A lettuce (Lactuca sativa) homolog of human Nogo-B receptor interacts with cis-prenyltransferase and is necessary for natural rubber biosynthesis

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Running title: Natural rubber biosynthesis in lettuce

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Background: Natural rubber biosynthesis in lettuce (Lactuca sativa) and other plants remains elusive.

Results: An unusual cis-prenyltransferase-like protein interacts with and tethers a cis-prenyltransferase on endoplasmic reticulum, and its RNAi-silencing eliminates natural rubber.

Conclusion: cis-prenyltransferase-like protein is a necessary component in natural rubber biosynthesis in lettuce.

Significance: The results presented here suggest hetero-protein complexes are involved in natural rubber biosynthesis.

ABSTRACT

Natural rubber (cis-1,4-polyisoprene) is an indispensable biopolymer used to manufacture diverse consumer products. Although a major source of natural rubber is the rubber tree (Hevea brasiliensis), lettuce (Lactuca sativa) is also known to synthesize natural rubber. Here, we report that an unusual cis-prenyltransferase-like 2 (CPTL2), which lacks the conserved motifs of conventional cis-prenyltransferase, is required for natural rubber biosynthesis in lettuce. CPTL2, identified from the lettuce rubber particle proteome, displays homology to a human NogoB receptor and is predominantly expressed in latex. Multiple transgenic lettuces expressing CPTL2-RNAi constructs showed that decrease of CPTL2 transcripts (3-15% CPTL2 expression relative to controls) coincided with the reduction of natural rubber as low as 5%. We also identified a conventional cis-prenyltransferase 3 (CPT3), exclusively expressed in latex. In subcellular localization studies using fluorescent proteins, cytosolic CPT3 was re-localized to endoplasmic reticulum by co-occurrence of CPTL2 in tobacco and yeast at the log phase. Furthermore, yeast two-hybrid data showed that CPTL2 and CPT3 interact. Yeast microsomes containing CPTL2/CPT3 showed enhanced synthesis of short cis-polyisoprenes, but natural rubber could not be synthesized in vitro. Intriguingly, a homologous pair CPTL1/CPT1, which displays ubiquitous expressions in lettuce, showed a potent dolichol biosynthetic activity in vitro. Taken together, our data suggest that CPTL2 is a scaffolding protein that tethers CPT3 on endoplasmic reticulum and is necessary for natural rubber biosynthesis in planta, but yeast-expressed CPTL2/CPT3 alone could not synthesize high molecular weight natural rubber in vitro.

Natural rubber (NR) is an important bio-polymer used in the manufacture of hundreds of industrial products (1). NR displays exceptional physicochemical properties (e.g., elasticity and abrasion resistance) required for many heavy-duty products. Despite progress in polymer chemistry,
NR belongs to the isoprenoid class of natural products, which use isopentenyl diphosphate (IPP) as a central building block. IPP and its isomer dimethylallyl diphosphate (DMAPP) undergo condensations to generate C10 geranyl diphosphate (GPP), C15 farnesyl diphosphate (FPP), and C20 geranyl geranyl diphosphate (GGPP) (4). In these reactions, DMAPP serves as a primer on which IPPs are sequentially condensed in trans-configuration, thereby the enzymes catalyzing these reactions are referred to as trans-prenyltransferases (TPTs). On the other hand, in NR biosynthesis, a number of IPP molecules are conjugated onto a FPP molecule in cis-1,4-configuration by cis-prenyltransferase (CPT) (Figure 1). About 15,000 IPP molecules are condensed to form the NR polymer, with average molecular weight \( M_w \) exceeding 1 million g/mol in rubber tree. The resulting NR bio-polymer shows unique polymer features, including exclusive cis-configuration, extremely long polymer length, and narrow polydispersity (the spread of polymers). Although TPT and CPT share a similar mechanism involving a carbocation intermediate, they share no sequence homology (5).

Crystal structures of TPT (FPP synthase) and CPT [undecaprenyl diphosphate synthase (UPPS)] further confirmed no structural similarity between these enzymes (5,6). Thus, TPT and CPT are entirely different classes of enzyme and likely evolved independently.

In all living organisms, cis-isoprenoids with carbon length of C55 to C110 are ubiquitously present. Undecaprenyl diphosphate (C55) and dolichyl diphosphate (C70-C110) function as essential polysaccharide carrier molecules during cell wall biosynthesis in bacteria and N- and O-glycosylation of proteins in eukaryotes, respectively (7,8). It was reported that \textit{E. coli} has a single \textit{CPT} (i.e., UPPS), whereas yeast (\textit{Saccharomyces cerevisiae}) has two \textit{CPTs}, \textit{RER2} and \textit{SRT1}, responsible for C70-110 dolichyl diphosphate biosynthesis (9-11). Null mutations of \textit{E. coli UPPS} and double knock-out of the two yeast \textit{CPTs} are lethal, confirming that they are essential genes (9,11). In plants, studies of well-annotated genomes showed that multiple copies of \textit{CPTs} are present (12,13), but the exact physiological role of each \textit{CPT} is not fully understood.

NR is distinguishable from undecaprenol and dolichol in that it has significantly longer polymer length and is not involved in essential primary metabolism. It is important to note that 2,500 plant species can produce NR with \( M_w \) ranging from \( 10^7 \)- to 2 million g/mol (14). Of particular interest is that several members of the Asteraceae family produce \textit{cis}-polyisoprenes qualitatively comparable to the NR from rubber tree. For example, dandelion (\textit{Taraxacum spp.}), lettuce (\textit{Lactuca sativa}), and guayule (\textit{Parthenium argentatum}) can synthesize NR of more than 1 million g/mol \( M_w \) (15-17). Although shorter than rubber tree NR, other Asteraceae plants, such as sunflower (\textit{Helianthus spp.}) and goldenrod (\textit{Solidago canadensis}), can also synthesize \textit{cis}-polyisoprenes with \( M_w \) of 30-230 x10\(^3\) g/mol (17,18). Therefore, studies of NR from these Asteraceae plants can help elucidate NR biosynthesis in plants.

At present, the molecular mechanism for NR biosynthesis, resulting in a wide range of \( M_w \), remains elusive. NR from rubber tree is synthesized in the cytoplasm of laticifer cells (latex), a pipe-like tissue which develops alongside the phloem. Classical biochemical studies using latex have proposed a model where NR is synthesized on a monolayer organelle, known as the rubber particle, suspended in the latex (19). A rubber synthase complex is proposed to reside on the membrane of rubber particles with its catalytic domain exposed to the cytosol. In this model, the newly synthesized NR polymer extends into the core of the rubber particle using IPP substrates from the cytoplasm. However, molecular evidence for this model is scarce, and the rubber synthase mystery remains unsolved in the past 50 years.

Considering the mechanistic commonality, it can be postulated that rubber synthases evolved from bacterial and eukaryotic \textit{CPTs}. Recently, three dandelion \textit{CPTs} displaying >96% amino acid identity were silenced by RNA interference (RNAi) (20). The resulting transgenic dandelions showed elimination of NR in the latex, thereby proving the
necessity of CPT in NR biosynthesis, although purified recombinant CPT activity was not reported. The in vitro synthesis of ~1 million g/mol NR was reported using E. coli-produced H. brasiliensis CPT co-incubated with latex fraction (21); however, yeast-produced H. brasiliensis CPT from the same clone only synthesized regular dolichol and not NR in comparable in vitro conditions (22), thereby questioning the in vitro NR-synthetic activity of H. brasiliensis CPT. Currently, the in vitro synthesis of cis-polyisoprenes longer than undecaprenol and dolichol has not been achieved using purified recombinant CPT enzymes.

It was recently identified that the human Nogo-B receptor (NgBR) shares weak sequence homology to CPT (<15% at the amino acid level), and it lacks all five motifs conserved in all CPTs (23,24). In humans, NgBR interacts with the Nogo-B peptide and Niemann-Pick type C2 (NPC2) protein to mediate signals for the proliferation of epithelial cells and to regulate intracellular cholesterol trafficking, respectively (23,25). Interestingly, it also interacts with the human CPT enzyme (24). In plant, its homolog in Arabidopsis (LEW1) was identified during genetic screening for a leaf wilting phenotype (26). The lew1 mutant is unable to efficiently synthesize dolichol, causing defects in O- and N-glycosylation of proteins. These results suggest that the CPT-like proteins have implications in dolichol biosynthesis in eukaryotes.

One difficulty of investigating rubber tree NR metabolism is that reverse genetic tools are not applicable to this perennial tropical tree. It was reported that an annual, transformable plant species, lettuce (Lactuca sativa), could synthesize and store NR in rubber particles in the same manner as that of the rubber tree (15). In this work, we established lettuce as a model plant for the study of NR biosynthesis. Proteomic analysis identified a CPT-like (CPTL) gene as a homolog of NgBR, and its involvement in NR biosynthesis was comprehensively studied by reverse genetic, cell biological and biochemical approaches. From these results, we conclude that CPTL2 is necessary but not solely responsible for NR biosynthesis in lettuce.

**EXPERIMENTAL PROCEDURES**

**Plant material and growth condition** - The seeds of L. sativa cv. Mariska and Ninja were obtained from the University of California Davis (Dr. Richard Michelmore). Lettuce plants were grown in a growth chamber at 20 °C and a 16 h photoperiod for 30 days. Plants were then transferred to 6-inch diameter pots and cultivated at the University of Calgary greenhouse at 23 ± 3 °C with a 16 h photoperiod.

