

Fluorescence Anisotropy-Based Tethering for Discovery of Protein–Protein Interaction Stabilizers

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Cite This: *ACS Chem. Biol.* 2020, 15, 3143–3148

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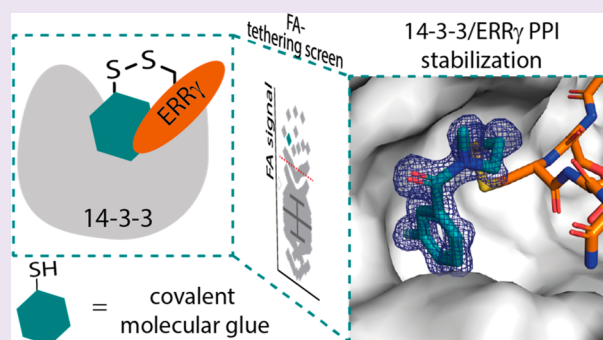
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ABSTRACT: Protein–protein interaction (PPI) networks are fundamental for cellular processes. Small-molecule PPI enhancers have been shown to be powerful tools to fundamentally study PPIs and as starting points for potential new therapeutics. Yet, systematic approaches for their discovery are not widely available, and the design prerequisites of “molecular glues” are poorly understood. Covalent fragment-based screening can identify chemical starting points for these enhancers at specific sites in PPI interfaces. We recently reported a mass spectrometry-based disulfide-trapping (tethering) approach for a cysteine residue in the hub protein 14–3–3, an important regulator of phosphorylated client proteins. Here, we invert the strategy and report the development of a functional read-out for systematic identification of PPI enhancers based on fluorescence anisotropy (FA-tethering) with the reactive handle now on a client-derived peptide. Using the DNA-binding domain of the nuclear receptor Estrogen Related Receptor gamma (ERR γ), we target a native cysteine residue positioned at the 14–3–3 PPI interface and identify several fragments that form a disulfide bond to ERR γ and stabilize the complex up to 5-fold. Crystallography indicates that fragments bind in a pocket comprised of 14–3–3 and the ERR γ phosphopeptide. FA-tethering presents a streamlined methodology to discover molecular glues for protein complexes.



Physical interactions between proteins have increasingly been recognized as an attractive means to manipulate protein function. As such, small-molecule modulators of protein–protein interactions (PPIs) have become highly sought after.¹ By modulating their interactions with regulatory proteins, disease-related proteins that were previously regarded as “undruggable” targets have become tractable for therapeutic intervention.^{2,3} Whereas most successes have been described for PPI inhibitors, great potential remains to be realized for enhancer molecules acting as “molecular glue” to stabilize protein complexes.^{4,5} The discovery of such compounds has proved to be challenging, as apparent from the scarcity of systematic small-molecule PPI enhancers in the literature. Yet, examples of natural product PPI stabilizers^{6,7} and the successes in using heterobifunctional probes for induced protein degradation (PROTACs)⁸ strongly support the value of molecules with a PPI stabilization mode of action and the development of screening technologies for their rational discovery and design.

We are developing systematic PPI stabilization strategies for the family of host/client complexes involving the host protein 14–3–3. These adaptor proteins regulate the activity, cellular localization, and/or the stability of client proteins by binding to phosphorylated motifs in their sequence. The important and widespread chaperone function and the large number of

cellular binding partners places 14–3–3 at critical nodes in disease-related signaling networks.^{9,10}

Recently, we reported a fragment-based discovery strategy based on disulfide trapping (“tethering”)^{11,12} for the identification of molecular glues, resulting in hit molecules that stabilize the 14–3–3/Estrogen Receptor α (ER α) complex while tethered to a cysteine residue on the 14–3–3 host.¹³ Herein, we demonstrate a variant of the tethering approach in which we target a native cysteine residue found in the 14–3–3 binding motif of the Estrogen Related Receptor gamma’s DNA binding domain (ERR γ DBD). In this approach, the ERR γ -phosphopeptide is labeled with a fluorescent probe, and the fragment-induced binding of ERR γ -phosphopeptide to 14–3–3 is monitored by an increase in fluorescence anisotropy (FA; Figure 1). The use of FA-tethering as a functional readout has previously been established in campaigns to identify PPI disruptors for KIX

