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A surface plasmon resonance-based assay for small molecule inhibitors of human cyclophilin A

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Abstract

A simple protocol for generating a highly stable and active surface plasmon resonance (SPR) sensor surface of recombinant human hexahistidine cyclophilin A (His-CypA) is described. The sensor surface was sensitive and stable enough to allow, for the first time, the screening and ranking of several novel small-molecule (M_r/C_24 250–500 Da) ligands in a competition binding assay with cyclosporin A (CsA). It also allowed us to accurately determine the kinetic rate constants for the interaction between His-CypA and CsA. His-CypA was first captured on a Ni²⁺–nitrilotriacetic acid (NTA) sensor chip and was then briefly covalently stabilized, coupling via primary amines. The significant baseline drift observed due to dissociation of weakly bound His-CypA from the Ni²⁺–NTA moiety was eliminated, resulting in a surface that was stable for at least 36 h. In addition, immobilized protein activity levels were high, typically between 85 and 95%, assayed by the interaction between His-CypA and CsA. The mean equilibrium dissociation constant for CsA (K_d/CsA) binding to the immobilized His-CypA was 23 ± 6 nM, with on and off rates of 0.53 ± 0.1 M⁻¹ s⁻¹ and 1.2 ± 0.1 (×10³) s⁻¹, respectively. These values agree well with the values for the corresponding binding constants determined from steady-state and kinetic fluorescence titrations in solution.
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Keywords: Surface plasmon resonance; Cyclophilin A; Cyclosporin A; Competition binding assay

Surface plasmon resonance (SPR)¹ is now a technique exploited regularly in the determination of equilibrium binding and kinetic rate constants of biomolecular interactions [1–4]. It is also increasingly useful in drug discovery/hit validation studies with small molecules [4–11]. A prerequisite for measuring the binding constants of an interaction by SPR is that the surface-immobilized molecule must be both stably attached and highly active. Most immobilization procedures are reliant on direct covalent linkage of a purified protein to chemically activated sensor surfaces. Such procedures generate stable surfaces. However, a significant proportion of proteins are not compatible with either the solution conditions or the surface chemistry used. As a result, many biomolecules have very low activity, or often become completely inactive, on immobilization. Over and above the loss of activity due to modification of critical residues involved in binding sites, the essentially random orientation resulting from

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Abbreviations used: SPR, surface plasmon resonance; NTA, nitrilotriacetic acid; His-CypA, hexahistidine cyclophilin A; CypA, cyclophilin A; PPIase, peptidyl-prolyl isomerase; CsA, cyclosporin A; SA, streptavidin; PCR, polymerase chain reaction; IPTG, isopropylthio-galactoside; DTT, dithiothreitol; EDTA, ethylenediamine tetraacetic acid; PMSF, phenylmethanesulfonyl fluoride; EDC, 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide hydrochloride; NHS, N-hydroxysuccinimide; RU, response units; DMSO, dimethyl sulfoxide; ITC, isothermal titration calorimetry; BSA, bovine serum albumin.
covalent coupling leads to subpopulations of immobilized molecules that are incorrectly oriented for binding. There are a number of capture methods available (e.g., antibodies, biotinylation, oligohistidine tags) that can be used as alternatives to direct covalent coupling. However, the sensor surface often has a relatively low binding capacity and/or exhibits significant baseline drift due to the relatively weakly bound captured molecule dissociating from the surface. For example, single hexahistidine-tagged proteins captured on Ni$^{2+}$–nitrilotriacetic acid (NTA) sensor surfaces have $K_d$ values in the low micromolar range [12]. Such leaching makes it difficult to assess binding kinetics accurately.

Here, we describe the generation of a highly stable and active sensor surface of an N-terminally tagged hexahistidine cyclophilin A (His-CypA), which has enabled a series of ligand binding studies with nonpeptide small molecules to be performed. The cyclophilins [13] have emerged recently as a potential drug target for a several diseases, including HIV and malaria infection [14,15]. The SPR-based binding assay described here provides the basis of a screen for novel small-molecule cyclophilin inhibitors of potential therapeutic interest. The in vivo role of human cyclophilin A (CypA) is poorly understood, but its ability to enhance the rate of folding (or unfolding) of proteins via its peptidyl-prolyl isomerase (PPIase) activity is likely to be important, particularly when the cell is stressed [13,16]. CypA was isolated as a complex with the immunosuppressive cyclic undecapeptide cyclosporin A (CsA) [17], which is used as a therapy for preventing organ rejection following transplants. CsA acts by complex formation with cytosolic CypA, followed by binding and inhibition of the phosphatase calcineurin, blocking the signal transduction pathway for immunostimulation [18]. Previous work has identified many cyclosporin derivatives as CypA inhibitors [19–25], but there are very few published examples that characterize nonpeptide inhibitor binding [26,27].

The development of an SPR-based assay for screening small-molecule ligands requires a stable and sensitive CypA sensor surface. Human CypA is not readily amenable to direct covalent immobilization on a CM5 sensor, or on a streptavidin (SA) sensor, using standard coupling chemistries. In our hands, these coupling protocols generated surfaces with very low levels of protein activity, typically only approximately 5%. Surfaces with such low levels of activity are neither active nor sensitive enough to facilitate the detection of small molecules. Although interactions between CypA, directly coupled to the sensor surface, and protein binding partners analyzed by SPR have been published [28–30], the activity of immobilized protein and the sensitivity of the sensor required to detect binding of these much larger molecules are significantly less than those needed to detect small-molecule ligands. In contrast, when His-CypA was first captured and oriented via its N-terminal hexahistidine tag on an NTA sensor surface and then was briefly covalently stabilized, using standard chemistries to activate the surface and couple via primary amines, the baseline drift was completely eliminated and the surface activity levels typically were in excess of 85%. These sensor surfaces allowed us, for the first time, to assess and rank the equilibrium dissociation constants ($K_d$) for several new small-molecule ($\sim$300–500 Da) inhibitors of CypA.

