Cytokine secretion requires phosphatidylcholine synthesis

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Abstract

Cytidine-cytidyltransferase (CCT) is the rate-limiting enzyme in the phosphatidylcholine biosynthetic pathway. Here, we demonstrate that CCTα-mediated phosphatidylcholine synthesis is required to maintain normal Golgi structure and function as well as cytokine secretion from the Golgi complex. CCTα is localized to the trans-Golgi region and its expression is increased in lipopolysaccharide (LPS)-stimulated wild-type macrophages. Although LPS triggers transient reorganization of Golgi morphology in wild-type macrophages, similar structural alterations persist in CCTα-deficient cells. Pro-tumor necrosis factor α and interleukin-6 remain lodged in the secretory compartment of CCTα-deficient macrophages after LPS stimulation. However, the lysosomal-mediated secretion pathways for interleukin-1β secretion and constitutive apolipoprotein E secretion are unaltered. Exogenous lysophosphatidylcholine restores LPS-stimulated secretion from CCTα-deficient cells, and elevated diacylglycerol levels alone do not impede secretion of pro-tumor necrosis factor α or interleukin-6. These results identify CCTα as a key component in membrane biogenesis during LPS-stimulated cytokine secretion from the Golgi complex.

Introduction

Membrane biogenesis is dependent on phospholipid synthesis, specifically phosphatidylcholine (PtdCho), the most abundant phospholipid component that forms the bilayer barrier structure in eukaryotic cells. Doubling of the membrane phospholipid during cell cycle progression is regulated in part by the phosphocholine cytidylyltransferase (CCT; Jackowski, 1994, 1996), a key enzyme in PtdCho synthesis (Jackowski and Fagone, 2005). Also, biochemical activation of CCT mediates expansion of the ER membrane in B lymphocytes that are activated by lipopolysaccharide (LPS; Fagone et al., 2007). Inactivation of CCT either genetically or pharmacologically in immortalized cells leads to cell death (Lykidis and Jackowski, 2001; Cui and Houweling, 2002). Genetic deletion in mice of the ubiquitous CCTα isoform is lethal in embryogenesis before the blastocyst stage (Wang et al., 2005). These results and the fact that the CCTα isoform is expressed at levels 10- to 30-fold higher than the β isoforms (Jackowski et al., 2004) suggest that CCTα activity is required to support membrane biogenesis. However, tissue-specific deletion of CCTα in mice does not severely restrict the proliferation or development of mouse macrophages (Zhang et al., 2000), hepatocytes (Jacobs et al., 2004), or lung epithelial cells (Tian et al., 2007), which demonstrates that either expression of the CCTβ2 or CCTβ3 isoforms in mice (Karim et al., 2003) is sufficient for development and differentiation, and/or that circulating lipoproteins supply substantial amounts of PtdCho to primary cells and tissues (Gunter et al., 2007). However, the loss of CCTα expression does alter the differentiated function of these cell types. CCTα-deficient macrophages have increased susceptibility to cell death by apoptosis after challenge with cholesterol (Zhang et al., 2000; Devries-Seimon et al., 2005). Deficient hepatocytes are larger, and secretion of high-density lipoprotein and very low-density lipoprotein from the liver is impaired (Jacobs et al., 2004). Conditional deletion...
of CCTα in lung epithelia results in insufficient synthesis and secretion of dipalmitoyl-PtdCho, the major surfactant phospholipid (Tian et al., 2007). These data obtained by inactivation of CCTα-deficient macrophages were labeled for 6 h with 1 μCi/ml [3H]choline, and incorporation into PtdCho was normalized to cellular protein. (B) Relative transcript levels for CCTα, CCTβ2, CCTβ3, or PEMT were determined by real-time qRT-PCR in wild-type (n = 3) or CCTα-deficient (n = 3) macrophages. (C) CCT enzyme activity in wild-type (○; n = 4) or CCTα-deficient (●; n = 4) macrophage lysates was determined in vitro as a function of protein concentration. (D) Protein expression levels of CCTα (42 kD) and CCTβ3 (39 kD) were determined in wild-type and CCTα-deficient macrophage lysates (50 μg) by immunoblotting with isoform-specific antibodies. Immunoblots are representative of macrophages from two mice of each genotype. Cells (HEK293T) transfected with expression plasmids encoding CCTα or CCTβ3 were lysed, and 5 μg of protein was loaded as a positive control. Multiple immunopositive bands in a lane correspond to different phosphorylation states of the CCT protein (Jackowski, 1994; lykidis et al., 1998). Error bars indicate mean ± SE. *, P < 0.05; **, P < 0.01.

