

Characterization of an evolutionarily distinct bacterial ceramide kinase from Caulobacter crescentus

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A common feature among nearly all gram-negative bacteria is the requirement for lipopolysaccharide (LPS) in the outer leaflet of the outer membrane. LPS provides structural integrity to the bacterial membrane, which aids bacteria in maintaining their shape and acts as a barrier from environmental stress and harmful substances such as detergents and antibiotics. Recent work has demonstrated that Caulobacter crescentus can survive without LPS due to the presence of the anionic sphingolipid ceramide-phosphoglycerate (CPG). Based on genetic evidence, we predicted that protein CpgB functions as a ceramide kinase and performs the first step in generating the phosphoglycerate head group. Here, we characterized the kinase activity of recombinantly expressed CpgB and demonstrated that it can phosphorylate ceramide to form ceramide 1-phosphate. The pH optimum for CpgB was 7.5, and the enzyme required Mg²⁺ as a cofactor. Mn²⁺, but no other divalent cations, could substitute for Mg²⁺. Under these conditions, the enzyme exhibited typical Michaelis-Menten kinetics with respect to NBD C6ceramide ($K_{m,app}$ = 19.2 ± 5.5 µM; $V_{max,app}$ = 2590 ± 230 pmol/min/mg enzyme) and ATP ($K_{m,app} = 0.29 \pm 0.07$ mM; $V_{max,app} = 10,100 \pm 996 \text{ pmol/min/mg enzyme}$). Phylogenetic analysis of CpgB revealed that CpgB belongs to a new class of ceramide kinases, which is distinct from its eukaryotic counterpart; furthermore, the pharmacological inhibitor of human ceramide kinase (NVP-231) had no effect on CpgB. The characterization of a new bacterial ceramide kinase opens avenues for understanding the structure and function of the various microbial phosphorylated sphingolipids.

Gram-negative bacteria have a three-layered cell envelope composed of the inner membrane, a thin layer of peptidoglycan cell wall, and an outer membrane. A key component of the outer membrane is lipopolysaccharide (LPS) (1). LPS is an essential molecule in nearly all gram-negative species due to its roles in barrier formation and membrane integrity (2). While the general structure of LPS is well conserved, there is considerable variation between and within species (3). LPS can be divided into three structural domains: (1) lipid A, a

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membrane-anchored multiacylated oligosaccharide, (2) the core oligosaccharide, often containing 3-deoxy-d-manno-oct-2-ulosonic acid (Kdo), which is generally conserved within a species, and (3) the polysaccharide O-antigen, which is highly variable, even among strains of the same species. In many organisms, like *Escherichia coli*, the lipid A portion of LPS is negatively charged due to the presence of phosphate groups on the glucosamine disaccharide (3). These phosphates are the binding sites for cationic antimicrobial peptides like polymyxins (4, 5). While LPS is generally considered to be essential, LPS-null mutants of several gram-negative organisms have been isolated including Acinetobacter baumannii (6), Moraxella catarrhalis (7), Neisseria meningitidis (8), and Caulobacter crescentus (9). The ability of C. crescentus to survive in the absence of LPS is, in part, due to the presence of the anionic sphingolipid ceramide-phosphoglycerate (CPG), as sphingolipid synthesis becomes essential in the LPS-null mutant (9). In contrast to E. coli, the mature lipid A molecule in C. crescentus is not phosphorylated; instead, the phosphate groups are hypothesized to be removed by the phosphatase CtpA (9) and replaced with galactopyranuronic acid (10). Whereas polymyxin antibiotics target the phosphorylated lipid A in E. coli, antibiotic sensitivity assays demonstrated that cationic antimicrobial peptides kill C. crescentus by interacting with the anionic CPG lipids (9). Synthesis of the CPG head group is sequentially catalyzed by the three proteins CpgABC (CCNA_01217-01219) (9) (Fig. 1A). Deletion of cpgB (ccna_01218) results in the loss of ceramide 1-phosphate (C1P), which is consistent with its annotation as a putative lipid kinase (9) (Fig. 1B).