Materials and methods

Materials

All chemicals used were of the highest grade available commercially.

Plasmid construction

The plasmids for expression of recombinant human CypA were created by polymerase chain reaction (PCR) using a whole tissue human lung cDNA library (Stratagene) as a template, with 5’-CCATGGTCAAC CCCACCGTGTTC-3’ as the forward primer and 5’-GGATCCTTTATTTCCAGGTTCACACAGTC-3’ as the reverse primer. The resulting PCR product was verified by sequencing in both directions, using ABI PRISM BigDye v3 Terminator Cycle Sequencing Ready Reaction Kit and an ABI PRISM 310 genetic analyzer (Applied Biosystems). For generation of the untagged CypA expression vector, pSW3-001, the PCR product was digested with NcoI and BamHI (New England Biolabs) and then ligated into a pET-5a vector (Novagen) digested with NcoI and BamHI. For generation of the N-terminal His-CypA expression vector, pSW3-002, the NcoI and BamHI digested PCR product was ligated into a pET-15b vector (Novagen) similarly digested. Correct insertion of the coding region of CypA was verified by restriction digest and by sequencing the entire coding region in both directions.

Protein expression and purification

All purification was performed on ÄKTA Prime (Pharmacia) equipment at 4 °C.

His-CypA purification

Recombinant human His-CypA was expressed and purified to homogeneity from BL21 (DE3) Escherichia coli (Novagen). LB media containing carbenicillin (100 µg ml$^{-1}$) were grown shaking (260 rpm) at 37 °C until the $A_{600nm}$ was approximately 0.6. Overexpression of His-CypA was induced by the addition of isopropyl-thiogalactoside (IPTG) to 1 mM and growth for a further 3 h at 37 °C. Lysis and purification, on NTA agarose resin (Qiagen), was performed according to...
standard protocols. Fractions containing His-CypA were pooled, concentrated to ≤1 ml, filtered through a 0.2-μm filter, and loaded onto a Sephacryl 200 HR (Pharmacia) gel filtration column (V_{t} ~ 1.6 × 60 cm) preequilibrated in 25 mM Tris (pH 7.5), 100 mM NaCl, 0.5 mM dithiothreitol (DTT), 0.5 mM ethylenediamine tetraacetic acid (EDTA), and 1 mM NaN₃. His-CypA was more than 95% pure as judged by SDS-PAGE.

Untagged CypA purification

Recombinant human CypA was expressed and purified to homogeneity from BL21 Star (DE3) E. coli (Invitrogen). 2 × TY liquid media containing carbenicillin (100 μg ml⁻¹) was grown at 37 °C until the A_{600nm} was approximately 0.6. Overexpression of CypA was induced by the addition of IPTG to 1 mM and growth for was approximately 0.6. Overexpression of CypA was induced by the addition of IPTG to 1 mM and growth for 3 h at 37 °C. Cells were harvested by centrifugation (3000g for 15 min) and washed once in 100 ml of lysis buffer (50 mM Hepes [pH 7.5], 1 mM DTT, 2.5 mM EDTA, 0.5 mM NaN₃). The cell pellet was resuspended at 10% weight per volume in ice-cold lysis buffer plus excess protease inhibitor cocktail (Sigma) and sonicated on ice for 6 × 30-s bursts with 30 s cooling in between. The cell lysate was subjected to centrifugation at 50,000g for 1 h at 4 °C. The high-speed supernatant was dialyzed exhaustively overnight against 50 mM Hepes (pH 6.8), 1 mM DTT, 2.5 mM EDTA, 1 mM NaCl, 100 μM phenylmethanesulfonyl fluoride (PMSF), and 100 μM benzamidine; filtered through a 0.2-μm filter; and applied to an SP Sepharose (Pharmacia) column (V_{t} ~ 50 ml, 2.6 × 10 cm) preequilibrated in the same buffer. Proteins were eluted with a 0- to 400-mM NaCl gradient in the same buffer over 200 ml and were analyzed by SDS-PAGE. Fractions containing CypA, eluting between 130 and 160 mM NaCl, were pooled, concentrated to approximately 1 ml, filtered through a 0.2-μm filter, and loaded onto a Sephacryl 200 HR gel filtration column (V_{t} ~ 120 ml, 1.6 × 60 cm) preequilibrated in 25 mM Tris (pH 7.5), 100 mM NaCl, 0.5 mM DTT, 0.5 mM EDTA, and 1 mM NaN₃. CypA was more than 95% pure as judged by SDS-PAGE.

SPR equipment and reagents

SPR measurements were performed on a Biacore 3000 instrument kindly provided on loan from Biacore. Research-grade CM5, SA, and NTA sensors were used. The reagents 1-ethyl-3-(3-diaminopropyl) carbodimide hydrochloride (EDC) and N-hydroxysuccinimide (NHS) were purchased from Biorad and used according to recommended protocols.

Immobilization and covalent stabilization of His-CypA

Pure His-CypA was immobilized on an NTA sensor chip. The sensor was primed and loaded with Ni²⁺ according to Biacore’s recommended protocols. His-CypA in 10 mM Hepes (pH 7.4), 150 mM NaCl, 0.005% surfactant P20, and 2% ethanol, at concentrations between 200 nM and 1 μM, was passed over the sensor surface at a flow rate of 5 μl min⁻¹. Following saturation of the response units (RU) signal, a 30-s injection (at 15 μl min⁻¹) of a mixture of NHS (115 mg ml⁻¹) and EDC (750 mg ml⁻¹), followed immediately followed by a 30-s injection (at 15 μl min⁻¹) of 1 M ethanolamine (pH 8.5), was performed. The final amount of His-CypA covalently immobilized on the surface was typically between 850 and 1600 RU.

