 and Coq2. These intermediates then undergo a series of ring modifications catalyzed by the enzymes denoted above each step to yield Q. Two steps in the pathway, denoted with *question marks*, remain uncharacterized. The step at which the amino group is converted to the hydroxyl group remains unclear and is denoted by *dashed arrows*. Two late stage intermediates, DMQ₆ and IDMQ₆, are readily detected in yeast. The hexaprenyl moiety is abbreviated as *R*.

(14–18). Although required for Q biosynthesis, the functions of Coq4, Coq8, and Coq9 are not fully understood, although Coq8 has been proposed to function as a protein kinase (19), and Coq9 is required for the replacement of the ring amino group with a hydroxyl group and for the efficient function of Coq6 and Coq7 (20). There are steps in the yeast Q biosynthetic pathway (Fig. 1), such as the decarboxylation reaction and subsequent ring hydroxylation, for which no enzyme has yet been characterized.

Genetic and biochemical experiments have demonstrated the interdependence of several of the Coq proteins and have shown the existence of a high molecular mass Coq protein complex. Each of the $coq_3 - coq_9\Delta$ mutants accumulate only the early intermediates 3-hexaprenyl-4-hydroxybenzoic acid (HHB) and 3-hexaprenyl-4-aminobenzoic acid (HAB), resulting from prenylation of 4-hydroxybenzoic acid (4HB) and paraaminobenzoic acid (pABA), respectively (20). Steady-state levels of Coq3, Coq4, Coq6, Coq7, and Coq9 are reduced in several $coq1-coq9\Delta$ mutants, although levels of Coq3 were stabilized in $coq4-coq9\Delta$ mutant samples prepared with phosphatase inhibitors (20-22). This has been observed in other mitochondrial complexes such as the cytochrome bc_1 complex and ATP synthase in which the absence or mutation of one component protein results in reduced steady-state levels of the other subunits (23, 24). Gel filtration chromatography and blue native-PAGE have shown that several of the Coq proteins exist in high molecular mass complexes, and the O-methyltransferase activity of Coq3 can be detected in these complexes (21, 25–27). Co-precipitation experiments have demonstrated the physical association of several of the Coq proteins; biotinylated Coq3 was shown to co-precipitate Coq4 and HA-tagged Coq9 co-purified Coq4, Coq5, Coq6, and Coq7 (21, 25). Although Coq8 has not been shown to be associated in a complex with the other Coq proteins, its putative role as a kinase is important for the stability of several other Coq proteins because overexpression of *COQ8* in various $coq\Delta$ mutants stabilizes several of the other Coq proteins, as well as multisubunit Coq polypeptide complexes, and leads to the accumulation of later stage Q biosynthetic intermediates diagnostic of the mutated step (20, 28, 29).

Overexpression of *COQ8* in the $coq1\Delta$ or $coq2\Delta$ mutants, which do not produce any polyisoprenylated intermediates, fails to stabilize steady-state levels of sensitive Coq polypeptides (20). When diverse polyprenyl diphosphate synthases from

prokaryotic species that do not synthesize Q are expressed in yeast $coq_1\Delta$ mutants, Q biosynthesis and steady-state levels of sensitive Coq polypeptides are restored (30), suggesting that a polyisoprenylated compound is essential for complex stability. Analysis of gel filtration fractions has also shown the association of demethoxy-Q₆ (DMQ₆) with the complexes (25). Supplementation of exogenous Q₆ was shown to stabilize the steady-state levels of Coq₃ and Coq₄ in the $coq_7\Delta$ mutant (27), enhance DMQ₆ production in the $coq_7\Delta$ mutant (29), and stabilize steady-state levels of Coq₉ in the $coq_3\Delta$, $coq_4\Delta$, $coq_6\Delta$, $coq_7\Delta$ mutants and steady-state levels of Coq₄ in the $coq_3\Delta$, $coq_6\Delta$, and $coq_7\Delta$ mutants (28), suggesting that associated Q₆ plays a role in the stability of the complex.

The functions of certain Coq polypeptides in Q biosynthesis remain unclear, and there are reactions in the Q biosynthetic pathway for which an enzyme has not been assigned. In this study, the composition of the Q biosynthetic complex is investigated through the use of C-terminal tandem affinity purification tags coupled to proteomic analysis. The presence of associated Q_6 and Q_6 intermediates with the complex is also assessed. Two new candidate proteins associated with the complex are characterized via analysis of their null mutants. One of these candidates, YLR290C, was further analyzed by tandem affinity purification and shown to be associated with the Q biosynthetic complex. This observation along with the decrease in *de novo* Q_6 synthesis in the *ylr290c* Δ mutant has led us to propose that YLR290C be designated Coq11.

EXPERIMENTAL PROCEDURES

Yeast Strains and Growth Media—S. cerevisiae strains used in this study are described in Table 1. Medium was prepared as described (31) and included YPD (2% glucose, 1% yeast extract, 2% peptone), YPgal (2% galactose, 0.1% glucose, 1% yeast extract, 2% peptone), and YPG (3% glycerol, 1% yeast extract, 2% peptone). Synthetic dextrose/minimal medium was prepared as described (32) and consisted of all components (SDcomplete) or all components minus histidine (SD–His). Drop out dextrose medium (DOD) was prepared as described (33), except that galactose was replaced with 2% dextrose. Plate medium was prepared with 2% bacto agar.

Construction of Integrated CNAP-tagged Yeast Strains—The coding sequence for the CNAP (consecutive nondenaturing affinity purification) tag, **HHHHHHHHHHHHH**GGAGG<u>EDQVD</u>-

Strain	Genotype	Source or reference	
W303-1A	Mat a ade2-1 his3-1,15 leu2-3,112 trp1-1 ura3-1	R. Rothstein ^a	
W303 $coq8\Delta$	Mat a ade2-1 his3-1,15 leu2-3,112 trp1-1 ura3-1 coq8::HIS3	Ref. 81	
CNAP3	Mat a ade2-1 his3-1,15 leu2-3,112 trp1-1 ura3-1 COQ3::COQ3-CNAP-HIS3	This study	
CNAP6	Mat a ade2-1 his3-1,15 leu2-3,112 trp1-1 ura3-1 COQ6::COQ6-CNAP-HIS3	This study	
CNAP9	Mat a ade2-1 his3-1,15 leu2-3,112 trp1-1 ura3-1 COQ9::COQ9-CNAP-HIS3	This study	
CA-1	Mat a ade2-1 his3-1,15 leu2-3,112 trp1-1 ura3-1 YLR290C::YLR290C-CNAP-HIS3	This study	
BY4741	MAT a $his3\Delta 1 \ leu2\Delta 0 \ met15\Delta 0 \ ura3\Delta 0$	Ref. 82	
BY4741 $coq3\Delta$	MAT a his $3\Delta 1$ leu $2\Delta 0$ met $15\Delta 0$ ura $3\Delta 0$ coq 3 ::kan $MX4$	Ref. 83	
BY4741 $ilv6\Delta$	MAT a his $3\Delta 1$ leu $2\Delta 0$ met $15\Delta 0$ ura $3\Delta 0$ ilv $\hat{6}$::kanMX4	Ref. 84 ^b	
BY4741 $vlr290c\Delta$	MAT a his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 ylr290c::kanMX4	Ref. 84 ^b	

TABLE 1 Genotype and source of yeast str

^a Dr. Rodney Rothstein, Department of Human Genetics, Columbia University.

^b GE Healthcare Yeast Knockout Collection, available online.

<u>PRLIDGK</u> (His₁₀ tag in bold, protein C (PC) epitope underlined) (34) and *HIS3* ORF were amplified from pFACNA-PHISMX (Dr. C. M. Koehler, Department of Chemistry and Biochemistry, UCLA) using primers with 5'-flanking regions corresponding to 50 bp upstream of the stop codon and 50 bp downstream of the stop codon of the gene of interest. Following PCR, the product was purified using the PureLink quick PCR purification kit (Life Technologies) and used to transform W303-1A yeast via the lithium acetate/PEG method (35) to allow integration of the CNAP and *HIS3* coding sequence into the endogenous stop codon of the target gene. Integrants were selected on SD—His and screened by colony PCR to verify the presence of the *CNAP-HIS3* insert at the correct locus; the wildtype ORF lacking the insert was not detected. Primer sequences are available upon request.

