Membrane Association of the PTEN Tumor Suppressor: Molecular Details of the Protein-Membrane Complex from SPR Binding Studies and Neutron Reflection

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Abstract

The structure and function of the PTEN phosphatase is investigated by studying its membrane affinity and localization on in-plane fluid, thermally disordered synthetic membrane models. The membrane association of the protein depends strongly on membrane composition, where phosphatidylinosine (PS) and phosphatidylinositol diphosphate (P(4,5)P2) act pronouncedly synergistic in pulling the enzyme to the membrane surface. The equilibrium dissociation constants for the binding of wild type (wt) PTEN to PS and P(4,5)P2 were determined to be $K_d \approx 12 \mu M$ and 0.4 $\mu M$, respectively, and $K_d \approx 50 nM$ if both lipids are present. Membrane affinities depend critically on membrane fluidity, which suggests multiple binding sites on the protein for P(4,5)P2. The PTEN mutations C124S and H93R show binding affinities that deviate strongly from those measured for the wt protein. Both mutants bind PS more strongly than wt PTEN. While C124S PTEN has at least the same affinity to P(4,5)P2 and an increased apparent affinity to P(3,4,5)P3, due to its lack of catalytic activity, H93R PTEN shows a decreased affinity to P(4,5)P2 and no synergy in its binding with PS and P(4,5)P2. Neutron reflection measurements show that the PTEN phosphatase “scouts” along the membrane surface (penetration $<5 \AA$) but binds the membrane tightly with its two major domains, the C2 and phosphatase domains, as suggested by the crystal structure. The regulatory C-terminal tail is most likely displaced from the membrane and organized on the far side of the protein, $\sim 60 \AA$ away from the bilayer surface, in a rather compact structure. The combination of binding studies and neutron reflection allows us to distinguish between PTEN mutant proteins and ultimately may identify the structural features required for membrane binding and activation of PTEN.

Introduction

Lipid-mediated cell signaling with phosphatidylinositol phosphates (PIPs) results in exquisite spatio-temporal control of vital cell functions. The rich functionality of the PIP headgroup with its two major domains, the C2 and phosphatase domains, as suggested by the crystal structure. The regulatory C-terminal tail is most likely displaced from the membrane and organized on the far side of the protein, $\sim 60 \AA$ away from the bilayer surface, in a rather compact structure. The combination of binding studies and neutron reflection allows us to distinguish between PTEN mutant proteins and ultimately may identify the structural features required for membrane binding and activation of PTEN.

Vacuole morphology [2]. In the plant Arabidopsis, PTEN is expressed in pollen grains and is required for pollen maturation and survival [3]. PTEN has an asymmetric distribution in the slime mold Dictyostelium discoideum, resulting in accumulation of P(3,4,5)P3 at the leading edge of migrating cells [4]. PTEN in honeybees plays a role in nutrient sensing and, thereby, quencher differentiation [5]. DAF-18 is a PTEN homolog in Caenorhabditis elegans worms that regulates aging [6]. In mice, loss of PTEN is embryonic lethal. PTEN affects many organs, but its roles in the nervous and immune systems have been particularly well studied [7–9]. With its function as a phosphatase, PTEN is the major antagonist to PI-3-kinase (PI3K), limiting basal levels of P(3,4,5)P3 in the inner leaflet of the plasma membrane. Because elevated P(3,4,5)P3 levels lead to unconditional cell survival and growth, PTEN inactivation is associated with multiple disease states, including many types of cancer [10]. In fact, PTEN is the second most commonly mutated protein in sporadic human...
tumors [11], and its activity as a tumor suppressor is essential [12]. More recently, roles of PTEN in human behavior have been identified. For example, the protein suppresses responses to drugs of abuse [13], and PTEN mutations were observed in autistic patients with macrocephaly [14–20]. Despite its importance, our current knowledge of the regulation of PTEN–PIP interactions is largely indirectly based on the analysis of cancer mutations [21]. Clearly, an assessment of the molecular origin of PTEN’s membrane association and phosphatase activity will benefit our understanding of key processes in organisms from yeast to humans.

From the N-terminus to the C-terminus, PTEN consists of a PI(4,5)P2-binding module (PBM), the predominantly α-helical phosphatase domain (PD), a C2-domain dominated by β-sheet, and a ~50 amino acid (AA) C-terminal tail that includes two PEST motifs and a PDZ motif. Plasma membrane binding of PTEN is partially electrostatically driven, with phosphatidylserine (PS) and PI(4,5)P2 being the main anionic phospholipids in the cytosolic leaflet that interact with PTEN [22]. The N-terminus forms the PBM and may thus initiate the binding process [21]. By binding to PS, the C2 domain helps target the protein to the bilayer where it hydrolyzes its substrate PI(3,4,5)P3. Membrane specificity is provided by selective binding of the PBM to PI(4,5)P2, targeting the protein to the plasma membrane, which is particularly rich in PI(4,5)P2 compared with other intracellular membranes. On the other hand, phosphorylation of the C-terminal tail abrogates membrane binding of the protein [23], thus providing another level of control of PTEN [21].

The structural arrangement and localization of PTEN at the membrane are currently poorly understood. The only crystal structure of PTEN available [24] shows that the C2 domain and the PD stabilize each other through interdomain contacts, but this structural model was developed with a truncated protein where parts of the N-terminus (AA residues 1–6), an interconnecting loop on the C2 domain (AAs 286–309) and the C-terminal tail (AAs 354–403) were deleted to allow crystallization. In addition, peptide segments adjacent to these deletions are not defined in the crystal structure. Hence, we do not have a structural basis to ask how does the complex network of interactions between lipid species and protein binding domains control PTEN’s enzymatic activity at the membrane surface? Which structural aspects are characteristic of PTEN membrane association, and how is membrane association altered by mutations at the protein-membrane interface? How do such structural alterations translate into changes in the membrane affinity of PTEN?

In this work, we develop methodologies to approach such questions. The key challenge here is to characterize the system—protein and membrane—at high resolution while the bilayer is in a thermally disordered, fluid state. We have recently demonstrated that this can be indeed achieved by analyzing neutron reflection (NR) [25] from substrate-supported, in-plane fluid, lipid membranes [26]. The membrane model systems used in these studies, sparsely-tethered bilayer lipid membranes (sBLMs), have been thoroughly studied in terms of their structural [26], functional [27] and dynamic properties [28]. In such studies, NR provides the one-dimensional (1D) neutron scattering length density (nSLD) profiles of the interfacial structures [29], i.e., projections of the material distribution along the surface normal. However, by accounting for molecular volumes and properties, chemical connectivities and the internal structure of the protein, the “missing dimensions” can be filled in by computer models [30–32]. In such cases, one can resolve with high resolution the spatial association [33] and orientation [34] of a protein on the bilayer. In the case of PTEN, the situation is more complex, because ~20% of the AAs of the wild-type protein are either omitted or not resolved in the crystal structure. In addition, a recent IR investigation showed that the protein undergoes conformational changes upon membrane binding [35]. Despite these complications, we describe here the first steps to resolving the structure of a carefully designed PTEN/membrane model system.

Materials and Methods

[36] Preparation of PTEN Protein.