Covalent immobilization of untagged CypA

Pure CypA was immobilized on a CM5 sensor chip. HBS-EP2 buffer (10 mM Hepes [pH 7.4], 150 mM NaCl, 3 mM EDTA, 0.005% surfactant P20, 2% ethanol) was used as running buffer. Activation of the sensor chip surface was performed with a mixture of NHS (115 mg ml⁻¹) and EDC (750 mg ml⁻¹) for 7 min at 5 μl min⁻¹. CypA was diluted with 10 mM acetate buffer (pH 4.9) to a final concentration of between 10 and 50 μg ml⁻¹. The amount of CypA immobilized on the activated surface was controlled by altering the contact time of the protein solution and was between approximately 1000 and 5000 RU. After the immobilization of the protein, a 7-min injection (at 5 μl min⁻¹) of 1 M ethanolamine (pH 8.5) was used to quench excess active succinimide ester groups.

Immobilization of biotinylated CypA

Pure CypA was biotinylated using the EZ Link NHS-LC-Biotin kit (Pierce) according to recommended protocols. The protein was then immobilized on an SA sensor chip, according to Biacore’s recommended protocols, in HBS-EP2 buffer at 5 μl min⁻¹. The final amount of Bio-CypA covalently immobilized on the SA surface was 1811 RU.

SPR binding experiments with CsA

SPR binding experiments with CsA were performed in HBS-EP2 buffer as the running buffer. The flow rate was 50 μl min⁻¹ in all experiments. The sensor surface was regenerated between experiments by dissociating any formed complex in HBS-EP2 buffer for 30 min, followed by a further 30-min stabilization period. CypA is not amenable to more stringent and rapid regeneration conditions (data not shown). Furthermore, CsA is a hydrophobic molecule with limited solubility in aqueous solutions and has a tendency to adhere to tubing and tips at concentrations greater than 1.2 μM. Ethanol (2%) was used in experimental running buffers to help alleviate solubility problems. Ethanol was used in
preference to dimethyl sulfoxide (DMSO) because this binds weakly to CypA [31]. The 60-min total regeneration time between experiments helped to eliminate any carryover of CsA [32]. A concentration series of CsA ranging from 0.5 nM to 1.2 µM was typically run in these experiments. The binding curves were analyzed for a one-to-one Langmuir binding model provided by with the Biacore 3000 instrument software.

**SPR competition binding experiments**

SPR competition experiments were performed in HBS-EP2 buffer. A fixed concentration (25 nM) of CsA in the presence of increasing concentrations of the respective small molecule was passed over a covalently stabilized His-CypA sensor surface for 5 min at 50 µl min⁻¹. The sensor surface was regenerated between experiments by dissociating any formed complex in HBS-EP2 buffer for 30 min, followed by a further 30-min stabilization period. The binding curves were analyzed using the heterogeneous analyte competition model (i.e., CsA and the small molecule compete for the same single binding site on CypA) supplied with the Biacore 3000 instrument software, where the on- and off-rate constants for CsA over the particular surface were predetermined and fixed for the fitting process.

**Intrinsic tryptophan fluorescence binding assays**

CypA possesses a single tryptophan residue (Trp121) that is near (~9 Å) the substrate proline binding pocket of the active site. The structure of CsA bound to CypA shows a strong H-bonded interaction between Trp121 and the carbonyl oxygen of MeLeu9 in CsA [33]. In addition to this interaction contributing significantly to the binding affinity and specificity of CsA for CypA, it accounts for the spectral changes in the fluorescence of Trp121 observed on complex formation [34]. Binding of CsA produces an enhancement of approximately two- to threefold and roughly an 8-nm blue shift in the tryptophan emission maxima (from 350 to 242 nm). Fluorescence emission spectra for CypA and His-CypA were obtained on a PTI Quantmaster spectrofluorometer (Photon Technology International) in a 3-ml cuvette at 25 °C under constant gentle stirring. Tryptophan fluorescence was excited at 295 nm (5 nm bandpass), and emission was measured at 342 nm (5 nm bandpass). CypA (0.2–1 µM) was incubated in the absence or presence of increasing amounts of CsA in 25 mM Tris (pH 7.5), 100 mM NaCl, 0.5 mM DTT, 0.5 mM EDTA, and 1% ethanol. Various concentrations of CsA were added to the CypA mixture, and the solution was mixed and placed in the fluorometer with a total dead time of approximately 3 s. The on- and off-rate constants (k₊CsA and k₋CsA, respectively) were determined by fitting the fluorescence enhancement at 342 nm as a function of time to reaction 1, indicated below, and least squares minimization fitting of the data using version 8.01 of the Berkeley Madonna package. CsA is the CsA concentration and CypA is the concentration of recombinant human CypA:

**Steady-state binding assays**

We assumed that any change in the fluorescence signal at 342 nm is proportional to the concentration of CypA:ligand complex. The observed fluorescence was buffer background subtracted and corrected for dilution and inner filter effects by Eq. (1):

\[
f_{\text{corr}} = \frac{(\text{vol}_0 + \text{vol}_{\text{ads}}/\text{vol}_0) \times f_{\text{obs}}}{e^{-2.303 \times 295 \text{nm} \times L \times [\text{Lig}]}}
\]

(1)

where \(f_{\text{corr}}\) is the corrected fluorescence signal, \(f_{\text{obs}}\) is the observed fluorescence signal, \(e_{295 \text{nm}}\) is the ligand extinction coefficient at 295 nm (CsA, 985 M⁻¹ cm⁻¹), \(L\) is the path length (in this case 0.5 cm), and \([\text{Lig}]\) is the molar ligand concentration. The corrected fluorescence signal, \(f_{\text{corr}}\) (in arbitrary units), on complex formation can be defined as

\[f_{\text{corr}} = f_i + (f_0 - f_i),\]