C1P has important physiological roles in eukaryotes including mast cell activation, phagocytosis, cellular proliferation, and survival (reviewed in (11)). Human ceramide kinase (hCERK) uses ceramide and ATP as substrates to produce C1P (12). The ceramide kinase (CERK) enzyme is part of a larger family of lipid kinases including sphingosine kinase and diacylglycerol (DAG) kinase. A bacterial dihydrosphingosine kinase has recently been identified in Porphyromonas gingivalis (13); however, to our knowledge, this is the first described bacterial CERK enzyme. In this study, we used purified C. crescentus CpgB to characterize its CERK

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Figure 1. Identification of CpgB as a putative ceramide kinase. *A*, previous genetic analysis of the *cpgABC* genes led to a proposed mechanism for ceramide-phosphoglycerate (CPG) synthesis (9). *B*, extracted-ion chromatograms show the presence or absence of ceramide and C1P. Total lipids were extracted from the indicated strains and analyzed by normal phase LC/ESI-MS in the negative ion mode. The signal for the C1P peak was magnified 10-fold since this lipid is only a minor component of the *Caulobacter crescentus* lipidome. This figure is a representative chromatogram (n = 2). C1P, ceramide 1-phosphate; CPG, ceramide-phosphoglycerate; LC/ESI-MS, liquid chromatography electrospray ionization mass spectrometry.

activity. Phylogenetic analysis comparing various lipid kinases suggests that bacterial CERK enzymes form a distinct clade from their eukaryotic counterparts.

Results

CCNA_01218 is a bacterial CERK

Most gram-negative bacteria require LPS in the outer membrane for survival. A recently isolated mutant of *C. crescentus* is capable of surviving without LPS, largely due to the presence of the anionic sphingolipid CPG (9). Genetic analysis identified three genes (ccna 01217-01219) that were required for synthesizing the phosphoglycerate head group. CCNA 01218 (CpgB) is annotated as a lipid kinase-related protein, and deletion of cpgB resulted in a loss of C1P (Fig. 1B) (9), consistent with cpgB encoding a bacterial CERK. To determine the enzymatic activity of CpgB, we purified the His-tagged recombinant protein from E. coli (Fig. 2) and performed kinase assays. CpgB could readily phosphorylate C16-ceramide (Fig. 3A) as well as a fluorescent NBD-C6ceramide (Fig. 3B). The identity of the phosphorylated NBDceramide product was confirmed by mass spectrometry (Fig. 3C). Since CpgB has a conserved LCB5 DAG kinase domain, we tested whether CpgB could phosphorylate DAG to produce phosphatidic acid (PA) and found comparable activity (Fig. 3, A and B). Although CpgB can produce PA in vitro, the C. crescentus lipidome contains only $\sim 1\%$ PA (14) and deletion of cpgB had no effect on PA levels (Fig. 3D). Therefore, we conclude that ceramide is the preferred in vivo substrate for CpgB. Owing to their ease of use, NBD-labeled lipid substrates have been used to characterize the activities of ceramide glycosyltransferases (15), PA phosphatase (16), hCERK (17), and bacterial dihydrosphingosine kinase (13); similarly, the remainder of the kinase assays described below use the NBDceramide substrate.

Influence on pH and divalent cations on CpgB activity

To characterize the requirements for CpgB activity, we measured C1P production over a pH range from 4.5 to 10; the optimal activity was observed at pH 7.5 (Fig. 4*A*). By contrast, hCERK has optimal activity at pH 6.5 (12, 17). Since hCERK activity increases strongly in the presence of magnesium or calcium (12), we tested CpgB's dependence on divalent cations. In the absence of any cations, we did not observe production of C1P (Fig. 4*B*). Both magnesium and manganese strongly increased CpgB activity, with smaller effects observed in the presence of zinc or cobalt (Fig. 4*C*). In contrast to hCERK, calcium did not stimulate CpgB activity (Fig. 4*C*).

Determination of CpgB kinetic parameters

Using the NBD-ceramide substrate, we measured C1P production over a 2-h period to identify the linear range of activity for subsequent determinations of enzyme kinetic parameters (Fig. 5A); unless otherwise noted, all remaining kinase assays were performed for 30 min in the presence of Mg^{2+} at pH 7.4. The enzyme exhibited typical Michaelis-Menten kinetics with respect to NBD-C6-ceramide ($K_{m,app}$ = 19.2 ± 5.5 μ M; V_{max,app} = 2590 ± 230 pmol/min/mg enzyme) and ATP ($K_{m,app} = 0.29 \pm 0.07 \text{ mM}$; $V_{max,app} = 10100 \pm 996 \text{ pmol}/$ min/mg enzyme) (Fig. 5, B and C). We are reporting apparent K_m and V_{max} values since CpgB has two substrates and performs a Bi-Bi reaction; under these conditions, the concentration of each substrate affects the apparent kinetic parameters of the other. Additionally, the kinetic parameters determined using the NBD-ceramide substrate are likely to differ from the true physiological constants and cannot be used