Received: August 10, 2020

Accepted: November 3, 2020

Published: November 16, 2020



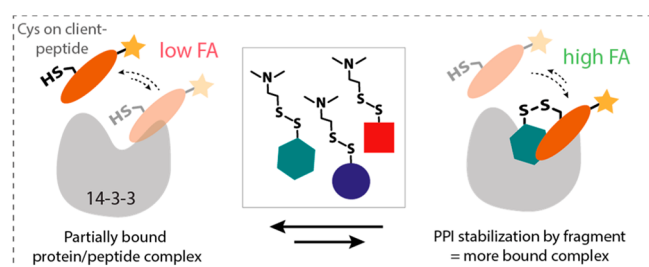


Figure 1. Schematic representation of FA-tethering for PPI stabilizers by fluorescence anisotropy (FA). A fluorescently labeled cysteine-containing client peptide is initially weakly bound to 14-3-3, as evidenced by low FA (low anisotropy). Stabilization of the protein-peptide complex upon incubation with a disulfide library, under reducing conditions, leads to increased 14-3-3 binding to the client peptide, observed by an increase in anisotropy.

and PDK1.^{14,15} The new method, in which the fragment binds to the client peptide to enhance PPI complex formation, contrasts with the previously published screens in which PPI complex formation was inhibited. Taken together, FA-tethering is a rapid, functional assay for directly screening for PPI inhibitors and stabilizers.

NOVEL 14-3-3 INTERACTION MOTIFS IN A NUCLEAR RECEPTOR

Members of the nuclear receptor (NR) superfamily of ligand-responsive transcription factors are master regulators of specific target genes and are implicated in a range of pathological conditions.¹⁶ On the basis of cellular and

proteomics studies, several NRs have been hypothesized to be regulated by 14-3-3.¹⁷⁻¹⁹ Whereas the phosphorylated 14-3-3 binding motif in ER α has been validated in cellular studies, most putative 14-3-3/NR interactions remain unvalidated, including for ERR γ , a relevant drug target in insulin metabolism.^{19,20} The phosphorylation of ERR γ at S179 by protein kinase B (PKB) has been proposed to drive 14-3-3 binding, resulting in decreased nuclear localization¹⁹ and providing endogenous inhibition of ERR γ activity by 14-3-3. Since the molecular mechanism for formation of this protein complex remains to be unraveled, we first focused on the biochemical and structural characterization of the 14-3-3/ERR γ interaction. In addition to the pS179 motif in the ERR γ DBD, a second motif in the ligand-binding domain (LBD) was identified by the 14-3-3 prediction tool developed by the MacKintosh and Barton groups.²¹ Binding of both phosphorylated ERR γ motifs to the seven human 14-3-3 isoforms was confirmed by FA and isothermal calorimetry (ITC) studies (Figure 2a,b, SI Figure S1).

14-3-3/ERR γ PPI: A CYSTEINE IN A DRUGGABLE POCKET

To characterize these two 14-3-3/peptide binding interactions, we obtained highly diffracting cocrystal structures using the 14-3-3 sigma isoform (14-3-3 σ). Crystals of 14-3-3 σ bound by the ERR γ LBD or DBD motif diffracted to 1.9 and 1.4 Å, respectively. The electron density for the ERR γ LBD motif allowed modeling for five of eight peptide residues (SL{pS³¹⁹}FE; SI Figure S2). The electron density for the ERR γ DBD motif allowed seven amino acids of a 9-mer