Human PTEN with a six-histidine tag at the C-terminus was expressed in Escherichia coli BL21 (DE3) bacteria as described [35]. We produced wild-type (wt) and an enzymatically disabled mutant (C124S PTEN) [37], an autism-related H–R mutant (H93R PTEN) [38] and a truncated PTEN whose crystal structure was reported [24]. The plasmid of the truncated PTEN was provided by Dr. N. R. Leslie (University of Dundee). PTEN proteins were purified with a HiTrap HP kit from GE Healthcare, a Superdex 200 column and a MonoQ anion-exchange column. For SPR or NR, the protein was centrifuged at 3,200 rpm for 10 min., dialyzed in 100 mM NaCl, 10 mM HEPES, 1 mM EDTA, 1 mM β-mercaptoethanol (βME), pH 7.0 (buffer 1) overnight using a Slide-A-Lyzer dialysis cassette (Thermo Scientific, mod. 66370), centrifuged again for 10 min., and the supernatant used. Unless otherwise stated, reagents were of p.a. grade and obtained from Sigma-Aldrich.

Preparation of sBLMs

3-1-1 nm glass slides from Fisher Scientific (for SPR) and 3, 580 μm thick, [100]-cut Silicon wafers (Silicon Quest International; for NR) were cleaned first in 5 vol% Hellmanex (Hellma GmbH) and then Norchoromix (35 g per 2.5 L of 10 M H2SO4; Goxlak Laboratories), followed by excessive rinsing with ultrapure water (Millipore) and absolute EtOH (Pharmco-Aaper) and drying in an N2 gas stream. They were coated with Cr (~20 Å) and Au (~150 Å and ~450 Å for NR and SPR, respectively) by sputtering in a high-energy magnetron (ATC Orion; AJA International) at 0.15 μTorr. Some of the samples produced for NR experiments incorporated an iron-nickel (“permalloy”) bonding layer instead of Cr. After the sputtering, x-ray reflectometry (X-ray/Brucker AXS) showed a typical RMS surface roughness of ~5 Å of the gold film. After sputtering, the gold-coated Si wafers were immediately transferred into a 3:7 (mol/mol) 0.2 mM (total concentration) ethanolic solution of ζ 20–20 octadec-9-enoic acid)-3,6,9,12,15,18,22-heptaoxatetracont-31-ene-1-thiol (HC18; structure, see figure in Information S1) and βME to form a self-assembled monolayer (SAM). HC18 was a kind gift from Dr. D. J. Vanderklip (Institute for Bioscience and Biotechnology Research, IBBR, Rockville, MD), synthesized similarly to related compounds described earlier [26,39]. sBLMs were completed by precipitation of phospholipids on the preassembled SAM by rapid solvent exchange (RSE) [26,40]. Dipalmitoyl-[DP]-PI(4,5)P2 and DPPG(3,4,5)P3 were from Cayman Chemical. Brain-derived PI(4,5)P2 and all other phospholipids and cholesterol were from Avanti Polar Lipids. At T = 45°C, ethanolic solutions of lipid mixtures (~5 mg/mL) of the appropriate compositions were used to incubate the SAM-covered substrates, followed by a rapid replacement with aqueous buffer 1. To avoid the formation of incomplete bilayers, phospholipid mixtures that contained ~20 mol% of anionic lipids were supplemented with 5 mol% cholesterol. For compositions that contain ~40 mol% total anionic lipids, this procedure leads to the formation of complete bilayers in the sBLMs, as observed with electrochemical...
impedance spectroscopy (EIS) and NR (Shenoy, Heinrich and Lösche, unpublished results).

**Surface Plasmon Resonance (SPR) and Electrochemical Impedance Spectroscopy (EIS)**

Glass slides with \(~450\) \(\text{Å}\) of gold were coupled to a prism in a custom-built SPR instrument from SPR Biosystems (Germantown, MD) in the Kretschmann configuration. A superluminescent LED (EXS7510, Exalos AG, Switzerland) excited surface plasmons in the gold film at \(\lambda = 763.8\) \(\text{nm}\). The LED emission was focused on the sample by means of a hemicylindrical prism, such that a range of incident angles were covered on the sample, reflected, and collected on a position-sensitive CCD detector. The illuminated sample area was \(10\ \text{mm} \times 100\ \text{µm}\) with the long side perpendicular to light propagation. The reflected light was collected on a CCD (Hamamatsu C10990) with 250 lines of 1024 pixels at 25 \(\mu\text{m}\) resolution. In the standard, single channel measurement mode, all 250 lines were binned into one line of 1024 pixels. The system had a time resolution of 0.1 s with a sensitivity of \(3 \times 10^{-7}\) reflectivity units (RU) or better. The dynamic range in refractive index was \(n = 1.33\) to 1.41. A temperature controller (Wavelength Electronics LFI-3751) was used with a range from ambient to \(50\) \(\text{ºC}\) with \(0.005\) \(\text{ºC}\) resolution. The sample cell was composed of a Teflon cylinder of 6 mm inner diameter with a volume of \(\sim 1\ \text{mL}\). A modified IKA (Wilmington, NC) RW11 overhead stirrer was used following injection of protein into the sample cell.

Simultaneously with the SPR measurement, the instrument allowed EIS measurements using a Solartron 1287A potentiostat and 1260 frequency response analyzer if the stirrer was not in use. In standard data acquisition mode, the position, \(R\), of the reflection minimum as a function of incident angle within the convergent incident beam was determined at 0.2 s intervals. The EIS response was measured with a saturated silver-silver chloride (Ag|AgCl|-NaClaq|sat) microelectrode (Microelectrodes, mod. M-401F) with the auxiliary electrode consisting of a 0.25 mm diameter Pt wire (99.99% purity, Aldrich) coiled around the barrel of the reference electrode and the gold film used as the work electrode. Results were fitted to equivalent circuit models [26,27] using ZView (Scribner Associates, Southern Pines, NC).

In a standard equilibrium binding assay in SPR, a given concentration of protein, \(c_p\), was added to a stable bilayer. The increase in the SPR signal \(R\), due to an increase in refractive index at the bilayer/buffer interface was attributed to protein binding at the bilayer. Once \(R\) ceased to change, equilibrium was assumed to have been attained at \(R_{eq}\). A higher concentration of protein was then added to repeat the adsorption process. The change of \(R_{eq}\) as a function of \(c_p\) was fitted to [41]

\[
R_{eq} = \frac{c_p B_{max}}{c_p + K_d} \tag{1}
\]

where \(K_d\) is the equilibrium dissociation constant and \(B_{max}\) is the saturation of the SPR response at \(c_p \rightarrow \infty\). The calibration of the instrument is described in Information S1.

**Neutron Reflection**

NR measurements were performed on the Advanced Neutron Diffractometer/Reflectometer (AND/R) [42] or the NG1 reflectometer at the NIST Center for Neutron Research (NCNR). The sample was assembled in a dedicated flow-through sample cell, such that the fluid adlayer in the sample that bathes the stBLM was continuously exchanged by means of a microfluidic pump (MP5 with MP-X controller, Bartels Mikrotechnik GmbH). The resilience of the stBLMs permitted the NR characterization of the membrane at various solvent contrasts with the same physical sample. For contrast variation, exchanges of the buffer phase bathing the stBLM were performed in situ on the instrument, so that the neutron spectra were taken on exactly the same footprints on the wafers. This ensured that the inorganic substrates, in particular the SiO\(_x\)/Cr/Au or SiO\(_x\)/Fe-Ni/Au surface layers that dominate the interference patterns in the data, contributed identically to subsequent measurements. For each contrast, the reflectivity was measured at \(T = 25\) \(\text{ºC}\) over a momentum transfer, \(q_z\), range of 0.008 to 0.25 \(\text{Å}^{-1}\), taking typically 6 h to obtain sufficient statistics. Protein adsorption to a preformed stBLM was accelerated by continuously pumping protein solution through the sample before starting the measurement scan.

