Multisite phosphorylation of oxysterol-binding protein regulates sterol binding and activation of sphingomyelin synthesis

Asako Goto, Xinwei Liu, Carolyn-Ann Robinson, and Neale D. Ridgway
Department of Pediatrics and Department of Biochemistry and Molecular Biology, Atlantic Research Centre, Dalhousie University, Halifax, NS B3H 4R2, Canada

ABSTRACT The endoplasmic reticulum (ER)-Golgi sterol transfer activity of oxysterol-binding protein (OSBP) regulates sphingomyelin (SM) synthesis, as well as post-Golgi cholesterol efflux pathways. The phosphorylation and ER-Golgi localization of OSBP are correlated, suggesting this modification regulates the directionality and/or specificity of transfer activity. In this paper, we report that phosphorylation on two serine-rich motifs, S381-S391 (site 1) and S192, S195, S200 (site 2), specifically controls OSBP activity at the ER. A phosphomimetic of the SM/cholesterol-sensitive phosphorylation site 1 (OSBP-SSE) had increased in vitro cholesterol and 25-hydroxycholesterol-binding capacity, and cholesterol extraction from liposomes, but reduced transfer activity. Phosphatidylinositol 4-phosphate (PI(4)P) and cholesterol competed for a common binding site on OSBP; however, direct binding of PI(4)P was not affected by site 1 phosphorylation. Individual site 1 and site 2 phosphomutants supported oxysterol activation of SM synthesis in OSBP-deficient CHO cells. However, a double site1/2 mutant (OSBP-S381A/S391S200A) was deficient in this activity and was constitutively colocalized with vesicle-associated membrane protein–associated protein A (VAP-A) in a collapsed ER network. This study identifies phosphorylation regulation of sterol and VAP-A binding by OSBP in the ER, and PI(4)P as an alternate ligand that could be exchanged for sterol in the Golgi apparatus.

INTRODUCTION
Mammalian oxysterol-binding protein (OSBP) and OSBP-related proteins (ORPs) constitute a large eukaryotic gene family characterized by a conserved C-terminal sterol-binding domain (Lehto and Olkkonen, 2003). Full-length OSBP/ORPs also have N-terminal pleckstrin homology (PH) domains that interact with phosphatidylinositol (PI) phosphates (Levine and Munro, 1998, 2002) and a two phenylalanines (FF) in an acidic tract (FFAT) motif that bind vesicle-associated membrane protein–associated protein A (VAP-A) on the cytoplasmic surface of the endoplasmic reticulum (ER; Wyles et al., 2002; Loewen et al., 2003). This domain organization suggests that a primary function of OSBP/ORPs is to transfer cholesterol, ergosterol, or oxysterols between target membranes and/or to transduce sterol-dependent signals at these points of contact (Prinz, 2007; Ngu et al., 2010).

Oxysterol-binding protein homologues (OSH) in Saccharomyces cerevisiae catalyze phosphatidylinositol 4,5-bisphosphate (PI(4,5)P2)-dependent ergosterol transfer in vitro and regulate cholesterol delivery from the ER to the plasma membrane (PM; Raychaudhuri et al., 2006). However, Osh4p and other Osh proteins are also implicated in the regulation of PM phosphatidylinositol 4-phosphate (PI(4)P) levels (Stefan et al., 2011), PM sterol distribution (Georgiev et al., 2011), and polarized endocytosis by mechanisms that evoke sterol-sensing activity (Alfaro et al., 2011). The integration of PI(4)P and ergosterol distribution and function by Osh4p could be due to
competitive, high-affinity binding and transfer of these two ligands (de Saint-Jean et al., 2011). OSBP family members bind cholesterol and oxysterols, and some have been implicated in sterol transfer between organelles (Suchanek et al., 2007; Ngo and Ridgway, 2009; Banerji et al., 2010). ORP9L and OSBP extract and transfer cholesterol between liposomes in vitro (Ngo and Ridgway, 2009). Both proteins localize to the ER and Golgi apparatus and influence the post-Golgi distribution of cholesterol in cultured cells, suggesting they directly or indirectly transfer cholesterol within the early secretory pathway. In the case of OSBP, sterol transfer between the ER and Golgi apparatus is coupled to activation of ceramide transport protein (CERT) and sphingomyelin (SM) synthesis through increased activity of PI4KIIIα, a cholesterol-sensitive PI4-kinase (Waugh et al., 2006; Banerji et al., 2010). On the other hand, OSBP and ORPs have been implicated in regulation of diverse signaling pathways that could be the result of sterol-sensing or sterol-transduction activity (Wang et al., 2005; Johansson et al., 2007; Lessmann et al., 2007; Lehto et al., 2008; Ngo et al., 2010).

The ER is cholesterol-poor compared with later secretory compartments, indicating that transport proteins must utilize a mechanism to deliver ligands against a concentration gradient. In this regard, OSBP is subject to a rapid phosphorylation cycle that is linked to Golgi localization and the cholesterol and SM content of cells (Ridgway et al., 1998b; Storey et al., 1998). Rapid depletion of SM in the PM results in dephosphorylation of OSBP and localization to the Golgi apparatus. Similarly, depletion of cholesterol by prolonged inhibition of synthesis causes OSBP dephosphorylation and Golgi localization. Because SM and cholesterol are essential components of PM rafts, OSBP phosphorylation appears to be dependent on the integrity of these membrane domains. A serine-rich motif situated between the FFAT and sterol-binding domain of OSBP (S381, S384, and S387; Figure 1) is responsive to changes in sterol and SM; S381 is required for subsequent phosphorylation of S384 and S387, indicating these later two sites are phosphorylated by casein kinase 1 (CK1; Mohammadi et al., 2001). A similar cholesterol and SM-sensitive phosphorylation site in CERT negatively regulates PH domain interaction with PI(4)P in the Golgi apparatus (Kumagai et al., 2007). This serine-rich motif consists of a protein kinase D (PKD) phosphorylation site (S132; Fugmann et al., 2007) that regulates phosphorylation of nine adjacent serine and threonine residues by CK1γ2 (Tomishige et al., 2009). Dephosphorylation in the ER by protein phosphatase 2Cε relieves inhibition of the PH domain (Saito et al., 2008). This identified two functional states for CERT; a dephosphorylated form that associates with the Golgi and mediates ceramide transfer from the ER, and an inactive PKD/CK1 phosphorylated form that binds ceramide but does not interact with the Golgi apparatus. The regulation of CERT and OSBP at the level of sterol- and SM-mediated phosphorylation and the sterol-dependent activation of CERT by OSBP (Perry and Ridgway, 2006; Banerji et al., 2010), demonstrate that this functional pair have an important role in regulating membrane homeostasis in the secretory pathway.