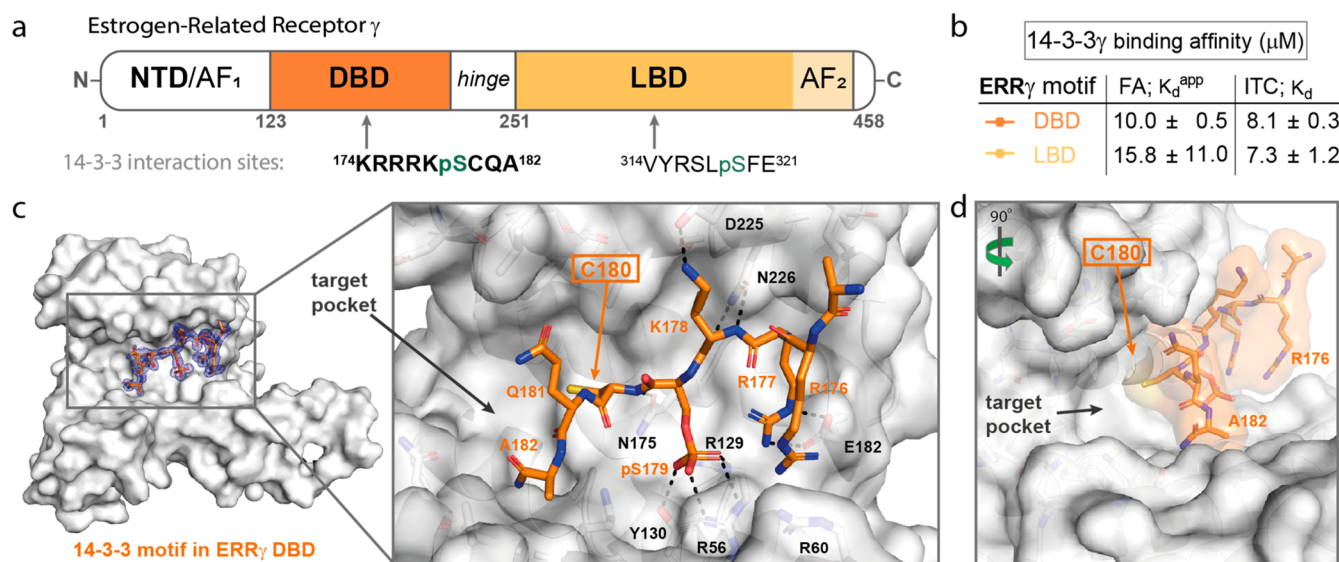


Figure 2. (a) Schematic representation of the ERR γ domain organization, including the N-terminal domain (NTD), DNA-binding domain (DBD), and ligand-binding domain (LBD). AF: activation function. The phosphorylated 14-3-3 interaction motifs studied in this work and their relative positions in the ERR γ protein sequence are indicated. (b) Validation of 14-3-3 binding affinity for the ERR γ DBD and LBD motifs, as determined by fluorescence anisotropy (FA) and isothermal titration calorimetry (ITC). (c) Co-crystal structure of 14-3-3 σ bound by phosphorylated ERR γ DBD motif (PDB: 6Y1D). Front view of a 14-3-3 monomer (white surface) bound by ERR γ phosphopeptide (orange sticks) and close-up view of the peptide in the primary 14-3-3 binding groove (white cartoon and sticks representation). Polar contacts (black dashed lines) were observed between the phospho-group of S179 (ERR γ) and the Arg-Arg-Tyr triad R56, R129, and Y130 (14-3-3). N175 and N226 of 14-3-3 interact with the backbone of ERR γ 's C180 and K178, respectively. Furthermore, one polar contact was observed between D225 (14-3-3) and K178 (ERR γ), and two polar contacts were observed between E182 (14-3-3) and the guanidinium of R177 (ERR γ). The guanidinium groups of R176, R177 (ERR γ), and R60 (14-3-3) form a triple stack. $2F_o - F_c$ electron density maps are contoured at 1σ . (d) Side view of 14-3-3 σ (white surface) bound by ERR γ phosphopeptide (orange sticks with transparent surface) revealing a target pocket for 14-3-3/ERR γ DBD with C180 of the ERR γ peptide motif oriented toward it.