NR data were analyzed using the ga\(_{n}\) software package developed at NCNR [see Appendix A in ref. [43]]. We adapted a recently developed model which describes the distribution of the stBLM components in terms of thermally broadened distributions of molecular fragments along the surface normal [44]. We refer to this parameterization as the continuous distribution (CD) approach to data modeling. Catmull-Rom splines [45] were used to parameterize the distribution of the protein normal to the bilayer with a constant protein nSLD derived from the PTEN crystal structure. The spline profiles were constrained to be single-peaked. The structural relation of the protein profile with the lipid bilayer was established by letting a parameter that describes position of the support point of the spline closest to the membrane surface adjust to a position within or outside of the lipid headgroup. The resulting compound nSLD profile, \(\rho(z)\) still depends on the isotopic solvent contrast used in a particular NR scan. Multiple data sets from distinct isotopic contrasts with and without protein were all simultaneously fitted, thus generating a set of nSLD profiles consistent in terms of the underlying molecular structure [30,31]. The quality of the fit is quantified as the normalized \(\chi^2\) deviation between the model and the experimental data. To determine the variability of the resulting nSLD profiles within the experimental uncertainties, to quantify confidence intervals and to track correlations between the fit parameters, a Monte-Carlo resampling procedure [39] was implemented in ga\(_{n}\).

For a comparison of experimental nSLD profiles with the putative nSLD distribution of the truncated PTEN, we aligned the crystal structure with the membrane as proposed by Lee and coworkers [24]. As in earlier work [33], we then determined the nSLD in 0.5 \(\text{Å}\) thick slices parallel to the interface according to their atomic contents and solvent-excluded partial protein volumes measured using Connolly’s method [46] with a probe radius of 1.4 \(\text{Å}\). The resulting nSLD distribution was subject to Gaussian smoothing with the \(\sigma\) value determined from the modeling of the experimental nSLD profile, because this parameter is essentially determined by the roughness of the gold surface. Finally, the resulting PTEN profile was scaled to the maximum of the protein contribution to the nSLD profile and its position adjusted along \(z\) for maximum overlap with the experimental protein distribution.

**Results**

In this work, we seek a solid understanding of the molecular factors that determine PTEN membrane binding. This was accomplished with SPR measurements on stBLMs of various binary and ternary compositions for which DOPC was the invariable majority component that quantified affinities and saturating protein concentrations for wt PTEN and various mutants. We also report NR measurements on the association of wt PTEN mutant proteins to stBLMs, which provide direct
evidence of the protein structure on an in-plane fluid, thermally disordered membrane.

The most abundant anionic component of the cytosolic leaflet of mammalian plasma membranes, where PTEN exerts its phosphatase activity, is phosphatidylethanolamine (PS), PI(4,5)P2 and phosphatidic acid (PA) contribute to electrostatic interactions to a lesser extent. The relevant concentrations of these lipids in the plasma membrane are approx. 25% (PS), <5% (PA) and ~1% (PI(4,5)P2). Finally, PTEN's substrate, PI(3,4,5)P3, contributes only minimally to membrane charge, even though it is highly charged, as it is present only in minuscule concentrations [47]. In distinction, the plasma membrane is the major cellular repository of PI(4,5)P2, with a global concentration estimated to be ~1%.

SPR experiments on stBLMs precisely determine protein binding affinities and are invaluable as precursor experiments, conveniently optimizing sample preparation for the scattering experiments. We prepared stBLMs of various compositions of DOPE, DOPE and PI(4,5)P2. For membranes with greater than 10 mol% anionic lipids, we routinely added 5 mol% of cholesterol (chol) to the lipid mixture in RSE preparations because bilayers without chol were less complete, typically 95% coverage of the surface by the membrane. While chol may affect the morphology of PI(4,5)P2-containing membranes at higher concentrations, such low concentrations of cholesterol apparently do not impact the lateral distribution of PI(4,5)P2 (Gericke and Jiang, unpublished data). To study PTEN interactions with PS, we compared wt PTEN with H93R PTEN, whose affinity to PS is significantly enhanced [38]. The association of these proteins was further studied on stBLMs that contained PI(4,5)P2 as the only anionic lipid and to membranes with both PS and PI(4,5)P2. We also studied the binding of C124S PTEN to PIP2-containing stBLMs to quantify interaction with its specific substrate. By investigating these point mutants, one may also obtain information on the plasticity of the protein. Furthermore, we investigated the truncated PTEN variant used in the crystallization study [24], which will allow us to use the crystal structure as a starting point for our NR-determined structures.

PTEN Membrane Affinities

Earlier work showed that the binding affinity of PTEN to DOPE bilayers is low (>500 μM) [35]. Here, we performed SPR in a custom-designed sample environment that permits the preparation of stBLMs by RSE in situ, as well as the simultaneous measurement of SPR and EIS. This enables characterization of stBLMs quality before the injection of protein into the sample chamber. Protein studies were only conducted with bilayers that were >99% complete. Figure 1 shows an SPR binding curve of wt PTEN to an stBLM composed of DOPE:DOPS:PIP2 (68.8:28.6:2.6) (+chol). This binding curve shows two components with distinct affinities, Kd1 = 0.04 μM and Kd2 = 10 μM. The solid line is the best fit to the data. Similar experiments with DOPE:DOPS only or DOPE:PIP2 only showed always single-component adsorption processes.

Two pieces of information are gleaned from such data. (1) The equilibrium binding constant Kd quantifies the affinity of the protein to the membrane and is determined as the midpoint concentration of an adsorption process. It can be determined with confidence if the highest protein concentration (cp) sampled is well above the nominal value of Kd where the binding curve reaches half-saturation. In cases where the cp doesn’t reach the midpoint, for example because of protein aggregation at high cp, Kd can still be estimated as described in Information S1. A range of Kd values, which is consistent with the truncated data, is given as the result in such cases. (2) The maximum SPR shift, Bmax at cp→∞, is a measure of the amount of adsorbed material. Complementary studies of protein adsorption to stBLMs showed that densely packed monolayers of protein on the membrane surface, as quantified with NR, show values in the range Rg = 175–300 ng/cm² on the custom-built SPR instrument used in this study, depending on the physical size—and therefore the monolayer thickness—of the adsorbed protein (Shenoy and Löffler, unpublished data). Note that Kd and Bmax may be uncorrelated, i.e., the affinity and the amount of adsorbed protein are distinct characteristics of an adsorption process. For example, we frequently observe that the absorption of PTEN to PIP2 (PI(4,5)P2) has high affinity (Kd ~ 0.1 μM) with low amounts of protein adsorbed (Bmax < 60 ng/cm²), while large amounts of PTEN protein bind to PS with low affinity (Kd ~ 10 μM; Bmax > 175 ng/cm²). Tables 1 and 2 summarize the equilibrium binding constants of the three PTEN constructs to stBLMs of various compositions. Specific results are presented below.