In proliferating cells, the ER is a major site of PtdCho synthesis (Henneberry et al., 2002; Fagone et al., 2007; Sriburi et al., 2007), and PtdCho is then mobilized to the Golgi compartment (Altan-Bonnet et al., 2004, 2006). DAG production is initiated by the breakdown of PtdCho by phospholipase D followed by a PtdOH phosphatase (P’tse), and this process promotes secretion through the Golgi apparatus (Kearns et al., 1997; Litvak et al., 2005; Lev, 2006). Maintaining the supply of DAG is critical to secretion in both yeast and mammalian cells, and the inhibition of PtdCho synthesis increases DAG (Kearns et al., 1997; Litvak et al., 2005). DAG is reincorporated into PtdCho by choline phosphotransferase (CPT) activity located at the Golgi apparatus (Henneberry et al., 2002; Gleichrist et al., 2006), but the limiting substrate for this reaction is cytidine diphosphocholine (CDP-Chol) produced by CCT (Jackowski and Fagone, 2005). In this study, we find that genetic inactivation of the major CCT isoform in murine macrophages resulted in reduced PtdCho synthesis and increased DAG levels, coupled with impaired secretion of cytokines from the Golgi apparatus of stimulated cells. The experiments illustrate that PtdCho biosynthesis is critical for vesicular traffic from the Golgi apparatus and that aberrantly high DAG levels do not impede vesicular transport.

Results

Mice with CCTα-null macrophages were derived as described previously (Zhang et al., 2000). In brief, mice carrying a floxed Pcyt1a gene were crossed with mice carrying a Cre recombinase driven by the macrophage-specific LysM promoter. Two flox sites flanked a 12.5-kb fragment of the Pcyt1a gene containing exons 5–9 (Karim et al., 2003) and the intervening introns. Deletion of the region between the flox sites resulted in expression of a
truncated CCTα protein that lacked the catalytic and regulatory domains located at the carboxy terminus. Mice that were homozygous for the floxed gene and heterozygous for the LysMCre gene were mated, and the pups were genotyped and used to obtain either CCTα knockout or littermate wild-type control macrophages. Peritoneal macrophages were elicited, and the CCTα-deficient macrophage population was confirmed to have reduced rates of de novo PtdCho synthesis as measured by [3H]choline incorporation into the lipid fraction of adherent cells (Fig. 1 A). The deletion in this macrophage model was not completely penetrant (Zhang et al., 2000). Analysis of the CCTα transcript level in CCTα-deficient macrophages by quantitative RT-PCR (qRT-PCR) revealed that the transcript level was significantly reduced to 30% of the wild-type level (Fig. 1 B). The CCTα-null cells were ≥95% macrophages as determined microscopically after staining with macrophage-specific anti–mouse CD11b. Cells were also stained for expression of CCTα protein, and 20–30% of the macrophages isolated from individual knockout mice (n = 4) remained positive for expression, which confirms that the action of the Cre recombinase was not 100% penetrant. Thus, the residual PtdCho synthesis measured by metabolic radiolabeling in the CCTα-null population was due not only to expression of the alternate isoform, CCTβ3 (Fig. 1 B), but also to the presence of cells that did not delete the CCTα gene. The qRT-PCR revealed that the CCTβ2 isoform was not expressed in macrophages, and that CCTβ3 expression was increased in the knockouts. CCTβ3 is equivalent to CCTα in its primary biochemical function, and expression of its activity can compensate for loss of CCTα to support bulk membrane synthesis in cultured cells (Karim et al., 2003). It was previously found that CCTβ2 expression increased in the CCTα-deficient cells (Zhang et al., 2000), but at the time, CCTβ3 had not been discovered, and both isoforms share the common epitope that was signaled in the previous immunoblots (Karim et al., 2003). Total CCT activity was significantly lower in cell lysates prepared from CCTα-null macrophages compared with wild-type populations (Fig. 1 C), and these data agreed with the previously published study (Zhang et al., 2000). The decrease in CCTα and the increase in CCTβ3 protein assessed by immunoblotting with isoform-specific antibodies reflected the changes in mRNA levels (Fig. 1 D). Macrophages were isolated from two knockout and two wild-type mice and pooled according to genotype, then immunoblots were prepared from six replicates of each genotype. The level of CCTα protein in the knockout population was 0.36 ± 0.08 compared with the wild-type population (level set at 1.00). The reduction in CCTα transcripts correlated with the protein level determined by immunoblots and with the number of macrophages devoid of CCTα expression as determined by microscopy of immunostained cell populations. These data confirm the loss of CCTα expression in the majority of cells and also provide the first example of CCTβ3 regulation as an adaptive response to PtdCho deficiency.