**RNA isolation and transcription analysis** - Fine powder (100 mg) ground in liquid nitrogen from various tissues, or fresh latex (100 mg), was immediately mixed with 1 mL of Trizol reagent (Invitrogen, Carlsbad, CA). RNA was isolated according to the manufacturer's protocol. First-strand cDNA was synthesized using SuperScript III reverse transcriptase (Invitrogen, Carlsbad, CA) and oligo (dT) 12-18 primer (Invitrogen, Carlsbad, CA) using 1-5 µg total RNA. qRT-PCR was performed (Step One Real-Time PCR System, Applied Biosystems, Carlsbad, CA) using Power SYBER Green PCR Master Mix (Applied Biosystems, Carlsbad, CA), 5 µM primer and cDNA template (equivalent to 5 ng total RNA) in a reaction volume of 10 µL. The qRT-PCR program was 1 cycle of 95 °C for 10 min and 40 cycles of 95 °C for 15 s and 58 °C for 1 min. The Critical Threshold (Ct) values were used to calculate the relative transcript abundance using actin as the internal control as described (27). The primer efficiency was calculated from qRT-PCR of the serial dilution of total cDNA, and the specificity of the primers was confirmed by the dissociation curve for each primer set. The primers used for qRT-PCR are listed in Table 3 (primer numbers 36-47). The transcript levels of CPTL2 and CPT3 in latex were directly compared using an external standard curve in which the Ct values were plotted against the corresponding known amount of either CPTL2 or CPT3 plasmid in qRT-PCR reactions.

**Isolation of rubber particles and proteomics** - Latex was collected as described (15) in ice-cold latex collection buffer. The samples were centrifuged at 10,000xg for 2 min at 4 °C. The floating rubber layer was washed twice with latex collection buffer. The rubber particles were resuspended in latex storage buffer. The particle proteins were resolved on 10% SDS-PAGE. The
gel was sliced into nine pieces. Trypsin digestion and subsequent LC-MS/MS analysis using MASCOT software (Matrix Science, Boston, MA) were carried out at the Southern Alberta Mass Spectrometry (SAMS) Center at the University of Calgary. Detailed methods were previously described (28). Reference sequence file used for the MASCOT analysis was CLS_S3_ESTs_Sat.assembly (The Genome Center, University of California, Davis, USA).

**Rubber quantity and quality analysis** - Fresh latex (50 mg) was mixed with 1 mL of acetone and centrifuged at 20,000g for 1 min. The pellet was allowed to dissolve in 1 mL of tetrahydrufuran (THF) overnight. The sample solution was filtered through a 0.45 µm polytetrafluoroethylene (PTFE) filter disk and subjected to HPLC (Waters Alliance HT 2795 Separations Module, Waters, Milford, MA) analyses. Samples (50 µl) were injected and separated (mobile phase: 0.6 mL min⁻¹ THF) in tandem-connected GPCs with a linear separation range of 2×10⁶ to 1×10⁸ Da (Styragel HR 3 and Syragel HR 5, Waters, Milford, MA) at 35 °C. The signal was detected by Waters 2420 ELS Detector (Waters, Milford, MA) at 36 °C for nebulizer and 50 °C for drift tube. Empower2 chromatography Data Software (Waters, Milford, MA) was used to analyze the data. Molecular mass and polydispersity were calculated based on cis-polyisoprene standards (Polymer Standards Service-USA, Amherst, MA). The relative rubber content was obtained from an external standard curve (ELSD peak area against a serial dilution of lettuce latex) and compared to that of the controls.

For visual rubber analysis in Figure 5A, 350 mg (fresh weight) latex was suspended in 1 mL latex collection buffer and centrifuged at 10,000xg for 5 min. For Figure 5B, 30 mg fresh weight of latex was suspended in 180 µL of 2% (w/v) SDS and centrifuged at 20,000g for 10 min at 4 °C.

**CPT transient expression in tobacco and confocal microscopy** - The binary vectors for fluorescent protein tagging (pSITE-0B, -2NB and -4NB with CPT/CPTL) were transformed into the Agrobacterium strain C58. Cells collected from an overnight culture were diluted in infiltration buffer (10 mM MES pH 5.5, 10 mM MgCl₂, 150 µM acetosyringone) to OD₆₀₀ of 0.8 and incubated at room temperature for 2 h before infiltration into N. benthamiana leaves. The localization of CPTs was visualized after 3-4 days using a confocal microscope (Leica TCS SP5 II, Leica Microsystems Inc., Concord, Ontario, Canada). The lipid substance was visualized by staining leaf samples with 0.5% (w/v) Nile Red in 70% ethanol for 20 min. The endoplasmic reticulum was visualized by co-infiltration with pBIN20-mCherry-HDEL. EGFP was detected by excitation at 488 nm and emission at 500-530 nm. RFP and mCherry were detected by excitation at 543 nm and emission at 590-650 nm. Nile red was detected by excitation at 543 nm and emission at 580-630 nm. Images from different excitations were sequentially collected.

**Localization of CPT in yeast** - The vectors (pUG23 and p415GPD with CPT/CPTL) were transformed into a yeast strain EY0987 (ATCC 201389, MATα his3Δ1 leu2Δ0 lys2Δ0 ura3Δ0) with CPT/CPTL overnight culture were diluted in infiltration buffer (10 mM MES pH 5.5, 150 µM acetosyringone) to OD₆₀₀ of 0.8 and incubated at 4 °C. The vectors (pUG23 and p415GPD with CPTL/CPTL) were transformed into a yeast strain JC1280 (MATα his3Δ1 trp1Δ0 leu2::proLEU2-lexAop6) (30). Multiple colonies were inoculated in 2 mL SC medium containing either 2% glucose or 2% galactose and incubated at 30 °C. Localization of CPT/CPTL was determined at 8 h (log phase) and 20 h (stationary phase) after inoculation using a confocal microscope.

**Yeast two-hybrid assay** - The vectors (pGAL-LexA/pJG4-6 with CPTL/CPTL and reporter plasmid pSH18-34 for β-galactosidase) were transformed into a yeast strain JC1280 (MATα his3Δ1 trp1Δ0 leu2::proLEU2-lexAop6) (30). Multiple colonies were inoculated in 2 mL SC media containing 2% raffinose. Cells (0.5 mL) from an overnight culture were inoculated into 2 mL SC media containing either 2% glucose or 2% galactose and incubated at 30 °C for 6 h. Yeast cells, equivalent to 1 mL culture with OD₆₆₀ between 0.6-1.0, were collected and resuspended in 1 mL of Z buffer (60 mM Na₂HPO₄, 40 mM NaH₂PO₄, 10 mM KCl, 1 mM MgSO₄, 50 mM β-mercaptoethanol, pH 7), followed by lysis by vortexing for 20 s with the addition of 20 µL chloroform and 20 µL 0.1% SDS. The mixture was incubated at 28 °C for 5 min. The reaction was started by addition of 0.2 mL of 4 mg mL⁻¹ orthonitrophenyl-β-galactoside (ONPG) in Z-buffer and was stopped by addition of 0.5 mL of 1 M Na₂CO₃ after 2.5-5 min. The mixture was centrifuged at 20,000g for 5 min and the OD₄₂₀ of
the supernatant was read. The β-galactosidase activity was calculated in Miller units.

Generation of DNA constructs - The primer sequences are listed in Table 3. Three RNAi-fragments of CPTL2 cDNA were sequentially amplified by primers (3/4, 5/6, 7/8) and universal Gateway-cloning primers (1/2). The amplicons were cloned into an RNAi binary vector pK7GW/IGW2D[II] (Plant Systems Biology, VIB-Ghent University, Belgium), sequentially using the Gateway BP Clonase and Gateway LR Clonase (Invitrogen, Carlsbad, CA). For GFP tagging and overexpression in plant, full-length CPTL2 was amplified with primers (3/9) and (3/10) and cloned into binary vector pSITE-2NB and pSITE-OB, respectively (31). For GFP- and RFP-tagging, full-length CPT3 was amplified with primers (11/12) and cloned into pSITE-2NB and pSITE-4NB vectors, respectively. For expression of GFP-tagged, FLAG-tagged, and non-tagged CPTL2 in yeast, CPTL2 was amplified from lettuce latex cDNA with primers (14/15 for GFP-tagging, first PCR with 16/17 and second PCR with 16/53 using the first PCR product as template for FLAG-tagging, 16/17a for non-tagged version) and cloned into pUG23 for GFP-tagging and in p415GPD for both FLAG-tagged and non-tagged versions. CPTL1 was also similarly cloned into p415GPD using primers (first PCR with 22/23 and second PCR with 22/53 using the first PCR product as template for FLAG-tagging and 22/23a for non-tagged version), and cloned into pUG23 for GFP-tagging and in p415GPD for both FLAG- and non-tagged versions. CPTL1 was similarly cloned into p415GPD using primers (first PCR with 18a/19a and second PCR with 18a/53 using the first PCR product as template for FLAG-tagging) and (18a/19b for non-tagged version). Similarly, CPT1 was amplified from lettuce latex cDNA with primers (51/52 for GFP-tagging, first PCR with 22/23 and second PCR with 22/53 using the first PCR product as template for FLAG-tagging and 22/23a for non-tagged version), and cloned into pUG23 for GFP-tagging and in p416GPD for both FLAG-tagged and non-tagged versions. The yeast RER2 was amplified from yeast genomic DNA and cloned into p416GPD for FLAG-tagging with primers (first PCR with 20/21 and second PCR with 20/51 using the first PCR product as template). For the Y2H assay, CPTL2 and synCPT3 were cloned into pGAL-LexA and pIJ46-6 vectors with primers (24/25, 28/29, 26/27, 30/31), respectively. For expression in E. coli, CPTL2 and synCPT3 were cloned into the pDONOR221/pDEST17 vector (Invitrogen, Carlsbad, CA) using primers (3/10 and 11/13), respectively. Recombinant SRPP4 and Caleosin were cloned into pDONOR221/pDEST17 using primers (32/33 and 34/35), respectively.