PTEN binding to single anionic lipid components in stBLMs with PC. Of the PTEN variants tested, wt PTEN shows the weakest binding affinity, Kd = 12 μM [48], to DOPE membranes with DOPS as the sole anionic lipid. On the other hand, the B values reached at high protein concentrations (Bmax = 155 ng/cm²) suggest that we will be able to extract structural information from NR measurements. Given the differences in methodology, we consider this result to be consistent with the value, Kd = 22 ± 0.5 μM, reported earlier for a similar lipid composition [35]. H93R PTEN has a 4-fold higher binding affinity to PS with Kd = 3.1 ± 0.3 μM, again in agreement with earlier measurements using different techniques [38]. This is interesting, as the point mutation is in the PD—distant from both the PBM and C2 domains that are thought to mediate membrane association. As with wt PTEN, we were unable to determine Kd for the truncated PTEN because we could not use large enough
This indicates a remarkable synergy in binding of any of the mutants for which this lipid composition was studied. Surprisingly, the binding of WT protein, is similar to those of the mutant proteins, its affinity to (PS + PI(4,5)P2), which the mutants apparently lack.

Table 1. Quantification of wt PTEN Binding to stBLMs by SPR.

| Composition | \(K_d\) (\(\mu\)M) | \(B_{max}\) (ng/cm²) |
|-------------|-------------------|---------------------|
| PC:PS (+chol) | 70:30          | 11.9±0.4             | 155±3   |
| PC:PI(4,5)P2 | 99:3:0:7        | 0.4±0.1              | 23±1    |
|              | 97:8:2:2        | 0.4±0.1              | 26±1    |
|              | 96:6:3:7        | 0.4±0.1              | 71±2    |
| DPPC:DPPPIP(4,5)P2 |         |                      |         |
|              | 96:6:3:7        | 1.9±0.3              | 73±2    |
| PC:PS:PI(4,5)P2 (+chol) |          |                      |         |

Component a Component b Component a Component b

| 69:7:29:1.3 | 0.04±0.01 | >5       | 22±1 | >175 |
| 68:8:26:2.6 | 0.04±0.01 | >5       | 29±12| >175 |
| 97:9:2:1.3  | 2.4±0.2   |          | 145±23|
| 96:4:1:8:1:8 | 1.0±0.1 |          | 93±35 |

Dissociation constants, \(K_d\) in \(\mu\)M for the binding, at room temperature, of wt PTEN to stBLMs prepared by RSE on SAMS composed of HC18:ME 70:30 from lipid solutions of the compositions shown. The lipid chain compositions are dioleoyl (DOPC, DOP3) and that of the natural brain extract for PI(4,5)P2 (mostly stearoyl-arachidonoyl), with the exception of the sample denoted as DPPC:DPPPIP(4,5)P2, for which the chain composition on both stBLM components is dipalmityl. With the exception of the system denoted as PC:PS:PI(4,5)P2 (+chol) which was clearly bimodal, all experiments were evaluated as single component fits.

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protein concentrations, as the protein aggregated above \(c_{PAN}\sim 3\) \(\mu\)M. \(K_d\) values between 2.5 and 4.9 \(\mu\)M are consistent with the partial binding isotherms that we obtained.

The four PTEN proteins included in this study have highly distinct binding characteristics. The H93R PTEN mutant shows the weakest binding affinity to membranes that contain PI(4,5)P2 as the sole anionic component (\(K_d\) = 1.3±0.2 \(\mu\)M for an stBLM with 2.2 mol% PI(4,5)P2). Although \(K_d\) is a factor of 3 smaller for PI(4,5)P2 than PS, \(B_{max}\), and therefore the amount of bound protein, is \(\sim 3\) larger for PS-containing membranes than for PI(4,5)P2-containing membranes. The truncated PTEN mutant shows a two-fold stronger affinity and wt PTEN shows a four-fold stronger affinity to PI(4,5)P2 with \(K_d\) = 0.77±0.07 \(\mu\)M and \(K_d\) = 0.4±0.1 \(\mu\)M, see Tables 1 and 2. A comparison of the affinities of wt and mutant PTEN shows that the wt protein has by far the lowest affinity to PS alone. While its affinity to PI(4,5)P2 is similar to that of the mutant proteins, its affinity to (PS+PI(4,5)P2) surprisingly is more than an order of magnitude greater than that of any of the mutants for which this lipid composition was studied. This indicates a remarkable synergy in binding of wt PTEN to (PS+PI(4,5)P2), which the mutants apparently lack.

To investigate the stoichiometry of PI(4,5)P2 binding to PTEN, we also studied the binding of wt PTEN to DPPC+DPPPIP(4,5)P2. Because we used lipids with saturated dipalmitoyl chains, the bilayer is expected to be in the gel phase at room temperature (DPPC has a melting point, \(T_m\sim 41^\circ\)C). As a result, the diffusivity of PI(4,5)P2 within the bilayer is low and prevents active recruitment of multiple PI(4,5)P2 molecules by PTEN, for example by electrostatic attraction. We observe that the binding affinity is approximately five-fold less (\(K_d\) = 1.9±0.3 \(\mu\)M) to a membrane with saturated lipids compared to that with unsaturated lipids with the same headgroup composition. The fact that the \(K_d\) values are distinct demonstrates the importance of bilayer fluidity in the binding, and argues against a 1:1 binding stoichiometry between PTEN and PI(4,5)P2 [49].

Catalytic activity requires PTEN binding to PI(3,4,5)P3, dephosphorylation and production of PI(4,5)P2. We investigated the binding of wt PTEN and catalytically dysfunctional C124S PTEN to stBLMs composed of PC and PI(3,4,5)P3. The affinity of wt PTEN is six-fold larger to membranes that contain PI(4,5)P2 than to those containing PI(3,4,5)P3 (\(K_d\) = 0.4±0.1 \(\mu\)M; \(K_d\) for binding to PI(4,5)P2-containing membranes is about the same for the wild type and the mutant (\(K_d\) = 0.32 \(\mu\)M vs. \(K_d\) = 0.4 \(\mu\)M). This suggests that PI(3,4,5)P3 attracts the protein with a binding strength comparable to that of PI(4,5)P2, but following hydrolysis, the phosphatase active site binds PI(4,5)P2 less avidly, the PTEN is released more readily and hence, has a lower binding constant.

PTEN binding to dual anionic lipid components in stBLMs with PC. In mammalian cells, both PS and PI(4,5)P2 anionic lipids are implicated in PTEN’s association with the lipid membrane. We prepared DOPC stBLMs containing 28.6 mol% PS and 2.6 mol% PI(4,5)P2. Both the H93R mutant and the truncated PTEN mutant show similar binding affinities with \(K_d\) = 0.7±0.1 \(\mu\)M and \(K_d\) = 0.9±0.2 \(\mu\)M. While \(K_d\) for H93R binding to (PS+PI(4,5)P2) is comparable to that for binding to PI(4,5)P2 alone [50], the amount of bound protein is by a factor of \(\sim 3\) larger. In fact, it is larger than that for binding to PS alone, showing that the presence of both lipids maximizes the localization of PTEN on the membrane. For both H93R and the truncated PTEN mutant, the binding curves are well fitted by a single-component association model. However, the binding curves for wt
PTEN show a strong bimodal signature, and while the higher $K_d$ is only approximate, due to a lack of high-concentration data, the lower one can be precisely determined to be $K_d = 0.04 \pm 0.01 \mu M$. This is an order of magnitude stronger than association to PI(4,5)P₂ alone—which in turn is nearly an order of magnitude stronger than association to PTEN, a continuous change of the single-component model

$$K_d(\text{PI}(4,5)\text{P}_2) = 0.32 \mu M$$

and

$$K_d(\text{PI}(3,4,5)\text{P}_3) = 0.12 \mu M$$

as the relative proportion of the two PIP components is tuned between PI(4,5)P₂ and PI(3,4,5)P₃ (with the overall PIP content in the membrane fixed at 2 mol%, see Table 2). This is consistent with non-competitive binding of the two PIP species, as expected, since the PBD targets PI(4,5)P₂ and the catalytic site in the PD targets PI(3,4,5)P₃.