Several macrophage functions were assessed to determine if they were compromised by reduced PtdCho synthesis. Phagocytosis was evaluated by quantifying the uptake of fluorescein-labeled Escherichia coli, and the values were normalized to cell number by counterstaining and measuring DAPI fluorescence. The uptake of the bacteria was linear up to 2 h and was the same in both wild-type and CCTα-deficient cells (Fig. 2 A). Random migration and chemotaxis of the macrophages in response to the attractant formyl-methionylleucylphenylalanine (fMLP) was determined in modified Boyden chambers. The numbers of cells that crossed a membrane into an adjacent chamber containing medium alone or medium plus 10 nM fMLP were determined by staining with Calcein AM, quantification of the fluorescent signal, and comparison with a calibration curve of increasing cell number. (C) Cytokine and PGE 2 secretion from wild-type (white; n = 8) and CCTα-deficient (black; n = 8) macrophages 18 h after LPS stimulation was measured with the luminex assay (BioRad Laboratories) or by individual ELISA assays and normalized to protein content in each sample. The mean values for cytokines in the medium from WT cells were: TNFα, 319.3 pg/μg cell protein; IL-6, 266.2 pg/μg cell protein; IL-1β, 15.3 pg/μg cell protein; and PGE 2 , 5.46 pg/μg cell protein. Error bars indicate mean ± SE. *, P < 0.05; **, P < 0.01.
from the wild-type cells by 24 h. TNF processing is concurrent with its movement from the Golgi to the cell surface and mediated by a metalloprotease called TNF-converting enzyme (TACE; Gearing et al., 1994, 1995). A TACE inhibitor, \( \text{N-(d,l-}[2\text{-hydroxyaminocarbonyl} \text{-methyl}]\text{-4-methylpentanoyl}]\text{-l-3-(2'naphthyl)-alanyl-l-alanine}, 2\text{-aminoethylamide} \) (TAPI; Crowe et al., 1995), was added to wild-type macrophages after LPS addition to inhibit TNF processing and confirm the identification of the pro-TNF on the blots (Fig. 3). TAPI abolished TNF release to the medium, but TAPI did not inhibit IL-6 secretion from wild-type cells (Fig. S2). The data demonstrated that LPS signaling and cytokine synthesis were intact in the CCT-deficient macrophages. Because IL-6 secretion was not compromised, the inhibition of TACE activity by PtdCho deficiency was ruled out as a mechanism for the TNF secretory defect.

Constitutive secretion of apolipoprotein E (ApoE) was not dependent on LPS stimulation and was not impaired in the CCT-deficient cells (Fig. 3D). The synthesis of ApoE is repressed at the transcript level after LPS stimulation (Gafencu et al., 2007), and, thus, the amount released to the medium was reduced in both wild-type and knockout cells at later times after LPS stimulation (Figs. 3D and S3A). Imaging of cellular TNF and ApoE secretion was measured by sampling the culture medium 18 h after addition of the ligand. Secretion of TNF, and interleukin-6 (IL-6) was reduced in the CCT-deficient population, but the secretion of interleukin-1β (IL-1β) was comparable to that in wild-type cells (Fig. 2C). Both TNF and IL-6 are delivered to the medium by vesicle-mediated secretion from the Golgi apparatus. IL-1β is transported by ATP-dependent secretion of specialized lysosomes that bypass the Golgi compartment (Andrei et al., 2004). The release of prostaglandin E₂ (PGE₂) was also normal in the CCTα-null cells (Fig. 2C). Altogether, these data indicated that LPS signaling was intact and that secretion of cytokines from the Golgi apparatus was impaired in the CCTα-deficient macrophages.

A time course revealed reduced release of both TNF and IL-6 from the knockout cells into the medium (Fig. 3A and C). Reduced cytokine secretion was reflected in reduced total production of the cytokines (Fig. S1, available at http://www.jcb.org/cgi/content/full/jcb.200706152/DC1). Immunoblotting of cell lysates at 12 and 24 h after stimulation showed that TNF accumulated in the CCTα-deficient macrophages as both the precursor and the mature form (Fig. 3B). In contrast, the membrane-bound precursor form of TNF (pro-TNFα) was quantitatively processed to the soluble mature form and completely released from the wild-type cells by 24 h. TNF processing is concurrent with its movement from the Golgi to the cell surface and mediated by a metalloprotease called TNF-converting enzyme (TACE; Gearing et al., 1994, 1995). A TACE inhibitor, \( \text{N-(d,l-}[2\text{-hydroxyaminocarbonyl} \text{-methyl}]\text{-4-methylpentanoyl}]\text{-l-3-(2'naphthyl)-alanyl-l-alanine}, 2\text{-aminoethylamide} \) (TAPI; Crowe et al., 1995), was added to wild-type macrophages after LPS addition to inhibit TNF processing and confirm the identification of the pro-TNF on the blots (Fig. 3). TAPI abolished TNF release to the medium, but TAPI did not inhibit IL-6 secretion from wild-type cells (Fig. S2). The data demonstrated that LPS signaling and cytokine synthesis were intact in the CCTα-deficient macrophages. Because IL-6 secretion was not compromised, the inhibition of TACE activity by PtdCho deficiency was ruled out as a mechanism for the TNF secretion defect. Constitutive secretion of apolipoprotein E (ApoE) was not dependent on LPS stimulation and was not impaired in the CCTα-null cells (Fig. 3D). The synthesis of ApoE is repressed at the transcript level after LPS stimulation (Gafencu et al., 2007), and, thus, the amount released to the medium was reduced in both wild-type and knockout cells at later times after LPS stimulation (Figs. 3D and S3A). Imaging of cellular TNF and ApoE