Figure 2. Purification of CpgB. His-tagged CpgB was expressed and purified from *Escherichia coli*. An SDS-PAGE gel of recombinant CpgB was stained with Coomassie blue to assess protein purity. CPG, ceramide-phosphoglycerate.

to make any definitive conclusions about intracellular substrate concentrations.

Bacterial and eukaryotic CERKs are phylogenetically distinct enzymes

Given the observed enzymatic differences between hCERK and CpgB, we considered whether these two enzymes are evolutionarily related. Sequence alignment shows limited agreement (12.5% identity and 22.5% similarity); four of the five sphingosine kinase conserved domains show some homology between the eukaryotic and bacterial kinases (Fig. 6A) (12). The two kinases also share a conserved GGDG motif, which is involved in ATP binding (18). However, the eukaryotic CERKs have an absolutely conserved CxxxCxxC motif that is required for enzyme activity (19) but is absent from CpgB.

To further assess the functional similarity between the CERKs, we treated CpgB with the hCERK inhibitor NVP-231 (20). NVP-231 is a competitive inhibitor of ceramide binding and inhibits 90% of hCERK activity at 100 nM (20). By contrast, 100 nM NVP-231 had no significant effect on CpgB activity (Fig. 4*B*). When the concentration was increased to 300 nM, we observed only a modest 20 to 25% inhibition (Fig. 6*B*), suggesting that CpgB may have a distinct active site from hCERK.

CpgB is a bacterial ceramide kinase

Several enzyme families are capable of phosphorylating sphingolipids and DAG. To visualize the similarity of CpgB to these enzymes, we performed a maximum-likelihood phylogenetic analysis and included representative proteins from the following families: hCERK, yeast DAG kinase Dgk1 (21), bacterial dihydrosphingosine kinase dhSphK1 (13), and bacterial phosphatidylglycerol kinase YegS (22). Each of these enzymes formed a distinct clade despite having overlapping activities (Fig. 6*C*). We did find several cyanobacterial enzymes with homology to hCERK as well as some green algae with homologs of YegS; CpgB homologs were only found in bacterial species. Further analysis of the CpgB-encoding organisms revealed that nearly all genera with the *cpgB* gene either produce or encode the genes required for sphingolipid synthesis (23) (Fig. 6*D*).

Discussion

Bacterial sphingolipids have a wide range of head groups including sugars (24–26), phosphoglycerol (27), phosphoglycerate (9), and phosphoethanolamine (28). These modifications likely determine the physiological functions of the respective sphingolipids. For example, phosphoglycerol dihydroceramide produced by *P. gingivalis* promotes osteoclastogenesis through its interactions with nonmuscle myosin II-A (27). In the case of *C. crescentus*, production of the anionic CPG enables survival in the absence of LPS (9). Genetic analyses using single-gene deletion mutants led to the identification of three enzymes required for CPG synthesis and suggested that the first step is catalyzed by CpgB, a putative CERK. In this report, we used recombinant CpgB expressed and purified from *E. coli* to confirm its CERK activity and analyze its enzymatic properties.

CpgB differs from the human CERK with regards to divalent cation specificity, susceptibility to the inhibitor NVP-231, and kinetic parameters. For comparison, the $K_{m,app}$'s for CpgB are 19 μ M and 0.29 mM for ceramide and ATP, respectively, whereas the reported K_m 's for hCERK are 187 μ M and 32 μ M (12).

From a structural perspective, hCERK activity is observed in cellular membrane fractions despite not having any predicted transmembrane domains; one explanation is that the N-terminal pleckstrin homology domain interacts with membrane phosphatidylinositol molecules (12). By contrast, CpgB purifies as a soluble protein without the use of detergents and is predicted to be a cytoplasmic protein (29). Consistent with these biochemical findings, phylogenetic analysis suggests that the bacterial CERK forms a unique subfamily of lipid kinases, distinct from eukaryotic CERK. Broad conservation of CpgB across many classes of bacteria suggests that phosphorylation may be a common sphingolipid modification.

