Disorder-to-order transition underlies the structural basis for the assembly of a transcriptionally active PGC-1 α /ERR γ complex

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Peroxisome proliferator activated receptor (PPAR) γ coactivator-1 α (PGC-1 α) is a potent transcriptional coactivator of oxidative metabolism and is induced in response to a variety of environmental cues. It regulates a broad array of target genes by coactivating a whole host of transcription factors. The estrogen-related receptor (ERR) family of nuclear receptors are key PGC-1 α partners in the regulation of mitochondrial and tissue-specific oxidative metabolic pathways; these receptors also demonstrate strong physical and functional interactions with this coactivator. Here we perform comprehensive biochemical, biophysical, and structural analyses of the complex formed between PGC-1 α and ERR γ . PGC-1 α activation domain (PGC-1 α^{2-220}) is intrinsically disordered with limited secondary and no defined tertiary structure. Complex formation with ERRy induces significant changes in the conformational mobility of both partners, highlighted by significant stabilization of the ligand binding domain (ERRyLBD) as determined by HDX (hydrogen/deuterium exchange) and an observed disorder-to-order transition in PGC-1 α^{2-220} . Small-angle X-ray scattering studies allow for modeling of the solution structure of the activation domain in the absence and presence of ERRyLBD, revealing a stable and compact binary complex. These data show that PGC-1 α^{2-220} undergoes a large-scale conformational change when binding to the ERRyLBD, LBD, leading to substantial compaction of the activation domain. This change results in stable positioning of the N-terminal part of the activation domain of PGC-1 α , favorable for assembly of an active transcriptional complex. These data also provide structural insight into the versatile coactivation profile of PGC-1 α and can readily be extended to understand other transcriptional coregulators.

Peroxisome proliferator activated receptor (PPAR) γ coactivator-1 α (PGC-1 α) is a powerful transcriptional coactivator with an established role in key metabolic functions, and has been implicated in the pathogenesis of obesity, type 2 diabetes, neurodegeneration, and cardiomyopathy (1, 2). PGC-1 α is unusual or even unique in its ability to respond to a wide variety of physiological signals, coactivate a broad range of transcription factors, and coordinate the regulation of oxidative metabolic gene programs in a tissue-specific manner (1-3). For example, PGC-1 α is induced by exercise in skeletal muscle. In this tissue PGC-1 α coactivates transcription factors such as nuclear receptors, nuclear respiratory factors, and myocyte-specific enhancer factors, and induces mitochondrial biogenesis, fatty acid oxidation, glucose and lipid uptake, angiogenesis, and resistance to atrophy (1, 4). A substantial portion of PGC-1a function in oxidative metabolism is mediated through the estrogen-related receptor (ERR) subfamily of nuclear receptors (3, 5). ERR γ is a member of the subfamily and plays a significant role in the regulation of energy homeostasis as well as in mitochondrial function (5, 6). PGC-1 α and ERRy share tissue specificity, have overlapping functions, and demonstrate a robust physical and functional interaction (7,8). Following a coactivator-receptor interaction model (9), the

interaction between PGC-1 α and ERR γ is mediated at least in part by characteristic nuclear receptor (NR) boxes or LXXLL motifs (denoted as L1, L2, and L3 in Fig. 1 A and B) located within the activation domain of PGC-1 α and the nuclear receptor ligand binding domain (LBD). The broad range of physiologic processes mediated by PGC-1 α and the large repertoire of nuclear receptors with which it partners necessitate a deeper understanding of how the coactivator achieves specificity despite the similarity in its interactions with members of the superfamily (10-16). Structural analyses of these interactions have been restricted to the use of small peptides representative of single LXXLL motifs. Unfortunately, these studies have not yielded sufficient insights into the overall structure of PGC-1 α and its correlation to function. An in-depth investigation of the PGC-1 α activation domain (the primary mediator of these interactions) in the context of its interaction with ERRyLBD could help unravel the molecular basis for PGC-1 α binding and specificity.

