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Phosphatidylycerine Stimulates Ceramide 1-Phosphate (C1P) Intermembrane Transfer by C1P Transfer Proteins*

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Genetic models for studying localized cell suicide that halt the spread of pathogen infection and immune response activation in plants include Arabidopsis accelerated-cell-death 11 mutant (acd11). In this mutant, sphingolipid homeostasis is disrupted via depletion of ACD11, a lipid transfer protein that is specific for ceramide 1-phosphate (C1P) and phyto-C1P. The C1P binding site in ACD11 and in human ceramide-1-phosphate transfer protein (CPTP) is surrounded by cationic residues. Here, we investigated the functional regulation of ACD11 and CPTP by anionic phosphoglycerides and found that 1-palmitoyl-2-oleoyl-phosphatic acid or 1-palmitoyl-2-oleoyl-phosphatidylglycerol (≤15 mol %) in C1P source vesicles depressed C1P intermembrane transfer. By contrast, replacement with 1-palmitoyl-2-oleoyl-phosphatidylserine stimulated C1P transfer by ACD11 and CPTP. Notably, “soluble” phosphatidylserine (dihexanoyl-phosphatidylserine) failed to stimulate C1P transfer. Also, none of the anionic phosphoglycerides affected transfer action by human glycolipid lipid transfer protein (GLTP), which is glycolipid-specific and has few cationic residues near its glycolipid binding site. These findings provide the first evidence for a potential phosphoglyceride headgroup-specific regulatory interaction site(s) existing on the surface of any GLTP-fold and delineate new differences between GLTP superfamily members that are specific for C1P versus glycolipid.

Sphingolipids (SLs) regulate key physiological processes, including cell mitogenesis, growth, migration, and differentiation as well as stress-induced programmed cell death responses (autophagy and apoptosis) (1–4). Because SL synthesis occurs at distinct sites in cells, SL distribution and localization to various membrane organelles involves vesicle- and non-vesicle-mediated transport processes. When ceramide is initially glycosylated in the Golgi, nonvesicular transfer can occur via proteins characterized by a glycolipid transfer protein fold (GLTP-fold) (5–7). The GLTP-fold consists of multiple α-helices arranged in a two-layer “sandwich” topology to form a single sphingolipid binding site (8–12). A newly discovered family within the GLTP superfamily uses a modified GLTP-fold to selectively transfer ceramide 1-phosphate (C1P) rather than glycosphingolipids between membranes (12–14). Human C1P transfer protein (CPTP; 214 amino acids) is encoded by a three-exon transcript from the single-copy CPTP gene on chromosome 1 (locus 1p36.33). By contrast, human GLTP (209 amino acids) originates from a five-exon transcript from single-copy GLTP on chromosome 12 ( locus 12q24.11) (15). The shared folding topology (but with different SL specificity generated from the limited sequence homologies of CPTP and GLTP) serves as a striking example of evolutionary convergence and emphasizes the structural premium placed on GLTP-fold conservation by eukaryotes.

CPTPs, like GLTPs, occur almost ubiquitously in eukaryotes but play key roles in controlling inflammation and programmed cell death processes (12–14). Arabidopsis thaliana contains a CPTP orthologue known as ACD11 that functions as a lipid transfer protein for C1P and phyto-C1P (13). The ACD11 name originates from the accelerated cell death (acd) phenotype.
observed upon disruption of the *acd11* gene (16). In the *Arabidopsis acd11* mutant, altered sphingolipid homeostasis manifests as moderately increased C1P and highly increased ceramide that help drive a programmed cell death response (13). *Arabidopsis acd* mutants provide genetic models for studying localized cell suicide that can halt the spread of pathogen infection and immune response activation in plants (2, 17, 18).

ACD11, CPTP, and GLTP are considered to be amphitropic proteins because their functionality involves translocation on/off membranes to bind and release the sphingolipid cargo (19–21). Amphitropic proteins often contain so-called lipid binding domains (LBDs) that bind specific phosphoglyceride headgroups within membranes to help target and tether to select cell membrane destinations (22–27). LBDs, such as the C1, C2, PH, PX, and FYVE domains, differ structurally from the GLTP-fold. In the case of ACD11, CPTP, and GLTP, the protein region surrounding the sphingolipid binding site contains differing numbers of tryptophans, tyrosines, lysines, and arginines, residues known to be concentrated in the membrane interaction regions of proteins (28–32). Clusters of positively charged Arg and Lys residues occur near the SL binding site in ACD11 and CPTP, but not in GLTP (12, 13). This raises the issue of whether the cationic residue clusters of ACD11 and CPTP might be topologically organized to engage specific phosphoglyceride headgroups during membrane interaction.