Until recently, the genes responsible for specific ceramide modifications were unknown. As a result, various studies broadly determined the importance of total sphingolipid production by knocking out the *spt* gene and assessing phenotypes related to survival or virulence (28, 30, 31). With the discovery of enzymes required for sphingolipid glycosylation,

CpgB is a bacterial ceramide kinase



Figure 3. Cpg has ceramide kinase activity. *A*, recombinant CpgB was used to phosphorylate C16-ceramide or DAG. ³²P incorporation was monitored by TLC and phosphorimaging. This TLC result is a representative image (n = 3). *B*, the substrate specificity of CpgB was analyzed using fluorescent NBD lipid substrates as indicated. This TLC is a representative result (n = 3). *C*, production of the phosphorylated NBD-ceramide product was confirmed by negative ion ESI/MS analysis. *D*, negative ion ESI/MS analysis of lipid extracts from WT and $\Delta cpgB$ strains shows no difference in phosphatidic acid (PA) levels. CPG, ceramide-phosphoglycerate; DAG, diacylglycerol; ESI/MS, electrospray ionization mass spectrometry.

phosphorylation, and other modifications (9, 13, 24, 26), we can now dissect the roles of specific head group modifications. The characterization of a new bacterial CERK opens avenues for understanding the structure and function of the various microbial phosphorylated sphingolipids.

Experimental procedures

Cloning His-tagged CpgB

The *cpgB* gene was amplified from *C. crescentus* genomic DNA using primers EK1462 (tatattcatATGCTTCGTCGTG CACGCCATCC) and EK1464 (tactgaattcTCATCCGACC AGGAACCGCAAGGC) and ligated into the NdeI/EcoRI site of plasmid pET28a to generate an N-terminal His-tagged fusion. The resulting plasmid was verified by Sanger sequencing and transformed into *E. coli* strain BL21(DE3) for expression and purification.

Purification of CpgB

A 1 l culture of *E. coli* BL21(DE3) cells carrying the pET28acpgB plasmid was grown in LB broth with 30 µg/ml kanamycin at 37 °C with shaking to an A_{600} of 0.6. IPTG was added to a final concentration of 0.5 mM, followed by induction at 16 °C for 18 h. The cells were collected by centrifugation at 10,000g and resuspended in 12.5 ml of buffer containing 0.5 M sucrose and 10 mM Tris, pH 7.5. Lysozyme was added to a final concentration of 144 µg/ml, and the suspension was stirred on ice for 2 min. A total of 12.5 ml of 1.5 mM EDTA was added with stirring for an additional 7 min to induce plasmolysis. The cells were collected by centrifugation at 10,000g for 10 min, and the pellet was resuspended in lysis buffer (20 mM Tris, pH 7.5, 0.5 M NaCl, and 10 mM imidazole) prior to lysis via 2 to 3 passages through a French press (20,000 psi). The lysate was centrifuged at 8000g for 10 min to remove unbroken cells. His-CpgB was purified using an ÄKTA start FPLC system and a





Figure 4. Characterization of CpgB pH and divalent cation requirements. A, CpgB kinase activity was determined over a range of pH's using the following buffers: pH 4.5 to 6 (100 mM citrate), pH 6.5 to 7.5 (100 mM Mops), pH 8 to 9 (100 mM Tris–HCl), and pH 10 (100 mM borate). Activity was quantified using the NBD-ceramide substrate (n = 3, error bars are the SD). *B*, the CpgB kinase assay was performed in the presence or absence of 15 mM Mg²⁺ using NBD-ceramide. Product formation was analyzed by TLC. *C*, the activity of CpgB was determined in the presence of 15 mM of the indicated divalent cations. Activities were normalized to magnesium (n = 3; error bars are the SD). CPG, ceramide-phosphoglycerate.

1 ml HisTrap HP column (Cytiva). After loading, the column was washed with lysis buffer prior to elution *via* a linear gradient to 1 M imidazole. Protein elution was monitored by A_{280} , and fractions were collected and analyzed by SDS-PAGE followed by Coomassie blue staining. Fractions containing the purified CpgB were combined and dialyzed into 10 mM Tris, pH 7.2, 0.1 M NaCl, 2 mM EDTA, 1 mM DTT over 48 h at 4 °C. The dialyzed protein was concentrated using an Amicon Ultra centrifugal filter (10 kDa molecular weight cutoff) (Millipore Sigma). The protein concentration was determined using the BCA Protein Assay Kit (Pierce).