To understand how phosphorylation regulates OSBP, we undertook a functional analysis of phosphorylation sites adjacent to the sterol-binding (site 1) and PH (site 2) domains. Results with phosphomimetic mutants indicate that the VAP-A–binding and sterol-binding phase of OSBP at the ER is regulated by phosphorylation at these two sites and is functionally linked to sterol-dependent activation of SM synthesis in the Golgi apparatus. We also demonstrate that the PI(4)P-binding activity of OSBP is competitive with cholesterol but insensitive to phosphorylation of site 1, indicating involvement in a Golgi-specific phase of the transfer cycle.

**RESULTS**

**OSBP site 1 phosphorylation specifically increases sterol-binding activity**

OSBP is phosphorylated between the sterol-binding and FFAT domains (site 1), as well as a PKD site at S242 (Mohammadi et al., 2001; Nhek et al., 2010). Recent phosphoproteome studies have extended site 1 to include phosphorylation at T379, S388, and S391, constituting a large CK1 motif that is activated by phosphorylation of S381 (Figure 1; Dephoure et al., 2008; Zahedi et al., 2008; Gauci et al., 2009). These studies also identified a cluster of three phosphoserine residues (positions 192, 195, and 200, designated site 2) in a potential casein kinase 2 motif next to the PH domain. To determine how phosphorylation at site 1 and site 2 regulates OSBP activity, we mutated serine and threonine residues to mimic fully dephosphorylated or phosphorylated species (summarized in Figure 1). In the case of site 1, OSBP-S381A prevents phosphorylation of downstream serine residues and mimics a dephosphorylated state with increased mobility on SDS–PAGE, while introduction of glutamine residues simulates a phosphorylated state with reduced mobility on SDS–PAGE (Figure 2 and Figure 5 later in the paper; Mohammadi et al., 2001).

To establish whether phosphorylation of site 1 affects OSBP activity in vitro, we expressed and purified OSBP and OSBP-S381A, -S3E, and -S5E from Sf21 cells and assayed them for sterol binding, extraction, and transfer activities (Figure 2). The migration of each protein on SDS–PAGE was consistent with its proposed phosphorylation state; S381A migrated as a dephosphorylated species, while OSBP-S3E and -S5E mimicked a phosphorylated state with reduced mobility (Figure 2A). Wild-type OSBP expressed in Sf21 cells is dephosphorylated at site 1, as indicated by comigration with OSBP-S381A. OSBP

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**FIGURE 1:** OSBP phosphorylation sites and mutations. Site 1, site 2, and PKD phosphorylation sites and mutations reported in this study are indicated in bold. PHD, PH domain; SBD, sterol-binding domain.
displayed saturable binding of [3H]25-hydroxycholesterol (25OH; $K_d$ 28 ± 5 nM; $B_{\text{max}}$ 0.63 ± 0.05) that was similar to OSBP-S381A ($K_d$ 51 ± 12 nM; $B_{\text{max}}$ 0.68 ± 0.11) and OSBP-S3E ($K_d$ 58 ± 14; $B_{\text{max}}$ 0.49 ± 0.15; Figure 2B). The $K_d$ for oxysterol binding to OSBP-S5E (35 ± 9 nM) was similar to wild-type and the other mutants, but binding capacity ($B_{\text{max}}$ 2.0 ± 0.26) was increased threefold. Specific binding of [3H]cholesterol by OSBPs was saturable, but calculated binding constants were inconsistent due to nonlinear Scatchard plots (Figure 2C). However, it is evident that maximal cholesterol binding by OSBP-S5E was also increased greater than twofold compared with OSBP and the other phosphomutants. The rate of sterol dissociation was determined by incubating [3H]25OH-loaded OSBP or OSBP-S5E in buffer containing unlabeled 25OH at 20°C and measuring the appearance of radiolabeled oxysterol in the supernatant (Figure 2D). Under these conditions, [3H]25OH exchanged more slowly from OSBP-S5E compared with wild-type OSBP.

We next determined whether OSBP-S5E had altered cholesterol extraction and transfer activity using liposomal substrates. Extraction was measured by incubating increasing amount of OSBPs with liposomes containing 1% [3H]cholesterol and measuring the appearance of the radiolabel in the supernatant after donor lipidome removal (Figure 2E). Similar to results with detergent dispersions of cholesterol (Figure 2C), OSBP-S5E extracted twofold more [3H]cholesterol from liposomes compared with OSBP and OSBP-S381A. The effect of site 1 phosphorylation on cholesterol transfer was determined used a modified assay in which OSBPs were first incubated with donor liposomes to initiate [3H]cholesterol extraction; this was followed by addition of acceptor liposomes (Figure 2F). The percentage of [3H]cholesterol transferred by all OSBPs to acceptor liposomes containing PI was negligible but was stimulated by the presence of 2 mol% PI(4)P. Although OSBP-S5E extracted twice as much cholesterol as OSBP and OSBP-S381A (Figure 2E), the percentage transferred to acceptor liposomes with PI(4)P was similar, indicating less efficient delivery of bound cholesterol. Transfer by a OSBP PH domain mutant (RR/EE) that does not interact with PI(4)P was reduced by 75%, confirming previous results indicating the PH domain has a stimulatory role in transfer (Ngo and Ridgway, 2009). Like Osh4p (de Saint-Jean et al., 2011), cholesterol transfer by OSBP is stimulated by PI(4)P in acceptor liposomes, but the N-terminal PH domain also has a positive role.