Herein, we report the discovery of stimulation of C1P transfer by ACD11 when phosphatidylcholine (PC) bilayer vesicles contain both phosphatidylserine (PS) and C1P. A PS stimulatory effect also was observed on human CPTP transfer of C1P. By contrast, slowdowns in the C1P transfer rate were observed in the presence of other anionic phosphoglycerides (phosphatic acid, phosphatidylglycerol). Inclusion of PS had no effect on glycolipid transfer by human GLTP. Our findings suggest the presence of a potential PS-specific headgroup interaction site on the surface of C1P-specific ACD11 and CPTP.

**Results**

X-ray crystallography of ACD11 and CPTP previously revealed a sphingolipid headgroup recognition site containing a positively charged Arg/Lys triad that binds C1P in “transfer-viable” fashion (13, 14). Co-crystallization with other phosphate headgroup lipids showed altered non-transfer-viable interactions involving the same binding site. We initially determined whether fluorescence emission of the intrinsic Trp residues in ACD11 serves as an indicator of transfer-viable protein-lipid complex formation. The two Trp residues in ACD11 are located on the protein surface (Fig. 1a). Trp-145 in helix-6 is at a highly conserved location in the membrane interaction region of the GLTP-fold (12, 13) and is 16–19 Å from the bound C1P headgroup. Trp-206, which forms the C terminus, is closer to the bound C1P headgroup (7–9 Å) and is more favorably positioned for detecting environmentally induced fluorescence emission changes associated with C1P binding. Fig. 1b illustrates a typical Trp emission response for ACD11 upon titrating with C1P. The Trp emission $\lambda_{\text{max}}$ undergoes progressive blue-shifting (i.e. 347 to 342 nm), whereas the fluorescence intensity decreases in response to stepwise additions of N-octanoyl C1P dissolved in ethanol (titration increments < 0.001 mol % total ACD11). A saturation response is eventually achieved, as $\lambda_{\text{max}}$ blue-shifts to ~342 nm and the total Trp emission intensity declines by ~20–30%. A similar but stronger response occurs during glycolipid uptake by GLTP and
related orthologs that differ from ACD11 by having a Trp residue within the glycolipid binding site (33–35). Titration of ACD11 with sphingosine 1-phosphate or phosphatidic acid, which contain phosphate headgroups but are not transferred (13), induced no significant change in Trp emission $\lambda_{\text{max}}$ (Fig. 1, c and d). Other sphingolipids, such as ceramide, which is not transferred (13), elicited almost no change (Fig. 1e), whereas sphingomyelin, which is transferred slowly (36), elicited a weak response (Fig. 1f).

Formation of the ACD11-C1P complex indicated by the intrinsic Trp emission changes was verified by electrospray ionization mass spectrometry (ESI-MS). Fig. 2a shows the raw spectra obtained by direct infusion of ACD11-C1P complex solution under nondenaturing conditions. Deconvolution reveals that the positive ions (+8 and +9) correspond to monomeric complex (ACD11 + N-octanoyl C1P; 23,186 Da) plus small amounts of monomeric, C1P-free ACD11 (22,681 Da) (Fig. 2b). Compared with ESI-MS analyses of human GLTP and HET-C2 fungal GLTP binding of monohexosylceramide (33, 34), ACD11 showed stronger complexation based on the energy needed to disrupt the complex. This could reflect involvement of the positively charged Arg/Lys triad in binding the negatively charged phosphate of the C1P headgroup. No complexation could be detected with sphingomyelin, which is transferred slowly by ACD11 (36).