(2)

where \(f_i\) is the fluorescence of free uncomplexed CypA and \(f_0\) is the fluorescence of the CypA:ligand complex at infinite concentration of ligand. At any total concentration of CypA [CypA], \(f\) depends on the total ligand [Lig] concentration and the dissociation equilibrium constant for the complex (\(K_d\)) according to Eq. (3). The data were least squares fit to Eq. (3) using Kaleidagraph v3.6 software (Synergy Software):

\[
f_{\text{corr}} = f_i + (f_0 - f_i) \times \left\{\frac{(K_d + [\text{CypA}] + [\text{Lig}]) - \sqrt{((K_d + [\text{CypA}] + [\text{Lig}]^2 - (4 \times [\text{CypA}]) \times [\text{Lig}] )/2}}{[\text{Lig}]}\right\}.
\]

(3)

**Determination of kinetic rate constants for CsA binding to CypA**

The interaction of CsA and CypA was followed by monitoring the enhancement in the intrinsic tryptophan fluorescence of CypA (reported in arbitrary units) at 25 °C in a 3-ml cuvette under constant gentle stirring on a PTI QuantaMaster spectrophuorometer, with excitation at 295 nm and emission at 342 nm (5 nm slit width for both). CypA was used at a final concentration of 0.25 µM in 25 mM Tris (pH 7.5), 100 mM NaCl, 0.5 mM DTT, 0.5 mM EDTA, and 1% ethanol. Various concentrations of CsA were added to the CypA mixture, and the solution was mixed and placed in the fluorometer with a total dead time of approximately 3 s. The on- and off-rate constants (k₊CsA and k₋CsA, respectively) were determined by fitting the fluorescence enhancement at 342 nm as a function of time to reaction 1, indicated below, and least squares minimization fitting of the data using version 8.01 of the Berkeley Madonna package. CsA is the CsA concentration and CypA is the concentration of recombinant human CypA:

**Reaction 1**: CsA + CypA ⇌ CsA : CypA \(k_{+\text{CsA}}\); \(k_{-\text{CsA}}\).

**Competition fluorescence experiments**

If a second nonfluorescent ligand, [LigB], that has negligible effect on the fluorescence intensity of CypA
itself competes with CsA. [CsA] for binding Cyp-A, [CypA], then a plot of the corrected fluorescence, \( f_{\text{corr}} \), versus the concentration of competing ligand, in the presence of a fixed amount of [CsA], can be fitted to

\[
f_{\text{corr}} = f_c + (f_b - f_c)/(\{(K_d \times (([\text{LigB}]+K_i))/K_i \\
\times [\text{CypA}]_0\} + 1)
\]

where \( K_d \) is the equilibrium dissociation constant for CsA binding to Cyp-A, [LigB] is the concentration of the competing ligand, \( K_i \) is the equilibrium dissociation constant of the competing ligand and CypA, and [CypA]_0 is the free concentration of CypA at [LigB] = 0.

### Miscellaneous

The molecular weights of CsA, CypA, and His-CypA are 1202.12, 18,012, and 20,307 Da, respectively. Protein concentration was determined by measurement of absorbance at 280 nm and was calculated using the extinction coefficient 8490 M\(^{-1}\) cm\(^{-1}\). The molecular weights of compounds KM19, KM184, KM198, and CD291/02 are 372.5, 412.5, 430.5, and 322.5 Da, respectively.

### Results

#### Generation of a stable NTA-His-CypA sensor surface

Between approximately 1000 and 3200 RU of pure His-CypA could be captured on an NTA sensor surface using recommended protocols (Fig. 1A). However, the affinity of the single hexahistidine tag for the Ni\(^{2+}\)–NTA moiety on the chip is relatively weak (\( K_d \sim 0.5–3 \) \( \mu \)M [12]), and immediately after the injection phase (Fig. 1A, phase a) is halted there is significant and steady baseline drift due to dissociation of the histidine tag from the chip surface (Fig. 1A, phase b). All of the initially immobilized protein could be dissociated from the surface by washing with running buffer for approximately 2 h (Fig. 1A, *). Such a sensor surface is not really stable enough for the determination of binding constants, especially for potentially weakly binding and low-molecular weight ligands.

Complete elimination of protein dissociation was achieved by rapidly following the capture of His-CypA on the NTA surface with a brief covalent stabilization phase (Fig. 1B). After saturation responses were achieved (Fig. 1B, phase a), a 30-s burst of surface activation and coupling (with a mixture of NHS/EDC, Fig. 1B, phase c) was quickly followed by a 30-s injection of ethanolamine (Fig. 1A, phase d) to quench the unreacted succinimide esters remaining on the sensor surface (see Materials and methods). The dissociation of His-CypA from the sensor surface was completely arrested. Even after washing the surface exhaustively (\( \geq 3 \) h) with running buffer, the response level remained constant (Fig. 1B).

Using standard protocols, significant levels (\( \sim 1000–5000 \) RU) of pure CypA and biotinylated CypA could also be covalently immobilized on activated CM5 and SA sensor surfaces, respectively (data not shown) [35] (see Materials and methods). These surfaces exhibited stable response levels with no baseline drift (data not shown).
**Activity of NTA-captured, covalently stabilized His-CypA**

Next, we wanted to assess the activity of the immobilized protein on the sensor surfaces. This was performed by passing saturating concentrations (>0.95 μM) of the naturally occurring tight binding ligand (K\textsubscript{d}/C\textsubscript{24} 10⁻⁴⁰ nM, Table 1) CsA over the sensor surfaces. CsA is a cyclic undecapeptide fungal metabolite with immunosuppressive properties that is widely used in transplant surgery [13]. The surface activity of various sensor surfaces is represented graphically in Fig. 2. His-CypA captured and covalently stabilized on the NTA surface retained very high levels of activity, typically in excess of 85% (Fig. 2 A). In comparison, the immobilized protein on the surface of the CM5 chips retained only approximately 5% activity (Fig. 2 B). Similarly, only 7% activity was retained by biotinylated CypA immobilized on an SA sensor surface (Fig. 2 B). Pure CypA was biotinylated using standard primary amine coupling chemistries before immobilization on the SA chip (see Materials and methods).