PI(4)P occupies the Osh4p sterol-binding pocket with its 4-phosphate residue associated with a phylogenetically conserved histidine pair, raising the possibility that other OSBP/ORPs bind both sterols and PI(4)P (de Saint-Jean et al., 2011). We therefore used competition and direct-binding assays to determine whether OSBP binds PI(4)P, and whether site 1 mutations affect this binding activity. First, OSBP extraction of [3H]cholesterol was assayed using liposomes that contained increasing amounts of unlabeled PI, PI(4)P, or PI(4,5)P$_2$ (Figure 3A). Inclusion of PI or PI(4,5)P$_2$ in donor liposomes

**FIGURE 2:** Identification of an OSBP site 1 phosphomutant with increased sterol-binding activity. (A) Purified OSBP, OSBP-S381A, OSBP-S3E, and OSBP-S5E (2 μg each) were resolved by SDS–6%PAGE and stained with Coomassie Blue. (B and C) Specific binding of [3H]25OH and [3H]cholesterol by 12 pmol of OSBP ( ), OSBP-S381A ( ▲ ), OSBP-S3E ( ■ ), or OSBP-S5E ( ♦ ) was assayed as described in Materials and Methods. (D) Following binding of 100 nM [3H]25OH, OSBP and OSBP-S5E were isolated on Talon resin and resuspended in binding buffer containing PI and PI(4,5)P$_2$. Binding of [3H]25OH to OSBP was assayed as described in **Materials and Methods**.

First, OSBP extraction of [3H]cholesterol between liposomes by OSBP, OSBP-S5E, OSBP-S381A, or OSBP-RR/EE was determined using a modified assay that involved preextraction of sterols from donor liposomes prior to addition of acceptor liposomes containing 2 mol% PI or PI(4)P. Results in all panels are the mean and SEM of three or more experiments using two to three different protein preparations.
did not inhibit \[^{3}H\]cholesterol extraction compared with liposomes without PIPs. In contrast, inclusion of PI(4)P caused a 75% reduction in \[^{3}H\]cholesterol extraction. We needed to exclude the possibility that this resulted from OSBP sequestration by PI(4)P-containing liposomes, so the distribution of OSBP between the supernatant and pellet fraction of the assay shown in Figure 3A was subjected to SDS–PAGE and protein staining (Figure 3B). It is evident that concentrations of PI(4)P that inhibited \[^{3}H\]cholesterol to OSBP did not increase OSBP association with donor liposomes in the pellet fraction.

To verify the results from competition assays shown in Figure 3A, we measured direct binding and extraction of radiolabeled PI and PI(4)P by OSBP (Figure 3, C and D). OSBP extracted significantly more \[^{32}P\]labeled PI(4)P from liposomes compared with \[^{3}H\]PI (Figure 3C). \[^{32}P\]PI(4)P extraction by OSBP was dose-dependent but became nonlinear after 50 pmol (Figure 3D). Interestingly, OSBP-S5E extracted similar amounts of \[^{32}P\]PI(4)P compared with wild type and OSBP-S381A, indicating that phosphorylation of site 1 specifically stimulates sterol-binding activity. Thus, in addition to binding PI(4)P via the PH domain, OSBP has an additional PI(4)P-binding site that is competitive with cholesterol.

OSBP site 1 phosphorylation does not affect oxysterol-activation of SM synthesis

Because OSBP-S5E has increased sterol-binding activity, and oxysterol and cholesterol binding is known to enhance OSBP association with the Golgi apparatus (Lagace et al., 1999), we tested whether OSBP-S5E and other site 1 phosphomutants have altered membrane-binding activity and Golgi localization (Figure 4). PH domain–dependent interactions of OSBPs were assessed by binding to phosphorylated PIs on nitrocellulose and in liposomes, although the later assay is complicated by the potential for PI(4)P to be bound and extracted by the sterol-binding domain (Figure 3). Wild-type OSBP and the site 1 phosphomutants had similar, preferential binding to PI(4)P that was immobilized on nitrocellulose filters (Figure 4A). Similarly, OSBP binding to liposomes containing 10 mol% PI(4)P or PI(4,5)P\(_2\) was unaffected by site 1 mutations, although this assay indicated a binding preference for PI(4,5)P\(_2\) (Figure 4B). For assessment of interaction of OSBP site 1 mutants with the Golgi apparatus, N-terminal mCherry fusions were transiently expressed in CHO cells, and cellular localization was determined by immunofluorescence microscopy in the presence and absence of 25OH (Figure 4C). OSBP and OSBP-S5E were primarily in a diffuse cytoplasmic/ER compartment in the absence of 25OH but became strongly localized to the perinuclear Golgi compartment in cells exposed to 25OH. OSBP-S381A had a slightly more pronounced perinuclear distribution in the absence of 25OH but translocated to the Golgi apparatus in oxysterol-treated cells. Untagged OSBP mutants showed similar behavior in response to 25OH treatment (unpublished data). Although the site 1 phosphomutants did not have altered ER-Golgi localization, the proximity of site 1 to the FFAT motif suggests that phosphorylation could increase negative charge in this region and influence VAP-A interaction. For assessing this in vitro, increasing amounts of recombinant OSBP site 1 mutants were incubated with GST-VAP-A and interaction was determined by pulldown and SDS–PAGE (Figure 4D). Binding of OSBPs by GST-VAP-A was relatively linear over the chosen concentration range, and there was no significant difference in the amount of OSBP-S381A or -S5E associated with GST-VAP-A compared with wild-type OSBP. These in vitro experiments show that site 1 phosphorylation specifically enhances sterol-binding capacity without affecting the FFAT and PH domain interactions of OSBP.