The X-ray structures for ACD11 and human CPTP (13, 14) also show the sphingolipid binding sites surrounded by positively charged residue clusters that are absent in human GLTP (supplemental Fig. 1, a–c). This raised the issue of whether anionic membrane phospholipids influence the function of ACD11 and CPTP more strongly than GLTP because the membrane interaction region of the GLTP-fold encompasses the sphingolipid headgroup recognition site (12). To evaluate this possibility, we tested the effect of the negatively charged phospholipids PG, PS, and PA on sphingolipid departure rates from 1-palmitoyl-2-oleyl-sn-glycero-3-phosphocholine (POPC) vesicles by ACD11, CPTP, and GLTP. A well established approach (Fig. 2c) based on loss of fluorescence resonance energy transfer (FRET) was used to obtain real-time kinetic insights into lipid intermembrane transfer (13, 14, 37–40). In the assay, fluorescent anthrylvinyl (AV)-sphingolipid (energy donor) and 3-perylonyl (Per)-PC (energy acceptor) are both incorporated into POPC vesicles either lacking or containing negatively charged phosphoglyceride. Fig. 2d illustrates the FRET response observed by excitation of AV-sphingolipid (AV-SL) at 370 nm, resulting in minimal AV emission (400 – 450 nm) and strong Per-PC emission (460 – 560 nm). The addition of protein plus excess POPC receiver vesicles (containing no lipid fluorophores) triggers a sudden, time-dependent loss of FRET as AV-SL departs while the nontransferable Per-PC remains in...
the SL source vesicles, consistent with AV-SL intermembrane transfer. With ACD11 or CPTP, the resulting increase in AV-C1P emission (Fig. 2d) enables monitoring of the AV-SL intermembrane transfer rate. If no transfer protein is added, almost no increase in AV emission is observed, confirming very slow spontaneous migration of AV-C1P to POPC receiver vesicles (41–43). If POPC receiver vesicles are omitted and protein amounts are increased by 20–30-fold, then only slight increases in AV-SL emission are observed, showing that AV-C1P binding to ACD11 does not account for FRET loss (e.g. see Ref. 14). With catalytic protein amounts, the "shuttle-like" protein action keeps AV-C1P transferring from the outer surfaces of the SL source vesicles to the receiver vesicles until dynamic equilibrium is reached (~10 min).

The in vitro assay design is intended to reflect certain aspects of the physiological situation. In mammals, C1P is initially produced by ceramide kinase at select sites, such as the cytosolic face of the trans-Golgi, and is then transported to certain intracellular membranes. For this reason, C1P is localized initially only in the donor vesicles rather than in both the donor and acceptor vesicles. Anionic phosphoglycerides also often localize to intracellular membrane cytosolic faces (72), where they can encounter CPTP. Our experimental set-up takes these physiological factors into account.

C1P Transfer by ACD11 and CPTP Is Slowed by PA and PG but Accelerated by PS—Studies of the regulation of GLTP and its homologs by anionic phosphoglycerides previously focused on mammalian GLTP (38, 44, 45), the fungal HET-C2 GLTP (46), and the GLTPH domain of FAPP2 (35). With GLTP, including anionic phosphoglycerides, such as PA, PG, PS, and PI, in sphingolipid source vesicles impedes GalCer intermembrane transfer at low ionic strength due to enhanced membrane association by GLTP (i.e. Ks). However, adding salt to raise the ionic strength to physiologic or higher levels abrogates the enhanced membrane partitioning and restores GLTP transfer to rates similar to those observed when sphingolipid source vesicles lack negatively charged phosphoglycerides. The earlier studies involved GLTP purified from bovine brain (38). Here, ACD11 and CPTP functional regulation by anionic phosphoglycerides at physiologic ionic strength was studied using human GLTP expressed in Escherichia coli as a control (Fig. 3, a–i). AV-C1P kinetic transfer rates decreased nonlinearly in response to increasing 1-palmitoyl-2-oleoyl-sn-glycero-3-phospho-(1′-rac-glycerol) (POPG) concentrations ranging from 2 to 15 mol % in the sphingolipid source vesicles for both ACD11 (Fig. 3, a and d) and CPTP (Fig. 3, b and e). C1P transfer by both transfer proteins became maximally suppressed when POPA reached ~4 mol %, whereas inclusion of only 2 mol % POPG resulted in C1P transfer reduction that remained low at the higher POPG molar fractions. Fig. 3, c and f, shows that increasing POPA and POPG concentrations (from 2 to 15 mol %) produced virtually no slowdown in the AV-GalCer transfer rate by human GLTP at physiologic ionic strength.