To address these questions, we performed a comprehensive biochemical and biophysical analysis of a binary complex formed between the activation domain of PGC-1 α [PGC-1 α^{2-220} /PGC1 α 220] and the ERR γ ligand binding domain [ERR γ LBD]. Combinations of several biophysical approaches (refer to *SI Text, Glossary of Terms*, for a summary of the terms and techniques used in the study) delineate properties of PGC1 α 220 that are responsible for its specificity and affinity for ERR γ . Small-angle X-ray scattering (SAXS) studies provide unique solution models of the binary complex. Our results in combination with existing data allow us to propose a structural model of association involving the activation domain and also develop a generalized theory explaining the functional adaptability of PGC-1 α .

Results

Biochemical Analysis of Specificity, Affinity, and Stoichiometry for PGC1 α /ERR γ . We first investigated the interaction between PGC-1 α and ERR γ to establish requirements for binding in vitro. GST pulldown studies demonstrated binding between ERR γ LBD and PGC1 α 220, and fragments nested within PGC1 α 220 (Fig. 1*A*–*D*). These results in conjunction with coactivation studies (Fig. S1) were in agreement with previous data (7, 8). Our results showed that although L1 is dispensable to the interaction, L2 and L3 are both capable of mediating this interaction independent of each other. Isothermal calorimetry (ITC) is a thermodynamic technique

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Fig. 1. Molecular determinants of specificity and affinity for the binary complex. A schematic representation of (*A*) ERR γ and PGC-1 α domains and (*B*) three NR boxes/LXXLL motifs (L1, L2, and L3). Interaction studies using GST-fusion fragments of (*C*) ERR γ and ³⁵S-labeled PGC-1 α or (*D*) PGC-1 α and ³⁵S-labeled ERR γ .

that directly measures the heat released or absorbed when two biomolecules interact and provides thermodynamic parameters including binding constants; hydrogen/deuterium exchange (HDX) mass spectrometry is a study of the rate and percentage of the mass increase of a protein when the amide hydrogens are exchanged with solvent deuterium. These changes are indicative of conformational mobility/stability in the regions studied and can provide insight into the local changes in structure upon complex formation. Analysis of ERR γ LBD interactions with PGC-1 $\alpha^{136-220}$ (a fragment containing L2 and L3) by ITC showed that wild type had the highest binding affinity ($K_d = 0.95 \ \mu M$) for the LBD (Fig. S1, Table S1), consistent with data from HDX analysis (Fig. 2). HDX suggests the functional equivalence of PGC-1 $\alpha^{136-220}$ and PGC1 α 220 in stabilizing the conformational mobility of ERRyLBD, as indicated by the reduction in deuterium exchange kinetics in helices 6, 7, 10/11, and 12 and the beta sheet region (Fig. 2). The noticeable decrease in both affinity and HDX kinetics of the LBD upon mutating L2 suggests that it is the major contributor to the interaction (Fig. 2, Table S1). The modest, yet important, contribution by L3 is evident in data from the L2 mutant; L3 shows micromolar affinity for the LBD in addition to muted stabilization of equivalent regions of the LBD as determined by HDX. The role of both L2 and L3 is underscored by the observation that only a combinatorial mutation involving both L2 and L3 has a dramatic effect.

The binary complex between ERRγLBD and PGC1α220 was then further characterized by size-exclusion chromatography in-line with multiangle light scattering (SEC-MALS) and sedimentation velocity (SV) analysis by analytical ultracentrifugation. Parameters determined from SEC-MALS as well as SV analysis provide solution properties of the molecules such as molecular mass, shape, and size. SEC-MALS analysis demonstrates that PGC1 α 220 exists as a monomer in solution, with a retention time consistent with an extended shape. In contrast, ERRyLBD exists exclusively as a compact dimer (Fig. 3A, Table 1). In agreement with ITC analysis (Table S1), SEC-MALS analysis of the binary complex suggests a molar ratio of 1:2 (PGC1 α 220:ERR γ LBD) (Table 1). A molecular mass of 82,747 Da also consistent with a 1:2 complex, was derived by combining parameters determined from SEC and SV analysis (Fig. S2) (17). Taken together, the 1:2 stoichiometry thus established for the interaction shows that a single coactivator molecule engages the ERRyLBD homodimer. Helix 12 stabilization seen from HDX data indicates that both L2 and L3 engage helix 12 of the LBD dimer, whereas the variation in binding constants for L2 and L3 (1.61 µM versus 8.69 µM), suggests an asymmetric interaction.