Sterol-mediated translocation of OSBP to the Golgi apparatus activates CERT-dependent ceramide delivery to the Golgi apparatus for conversion to SM (Perry and Ridgway, 2006). Reconstitution of SM synthesis (measured by \[^{1}H\]serine incorporation) in OSBP-deficient CHO cells by cDNA transfection can be used as a functional readout to assess the influence of OSBP mutations on this pathway. Using this approach, OSBP site 1 mutants were tested for their ability to restore 25OH-activated SM synthesis in CHO cells depleted of endogenous OSBP (Figure 5). Compared with mock-transfected cells, OSBP expression in deficient CHO cells did not affect synthesis under unstimulated conditions but restored a 2.5-fold activation of SM synthesis by 25OH (Figure 5A). Expression of OSBP-S381A or the phosphomimics OSBP-S3E or -S5E also restored SM synthesis to a level that was not significantly different...
OSBP is phosphorylated at site 2 adjacent to the PH domain

Phosphoproteome studies identified additional phosphoserines at positions 192, 195, and 200 that are embedded in a negatively charged region adjacent to the PH domain (Figure 1). For confirmation from wild type. T379 phosphorylation was also evaluated as a potential regulatory site of SM synthesis. OSBP-T379A migrated as a doublet on SDS–PAGE (Figure 5C), indicating that site 1 phosphorylation is intact and phosphorylation of S381 alone triggers phosphorylation of the adjacent four serine residues. Expression of OSBP-T379A or a mutant with all six threonine and serine residues changed to glutamate (OSBP-TS6E) restored SM synthesis (Figure 5A). [3H]serine incorporation into ceramide in mock- and OSBP-transfected CHO cells was similar under control and 25OH-treated conditions (Figure 4B). Collectively, this suggests that phosphorylation at site 1 is sufficient to enhance sterol binding but not subsequent steps in OSBP regulation of SM synthesis at the Golgi apparatus.

OSBP is phosphorylated at site 2 adjacent to the PH domain

Phosphoproteome studies identified additional phosphoserines at positions 192, 195, and 200 that are embedded in a negatively charged region adjacent to the PH domain (Figure 1). For confirmation of
phosphorylation at site 2, OSBP and single or double site 1/site 2 mutants that prevent phosphorylation (S3A and S381A) were transiently transfected into CHO cells, and phosphorylation status was monitored by immunoblotting and phosphopeptide analysis (Figure 6). OSBP-S3A had similar mobility as OSBP on immunoblots and reduced 32P incorporation into the lower band but not the upper band, indicating that site 1 was unaffected (Figure 6A). OSBP-S381A lacked the upper phosphorylated species corresponding to site 1 phosphorylation but had similar incorporation of 32P into the lower band, suggesting that site 2 phosphorylation was unaffected. The double mutant lacked the upper band and had reduced incorporation into the lower species. Tryptic digestion of immunoprecipitated OSBP double mutant lacked the upper band and had reduced incorporation band, suggesting that site 2 phosphorylation was unaffected. The phosphorylation but had similar incorporation of

A site 1/site 2 double mutant is nonfunctional and aggregates in the ER

Restoration of 25OH-activated SM synthesis in OSBP-depleted CHO cells was used to assess the role of site 2 in OSBP activity (Figure 7). Expression of the site 2 phosphorylation mutants S3A and S3D restored 25OH-activated [3H]serine incorporation into SM and ceramide to the level of cells expressing OSBP (Figure 7A). Immunoblots of total cell lysates further established that site 2 mutations did not affect the slower migrating site 1 phosphospecies (Figure 7C). Immunofluorescence localization experiments confirmed that OSBP-S3A and -S3D localization was similar to wild type in control and 25OH-treated cells (Supplemental Figure S1).

Because individual phosphorylation site mutations did not affect OSBP function, we decided to test a series of four site 1/site 2 double mutants (Figure 7, B and D). Three of these mutants (S381A/S3A, S5E/S3D, and S5E/S3A) restored a 2.5-fold activation of SM synthesis by 25OH without affecting basal synthesis. In contrast, expression of OSBP-S381A/S3D failed to restore oxysterol activation of SM synthesis without affecting basal SM or ceramide synthesis. OSBP-S381A/S3D was highly expressed compared with mock cells, in which RNA interference silencing is >95% effective, but was only 20–25% compared with the other double mutants (Figure 7D).

To determine why OSBP-S381A/S3D was defective with respect to sterol activation of SM synthesis, we examined its intracellular localization using immunofluorescence and transmission electron microscopy (TEM) in OSBP-depleted CHO cells (Figure 8). Site 1/site 2 double mutants that restored SM synthesis in CHO cells (Figure 7) showed the expected translocation from an ER/cytoplasmic compartment to the Golgi apparatus in response to oxysterol treatment (Figure S1). Interestingly, OSBP-S381A/S3D was exclusively localized to unusual filamentous structures around the nucleus of CHO cells (Figure 8A). These structures were absent in adjacent nonexpressing cells and unaffected by 25OH treatment and colocalized with VAP-A, which is normally present in an extended reticular network. These aggregated structures did not localize with the trans-Golgi enzyme PI4KIIIβ, but cells expressing OSBP-S381A/S3D had a dispersed PI4KIIIβ staining pattern (Figure 8B). A similar pattern of ER aggregation and colocalization of OSBP-A381A/S3D with VAP-A was observed in HeLa cells (unpublished data). To determine whether OSBP-A381A/S3D was defective with respect to interaction with VAP-A, we mutated the FFAT motif (FF-AA), expressed the triple mutant in CHO cells, and determined its location relative to endogenous VAP-A (Figure 8C). Unlike the double phosphomutant, OSBP-S381A/S3D/FF-AA was dispersed in the cytoplasm or in small punctate structures and did not colocalize with VAP-A or cause its aggregation. Interestingly, OSBP-S381A/S3D/FF-AA did not localize to the Golgi apparatus when cells were treated with 25OH, suggesting a nonfunctional PH domain.