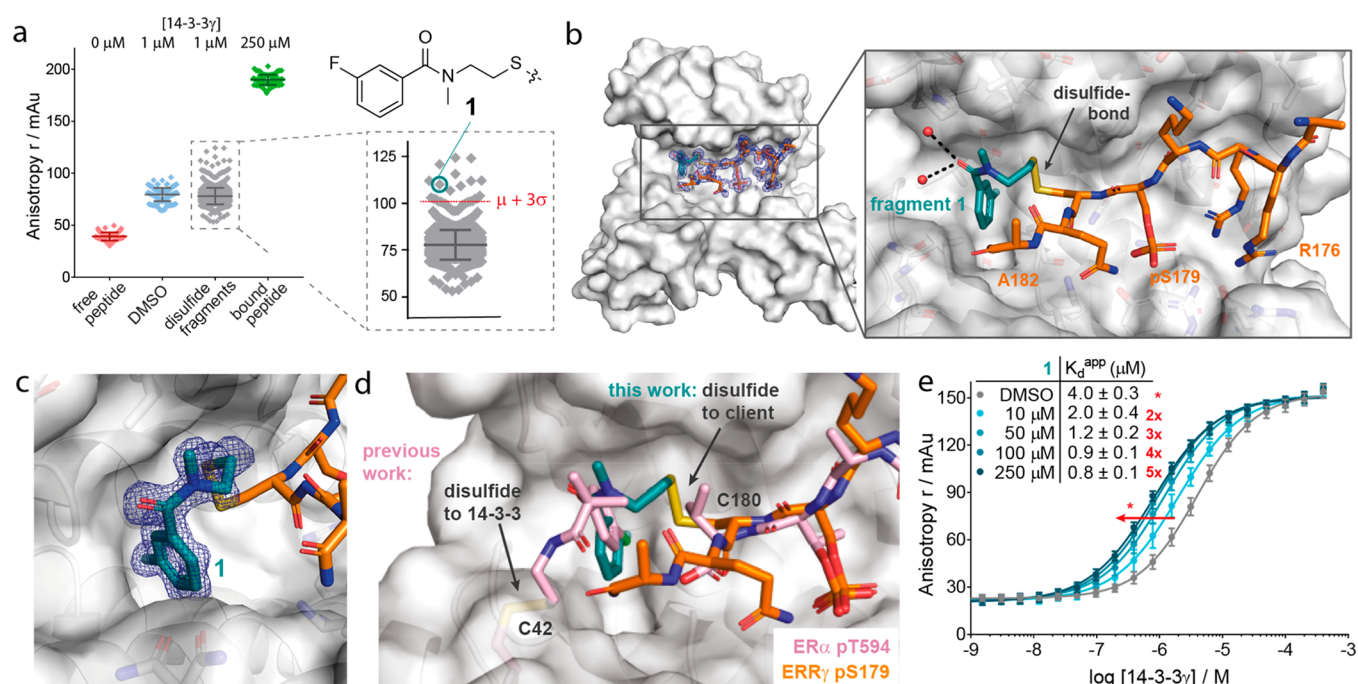


Figure 3. (a) FA-tethering screening results showing anisotropy values for fluorescently labeled ERR γ DBD phosphopeptide binding to 14–3–3 γ . Anisotropy values for free peptide (0 μM 14–3–3 γ , negative control) are shown in red and bound peptide (250 μM 14–3–3 γ , positive control) in green. Screening data of individual disulfide fragments (gray) is obtained with 1 μM 14–3–3 γ (~25% bound) with DMSO (blue) as control. Bars represent mean (μ) and standard deviation (σ). A hit selection cutoff value of $\mu + 3\sigma$ was used. Selected hit fragment 1 is indicated (blue circle). (b) Front view of a 14–3–3 σ monomer (white surface) bound by ERR γ DBD phosphopeptide (orange sticks) with view-of-bound 1 (turquoise sticks). Depicted are polar contacts (black dotted lines) and water molecules (red spheres). PDB: 6Y3W (c) Close-up view of ERR γ DBD-1 in the 14–3–3 binding groove. $2F_o - F_c$ electron density maps are contoured at 1σ . (d) Crystallographic overlay of 14–3–3/ERR γ DBD (orange sticks) with ERR γ (C180)-trapped-1 (teal sticks) and 14–3–3/ER α (pink sticks) with 14–3–3(C42)-trapped-stabilizer (pink sticks; PDB: 6HMT). (e) Dose–response anisotropy data for 14–3–3 γ titrations to the ERR γ DBD phosphopeptide in the presence of increasing concentrations of 1, with resulting apparent K_d values and fold stabilization in red.

phosphopeptide to be built (RRK{pS¹⁷⁹}CQA; Figure 2c). This ERR γ (pS179) peptide motif resides in the second zinc finger motif of the DBD, and C180 is a part of the Zn²⁺-coordinating Cys₄ structural fold.²² Interestingly, the cocrystal structure revealed that the cysteine residue (C180) at the +1 position with respect to the phosphorylated serine pointed directly into a pocket formed by the 14–3–3/peptide complex (Figure 2d). This orientation of C180 in the 14–3–3 binding groove inspired our efforts to discover disulfide-bound fragments to stabilize this complex.