The responses of ACD11 and CPTP to 1-palmitoyl-2-oleoyl-sn-glycero-3-phospho-1′-serine (POPS) sharply contrasted to those elicited by POPA and POPG. Increasing the POPS in sphingolipid source vesicles over the same range (2–15 mol %) strongly stimulated C1P transfer rates (Fig. 3, g and h). With ACD11, the transfer rate more than doubled when POPS concentrations reached 10 mol % (Fig. 3j). With CPTP, the rate increases were slightly less at equivalent POPS molar fractions (Fig. 3k). By contrast, with human GLTP, no increase in the AV-GalCer transfer rate was observed over the same POPS concentration range (2–15 mol %) in the sphingolipid source vesicles (Fig. 3i). Neither calcium nor other divalent cations were needed to elicit the POPS stimulatory effect. Notably, however, when the POPC receiver (acceptor) vesicles contained the POPS instead of the C1P source (donor) vesicles, the increase in C1P was not observed for either ACD11 or CPTP (Fig. 4, a and b). To explain our findings, we hypothesized that the membrane interaction regions of ACD11 and CPTP possess a PS-specific headgroup interaction site but with no associated hydrophobic pocket for accommodating the PS acyl chains. Such an arrangement would enable the PS acyl chains to remain embedded in the membrane while the PS headgroup acts as a selective tethering/activation site for ACD11 and CPTP. To test this idea, we assessed ACD11 and CPTP for their ability to transfer AV-PS between vesicles, but none was observed (Fig. 4c). This lack of PS transfer is noteworthy because other lipid transfer proteins exist that can transfer more than one lipid type. Yeast Sec 14 phosphatidylinositol transfer protein contains overlapping binding/transfer sites for either phosphatidylinositol or phosphatidylcholine (47). Also, certain oxysterol-binding protein family members that originally were thought to bind/transfer only sterol derivatives have since been shown to also bind/transfer of PS and phosphatidylinositol 4-phosphate (48–53).

Membrane Partitioning of ACD11.—We initially evaluated using surface plasmon resonance at physiological ionic strength. After adsorbing vesicles of differing lipid composition to the lipophilic sensor chip, protein was introduced into the flow cell. Fig. 5a shows the enhanced partitioning to POPC/C1P (85:5) vesicles containing 10 mol % POPS or POPG by ACD11 compared with either POPC/C1P (95:5) or POPC vesicles. Using FRET, we further assessed ACD11 interaction with membranes of differing phosphoglyceride composition. Fig. 5 (b, c, and d) shows the spectral responses observed when ACD11 was titrated with increasing amounts of POPC/AV-PC (98:2) vesicles. Membrane interaction by ACD11 resulted in a nonlinear decline in the Trp emission intensity with increasing POPC/AV-PC vesicles in saturation-like fashion due to the spectral overlap of the intrinsic Trp emission of ACD11 (energy donor) and the AV excitation signal associated with PC (energy acceptor). Comparison of the interaction curves (Fig. 5d) shows that having POPS (10 mol %) in the PC membrane interface slightly enhanced FRET compared with the diminished FRET elicited by POPG. A similar outcome was observed when AV-C1P replaced the AC-PC in the POPC vesicles (data not shown).

No C1P Transfer Acceleration by “Soluble” PS—To test whether POPS stimulates C1P transfer by ACD11 and CPTP by helping to optimally orient the C1P binding site of CPTP during membrane interaction, we investigated whether soluble PS with short acyl chains (dihexanoyl-PS) stimulates C1P transfer by ACD11 and CPTP. We reasoned that little or no activation
effect of soluble PS on C1P transfer by ACD11 and CPTP. The C1P transfer rates are expressed as pmol/min transferred from SL source to POPC vesicles as a function of different amounts of anionic phosphoglycerides (0–15 mol %) present in the SL source vesicles for ACD11, CPTP, and GLTP (Fig. 6a–c, d–f). Per-PC along with AV-C1P or AV-GalCer is present at 1 mol % in the SL source vesicles at time 0. g–i, POPS effects (0–15 mol %) on ACD11, CPTP, and GLTP (2 μg), respectively. Per-PC (nontransferable) along with AV-C1P or AV-GalCer is present at 1 mol % in the SL source vesicles at time 0. j and k summarize the C1P transfer rate changes of ACD11 and CPTP induced by POPS, POPA, and POPG. The C1P transfer rates are expressed as pmol/min transferred from SL source to POPC vesicles as a function of different amounts of anionic phosphoglycerides (0–15 mol %) present in the SL source vesicles for ACD11 (a) and CPTP (b). Red, POPA; green, POPG; blue, POPS; gray, control. Error bars, S.D.