We used the covalently stabilized His-CypA sensor surfaces to further characterize the interaction of CypA with CsA. Globally fitting a kinetic model where a 1:1 complex is formed between His-CypA and CsA to data similar to those illustrated in Fig. 3A gave very good fits. The mean on-rate constant (k\textsubscript{+CsA}) was 0.53 ± 0.1 μM⁻¹ s⁻¹, and the mean off-rate constant (k\textsubscript{−CsA}) was 0.012 ± 0.01 s⁻¹, giving an equilibrium dissociation constant (K\textsubscript{dCsA}) of 23 ± 6 nM (Table 1). The inclusion of mass transport considerations had only a very minor effect on the kinetic constants extracted from the data. Very similar values for K\textsubscript{dCsA} were obtained from steady-state response calculations, where the mean K\textsubscript{dCsA} value was 18.5 ± 6 nM (Fig. 3B and Table 1). These values determined by SPR for the equilibrium dissociation constant for CsA binding to His-CypA are in good agreement with those determined from solution fluorescence titration experiments (K\textsubscript{dCsA} = 29 ± 6 nM, Fig. 4) and with values determined by other methods in the literature (for a representative comparison, see Table 1).

Our values for the apparent kinetic rate constants, determined from SPR experiments, also agree well with those determined from modeling the time course of the CsA binding-induced fluorescence enhancement (Fig. 4B and Table 1) and with those recently published for the interaction of the CypD isoform and CsA ana-

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**Table 1**

Representative equilibrium dissociation (K\textsubscript{dCsA}), on-rate constant (k\textsubscript{+CsA}), and off-rate constant (k\textsubscript{−CsA}) for the interaction of CsA with CypA, His-CypA, or CypD, determined from this study and other studies

| Technique      | K\textsubscript{dCsA} (nM) | K\textsubscript{Km} (nM) | k\textsubscript{+CsA} (μM⁻¹ s⁻¹) | k\textsubscript{−CsA} × 10⁻³ (s⁻¹) |
|----------------|-----------------------------|---------------------------|---------------------------------|----------------------------------|
| SPR            |                             |                           |                                 |                                  |
| His-CypA\textsuperscript{a} | 18.5 ± 6                    | 23 ± 6                    | 0.53 ± 0.1                      | 12 ± 1                           |
| CypA\textsuperscript{a}     | 38.5 ± 10.4                 | —                         | —                               | —                                |
| CypD\textsuperscript{a}     | —                           | 12.5 ± 1                  | 0.25 ± 0.1                      | 3.9 ± 1                          |
| TcyPd-hAGT\textsuperscript{e} | —                           | 12.8 ± 1                  | 0.39 ± 0.1                      | 4.8 ± 1                          |
| Tryptophan fluorescence    |                             |                           |                                 |                                  |
| CypA\textsuperscript{a}     | 26 ± 9                      | 16 ± 9                    | 0.45 ± 0.13                     | 7 ± 2                            |
| His-CypA\textsuperscript{a} | 29 ± 13                     | —                         | —                               | —                                |
| CypA\textsuperscript{d}     | 46                          | —                         | —                               | —                                |
| CypD\textsuperscript{d}     | 12.5 ± 4                    | —                         | —                               | —                                |
| ITC             |                             |                           |                                 |                                  |
| CypA\textsuperscript{e}     | 11.4 ± 3.9                  | —                         | —                               | —                                |
| CypA\textsuperscript{f}     | 47.6                        | —                         | —                               | —                                |
| PPIase assay     |                             |                           |                                 |                                  |
| CypA\textsuperscript{a}     | 6.3 ± 3.8                   | —                         | —                               | —                                |
| CypA\textsuperscript{g}     | 1.6 ± 0.4                   | —                         | —                               | —                                |
| CypA\textsuperscript{h}     | 20                          | —                         | —                               | —                                |

Note. K\textsubscript{Km} values were determined from steady-state experiments using the indicated technique. K\textsubscript{dCsA} values were calculated from kinetic assays using the determined on-rate (k\textsubscript{+CsA}) and off-rate (k\textsubscript{−CsA}) constants from SPR experiments or from intrinsic tryptophan fluorescence time course experiments using the formula K\textsubscript{dCsA} = k\textsubscript{−CsA}/k\textsubscript{+CsA}.

\textsuperscript{a} Values (means ± SE, where n ≥ 3) determined from this study.
\textsuperscript{b} [39].
\textsuperscript{c} Values determined by SPR in [32] using CsA and either human CypD or a fusion of O\textsuperscript{6}-alkylguanine-DNA-alkyltransferase and CypD (TCypd-hAGT).
\textsuperscript{d} [43].
\textsuperscript{e} Values determined using ITC in [36].
\textsuperscript{f} Values determined using ITC in [44].
\textsuperscript{g} K\textsubscript{i,app} values determined using a PPIase enzymatic assay performed as described in [45,46].
\textsuperscript{h} K\textsubscript{i,app} values determined using a PPIase enzymatic assay performed as described in [45,47].
lyzed by SPR [32]. In the study by Huber and coworkers [32], the apparent on and off rates for CsA binding to CypD ranged between 0.2 and 0.5 \( \text{M}^{-1} \text{s}^{-1} \) (on rate) and between 0.003 and 0.006 \( \text{s}^{-1} \) (off rate), giving a mean \( K_{dCsA} \) of 12 nM [32] (Table 1).