Summary. In combination, the results presented above suggest distinct essential roles for all three anionic lipid components—studied—PS, PI(4,5)P₂, and PI(3,4,5)P₃—in localizing PTEN to the membrane. At low PTEN concentrations, PI(4,5)P₂ attracts PTEN to the bilayer by virtue of its rather high equilibrium binding constant. Whether PS merely provides a general background of electrostatic attraction or contributes an element of specific binding on its own is not clear; in any case, it leads to high protein coverage of the membrane surface at large ($>1 \mu M$) protein concentrations. Obviously, this suggests experimental conditions for preparing samples well suited for NR, i.e., $B_{\text{max}} > 60 \text{ng/cm}^2$. Despite its strong interaction with PTEN, PI(3,4,5)P₃ alone is insufficient to attract the phosphatase on its own, given its low concentration in the plasma membrane.

Most interestingly, the cooperativity between anionic membrane components enhances binding of PTEN to membranes, as shown by the decrease of $K_d$ values measured for the same lipids in single and dual anionic component membranes. This effect is easily recognized by comparing PTEN binding to PI(4,5)P₂ in membranes with and without PS. However, such cooperativity is not observed between PTEN binding to PI(4,5)P₂ and PI(3,4,5)P₃, or cannot be resolved because the individual $K_d$ values are too similar. Even if the affinity of PTEN to PI(3,4,5)P₃ is slightly higher than to PI(4,5)P₂, a threshold concentration of the latter may be required to attract the phosphatase to its target location, as indicated by the reduction of the affinity to membranes prepared from saturated lipids. It is therefore a pool of PI(4,5)P₂, which distinguishes the composition of the inner plasma membrane from that of other cellular loci, that attracts PTEN to its interaction site.

Neutron Reflection Results

With experimental conditions for the binding of the PTEN mutants to stBLMs optimized by SPR, we began structural characterization of the membrane-associated proteins. NR experiments were performed on wt PTEN bound to (1) DOPC:DOPS = 70:30 (+chol) and (2) DOPC:DOPS:PI(4,5)P₂ = 68:8.28:6.26 (+chol). In addition, we examined the binding of H93R to an stBLM of the same lipid composition as for sample (1). Figure 2 shows the NR data for sample (1). The main panel compares the Fresnel-normalized reactivities of the as-prepared stBLM and the membrane after incubation with the protein in H₂O-based buffer I. Error bars are based on neutron counting statistics and represent 68% confidence intervals. While the differences between the two NR spectra are small, the error-weighted residuals at the bottom of Fig. 2 show that they are significant. Similar spectra were measured for the as-prepared and protein-loaded membrane bathed in buffers based on D₂O and on a mixture of H₂O and D₂O with a nSLD of $\approx 4 \times 10^{-6} \text{ Å}^{-2}$, denoted as “CM4”. The inset displays nSLD profiles with and without protein for H₂O-based buffer I. Error bars are based on neutron counting statistics and represent 68% confidence intervals. While the differences between the two NR spectra are small, the error-weighted residuals at the bottom of Fig. 2 show that they are significant. Similar spectra were measured for the as-prepared and protein-loaded membrane bathed in buffers based on D₂O and on a mixture of H₂O and D₂O with a nSLD of $\approx 4 \times 10^{-6} \text{ Å}^{-2}$, denoted as “CM4”. The inset displays nSLD profiles with and without protein for H₂O-based buffer I. Error bars are based on neutron counting statistics and represent 68% confidence intervals. 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The coloration is the same as in Fig. 3. In addition, we show that in similar studies on the incorporation of lipid headgroups and associated PTEN protein, shown in close-up view in Fig. 4. The signal-to-noise in these measurements is comparable to the magnitude of the experimental errors, at the bottom. Lines in the main panel show the computed NR of the nSLD profiles shown in the inset. Note that these profiles are derived by fitting multiple data sets simultaneously (neat stBLM and stBLM with PTEN, each measured at different isotopic buffer contrasts) by sharing model parameters as appropriate. A dashed box in the inset indicates the region of the distal lipid headgroups and associated PTEN protein, shown in close-up view in Fig. 4. The signal-to-noise in these measurements is comparable to that in similar studies on the incorporation of α-hemolysin into stBLMs [33] and the binding of the HIV-1 matrix protein to bilayer surfaces [34].

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within experimental uncertainty. As described in detail in a recent publication [39], we resampled the experimental data within their neutron-flux limited errors with a Monte-Carlo algorithm to obtain a family of nSLD profiles (typically, 1000 iterations) that are consistent with the data given the error bars. As an example, Fig. 3 shows the resulting ensemble of nSLD profiles of the interfacial region that encompasses the stBLM and adsorbed PTEN protein for the same system as in Fig. 2. Darker colors indicate more likely values for the nSLD profile, while lighter coloration indicates values with a low likelihood. This depiction provides a visual assessment of the spread in the nSLD distribution that is consistent with the experimental result. In addition, black and red solid lines show the most likely nSLD profiles for the stBLM with and without adsorbed wt PTEN protein, respectively. (For clarity, the spread of nSLD profiles for the as-prepared stBLM is not shown.) The model parameters that give rise to the nSLD ensembles are narrowly distributed about their most likely values. The widths of these parameter distributions were used to determine their confidence limits [39], listed in Table 2 (and in more detail in Information S1).

Discussion

This study serves two major objectives. (1) Map out the contributions of various anionic lipids of the inner plasma membrane to PTEN membrane binding and (2) establish experimental conditions for structural studies of the PTEN/membrane complex in the physiologically relevant, disordered state of lipid bilayers. The stBLM system is well suited both for precise binding affinity determinations via SPR and NR measurements of thermally disordered interfacial structures.

PTEN is a cytosolic protein that associates with membranes only if specific conditions are met [21]. As a result, only a small fraction of the pool of cellular PTEN is bound to the membrane at any time, and that pool is exquisitely controlled with respect to...
Membrane Association of the PTEN Tumor Suppressor