Agrobacterium-mediated plant transformation - Leaf explants for transformation were obtained from seedlings (1-2 months old), cultivated at 25°C, 16 hr photoperiod and 45% relative humidity (RH) in Murashige and Skoog (MS) medium. Overnight culture of A. tumefaciens strain LBA4404 carrying the specific binary plasmid was used to infect the leaf explants, and thereafter co-cultivated in MS media supplemented with 0.2 mg L⁻¹ 6-benzylaminopurine (BA, Sigma-Aldrich, St Louis, MO) and 0.2 mg L⁻¹ 1-naphalene acetic acid (NAA, Sigma-Aldrich, St Louis, MO) for 2 d at 25 °C, 16 h day/ 8 h night photoperiod and 45% RH (the growth condition remained the same unless specified). Subsequently, the explants were transferred to MS media supplemented with 0.2 mg L⁻¹ BA, 0.2 mg L⁻¹ NAA, 25 mg L⁻¹ kanamycin (Amresco, Solon, OH) and 250 mg L⁻¹ cefotaxime (Duchefa Biochemie, Haarlem, Netherland). The putative shoots were placed in rooting medium containing 0.1 mg L⁻¹ NAA, 12.5 mg L⁻¹ kanamycin and 250 mg L⁻¹ cefotaxime. After rooting, the shoots were screened for GFP expression through confocal microscopy, transferred to potted soil in growth chamber (for 1 month), and then transferred to the greenhouse for further growth and analyses.

CPT and CPTL expression in E. coli - For CPTL2, E. coli BL21-AI cells (Invitrogen, Carlsbad, CA) with GST-CPTL2 in pGEX-6p-2 vector (GE Healthcare Life Sciences, Piscataway, NJ) were cultured at 37°C in 1 L LB medium until an OD₆₀₀ of 0.5 was reached, then cells were induced with 0.1% arabinose and 0.4 mM IPTG. Cells were grown at 15°C for 24 h. The cells were collected and resuspended in lysis buffer [25 mM Tris-HCl pH 7.5, 100 mM NaCl, 10% (v/v) glycerol, 1 mM
After sonication, the soluble fraction was incubated for 1 h with 1 mL of pre-equilibrated glutathione resin. The column was washed with wash buffer [50 mM Tris-HCl pH 8, 100 mM NaCl, 1 mM EDTA, 1 mM DTT, 0.1% (v/v) Triton X-100], and eluted with elution buffer (50 mM Tris-HCl pH 8, 100 mM NaCl, 1 mM EDTA, 1 mM DTT, 10 mM reduced glutathione). For CPT3, *E. coli* BL21AI cells with 6x Histidine-CPT3 in pDEST17 vector were cultured at 37°C in 0.3 L LB media until an OD$_{600}$ of 0.6 was reached, then induced with 0.2% arabinose at 37°C for 0.5 h. The cells were collected and resuspended in lysis buffer [25 mM Tris-HCl pH 7.5, 100 mM NaCl, 10% (v/v) glycerol, 1 mM PMSF, 1 mM DTT]. After sonication, the soluble fraction was incubated for 1 h with 1 mL Ni-NTA resin. The column was washed with 30 mL wash buffer [25 mM Tris-HCl pH 7.5, 100 mM NaCl, 10% (v/v) glycerol, 1 mM PMSF, 0.01% (v/v) Triton X-100, and 25 mM imidazole], and was eluted with elution buffer [25 mM Tris-HCl pH 7.5, 100 mM NaCl, 10% (v/v) glycerol, 1 mM PMSF, and 500 mM imidazole].

**CPT expression in yeast and microsome isolation** - The p415GPD or p416GPD vector containing CTPL1/2 or CPT1/3 respectively either with C-terminal FLAG or without any tagging was transformed into YPH499 yeast strain using a standard lithium acetate transformation protocol. Transformed yeast cells were grown to mid-late logarithmic phase in synthetic complete medium with glucose lacking uracil and leucine. The cell pellet was broken open by glass bead-beating method (BioSpec Products) in TES-B buffer (50 mM Tris-HCl pH 7.4, 1 mM EDTA, 500 mM D-sorbitol). For microsome preparation, the cell lystate was centrifuged at 10,000 g for 10 min at 4 °C. The supernatant was further centrifuged at 100,000 g for 1 hour at 4 °C. The microsome pellet was resuspended in TG buffer (50 mM Tris-HCl pH 7.4, 20% (v/v) glycerol) and stored at -80 °C.

**Immunoblot analysis** - Yeast microsomal protein or total protein was separated by SDS-PAGE on 12% SDS-PAGE and transferred to PVDF membrane. The membrane was hybridized with 1:2000 diluted anti-FLAG antibody (Sigma-Aldrich, St Louis, MO) and 1:5000 diluted anti-mouse secondary antibody (Sheep anti-mouse, GE Healthcare Biosciences, Pittsburgh, PA), developed with Amersham ECL Plus Western Blotting Detection Reagent (GE Healthcare Life Sciences, Piscataway, NJ), and exposed to a film. For DPM1 detection, 1:1000 diluted anti-DPM1 antibody (Life technologies, Carlsbad, CA) and 1:10,000 diluted anti-mouse secondary antibody were applied to the same blot after stripping.

**CPT enzyme assay and TLC** - Yeast microsomes (50 µg protein) containing recombinant CPT/CPTL was assayed as described (24). The 50 µl reaction mixture contained: 50 µM FPP (Echelon, Biosciences Inc., Salt Lake City, UT), 175 µM IPP (Echelon, Biosciences Inc., Salt Lake City, UT), 25 µM $^{14}$C-IPP (1.1×10$^5$ dpm, Perkin Elmer, Waltham, MA), 50 mM HEPES pH7.5, 5 mM MgCl$_2$, 2 mM DTT, 2 mM Na$_3$VO$_4$ and 2 mM NaF. The reaction was incubated at 30 °C for 2 h, then extracted with 1 mL chloroform:methanol (2:1 v/v) and 0.4 mL 0.9% (w/v) NaCl. After partitioning by centrifugation, the chloroform fraction was washed 3 times with 0.5 mL chloroform: methanol: water (39:1 v/v), and a 10% aliquot was assayed for radioactivity by scintillation counting. The remaining extraction was dried down completely and hydrolyzed in 1M HCl for dephosphorylation by incubating at 85 °C for 1 h, followed by equal volume extraction twice with benzene. Benzene was dried down to a small volume for spotting on TLC plate. C18 reverse silica plate (Whatman, England) was used for separation in acetone:water (39:1 v/v) solvent. The TLC plate was exposed to a Phosphimager Screen (GE Healthcare) for indicated number of days before scanning in Molecular Imager FX (Bio Rad, Hercules, CA).

**Multiple reaction monitoring** - SRPP4 and caleosin were used as the internal calibrators for CPTL2 and CPT3 quantitation. Full-length SRPP4 and calsosin were amplified with primers (32/33 and 34/35, respectively, Table 3) and extended with primers (1/2). Both genes were cloned with Gateway cloning into pDEST17 vector (Invitrogen, Carlsbad, CA), according to the manufacture’s protocol. The vectors were transformed to BL21AI cells (Invitrogen, Carlsbad, CA). The cells were cultures at 37°C in 2 mL LB medium until an OD$_{600}$ of 0.6 was reached, then was induced with...
0.2% arabinose at 37°C for 0.5 h. The cells were collected and resuspended in lysis buffer [25 mM Tris-HCl pH 7.5, 100 mM NaCl, 10% (v/v) glycerol, 1 mM PMSF, 1 mM DTT]. After sonication, the total pellets were collected, resuspended in lysis buffer, and resolved in 12% SDS-PAGE. The recombinant protein bands were cut from the gel and subjected to MRM. Multiple reaction monitoring by LC-MS/MS was performed by UC Davis Proteomics Core Facility. Peptides and daughter ions used were shown in Supplemental Table 2. For in-gel digestion, gel pieces were cut into ~1 mm cubes and washed two times with 50 mM ammonium bicarbonate (AMBIC), pH 8, and then dehydrated with 100% acetonitrile (ACN). The gel pieces were then dehydrated with 10% ACN and alkylated with 55 mM iodoacetamide for 20 min. The gels were then washed with 50 mM AMBIC, dehydrated one last time, and digested overnight at 37 °C using modified sequencing-grade trypsin (Promega BioSciences, San Luis Obispo, CA). For extraction, the supernatant was collected and put into a fresh tube. Next, 60% ACN:0.1% trifluoroacetic acid was added to the gel pieces, sonicated for 10 min, and centrifuged for 8 min. The supernatant was then removed and added to the fresh tube. The supernatant was then vacuum-centrifuged until dry.