AQ: B

Mitochondrial Purification from Yeast—Yeast strains were grown in 5 ml of YPgal precultures and inoculated into 600-ml YPgal cultures for overnight growth in a shaking incubator (30 °C, 250 rpm). Cells were harvested at an A_{600} of 3.5–4.0, and mitochondria were purified as described (36), with the addition of Complete EDTA-free protease inhibitor mixture (Roche) and phosphatase inhibitor cocktails I and II (Calbiochem). Purified mitochondria were flash frozen in liquid nitrogen and stored at -80 °C. Protein concentration was measured with a BCA assay using bovine serum albumin as the standard (Thermo Scientific).

Tandem Affinity Purification of CNAP-tagged Proteins-Purified mitochondria (15 mg of protein) were pelleted by centrifugation at 12,000 \times g for 10 min and solubilized at 2 mg/ml with solubilization buffer (20 mM HEPES-KOH, pH 7.4, 100 mM NaCl, 20 mM imidazole, 10% glycerol, 1 mM CaCl₂, Complete EDTA-free protease inhibitor mixture (Roche), phosphatase inhibitor cocktails I and II (Calbiochem), 6 mg/ml digitonin (Biosynth)) for 1 h on ice with mixing every 10 min. The soluble supernatant fraction was then separated from the insoluble pellet by centrifugation at 100,000 \times g for 10 min in a Beckman Airfuge. The soluble fraction was then mixed with 8 ml of lysis buffer (50 mM NaH₂PO₄, pH 8.0, 300 mM NaCl, 10 mM imidazole) and 800-µl bed volume Ni-NTA resin (Qiagen) pre-equilibrated with lysis buffer and incubated at 4 °C for 90 min with mixing by rotation. The Ni-NTA slurry was then applied to a flowthrough column and washed twice with 12 ml of Ni-NTA wash buffer (50 mм NaH₂PO₄, pH 8.0, 300 mм NaCl, 20 mм imidazole, 1 mg/ml digitonin), followed by elution of bound protein with 8 ml of Ni-NTA elution buffer (20 mM HEPES-

TABLE 2

Description and source of antibodies

Antibody	Working dilution	Source			
Atp2	1:2000	C. M. Koehler ^a			
Coq1	1:10,000	Ref. 30			
Coq2	1:1000	Ref. 21			
Coq3	1:1000	Ref. 85			
Coq4	1:1000	Ref. 86			
Coq5	1:5000	Ref. 87			
Coq6	1:250	Ref. 88			
Coq7	1:500	Ref. 27			
Cog8	(Affinity-purified; 1:30)	Ref. 21			
Coq9	1:1000	Ref. 89			
Coq10	(Affinity-purified; 1:15)	Ref. 90			
Mdĥ1	1:10,000	L. McAlister-Henn ^b			
Protein C	0.7 μg/ml	Antibodies-online Inc.			

^a Dr. C.M. Koehler, Department of Chemistry and Biochemistry, UCLA.
^b Dr. L. McAlister-Henn, Department of Molecular Biophysics and Biochemistry, University of Texas Health Sciences Center, San Antonio, TX.

KOH, pH 7.4, 100 mM NaCl, 300 mM imidazole, 10% glycerol, 1 mM CaCl₂, 1 mg/ml digitonin). The Ni-NTA eluate was applied directly to 1 ml of anti-PC resin (Roche) pre-equilibrated with anti-PC equilibration buffer (20 mM HEPES-KOH, pH 7.4, 100 mM NaCl, 1 mM CaCl₂) and incubated overnight at 4 °C with mixing by rotation. The anti-PC slurry was then applied to a flowthrough column and washed twice with 15 ml of anti-PC wash buffer (20 mM HEPES-KOH, pH 7.4, 100 mM NaCl, 1 mM CaCl₂, 1 mg/ml digitonin). Bound protein was eluted twice with 1 ml of anti-PC elution buffer (20 mM HEPES-KOH, pH 7.4, 100 mM NaCl, 5 mM EDTA, 1 mg/ml digitonin). In the first elution (E1), the anti-PC resin was incubated at 4 °C for 15 min followed by incubation at room temperature for 15 min. In the second elution (E2), the anti-PC resin was incubated for 15 min at room temperature. Eluates were stored at -20 °C.

SDS-PAGE and Immunoblot Analysis—Protein samples incubated with SDS sample buffer (50 mM Tris-HCl, pH 6.8, 10% glycerol, 2% SDS, 0.1% bromphenol blue, 1.33% β -mercaptoethanol) were separated on 12% Tris-glycine SDS-polyacrylamide gels by electrophoresis (37) followed by transfer to Immobilon-P PVDF membranes (Millipore) at 100 V for 1.5 h. Membranes were then blocked overnight in 3% nonfat milk, phosphate-buffered saline (140.7 mM NaCl, 9.3 mM Na₂HPO₄, pH 7.4), 0.1% Tween 20. Membranes were then probed with primary antibodies (Table 2) in 2% nonfat milk, phosphatebuffered saline, 0.1% Tween 20 at the dilutions stated (Table 2). Goat anti-rabbit secondary antibody conjugated to horseradish peroxidase (Calbiochem) was used at 1:10,000 dilutions. Blots were visualized using Supersignal West Pico chemiluminescent substrate (Thermo).



T2



FIGURE 2. Expression of CNAP-tagged Coq3, Coq6, or Coq9 proteins preserves growth on a nonfermentable carbon source and *de novo* Q biosynthesis. *A*, designated yeast strains were grown overnight in 5 ml of YPD, diluted to an A_{600} of 0.2 with sterile PBS, and 2 µl of 5-fold serial dilutions were spotted onto each type of plate medium, corresponding to a final A_{600} of 0.2, 0.04, 0.008, 0.0016, and 0.00032. Plates were incubated at 30 °C, and growth is depicted after 2 days for both YPD and SD – His and 3 days for YPG. *B* and *C*, total ¹³C₆-Q₆ + ¹³C₆-Q₆ + ¹³C₆-Q₆H₂) (*B*) and total ¹²C-Q₆ + ¹³C₆-Q₆H₂) (*C*) were measured in the designated yeast strains by HPLC-MS/MS. Yeast strains were grown overnight in 5 ml of SD-complete diluted to an A_{600} of 0.5 in 50 ml of DD-complete, labeled with either ¹³C₆-4HB or ¹³C₆-pABA at an A_{600} of 0.5, and harvested after 3 h of labeling. Lipid measurements were normalized by the wet weight of extracted cells. Each *bar* represents the mean of four measurements from two biological samples with two injections each. *Error bars* represent standard deviations. Statistical significance was determined with the two-tailed Student's *t* test, and the *lowercase letters* above the *bars* are indicative of statistical significance in *B*, the content of total ¹³C₆-QaBA in CNAP9, CNAP9, and *coq8*Δ was compared with wild type with the corresponding ¹³C₆-labeled precursor; ¹³C₆-Q₆ synthesized from ¹³C₆-PABA in CNAP6 was 120% of wild type, whereas ¹³C₆-Q₆ synthesized from ¹³C₆-PABA labeling compared with type under ¹³C₆-4HB labeling, 120% in CNAP6 under ¹³C₆-PABA labeling compared with wild type with the corresponding ¹³C₆-labeled precursor; accumulated ¹²C-Q₆ in CNAP6 was 80% of wild type under ¹³C₆-4HB labeling, 120% in CNAP6 under ¹³C₆-PABA labeling compared with wild type with the corresponding ¹³

SYPRO Ruby Staining—Protein samples incubated with SDS sample buffer were separated on an 8–16% Criterion SDS-PAGE gel (Bio-Rad). Following electrophoresis, the gel was washed twice with water and fixed for 1 h with 100 ml of 10% methanol, 7% acetic acid. The gel was washed twice with water and then stained overnight at room temperature in the dark with 80 ml of a 1:1 mixture of fresh and used SYPRO Ruby (Life Technologies). Following staining, the gel was washed twice with water and destained for 4 h with 10% methanol, 7% acetic acid. Proteins were visualized with an FX Pro Plus Molecular Imager (Bio-Rad) with 532-nm excitation, and emission was measured with a 555-nm long pass filter.