DISULFIDE TRAPPING BY FA-TETHERING TO FIND STABILIZERS

We have previously shown that a natural or engineered cysteine residue in close proximity to a ligandable pocket can aid the discovery of PPI stabilizers through the use of disulfide trapping (“tethering”).^{11,13} This approach was demonstrated for the interaction of 14–3–3 with ER α ,¹³ where we incubated a cysteine-containing isoform of 14–3–3 with a disulfide-containing fragment library in the *apo* or phosphopeptide-bound state and quantified the protein-fragment conjugate by intact protein mass spectrometry (MS). We envisioned that an inversion of this paradigm, wherein a cysteine residue was located on the client-derived peptide, could also be useful for disulfide trapping. Furthermore, FA could be used as the primary read-out for induced complex formation. The placement of C180 in the ERR γ DBD motif presented an opportunity for the application of this site-directed approach,

based on client FA-tethering to identify novel chemical starting points for PPI enhancers.

For the FA-tethering screen, we used 14–3–3 γ , the isoform with the highest affinity for ERR γ DBD (Figure 2b; SI Figure S1) and a fluorescein-labeled ERR γ DBD phosphopeptide. The complex was incubated with 1600 individual disulfide fragments under reducing conditions (250 μM β -mercaptoethanol), at a protein concentration where initially ~25% of the peptide was bound (25% of the maximal FA signal; 1 μM 14–3–3 γ). The read-out of this screen was functional stabilization as a result of fragment binding, as observed by an increase in the anisotropy of the labeled phosphopeptide. The free and fully protein-bound peptides were included as controls (Figure 3a). Hit selection was based on an anisotropy value of three standard deviations above the mean of the test compounds ($\mu + 3\sigma$). On the basis of this, a total of 30 disulfide fragments were selected for further validation experiments.

MOLECULAR MECHANISM OF PPI STABILIZATION

To validate the stabilizing effect observed during the screen, fragments were soaked into cocrystals of 14–3–3 σ and the ERR γ DBD phosphopeptide. Only one of the costructures revealed additional electron density, allowing unambiguous modeling of fragment 1 in the continuous density (Figure 3b,c), indicating covalent binding of the fragment to C180 of ERR γ . A side chain flip for Q181 of ERR γ was observed compared to the protein/peptide binary crystal structure, enabling accommodation of the fragment in the pocket. The phenyl group of the fragment engaged the 14–3–3 binding