Discussion

Our investigation provides the first evidence for the existence of a phosphoglyceride regulatory interaction site(s) on the GLTP-fold surface of a GLTP superfamily member. The GLTP-fold is a structural motif that uses an all-α helical, two-layer “sandwich” topology to form a single, distinct sphingolipid binding site (8, 11–14). Our data indicate that PS stimulates the sphingolipid transfer activity of C1P-specific GLTP-folds (i.e. plant ACD11 and human CPTP). This stimulation by PS is a special feature of the C1P-specific GLTP-fold and is not duplicated by the glycolipid-specific GLTP-fold (i.e. human GLTP).
Other negatively charged phosphoglycerides (e.g. PA and PG) exert an opposing effect on ACD11 and CPTP function and slow the C1P transfer rate from sphingolipid source vesicles at physiological ionic strength. These same phosphoglycerides show no inhibitory effect on recombinant, glycolipid-specific human GLTP, consistent with earlier studies of GLTP purified from bovine brain (38). The findings indicate that potential post-translational processing differences are not responsible for the lack of response by GLTP to anionic phosphoglycerides. Previously, the essential role of FAPP2 in glycosphingolipid biosynthesis in the Golgi was shown to involve its C-terminal, glycolipid-specific GLTP homology domain (5–7, 35). However, the regulatory features controlling the membrane interaction of FAPP2 focused only on its N-terminal pleckstrin homology domain that interacts with phosphatidylinositol 4-phosphate without attention to the GLTP domain.

We expected that negatively charged phosphoglycerides embedded in PC membranes would down-regulate ACD11 and CPTP transfer activity by increasing protein affinity for the SL source vesicles because of the high density of positively charged residues in the membrane interaction regions surrounding the C1P recognition centers (13, 14). Indeed, surface plasmon resonance indicates enhanced partitioning of ACD11 to POPC vesicles containing either POPG or POPS compared with POPC or POPC/C1P (95:5) vesicles. However, ACD11 Trp-to-POPG transfer evident at higher POPA concentrations. When evaluating lipid transfer processes, factors pertaining to the membrane also need consideration besides the protein structural features and simple charge-charge effects. The membrane serves as a matrix for both the “substrate” C1P and the “effector” anionic phosphoglycerides. In bilayer vesicles, the C1P located in the bilayer outer leaflet represents the sphingolipid pool that is accessible to the transfer protein. Spontaneous C1P transbilayer migration is expected to be highly restricted due to the unfavorable energetics of moving the phosphate charged headgroup through the nonpolar hydrocarbon matrix. The interactions of GLTP homologs with membranes occur in a nonperturbing and weakly penetrating manner that leaves the sphingolipid pool in the inner leaflet of the vesicle bilayer undisturbed and inaccessible to protein (14, 37, 44, 55). However, a condition known to influence the initial sphingolipid transbilayer distribution and expected to impact the C1P pool size in the bilayer outer leaflet is the lipid composition during formation of high curvature, small unilamellar vesicles (e.g. see Refs. 56 and 57). When the sphingolipid source vesicles are POPC, the observed 35% C1P transfer equilibrium value suggests that 35% of C1P localizes in the vesicle outer leaflet. This value is similar to that of GalCer (39, 56). When anionic phosphoglycerides are added to the mix, their influence also needs consideration. Because PA shares the same small phosphate headgroup as C1P (58), minimal alteration of the C1P transbilayer distribution and the outer leaflet pool size is expected compared with C1P source vesicles lacking PA. By contrast, the PG headgroup volume is significantly larger than that of C1P and only slightly smaller than that of PC (59). Thus, the PG transbilayer distribution, like that of PC, tends to remain mass-distributed with ~65% in the outer leaflet of the donor vesicle.