In-gel Trypsin Digestion and Proteomic Analysis—Sample lanes from a SYPRO Ruby-stained gel were cut into 5-mm slices and subjected to in-gel trypsin digestion as described previously (38), with a few modifications. Gel slices were washed with 50 mM NH₄HCO₃, 50% acetonitrile followed by 100% acetonitrile, three alternating cycles total. Samples were then incubated with 10 mM dithiothreitol (MP Biomedicals) at 60 °C for 1 h to reduce disulfide bonds, followed by treatment with 50 mM iodoacetamide (Calbiochem) at 45 °C for 45 min in the dark to alkylate-free sulfhydryl groups. Gel slices were then washed once with 50 mM NH₄HCO₃, followed by three alternating washes with 100 mM NH₄HCO₃ and 100% acetonitrile. Gel slices were then dried by vacuum centrifugation (SpeedVac) at 60 °C for 10 min. Gel-entrapped proteins were then digested overnight at 37 °C using 20 ng/ μ l trypsin (Promega) in 50 mM NH₄HCO₃. Following digestion tryptic peptides were extracted from the gel slices with 50% acetonitrile, 0.1% TFA and dried by SpeedVac at 30 °C for 2 h.

LC-MS/MS analysis of the trypsin-digested peptides was performed as follows: Peptides were resuspended in 30 μ l of 3% acetonitrile, 0.1% formic acid. The peptides were measured using a nanoACQUITY UPLC system coupled to a Xevo QTof (quadrupole time of flight) mass spectrometer (Waters). Peptides were eluted from the UPLC to the electrospray ionization mass spectrometer using a 5-µm 2G-VM Symmetry C18 180- μ m \times 20-mm trap column in-line with a 1.8- μ m HSS T3, 75- μ m \times 150-mm C18 analytical column (Waters). The digested peptides were separated using a 0-60-min gradient, beginning at 3% acetonitrile, 0.1% formic acid to 40% acetonitrile, 0.1% formic acid, followed by a 40-95% gradient between 60 and 62 min at a flow rate of 0.3 μ l min⁻¹ (90-min total run time). [Glu¹]-Fibrinopeptide B was used as a mass calibration standard (100 fmol/µl) and was infused via a separate electrospray ionization sprayer, and the standard peptide was measured every 45 s during data collection.



The mass spectrometer was operated in the MS^E data-independent acquisition mode (39-41). ProteinLynx Global Server (version 3; Waters) was used for protein identification. The MS data were searched against the Uniprot S. cerevisiae (strain ATCC 204508/S288c) reference proteome data set (release date July 9, 2014, containing both Swiss-Prot (6718) and TrEMBL (9) entries). The database was further supplemented with sequences for keratin and trypsin. Low energy, elevated energy, and intensity thresholds were set to 250/100/1000 counts, respectively. The MS data files were searched following ProteinLynx Global Server workflow parameters: trypsin digestion with allowed missed cleavages (2), peptide and fragment tolerances set automatically (~11 ppm and 28 ppm, respectively), minimum number of peptide matches (1), minimum number of product ions per peptide (3), minimum number of product ions per protein (7), and maximum false positive rate (4%). Carbamidomethylation of Cys residues was set as a fixed modification, and Met oxidation, Ser, Thr, and Tyr phosphorylation, and Cys propionamidation resulting from iodoacetamide treatment were set as variable modifications.

AO: C

Metabolic Labeling of Q_6 with ${}^{13}C_6$ -labeled Precursors— Yeast strains were grown overnight in 5 ml of SD-complete and diluted to an A_{600} of 0.05 in 50 ml of DOD-complete. Cultures were grown to an A_{600} of 0.5, corresponding to early log phase and then labeled with either ${}^{13}C_6$ -4HB or ${}^{13}C_6$ -pABA at a concentration of 5 μ g/ml (250 μ g total). Cells were grown an additional 3 h with label, harvested by centrifugation at 2000 × g for 10 min, and stored at -20 °C. Wet weights were determined for yeast cell pellets.

Analysis of Q_6 and Q_6 Intermediates—Lipid extraction of eluate samples and cells grown in DOD-complete was performed with methanol/petroleum ether and Q_4 as an internal standard, and LC-MS/MS analysis of extracts was performed as described (33). Briefly, a 4000 QTRAP linear MS/MS spectrometer (Applied Biosystems, Foster City, CA) was used, and Analyst version 1.4.2 software (Applied Biosystems) was used for data acquisition and processing. A binary HPLC delivery system was used with a phenyl-hexyl column (Luna 5u, 100×4.60 mm, 5- μ m; Phenomenex), with a mobile phase consisting of solvent A (methanol/2-propanol, 95:5, 2.5 mM ammonium formate) and solvent B (2-propanol, 2.5 mM ammonium formate). The percentage of solvent B was increased linearly from 0 to 5% over 6 min, and the flow rate was increased from 600 to 800 μ l/min. The flow rate and mobile phase were changed back to initial conditions linearly over 7 min. All samples were analyzed in multiple reaction monitoring mode. Multiple reaction monitoring mode precursor to product ion transitions were detected as follows: ¹³C₆-HAB m/z 552.4/156.0, ¹²C-HAB m/z 546.4/ 150.0, ¹³C₆-HHB *m*/*z* 553.4/157.0, ¹²C-HHB *m*/*z* 547.4/151.0, IDMQ₆ *m*/*z* 560.6/166.0, ¹³C₆-DMQ₆ *m*/*z* 567.6/173.0, ¹²C-DMQ₆ m/z 561.6/167.0, ${}^{13}C_6$ -Q₆ m/z 597.4/203.0, ${}^{13}C_6$ -Q₆H₂ *m*/*z* 616.4/203.0, ¹²C-Q₆ *m*/*z* 591.4/197.0, and ¹²C-Q₆H₂ *m*/*z* 610.4/197.0.

Bioinformatic Analyses—Evolutionary relationships of amino acid sequences were inferred using the neighbor-joining method (42) with MEGA6 (43). The protein similarity network (44) was constructed as described previously (45) from all-*ver-sus*-all blastp alignments using an *E* value cutoff of 1×10^{-21} .



FIGURE 3. **CNAP-tagged Coq proteins co-precipitate several other Coq proteins.** W303, CNAP3, CNAP6, and CNAP9 purified mitochondria (15 mg of protein) were solubilized with digitonin and subjected to tandem affinity purification with Ni-NTA resin (Qiagen) followed by anti-PC-agarose (Roche). Samples were separated on 12% SDS-PAGE gels followed by transfer to PVDF membranes for immunoblotting with antisera to the designated yeast polypeptides. 25 μ g of mitochondria protein were analyzed for each strain (*M*), and 2.5% of the first anti-PC elution (*E1*) volume was loaded per strain (25 μ). *Arrows* denote each tagged protein in their respective blots. The predominant band in the Coq3 blot detected in the mitochondrial samples represents a background protein and not Coq3, accounting for its presence in CNAP3 mitochondria.

Protein IDs analyzed are available in supplemental Data Set 1. Protein sequences were retrieved through BLAST of the Uniprot database (46). The multiple sequence alignment was built with MUSCLE (47) and visualized with EsPript (48).

RESULTS

Effects of CNAP Tag Integration on Respiration and Q Biosynthesis—The CNAP tag was integrated into the 3' end of the COQ3, COQ6, and COQ9 ORFs, generating the CNAP3, CNAP6, and CNAP9 strains, respectively. The ability of these strains to respire as measured by growth on a nonfermentable carbon source was assessed by plate dilution assay, using W3031B as the wild-type control and $coq8\Delta$ as the Q-less, respiratory defective control (Fig. 2A). All three tagged strains F2 grew as well as wild type on the nonfermentable carbon source, YPG, whereas the Q-less mutant failed to grow. The three tagged strains and the Q-less mutant were generated by chromosomal integration of the *HIS3* ORF and grew on SD—His, whereas the wild type did not. All five strains were able to grow on the permissible carbon source YPD.