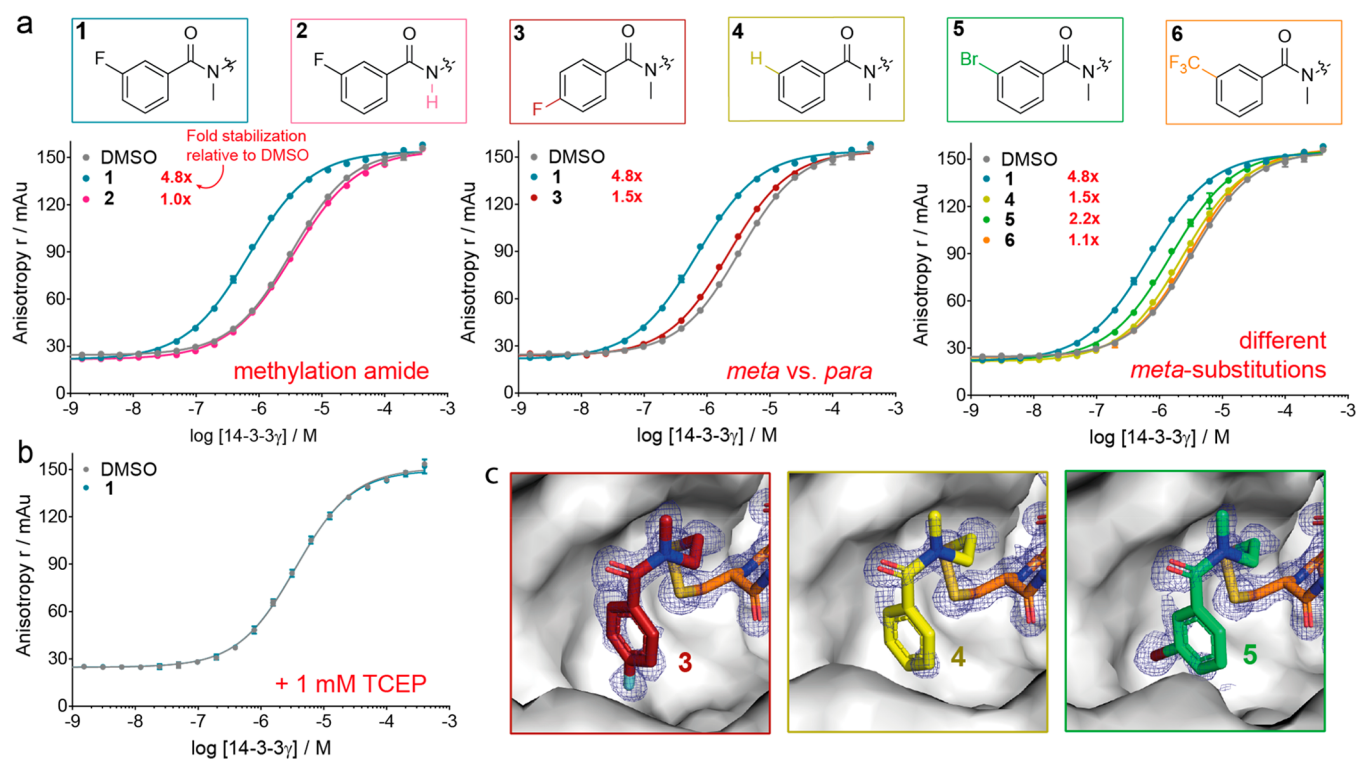


Figure 4. (a) Dose–response anisotropy data for 14–3–3γ titrations to 100 nM fluorescein-labeled ERRγ DBD phosphopeptide in the presence of either DMSO (gray) or 100 μM of fragment 1–6 (chemical structures with structural differences compared to 1 given in different colors). (b) Dose–response anisotropy data for 14–3–3γ titrations to 100 nM fluorescein-labeled ERRγ DBD phosphopeptide in the presence of either DMSO or 250 μM of fragment 1, whereby 1 mM TCEP is added to the assay buffer. (c) Close-up views of ERRγ DBD phosphopeptide (orange sticks) in the 14–3–3 binding groove (white surface) with disulfide-bound fragments 3, 4, and 5. $2F_o - F_c$ electron density maps are contoured at 1σ . PDB: 6Y18, 6XXC, 6XYS.

groove to bury the *meta*-fluorine substitution fully in a hydrophobic cavity. Additionally, the methylated amide was oriented so that the methyl group pointed upward, toward the hydrophobic residues in helix 9 of 14–3–3 that form the “roof” of the pocket.

Interestingly, a crystallographic overlay with the costructure of 14–3–3 bound by an ERα phosphopeptide and 14–3–3σ(C42)-tethered stabilizer reported previously¹³ reveals the strikingly similar subpocket engagement by the two distinct fragments (Figure 3d). Whereas both fragments are disulfide-trapped to cysteines, the trapped residue is localized at different sites and on different proteins—C42 of 14–3–3 versus C180 on the ERRγ client peptide. Despite this, a similar orientation of the fragments is observed. Remarkably, even though the halogen substituent is not at the same position (*meta*-fluorine versus *para*-chloro), the phenyl rings are tilted in such a way that the halogens nearly overlap. An additional similarity is the presence of a methyl group pointing upward in the direction of hydrophobic residues on the “roof” of the binding site.

The stabilizing activity of this fragment toward the 14–3–3/ERRγ DBD (C180) protein complex was further studied in a protein titration experiment where 14–3–3γ was titrated against a fixed concentration of fluorescently labeled ERRγ DBD phosphopeptide in the presence of DMSO (control) or increasing concentrations of 1. Saturating concentrations of 1 resulted in a 5-fold PPI stabilization of the 14–3–3/ERRγ-phosphopeptide complex (Figure 3e).

DISSECTING THE CRUCIAL ELEMENTS FOR STABILIZATION

To investigate which chemical features were important for the stabilization of the 14–3–3/ERRγ phosphopeptide complex, several variants of hit fragment 1 were synthesized. Stabilizing activity of these fragments was analyzed by FA where 14–3–3γ was titrated against a fixed concentration of fluorescein-labeled ERRγ DBD phosphopeptide in the presence of DMSO (control) or 100 μM of disulfide (Figure 4a). The ability of the fragment to stabilize the protein–peptide complex was indicated by the left shift of the binding curve (fold-stabilization relative to DMSO control).