PGC1 α **220 is Intrinsically Disordered.** Given the importance of the PGC1 α 220 domain for nuclear receptor coactivation, we examined the solution properties of this domain alone. Bioinformatic analysis of the PGC1 α 220 sequence indicates the presence of ordered regions in the N-terminal half of the protein, but clearly predicts significant disorder in the regions following L1 (Fig. S3).



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Fig. 2. HDX analysis of the ERR_YLBD/PGC-1 α activation domain interaction. HDX analysis of ERR_YLBD interaction was performed with five distinct constructs of PGC-1 α activation domain. (A) ERR_YLBD \pm PGC1 α 220. (B) ERR_YLBD \pm PGC1 α (136–220) WT, L2A, L3A, and L2L3A. The differential HDX between apo ERR_YLBD and PGC-1 α bound LBD is mapped onto PDB ID code 1KV6. The difference in the mean HDX across six time points for coactivator bound and unbound LBD is represented as percent change and colored according to the key. Gray, no change in HDX between bound and unbound LBD; light to dark blue, slower rates of HDX between compared conditions; yellow to red, faster rates of HDX between compared conditions.

A protein of this mass is predicted to have Stokes radii (R_s) of 39.1, 41.1, and 52.6 Å, in the native, molten globule, and unfolded states, respectively (18). Despite being monomeric by mass as indicated by SEC-MALS, a Stokes radius of R_s of 42.9 Å \pm 0.46 was determined, indicative of a molten-globular, extended conformation in solution (Fig. 3A, Table 1). This extended confor-



Fig. 3. Biophysical properties of the binary complex. (A) Representative SEC-MALS analyses showing elution profiles for PGC1 α 220 (red), ERR γ LBD (blue), and the binary complex (pink) on a Superdex 200 10/300 column at room temperature. Absorbance profiles are shown in relative units at 280 nm (left y axis). Shown as colored circles on each elution profile are the masses determined from MALS analysis; the right y axis denotes mass in daltons. (*B*) Sedimentation coefficient distribution [c(S)] analysis of sedimentation to efficient distribution [c(S)] analysis of sedimentation to a 20 °C), and the binary complex (pink line, 28 μ M at 20 °C).

Table 1. Biophysical properties of PGC1 α 220/ERR γ LBD binary complex and its components

	SEC*	MALS ⁺		SV		Siegel and Monty	QELS [†]
Protein	R _s (Å)	Molar Mass [‡]	Stochiometry [§]	S _{20,w}	f/f_o	Molar Mass [‡]	<i>R_h</i> (Å)
PGC1α220	42.9 ± 0.5	23,945 ± 3,434	Monomer (24,456)	1.3	2.0	22,598	ND ¹
ERRyLBD	33.2 ± 0.13	53,673 ± 3,094	Dimer (52,928)	3.8	1.2	51,119	25.7 ± 1.2
Complex	51.7 ± 1.8	74,175 ± 473	1:2 (77,386)	4.0	1.5	82,747	ND

*The results presented are the average of 2-4 replicates.

[†]SEC-MALS-quasi-elastic light scattering (QELS) measurements were made at room temperature.

*Molar masses are presented in units of daltons.

[§]Stoichiometry as inferred from experimentally determined molecular mass. In parentheses, theoretical values as computed from primary sequence using SEDNTERP are presented.

¹ND, not determined

mation is recapitulated in SV analysis, which at 4 °C yields a frictional coefficient ratio (f/f_o) of 2.7 (a globular particle has an f/f_o of approximately 1.2, whereas a value greater than 1.8 indicates asymmetry in shape), indicating an extremely elongated shape (Fig. 3B, Table 1). HDX analysis of PGC1 α 220 revealed that most of the backbone amides were fully exchanged after 10 s incubation with deuterium, consistent with the predicted disorder (Table S2). CD spectroscopy provides a qualitative and quantitative estimation of secondary structure in a macromolecular species. CD analysis of PGC1 α 220 estimated only 27% native secondary structure in the activation domain (Fig. 4A, Table 2).