Improvements in Quantitative Analysis of PTEN Membrane Binding

For SPR, carboxymethylated dextran chips are the conventional method to characterize protein binding to membranes and have been used to determine PTEN binding constants [22]. However, $K_D$ values obtained with that technology were by a factor of $\approx 1,000$ smaller than those determined by other methods [35]. Narayan and Lemmon offer an explanation in their discussion of SPR analysis of phosphoinositide binding domains [51]. $K_D$ values are often obtained through a kinetic analysis from the calculation of $k_{on}/k_{off}$ ratios. Such a kinetic analysis of SPR data yielded apparent affinities which were 10–1000-fold higher than those obtained by other equilibrium-based methods (e.g., titration calorimetry or sedimentation analysis) [51]. While the reasons for these discrepancies remain unknown, unspecific protein binding to the carboxymethylated dextran chip, due to incomplete lipid coverage, might be at least part of the problem. In contrast, an SPR saturation analysis of steady state values reached for each protein concentration yielded equilibrium binding values that match the data obtained by alternative methods (for a detailed discussion see ref. [51]). In this work, we adopt a similar approach for the evaluation of $K_D$ values, however, instead of using a carboxymethylated dextran chip with an unknown lipid coverage, we used stBLMs that cover the chip surface with virtually defect-free single bilayers, as determined by NR [26,39]. For the SPR analysis of phosphoinositide binding domains, including PTEN, stBLMs offer significant advantages: First, the combination of SPR and EIS allows us to monitor the quality of the lipid bilayer in situ. Specifically, EIS has the sensitivity of determining the specific resistivity of a bilayer over many orders of magnitude and showed that stBLMs are virtually defect-free [26]. Second, because bilayers within stBLMs are complete (>99%), as shown by EIS and NR [26,33,39] and defect-free, non-specific binding to the chip surface can be ruled out, and therefore SPR measurements do not need to be referenced to a phosphatidylcholine covered chip surface. It was noted that protein binding to PC cannot be determined using the carboxymethylated dextran chips because of the lack of an appropriate reference for such a measurement [51]. In our case, it is possible to obtain binding data for any lipid system, including PC. Third, the long-term stability of lipid-covered carboxymethylated dextran chips is an issue [51], which limits their use to less than 8 hours. In contrast, stBLMs have been shown to be stable for months [52]. An additional advantage that stBLMs offer in comparison to carboxymethylated dextran chips is the cross-platform utility of the model system. stBLMs can be used to characterize PTEN/membrane interactions using fluorescence temporal and spatial distributions. PTEN’s targeting and activation requires binding to a distinct ensemble of lipids, and the level of binding is a major determinant of its activity. Our well-controlled model membrane systems allow comparison of binding of wt protein with that of mutant proteins to gain insights into the role of membrane composition in PTEN regulation. The methodology used to quantify PTEN binding in this work offers significant improvements that derive from the stBLM sample format. These improvements include continuous, virtually defect-free single bilayer interfaces that permit quantitative measurements without the need for reference surfaces; suppression of unspecific protein binding; long-term stability; and the compatibility of the sample format with multiple complementary characterization techniques such as electrochemical impedance spectroscopy, fluorescence correlation, fluorescence microscopy, and reflectivity measurements, harnessed here to obtain a first glimpse on structure.

Figure 4. Comparison of nSLD profiles of PTEN Interfacially Bound to Lipid Bilayers and Their Correspondence with the Crystal Structure of the Truncated PTEN. The protein contributions (d = 60–120 Å from the Au surface) to the overall nSLD profile correspond to wt PTEN adsorbed to stBLMs (A) with 30 mol% DOPS (+chol), (B) with 28.6 mol% DOPS and 2.6 mol% P(4,5)P 2 (+chol), and to H93R PTEN adsorbed to an stBLM with 30 mol% DOPS (+chol). The peak near $d = 50$ Å originates from the lipid headgroups. The color coding is the same as in Fig. 3, and panel (A) is a close-up of the results shown there. Dashes black lines represent the best-fit overlay of a scaled nSLD distribution calculated from the crystal structure of the truncated PTEN variant. Because different amounts of protein was adsorbed in the individual experiments, the ratios between the protein and lipid headgroup peaks varies from sample to sample. doi:10.1371/journal.pone.0032591.g004
techniques (e.g., imaging, FCS or single molecule measurements) or to study membrane structure by NR, as described in this paper.

**PTEN Affinities to Membranes of Different Lipid Compositions**

It was recently shown that PTEN requires a distinct lipid environment for membrane association and activation. PTEN binds much more strongly to bilayers containing PS and PI(4,5)P₂ than to bilayers that contain only one of these components [35]. From these data, we argued that the binding of PTEN to PI(4,5)P₂ and PS is synergistic rather than competitive. The highly quantitative data presented here dramatically underscore this point (see Tables 1 and 2). For example, we find for wt PTEN that the affinity of the protein to bilayers with both lipids is an order of magnitude larger than the affinity to PI(4,5)P₂ alone, and more than 2 orders of magnitude larger than to PS alone. Finally, this very high affinity binding was not observed for the H93R and truncated PTEN proteins, suggesting that this synergy requires multiple PTEN domains.

We also assessed the binding of PTEN to PI(3,4,5)P₃-containing membrane. wt PTEN apparently shows weaker binding to PI(3,4,5)P₃-bearng membranes than to PI(4,5)P₂, both with the same PC bilayer background. Surprisingly, the binding is slightly lower for the ternary (PC+PI(4,5)P₂+PI(3,4,5)P₃) system than for the binary (PC+PI(4,5)P₂) system. We can only speculate that structural adjustments imposed by PI(3,4,5)P₃ on the protein that may prevent more effective binding. It is also likely that some of the PI(3,4,5)P₃ substrate is converted to PI(4,5)P₂ during the experiment, which prevents a clean analysis of the binding data. Because of this latter complication, we complemented the wt PTEN measurements with binding studies of C124S PTEN where the mutation, located in the catalytic binding pocket abolishes the protein's hydrolytic activity. Indeed, for C124S PTEN, we find that the binding to PI(4,5)P₂ is similar to that of wt PTEN, i.e., the mutation in the binding pocket does not affect the interaction of PI(4,5)P₂ with the protein. However, the association of C124S PTEN with the PI(3,4,5)P₃-containing stBLMs is significantly stronger than that of wt PTEN. There are two possible explanations. First, wt PTEN has an apparently weaker binding for PI(3,4,5)P₃ because it converts it to PI(4,5)P₂, which binds only weakly to the active site. Second, the C124S mutation may affect the PTEN-PI(3,4,5)P₃ interactions at the active site. While an amino acid switch from Cys to Ser is neutral with respect to the headgroup. PI(4,5)P₂ lacks the phosphate group in the 3-position, which explains why its binding is not affected by the mutation. It is also interesting that the binding of wt PTEN to PI(3,4,5)P₃-containing membranes is significantly weaker than its binding to PI(4,5)P₂. Binding studies with peptides derived from PTEN's N-terminus, which contains a PI(4,5)P₂ binding site, showed that PI(3,4,5)P₃ binding to PTEN's N-terminal end is minimal [35], suggesting that the observed PI(3,4,5)P₃/PTEN interaction is largely at the active site. However, this binding is weaker than PTEN binding to PI(4,5)P₂, even though the concentrations of the anionic lipids were the same in these experiments. We reported previously that the interaction with PI(4,5)P₂ leads to an allosteric activation of PTEN [35,53] that is not observed in the presence of PI(3,4,5)P₃. Taken together, these observations suggest that PI(4,5)P₂ binds to the N-terminal end of the protein, which augments the affinity of the protein to the membrane, possibly by locking in the bound state by a conformational change, and that this interaction does not occur with PI(3,4,5)P₃. Therefore, as confirmed by the data presented here, the cellular concentration of PI(3,4,5)P₃ is too low to attract PTEN to the plasma membrane in appreciable amounts.

The H93R and C124S PTEN mutants showed approximately 4-fold stronger binding to PS-bearing membranes compared with wt PTEN (Table 2). We have previously suggested that the H93R mutant might bind PS more avidly because this mutation may affect the PS-binding to the C2 domain [38]. Another possibility, however, is that these mutations enhance binding of the phosphatase active site to PS. Since PS has a smaller headgroup than PI(3,4,5)P₃, it will fit into the active site. However, because it has fewer negative charges and less hydrogen bonding capability, PS would likely have a lower affinity than PI(3,4,5)P₃. Although speculative at this point, the stronger PS-binding of the mutant proteins in comparison with wt PTEN might be due to the fact that PS enters the substrate binding pocket, with the implication that stronger hydrogen bonding (C124S) and stronger ionic interactions as well as hydrogen bonding (H93R) increases the membrane affinity of these mutants.