To better visualize ultrastructural changes to the ER induced by OSBP-S381A/S3D, we used TEM to analyze CHO cells expressing the mutant (Figure 8, D–I). CHO cells expressing OSBP displayed

![Figure 5](image-url)

**Figure 5:** OSBP site 1 phosphomutants restore 25OH-activated SM synthesis in OSBP-depleted CHO cells. (A and B) CHO cells deficient in endogenous OSBP were transiently transfected for 48 h with wild-type OSBP and site 1 phosphomutants. SM (A) and ceramide (B) synthesis was measured by [3H]serine incorporation after treatment with 25OH (6 μM, black bars) or solvent control (empty bars) for 6 h. Results are the mean and SEM of three experiments. (C) Total cell lysates from mock and OSBP-transfected cells were resolved by SDS–6%PAGE and immunoblotted with an OSBP polyclonal antibody.
normal organelle morphology with many ribosome-studded ER tubules (Figure 8D, arrowheads). In contrast, CHO cells expressing OSBP-S381A/S3D contained extended aggregates of membrane tubules that were close to the nucleus and devoid of ribosomes (Figure 8E–G). Cells containing these ER aggregates were also devoid of normal ER tubules elsewhere in the cell. Immunostaining for OSBP-S381A/S3D showed the presence of 5-nm colloidal gold particles in areas enriched in tubule aggregates (Figure 8, H and I). Thus OSBP-S381A/S3D caused the ER to collapse into filamentous bundles and dispersed the Golgi apparatus, perhaps contributing to its poor expression compared with other mutants (Figure 7D).

To assess whether the colocalization of OSBP-S381A/S3D and VAP-A observed by immunofluorescence was the result of enhanced physical interaction, and whether other phosphomutants might display altered binding, we assessed interaction with VAP-A using coimmunoprecipitation from OSBP-transfected CHO cells (Figure 9). Detergent extracts from cells expressing OSBP site 1 and site 2 phosphomutants were immunoprecipitated using a VAP-A antibody and probed for OSBP. There were no reproducible differences in the amount of single OSBP mutants coimmunoprecipitated with VAP-A under control or 25OHTreated conditions (Figure 9A). Compared with wild-type, OSBP-S381A/S3A, -S5E/S3D, and -S5E/S3A double mutants had similar levels of expression and interaction with VAP-A (Figure 9B). Despite reduced expression, OSBP-S381A/S3D showed similar levels of coimmunoprecipitation with VAP-A compared with the other double mutants. When quantified relative to input, approximately sixfold more OSBP-S381A/S3D was coimmunoprecipitated with VAP-A compared with the other three double mutants.

FIGURE 6: A site 2 phosphorylation motif is adjacent to the PH domain. (A) OSBP and the indicated site 1 and site 2 mutants were transiently transfected into CHO cells for 48 h. Cells were then harvested for immunoblotting or labeled with $^{32}$PO$_4$ (0.3 mCi/ml) for 4 h. $^{32}$PO$_4$-labeled OSBPs were immunoprecipitated with monoclonal 11H9, resolved by SDS–PAGE, and subject to autoradiography for 12 h at −80°C. (B) $^{32}$PO$_4$-labeled OSBPs were digested with trypsin and resolved by two-dimensional thin-layer electrophoresis and chromatography on cellulose-coated plates (Mohammadi et al., 2001). Arrows indicate the position of the site 2 phosphopeptide. Asterisks indicate the positions of site 1 phosphopeptides.

FIGURE 7: Identification of a site 1/site 2 double mutant that does not restore sterol activation of SM synthesis. (A) OSBP site 2 mutants (S3A and S3D) were transiently expressed in OSBP-depleted CHO cells, and SM and ceramide synthesis was measured by $[^{3}H]$serine incorporation as described in the legend to Figure 5. (B) A series of four OSBP site1/site2 double phosphomutants were transiently expressed in OSBP-depleted CHO cells, and SM and ceramide synthesis was measured by $[^{3}H]$serine incorporation as described in the legend to Figure 5. (C and D) Immunoblot analysis of site 2 (C) and site 1/site 2 double phosphomutants (D) expressed in OSBP-depleted CHO cells.
DISCUSSION

The differential localization of OSBP between the ER and Golgi apparatus in response to exogenous and endogenous sterol ligands and the resultant effects on sterol-responsive activities in the Golgi apparatus suggest that OSBP transfers cholesterol and/or oxysterols between these organelles. This activity can be partially reconstituted in vitro if cholesterol extraction and transport occurs along a concentration gradient. This is unlikely to occur in vivo, since the ER is cholesterol-poor relative to the Golgi and other organelles (Ikonen, 2008). In this study, we identify two factors that could facilitate sterol loading and transport to the Golgi apparatus by OSBP: phosphorylation on two serine-rich motifs (sites 1 and 2) that control sterol binding and interaction with VAP-A in the ER and competition for sterol binding by PI(4)P. OSBP phosphorylation regulates the ER-specific sterol/VAP-A binding but not the binding of PI(4)P, a ligand highly enriched in the Golgi apparatus, where it could exchange with cholesterol.