SAXS is an established method for the characterization of biological macromolecules in solution and is extremely wellsuited to study flexible and extended macromolecules. Structural parameters such as the mean particle size (radius of gyration, R_g) and maximal intramolecular distance (D_{max}) can be derived from SAXS analysis. These data can then be used to determine molecular characteristics such as mass, shape, and volume; furthermore, recent innovations allow for the three-dimensional modeling of molecular shape in solution at low resolution.

SAXS analysis of PGC1a220 yielded consistent data and indicated a monodisperse particle (Table 3, Table S3). By Flory's law (19) a protein of this size is expected to have an R_p of 18.6 Å when natively folded and an R_g of 78.2 Å as an extended polymer. SAXS analysis of PGC1 α 220 determined an R_g of 60 Å and is consistent with other data that indicate a protein with high flexibility and low residual structure. This low level of compactness is further underscored by an R_s/R_g value of approximately 1.45 for PGC1 α 220, similar to that expected for a random coil (~1.5), rather than that for a globular protein (~ 0.8) or a premolten globule (~ 0.9) (20, 21). The asymmetry in the P(r) profile derived from SAXS data for PGC1 α 220 with a D_{max} of approximately 240 Å correlates to anisotropy in molecular shape (Fig. 5A). The Kratky and Porod–Debye plots (22–26) for PGC1α220 are clearly indicative of a largely disordered species lacking a well-packed core (Fig. 5 B and C). In summary, these orthogonal measurements clearly establish PGC1 α 220 as a predominantly disordered species with limited secondary structure



Fig. 4. Thermodynamic and structural stability of the binary complex. (*A*) Far-UV CD spectra of PGC1 α 220 (red), ERR γ LBD (blue), and binary complex (pink). Spectra were normalized to molar ellipticity, secondary structure was quantified using CDFIT and summarized in Table 2. (*B*) Thermal denaturation of ERR γ LBD (•) and binary complex (**n**). The CD signal at 222 nm was monitored as a function of temperature. (*C*) Chemical denaturation ERR γ LBD (•) and binary complex (**n**). The CD signal at 222 nm was monitored as a function of temperature. (*C*) Chemical denaturation ERR γ LBD (•) and binary complex (**n**). Intrinsic tryptophan fluorescence was monitored as a described in *Materials and Methods* and are summarized in Table 3.

PGC1α220 Undergoes a Disorder-to-Order Transition upon Complex Formation with ERRγLBD. Next we investigated the nature of complex assembly. HDX studies showed that despite the absence of HDX perturbations in PGC1α220 upon complex formation (Table S4), docking of the coactivator results in significant stabilization of the conformational mobility of ERRγLBD (Fig. 2). This result is in agreement with secondary structure analysis, which estimated 52% alpha-helical content in the binary complex (Table 2), illustrating an increase in total secondary structure induced by protein interaction. This phenomenon translates into increased stability of the binary complex, as seen in thermal and chemical denaturation studies (Fig. 4 *B* and *C*, Table 2). Despite the disordered nature of PGC1α220 alone, the increase in stability indicates that the interaction confers conformational stability to the complex.

The properties of this complex were further investigated using SV analysis (Table 1). The differences $in f/f_0$ between ERR γ LBD alone and the binary complex (1.2 versus 1.5) indicate a change from a compact to a more elongated state upon complex formation. The reduction in f/f_0 between PGC1 α 220 and the binary complex (2.0 versus 1.5) is suggestive of coactivator compaction upon complex formation.