Fragment 2 revealed the significance of the methylated amide as removal of the methyl group destroyed the stabilizing activity of the fragment. This was likely caused by the earlier described interactions between the methyl group of 1 and the hydrophobic amino acids on the “roof” of the 14–3–3 binding groove.

Replacement of *meta*-fluorine substitution with a *para*-fluorine substituent, as in fragment 3, led to a strong decrease in activity. Given the overlap of halogen positioning between these fragments and fragments from earlier published work (Figure 3d), we expected the exact positioning of the *meta*-fluorine in the 14–3–3 binding groove to be crucial for the stabilization. This was further illustrated by the comparison of fragment 1 to fragments 4–6. When the *meta*-fluorine substituent was removed (fragment 4), the stabilization fell dramatically. When the *meta*-fluorine was substituted for a bromine substituent (fragment 5), the fragment retained some stabilization; however, the increase in halogen size seemed to

be less favorable for stabilization. This size effect was also apparent in the lack of stabilization by fragment **6**, containing the larger CF₃ substituent. Overall, this concise SAR revealed the impact of small structural changes on the stabilizing effect.

The significance of fragment-peptide disulfide formation for stabilization has been shown by performing the 14–3–3 γ titrations in the presence of a large amount of reducing agent (1 mM TCEP). TCEP removed all stabilizing effect by fragment **1** on the PPI as the disulfide formation between the fragment and the peptide was impaired (Figure 4b). The importance of disulfide formation for stabilization was further confirmed as fragment **1** did not stabilize the binding of 14–3–3 γ to ERR γ C180S or two other client-derived phosphopeptides that lacked a cysteine (ER α and TAZ, SI Figure S3).

In addition to the FA experiments, fragments were soaked in cocrystals of 14–3–3 σ and ERR γ , providing electron densities for three additional fragments (3–5) with similar binding motifs to those of **1** (Figure 4c). These crystal structures showed weaker electron densities in comparison to fragment **1**. However, the strong electron densities were consistently observed for the disulfide, methylated amide, and halogens, which we postulated were the most crucial elements for 14–3–3/ERR γ PPI stabilization. Interestingly, conclusions drawn from FA data about the importance of halogen positioning and size were confirmed by the crystal structures. Structural overlay of fragment **1** with fragments **3** and **5** shows small ring displacements upward and sideways, respectively (SI Figure S4).

The identification of molecular starting points for the development of PPI stabilizers has been a major hurdle hampering widespread adoption of the paradigm for disease modulation. In our previous work, we showed that disulfide tethering, using an MS based screen, identified fragment molecules that stabilized a 14–3–3 PPI. Herein, we explored a variant of the screen using FA as readout and targeting a native cysteine residue in the client peptide to enhance PPI complex formation. Beginning with a diverse disulfide library, we identified stabilizing chemical matter and validated the screen with a concise SAR, thus illustrating the effectiveness of FA-tethering as a primary screen. In addition to directly selecting for compounds that enhance the PPI, this method has proven itself as a fast, inexpensive, and robust methodology leading to novel chemical matter for PPI modulation.

■ ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at <https://pubs.acs.org/doi/10.1021/acscchembio.0c00646>.

FA and ITC 14–3–3/ERR γ motif binding data, cocrystal structure of 14–3–3/ERR γ LBD motif, dose–response curves for FA tethering hits, FA selectivity data, and crystallographic overlay of costructures (Figures S1–S4); experimental methods, synthetic procedures, and compound characterization; XRD data collection and refinement statistics (Table S1) (PDF)

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Notes

L.B., M.R.A. and C.O. are founders of Ambagon Therapeutics. L.B. is a member of Ambagon & scientific advisory board, M.R.A. is a director, C.O., E.S., and G.P.M., are employees of Ambagon.

■ ACKNOWLEDGMENTS

We thank the Renslo laboratory for synthesis of the SMDC disulfide library, M. Zhong and G.M. Lee for their help with the FA-tethering screen, and R.J. Neitz and P. Burroughs for sharing synthesis routes and discussing hit optimization. The research described was funded by The Netherlands Organization for Scientific Research (NWO); Gravity Program 024.001.035, Vici grant 016.150.366 and Echo grant 711.017.014) and a Breakthrough Award from the Ono Pharma Foundation.

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