De novo Q biosynthesis was measured by LC-MS/MS of cells grown in DOD-complete medium with the stable isotopically



labeled ring precursors $^{13}C_6$ -4HB or $^{13}C_6$ -pABA (Fig. 2*B*). Levels of *de novo* $^{13}C_6$ -Q in CNAP3 synthesized from either precursor were not significantly different compared with wild type. $^{13}C_6$ -Q synthesized *de novo* from $^{13}C_6$ -4HB in CNAP6 and

 $^{13}\mathrm{C}_6\text{-}\mathrm{Q}$ synthesized *de novo* from $^{13}\mathrm{C}_6\text{-}\mathrm{pABA}$ in CNAP9 were not significantly different compared with wild type. $^{13}\mathrm{C}_6\text{-}\mathrm{Q}$ synthesized *de novo* from $^{13}\mathrm{C}_6\text{-}\mathrm{pABA}$ in CNAP6 was higher than wild type, and $^{13}\mathrm{C}_6\text{-}\mathrm{Q}$ synthesized *de novo* from $^{13}\mathrm{C}_6\text{-}4\mathrm{HB}$



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in CNAP9 was lower than wild type. The $coq8\Delta$ strain did not produce detectable *de novo* ¹³C₆-Q₆. Total Q accumulation was also measured by analyzing unlabeled ¹²C-Q₆ by LC-MS/MS (Fig. 2*C*). Levels of accumulated Q₆ in CNAP3 grown with either precursor were not significantly different compared with wild type. The accumulated Q₆ in CNAP9 grown with ¹³C₆pABA was not significantly different compared with wild type. Accumulated Q₆ in CNAP6 was lower than wild type under ¹³C₆-4HB labeling and higher under ¹³C₆-pABA labeling when compared with wild type. Accumulated Q₆ in CNAP9 labeled with ¹³C₆-4HB was lower than wild type. The *coq8*Δ strain did not accumulate detectable Q₆. The results indicate that CNAPtagged Coq3, Coq6, or Coq9 polypeptides expressed from the integrated loci remain functional.

Co-precipitation of Coq Proteins with CNAP-tagged Coq Proteins-Solubilized mitochondria from W303, CNAP3, CNAP6, and CNAP9 were subjected to tandem affinity purification of the CNAP tag. Purified mitochondria (25 μ g of protein) and fractions of the first anti-PC eluate (E1) were analyzed by immunoblot to identify interacting Coq proteins (Fig. 3). Membranes were probed with antibodies to Coq1-Coq10, the protein C epitope to identify the tagged protein, and Atp2 and Mdh1 to assess the purity of the eluates. CNAP3 co-precipitated Cog4, Cog5, Cog6, Cog7, and Cog9; CNAP6 co-precipitated Coq4, Coq5, Coq7, Coq8, and Coq9; and CNAP9 co-precipitated Coq4, Coq5, Coq6, and Coq7. The signals for Coq5 and Coq9 were much weaker with CNAP3 and CNAP6 compared with CNAP9. The predominant band in the Cog3 blot is a nonspecific band recognized by the antisera at the same molecular mass as Coq3 and is present in the $coq3\Delta$, accounting for the apparent wild-type Coq3 band in the CNAP3 mitochondrial protein (M) sample; the primary purpose of the Coq3 blot is to visualize the shifted band in the CNAP3 E1 sample. All three tagged proteins are present in their respective blots at shifted molecular masses because of the tag (and are denoted by arrows in Fig. 3). The three tagged proteins are also present on the anti-PC blot at their appropriate molecular masses. No signal is visible in the eluate fractions for the nonspecific mitochondrial proteins Atp2 and Mdh1, which are 2-3 orders of magnitude more abundant than the Coq proteins, indicating the specificity of the co-precipitation.

F3

F4

Identification of Associated Lipids and Proteins by Mass Spectrometry—Aliquots of anti-PC eluates (29% E1 and 29% E2) were combined for each strain and subjected to lipid extraction (Fig. 4). Extracts were then analyzed by LC-MS/MS for Q_6 and Q_6 biosynthetic intermediates. The CNAP3, CNAP6, and CNAP9 eluates were all found to have significantly higher levels of associated Q_6 compared with wild type (Fig. 4A). The late stage intermediate DMQ₆ was also measured and found to be significantly higher in CNAP3, CNAP6, and CNAP9 compared



FIGURE 5. SYPRO Ruby staining for total protein reveals unique bands in CNAP tagged eluates. For each strain designated, aliquots of the anti-PC eluates (20% of E1 and 20% of anti-PC eluate two (E2)) were combined and dried with a SpeedVac at 60 °C for 2 h. Samples were resuspended in 42 μ l of SDS sample buffer and separated with an 8–16% Criterion SDS-PAGE gel (Bio-Rad). Following separation proteins were fixed, and the gel was stained overnight with a 1:1 mixture of fresh and used SYPRO Ruby (Life Technologies). The gel was subsequently washed and visualized with an FX Pro Plus Molecular Imager (Bio-Rad) at 532-nm excitation, and emission was measured with a 555-nm long pass filter. The *ladder* denotes protein masses in kDa.

with wild type (Fig. 4*C*). The late stage intermediate iminodemethoxy- Q_6 (IDM Q_6) was also found in measurable quantities in the CNAP3, CNAP6, and CNAP9 eluates, whereas no IDM Q_6 was detected in the wild-type eluate (Fig. 4*E*). Representative HPLC traces of Q_6 , DM Q_6 , and IDM Q_6 are shown (Fig. 4, *B*, *D*, and *F*). The early intermediates HHB and HAB were not detectable in any of the analyzed eluates (data not shown).

Aliquots of the anti-PC eluates (20% E1 and 20% E2) were combined for each strain, concentrated, and resuspended in 42 μ l of SDS sample buffer. Protein was then separated on an 8-16% SDS-PAGE gel and stained with SYPRO Ruby to visualize total protein (Fig. 5). CNAP3, CNAP6, and CNAP9 each contain several unique protein bands compared with wild type, specifically at 25-37 and 50 kDa. Each sample lane was cut into 5-mm slices and subjected to in-gel trypsin digestion. Tryptic peptides were analyzed by LC-MS/MS, and the results from five trials (performed for W303, CNAP3, and CNAP9) or two trials (performed for CNAP6) are shown in Table 3. The results are T3 ranked by sequence coverage of detected peptides and also depict the number of peptides recovered, which represent the total number of peptides detected for a given polypeptide across all trials. Several of the Coq proteins observed by immunoblotting were detected by mass spectrometry of the CNAP3, CNAP6, and CNAP9 eluates and are highlighted. Multiple incidents of recovery of Ilv6 in the CNAP6 and CNAP9 eluates and YLR290C in CNAP3 eluates seemed particularly noteworthy and were investigated further.

Analyses of the $ilv6\Delta$ and $ylr290c\Delta$ Mutants—Because of the high incidence of recovery of the Ilv6 and YLR290C polypep-



FIGURE 4. **Tandem affinity-purified Coq complexes co-purify with Q₆ and late-stage Q₆ intermediates.** For each strain, aliquots of the anti-PC eluates (29% of E1 and 29% of E2) were combined and subjected to lipid extraction with methanol and petroleum ether. Lipid extracts were analyzed by HPLC-MS/MS for Q₆ (*A*), DMQ₆ (*C*), and IDMQ₆ (*E*). Measured lipids were normalized by the extracted eluate volume. *Bars* represent the means of two measurements, and *error bars* represent the standard deviation. Statistical significance was determined with the two-tailed Student's t test, and the *lowercase letters* above the *bars* are indicative of statistical significance. In *A*, the content of Q₆ in CNAP3, CNAP6, and CNAP9 was 11.7-, 8.6-, and 22.6-fold higher, respectively, compared with wild type (*a*, *p* = 0.0001; *b*, *p* = 0.0006). In *C*, the content of DMQ₆ in CNAP3, CNAP6, and CNAP9, and CNAP9 was 30-, 24-, and 97-fold higher, respectively, compared with wild type (*a*, *p* = 0.0110; *b*, *p* = 0.0113; *c*, *p* = 0.0010). In *E*, there was no detectable IDMQ₆ in the wild type. *ND*, not detected. Representative overlaid traces of all four strains (W303, *purple*; CNAP3, *green*; CNAP6, *red*; CNAP9, *blue*) are shown for Q₆ (*B*), DMQ₆ (*D*), and IDMQ₆ (*F*).