SAXS analysis provides structural insight into the nature of these differences in shape. Structural parameters derived from SAXS measurements on the binary complex were constant across the range of concentrations examined (Fig. 5*A*, Table S3). A comparison of the R_g and D_{max} values for the binary complex upon extrapolation to infinite dilution (45 Å and 175 Å) and PGC1 α 220 (60 Å and 240 Å) shows a contraction of the maximum dimension upon complex formation, whereas the Kratky and Porod–Debye plots for the complex are indicative of a well-folded globular particle (Fig. 5 *B* and *C*). If the PGC1 α 220 component remained intrinsically disordered upon complex formation, we would have expected to observe an even larger spatial extent for the complex than seen with either component alone. Combined, these experiments indicate that PGC1 α 220 undergoes a disorder-to-order transition (27, 28) upon docking onto the

Table 2. Characterization of secondary structure and thermal and chemical stability of the complex by CD and fluorescence

_		PGC1α220	ERRγLBD	Binary complex
Secondary	% alpha	21.69	68.06	52.38
structure*		(47-aa)	(157-aa)	(236-aa)
	% beta	5.39	2.35	4.76
	% coil	72.91	29.59	42.86
Thermal denaturation Chemical denaturation	T_m^{\dagger} [K] D_m^{\dagger} [M] _{urea}	Inde- terminate Inde- terminate	324.42 ± 0.059750 3.25 ± 0.095	328.51 ± 0.055400 3.47 ± 0.075

*Secondary structure estimation was calculated using CDFIT.

 ${}^{\dagger}T_m$ and D_m values reported for each species are obtained from an average of three different experiments. The error reported is the standard deviation.

Table 3. Table of structural parameters derived from SAXS analysis

Sample	Range of concentrations examined	R _g (Å)	D _{max} (Å) *	
	mg/mL			
PGC1α220	2.5-4.5	61.3 ⁺	240	
ERRyLBD	1.6–11.5	25.3 ^{+,‡}	90	
PGC1α220—ERRγLBD	1.2–5.6	42.3 ^{+,‡}	175	

A complete listing of structural parameters derived from SAXS measurements is available in Table S3.

*As determined by GNOM analysis.

[†]By Guinier analysis.

*Extrapolated to infinite dilution.

LBD, resulting in a binary complex, which is significantly more compact and structured.

Shape Reconstruction of PGC1 α 220/ERR₇LBD Binary Complex from SAXS Data. For well-folded globular species, SAXS data can be used to reconstruct solution shape at low resolution (29, 30). However, these methods are not suitable for modeling intrinsically disordered proteins such as PGC1 α 220. Instead, these species are better described as ensembles in solution. Using the ensemble optimization method (EOM) (31, 32) (see *Materials and Methods*) we identified a pool of conformers that collectively reproduce the experimental solution data (Fig. 6 *A* and *B*). A gallery of EOM-generated models for PGC1 α 220 clearly illustrates the lack of globularity and the conformational diversity that is compatible with the intrinsic properties of the protein (Fig. 6*C*).

Shape reconstructions for $ERR\gamma LBD$ were generated using the programs DAMMIN and GASBOR (29, 30). Both approaches reproducibly yielded envelopes with good correlations between experimental and calculated scattering data [sqrt(χ) ~1.5; normalized spatial discrepancies (NSDs) ranging from 1.0 to 1.3] (Fig. 7A). The averaged envelope is a globular particle with a dimension of $76 \times 52 \times 48$ Å (Fig. S4). Shape reconstructions of the binary complex were also performed in the same manner and the GASBOR results are discussed here. Reproducible envelopes with good correlations between experimental and calculated scattering data were obtained [sqrt(χ) ~1.2, NSDs ranging from 1.4 to 1.5]; because of the inherent asymmetry suggested by its 1:2 stochiometry, no symmetry restraints were applied (Fig. 7B). The final averaged envelope is an asymmetric and elongated prolate ellipsoid with a dimension of $83 \times 150 \times 69$ Å (Fig. S4). The hydrodynamic properties calculated for these shapes also closely resemble those determined experimentally (Table S5).