Our sensor surfaces generated by this method gave responses that were very reproducible. Three independent runs of 25 nM CsA over the same sensor surface 4 h apart gave sensorgrams that are virtually superimposable (Fig. 3C). Locally fitting each curve gave essentially identical on- and off-rate constants (data not shown). In addition to a high level of run reproducibility, the saturation responses with CsA varied minimally over the course of many hours (Fig. 3D). Essentially the same response was observed for \( P_{38} \).

These results indicate that the simple protocol of first capturing His-CypA on a Ni\(^{2+}\)-NTA chip and then briefly covalently stabilizing it, using standard primary amine coupling chemistries, generates a very stable and active sensor surface. They further suggest that in terms of binding affinities and kinetics, His-CypA is interacting with CsA in a manner similar to that in free solution.

**Competition SPR binding assay**

We have previously generated several novel combinatorial libraries of small molecule inhibitors of human CypA (K. Malone, C. Dunsmore, N.J. Turner, unpublished results). We wanted to test whether the His-CypA sensor surfaces described in the above sections could be used to facilitate the primary screening of these small-molecule (\( M_r/24 \sim 300–500 \text{ Da} \)) ligand libraries. Fig. 5A graphically illustrates the equilibrium response signal attained for 120 \( \text{nM} \) of the respective ligands. These compounds were chosen as positive controls to assess the sensor surface because they have previously been shown to bind to CypA with \( K_d \) values in the 10- to 100-\( \text{nM} \) range (Table 2). One problem encountered was that the ligands tested have limited solubility in aqueous buffers, and this led to significant variation in the response signals for repeat runs of the same ligand. In addition, large “spikes” in the RU signal at the beginning and end of the injection phases further hampered reliable direct detection (data not shown). These can likely be attributed to a bulk phase shift given that there are large amounts of insoluble material present during the mobile phase as it passes over the chip surface with protein attached. Ethanol (2%) was added to the running buffer to help ligand solubility for all runs. Higher concentrations of either ethanol or methanol resulted in protein denaturation and rapid loss of activity on the sensor surface on subsequent runs (data not shown). DMSO was avoided as a solvent because this binds to CypA itself, albeit very weakly [31]. Nevertheless, direct binding to His-CypA could be detected for the ligands KM19, KM184, KM198, and C291/02 (Fig. 5A).

We also used a competition binding assay in which a fixed concentration of CsA (25 nM, a concentration near the \( K_{dCsA} \) value, Table 1) was passed over the
sensor surface in the absence or presence of increasing concentrations of the small-molecule inhibitors. Fig. 5B illustrates a representative set of sensorgrams from such experiments. The sensorgrams obtained from such assays were generally less noisy than those obtained in direct binding experiments. Three small molecules (KM184, KM19, and KM198) were tested and competed with CsA for binding to CypA with \( K_d \) values in the micromolar range (Table 2). The rank ordering of the compounds from tightest to weakest binding was as follows: KM184, KM19, and KM198 with \( K_d \) values of 36, 46.9, and 73.6 \( \mu M \), respectively (Table 2). Relatively similar \( K_d \) values, and (more important) the same rank order, were obtained for these three compounds in solution by a fluorescence competition assay (data not shown) (Table 2).

**Discussion**

Covalent coupling of proteins via primary amines is frequently the method of choice for immobilization in SPR assays, generating sensor surfaces with high ligand density and no baseline drift. Our results here illustrate, however, that CypA is not amenable to direct covalent linkage using primary amines. Protein activity levels...
for CypA directly immobilized on CM5 surfaces were only approximately 5%. This activity was unaffected by the initial immobilization levels; immobilization of between 1000 and 5000 RU of untagged CypA gave essentially the same low levels of activity (Fig. 2B). The X-ray structure indicates that the majority of CypA’s 16 primary amines (15 lysine residues and the N terminus) are surface exposed [33] and probably available for direct covalent linkage. However, the surface electrostatic potential of CypA (Fig. 6) illustrates that the front face (the CsA binding face) is strikingly basic, whereas the rear face is much more acidic in character.

Fig. 4. Binding of CsA to CypA in solution. (A) At saturating concentrations of CsA, the intrinsic tryptophan fluorescence emission spectra of CypA shows an enhancement of approximately twofold and a blue shift of approximately 8 nm in the emission maxima, from 350 to 342 nm, compared with protein alone [34]. The steady-state fluorescence enhancement at 342 nm (ΔF342nm) of 0.25 μM CypA is plotted versus the concentration of CsA in nanomolars. Each point is the mean of three separate measurements ±SE. The data were least squares fit (solid black line) to Eq. (3) (see Materials and methods), giving an apparent K_{dCsA} of 25 ± 6 nM (n = 9) for the binding of CsA to CypA. Essentially identical values were obtained with His-CypA (Table 1). (B) The time course of the fluorescence enhancement of 0.25 μM CypA (black lines) on the addition of various concentrations of CsA (indicated) is shown. The red lines are a least squares fit of the data to a kinetic model, as described in Materials and methods. The 3-s dead time for recording the data is indicated, whereas the fitted lines start at t = 0. The mean apparent on and off rates for CsA binding to CypA are 0.45 ± 0.13 μM⁻¹ s⁻¹ and 0.007 ± 0.002 s⁻¹, respectively, giving a K_{dCsA} of 16 ± 8.6 nM (Table 1). a.u., arbitrary units. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