TABLE 3

Top protein hits from proteomic analysis of tandem affinity purification eluates

Only proteins that were observed in duplicate injections of samples were considered top hits. Results are compiled from five replicates (W303, CNAP3, and CNAP9) or from two replicates (CNAP6).

	W303			CNAP3			CNAP6			CNAP9	
Proteir	Function (MW)	Sequence coverage (#pep)	Protein	Function (MW)	Sequence coverage (#pep)	Protein	Function (MW)	Sequence coverage (#pep)	Protein	Function (MW)	Sequence coverage (#pep)
Rga2	GTPase-activating protein (113 kDa)	50.3% (130)	Htb1	Histone H2B (14.3 kDa)	79.4% (27)	Coq6	Q biosynthesis (53.5 kDa)	41.5% (102)	Coq6	Q biosynthesis (53. 5 kDa)	54.1% (251)
Pfd4	Chaperone prefolding complex (15.2 kDa)	43.4% (7)	Coq4⁵	Q biosynthesis (38.6 kDa)	73.1% (166)	llv6	Regulatory subunit of acetolactate synthase (34.0 kDa)	35.9% (14)	Caj1	Heat shock protein (44.9 kDa)	48.6% (23)
Act1	Actin (41.7 kDa)	42.1% (40)	Coq3	Q biosynthesis (36.3 kDa)	56.4% (113)	Dld3	D-lactate dehydrogenase (55.2 kDa)	24.4% (12)	Coq5	Q biosynthesis (34.7 kDa)	37.8% (62)
llv6	Regulatory subunit of acetolactate synthase (34.0 kDa)	22.7% (8)	Coq7	Q biosynthesis (26.1 kDa)	52.4% (31)	Hsp60	Heat shock protein (60.8 kDa)	23.3% (17)	Mrp21	Mitochondrial ribosome (20.4 kDa)	37.3% (7)
Flo8	Transcription factor (86.8 kDa)	16.6% (25)	YLR290	C Unknown (31.5 kDa)	48.4% (73)	Coq9	Q biosynthesis (30.0 kDa)	19.6% (13)	llv6	Regulatory subunit of acetolactate synthase (34.0 kDa)	37.2% (40)
Spt16	Associated with chromatin (119 kDa)	2.0% (9)	Ssa1	Heat shock protein (69.7 kDa)	48.0% (27)	SId3	Initiation of DNA replication (77.3 kDa)	17.2% (9)	Coq7	Q biosynthesis (26.1 kDa)	30.5% (11)
			Coq5	Q biosynthesis (34.7 kDa)	46.9% (75)	Coq5	Q biosynthesis (34.7 kDa)	14.7% (6)	Coq4	Q biosynthesis (38.6 kDa)	29.9% (33)
			Ssa3	Heat shock protein (70.5 kDa)	46.5% (34)	His6	Histidine biosynthesis (29.6 kDa)	14.6% (4)	Coq9	Q biosynthesis (30.0 kDa)	29.2% (49)
			Coq6	Q Biosynthesis (53.5 kDa)	45.9% (121)	Mmt1	Metal transporter (56.2 kDa)	14.5% (8)	Rga2	GTPase-activating protein (113 kDa)	24.2% (31)
			Tim11	ATP synthase (10.9 kDa)	45.8% (6)	Npl4	Regulation of polyubiquitinated proteins (65.8 kDa)	12.9% (9)	Mmt2	Metal transporter (52.4 kDa)	18.8% (20)
			Ssa2	Heat shock protein (69.5 kDa)	44.4% (26)	Mmt2	Metal transporter (52.4 kDa)	12.8% (5)	Msh1	DNA binding protein (109 kDa)	12.4% (15)
			Bmh1	14-3-3 protein (30.1 kDa)	41.9% (17)	Yhm2	Citrate/oxoglutarate carrier (34.2 kDa)	7.6% (3)	Mmt1	Metal transporter (56.2 kDa)	10.8% (7)
			Ssa4	Heat shock protein (69.7 kDa)	37.1% (30)	Ura2	Pyrimidine biosynthesis (245 kDa)	2.2% (10)		()	
			Act1	Actin (41. 7 kDa)	35.7% (18)						
			Bmh2	14-3-3 protein (31.1 kDa)	34.1% (15)						
			Coq9	Q biosynthesis (30.0 kDa)	32.3% (36)						
			Sas4	Histone acetylation (55.4 kDa)	23.5% (16)						
			Mrp21	Mitochondrial ribosome (20.4 kDa)	21.5% (4)						
			Rga2	GTPase-activating protein (113 kDa)	20.5% (33)						
			Por1	Porin (30.4 kDa)	19.8% (10)						
			llv6	Regulatory subunit of acetolactate synthase (34.0 kDa)	18.8% (8)						
			Pef1	Polar bud growth (38.3 kDa)	14.0% (9)						
			Mas2	Mitochondrial processing protease (53.3 kDa)	13.3% (8)						
			Hpc2	Nucleosome assembly complex (67.8 kDa)	9.9% (20)						

^{*a*} Highlighted hits denote proteins confirmed by immunoblotting.



FIGURE 6. Yeast *ilv6* and *ylr290c* mutants retain the ability to grow on a nonfermentable carbon source. Designated yeast strains were grown overnight in 5 ml of YPD and diluted to an A_{600} of 0.2 with sterile PBS, and 2 μ l of 5-fold serial dilutions were spotted onto each type of plate medium, corresponding to a final A_{600} of 0.2, 0.04, 0.008, 0.0016, and 0.00032. Plates were incubated at 30 °C, and growth is depicted after 2 days for YPD and 3 days for YPG.

tides in the CNAP-tagged Coq protein eluates (Table 3), the ilv6 Δ and ylr290c Δ mutants in the BY4741 background were purchased from the GE Healthcare yeast knock-out collection for phenotypic analysis. The ability of these null mutants to grow on a nonfermentable carbon source was assessed by plate dilution assay using BY4741 as the wild-type control and coq3 Δ as the Q-less, mutant control (Fig. 6). Both the ilv6 Δ and ylr290c Δ mutants grew as well as wild type on the nonfermentable carbon source YPG, whereas the coq3 Δ mutant failed to grow. All strains grew on the permissible carbon source YPD.

The levels of *de novo* ¹³C₆-Q₆ biosynthesis, as well as accumulation of Q₆ and Q₆ intermediates, were measured by LC-MS/MS of yeast cultures grown in DOD-complete medium labeled with the stable isotopically labeled precursors ¹³C₆-4HB or ${}^{13}C_6$ -pABA (Fig. 7). The *ilv*6 Δ mutant produced high levels of de novo ¹³C₆-HAB and ¹³C₆-HHB relative to wild type, similar to the $coq3\Delta$ mutant (Fig. 7, A and C). The *ilv*6 Δ mutant also accumulated higher levels of ¹²C-HAB and ¹²C-HHB compared with wild type (Fig. 7, *B* and *D*). The *ilv* Δ mutant had increased levels of the late stage intermediate ¹³C₆-DMQ₆ when labeled with ¹³C₆-pABA (Fig. 7E), and accumulated higher levels of $^{12}\text{C-DMQ}_6$ relative to wild type when labeled with $^{13}\text{C}_6\text{-pABA}$ (Fig. 7*F*). The late stage intermediate ${}^{13}C_{6}$ - or ${}^{12}C$ -IDMQ₆ was not detected in this analysis (data not shown). The $ilv6\Delta$ mutant produced decreased *de novo* ¹³C₆-Q₆ compared with wild type when labeled with ¹³C₆-4HB (Fig. 7G) and accumulated decreased 12 C-Q₆ relative to wild type when labeled with 13 C₆-4HB (Fig. 7H). ILV6 encodes the regulatory subunit of acetolactate synthase; Ilv2 catalyzes this committed step of branched chain amino acid biosynthesis in yeast mitochondria (49). Ilv6 is not an essential protein and is not required for the catalytic activity of Ilv2 but regulates Ilv2 through feedback inhibition by valine, a final product of the branched chain amino acid biosynthetic pathway, an effect reversed by binding of ATP to Ilv6 (49, 50). There are functional homologs of Ilv6 in prokaryotes and