FIGURE 4. POPS in acceptor vesicles does not stimulate C1P transfer and POPS itself is not transferred. Traces for a and b represent AV-C1P emission intensity measured at 415 nm as a function of time resulting from the loss of AV-SL/Per-PC FRET as AV-SL is transferred to POPC vesicles. The presence of POPS (10 mol %) in POPC receiver (acceptor) vesicles (10-fold excess) does not stimulate AV-C1P transfer by ACD11 (a) or CPTP (b). c, traces represent AV-PS emission intensity measured at 415 nm as a function of time resulting from the loss of AV-PS/Per-PC FRET. The AV-PS transfer to POPC receiver (acceptor) vesicles (10-fold excess) by ACD11, CPTP, and GLTP is virtually nil.

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C1P Transfer Proteins and Phosphatidylserine
The surprising and large stimulation in C1P transfer activity by anionic PS raises the question of what makes PS such a special regulator. This led us to look beyond simple positive charge density in the surface region near the C1P binding sites of ACD11 and CPTP by virtue of C1P localization to the inner leaflet of the vesicles. The sudden and dramatic drop in the AV-C1P rate produced by low POPG concentrations in the donor vesicles seems consistent with this reasoning. These factors that influence C1P pool size and transbilayer distribution are noteworthy because the sphingolipid concentrations in the donor vesicles seems consistent with this reasoning. The phosphoglyceride headgroup and imply that the PS site could remain firmly associated with the C1P source membrane. It also implies that the interaction mode is not “transfer-compatible” for PS. By contrast, C1P interaction with CPTP and ACD11 involves recognition of the phosphate-amide headgroup region via a mechanism that triggers uptake of the nonpolar aliphatic chains into a hydrophobic pocket, thus shielding much of the sphingolipid cargo from the aqueous milieu (13, 14). The lack of PS transfer by ACD11 or CPTP is consistent with the phosphoserine polar headgroup being the interaction focal point while the PS acyl chains remain embedded in the membrane matrix.

Notably, replacement of POPS with soluble PS (dihexanoyl-PS) results in no stimulation of C1P transfer by ACD11 and CPTP. Also, simply including soluble PS in the buffer of the transfer reaction at up to 1000-fold excess compared with protein, rather than as a component of the SL source vesicles, fails to stimulate C1P transfer activity. Thus, stimulation of C1P transfer activity requires that the phosphoserine headgroup remain firmly associated with the C1P source membrane. It also is noteworthy that inclusion of POPS in the receiver membrane vesicles does not enhance the C1P transfer rate by ACD11 or CPTP. Taken together, the findings suggest the presence of a PS-specific site on the ACD11 and CPTP surface that targets the phosphoserine headgroup and imply that the PS site could help optimize orientation of ACD11 or CPTP for C1P uptake during membrane interaction. Thus, we predict that the mem-
brane interaction region surrounding the C1P binding site findings contains a PS headgroup binding site, as depicted in Fig. 7. Future studies will be needed to test this idea by structural approaches involving X-ray diffraction, NMR, or modeling coupled with functional analyses via point mutation. It will also be interesting to learn whether the interaction of ACD11 or CPTP with long-chain PS embedded in C1P-containing membranes facilitates protein conformational changes to enhance C1P uptake.

From the physiological perspective, the discovery of PS-induced enhancement of C1P-specific GLTP homolog action provides insights into how ACD11 and CPTP might be targeted and site-specifically stimulated by certain membranes in plant and animal cells, respectively. PS plays important roles in cells. Exofacial PS exposure in activated blood platelets induces binding and activation of clotting factors, including factors V, VIII, and X and prothrombin (62). Intracellularly, PS occurs in the cytosol-facing surfaces of the plasma membrane, endosomes, and lysosomes, enabling docking and activation by important cytoplasmic signaling and fusogenic proteins with specific PS-binding domains (63, 64). Such proteins include the E3 ubiquitin-protein ligase NEDD4, various protein kinase C isoforms, several phospholipase C and D isoforms, phosphatidylinositol 3,4,5-trisphosphate phosphatase (PTEN), spectrin, and dysferlin (important in muscle repair) as well as certain synaptotagmin isoforms that participate in vesicular trafficking and fusion. ACD11, which regulates accelerated cell death in plants, and human CPTP, a regulator of pro-inflammatory eicosanoid production, can now be added to this growing list of proteins. It is noteworthy that CPTP intracellular enrichment occurs not only at the trans-Golgi but also on the cytoplasmic surfaces of endosomes and the plasma membrane, sites where PS also is enriched (14, 73).