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**Figure 3. Secretion and processing of TNFα.** (A) TNFα in the culture medium from wild-type (○; \( n = 5 \)) or CCTα-deficient (●; \( n = 10 \)) macrophages was measured at indicated times after LPS stimulation. The anti-TNFα antibody used in the EUSA recognized both the pro-TNFα and mature TNFα. (B) Synthesis and processing of TNFα was evaluated by immunoblotting cell lysates from wild-type (WT) or CCTα-deficient (KO) macrophages. Cells were treated with LPS for either 12 or 24 h, or with an inhibitor of TACE, TAPI, which was added to selected cultures to confirm the identification of the unprocessed pro-TNFα. Purified recombinant mouse TNFα (Leinco Technologies, Inc.) was a positive control (lane 1), and anti-GAPDH mAb (Ambion) was used to confirm equal protein loading (50 μg) in each lane. The data are representative of three independent experiments. (C) IL-6 in the culture medium from the same macrophages used in A. (D) Apo E secretion was evaluated by immunoblotting the culture medium from wild-type or CCTα-deficient macrophages incubated for 18 h with and without LPS. Equal volumes of the culture medium were loaded onto the gel, and the immunoblot densities were quantified. The data are the mean ± SE from two independent determinations using macrophages from a total of four mice of each genotype.
in the same cells indicated that these secretory products were located in different compartments (Fig. S3 B), which supports the view that ApoE exited from the Golgi compartment via different vesicles. New PtdCho biosynthesis was not required for secretion of ApoE but was necessary for the release of TNFα and IL-6 after LPS stimulation.

The process of cytokine synthesis and secretion in the macrophages was investigated using immunocytochemistry. Wild-type and knockout cells were treated with LPS for 6 and 18 h before fixation and then stained with anti-TNFα or anti-IL-6 antibodies (Fig. 4, A and B). The images confirmed that the CCTα-deficient cells were able to synthesize TNFα and IL-6. Intracellular TNFα and IL-6 were detected at 6 h, whereas wild-type cells were negative for these cytokines by 18 h after LPS stimulation. These results correlated with cellular cytokine synthesis and secretion to the medium. However, both cytokines were still retained within the CCTα-deficient cells up to 48 h after LPS stimulation.

The results described thus far were obtained using thioglycollate-elicited macrophages that were isolated by peritoneal lavage and cultured in vitro. The physiological relevance of the phenotype was tested in vivo to ensure that the findings were not artifacts of manipulation of the cells or the in vitro system. The numbers of circulating monocytes, peritoneal macrophages, and bone marrow macrophages in the knockout animals were the same or slightly greater than those from the wild-type mice (Fig. S4, available at http://www.jcb.org/cgi/content/full/jcb.200706152/DC1). We subjected both wild-type and knockout mice to infection with an intranasal dose (10^7 colony-forming units) of luminescent Streptococcus pneumoniae strain T4X and followed the course of the infection in five to six individual littermate mice of each genotype (Fig. 5 A). Wild-type mice developed mild pneumonia and between 10 and 50% succumbed in three independent experiments. In contrast, mice with CCTα-deficient macrophages developed more serious diffuse pneumonia, with higher bacterial counts, and the disease quickly progressed to sepsis (Fig. 5 A), with a mean 70% mortality rate (Fig. 5 B). In lungs of knockout animals, there were extensive areas of interstitial pneumonia with consolidation, necrosis, and marked fibrinopurulent pleuritis (Fig. 5 C, top). The inflammatory infiltrate in the pulmonary areas was composed of viable and degenerate neutrophils, macrophages, and perivascular cuffs of lymphocytes and plasma cells. In lungs of wild-type mice, there were patchy mild interstitial inflammatory infiltrates composed primarily of...
neutrophils with fewer macrophages and perivascular and peribronchiolar infiltrates of lymphocytes and plasma cells. A Gram’s stain to detect bacteria in the lung tissue revealed abundant cocci in the knockout animals (Fig. 5 C, bottom), particularly embedded in the layers of fibrin along the pleural surface (arrows), whereas Gram-positive cocci were rarely observed in the wild-type lungs. The numbers of macrophages in the knockout model were equivalent to the numbers in wild-type animals (Fig. S4, A–C), and the knockout macrophages at the site of infection accumulated and retained TNFα, in contrast to the wild-type macrophages (Fig. 5 D). The macrophages that infiltrated the lungs 48 h after infection were identified in cryosections with an anti-MAC-1 antibody and counterstained with an anti-TNFα antibody. The wild-type macrophages were devoid of the cytokine by this time, but the majority of knockout cells still retained a strong signal, which indicates that the cytokine had not yet been released into the extracellular space (Fig. 5 D). These data supported the hypothesis that CCTα deficiency caused the macrophages to accumulate TNFα and impaired cytokine secretion in vivo.