Peptides from candidate proteins were selected based on prior shotgun LC-MS/MS data. All peptides were selected based on the following criteria: uniqueness to the targeted protein; fully tryptic (i.e. no missed cleavages or ragged ends); and absence of potentially modified amino acids. Cysteine-containing peptides were presented as carbamidomethylated derivatives. Acquisition scheduling was performed using the Skyline Retention Time (iRT) predictor calculator. A heavy peptide retention time calibration mixture (Thermo Scientific) was spiked into each LC-MRM-MS run to monitor for retention time shift. LC-MRM-MS analysis was performed on a TSQ Vantage Triple Quadrupole mass spectrometer (Thermo-Fisher Scientific) in conjunction with an Advance Splitless Nano UHPLC and an Advance Autosampler (Bruker-Michrom, Auburn, Ca). 10 µL of the digested peptides and 1 µL of the retention time mixture were loaded onto a Michrom C18 trap and desalted before they were separated using a Michrom 200 µm x 150 mm Magic C18AQ reverse phase column. A flow rate of 2 µL min⁻¹ was used. Peptides were eluted using a 60 min gradient with 5% B to 35% B for 50 min, 35% B to 80% B for 1 min, 80% B for 1 min, 80% B to 5% B for 1 min, and finally held at 5% B for 7 min (A=0.1% formic acid, B= 100% acetonitrile). The mass spectrometer was operated with a spray voltage of 1.8 kV, a capillary temperature of 200°C, and a normalized collision of 35%. Peak areas for each peptide were extracted and integrated using Skyline v1.4.

Nucleotide sequence deposition - The nucleotide sequences of the cDNAs reported in this paper have been submitted to GenBank with the following accession numbers: CPTL1, KF752484; CPTL2, KF752485; CPT1, KF752486; CPT2, KF752487; CPT3, KF752488.

RESULTS

Identification of a distinct CPT from lettuce - NR with Mᵦ of 1.27 million g/mol and polydispersity (Pd) value of 1.1 was previously reported from L. sativa, cv. Salinas (15). Using an HPLC-Gel Permeation Column (GPC), these literature data were corroborated in two different lettuce cultivars, Mariska and Ninja. The Mᵦ and Pd from Mariska and Ninja were comparable to each other and slightly higher than the reported values in Salinas (Table 1). Since NR accumulates in the stem, early bolting and flowering facilitate NR studies in lettuce. Mariska was used initially in this study. However, as Ninja flowers ~50 days earlier than Mariska, Ninja was primarily used to expedite the research progress.

Lettuce (cv. Mariska) liquid latex samples were centrifuged and separated into three fractions: floating rubber particles, soluble fraction, and precipitated bottom fraction. The floating rubber particle layer was collected and fractionated on SDS-PAGE, and the protein gel containing proteins from 10-kD to ~400-kD was divided into nine pieces. Each section was subjected to LC-MS/MS-based protein identification using MASCOT software to interrogate a reference lettuce transcriptome comprised of 29,417 unigenes. As a result, 372 unigenes were identified by at least 2 peptides with P<0.05 or 1 peptide with P<0.01. The proteins identified from the
lettuce rubber particles are listed in Supplemental Table 1.

Small rubber particle protein (SRPP) homologs, previously shown to be localized on rubber particles, were amongst the proteins identified; however, cis-prenyltransferase (CPT) was not initially identified. Upon careful examination, two lettuce unigenes were identified, which were annotated as homologs of Lew1 in Arabidopsis and Nogo-B receptor (NgBR) in human. Both genes have been implicated in dolichol metabolism (24, 26).

Sequence analysis of these two lettuce unigenes revealed that both are transcribed from the same gene, and the full-length cDNA, encoding a polypeptide of 244 amino acids, was isolated from lettuce cv. Mariska. Due to its low sequence homology to the conventional CPT sequences, this newly identified lettuce cDNA was named CPT-Like2 (CPTL2). One additional homolog, displaying 68.9% amino acid identity to CPTL2, was also identified from the lettuce transcript database and was named as CPTL1. The deduced protein sequences from the CPTL1/2 cDNAs show ~45% and ~25% identity to Lew1 and NgBR, respectively.

As the focus of this work is NR biosynthesis in plants, CPTL homologs were also searched and identified from the rubber tree and dandelion. The rubber tree has one CPT while the dandelion has two CPTs in the public database. To gain a better insight into the CPTL sequences, the protein sequences of these CPTLs were aligned with other CPTs including the CPTs from E. coli and yeast (Figure 2). The three-dimensional crystal structure and site-directed mutagenesis of the prokaryotic CPTs have revealed five conserved motifs, most of which are involved in substrate-binding, catalysis, or structural interactions (32, 33 and references therein). Although weak sequence similarities between CPTLs and CPTs were found, none of the five motifs of CPTs was conserved in the CPTLs (Figure 2), implying that CPTLs are not likely to have the catalytic functions found in the conventional CPTs.

**Latex-specific expression of CPTL2** - The relative transcript abundance of CPTL2 was examined in different lettuce tissues, including latex isolated from three different developmental stages of lettuce by quantitative real-time PCR (qRT-PCR). For comparison, two conventional CPTs (CPT1/2), possessing conserved catalytic motifs, were also identified from the lettuce transcript database and included in the qRT-PCR analysis. CPTL2 was exclusively expressed in latex with the highest expression in the latex of the flowering stage (up to 32-fold higher in latex relative to leaves; Figure 3A). This latex-specific expression was not observed for its closest homolog, CPTL1, and two other CPT genes (CPT1 and CPT2). CPTL1 and CPT1 displayed ubiquitous expression in all tissues while CPT2 showed a higher level of expression in green leaves than other tissues. This qRT-PCR result suggests the possible involvement of CPTL2 in NR biosynthesis in lettuce.

**Silencing of CPTL2 in lettuce causes a decrease in NR** - Next, we examined the function of CPTL2 using RNA-interference (RNAi) in lettuce. Three independent CPTL2-RNAi constructs, driven by the cauliflower mosaic virus 35S promoter, were designed from different regions of the CPTL2 transcript (denoted as RNAi-a, -b, and -c; Figure 4A). The use of three independent constructs in the silencing experiments reduces the occurrence of false positives arising from off-target effects. A total of 13 transgenic lettuce lines were established using Mariska (for RNAi-a) and Ninja (RNAi-b and RNAi-c) cultivars. For all 13 transgenic lines, three individual T2 plants from an independent T1 transgenic line were used to measure the relative abundance of NR and CPTL2 transcripts. Non-transformed lettuce, vector-transformed lettuce, and wild-type segregant from the T2 generation were used as control plants. Of these, six transgenic lettuces (RNAi-a1, a3, b1, b3, c2, and c4) derived from three distinct CPTL2-RNAi constructs showed less than 5% of NR in comparison to controls (Figure 4B, Table 2 in bold characters). The relative CPTL2 transcript levels of these six lines were reduced accordingly to 3-15%. Conversely, two transgenic lines without NR reduction (RNAi-a2 and c8) did not show any corresponding reduction of CPTL2 transcripts. Overall, the NR abundance and CPTL2 transcripts in 13 transgenic lettuces showed a strong linear correlation (Figure 4B).

To further confirm that a decrease in CPTL2 transcript also reduces CPTL2 protein levels in transgenic lettuce, CPTL2 antibodies were generated; however, the antibodies produced were
not suitable for latex immunoblot assays due to cross-reactivity. Therefore, CPTL2 proteins from two independent transgenic lettuces (RNAi-a1 and RNAi-b1) were quantified by multiple reaction monitoring (MRM) using LC-MS/MS. Latex calcojen (25.5 kD) and small rubber particle protein 4 (SRPP4, 25.4 kD), both of which migrate closely with CPTL2 (28.4 kD) were used as internal calibration proteins. Prior to the experiment, recombinant CPTL2, calcojen, and SRPP4 proteins were generated and used to select the most suitable peptides for MRM. The optimized MRM method was then used to calculate the relative quantity of CPTL2 in reference to calcojen and SRPP4 in latex. The relative CPTL2 protein abundance in the RNAi-a1 and RNAi-b1 latex lines were determined to be 19.2% and 18.3% of the controls, respectively (two arrows in Figure 4B and Supplemental Table 2). Therefore, significant decreases of CPTL2 transcript, CPTL2 protein, and NR were observed in the transgenic plants. Although NR was markedly decreased in transgenic lettuces, the $M_w$ and Pd values of NR from CPTL2-silenced lettuce were not different from those of the controls (Table 2). Therefore, we concluded that the silencing of CPTL2 in lettuce results in marked reduction of NR in latex without altering the NR polymer properties.

The chemical phenotype of latex from transgenic plants was apparent in GPC-chromatographic separation (Figure 4C). However, the reduction of the floating rubber particles using a centrifugal method was only noticeable upon close observation (Figure 5A). The thickness and dry weight of the rubber particle layers from the CPTL2-silenced lettuce were approximately 60% of those from the controls. We reasoned that the dramatic decrease of NR from CPTL2-silenced lettuce should be observable in latex. Therefore, we attempted to develop a simple method to sufficiently visualize the NR deficiency in latex. It was found that the addition of an appropriate amount of sodium dodecyl sulfate to latex allows the formation of optically dense NR colloids in NR-rich controls after centrifugation (Figure 5B; the first three samples). Such NR colloids were not detected in strong CPTL2-silenced lettuce lines, due to insufficient amounts of NR. Also, GPC-HPLC analysis was unable to detect NR from the non-colloidal, transparent portion, as expected (Figure 5C). Taken together, these results suggest that CPTL2-silencing causes a significant reduction of NR in lettuce latex, and the corresponding chemical phenotype can be visualized by simple centrifugation. We conclude that CPTL2 is necessary for NR biosynthesis in lettuce.