Site 1 phosphorylation is initiated at S381, which triggers phosphorylation of four adjacent serine residues by a CKI-like activity (Mohammadi et al., 2001). The organization of site 1 is reminiscent of the serine-rich motif in CERT, which is initiated by PKD phosphorylation and sensitive to cellular cholesterol and SM content (Kumagai et al., 2007). However, OSBP-S381 is not phosphorylated by PKD, nor is PKD phosphorylation of S242 involved in site 1 phosphorylation (unpublished data). While the PH- and FFAT-domain activity was unaffected by site 1 mutations, introducing a constitutive negative charge in OSBP-S5E, which is proximal to the α-helical lid of the sterol-binding domain, caused a significant increase in the maximal binding of aqueous dispersions of 25OH without significantly affecting the apparent $K_d$. The partially phosphorylated mimetic OSBP-S3E had sterol-binding properties similar to OSBP and OSBP-S381A, indicating that maximal phosphorylation of site 1 is necessary to increase sterol binding. The cholesterol-binding capacity of OSBP-S5E for cholesterol dispersions was similarly increased relative to dephosphomimics, but dissociation constants could not be reproducibly measured due to irregular binding curves, possibly related to the poor ligand solubility and nonequilibrium conditions that were
used to avoid protein inactivation. OSBP-S5E also extracted more cholesterol from liposomes compared with dephosphomimics, but the extracted cholesterol was not efficiently transferred to acceptor liposomes. This implies that phosphorylation at site 1 increases the initial capacity of OSBP to bind sterols at a donor membrane, and subsequent dephosphorylation facilitates the release of the sterol ligand to an acceptor membrane. In intact cells, $^{32}$PO$_4$ pulse-chase experiments showed that phosphorylation of site 1 has a half-life of $\sim$20 min (Ridgway et al., 1998a). This is consistent with a dynamic phosphorylation cycle at this site that, in conjunction with other protein/lipid interactions and phosphorylation sites, regulates sterol binding and release.

Site 1 phosphorylation is correlated with changes in OSBP localization, particularly in response to SM and cholesterol metabolism and content (Ridgway et al., 1998b; Mohammadi et al., 2001). However, individual site 1 mutations did not affect in vitro interactions with VAP-A or Pi(4)P or the distribution of OSBP between the ER and Golgi apparatus in CHO cells, indicating other variables are involved. Site 2 was of interest in this regard, since it encompasses three serine residues in a glutamate/aspartate-rich region located next to the PH domain. $^{32}$PO$_4$ labeling and peptide mapping of site 1 and site 2 mutants indicated no interdependence between the sites and, similar to site 1, individual site 2 phosphomutants were normal with respect to cellular localization and restoration of 25OH-activated SM synthesis. However, expression of a mutant mimicking dephosphorylation of site 1 and phosphorylation of site 2 (OSBP-S381A/S3D) collapsed the peripheral ER into membrane bundles and whorls. The similarity between OSBP W174A and OSBP-S381A/S3D with respect to ER perturbation and VAP-A interaction suggests that the latter mutant also has a nonfunctional PH domain. In support of this conclusion, mutation of the FFAT domain in OSBP-S381A/S3D prevented interaction with VAP-A and collapse of the ER, but the mutant did not regain the ability to localize to the Golgi in the presence of 25OH. This suggests a phosphorylation state that causes reduced cholesterol-binding activity, increased VAP-A binding, and reduced affinity for PH domain ligands in the Golgi apparatus. Based on the normal activity of other single and double site 1/site 2 mutants, subsequent changes in the phosphorylation status of either site 1 or 2, by as-yet-unknown kinases and/or phosphatases, would disengage OSBP from the ER and promote interaction with the Golgi apparatus.

OSBP homologues in diverse phyla are distinguished by two conserved histidine residues (H522 and H523 in human OSBP) situated at the entrance of the sterol-binding pocket (Lehto and Olkkonen, 2003; Im et al., 2005). This histidine pair is essential for Osh4p function but dispensable for in vitro sterol binding and transfer activity (Im et al., 2005; Raychaudhuri et al., 2006). Instead, these histidine residues form an essential contact with the 4-phosphate residue of Pi(4)P that mediates competitive binding with dehydroergosterol (de Saint-Jean et al., 2011). This suggested that Osh4p transport of ergosterol to Pi(4)P-enriched compartments is coupled to exchange and transport of Pi(4)P in the opposite direction. This model is particularly suited to OSBP, which could transport cholesterol against a gradient to the Pi(4)P-enriched trans-Golgi network in conjunction with Pi(4)P transport in the opposite direction. Initial in vitro experiments lend support to this model; cholesterol extraction from liposomes by OSBP was specifically competed by Pi(4)P, and radiolabeled Pi(4)P was extracted by OSBP from cholesterol-free liposomes. Based on the percent of extraction of cholesterol and Pi(4)P from liposomes by increasing amounts of OSBP (compare Figure 2E with Figure 3D), the relative affinity of OSBP for Pi(4)P and cholesterol is comparable. Interestingly, the increased sterol-binding activity of OSBP-S5E toward cholesterol was not evident with Pi(4)P. This indicates that phosphorylation events that affect the unique electrostatic and hydrogen-bonding contacts for sterols and Pi(4)P could regulate an organelle-specific interaction with these ligands.