The reorganization of the Golgi apparatus after LPS stimulation was compared between wild-type and CCTα-deficient macrophages using an antibody that recognizes the ER–Golgi intermediate compartment 53-kD protein (ERGIC-53), a protein specifically localized in the ER and cis-Golgi compartment (Litvak et al., 2005). The images indicated only limited rearrangement of the marker protein after LPS stimulation and a similar distribution in both wild-type and knockout cells (Fig. 6). Next, the trans-Golgi compartment was imaged with an antibody that recognizes the K58 marker protein (Bloom and Brashear, 1989). K58 was located proximal to the cell nucleus and surrounded 30–50% of the nucleus before LPS stimulation. In wild-type cells, the K58 protein coalesced into a focused structure at a distinct perinuclear site by 6 h and then returned to the more diffuse distribution characteristic of unstimulated cells at 18 h after LPS activation (Fig. 6). In the CCTα-deficient...
ganelle by a PLD, which is activated by secretory stimuli. PtdOH is in turn rapidly converted to DAG by a PtdOH P’tse. PtdCho is also the metabolic precursor to sphingomyelin (SM), a lipid that is synthesized in the Golgi compartment and then transported to the cell surface. Measurement of the amount before and 18 h after LPS stimulation revealed that the PtdCho was significantly reduced in the CCT/H9251-deficient cells after LPS, in contrast to the wild-type cells, where the PtdCho level was maintained after stimulation (Fig. 7A). In contrast, the amount of total protein per cell was the same in wild-type and knockout cells both before and after LPS (Fig. S4F). Phosphatidylethanolamine (PtdEtn), the second most abundant phospholipid in intracellular membranes, did not change significantly in the knockout cells after LPS stimulation (Fig. S5A, available at http://www.jcb.org/cgi/content/full/jcb.200706152/DC1). The DAG levels in CCTα-deficient cells increased significantly but remained the same in wild-type after LPS stimulation (Fig. 7B). SM was reduced to the same apparent level with no statistical difference between the wild-type and knockout cells (Fig. 7C). These data

Figure 6. Distributions of Golgi markers in LPS-treated macrophages. Wild-type (WT) and knockout (KO) peritoneal macrophages were treated with LPS at 10 ng/ml for 0, 6, and 18 h. After immediate fixation in 4% paraformaldehyde, cells were permeabilized and stained with antibodies against CCTα (green), the trans-Golgi marker K58 (green), or the cis-Golgi marker ERGIC-53 (green). Cell nuclei were counterstained with DAPI (blue). Each micrograph shown in the figure is representative of macrophages from at least four independent mice. Bar, 15 μm.
We confirmed the data using real-time qRT-PCR on total RNA extracted from wild-type macrophages before and 18 h after LPS stimulation, and found a statistically significant increase in the expression of these genes (Fig. S5 E). This information indicated the isoforms of each enzyme that worked together with the CCT/H9251 to control phospholipid metabolism during LPS-stimulated cytokine secretion. Taken together, the data support the existence of a cycle of PtdCho degradation and resynthesis that accompanies cytokine secretory vesicle formation and budding from the Golgi complex (Fig. 8).

A pharmacological approach was used to test several aspects of this hypothesis. Et-18-OCH₃ (edelfosine), an inhibitor of CCT (Boggs et al., 1995a), dramatically reduced TNF/H9251 secretion in cells of both genotypes (Fig. 9 A). Incubation with lysophosphatidylcholine (lysoPC), which is rapidly converted to PtdCho (Baburina and Jackowski, 1999), bypassed the CCT/H9251 genetic defect (Esko et al., 1982; Boggs et al., 1995b; Baburina and Jackowski, 1998) and partially restored TNFα secretion from CCTα-deficient cells (Fig. 9 A). The LysoPC pathway would replenish PtdCho but would not remove DAG. In fact, measurement of the DAG level in lysoPC-treated knockout cells revealed that the DAG level increased (Fig. S5 F). This result was probably caused by inhibition of CCTβ3 by lysoPC, which has been shown to interfere with the membrane association and activation of the enzyme (Boggs et al., 1995a; Attard et al., 2000). Treatment of cells with either 1-butanol to inhibit phospholipase D activity, or propranolol to inhibit PtdOH P'tse activity, also inhibited TNFα secretion (Fig. 9 B). These results support the involvement of the phospholipase D pathway in secretion. However, fumonisin B1, an inhibitor of ceramide and SM synthesis, had no effect.

Both CDP–Cho and DAG are substrates for the CPT that makes PtdCho. Limitation of the CDP–Cho supply by genetic inactivation of the CCTα accounted for the reduced PtdCho and for the metabolic accumulation of DAG after LPS. Affymetrix microarray data, available at the National Center for Biotechnology Information website (Gene Expression Omnibus; http://www.ncbi.nlm.nih.gov/; Shell et al., 2005), revealed that expression of a select group of lipid metabolic genes increased 18 h after LPS stimulation of wild-type macrophages, including those encoding the PLD1, the CCTα, and the choline/ethanolamine phosphotransferase (C/EPT; Table S1, available at http://www.jcb.org/cgi/content/full/jcb.200706152/DC1; Shell et al., 2005). We confirmed the data using real-time qRT-PCR on total RNA extracted from wild-type macrophages before and 18 h after LPS stimulation, and found a statistically significant increase in the expression of these genes (Fig. S5 E). This information indicated the isoforms of each enzyme that worked together with the CCTα to control phospholipid metabolism during LPS-stimulated cytokine secretion. Taken together, the data support the existence of a cycle of PtdCho degradation and resynthesis that accompanies cytokine secretory vesicle formation and budding from the Golgi complex (Fig. 8).