Fig. 5. Binding of small-molecule inhibitors to His-CypA. (A) Graphical representation comparing the mean (from three repeat runs, with error bars showing ±SE) equilibrium response signal for a sensor in contact with 120 μM of the respective small-molecule CypA inhibitors. ATP (120 μM), an unrelated molecule of similar molecular weight (505 Da), is shown as a control for nonspecific interactions with the surface. (B) KM19 competes with CsA for binding to His-CypA. Reference-corrected SPR binding curves (black) for various concentrations of KM19 in the presence of a fixed concentration of CsA (25 nM) were monitored on an NTA surface with 952 RU of covalently stabilized His-CypA. The data were globally fitted (red) using the competing analyte, single binding site model supplied with the Biacore 3000 machine software, where the on- and off-rate constants for CsA were calculated from a concentration series similar to that in Fig. 3A and fixed at 0.49 μM⁻¹ s⁻¹ and 0.01 s⁻¹, respectively (red line over gray sensorgram, K_{dCsA} = 21 nM). The apparent equilibrium dissociation constant (K_d) of KM19 for His-CypA is 46.9 μM, with on and off rates of 6.6 × 10⁻³ μM⁻¹ s⁻¹ and 0.031 s⁻¹, respectively. Curve a, 25 nM CsA alone; curve b, 25 nM CsA + 1.6 μM KM19; curve c, 25 nM CsA + 3.12 μM KM19; curve d, 25 nM CsA + 6.25 μM KM19; curve e, 25 nM CsA + 1.2 μM KM19; curve f, 25 nM CsA + 25 μM KM19; curve g, 25 nM CsA + 62.5 μM KM19. The R_max-CsA for this particular sensor surface was 51 RU. The theoretical maximum RU value for this surface is 56 RU. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
Under the solution conditions (pH 4.9) used during the immobilization process, the surface charge potential of CypA seems likely to force the molecule to orient itself with the CsA binding surface oriented downward toward the activated chip surface. Thus, most of the covalent linkage would occur via residues on the CsA binding surface, resulting in severe steric occlusion of the binding of CsA. The activity levels of CypD immobilized by direct covalent linkage, under similar conditions and with similar final protein immobilization levels, were approximately 45 to 65% [32]. However, CypD does not possess the same markedly polarized surface charge potential of CypA; thus, coupling via primary amines is unlikely to take place predominantly on the CsA binding face.

In addition to these structural/steric considerations, the pH of 4.9 used during the immobilization process, the surface charge potential of CypA seems likely to force the molecule to orient itself with the CsA binding surface oriented downward toward the activated chip surface. Thus, most of the covalent linkage would occur via residues on the CsA binding surface, resulting in severe steric occlusion of the binding of CsA. The activity levels of CypD immobilized by direct covalent linkage, under similar conditions and with similar final protein immobilization levels, were approximately 45 to 65% [32]. However, CypD does not possess the same markedly polarized surface charge potential of CypA; thus, coupling via primary amines is unlikely to take place predominantly on the CsA binding face.

In addition to these structural/steric considerations, the pH of 4.9 used during the initial immobilization phase likely further contributes to the low protein activity on CM5 chips. The \( K_{\text{dCsA}} \) at pH 4.9 was 150 nM ± 34 nM (determined from fluorescence titration experiments [data not shown]), compared with 26 ± 9 nM at pH 7.5 (Fig. 4 and Table 1). Similar acidic pH shift-dependent affinity loss has been observed in isothermal titration calorimetry (ITC) experiments with CypA and CsA [36], attributed to protein protonation effects. The protein recovers only partially; when the pH was raised again to 7.5, the \( K_{\text{dCsA}} \) for the same sample of CypA as above was 57 ± 12 nM (data not shown). It is unclear why protonation would not be reversible other than some partial irreversible denaturation due to the acidic solution conditions that for CypA does occur at a pH below 5 [17,36].

In contrast to these observations, the simple protocol described here for immobilizing and covalently stabilizing His-CypA yields a very stable (\( t_\text{P} \geq 38 \text{ h} \)), highly active (\( \geq 85\% \) activity), and sensitive sensor surface. Using the N-terminal histidine tag to first capture CypA on the NTA sensor surface seems to provide a measure of orientation for the immobilized protein. The 10-amino acid linker between the hexahistidine tag and the start of the CypA sequence likely lifts CypA away from the active chip surface, preventing coupling to random lysine residues that hinder/prevent CsA binding. The only primary amine near enough to the activated sensor surface is the N terminus, and the entropic effects of being immobilized near the surface via the histidine tag may provide specificity for coupling essentially only to this primary amine. Thus, the protein is covalently immobilized but is kept active and sterically competent for CsA binding.

It is interesting to note that the apparent on-rate constants for CsA binding to either CypA or CypD

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### Table 2

| Ligand  | Molecular weight (Da) | \( k_+ \) (M⁻¹ s⁻¹) | \( k_- \) (s⁻¹) | \( K_{\text{dBi}} \) (µM) | \( K_{\text{dFluor}} \) (µM) |
|---------|-----------------------|---------------------|-----------------|------------------------|------------------------|
| KM184   | 412.5                 | 0.58 × 10⁻³         | 0.021           | 36                     | 12.2 ± 9.2 (n = 5)      |
| KM19    | 372.5                 | 0.66 × 10⁻³         | 0.031           | 46.9                   | 15.1 ± 8.2 (n = 3)      |
| KM198   | 430.5                 | 0.53 × 10⁻³         | 0.039           | 73.6                   | 29 ± 21.3 (n = 3)       |

Note. \( K_{\text{dBi}} \) values were calculated from the off- and on-rate constants \( (k_- \) and \( k_+ \), respectively) determined from SPR competition experiments in the presence of 25 nM CsA and using the formula \( K_{\text{dBi}} = k_-/k_+ \). Mean \( K_{\text{dFluor}} \) values were determined from competition fluorescence titration experiments (see Materials and methods) (data not shown).