F6







FIGURE 8. **Expression of CNAP-tagged YLR290C preserves** *de novo* **Q biosynthesis.** Levels of total ¹³C₆-Q₆ ($^{13}C_6$ -Q₆ + $^{13}C_6$ -Q₆H₂) (*A*) and total ¹²C-Q₆ (^{12}C -Q₆ + ^{12}C -Q₆H₂) (*B*) were measured in the designated yeast strains by HPLC-MS/MS. Yeast strains were grown overnight in 5 ml of SD-complete, diluted to an A₆₀₀ of 0.05 in 50 ml of DOD-complete, labeled with either ¹³C₆-4HB or ¹³C₆-pABA at an A₆₀₀ of 0.5, and harvested after 3 h of labeling. Lipid measurements were normalized by the wet weight of extracted cells. Each *bar* represents the mean of four measurements from two biological samples with two injections each. *Error bars* represent standard deviations. Statistical significance was determined with the two-tailed Student's *t* test, and the *lowercase letters* above the *bars* are indicative of statistical significance. In *A*, the content of ¹³C₆-Q₆ in CA-1 and *coq8*Δ was compared with wild type with the corresponding ¹³C₆-labeled precursor; *b*, *p* = 0.0196). *ND*, not detected. In *B*, the content of ¹²C-Q₆ in CA-1 and *coq8*Δ was compared with wild type with the corresponding ¹³C₆-labeled precursor; accumulated ¹²C-Q₆ in CA-1 was 70% of wild type and 80% of are optimaled with wild type with the corresponding ¹³C₆-labeled precursor; accumulated ¹²C-Q₆ in CA-1 was 70% of wild type when labeled with either ¹³C₆-pABA (*a*, *p* = 0.0202; *b*, *p* = 0.0001). *ND*, not detected.

plants but not other eukaryotes, making this regulatory scheme unique to prokaryotes, fungi, and plants (49, 51, 52).

The *ylr290c* Δ mutant synthesized high levels of *de novo* ¹³C₆-HAB and ${}^{13}C_6$ -HHB compared with wild type (Fig. 7, A and C) and also accumulated higher levels of ¹²C-HAB and ¹²C-HHB compared with wild type (Fig. 7, B and D). The late stage intermediate ¹³C₆- or ¹²C-IDMQ₆ was not detected in this analysis (data not shown); however, the $ylr290c\Delta$ mutant labeled with ¹³C₆-pABA was found to have reduced levels of ¹³C₆-DMQ₆ relative to the wild type (Fig. 7*E*). In contrast, the *ylr290c* Δ mutant labeled with either precursor accumulated higher levels of ¹²C-DMQ₆ relative to wild type (Fig. 7F). The $ylr290c\Delta$ mutant produced severely decreased de novo ¹³C₆-Q₆ when labeled with either ${}^{13}C_{6}$ -4HB or ${}^{13}C_{6}$ -pABA (Fig. 7G) and also accumulated significantly less ¹²C-Q₆ when labeled with ¹³C₆-4HB or ${}^{13}C_6$ -pABA compared with the wild type (Fig. 7*H*). These results indicate that the $ylr290c\Delta$ mutant accumulates higher levels of Q₆ intermediates and has defective Q synthesis compared with wild type. Because of its dramatic effect on Q synthesis, we chose to focus our efforts on further characterization of YLR290C.

Analysis of CNAP-tagged YLR290C—The CNAP tag was integrated into the 3' end of the YLR290C open reading frame in W303-1A to yield the strain designated CA-1. The levels of *de novo* and accumulated Q_6 were measured in this strain to assess the impact of the tag on Q_6 biosynthesis (Fig. 8). The CA-1 strain produced modestly reduced levels of *de novo* ${}^{13}C_6$ - Q_6 with both ${}^{13}C_6$ -4HB and ${}^{13}C_6$ -pABA. Levels of accumulated ${}^{12}C$ - Q_6 were also decreased in CA-1 when labeled with either ${}^{13}C_6$ -4HB or ${}^{13}C_6$ -pABA.

Solubilized CA-1 mitochondria (15 mg of protein) were subjected to tandem affinity purification of the CNAP tag (Fig. 9). F9 Purified mitochondria (25 μ g of protein) and fractions of the first anti-PC eluate (E1) were analyzed by immunoblotting to identify interacting Coq proteins. Membranes were probed with antibodies to Coq1–Coq9, the protein C epitope to identify the tagged protein, and Atp2 and Mdh1 to assess the purity of the eluates. Coq4, Coq5, and Coq7 were observed to co-precipitate with YLR290C-CNAP in the CA-1 strain, whereas the nonspecific mitochondrial proteins Atp2 and Mdh1 were not present in the eluate. Although yielding the weakest signal, Coq7 was observed in the CA-1 eluate in independent blots.



FIGURE 7. **The yeast** *ylr290c* **mutant**, **but not the** *ilv6* **mutant**, **shows impaired** *de novo* **Q biosynthesis**. ¹³C₆-HAB (*A*), ¹²C-HAB (*B*), ¹³C₆-HHB (*C*), ¹²C-HHB (*D*), ¹³C₆-DMQ₆ (*F*), total ¹³C₆-Q₆ (¹³C₆-Q₆ + ¹³C₆-Q₆+Q₁) (*G*), and total ¹²C-Q₆ (¹²C-Q₆ + ¹²C-Q₆+Q₁) (*H*) were measured in the designated yeast strains by HPLC-MS/MS. Yeast strains were grown overnight in 5 ml of SD-complete, diluted to an A_{600} of 0.05 in 50 ml of DDD-complete, labeled with either ¹³C₆-4HB or ¹³C₆-pABA at an A_{600} of 0.5, and harvested after 3 h of labeling. Lipid measurements were normalized by the wet weight of extracted cells. Each *bar* represents the mean of four measurements from two biological samples with two injections each. *Error bars* represent standard deviations. Statistical significance was determined with the two-tailed Student's *t* test, and the *lowercase letters* above the *bars* are indicative of statistical significance. In *A*, the relative content of ¹³C₆-HAB in the three null mutants was compared with wild type with the corresponding ¹³C₆-labeled precursor (*a*, *p* = 0.0001; *b*, *p* = 0.0001; *b*, *p* = 0.0003; *c*, *p* = 0.0011; *b*, *p* < 0.0003; *c*, *p* = 0.0001; *b*, *p* = 0.0002; *b*, *p* = 0.0004; *c*, *p* = 0.0041; *b*, *p* < 0.0001). ND, not detected. In *F*, the relative content of ¹³C₆-labeled precursor (*a*, *p* = 0.0043; *b*, *p* = 0.0002; *b*, *p* = 0.0002; *b*, *p* = 0.0004; *c*, *p* = 0.0041; *b*, *p* < 0.0001). ND, not detected. In *G*, the content of ¹³C₆-Q₆ in the three null mutants was compared with wild type with the corresponding ¹³C₆-labeled p

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FIGURE 9. **Coq4, Coq5, and Coq7 co-precipitate with YLR290C-CNAP.** Purified mitochondria from W303 and CA-1 (15 mg of protein) were solubilized with digitonin and subjected to tandem affinity purification using Ni-NTA resin (Qiagen) followed by anti-PC-agarose (Roche). Samples were separated on 12% SDS-PAGE gels followed by transfer to PVDF membranes for immunoblotting. Mitochondria (25 μ g of protein) (*M*) and 2.5% of the first anti-PC elution (*E1*) were analyzed for each of the two strains.

The anti-PC blot verifies the presence of the CNAP-tagged protein.

Aliquots of anti-PC eluates (29% E1 and 29% E2) were combined for the wild-type and CA-1 strains and subjected to lipid extraction (Fig. 10). Extracts were then analyzed by LC-MS/MS for Q_6 and Q_6 biosynthetic intermediates. The CA-1 eluate was found to have higher levels of associated Q_6 compared with wild type (Fig. 10*A*). The late stage intermediate DMQ₆ was also measured and found to be higher in CA-1 relative to wild type (Fig. 10*C*). The late stage intermediate IDMQ₆ was also found in measurable quantities in the CA-1 eluate, whereas no IDMQ₆ was detected in the wild-type eluate (Fig. 10*E*). Representative traces of Q_6 , DMQ₆, and IDMQ₆ are shown (Fig. 10, *B*, *D*, and *F*). The early intermediates HHB and HAB were not detectable in any of the analyzed eluates (data not shown).