To assess the role of membrane dynamics in protein binding, we compared the affinities of wt PTEN to stBLMs formed from phospholipids with different fatty acid constitutions under otherwise identical conditions (Table 1). For the membrane compositions tested, the results show convincingly that the binding affinity is significantly lower, by a factor of ~5, for protein binding to DPPC/DPP(4,5)P₂ bilayers with low in-plane fluidity than to bilayers with unsaturated chains and, therefore, higher in-plane fluidity. The B_max values for these two situations agree within experimental error, as expected for protein binding in a one-to-one stoichiometry to PI(4,5)P₂. However, the observed B_max values correspond to protein number densities at the membrane surface that are much lower than the number density of PI(4,5)P₂. This suggests that either there is not a simple one-to-one stoichiometry in the binding of PTEN to PI(4,5)P₂ [49] or that local accumulation of PI(4,5)P₂ in the bilayer, for example driven by electrostatic attraction to adsorbed PTEN molecules, augments the k_off rate of protein binding by increasing the probability for a stable PTEN-PI(4,5)P₂ interaction in a one-to-one stoichiometry. The latter reasoning follows an argument made by McLaughlin and coworkers to rationalize a similar increase of binding affinity of MARCKS to PI(4,5)P₂-containing membranes [54] upon membrane fluidization which they attributed to the lateral sequestration through electrostatic interactions of the lipid with protein upon binding. In any case, these results suggest that lateral accumulation of PI(4,5)P₂, possibly in the form of protein-induced phase-separation, may play a vital role in the binding of the enzyme to the membrane surface. Clearly, more experiments are needed to clarify these details of the binding process, but the binding preference for fluid membranes may very well play an important role for the spatial control of membrane binding in vivo, as biomembranes are thought to be mosaics of patches with higher and lower fluidity [55].

Studies of the truncated PTEN variant were primarily undertaken to optimize sample preparation for NR experiments, where the crystal structure [24] will facilitate interpretation of the results. Because of multiple distinctions between wt and truncated PTEN, differences in their binding properties are difficult to interpret. In addition, the truncated PTEN has a propensity to aggregate in solution, which precluded binding studies at high protein concentrations. The interaction of the truncated PTEN with PS-containing membranes is more than a factor of two larger than that of the wt protein, which could be related to the clipping
of the C-terminal tail, 50 AAs in length, which is structurally ill-defined and may tend to obstruct the placement of the wt protein at the membrane. Also, the clipped portions of the protein bear a net charge of $-14$, and therefore their removal in the truncated protein may considerably decrease electrostatic repulsion from the negatively charged membrane surface. Conversely, the association of the truncated PTEN with membranes that contain PI(4,5)P$_2$ is somewhat reduced from that observed for the wt protein to the same membrane compositions, most likely because of the truncation of the N-terminal segment. Note however, that a key element of the PBM, K13 [35], is present on the truncated PTEN, which lacks only the first 6 AAs [24]. In terms of the amount of bound protein, all of the three membrane compositions studied with the truncated PTEN should be suitable for NR studies, as the $B_{\text{max}}$ values comfortably exceed 60 ng/cm$^2$. The main concern for such studies is therefore to have the protein bind in sufficiently high density at the membrane to observe structural differences, while avoiding protein aggregation.

### PTEN Structure on Membranes

Figure 2 shows that the adsorption of wt PTEN to stBLMs leads to significant changes of NR from interface, which can be quantified (Figs. 3 and 4) in terms of contributions of the proteins to the nSLD profiles. It is immediately clear from the protein nSLD distributions that PTEN associates superficially with the membrane and forms monomolecular layers in all cases studied, since the extension of the nSLD increment between the bilayer surface and the bulk buffer corresponds well with the protein size determined in the crystal structure. As quantified in Table 3 and schematically depicted in Fig. 5, any penetration of amino acid sidechains is limited to a few Ångstroms at most. This suggests that the phosphatase goes about its business by scooting peripherally on the membrane surface without penetrating the lipid headgroups. For the interaction of the phosphatase with its substrate, this implies that access of the inositol ring to the PD binding pocket is remarkably little change due to protein association.

Both experiments with wt PTEN–adsorption to an stBLM that contains only PC and PS and to an stBLM that also contains 2.6 mol% PI(4,5)P$_2$ show almost identical nSLD distributions [56]. If there is any distinction, then the protein hugs the membrane more closely on the PS-only bilayer and also, the nSLD distributions at $d>100$ Å appear slightly different. In both cases, the correspondence of the free-form fit of the experimental data with the nSLD distribution derived from the crystal structure of the truncated PTEN is striking. The major portion of the protein nSLD near the bilayer surface is almost perfectly described by the crystal structure, while there are ~15-20% of the nSLD, distally located from the membrane surface, unaccounted for in the experimental nSLD distributions. We draw the following conclusions for the structural interpretation of these results: (1) Because the putative nSLD distribution in the crystal structure was determined by assuming the orientation of the protein on the membrane as proposed by Lee and coworkers [24], and different orientations would result in significantly different nSLD distributions, the close correspondence suggests strongly that the C2 and phosphatase domains are organized as determined in the crystal structure and are both in close association with the bilayer of the stBLM. The resulting membrane positioning consistent with the nSLD profiles is schematically shown for wt and H93R PTEN in Figs. 5A and B, respectively. (2) FTIR characterization of dissolved and membrane-bound wt PTEN showed changes in the secondary structure upon membrane binding [35]. Our NR data suggest that these structural changes are indeed minor if they affect the C2 and phosphatase domains or that they occur in those parts of the protein that were clipped in the crystal structure, for example, on the ~50 AA long C-terminal tail. (3) Given the close correspondence between the protein region proximal to the bilayer and the crystal structure, it is tempting to assume that the major portion of the nSLD which is unaccounted for in the distal region is due to the truncated stretches on the crystal structure. While the truncation on the N-terminal tail is short and the clipped stretch within the C2 domain is fixed in its location on the protein, the long stretch of C-terminal tail is not similarly confined and could be rather mobile. We propose that the extra nSLD in the experimental $\rho_n$ profile is associated with this C-terminal stretch which points away from the membrane-bound protein with an overall conformation that is still in close vicinity to the two major protein domains, C2 and PD (Fig. 5). While the sheer length of the
C-terminal tail would permit it to reach the membrane surface and interact with the lipids, the observed nSLD profile suggests that this is not the case. However, it will require further efforts to establish the conformation of the C-terminal tail more rigorously, for example with NR experiments using wt PTEN protein with the C-terminus specifically deuterated.

The analysis of the reflectivity results is somewhat preliminary because we haven’t yet had the opportunity to conduct NR investigations with varying protein concentrations or studies using specific deuterium. In particular, experiments with deuterated PTEN will unmistakably show the spatial organization of, for example, the C-terminal protein stretch. The current data refinement yields envelopes of nSLD for PTEN, without any direct reference to the protein’s internal structure. Because the nSLD distribution of the truncated PTEN is not pronouncedly asymmetric about its peak density, there is the possibility that the bound protein is not unidirectionally oriented at the membrane surface. The two orientations would likely have different distances from the membrane surface, and one would expect a significantly broadened distribution from that predicted by the crystal structure, which we do not observe.