CpgB kinase assay using C16-ceramide

CpgB kinase activity was measured for 30 min at 30 °C as described previously for *E. coli* DAG kinase (32). The reaction mixture contained 50 mM imidazole–HCl, pH 6.6, 50 mM octyl- β -D-glucopyranoside, 50 mM NaCl, 12.5 mM MgCl₂, 1 mM EGTA, 10 mM β -mercaptoethanol, 1 mM cardiolipin, 0.1 mM ATP (1000 cpm/pmol), and 0.8 mM ceramide or DAG in a total volume of 20 µl. The radioactive products (PA or C1P) are chloroform soluble and were separated from the remaining radioactive substrate by a nonacidic chloroform/ methanol/MgCl₂ (1 M) phase separation. The chloroform soluble products were separated by TLC using a chloroform/



Figure 5. CpgB enzyme kinetics. The kinetic parameters of CpgB were measured using the C6-NBD ceramide substrate. *A*, CpgB activity was measured as a function of time (n = 3, error bars are SD). *B* and *C*, Michaelis–Menten kinetic parameters were determined for CpgB (n = 2, error bars are SD). *B*, to determine the K_{m,app} for ceramide, ATP concentration was held constant (1 mM) while NBD-ceramide concentration varied. *C*, the K_{m,app} for ATP was determined by holding the NBD-ceramide constant at 160 μ M while varying the ATP concentration. K_{m,app} values were 19.2 \pm 5.5 μ M and 0.29 \pm 0.07 mM for NBD-ceramide and ATP, respectively. V_{max,app} values were 2590 \pm 230 pmol/min/mg enzyme and 10,100 \pm 996 pmol/min/mg enzyme for NBD-ceramide and ATP, respectively. CPG, ceramide phosphoglycerate.

CpgB is a bacterial ceramide kinase



Figure 6. Bacterial CERK is a unique class of lipid kinases. *A*, sequence alignment of CpgB and hCERK shows limited homology. The hCERK sphingosine kinase domains are indicated by the *red boxes*. The conserved GGDG motif is indicated with *asterisks*. hCERK has a conserved CxxxCxxC which is absent in CpgB. *B*, CpgB activity was measured in the presence of the hCERK inhibitor NVP-231 using the NBD-ceramide substrate. Activities were normalized to the control sample (n = 3; error bars are SD, ANOVA F (4, 10) = 10.22, p < 0.0015; * post hoc comparisons using Tukey test, p < 0.05). *C*, phylogenetic analysis of various lipid kinases was performed using the maximum-likelihood method. The branch-tip color indicates the lipid kinase family, and the line colors designate proteins of bacterial or eukaryotic origin. *D*, the phylogenetic ree of the bacterial CpgB homologs is color coded to indicate which genera have members with either experimental evidence (*pink*) or genetic evidence (*green*) of sphingolipid production. Genetic evidence indicates that the genus has members with all three required enzymes for sphingolipid production: Spt, bCerS, and CerR. CPG, ceramide-phosphoglycerate; hCERK, human ceramide kinase.

methanol/water (65:25:4, v/v) solvent system and visualized by phosphorimaging.

NBD-CERK assay

The NBD-CERK assays were carried out largely as previously described for hCERK (12, 17). Briefly, the reaction was carried out in a buffer containing 20 mM Hepes (pH 7.4), 10 mM KCl, 15 mM MgCl₂, 10% glycerol, 1 mM DTT, 1 mM ATP, 0.2 mg/ml fatty acid-free bovine serum albumin, and 10 μ M C6-NBD ceramide (added from a 10 mM ethanol stock) (Thermo Fisher Scientific). The reaction was started by adding 0.025 μ g/ μ l of the CpgB enzyme. Tubes were incubated in the dark at 30 °C for the indicated times. After the incubation, 1 μ l of the reaction mixture was spotted onto silica gel 60 TLC plates. The spots were resolved in a solvent

system containing butanol/acetic acid/water (3:1:1, v/v). The dried TLC plates were visualized using the GFP filter set on a Bio-Rad ChemiDoc. To test the specificity of CpgB, we performed the reaction under identical conditions using 1-NBD-decanoyl-2-decanoyl-sn-Gly (NBD-DAG) (Cayman Chemical) as the substrate. Inhibition of CpgB activity was performed by adding the indicated concentrations of NVP-231 (Cayman Chemical) to the reaction prior to addition of the enzyme.