Solution Data-Based Structural Model for the Binary Complex. Finally, we combined our results from SAXS analysis with the existing structural data on ERR γ LBD [Protein Data Bank (PDB) ID code 1KV6] to model the binary complex. We first evaluated the scattering data and solution parameters for the LBD against the available crystal structure using CRYSOL and HYDROPRO



Fig. 5. Solution properties of the binary complex and its individual components determined using SAXS. (A) Shape distribution [P(r)] functions derived from SAXS analysis for PGC1a220 (red), ERR_YLBD (blue), and the binary complex (pink). (B) Kratky plot analysis for the proteins examined, where the intensity of scattering is plotted as IQ^2 versus Q. I is the scattering intensity and Q is scattering angle ($Q = 4\pi \sin \theta / \lambda$). (C) Porod–Debye plot of the SAXS data for the samples examined in the study, shown as IQ^4 versus Q^4 .



Fig. 6. Shape reconstruction for PGC1 α 220 from SAXS data using EOM analysis. (A) EOM fit (red line) to the SAXS data for PGC1 α 220 (open squares), with χ^2 of 1.14 for the best selected pool solution. (B) R_g distributions for the pool (black) and optimized (red) ensembles generated by EOM analysis. (C) Representative gallery of bead models for PGC1 α 220 derived from EOM analysis. The bead radius used in these models is 3.4 Å, and this figure was generated using PyMOL.

(see *Materials and Methods*), respectively, demonstrating general concordance, with some variations observed between the crystallographic model and its solution state (Table 3, Fig. S4). Superposition of the crystal structure into the molecular envelopes reveals good spatial agreement with the solution data (Fig. 7*A*) and allows inference of the position of PGC1 α 220 relative to the ERR γ LBD domain (Fig. 7*B*). We then implemented the program MONSA, which derives the relative positions of the different components of a composite particle by simultaneously fitting multiple scattering profiles from both the overall particle and its component parts (29, 33). By this approach, the shape determined recapitulates the shapes observed by the DAMMIN and GASBOR approaches, and corroborates our assignment of locating ERR γ LBD (Fig. 7*C*).



Fig. 7. Shape reconstruction of ERR_YLBD and its binary complex with PGC1 α 220. (*A, Left*) GASBOR fit (blue line) to primary scattering data (black squares) for ERR_YLBD. (*Right*) SAXS envelope calculated using GASBOR and rigid body docking of structural model of ERR_YLBD into the envelope (*B, Left*) GASBOR fit (red line) to primary scattering data (black squares) for binary complex. (*Right*) SAXS envelope of the binary complex calculated using GASBOR It to primary data (ERR_YLBD, blue line and black squares; binary complex, red line and black squares), (*Right*) SAXS envelope of the binary complex. (*Right*) SAXS envelope of the binary complex.

This analysis provides important insights into the solution structure and mechanism of interaction between PGC1a220 and ERRyLBD. SAXS analysis shows how the highly disordered region engages the well-structured LBD to form an elongated and asymmetric, but overall more compact and globular structure (Fig. 7 B and C). Although SAXS data alone cannot provide highresolution structural details of the binding interface, a comprehensive experimental approach allows us to propose an interaction model for the binary complex (Fig. 8). Our studies reveal that PGC1 α 220 is involved in a bipartite interaction with the LBD dimer through high affinity L2 and low affinity L3 interactions. The binding of a single PGC1α220 molecule to the LBD dimer is facilitated by the intrinsic disorder in the coactivator. PGC1a220 simultaneously occupies coactivator grooves on both LBD units and stabilizes the dimerization interface by navigating the region via the flexible spacer. Such an asymmetric interaction clearly distinguishes PGC-1a/ERRy complex from those with other receptors such as PPAR γ , hepatocyte nuclear factor-4 α (HNF4 α), and ERR α (Fig. S5) (13, 15, 16) and has important functional consequences.