As functionality assays and our SPR results show, the point mutation in H93R PTEN affects its catalytic affinity and membrane binding to PS- and (PS+PIP2)-containing membranes to a surprisingly high extent, given that it differs from the wt protein only by one single AA. It may therefore be less surprising that the H93R mutant also shows a significantly different nSLD distribution at the membrane surface. There are several reasons that could lead to these differences. In line with our interpretation of the wt PTEN nSLD distributions discussed above, the lack of nSLD in the distal region of the H93R protein may reflect a distinct organization of the C-terminal tail. As Fig. 4c shows, the distribution at the membrane surface. There are several reasons that could lead to these differences. In line with our interpretation of the wt PTEN nSLD distributions discussed above, the lack of nSLD in the distal region of the H93R protein may reflect a distinct organization of the C-terminal tail. As Fig. 4c shows, the extension of H93R PTEN at the membrane matches the overall width of the nSLD distribution derived for the truncated PTEN crystal structure quite well—better, in fact, than the nSLD distributions of wt PTEN. On the other hand, the detailed profile of the truncated PTEN is not well matched by the H93R PTEN nSLD distribution. This would be expected if the C-terminal tail, which provides the bulk of nSLD unaccounted for in the crystal structure, is located at the same distance from the membrane surface as the major domains of the membrane-bound PTEN, e.g., at d~90 Å, where as it is further away from the membrane for the wt protein, at d>100 Å. In any case, without further information about the conformation of this part of the protein, for example from NR investigations of specifically deuterated PTEN, such structural details cannot be resolved with confidence.

While the differences between the structures of wt and H93R PTEN on the membrane deserve further investigation, we summarize our NR results as showing conclusively that the membrane association of the PTEN phosphatase scoots along the membrane in search of its PI(3,4,5)P3 substrate molecules and most likely pulls their headgroups slightly out of the membrane surface to gain access with its catalytic binding pocket. Electrostatic, and possibly also chemically selective interactions of the C2 domain with PS help pull the phosphatase tight to the membrane surface to augment its interaction with the substrate. The PTEN protein goes about its business without

### Table 3. Parameters of the best-fit model for the neutron reflection from stBLMs with associated PTEN proteins.

| Substrate | wt PTEN on PC:PS = 7:3 (+chol) | wt PTEN on PC:PS:PIP2 = 68.8:28.6:2.6 (+chol) | H93R PTEN on PC:PS = 7:3 (+chol) |
|-----------|---------------------------------|---------------------------------------------|----------------------------------|
| thickness of Au layer / Å | 132.5±2.7 | 157.7±0.3 | 129.5±1.4 |
| nSLD of Au layer / 10^-6 Å^-2 | 4.44±0.01 | 4.47±0.02 | 4.50±0.01 |
| substrate roughness / Å | 4.5±0.5 | 4.3±0.4 | 3.9±0.6 |
| Sparsely-Tethered Bilayer Lipid Membrane | | | |
| area per phospholipid (as-prepared stBLM) / Å² | 66.2±1.8 | 59.8±1.0 | 61.1±0.9 |
| thickness of tether layer / Å | 18.8±0.4 | 16.6±0.4 | 19.1±0.4 |
| thickness of lipid bilayer core / Å | 28.9±0.4 | 29.6±0.5 | 30.1±0.3 |
| thickness of lipid headgroup / Å | 9.6 (fixed) | 9.6 (fixed) | 9.6 (fixed) |
| thermal bilayer roughness / Å | 3.4±0.4 | 5.0±0.3 | 3.0±0.1 |
| PTEN Protein | | | |
| PTEN penetration into lipid bilayer / Å | 4.3±2.7 | 0.1±2.0 | 2.4±3.3 |
| total amount of adsorbed protein in H2O-based buffer (volume per surface area) / Å | 6.8±1.0 | 7.3±0.9 | 3.0±0.8 |
| fraction of isotopically exchanged protons | 0.69±0.13 | 0.68±0.14 | 0.73±0.18 |
| distance of center of mass of protein from bilayer interface / Å | 23.9±1.9 | 29.6±1.4 | 25.1±5.2 |
| Global Properties | | | |
| quality of best-fit (χ²) | 1.68 | 2.86 | 2.22 |

Error limits indicate 68% confidence intervals derived from Monte-Carlo data resampling of the data [39].

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penetrating the lipid headgroups in the bilayer. In doing so, the two major domains of the protein sit flush on the membrane surface and the regulatory C-terminal tail is most likely pushed out of the way, located on the distal side of the protein core.

This work provides a wealth of new information about the membrane association and likely mode of action of the PTEN phosphoinositide phosphatase. We show that binding of the enzyme to the membrane surface is synergistically affected by two membrane components, PS and PI(4,5)P2. Membrane fluidity is an important factor in determining the protein’s affinity to the membrane, which we suggest is due to the binding of multiple PI(4,5)P2 lipids to the protein which actively recruits them through attractive interactions. The mutations C124S and H93R both affect the PTEN membrane affinity significantly. For C124S that can be rationalized by the fact that this mutant is incapacitated in its catalytic activity. The reasons for the rather dramatic effects documented for H93R, which binds with much higher affinity to PS but shows a reduction of its affinity to PI(4,5)P2, are less clear.

NR studies of the PTEN phosphatase on the in-plane fluid, thermally disordered bilayer membrane provide a first glimpse into the structural details of an active phosphoinositide phosphatase at the membrane surface and start to fill in an emerging picture of the structure-function relationship. We observe that PTEN scoots along the membrane superficially without penetration the lipid headgroup. Because of the lack of membrane penetration, the phosphatase probably does not even reach the position of the lipid headgroup phosphates while surfing the membrane. In view of the depth of its catalytic binding pocket, it is likely that the substrate is somehow pulled out of the membrane, presumably through electrostatic interactions, to promote access to the mostitol ring. Our current interpretation of the overall nSLD distribution of the protein at the interface is that the regulatory C-terminal tail is pushed away from the bilayer to the distal region of the membrane-bound protein.

Supporting Information

Information S1 The Information S1 provides a schematic structure of the tether lipid, HC18; the procedure used to calibrate the custom-made SPR instrument; the procedure used to estimate Ka ranges from insufficient SPR data sets; and the full sets of neutron reflectivity data measured for the various systems that have been investigated, as well as the derived nSLD profiles. A full set of model parameters used to describe the neutron reflectivity data sets is also given.

(PDF)

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Author Contributions

Conceived and designed the experiments: ML AHR. Performed the experiments: SS PS FH M-CD. Analyzed the data: SS PS FH AG AHR ML. Contributed reagents/materials/analysis tools: M-CD AHR. Wrote the paper: ML AHR AG SS FH.

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48. The ratios of integrated nSLD contributions of lipid headgroups and of protein—and hence also the ratios of the nSLD peaks near d = 50 Å and 75 Å—vary from experiment to experiment due to different amounts of adsorbed protein (see Table 3) Note also that the amount of adsorbed protein varied in successive NR scans with different isotopic buffers because the buffer exchange removed some protein These variations were accounted for in the model, and thus the nSLD profiles for CM4 or D2O-based buffers show smaller amplitudes in the protein region although the overall shapes are identical to those shown for the H2O-based buffers in Fig 4.