To test whether Pi(4)P stimulated the transfer of cholesterol to acceptor liposomes, we used a modified assay that involved pre-extraction of cholesterol from donor liposomes. In this context, Pi(4)P stimulated the transfer of cholesterol by OSBPs, suggesting an exchanged-based mechanism similar to Osh4 (de Saint-Jean et al., 2011). However, this is not the only explanation, since efficient cholesterol transfer by OSBP also required its N-terminal

**FIGURE 9:** Enhanced interaction between VAP-A and OSBP-S381A/S3D. (A) Triton X-100 extracts were prepared from OSBP-depleted CHO cells transiently expressing the indicated single OSBP and site 1 or site 2 mutants and treated without (−) or with (+) 25OH (6 μM for 2 h). VAP-A was immunoprecipitated from extracts with a polyclonal antibody and immunoblotted for OSBP and VAP-A. (B) OSBP was coimmunoprecipitated with VAP-A from Triton X-100 extracts of OSBP-depleted CHO cells expressing site 1/site 2 double phosphomutants. (C) The amount of OSBP coimmunoprecipitated with VAP-A in (B) was quantified as a percentage of total OSBP input (average and range of two experiments).

and Nir proteins, Pi-binding/transfer proteins that interact with VAP-A via FFAT domains (Amarilio et al., 2005; Peretti et al., 2008). In these instances, enhanced interaction with VAP-A causes cross-linking of ER tubules and collapse of the reticular network into membrane bundles and whorls. The similarity between OSBP W174A and OSBP-S381A/S3D with respect to ER perturbation and VAP-A interaction suggests that the latter mutant also has a nonfunctional PH domain. In support of this conclusion, mutation of the FFAT domain in OSBP-S381A/S3D prevented interaction with VAP-A and collapse of the ER, but the mutant did not regain the ability to localize to the Golgi in the presence of 25OH. This suggests a phosphorylation state that causes reduced cholesterol-binding activity, increased VAP-A binding, and reduced affinity for PH domain ligands in the Golgi apparatus. Based on the normal activity of other single and double site 1/site 2 mutants, subsequent changes in the phosphorylation status of either site 1 or 2, by as-yet-unknown kinases and/or phosphatases, would disengage OSBP from the ER and promote interaction with the Golgi apparatus.
PH domain (Figure 2F; Ngo and Ridgway, 2009). The PH domains could facilitate exchange by targeting OSBP to PI(4)P-enriched membranes or to regulate ligand transfer by inter/intramolecular interactions.

This study provides a new perspective on OSBP activity in the ER-Golgi secretory pathway through identification of serine phosphorylation motifs that regulate the ER-specific phase of the OSBP transfer cycle involving VAP-A and sterol binding. The subsequent release of sterols in the Golgi apparatus could involve ligand exchange with PI(4)P. This exchange could be driven by OSBP-dependent activation of PI(4)P synthesis by the sterol-regulated PI4KIIx (Banerji et al., 2010). Alternatively, PKD could also regulate OSBP at the Golgi apparatus via direct phosphorylation (Nhek et al., 2010) or activation of PI(4)P synthesis by PI4KIIIb (Hauser et al., 2005).

MATERIALS AND METHODS

Materials
Cholesterol and 25OH were purchased from Steraloids (Newport, RI). [3H]cholesterol, [3H]25OH, [32P]PO4, [3H]serine, and [3H]serine were from Perkin Elmer-Cetus (Waltham, MA). PI(4)P (dipalmitoyl and bovine brain) and PI(4,5)P2 (dipalmitoyl) were from Echelon (Salt Lake City, UT). Phosphatidylcholine (PC) was purchased from Avanti Polar Lipids (Alabaster, AL). OSBP polyclonal and monoclonal antibodies were as previously described (Ridgway et al., 1992; Mohammadi et al., 2001). A VAP-A–specific antibody was previously described (Wyles et al., 2002). A colloidal, gold-conjugated secondary antibody was purchased from Sigma-Aldrich (St. Louis, MO). Alexa Fluor 488– and 594–conjugated secondary antibodies were from Molecular Probes (Eugene, OR). Secondary antibody was purchased from Sigma-Aldrich (St. Louis, MO). Alexa Fluor 488– and 594–conjugated secondary antibodies were as previously described (Ridgway et al., 1992; Mohammadi et al., 2001). Inhibitors were purchased from EMD Chemicals (Gibbstown, NJ). Sf21 insect cells (Burlington, ON). IRDye 800– and 680–conjugated secondary anti–

Sterol binding, extraction, and transfer assays
The specific binding of 25OH by purified OSBP and OSBP mutants (12 pmol protein) was carried out in 75 μl of 10 mM HEPES (pH 7.4), 100 mM KCl, 2% (wt/vol) polyvinyl alcohol, and [3H]25OH at 4°C for 16 h. A charcoal-dextran slurry (45 μl) was added for 30 min, after which the mixture was sedimented by centrifugation at 10,000 × g for 10 min, and protein-bound sterol in the supernatant was measured by liquid scintillation counting. Specific binding was defined as total [3H]25OH binding minus binding in the presence of a 40-fold excess of unlabeled 25OH.

[3H]cholesterol-binding assays were similar to those described above, except 0.05% Triton X-100 was included. After incubation at 20°C for 2 h, OSBP-bound [3H]cholesterol was isolated by incubation with 12 μl of TALON affinity resin for 30 min with constant mixing. The resin was washed four times with 500 μl of 10 mM HEPES/100 mM KCl and collected by centrifugation at 5000 × g for 1 min. OSBP-bound [3H]cholesterol was released from the resin with 150 mM imidazole and quantified by liquid scintillation counting.