Lipidomic profiling and confirmation of ceramide-phosphate production by LC/MS/MS

Lipids were extracted from bacterial cells or the NBDceramide CpgB reaction using the method of Bligh and Dyer with minor modifications (33). The lipid extracts were analyzed by normal phase LC/MS/MS in the negative ion mode as previously described (34, 35).

Kinetic analysis of CpgB

To determine the kinetic constants for CpgB, activity assays were performed for 30 min as described above while varying substrate concentrations. To determine the $K_{m,app}$ for ceramide, ATP concentration was held constant (1 mM) while NBD-ceramide concentration ranged from 0.625 to 160 μ M. The $K_{m,app}$ for ATP was determined by holding the NBD-ceramide constant at 160 μ M while varying the ATP concentration from 0.031 to 1 mM. Product formation was measured from the fluorescent images using ImageJ (36) and quantified using a standard curve of NBD-ceramide spotted onto the TLC plates. The enzyme activity was fit to the Michaelis–Menten equation using OriginPro (OriginLab).

Assessing the pH optimum and the requirement for divalent cations

To test the effect of pH on CpgB activity, a standard reaction mix was made containing 10 mM KCl, 15 mM MgCl₂, 10% glycerol, 1 mM DTT, 1 mM ATP, 0.2 mg/ml fatty acid-free bovine serum albumin, and 10 μ M C6-NBD ceramide. The pH was controlled by adding the following buffers: pH 4.5 to 6 (100 mM citrate), pH 6.5 to 7.5 (100 mM Mops), pH 8 to 9 (100 mM Tris–HCl), and pH 10 (100 mM borate). The reactions were started with the addition of 0.025 μ g/ μ l of CpgB and allowed to run for 30 min. Phosphorylated product was quantified as above. The efficacy of various divalent cations was tested by replacing the MgCl₂ with 15 mM CaCl₂, ZnCl₂, MnCl₂, CuCl₂, or CoCl₂ and determining CpgB activity at pH 7.4 as described above.

Phylogenetic analysis of lipid kinase enzymes

Using CCNA 01218 (CpgB; Accession YP 002516591.3) protein as a query, we performed BLASTP searches to find related proteins in the NCBI database (37). The top hits were all from species closely related to C. crescentus, so we repeated the search excluding Alphaproteobacteria to get a wider range of organisms. Candidate hits were chosen using an E-value cutoff of 1E-20 and we manually curated the list to select the top ~ 60 hits from different genera. A similar method was used to find homologs of hCERK (Accession NP_073603.2), P. gingivalis dihydrosphingosine kinase (Accession AAQ66413), E. coli YegS (Accession NP_416590), and Saccharomyces cerevisiae Dgk1 (Accession QHB11896.1). A total of 397 protein sequences were aligned using MUSCLE aligner (38). Phylogenetic trees were prepared using Randomized Axelerated Maximum Likelihood (RAxML, version 8.2.12) (39) with 100 bootstraps and a maximum-likelihood search. RAxML was run on the CIPRES Portal at the San Diego Supercomputer Center (40). Phylogenetic trees were visualized in R using the packages ggtree (41), ape (42), treeio (43), and ggplot2 (44). To determine which *cpgB*-encoding bacterial genera produce sphingolipids, we performed a literature search as well as used the RIKEN JCM

catalog (https://jcm.brc.riken.jp/en/). For genera with no experimental evidence of sphingolipids, we used BLASTP to determine whether these genera encode all three key enzymes for sphingolipid synthesis: Spt (Accession A0A0H3C7E9.1), bCerS (Accession A0A0H3C8X0.1), and CerR (Accession A0A0H3C8X7.1).

Data availability

All of the data for this work is contained within the manuscript.

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Abbreviations—The abbreviations used are: C1P, ceramide 1-phosphate; CERK, ceramide kinase; CPG, ceramide-phosphoglycerate; DAG, diacylglycerol; hCERK, human ceramide kinase; LPS, lipopolysaccharide; PA, phosphatidic acid.

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