Conclusions

C1P-specific lipid transfer proteins are known regulators of inflammation and programmed cell death, but mechanisms by which these proteins can be regulated have remained unknown.
until now. In the present study, significant stimulation of two C1P-specific lipid transfer proteins, at physiological ionic strength and in the absence of calcium, has been discovered when POPS, but not POPA or POPG, is embedded in POPC vesicles along with C1P. Notably, soluble PS that does not remain firmly embedded in the bilayer matrix produces no stimulatory effect on C1P transfer. By contrast, glycolipid-specific human GLTP activity is unaffected by all three anionic phosphoglycerides. The existence of a specific PS headgroup interaction site on the ACD11 surface is suggested where the long acyl chains of PS remain embedded in the bilayer during membrane interaction (Fig. 7). In this way, POPS could help orient and tether ACD11 and CPTP on the membrane surface to enhance interaction with C1P with PS-enriched membrane sites serving as ACD11 and CPTP activity “hot spots” in cells.

Experimental Procedures

Materials—POPC, POPS, 6:0 PS, POPA, and POPG were purchased from Avanti Polar Lipids and used without further purification. Lipids containing Per or AV fluorophore (e.g. Per-PC, AV–C1P, AV–GalCer, and AV–PS) were synthesized by lypo-lipid reacylation with ω-labeled 9-(3-pyrenoyl)-nonanoyl or (11E)-12-(9-anthryl)-11-dodecenoyl chains and then purified (65–67).

Recombinant Protein Purification—Cloning, expression, and purification of ACD11, CPTP, and GLTP have been described previously (13, 14, 68–70). Briefly, Arabidopsis acd11 (NCBI NP_181016.1) and human CPTP (GenBankTM JN542538 and NP_077792.2) open reading frames were ligated into pET-30 Xa/LIC (GenBankTM JN542538 and GenBankTM AF209704) that was ligated into pET-30 Xa/LIC expression vector (Novagen) was used to transform BL21 cells for expression of GLTP N-terminally tagged with His$_{6}$-SUMO vector (Invitrogen). Transformation of BL21 (DE3) plasmid cells enabled expression of proteins N-terminally tagged with His$_{6}$-SUMO (13, 14). Human GLTP ORF (GenBankTM AF209704) that was ligated into pET-30 Xa/LIC expression vector (Novagen) was used to transform BL21 cells for expression of GLTP N-terminally tagged with His$_{6}$-S-peptide. Transformed cells were grown in Luria-Bertani medium at 37 °C for 6 h, induced with 0.1 mM isopropyl 1-thio-galactoside, and then incubated for 16–20 h at 15 °C. Affinity purification of soluble lysate was accomplished by nickel-nitrilotriacetic acid affinity chromatography. Cleavage of the N-terminal His$_{6}$-SUMO tag was carried out with SUMO protease, Ulpl, overnight at 4 °C, whereas the His$_{6}$-S-tag was removed from GLTP by factor Xa. Affinity repurification by nickel-nitrilotriacetic acid chromatography followed by FPLC gel filtration chromatography (HiLoad 16/60 Superdex-75 prep grade column; GE Healthcare), equilibrated in 25 mM Tris-HCl (pH 8.0) containing 100 mM NaCl and 1 mM DTT, yielded proteins with native properties. Pooled peak fractions were concentrated by centrifugal concentrators (Vivaspin; 10 kDa cut-off). Protein purity was confirmed by SDS-PAGE (69, 71) before flash-freezing the pure proteins in buffer containing 15% glycerol and storing at −80 °C.

Sphingolipid Binding Specificity Assessment by Trp Emission Changes of ACD11—SL titrations were performed by adding aliquots (1–μl increments; C1P step concentration of 0.08 μM) dissolved in ethanol to ACD11 (1 μM, 2.5 ml) in sodium phosphate (pH 6.6) containing 150 mM NaCl with constant stirring as described previously (33). Measurements were performed using a SPEX FluoroLog-3 spectrofluorimeter (Horiba Scientific). Band passes for excitation and emission were 2 nm. The cuvette was temperature-controlled to ±0.1 °C (NesLab RTE-111, ThermoFisher). Emission spectra (305–500 nm) were corrected by subtraction of buffer and vesicle blanks. Inner filter effects were avoided by using low protein concentration (optical density at 295 nm < 0.1). Excitation at 295 nm also eliminated fluorescence contributions from residues other than Trp.