**Identification of latex-specific conventional CPT (CPT3)** - During the course of this work, RNAi-mediated silencing of three highly homologous CPTs (TbCPT1-3; >96% amino acid identity to each other) in dandelion (T. brevicorniculatum) showed the reduction of NR in this plant (20). TbCPTs display very low homology to CPTL2 (17% amino acid identity), and importantly they possess all five conserved motifs required for CPT activity. Thus, these three TbCPTs are conventional CPTs and are clearly distinguishable from the CPTL2-type. As lettuce and T. brevicorniculatum belong to the same subfamily, Cichorioideae, in the Asteraceae family, we examined the presence of the TbCPT ortholog in lettuce. A TbCPT ortholog was not identified in the original lettuce transcript dataset used for the LC-MS/MS proteomic analysis, which explains the failure of locating a TbCPT ortholog from our proteomics data. However, a TbCPT ortholog in lettuce, which shares >82% amino acid identity with the TbCPT1-3 proteins, was identified from an updated lettuce transcriptomics dataset (updated at NCBI in April 2011 by Dr. Michelmore, UC Davis). This CPT was denoted lettuce CPT3 to distinguish it from the two previously described CPTs (lettuce CPT1/2). qRT-PCR analysis of CPT3 in various lettuce tissues showed that CPT3 expression closely resembled that of CPTL2. CPT3 was exclusively expressed in latex with the highest level of expression in the latex of the flowering stage (Figure 3B).

We questioned whether the abundance of CPT3 protein would be negatively affected in CPTL2-silenced lettuce. MRM analysis showed that the relative abundance of CPT3 in RNAi-a1 and RNAi-b1 transgenic lettuce decreased to 31.8% and 46.5% of that in control lettuce, respectively (Supplemental Table 2). On the other hand, CPT3 transcripts in CPTL2-silenced lines were not significantly affected (RNAi-a1, 0.83 ± 0.04; RNAi-b1, 1.08 ± 0.08). These data implied that...
CPT3 protein levels decreased when CPTL2 is silenced in transgenic lettuce.

**Co-localization of CPTL2 and CPT3 on ER in tobacco** - To examine the subcellular localization of CPTL2, green fluorescent protein (GFP) was fused to the C-terminus of CPTL2, and its localization was examined by confocal microscopy after transient expression in tobacco leaves. CPTL2-GFP was often localized to the perimeters of the round particle, and Nile red dye, known to stain lipophilic substances, stained the core of the round particles labeled by CPTL2-GFP (Figure 6A-C). Additionally, faint ER-signals from CPTL2-GFP were momentarily detected (60-72 hr after infiltration; Figure 6D-F), but this signal rapidly disappeared after 72 hr and only round-particles were detected. On the other hand, when the CPT3-GFP construct was expressed in tobacco leaves, most of the transfected cells showed diffused cytosolic patterns with occasional observations of a blurred ER-pattern (Figure 6G-I). Therefore, by individually expressing CPTL2 or CPT3, their subcellular locations do not entirely overlap with each other.

To allow CPTL2 and CPT3 encounter in the same cells, CPTL2-GFP and CPT3-RFP (red fluorescent protein) were co-expressed in tobacco epidermis. Surprisingly, CPT3 relocated to ER, resulting in clearly refined ER-patterns in both green- and red-fluorescent channels, and these two signals overlap entirely with each other (Figure 6J-L). To prove that the GFP signals are associated with ER, untagged CPTL2 and CPT3-GFP were transiently co-expressed with the mCherry-HDEL ER-marker. The GFP-signals from CPT3-GFP co-localize with the ER-marker (Figure 6M-O).

**Subcellular localization of CPTL2 and CPT3 in yeast** - A yeast strain endogenously expressing ERG6-RFP (sterol Δ24 methyltransferase fused with RFP) was used to examine the subcellular localization of CPTL2 and CPT3 in yeast. ERG6 is known to localize to ER/lipid-bodies in the log-phase and on lipid-bodies in the stationary phase. GFP was fused to the C-terminus of CPTL2 and CPT3, and these fusion constructs were individually expressed in a yeast strain endogenously expressing ERG6-RFP (Figure 7A-C). Also, by careful observations, we were able to detect the ER-pattern around the nucleus (Figure 7 arrows), although the signals were easy to miss because strong GFP signals from the lipid-bodies masked the much weaker ER signals. On the other hand, CPT3-GFP was localized in the cytosol (Figure 7D-F). By co-expressing CPTL2 and CPT3 (one GFP tagged and the other non-tagged), CPT3 relocated and clear yeast ER-pattern was observed for both GFP and RFP channels (Figure 7G-L). The observed signals on nuclear and periplasmic membranes (nuclear ER and cortical ER, respectively) are typical of ER-patterns in yeast (34). The localization data in yeast during the log phase closely mirrored those in tobacco cells.

At the stationary phase, yeast develops larger lipid-bodies as shown by an ERG6-RFP marker in Figure 8. At this stage, CPTL2-GFP was detected in lipid-bodies, of which the perimeters were labelled by ERG6-RFP (Figure 8A-C), but CPT3-GFP showed a diffused pattern in the cytosol (Figure 8D-F). Such a diffused CPT3-GFP localization pattern was, however, drastically altered by co-expression of non-tagged CPTL2. Clear re-localization of CPT3-GFP from cytosol to lipid-bodies were observed by their co-expression (Figure 8G-I).

In summary, as observed in tobacco cells, the co-expression of CPTL2 and CPT3 in yeast facilitates the co-localization of these two proteins on the ER during the log phase and on the lipid-bodies in the stationary phase.

**Evidence for CPTL2 and CPT3 interaction by yeast two-hybrid assay** - To examine whether the alterations of the localization are the result of the physical interaction between CPTL2 and CPT3, yeast two-hybrid assays (Y2Hs) were performed. To mask the putative signal-anchor domain, CPTL2 was fused to the C-terminus of the DNA binding domain (DB) as bait, whereas CPTL2 or CPT3 was fused to the C-terminus of the transcription activation (TA) domain as prey. When β-galactosidase assays were performed using yeast expressing the fusion constructs in all combinations, only yeast expressing DB-CPTL2 (bait) and TA-CPT3 (prey) showed reporter enzyme activity (Figure 9A). Basal level activities
of reporter enzyme were detected when yeasts were grown in non-inductive glucose medium. When the same experiments were conducted using the reverse orientation (e.g., CPT3 as bait and CPTL2 as prey), again only yeast expressing DB-CPT3 and TA-CPTL2 showed reporter enzyme activity, although the reporter activity from this combination was 4-fold lower than that from reversed combination (Figure 9A). In addition, the yeast transformants were tested for the ability to grow on amino acid drop-out medium, in which the essential amino acid leucine is absent and only produced when the prey and bait interact. Only yeast containing the DB-CPTL2/TA-CPT3 pair was able to grow (Figure 9B). No growth was observed from the yeast expressing DB-CPT3/TA-CPTL2, perhaps due to their weak interaction in this orientation, as shown in the reporter enzyme assays. These Y2H assay results further support the interaction between CPTL2 and CPT3.

**Biochemical assays using CPTL2 and CPT3 recombinant enzymes** - Catalytic activities of CPTL2 and CPT3 were examined by heterologous expression in *E. coli*. *E. coli*-produced, partially purified CPTL2, CPT3, and their mixture showed no CPT activity *in vitro* (data not shown) while the control enzyme, *E. coli* UPPS, prepared using the same method displayed potent activity (44.7 nmol IPP incorporation µg purified protein⁻¹ h⁻¹). As an alternative approach, these cDNAs were expressed in the yeast YPH499 strain. FLAG-epitope tagged CPTL2/CPT3 and their respective non-tagged versions were generated. In comparison, expression cassettes of an analogous pair, CPTL1/CPT1, were also constructed (CPT2 encodes a chloroplast targeting peptide and thus was excluded from this analysis). These cDNAs were expressed individually or simultaneously in yeast. Immunoblot analyses confirmed the presence of respective recombinant proteins in microsomes and total protein extract (Figure 10 A/C). To evaluate the biochemical activities of CPTL/CPT, microsomes containing CPTL/CPT were incubated with ¹⁴C-labelled IPPs and FPP, and radioactivities of the chloroform extract from the enzyme reactions were measured and separated by reverse phase-TLC (Figure 10B/D/E). In these assays, FLAG-tagged enzymes failed to show ¹⁴C-IPP incorporation activity above the vector-control activity for both CPTL1/CPT1 and CPTL2/CPT3.

However, when their respective non-tagged versions were co-expressed, discernable ¹⁴C-IPP incorporated polyisoprene products were observed on TLC in comparison to those from vector-control and FLAG-tagged enzymes (Figure 10B/D). Chloroform extract of the enzyme reactions showed that CPTL1/CPT1 (or CPTL1/CPT1-FLAG) showed a strong IPP incorporation activity (32.9 pmol IPP h⁻¹ µg⁻¹ of microsome), which is 13-fold (±0.4, n=3, p<0.001) higher activity than the vector-control activity (Figure 10E). Although marginal, CPTL2/CPT3 (or CPTL2/CPT3-FLAG) also showed 2.5-fold (±0.3, n=4, p<0.05) higher IPP incorporation activity (5.9 pmol IPP h⁻¹ µg⁻¹ of microsome) than the vector control activity. Notably, microsomes containing CPTL1/CPT1 incorporated 32% of the total IPP added in the reactions into *cis*-polyisoprenes in 1-hr incubation while no ¹⁴C-IPP incorporation above the background was detected from CPT1 alone, even though its protein was detected on microsomes by immunoblot. It should be also noted that only FLAG-tagged CPTL1 or CPTL2 effectively abolished CPT activity, but FLAG-tagged CPT1 or CPT3 had no influence on CPT activity when paired with its corresponding non-tagged CPTL (see mix in Figure 10B/D), indicating that C-terminal FLAG-epitope tag in the CPTL protein blocks CPT activity in the assays. This serendipitous finding provides additional evidence for the interaction and modulation of CPTL on CPT.