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DAG levels were increased to test whether impaired secretion would be the result. At 3 h after LPS stimulation, wild-type cells were incubated with increasing concentrations of a phospholipase C isolated from *Bacillus cereus*. The phospholipase C in the medium converted surface membrane PtdCho to DAG. We reasoned that at least a portion of the DAG would relocate intracellularly to the Golgi apparatus, similar to the supplemental lysoPC in the medium that resulted in rescue of the phenotype (Fig. 9). At 6 and 18 h after addition of the enzyme, the amount of TNFα in the medium and the amount of DAG in the cells was quantified (Fig. 10). Despite substantial increases in DAG that correlated with the amount of phospholipase C added (Fig. 10 A), TNFα secretion was not impaired (Fig. 10 B). These data argued that increased DAG was not the critical biochemical factor leading to the knockout phenotype. Rather, depletion of the PtdCho supply impeded vesicular transport, resulting in accumulation of cytokines in the Golgi apparatus.

**Discussion**

Our interpretation of the inhibition of induced cytokine secretion from the knockout cells is that the PtdCho requirement for cytokine trafficking arises from the inability to supply PtdCho

These results confirmed that the cellular defect was caused by reduced PtdCho synthesis and did not extend to an imbalance in SM synthesis (Fig. 9 B). SM synthesis and TNFα and IL-6 release to the medium were measured in the same experiment in the presence and absence of fumonisin B1 in wild-type cells. Although SM synthesis was severely reduced, the amount of either cytokine in the medium was the same with or without inhibitor treatment (Fig. S5, C and D). Nor did 2-butanol have an effect, which was a treatment control for the specificity of the phospholipase D inhibition by 1-butanol (Fig. 9 B). Taken together, these data support the idea that de novo PtdCho synthesis was required to maintain cytokine secretion from the Golgi apparatus in LPS-stimulated macrophages.

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**Discussion**

Our interpretation of the inhibition of induced cytokine secretion from the knockout cells is that the PtdCho requirement for cytokine trafficking arises from the inability to supply PtdCho
to the Golgi apparatus. The PtdCho degradation mediated by phospholipase D that accompanies cytokine secretion in LPS-stimulated cells depletes PtdCho in the absence of CCTα-dependent de novo synthesis, resulting in a lipid imbalance exemplified by an increase in DAG. The activities and Golgi localizations of both CCTα and the C/EPT in wild-type cells enable rapid resupply of PtdCho. The residual CCT activity in the knockout cells, caused by CCTβ3 expression, is not sufficient or not located in the appropriate cellular compartment (Fig. S4 D) to provide the PtdCho necessary to support cytokine secretion. Two datasets support this interpretation: the lower rate of PtdCho synthesis in the knockout cells compared with the wild type (Figs. 1 A and S5 A) and the lower amount of PtdCho per cell in the CCTα knockouts after LPS stimulation (Fig. 7 A). In contrast, ApoE secretion is not dependent on de novo PtdCho synthesis (Figs. 3 D and S3). The data indicate that ApoE protein exits the Golgi compartment by a pathway distinct from the TNFα and IL-6 cytokines. ApoE is concentrated in a subcellular of Golgi-derived vesicles distinct from those vesicles that contain secretory albumin (Gusarova et al., 2007). ApoE secretion is linked to cellular sphingolipid synthesis (Lucic et al., 2007), and neither process is impaired in the CCTα knockout macrophages (Figs. 7 C and S5 B).

CCTα participates in a cycle of PtdCho degradation and resynthesis illustrated in Fig. 8. The ER is a major cellular site for membrane biogenesis (Fagone et al., 2007; Sriburi et al., 2007), and new phospholipid is distributed throughout the cell via membrane lateral diffusion, vesicular transport, or protein carriers to other membrane-delimited organelles (Trotter and Voelker, 1994). The replenishment of PtdCho and the removal of DAG in the Golgi can occur via these transport processes. However, our experiments suggest that CCTα expression (Fig. S5 E and Table S1) and association with the trans-Golgi compartment (Fig. 6) are required to provide new CDP–Cho that is used by the Golgi-localized C/EPT (Fig. S5 E and Table S1) to resupply PtdCho. The heightened activity of phospholipase D (Bi et al., 1997; Corda et al., 2002; Freyberg et al., 2003; Jenkins and Frohman, 2005) during LPS-stimulated cytokine secretion degrades the Golgi-associated PtdCho and yields DAG. A threshold level of DAG is required for secretion in yeast (Kearns et al., 1997) and HeLa cells (Litvak et al., 2005; Lev, 2006). The DAG recruits specific proteins to the Golgi apparatus that function in vesicular transport (Patton-Vogt et al., 1997; Roth, 1999; Baron and Malhotra, 2002; Corda et al., 2002) and induces curvature in the phospholipid bilayer that is postulated to promote membrane fusion and fission events (Shemesh et al., 2003). However, our results indicate that aberrantly high DAG does not impede vesicular transport from the Golgi complex (Fig. 10, A and B; and Fig. S5 F).