The liposomal [3H]cholesterol extraction assay was previously described (Ngo and Ridgway, 2009). The cholesterol transfer activity of OSBP and phosphomutants was assayed by a modified method (Ngo and Ridgway, 2009). Donor and acceptor liposomes composed of [34C]PC:PE:PS:[3H]cholesterol (69:20:10:1, mol/mol) and PC:PS:PI:PI:PI with lactosyl-PE (58:20:10:2:10, mol/mol), respectively, were prepared by extrusion and sedimented at 15,000 × g for 5 min prior to use. [34C]PC was included in donor liposomes to correct for contamination of acceptor liposomes after sedimentation. The assay was modified to include a preextraction step prior to addition of acceptor liposomes. Donor liposomes (10 nmol) were incubated with 100 pmol of recombinant OSBPs and 3 μg fatty acid–free bovine serum albumin (BSA) in 80 μl of liposome buffer (25 mM HEPES, 150 mM NaCl, and 1 mM EDTA, pH 7.4) at 25°C. After 20 min, the reaction was stopped by addition of ice (to measure transfer). Acceptor liposomes were precipitated with 100 μg of a VAP-A–specific antibody (Ngo et al., 2005) and stored at -80°C.

PI(4)P competition and extraction assays
[32P]PI(4)P was isolated from HeLa cells incubated for 24 h in phosphate-free media containing 0.5 μCi/ml of [32P]PO4. Cells were harvested in 1 ml of phosphate-buffered saline (PBS) and four 4 ml of CHCl3/MeOH/12N HCl (2:4:0.1, vol/vol), 1.2 ml of CHCl3, and 1.2 ml of water were added. The organic phase was dried under nitrogen and resolved by TLC in CHCl3/MeOH/4M NH4OH (90:70:20, vol/vol), and [32P]PI(4)P was identified by autoradiography and comigration with an authentic standard (purification was repeated once more). Competitive extraction assays were performed using liposomes containing 1 mol% [3H]cholesterol and increasing mol% of PI, PI(4)P, or PI(4,5)P2. Direct extraction of 0.5% [3H]PI or [32P]PI(4)P by OSBPs from cholesterol-free liposomes was assayed as described for [3H] cholesterol.
Immunoblotting and immunoprecipitation

Total cell lysates were prepared in SDS–PAGE sample buffer (12.5% SDS, 30 mM Tris–HCl, 12.5% glycerol, and 0.01% bromophenol blue, pH 6.8), heated at 90°C for 5 min, resolved on SDS–PAGE, and transferred to nitrocellulose. Proteins were visualized by incubation with primary antibodies for 1–2 h at 20°C, which was followed by incubation with secondary IRDye 800- or 680-conjugated antibodies for infrared detection using an Odyssey Infrared Imaging System and quantification using Odyssey Application Software version 3.0 (LI-COR Biosciences, Lincoln, NE).

Immunoprecipitation of VAP-A and OSBP was carried out as previously described (Wyles et al., 2002). Briefly, CHO cells expressing OSBP phosphorylation mutants were lysed in PBS containing 5 mM KCl, 2 mM EDTA, 2 mM EGTA, and 0.5% Triton X-100, and were precleared by centrifugation at 14,000 × g for 15 min at 4°C. VAP-A was immunoprecipitated from supernatants using a polyclonal antibody and protein A-Sepharose, washed three times with PBS/0.1% Triton X-100, and resuspended in SDS–PAGE sample buffer. Immunoprecipitates were resolved by SDS–PAGE and immunoblotted for OSBP using monoclonal 11H9. The degree of VAP-A-OSBP communoprecipitation was quantified relative to OSBP input.

Immunofluorescence and transmission electron microscopy

CHO cells cultured on glass coverslips were fixed with 4% (wt/vol) paraformaldehyde in PBS for 15 min and permeabilized with 0.5% (wt/vol) Triton X-100 for 20 min at 4°C. Coverslips were incubated sequentially with a primary antibody in PBS containing 1% (wt/vol) BSA at 20°C for 1 h, which was incubation with followed by an Alexa Fluor–conjugated secondary antibody for an additional 1 h at 20°C. Coverslips were mounted on glass slides with Mowiol 4–88, and images were captured using a Zeiss LSM510/AxioVert 200M inverted microscope with Plan-Apochromat 100×/1.40 numerical aperture (NA) oil-immersion objective or a Zeiss LSM510 laser-scanning confocal upright microscope with a Plan-Apochromat 63×/1.40 NA oil immersion objective (Jena, Germany).

Thin-section (80–100 nm) TEM of CHO cells was performed as described previously (Lagace and Ridgway, 2005). For immunoelectron microscopy, CHO-K1 cells were fixed with 4% (wt/vol) paraformaldehyde/0.5% (vol/vol) glutaraldehyde in 0.1M sodium cacodylate buffer for 30 min. Thin sections mounted on nickel grids were incubated with OSBP monoclonal 11H9 overnight at 4°C, which was followed by an 1-h incubation with 5-nm colloidal, gold-conjugated goat anti-mouse secondary antibody at 20°C. Grids were fixed with 2.5% glutaraldehyde and stained with 2% (wt/vol) uranyl acetate and lead citrate. Images were captured using a JEOL (Tokyo, Japan) JEM 1230 transmission electron microscope at 80 kV and a Hamamatsu (Hamamatsu, Japan) ORCA-HR digital camera.

Analysis of PIP binding by OSBP phosphomutants

Liposomes composed of PC and PE and containing either PI, PI(4)P, or PI(4,5)P2 (70:20:10, mol/mol) were incubated with 100 pmol of purified protein for 25 min at 25°C. Samples were centrifuged at 100,000 × g for 30 min, and the supernatant and pellet fractions were analyzed by SDS–PAGE and stained with Coomassie Blue, and OSBP distribution was quantified using a LI-COR Odyssey IR imaging system.

Lipids (100–300 pmol) and solvent control were spotted onto a Hybond-C nitrocellulose membrane, which was incubated in blocking buffer (20 mM Tris HCl, pH 7.4, 150 mM NaCl, 3% [wt/vol] fatty acid–free BSA and 0.1% [vol/vol] Tween-20) at 4°C for 50 nM OSBPs for 1 h. Bound OSBPs were visualized with 11H9 monoclonal and IRDye 800-conjugated antibodies for infrared detection using an Odyssey Infrared Imaging System.

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