We further tested the alternatively paired co-expressions in the enzyme assay (i.e., CPTL1/CPT3 or CPTL2/CPT1). When CPT1 and CPT3 were paired with CPTL2 and CPTL1, respectively, about 70% catalytic activities were obtained in reference to their natural pairs (Figure 10 F/G). In particular, efficient dolichol synthesis could be still obtained from the CPT1/CPTL2 pair, implying both CPTL1 and CPTL2 support the CPT1 dolichol biosynthetic activities. On the other hand, CPT3 activity was still low when it was paired with CPTL1.

As shown in Figure 8, the CPT re-localization to the lipid-bodies in the presence of CPTL1 is a simple and reliable method to assess the influence of CPTL on CPT localization *in vivo*. We examined if CPTL1 or CPTL2 can influence the localization of CPT1 or CPT3 by expressing their cDNAs in a pairwise combination. In these...
sequence features in mind, these conventional CPTs (Figure 2). With such aberrant CPTL2 lacks all five conserved motifs of the polyisoprene synthesis. It is worth noting that CPTL2 itself is not sufficient for NR and demonstrating in a dramatic decrease of NR in latex, NR synthetic activity containing CPTL2 from yeast did not show any CPTL2 from the CPTL2/CPT3 assays (Figure 10B). In addition, overall IPP incorporation to polyisoprene was significantly weaker for CPTL2/CPT3 in comparison to CPTL1/CPT1, and the change in CPTL partner did not alter the status of the CPT activity significantly.

**DISCUSSION**

**CPTL2 is necessary but not sufficient for NR biosynthesis in lettuce** - NR biosynthesis remains largely unknown in plants. One key finding from this work is that CPTL2-silenced lettuce resulted in a dramatic decrease of NR in latex, demonstrating CPTL2 is necessary for NR biosynthesis. However, purified recombinant CPTL2 from E. coli and the microsomes containing CPTL2 from yeast did not show any NR synthetic activity in vitro. These data suggest that CPTL2 itself is not sufficient for NR and cis-polyisoprene synthesis. It is worth noting that CPTL2 lacks all five conserved motifs of the conventional CPTs (Figure 2). With such aberrant sequence features in mind, these in vitro assay data are not unexpected and are indicative of the non-enzymatic nature of CPTL2 in NR biosynthesis.

Similarly, the most studied CPTL2 homolog, human NgBR, also has no known catalytic activity. Lacking all enzymatic activity, NgBR is able to interact with human CPT (24). Recently, NgBR and its homologs in mice and yeast (NUS1) were identified as necessary subunits for CPT activities in respective organisms (35). NgBR also interacts with NogoB and NPC2 for entirely unrelated physiological events, such as remodeling vascular epithelial cells and controlling intracellular trafficking of cholesterol, respectively (23,25). Therefore, NgBR operates as an ER-residing, multifaceted scaffolding protein in humans. Considering the known NgBR functions and CPTL2 data from this work, we proposed that CPTL2 is a scaffolding protein for CPT3 in lettuce.

**CPTL homologs in Arabidopsis and yeast are LEW1 and NUS1 (YDL193W), respectively, and their null mutations in each organism are known to be lethal (26,36).** However, no apparent developmental abnormality was observed in the CPTL2-silenced lettuce except for NR reduction. This observation indicates that CPTL2 is not essential for survival but has a specialized role for NR biosynthesis in lettuce latex. On the other hand, the second homologous pair, CPTL1 and CPT1, showed ubiquitous expression patterns in various lettuce tissues (Figure 3), and the microsomes containing both proteins exhibited an efficient synthesis of short cis-polyisoprenes, equivalent to dolichols (Figure 10D). Therefore, the CPTL1/CPT1 pair is responsible for the biosynthesis of the primary metabolite, dolichols, in lettuce. Lettuce appears to have undergone a gene-duplication resulting in the CPTL1/CPT1 pair for dolichol synthesis in all cells and a specialized CPTL2/CPT3 pair for NR biosynthesis in latex.

**CPTL2 and CPT3 forms a protein complex on ER** - A rubber synthase complex has been suggested on rubber particles, yet no tangible experimental evidence has so far been reported. From this work, three lines of evidence support the interaction between CPTL2 and CPT3 to form a protein complex. First, in tobacco epidermis and log phase yeast, we showed that CPT3 relocates from cytosol to ER, when CPTL2 is co-expressed (Figure 6J-L; Figure 7G-L). Second, Y2H assays support direct CPTL2 and CPT3 interaction in yeast. Third, in CPTL/CPT-overexpressed yeast, FLAG-tagged CPTL1 or CPT2 effectively blocked CPT activities, whereas non-tagged CPTL1 or CPT2 allowed CPT activities. These results suggest that C-terminal modified CPTL1/2 can render the CPTL/CPT complex inactive. Collectively, these data indicate that CPTL2 and CPT3 form a hetero-protein complex.

Although the observed protein complex is conceptually new in NR, hetero-protein complexes have been previously reported for enzymes involved in trans-isoprenoid biosynthesis. Specifically, in order to acquire catalytic activities, GPP synthases from three plants (peppermint, snapdragon, and Clarkia breweri) (37-39) and heptaprenyl pyrophosphate synthase from M. luteus (40) require heterodimer formation. In these cases, a catalytic TPT subunit needs to form a complex with a non-catalytic subunit that displays overall sequence homology to the catalytic unit.

In conclusion, efficient in vitro synthesis of the dolichol equivalents was demonstrated from the CPTL1/CPT1 assays (Figure 10D), but no evidence of high molecular weight NR synthesis was obtained from the CPTL2/CPT3 assays (Figure 10B). In addition, overall IPP incorporation to polyisoprene was significantly weaker for CPTL2/CPT3 in comparison to CPTL1/CPT1, and the change in CPTL partner did not alter the status of the CPT activity significantly.
These TPT complexes, composed of a non-catalytic and a catalytic protein, closely resembles the CPTL2 and CPT3 data in this work.

Yeast shows distinct physiological features at a stationary phase, and one of the main changes is the formation of lipid-bodies (also called lipid-droplets) originating from the ER (41). When nutrition becomes depleted and cell density increases, ER-leaflet of the yeast creates monolayered lipid-bodies, where triacylglycerols are stored. It is intriguing to observe the localization of the CPTL2-CPT3 complex on lipid-bodies during the yeast stationary phase. In many aspects, the rubber particles in the latex of rubber tree resemble lipid-bodies in yeast because both have monolayer membranes encapsulating lipophilic substances - triacylglycerol in the lipid-body and NR in rubber particles (42-44). Their membranes have abundant integral membrane proteins (oleosin in the lipid-body and SRPP in the rubber particle) which represent the major components of the membrane proteins. These proteins are known to have structural roles by preventing the particle aggregation via charge-repulsion (45,46). Although yeast is a heterologous system for NR metabolism, from the observation of localization of CPTL2/CPT3 on ER in the log phase and on lipid-bodies in the stationary phase, this may explain how ER-synthesized NR deposited in rubber particles, analogous to the triacylglycerol deposition in lipid-bodies that bud off from the ER (43).

Based on our localization data and the similarities of rubber particles to lipid-bodies, we propose a model that CPTL2 anchors CPT3 on ER to form a stable protein complex with its catalytic domain facing cytosol (Figure 12). It can be inferred from this model that substantial NR could be synthesized on ER, and the monolayer rubber particles filled with NR may bud off from ER. Some residual enzymes and their activities may be transferred to and retained on the rubber particles during rubber particle biogenesis from the ER, therefore accounting for the reported biochemical competency of rubber particles (19). As demonstrated here, the localization of CPTL2 and CPT3 on yeast lipid-bodies could offer a convenient experimental system to study NR biosynthesis and storage in yeast. For example, synthesis and accumulation of NR or shorter cis-isoprenes in yeast lipid-bodies could be traced by feeding radio-labelled mevalonic acid. However, production of NR from the recombinant yeast will be ultimately limited by the low abundance of lipid-bodies in yeast, and thus genetically altering the plants that synthesize low quality NR to produce high quality NR is likely to be a more realistic avenue, once NR biosynthetic mechanism is fully elucidated.

Biochemical activities of CPTL and CPT - Since the discovery of a prokaryotic CPT, which displays an entirely different sequence and tertiary structure from TPT (5), NR research has focused on identifying distinct CPTs from NR-producing species (e.g., rubber tree and dandelion) with an aim of synthesizing NR in vitro. However, NR biosynthetic activity by purified CPTs has not been reported. In one study, the addition of a latex part to the purified CPT from E. coli allowed synthesis of high molecular weight NR (21). However, using yeast recombinant CPT, this NR-synthetic activity was not replicated in similar conditions, and only regular dolichols (C80-C100) were produced (22). From dandelion, CPT activity was not directly examined in vitro using recombinant CPT but was inferred from the complementation of a rer2 mutant (16). It is important to note that, to date, all attempts to synthesize NR in vitro have centered on conventional CPT, and the NR-synthetic activity of “conventional CPT together with CPTL” has not been investigated.