The association of CCTα with the trans-Golgi compartment during TNFα secretion (Fig. 6) is consistent with its ability to translocate to membranes enriched in DAG and become activated. CCTα reversibly binds to biological membranes via its amphipathic α-helical domain, and DAG is among the most potent membrane components that promote CCTα binding in vitro and in vivo (Kent, 1997; Cornell and Northwood, 2000; Jackowski and Fagone, 2005; Cornell and Taneva, 2006). CCTα membrane association is not specific for DAG but rather correlates with the membrane curvature elastic stress that is induced by accumulation of conically shaped membrane lipids (Attard et al., 2000). The binding of CCTα to membranes increases CDP–Cho production at the site of DAG formation to promote PtdCho synthesis by the Golgi-resident CPT or C/EPT enzymes (Henneberry et al., 2001). The association of the C/EPT isoform with the secretary compartment is suggested by its increased expression after LPS stimulation (Fig. S5 E and Table S1). Overexpression studies placed the CPT isoform at the Golgi apparatus and the C/EPT isoform with the ER (Henneberry et al., 2002). However, a later proteomics analysis of the endogenous components of the secretory pathway identified C/EPT rather than CPT in the Golgi cisterneae (Gilchrist et al., 2006). In addition, increased expression of CPT is associated with ER biogenesis in B cells (Fagone et al., 2007) and fibroblasts (Sriburi et al., 2004). Thus, our data (Fig. S5 E) suggest that the C/EPT is the likely participant in PtdCho synthesis associated with secretion from the Golgi apparatus.

Our findings in macrophages extend the relationship between PtdCho synthesis and vesicular trafficking uncovered in lower organisms. Recent studies in *Drosophila melanogaster* eye development indicate a role for CCT in regulation of the endocytic pathway that results in an altered subcellular localization of the EGF and Notch receptors (Weber et al., 2003). Similarly, the defects in ovarian development in *D. melanogaster* Cct1 mutants are explained by defects in the secretion of signaling molecules through the Golgi apparatus (Gupta and Schüpbach, 2003). The relationship between Golgi function and PtdCho synthesis is most clearly revealed in the yeast system, where the arrest of Golgi-mediated secretion by mutations in sec14, a PtdCho/ phosphatidylinositol transfer protein, is overcome by mutations in the CDP–Cho pathway (Clevés et al., 1991; Hujbregts et al., 2000). Yeast synthesize the majority of their PtdCho through an alternate pathway; thus, inactivation of the CDP–Cho pathway is not lethal in that system. The importance of DAG production to mammalian Golgi function is also highlighted by the recent work with the Nir2 protein. Depletion of the Nir2 protein limits the production of DAG in the Golgi, inhibits protein transport from the trans-Golgi to the plasma membrane, and affects the structural integrity of the Golgi apparatus (Litvak et al., 2005). There is a distinct difference in CCTα localization, and perhaps its metabolic role, in transformed cultured cells compared with primary cells. Although CCTα is located in association with the Golgi complex and outside the nucleus in primary macrophages (Fig. 6), it is found predominantly within the nucleus of transformed macrophages and HeLa cells (Lykidis et al., 1999). In all of the primary cells examined thus far, including macrophages (this paper), lung secretory type II cells (Ridsdale and Post, 2004), and B lymphocytes (Fagone et al., 2007), the CCTα is located outside the nucleus and often in association with organellar membranes.

**Materials and methods**

CCTα-deficient macrophages

Homozygous *Pcyt1α<sup>−/−</sup>* mice (Zhang et al., 2000) were obtained from I. Tabas (Columbia University, New York, NY) and bred with *lysMCre* mice (obtained from I. Förster, Technical University of Munich, Munich, Germany; courtesy of P. Murray, St. Jude Children’s Research Hospital, Memphis, TN) as described previously (Clausen et al., 1999; Zhang et al., 2000) to obtain...
knockout and littermate wild-type macrophages. The two flox sites flanked a 12.5-kb fragment of the gene containing exons 5–9 (renumbered; Karim et al., 2003), the intervening introns, and parts of introns 4 and 9. Deletion of the region between the flox sites resulted in loss of the catalytic domain of the CCTα protein. The resulting transcript encoded a truncated protein that lacked the catalytic domain and the regulatory domains located at the carboxy terminus. The Neo cassette and a third flox site that were originally inserted into the gene and used for embryonic stem cell selection of recombinants was removed by mating with an Ella Cre mouse (The Jackson Laboratory) to avoid hypomorphic effects on CCTα expression in the undelated macrophages from littermates (Zhang et al., 2000). Genotyping of tails and macrophages was performed as described previously (Zhang et al., 2000). Macrophages were elicited by injection of 3 ml thioglycolate (Becton Dickinson) followed by peritoneal lavage with PBS containing 5 U/ml heparin. After hypertonc lysis of the erythrocytes using 10 mM HKOSO4, 150 mM NaCl, and 0.1 mM Na2EDTA, pH 7.6, macrophages were cul-
tured in L cell-conditioned medium (LCM) as described previously (Tang et al., 1999). LPS from E. coli (Sigma-Aldrich) was dissolved in PBS, and optimized concentrations were 10 or 100 ng/ml and lot number–dependent.

**CCT activity and expression**

CCT activities in macrophage cell lysates were determined essentially as described previously (Luche et al., 1993). Protein concentration was det-
termined according to the Bradford method using y-globulin as a stan-
dard (Bradford, 1976). Total RNA was isolated from cells using TRizol reagent (Invitrogen) as described previously (Wang et al., 2003). Real-
time PCR was used to determine the relative expression levels of the CCA

ta, CCTβ2, CCTβ3, and phosphatidylethanolamine methyltransferase

(PEMT) isosform transcripts encoded by the Pcyt1α, Pcyt1β, and Pemt genes as described previously (Jankowski et al., 2004). Immunoblotting with CCT isoform-specific antibodies has been described previously (Lykidis et al., 2001).