Mass Spectrometry—The ACD11-C1P complex was prepared using the titration approach described above and concentrated by centrifugation (Vivaspin; 10 kDa cut-off). WT ACD11 and ACD11-C1P complex (10 μM) were analyzed using an Agilent 6210 LC/MS-TOF mass spectrometer in 5 mM NH$_{4}$ acetate plus 5% methanol and infusing directly into the electrospray source, as described earlier for GLTP and HET-C2 (33, 34). Spectra were collected in positive mode over a 500–5000 m/z range using parameters optimized for complex stability (e.g. capillary, 3000 V; fragmentor, 300 V; skimmer, 60 V; octopole radio frequency, 300 V; octopole direct current, 32 V). Raw data were transformed into relative molecular masses using Agilent Time-of-Flight Protein BioConfirm software.

Protein-mediated Sphingolipid Intermembrane Transfer—To monitor sphingolipid intervesicular transfer, we used an established FRET approach (37). Sphingolipid source vesicles composed of POPC and containing 1 mol% AV-lipid (acyl chain ω-labeled with anhydroylinyl fluorophore, (((11E)-12-(9-anthryl)-11-dodecenoyl)) and 1 mol% 1-acyl-2-[9-(3-perylenoyl)-nonanoyl]-3-sn-glycero-3-phosphocholine (Per-PC) were prepared by rapid ethanol injection, as described previously (37, 38). When AV is excited at 370 nm, Per-PC (energy acceptor) emits because of energy transfer, whereas AV emission (energy donor) is minimal. Sonicated POPC vesicles that receive the transferred AV-sphingolipid are added to the stirred cuvette to establish the “no protein” baseline response in buffer consisting of 10 mM potassium phosphate (pH 6.6), 150 mM NaCl, and 0.2% EDTA. The addition of C1P transfer protein results in time-dependent FRET loss between AV and Per and results in an exponential increase in AV emission intensity as the protein transports AV–C1P away from the sphingolipid source vesicles and delivers to the POPC receiver vesicles present at 10-fold excess. The AV emission increase at ~415 nm, relative to baseline fluorescence produced without protein, yields the AV–C1P transfer kinetics. The addition of Tween 20 detergent late in the kinetic time course provides the maximum AV intensity achievable upon “infinite” separation from 3-pyrenoyl fluorophore. Maximum transfer, ΔF, represents the difference in emission intensity in the absence and presence of C1P late in the kinetic time course (>15 min). The initial lipid transfer rate, νo, is obtained by nonlinear regression analyses (Origin version 7.0, OriginLab, Northampton, MA). The S.D. values are calculated at 95% confidence intervals, and $R^2$ values are >0.96.

Protein Partitioning to Phosphoglyceride Membranes—Membrane partitioning by the sphingolipid transfer proteins was assessed by FRET and surface plasmon resonance. FRET from intrinsic Trp of ACD11 and CPTP (0.5 μM) to AV–PC (2 mol%) in POPC vesicles was measured before and after the addition of the AV–PC/POPC vesicles lacking or containing the indicated
amounts of anionic phospholipid. Trp residues were selectively excited at 295 nm, and fluorescence spectral emission (304–525 nm) was monitored at 25 °C. Surface plasmon resonance was performed using a Biacore T200 system. Lipid vesicles of specified composition (0.5 mM), prepared by extrusion using 30 nm pore size membranes (33), were captured to a final surface density of 4000–7000 response units on a Sensor Chip L1 to establish the baseline prior to protein addition. Injections of increasing protein amounts or buffer were performed at 2 μl/min flow rates.

Author Contributions—X. Z. designed, performed, and analyzed all of the experiments and wrote the paper. Y.-G. G. and S. K. M. helped assess protein partitioning to membranes. D. K. S. designed and constructed ACD11 and CPTP expression vectors and performed the X-ray structure determinations essential for the experimental conception. I. A. B. and J. G. M. synthesized and purified the fluorescent lipids used in the experiments. L. M. B. and H. R. B. designed, performed, and analyzed the ESI-MS experiments. L. M. analyzed the structural data and contributed to the preparation of Figure 7. J. M., J. G. M., D. J. P., and R. E. B. analyzed and critically evaluated the results. R. E. B. conceived and designed the experiments, coordinated the study, and wrote the paper. All authors reviewed the results and approved the final version of the manuscript.

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