In this work, CPT activities from the two pairs of CPTL/CPT (i.e., CPTL2/CPT3 and CPTL1/CPT1) were examined as part of microsomes. We found that only the microsomes from the CPTL/CPT co-expressed yeast showed cis-polyisoprene patterns distinct from the vector-control activity (Figure 10B/D). Notably, robust dolichol synthesis was reconstituted in vitro using CPTL1/CPT1, but CPTL1 or CPT1 alone did not show activity above the background, despite the presence of these proteins in microsomes. These biochemical data support our proposed role of CPTL1 as a “CPT-scaffolding protein” that can modulate CPT activity, and it becomes apparent that both CPTL1 and CPT1 are required for efficient cis-polyisoprene synthesis. This finding is further supported by recent CPT activity assays, co-expressing human NgBR/CPT or yeast NUS1/CPT (35). Regrettably, in CPTL2/CPT3
assays, isoprene polymers larger than the regular dolichols could not be synthesized. Also, the rate of IPP incorporation into dolichols from CPTL2/CPT3 assay was markedly lower than that from CPTL1/CPT1 assay. It is evident that we were not able to synthesize high molecular weight NR by co-expressing CPTL2 and CPT3 in yeast. Therefore, present studies showed that CPTL2 is required for NR biosynthesis in vivo, but a simply blending CPTL2 and CPT3 in vitro did not result in NR biosynthesis. Although clear explanation cannot be given from the present data, we speculate that undesirable modifications of CPTL2/CPT3 in yeast could inhibit the NR synthetic activity of the CPTL2/CPT3 complex, or other latex component may be required to render the CPTL2/CPT3 complex fully active for NR synthesis. Alternatively, a higher dimensional nano-structure, as shown in the cellulose synthase complex (47), could be a prerequisite for NR biosynthesis in plant. Apparently, full in vitro synthesis of NR still needs further investigation, and we showed here that lettuce can serve as a powerful system to apply molecular genetics tools to study NR metabolism.

**CPTL in the rubber tree** – We questioned if the CPTL is present in other NR-producing plants and, if so, the CPTL gene duplication also occurred. In the public database, two CPTLs were found in dandelion which belongs to the same subfamily of lettuce (Chicorioideae), supporting the necessity of a specialized CPTL for NR synthesis by a gene duplication (Figure 2). However, only one member of the CPTL was identified in H. brasiliensis from 14.3 million transcript reads (Figure 2). Considering that two copies of CPTLs were identified from a relatively small number of transcript reads (<80,000 ESTs) in lettuce and dandelion, insufficient sequencing is not likely the reason for failing to identity the second copy of CPTL in the rubber tree. Instead, this analysis suggests that CPTL duplication is a recent event in the Chicorioideae subfamily of the Asteraceae family to which both lettuce and dandelion belong.

The rubber tree, on the other hand, did not undergo a gene duplication to develop a specialized CPTL in latex. Since CPTL is clearly required in eukaryotes for cis-polyisoprene synthesis (whether it is dolichol or NR) (35 and this work), the rubber tree should also require CPTL for dolichol and NR biosynthesis. We propose that the single copy of CPTL in the rubber tree has a dual role in supporting both the NR synthesis in latex and the ubiquitous dolichol synthesis in other cells. This prediction can be further examined by biochemical and gene expression studies of the rubber tree CPTL in the future.

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FIGURE LEGENDS

Figure 1. Enzymatic synthesis of *cis*-polyisoprenes.
FPP: Farnesyl diphosphate; IPP: Isopentenyl diphosphate

Figure 2. Sequence alignment of various CPT and CPTL proteins.
Green boxes are amino acids conserved in CPTs. Detailed information can be found in references 32 and 33. Abbreviations used are as follows. To: *Taraxacum officinallis*; Hb: *Hevea brasiliensis*; Ec: *Escherichia coli*; Sc: *Saccharomyces cerevisiae*; Ls: *Lactuca sativa*; Tb: *Taraxacum brevicorniculatum*. NogoBR (BC150654), AtLew1 (At1G11755), HbCPTL (JT945746), ToCPTL1 (DY820019), ToCPTL2 (DY832694), TbCPT (JQ991925), HbCPT1 (AB061234), HbCPT2 (AB064661), EcUPPS (EU906097), ScRER2 (AB013497).

Figure 3. Relative quantitation of CPT and CPTL transcripts in lettuce.
A, Relative *CPT1/2* and *CPTL1/2* transcripts from various lettuce (cv. Mariska) tissues were measured by qRT-PCR, relative to the respective transcripts in leaf (1st true leaf). B, Relative *CPT3* transcript levels were measured in the same tissues as in A relative to the leaf transcript level (cv. Mariska). Data are means ± SD from three biological replicates, each including three technical replicates. Actin was used to normalize the data.

Figure 4. Experimental design and characterization of CPTL2-silenced lettuce.
A, The positions of the different RNAi-silencing constructs (RNAi-a,b,c) in the *CPTL2* are shown and color-coded. B, Relative abundance of the *CPTL2* transcript is plotted against relative NR abundance from 13 independent transgenic lettuces. Data points in red, green, and blue are from RNAi-a, -b, and -c constructs, respectively. Each data is calculated from three independent T2 plants derived from an independent T1 transgenic line. NR abundances are from three biological replicates, and relative transcripts are from three technical and three biological replicates. Two transgenic lettuces designated by red and green arrows were subjected to the relative CPTL2 protein quantification by multiple-reaction monitoring. Numerical data and lettuce cultivars are given in Table 2. C, Chromatographic separation of the NR by HPLC-GPC. Wild-type lettuces (Ninja and Mariska) were used as controls.

Figure 5. Latex phenotypes from CPTL2-silenced lettuces.
A, Latex layers after centrifugation are shown from CPTL2-silenced and control lettuces (Mariska, vector-transformed, and segregant sibling). 350 mg latex (fresh weight) was dissolved in 1 mL extraction buffer as described in the Experimental Procedures. B, Presence or absence of NR in latex from control or CPTL2-silenced lettuces is visualized after SDS addition by centrifugation. C, HPLC-GPC analysis of transparent (RNAi-a1) and colloidal (Mariska) portions after centrifugation. 30 mg latex (fresh weight) was used for each experiment.

Figure 6. Subcellular localization of CPTL2 and CPT3 in tobacco.
Confocal microscopy was used to capture fluorescent signals from tobacco epidermal cells after transient expression of the various constructs as described below. A-C, a single expression of CPTL2-GFP; D-F, double expression of CPTL2-GFP and mCherry-HDEL; G-I, double expression of CPT3-GFP and mCherry-HDEL; J-L, double expression of CPTL2-GFP and CPT3-RFP; M-O, triple expression of CPTL2 (untagged), CPT3-GFP, and mCherry-HDEL. GFP channels are shown in A, D, G, J, and M; RFP channels are shown in E, H, K, N; Nile red channel is shown in B. Images from fluorescent channels, as well as those of bright field, were combined in C, F, I, L, and O. The bar in A indicates 25 μm.

Figure 7. Subcellular localization of CPTL2 and CPT3 in the log phase yeast.
Yeast cells at log phase (8-hr after inoculation) were observed by confocal microscopy. CPTL2-GFP (A-C) or CPT3-GFP (D-F) was expressed alone in yeast endogenously expressing ERG6-RFP (ER and lipid-
body marker). Arrows indicate nuclear ER in A. *CPT3-GFP* and untagged *CPTL2* (G-I) or *CPTL2-GFP* and untagged *CPT3* (J-L) were co-expressed. The bar in A indicates 5 μm.

**Figure 8. Subcellular localization of CPTL2 and CPT3 in the stationary phase yeast.**
Yeast cells at stationary phase (20-hr after inoculation) were observed by confocal microscopy. *CPTL2-GFP* (A-C) or *CPT3-GFP* (D-F) was expressed alone in yeast endogenously expressing *ERG6-RFP*. *CPT3-GFP* and untagged *CPTL2* was co-expressed to examine the change of *CPT3-GFP* localization by *CPTL2* (G-I). The bar in A indicates 5 μm.

**Figure 9. Yeast 2-hybrid assays.**
A, β-galactosidase activity from different combinations of CPTs as bait and prey. Data represents mean ± SD from at least three individual transformants. B, Growth of corresponding yeast on SC medium lacking leucine.

**Figure 10. CPT enzyme assays and immunoblots using CPT and CPTL recombinant proteins**
A/C, Immunoblots of C-terminal FLAG-tagged CPTL1/2 and CPT1/3 by anti-FLAG antibodies using microsomes and total proteins. The blots were stripped and re-hybridized with anti-DPM1 (Dol-P-Man synthase) antibodies as controls. Reaction products from the enzyme assays using recombinant CPTL2/CPT3 (B), CPTL1/CPT1 (D) or alternative parings (F) were separated by reverse phase-TLC. Total activities extracted by chloroform are shown in E and G. Data represent mean ± S.D. (n ≥ 3)

**Figure 11. Altered subcellular localization of CPT1 or CPT3 with co-occurrence of CPTL1 or CPTL2 in the stationary phase of yeast.**
Yeast cells at stationary phase (20-hr after inoculation) were observed by confocal microscopy. *CPT1-GFP* (A-C) or *CPT1-GFP* (D-F) was expressed alone in the yeast expressing *ERG6-RFP* (a lipid-body marker). CPT1-GFP re-localized to lipid-bodies when co-expressed with CPTL1 (G-I) or CPTL2 (J-L). CPT3-GFP also re-localized to lipid-bodies when co-expressed with CPTL1 (M-O). Bar = 5 μm.

**Figure 12. A schematic model of the NR biosynthesis on ER.**
This proposed model illustrates the NR biosynthesis on ER from which rubber particles may bud off. CPTL2 resides on ER where it recruits CPT3 to form a stable protein complex. NRs are shown in red color between the membrane monolayers.
Table 1. Weight average molecular weight ($M_w$) and polydispersity of NR from *Lactuca sativa* cv. Mariska and Ninja.

|                     | *L. sativa* cv. |               |               |
|---------------------|-----------------|---------------|---------------|
|                     | *Mariska*       | *Ninja*       |               |
| $M_w$ (million g/mol)$^1$ | 1.50 ± 0.07    | 1.45 ± 0.03   |               |
| Polydispersity$^1$   | 1.53 ± 0.06     | 1.49 ± 0.10   |               |
| Flowering time$^2$   | 135 ± 4         | 84 ± 9        |               |

$^1$Values are mean ± SD from three individual plants.