**Lipid analysis**

2.5 x 105 macrophages were plated in 60-mm dishes and incubated over-
night in LCM. For radiolabeling experiments, adherent cells were incu-
bated in fresh LCM containing either 1 μCi/ml [methyl-3H]choline chloride or 10 μCi/ml [14C]palmitic acid (American Radiolabeled Chemicals, Inc.) ± 10 ng/ml LPS for the time indicated in the figure legends. Lipids were ex-
tracted as described previously (Bligh and Dyer, 1959) and fractionated on preadsorbent H silica thin-layer plates (Analtech) developed with

chloroform/methanol/acetet acid (98.2:1) or chloroform/methanol/

ammonium hydroxide (60:35:8). Areas corresponding to PtdCho, PtdEtn, and SM standards were counted. For measurements of lipid quantity, ex-
tracted lipid mixtures were fractionated on silica gel rods using the same solvent systems. The amount of each identified lipid was quantified by

flame ionization using an Iatroscan (Iatron Instruments). DAG was quanti-
fied in lipids extracted from cells as described previously (Press et al., 1987) and normalized to cell number. Protein was quantified by the Bradford method (Bradford, 1976).

**Phagocytosis assays**

Macrophages were plated at a density of 105 pm well in 96-well black plates (Costar) and cultured overnight in LCM. A bacterial suspension con-
taining ~1 x 108 fluorescent bacteria from the Vybrant phagocytosis assay kit (Invitrogen) was added and incubated for 2 or 4 h at 37°C. Quantifi-
cation was performed according to the manufacturer’s instructions on a Fusion microplate (Corning) and cultured overnight in LCM. A bacterial suspension con-
tracted as described previously (Bligh and Dyer, 1959) and fractionated on preadsorbent H silica thin-layer plates (Analtech) developed with

chloroform/methanol/acetet acid (98.2:1) or chloroform/methanol/

ammonium hydroxide (60:35:8). Areas corresponding to PtdCho, PtdEtn, and SM standards were counted. For measurements of lipid quantity, ex-
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fied in lipids extracted from cells as described previously (Press et al., 1987) and normalized to cell number. Protein was quantified by the Bradford method (Bradford, 1976).

**Chemotaxis assays**

Cell migration assays were performed in 96-well MultiScreen-MIC plates (Millipore) with 5-mm-pore-sized polycarbonate membrane. 10 nM IMLP

(EMD) was or was not added to the lower wells. Cells that migrated after

1 h were stained with Calcein AM quantified using a Fusion Universal Microplate Analyzer plate reader.

**Cytokine and prostaglandin assays**

Macrophages were cultured with LCM containing 10–100 ng/ml LPS, and, as indicated, 200 μl aliquots of culture medium were removed and assayed for TNFα, IL-6, IL-1β, or PGE2; content by ELISA (Quantikine R&D Systems; Cytokine Chemical) according to the manufacturer’s instructions. The anti-
body used in the TNFα ELISA recognized both the precursor pre-TNFα and the mature TNFα isoforms. Where indicated, cells were harvested and

lysates were prepared as described previously (Mebel et al., 2005). In brief,
imaged for bioluminescence using the Isis Imaging System (Caliper Life Sciences) to document differences in the course of disease at 24, 48, 72, and 96 h after infection as described previously (Orhiuela et al., 2004). Lung was isolated just before death for histopathology analysis. Frozen sections were used for histochromy with TNFα and MAC-1 (Abcam), and paraffin sections were used for hematoxylin and eosin and Gram’s staining. All procedures were approved by the Animal Care and Use Committee of St. Jude Children’s Research Hospital.

Statistical analysis
All data were reported as the mean ± SE as calculated using GraphPad Prism 4 software [GraphPad Software]. The unpaired student t test was used to determine comparative significance. Results were considered significant at a p-value of <0.05 (*) or <0.01 (**).

Online supplemental material
Fig. S1 shows the total production and cellular cytokine content of wild-type and Cctα-null cells after LPS stimulation. Fig. S2 shows the inhibition by iAIP on total production of TNFα but not IL-6. Fig. S3 illustrates the secretion of ApoE from wild-type and Cctα-null cells. Fig. S4 characterizes the macrophage populations in mice, quantifies the protein content of macrophages, and illustrates the cellular distributions of Cctα, CTβ6, and TNFα in wild-type and knockout macrophages. Fig. S5 illustrates the PtdEtn, PtdCho, and SM levels in cells treated with LPS ± fumonisin B1; the lipid gene expression levels quantified by qRT-PCR after LPS stimulation of macrophages, and the DAG levels in knockout macrophages. Table S1 lists the lipid genes up-regulated in LPS-treated macrophages as determined by Affymetrix microarray analysis. Table S2 lists the qRT-PCR primers and probes used to obtain the data in Fig. S5 E. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.200706152/DC1.

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