In Silico Analysis of YLR290C and Related Proteins—Based on RPS-BLAST to the NCBI Conserved Domain Database (53), YLR290C and closely related proteins (referred to as YLR290Clike) belong to subgroup 5 of the atypical short chain dehydrogenase/reductase (SDR) superfamily. A protein similarity network reveals that YLR290C-like proteins can be divided into three distinct clusters that are delineated largely by taxonomy with the NDUFA9 subfamily of SDR proteins representing the closest distinct similarity cluster (Fig. 11). YLR290C-like proteins were commonly found in fungal genomes and either photosynthetic organisms such as land plants and algae (green algae, stramenopiles, and *Guillardia theta*) or in the genomes of

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organisms that may have a photosynthetic ancestor such as the alveolates. The genome of the land plant *Arabidopsis thaliana* encodes three homologs of YLR290C; based on proteomics data, the ortholog of YLR290C, AT1G32220, is likely localized to the chloroplast and specifically to plastoglobules of the plastid, which are lipid bodies that contain a large amount of prenylquinones among other lipidic compounds (54). We were intrigued to find that YLR290C-like proteins in the fungi *Ustilago maydis, Ustilago hordei, Pseudozyma flocculosa, Pseudozyma antarctica, Sporisorium reilianum, Pseudozyma hubeiensis, Pseudozyma aphidis*, and *Melanopsichium pennsylvanicum* were fused to Coq10 (Figs. 11 and 12), suggesting a functional link between YLR290C-like proteins and Coq10.

DISCUSSION

This study characterized the composition of the Q biosynthetic complex (termed the CoQ-synthome) in yeast mitochondria. We used tandem affinity purification of C-terminally tagged Coq polypeptides. The CNAP tag was used to purify the complex from digitonin-solubilized mitochondria under gentle conditions to preserve the noncovalent associations between protein constituents of the complex to allow for identification by both immunoblotting and mass spectrometry analyses. Tandem affinity purification of tagged Coq3, Coq6, and Coq9 confirmed the association of Coq3, Coq4, Coq5, Coq6, Coq7, and Coq9 in one or more complexes (Fig. 3). These findings recapitulate the results from previous affinity purification experiments (21, 25), with the additional observation that Cog3 is now shown to be associated with the entire complex and not just Coq4. In addition Coq8 was observed to co-purify with a Coq6-containing complex, but not with tagged Coq3 or Coq9, suggesting a closer association with Coq6. Although Coq8 is required for the stability of the Coq polypeptides and the biosynthetic complex (20, 28), this is the first evidence showing direct association of Coq8 with the complex.

The eluates from tandem affinity purification were also subjected to proteomic analyses to identify potentially novel binding partners beyond the known Coq proteins, the results of which are shown in Table 3. Several of the Coq proteins were observed in the eluates for the CNAP-tagged Coq proteins, consistent with results observed by immunoblotting in Fig. 3. In addition to the Coq proteins, a few other proteins were detected at levels comparable with the Coq proteins: YLR290C in the CNAP3 eluate and Ilv6 in the CNAP9 eluate. Ilv6 was also detected in the wild-type control and the CNAP3 and CNAP6 eluates but was much more abundantly detected with the CNAP9 eluate. Although there were other candidate proteins found specifically in eluates of CNAP-tagged samples, Ilv6 is a known mitochondrial protein, and ILV6 was also found to have negative genetic interactions with YLR290C in a high throughput study (55). The potential function of these two proteins in Q biosynthesis was assessed by monitoring growth on a nonfermentable carbon source and measuring de novo Q synthesis using stable isotopically labeled ring precursors for the corresponding null mutants. The *ilv* Δ and *ylr290c* Δ mutants did not display impaired growth on a nonfermentable carbon source (Fig. 6); however, both strains contained higher levels of Q intermediates compared with wild type. Moreover, the

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FIGURE 10. **Tandem affinity-purified YLR290C-CNAP co-purifies with Q and late-stage Q intermediates.** For each strain, aliquots of the anti-PC eluates (29% of E1 and 29% of E2) were combined and subjected to lipid extraction with methanol and petroleum ether. Lipid extracts were analyzed by HPLC-MS/MS for total $Q_6 (Q_6 + Q_6 H_2)$ (*A*), DMQ₆ (*C*), and IDMQ₆ (*E*). Measured lipids were normalized by the extracted eluate volume. The *bars* represent the mean of two measurements, and the *error bars* represent the standard deviation. Statistical significance was determined with the two-tailed Student's *t* test, and the *lowercase letters* above the *bars* are indicative of statistical significance. In *A*, the content of Q_6 in CA-1 was 5.4-fold higher compared with wild type (*a*, *p* = 0.0025). In *E*, there was no detectable IDMQ₆ (*D*), and IDMQ₆ (*F*). ND, not detected. Representative overlaid traces of both strains (W303, *magenta*; CA-1, *green*) are shown for Q_6 (*B*), DMQ₆ (*D*), and IDMQ₆ (*F*).





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FIGURE 11. **Sequence analysis of YLR290C and related proteins.** In *A*, YLR290C resides in a protein cluster populated by homologous fungal sequences including proteins that are fused to Coq10. Plant-like homologs form a distinct cluster composed of proteins from photosynthetic organisms. Prokaryotic sequences form a third independent cluster. The closest identified SDR subfamily distinct from YLR290C-like proteins includes human NDUFA9, a subunit of complex I, and orthologs in animals, fungi, and plants. In *B*, sequences representing each cluster from the protein similarity network in *A* were used to build a neighbor-joining tree. The tree is drawn to scale, with branch lengths in the same units (amino acid substitutions per site) as those of the evolutionary distances used to infer the phylogenetic tree. A branch length *scale bar* is shown below. A *number* after an organism name signifies the presence of multiple homologs in the tree. Protein identifications can be found in the supplemental Data Set 1. A schematic of the YLR290C-Coq10 fusion found in *U. maydis* is shown in *C*.

 $ylr290c\Delta$ mutant was found to have severely decreased *de novo* Q synthesis and decreased total Q as measured by ${}^{13}C_6$ -Q₆ and ${}^{12}C$ -Q₆, respectively (Fig. 7). The ability of the $ylr290c\Delta$ mutant to grow on a nonfermentable carbon source despite its decreased Q synthesis is not surprising, because other mutants with decreased Q synthesis have been characterized and retain the ability to grow on a nonfermentable carbon source. The $coq10\Delta$ mutant and particular yeast coq5 mutants expressing human COQ5 retain the ability to grow on a nonfermentable carbon source. The $cig10\Delta$ mutant and particular yeast coq5 mutants expressing human COQ5 retain the ability to grow on a nonfermentable carbon source despite having significantly decreased Q synthesis (56, 57). However, the $ylr290c\Delta$ mutant appeared to synthesize wild-type levels of Q₆ when grown in rich medium.³ Unlike several of the $coq\Delta$ mutants, steady-state levels of Coq4, Coq7, or Coq9 polypeptides were not decreased in the $ylr290c\Delta$ mutant in either rich or minimal medium.³

Because the *ylr290c* Δ mutant was partially defective in Q biosynthesis, the CNAP tag was integrated into the 3' end of the *YLR290C* ORF to generate a C-terminal fusion protein, which was subjected to tandem affinity purification to identify co-purifying proteins. Immunoblotting showed that YLR290C co-purified with Coq4, Coq5, and Coq7, confirming the results of the proteomic analysis and demonstrating association of YLR290C with the CoQ-synthome (Fig. 9). The levels of co-purified proteins were less than observed for CNAP-tagged Coq3, Coq6, and Coq9, which is consistent with the observation that proteomic analysis of the CA-1 eluate did not identify addi-



tional binding partners (not shown). It may be that YLR290C is more transiently associated with the Q biosynthetic complex or is less stable in digitonin extracts of mitochondria, leading to a reduced yield of associated proteins. Alternatively, the C-terminal CNAP tag may disrupt the interactions of YLR290C with other constituents of the CoQ-synthome, accounting for these observations.