$^2$Number of days from germination to the first flowering. Mean ± SD, Mariska, n=14; Ninja, n=15.
Table 2. Weight average molecular weight ($M_w$) and polydispersity (Pd) of cis-polyisoprene from CPTL2-silenced lettuce

| plant          | $^1$$M_w$ (million g/mol) | $^1$Pd | $^1$Relative abundance of cis-polyisoprene |
|----------------|--------------------------|--------|-----------------------------------------|
| Mariska control| 1.29 ± 0.04              | 1.22 ± 0.04 | 1.00 ± 0.12                        |
| CPTL2-a1       | 1.20 ± 0.09              | 1.31 ± 0.03 | **0.05 ± 0.01**                      |
| CPTL2-a2       | 1.28 ± 0.05              | 1.50 ± 0.02 | 0.85 ± 0.22                         |
| CPTL2-a3       | 1.15 ± 0.07              | 1.32 ± 0.08 | **0.05 ± 0.01**                      |
| CPTL2-a1       | 1.39 ± 0.06              | 1.22 ± 0.04 | **0.04 ± 0.01**                      |
| CPTL2-b1       | 1.36 ± 0.12              | 1.39 ± 0.07 | 0.13 ± 0.08                         |
| CPTL2-b2       | 1.32 ± 0.06              | 1.29 ± 0.12 | **0.04 ± 0.01**                      |
| CPTL2-c1       | 1.13 ± 0.15              | 1.43 ± 0.05 | 0.09 ± 0.05                         |
| CPTL2-c2       | 1.28 ± 0.08              | 1.32 ± 0.13 | **0.04 ± 0.01**                      |
| CPTL2-c3       | 1.25 ± 0.04              | 1.40 ± 0.04 | 0.10 ± 0.02                         |
| CPTL2-c4       | 1.26 ± 0.02              | 1.44 ± 0.03 | **0.03 ± 0.01**                      |
| CPTL2-c5       | 1.31 ± 0.04              | 1.37 ± 0.07 | 0.15 ± 0.09                         |
| CPTL2-c7       | 1.24 ± 0.06              | 1.43 ± 0.01 | 0.29 ± 0.04                         |
| CPTL2-c8       | 1.24 ± 0.06              | 1.44 ± 0.01 | 0.77 ± 0.14                         |

$^1$The values of the RNAi-lines are means ± SD (n=3) and were measured from three T2 transgenic plants derived from independent T1 transgenic lines. $^2$Data for controls were generated from two of non-transformed lettuce (Mariska for CPTL2-a line or Ninja for CPTL2-b/c lines), two of vector-transformed lettuce, and two of wild-type segregants from the T2 population (n=6).
### Table 3. A list of primers used in this work.

| Primer name | Primer Sequence |
|-------------|-----------------|
| 1 | Gateway-Universal-F | GGGGACAAGTTTGTACAAAAAAGCAGGCTTC |
| 2 | Gateway-Universal-R | GGGGACCACCTTTGTACAAAAAAGCAGGCTTC |
| 3 | RNAi-a-F | AAAAAGCAGGCTCTATGGATCTCGTAAAG |
| 4 | RNAi-a-R | CAAGAAGGCTGGGTCACATACACACCAAA |
| 5 | RNAi-b-F | AAAAAGCAGGCTCTATGGATCGTAAAG |
| 6 | RNAi-b-R | CAAGAAGGCTGGGTCACATACACACCAAA |
| 7 | RNAi-c-F | AAAAAGCAGGCTGGGTCACATACACACCAAA |
| 8 | RNAi-c-R | CAAGAAGGCTGGGTCACATACACACCAAA |
| 9 | Gateway-CPTL2-R | CAAGAAGGCTGGGTCACATACACACCAAA |
| 10 | Gateway-CPTL2-TGA-R | CAAGAAGGCTGGGTCACATACACACCAAA |
| 11 | Gateway-CPT3-F | AAAAAGCAGGCTCTATGGATCTCGTAAAG |
| 12 | Gateway-CPT3-R | CAAGAAGGCTGGGTCACATACACACCAAA |
| 13 | Gateway-CPT3-TGA-R | CAAGAAGGCTGGGTCACATACACACCAAA |
| 14 | pUG23-CPTL2-F | GTTCATCTGGTATGGATCTCGTAAAG |
| 15 | pUG23-CPTL2-R | GTTCATCTGGTATGGATCTCGTAAAG |
| 16 | p415GPD-CPTL2-F | TAGAATTCTGGAGTCACATATGCTGATGGTGAAC |
| 17 | p415GPD-CPTL2Flag-R | CGTCATCTGGTATGGATCTCGTAAAG |
| 18 | pUG23-synCPT3-F | GTTCATCTGGTATGGATCTCGTAAAG |
| 19 | pUG23-synCPT3-R | GTTCATCTGGTATGGATCTCGTAAAG |
| 20 | p416GPD-synCPT3Flag-R | TAGAATTCTGGAGTCACATATGCTGATGGTGAAC |
| 21 | p416GPD-synCPT3Flag-R | GTTCATCTGGTATGGATCTCGTAAAG |
| 22 | p416GPD-synCPT3Flag-R | GTTCATCTGGTATGGATCTCGTAAAG |
| 23 | pGAL-LexA-CPTL2-F | TAGCGGCCGCTTATGGATCTCGTAAAG |
| 24 | pGAL-LexA-CPTL2-R | ACTCCTCGAGTTAAGAACCGTAGTTCTGCTTC |
| 25 | pJG4-6-CPTL2-F | GTTCGCGGCCGCTGATAGGACCCTATCATCAG |
| 26 | pJG4-6-CPTL2-R | GTTCGCGGCCGCTGATAGGACCCTATCATCAG |
| 27 | pGAL-LexA-SynCPT3-F | TAGCGGCCGCTGATAGGACCCTATCATCAG |
| 28 | pGAL-LexA-SynCPT3-R | ACTCCTCGAGTTAAGAACCGTAGTTCTGCTTC |
| 29 | pJG4-6-SynCPT3-F | GTTCGCGGCCGCTGATAGGACCCTATCATCAG |
| 30 | pJG4-6-SynCPT3-R | GTTCGCGGCCGCTGATAGGACCCTATCATCAG |
| 31 | Gateway-SRPP4-F | AAAAAGCAGGCTGTATGGATCGAAGGGGAGGAGG |
| 32 | Gateway-SRPP4-R | AAAAAGCAGGCTGTATGGATCGAAGGGGAGGAGG |
| 33 | Gateway-Caleosin-F | AAAAAGCAGGCTGTATGGATCGAAGGGGAGGAGG |
| 34 | Gateway-Caleosin-R | AAAAAGCAGGCTGTATGGATCGAAGGGGAGGAGG |
| 35 | qPCR-Actin7-F | GAGATGAGGACAAATCCAAAAGAGG |
| 36 | qPCR-Actin7-R | GAGATGAGGACAAATCCAAAAGAGG |
| 37 | qPCR-CPT1-F | GAGATGAGGACAAATCCAAAAGAGG |
|   | Primer    | Sequence                        |
|---|-----------|---------------------------------|
| 39 | qPCR-CPT1-R | TCTAGAACAGCATGGAAGATTCATCGGT |
| 40 | qPCR-CPT2-F | TCGCCTTTCATCTGACAACCTG   |
| 41 | qPCR-CPT2-R | ACTCGATTTCGCTCCCTGACATA      |
| 42 | qPCR-CPT3-F | GCTGAAACAAGCGGAAGACGAAAG     |
| 43 | qPCR-CPT3-R | CTGGAATCGGGCGTGAAGAATGA       |
| 44 | qPCR-CPTL1-F | ATGAATGTGGCAGAGGATTCCAACAGA |
| 45 | qPCR-CPTL1-R | CCCTAAGAATCAAGTGGAGAACATGCCATA |
| 46 | qPCR-CPTL2-F | GGCCTATTCGTTATACAGAGATGGTACAC |
| 47 | qPCR-CPTL2-R | AGAACGTCATGGCTCCTCACCTTG      |
| 48 | p415GPD-CPTL1-F | TAGAACTAGTGGAATCCCAACATGAATGTCAGAGGATTTTC |
| 49 | p415GPD-CPTL1Flag-R | CTCAATCCCTGTAATCCCAACATGAATGTCAGAGGATTTTCGCCCTG |
| 50 | p415GPD-CPTL1N-R | CGAATTCCTGCAGCCCTTATGAACCATAATGTTTTTGCTTGACC |
| 51 | pUG23-CPT1-F | TCGAGGATCCATGGAATTTAAAAAAAG   |
| 52 | pUG23-CPT1-R | GCGTGTGACTACATCGCTCCATACATCTTTTTG |
| 53 | Flag-R        | TTTCAAAGCCTGAAATTCACATCTGTCGTCATCTGGTAATCCCC |
Figure 1
Figure 2
Figure 3
Figure 4
Figure 5
Figure 6
Figure 7

GFP

RFP (Erg6)

Overlay

CPTL2-GFP

D

E

F

CPT3-GFP

G

H

I

CPT3-GFP + CPTL2

J

K

L

CPTL2-GFP + CPT3

Figure 7

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Figure 8
Figure 9
Figure 10
CPTL1-GFP
CPT1-GFP
CPTL2
CPT3-GFP
CPT1-GFP + CPTL1
CPT1-GFP + CPTL2
CPT3-GFP + CPTL1

GFP
RFP (Erg6)
Overlay

Figure 11
A lettuce (*Lactuca sativa*) homolog of human Nogo-B receptor interacts with cis-prenyltransferase and is necessary for natural rubber biosynthesis

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