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Fig. 6. Electrostatic potential surfaces for CypA. The surfaces are viewed from the front CsA binding face (left) and are rotated 180º from the rear face (right). The CsA binding pocket is indicated on the left panel. The electrostatic potential at the surface is color coded from most positive (blue) through to most negative (red).
isoforms determined in this study (~0.5 μM\(^{-1}\) s\(^{-1}\)) and in that by Huber and coworkers (~0.3 μM\(^{-1}\) s\(^{-1}\)) [32] are somewhat slower than a diffusion limited on rate (~10 μM\(^{-1}\) s\(^{-1}\)), as one might have expected for a relatively small molecule such as CsA (M, ~1200 Da). The hydrophobic cyclic undecapeptide CsA is in multistate equilibrium between two extreme conformations influenced by the solution conditions. In aqueous solution, CsA tends more to adopt a compact closed conformation [37], whereas cyclophilins bind tightly only to an open conformation. The slower than diffusion limit on-rate constants might reflect the CsA conformational equilibrium. The off-rate constant for the CypA:CsA interaction appears to be two- to threefold quicker than equilibrium. The off-rate constant for the CypA:CsA interaction might reflect CsA conformational equilibrium. The slower than diffusion limit on-rate constants for these two differentially localized, functionally distinct cyclophilin isoforms. CypA is ubiquitously localized in mammalian cytoplasm [13], whereas CypD is a mitochondrial-targeted PPIase [38]. Nevertheless, the binding constants obtained in this study for the equilibrium dissociation constant for CsA binding to His-CypA by SPR agree well with the values for the same interaction determined by a variety of experimental approaches, including ITC, intrinsic tryptophan fluorescence, and PPIase enzymatic assays (Table 1). This slight difference may reflect genuine differences in the molecular details of the interaction between CsA and these two differentially localized, functionally distinct cyclophilin isoforms. CypA is ubiquitously localized in mammalian cytoplasm [13], whereas CypD is a mitochondrial-targeted PPIase [38]. Nevertheless, the binding constants obtained in this study for the equilibrium dissociation constant for CsA binding to His-CypA by SPR agree well with the values for the same interaction determined by a variety of experimental approaches, including ITC, intrinsic tryptophan fluorescence, and PPIase enzymatic assays (Table 1). This does suggest that the protein molecules immobilized on the chip surface behave in a manner comparable to that of protein molecules in solution, at least in terms of binding to CsA.

The results in this study illustrate the sensitivity of SPR in that we were able to reliably detect the direct binding of small-molecular weight ligands (300–500 Da) (Fig. 5A). However, the on rates for these small-molecule inhibitors are very slow (~0.4–0.7 × 10\(^{-3}\) μM\(^{-1}\) s\(^{-1}\)) (Table 2), orders of magnitude slower than even CsA itself. Although they appear to bind specifically to CypA, there may be some conformational constraint on these small molecules that requires a degree of “induced fit” in terms of their binding mechanism. Structural data of these cyclophilin–ligand complexes are not available, and the bound conformation of the ligands is not known. Poor solubility, aggregation, and/or nonspecific binding of the ligands may also play a role in the calculation of their apparently slow on rate, and it is possible that the functional ligand concentration, competent to interact with the immobilized His-CypA, is less than the expected measured total. This would result in the apparent on rates being faster and the equilibrium dissociation constants being tighter. However, this is unlikely to cause orders of magnitude changes in these kinetic constants. It is reassuring to note that despite the slow calculated on rates, the apparent \(K_d\) values for these ligands and the rank order of their relative affinities determined from the competition experiments agree with the values determined by intrinsic tryptophan fluorescence experiments.

This study is the first to use SPR to analyze small-molecule–CypA binding interactions. Indeed, this article also provides the first published values for the on and off rates of the CypA–CsA interaction. The protocol is simple and yields a sensor with high levels of immobilized protein activity that leads to sensors with a high degree of sensitivity. There are relatively few studies in the literature that report the use of SPR in the study of CypA and its binding partner interactions [28–30,32,39–42]. In early work analyzing the interaction of CypA and CsA and the conformational requirements required for this interaction [39–41], CsA (conjugated to bovine serum albumin [BSA]) and not CypA, was immobilized on the sensor surface [39]. Although interactions between CypA, directly coupled to the sensor surface, and protein binding partners have been published (e.g., HIV-1 capsid [28], HIV-1 Vpr [42], SARS coronavirus [30]), the activity and sensitivity of a sensor surface needed to detect binding in these much larger molecules are significantly less than those required to detect small molecules.

A further advantage of the surfaces generated in this study is that they are very unlikely to exhibit problems with steric hindrance due to the “overcrowding” of molecules on the surface, a potential problem with immobilization of relatively high amounts of protein. The average level of His-CypA finally stabilized on the NTA sensor surfaces in this study was approximately 1000 RU. This corresponds to approximately \(3 \times 10^{10}\) protein molecules within a volume of approximately \(1 \times 10^{14}\) nm\(^3\) and provides an average intermolecular spacing between each CypA molecule of more than 100 Å. This rather sparse but stable binding arrangement may explain the good agreement in the values for the kinetic and equilibrium constants among the SPR data, the solution fluorescence data, and other published data determined by other techniques.

Conclusions

The results in this study clearly indicate that CypA is not amenable to direct covalent coupling to SPR sensors via primary amines. Conversely, our study provides a good general methodology for generating a highly sensitive, stable, and reusable sensor surface of His-CypA that interacts with CsA in a manner essentially indistinguishable from that determined by other experimental approaches. Furthermore, it provides a good methodology that would allow the development of a medium-throughput screen for small molecules ligands/inhibitors of CypA that may provide useful leads in developing new drugs for the treatment of diseases such as HIV and malaria.
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