The function of YLR290C is presently unknown, but it was previously determined to be a mitochondrial protein in a high throughput study of Perocchi *et al.* (58). Although we were unable to find homologs in available animal genomes, YLR290C-like proteins are common to photosynthetic organisms, including those with both primary and secondary plastids (Fig. 11). The *A. thaliana* ortholog of YLR290C is localized to the plastid based on proteomics data (59), and several other reports have identified it in plastoglobules (54, 60–63), which are sites of prenylquinone synthesis and storage (64). Indeed, the finding of YLR290C homologs in plastoglobules is reminiscent of the Coq8/Abc1 atypical protein kinase family members (ABC1K), also present in plastoglobules (65).

Sequence analyses identified YLR290C as part of the SDR superfamily (Fig. 11), which includes a diverse family of oxidoreductases that catalyze reactions such as isomerization, decarboxylation, epimerization, imine reduction, and carbonyl-alcohol oxidoreduction (66). SDR proteins contain a conserved Rossmann fold, a structural motif found in proteins that bind nucleotide co-factors such as NAD(P), FAD, and FMN (67). The crystal structure of *Pseudomonas aeruginosa* UbiX,

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³ C. Allan, unpublished observations.



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FIGURE 12. Amino acid alignment of selected fungal YLR290C fusion proteins with YLR290C and Coq10 from S. cerevisiae. The N-terminal YLR290C-like portion is highlighted in *yellow*, and the Coq10 portion near the C terminus is highlighted in *blue*. The poorly conserved N-terminal sequence found in *P. antarctica* (res 1–100) is not shown. The *numbering* at the top of the alignment corresponds to the *P. antarctica* sequence. *Pan*, *P. antarctica*; *Sre*, *S. reilianum*; *Pfl*, *P. flocculosa*; *Mpe*, *M. pennsylvanicum*; *Uma*, *U. maydis*.

which catalyzes the decarboxylation step in Q biosynthesis, was shown to contain a Rossmann fold with a bound FMN (68). Similarly, the structure of E. coli Pad1, a paralog with 51% identity to E. coli UbiX, was shown to contain a typical Rossmann fold with a bound FMN (69). Although the function of the Rossmann fold in YLR290C remains unclear, it is tempting to speculate that it may also function to catalyze an FMN-dependent decarboxylation in yeast Q biosynthesis, potentially catalyzing the first step denoted by question marks in Fig. 1. The observation that the *ylr290c* Δ mutant still produces small amounts of Q_6 is not fully consistent with this hypothesis; however, there may be redundant decarboxylases capable of bypassing the $ylr290c\Delta$ mutant such as Pad1 and Fdc1, two phenylacrylic acid decarboxylases with homology to the E. coli Q biosynthetic decarboxylases UbiD and UbiX (70, 71). Expression of yeast PAD1 in an E. coli ubiX mutant restored Q₈ synthesis; however,

yeast mutants lacking either *PAD1*, *FDC1*, or both produce wild-type levels of Q_6 (71, 72), potentially because of their functions as redundant decarboxylases.

Yet another link connecting YLR290C to Q biosynthesis is the presence of YLR290C-Coq10 fusions encoded in five fungal genomes (Figs. 11 and 12). Such fusion proteins, also known as Rosetta Stone proteins, are often indicative of a functional link between two proteins including a physical interaction or residency in the same pathway (73). Coq10 was shown to be required for efficient *de novo* Q biosynthesis (56), and several studies indicate that yeast Coq10 and orthologs bind Q or DMQ via a hydrophobic START domain (56, 74, 75), leading to the hypothesis that Coq10 acts a Q chaperone, necessary for delivery of Q to the CoQ-synthome for efficient *de novo* Q synthesis and respiration (56). The fusion of YLR290C with Coq10 strongly suggests a functional interaction of YLR290C with this

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FIGURE 13. **Model of the Q biosynthetic complex.** The organization of the complex is based on co-precipitation experiments performed in previous work (21) and this study, as well as two-dimensional blue native-PAGE analysis (28). Coq1, Coq2, and Coq10 have not been shown to associate with the complex. Coq10 binds Q and particular Q intermediates and is postulated to function as a Q chaperone for efficient respiration and *de novo* Q biosynthesis (56). Coq8 is required for the phosphorylation of Coq3, Coq3, and Coq7 (19), and co-precipitation of CNAP-tagged Coq6 demonstrates their physical association. Co-precipitation of tagged YLR290C, here designated Coq11, showed its association with Coq4, Coq5, and Coq7. The number of copies of each Coq polypeptide depicted in the CoQ-synthome is hypothetical because the stoichiometry has not been determined. Lipid analysis of co-precipitation eluates demonstrated the association of Q₆ and the late stage intermediates DMQ₆ and IDMQ₆ with the complex, illustrated as small molecules in association with Coq4.

Q-binding protein. Interestingly, *YLR290C* was found to have a genetic correlation with both *COQ2* and *COQ10* in high throughput analyses (55). *YLR290C* was also identified along with *COQ1*, *COQ2*, *COQ3*, *COQ4*, *COQ6*, *COQ7*, and *COQ10* as part of a set of mitochondrial genes required for doxorubicin resistance (76).

To further explore the association of associated lipids with the CoQ-synthome, eluates from tandem affinity purification were subjected to lipid extraction and LC-MS/MS to measure Q_6 and Q_6 intermediates. Q_6 and the Q_6 intermediates DMQ_6 and IDMQ_6 were all detected at levels significantly greater than the wild-type control in eluates recovered from the CNAP3, CNAP6, and CNAP9 strains (Fig. 4) and also in the CA-1 strain with the CNAP-tagged YLR290C-captured complex (Fig. 10). The early intermediates HHB and HAB were not detected in any of the eluates (data not shown). The observation that YLR290C co-purified with Q and late-stage Q intermediates is consistent with the immunoblot results showing the association of YLR290C with Coq4, Coq5, and Coq7 and supports the role of YLR290C as a component of the Q biosynthetic complex.

These results are consistent with the previous association of DMQ₆ with the Q biosynthetic complex and with the ability of exogenous Q_6 to stabilize several Coq proteins in certain *coq* mutants. For example, the late stage Q intermediate DMQ_6 co-purifies with the Q biosynthetic complex (25), and exogenous Q₆ has been shown to stabilize particular Coq proteins and lead to production of later stage biosynthetic intermediates in certain coq mutants (27–29), suggesting a function for Q or a Q intermediate in the stability of the CoQ-synthome. Coq4 was previously hypothesized to serve as a scaffolding protein for the complex, and the crystal structure of the Coq4 homolog Alr8543 from Nostoc sp. PCC7120 revealed a bound geranylgeranyl monophosphate, suggesting that Coq4 functions to stabilize the Q biosynthetic complex through its interactions with other Coq proteins and a polyisoprenoid lipid (26, 77). The crystal structure of human COQ9 was recently solved and shown to have a lipid-occupied cavity; mass spectrometry analyses of purified human COQ9 identified its association with phospholipids and Q (78). Moreover, human COQ9 was shown

to bind human COQ7, indicating that a complex is required for Q biosynthesis in human cells (78). Finally, a recent study showed that a mutated form of ADCK3, a human Coq8 ortholog, possessed autophosphorylation activity (79), a finding corroborated by another recent study showing that a purified MBP-ADCK3 fusion exhibited ATPase activity *in vitro* (80). The authors suggested that ADCK3 may act in concert with Coq9, a hypothesis made particularly compelling by their observation that a fusion of COQ9 and ADCK3 is present in the protozoan *Tetrahymena thermophila* (79).

The findings presented here strongly indicate that YLR290C functions in Q biosynthesis, and we propose that this protein be designated Coq11. These results have been incorporated into an updated model of the Q biosynthetic complex (Fig. 13), which now depicts Coq8 in association with Coq6; Coq11 is shown in association with Coq4, Coq5, and Coq7. There were additional protein candidates from proteomic analysis of the tandem affinity purification eluates. Further characterization of these proteins may yield a more complete understanding of the proteins associated with the Q biosynthetic complex and the regulation of this pathway.

Acknowledgments—We thank Dr. Carla Koehler (UCLA) for advice on generating the CNAP-tagged constructs used in this study. We thank Letian Xie and Kelly Quinn for help with analyses of steadystate levels of Coq polypeptides in the ylr290c Δ mutant yeast. We thank Dr. A. Tzagoloff (Columbia University) for the original yeast coq mutant strains. We acknowledge the UCLA Molecular Instrumentation Core proteomics facility for the use of QTRAP 4000 and the gel imager.

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