The solution model also illustrates the structure-function relationship of the activation domain with consequences to PGC-1 α mediated transcription complex assembly. Our data show that PGC1 α 220 is tethered to the LBD via LXXLL-mediated interactions. With the C-terminal portion of PGC1 α 220 (PGC-1 α ¹³⁶⁻²²⁰) wrapped across the LBD, it is evident that the N-terminal half (PGC-1 α ²⁻¹³⁵) contributes to the asymmetry in shape seen in the envelope structure. This region contains the CREB binding protein (CBP)/p300 and steroid receptor coactivator-1 (SRC-1) interaction domain and appears to be locked into a stable position in the binary complex in comparison to the isolated structure. The compaction and securing of the LXXLL motif-containing region stably positions the N-terminal portion unhindered by the C-terminal disorder, in an orientation accessible for CBP/p300 recruitment, a crucial step in the assembly of an active transcriptional complex.

Discussion

PGC-1 α is a highly versatile coactivator, featuring an extremely broad repertoire of transcriptional partners. This versatility is presumed to be the biochemical basis for a huge array of physiological processes under control or modulated by PGC-1 α . The ERRs are key PGC-1 α partners with the PGC-1 α /ERR interaction complexes regulating energy homeostasis in metabolically active tissues. These functions make them attractive drug design targets for metabolic disorders, necessitating a thorough understanding of the molecular nature of these interactions. Despite the existing body of work examining PGC-1 α interactions with nuclear receptors, the paucity of structural data has left important questions about its function unanswered. The work presented



Fig. 8. Predicted interaction model for the binary complex. A structural model for ERR_YLBD and PGC1 α 220 interaction proposed based on a summary of our data. HDX data is mapped onto the ERR_YLBD dimer (PDB ID code 1KV6) to indicate regions of the LBD affected by the interaction and PGC1 α 220 is shown in orange.

here provides a unique structural model for PGC-1 α interaction with ERR γ and establishes a foundation for its functional adaptability.

Our multipronged approach unravels features of the PGC-1 α / ERR γ interaction elusive to traditional structural methods and allows us to propose an interaction model for the binary complex. This model is distinguished from those proposed for SRC-1 binding to PPAR γ LBD (34) and 1,25-dihydroxyvitamin D₃ receptor (VDR)-retinoid X receptor heterodimer (35), in that it involves the entire activation domain of PGC-1 α and is supported by a low resolution solution model.

The results presented here firmly establish that intrinsic disorder in PGC-1 α is the principal physical basis for its versatility. For example, despite the high degree of sequence similarity, PGC-1 α is able to distinguish between ERR γ and ERR α by engaging distinct interaction interface and receptor:coactivator stoichiometry; this ability is highly dependent on its structural flexibility. These differences have important consequences visà-vis drug design targeting PGC-1a interactions with the ERRs and provide a framework for the design of subtype specific drugs. Furthermore, such structural flexibility presumably allows PGC- 1α to simultaneously engage transcription factors as well as chromatin remodeling proteins (CBP/p300). We also provide a biophysical and spatial illustration of transcription factor-induced conformational changes in PGC-1a and can infer the functional correlation between PGC-1a structure and its coactivation. Interaction between the two partners restricts conformational flexibility of the coactivator and provides structural stability to the N-terminal region of the activation domain; this series of events then primes the coactivator for CBP/p300 recruitment, an important step in transcriptional activation.

These results demonstrate the utility of combining different biophysical methods to study PGC-1 α interactions and expose the absence of equivalent data from traditional structural analysis; the lack of electron density for these regions in crystal structures is directly attributable to the intrinsic disorder (13, 15). Transcriptional coregulators are typically characterized by large molecular size and small structured domains separated by long stretches of disorder (36, 37). Recent studies including our current work demonstrate that SAXS is well-suited for the study of coregulator structure (16), provide a strong case for the continued application of SAXS as a complementary technique to traditional structural studies of transcriptional coregulators, and could contribute to a comprehensive understanding of these interactions.

Materials and Methods

Detailed methods are available in SI Text.

Cloning, Protein Expression, and Purification. Protein domains were cloned as GST-fusion or His-tagged proteins and purified by affinity chromatography using the manufacturer's protocols (GE Healthcare and Qiagen, respectively).

Domain-Mapping Studies. GST pull-down studies were performed by incubating Sepharose bead-bound GST-fusion protein with in vitro translated labeled protein (using a Promega T7 TNT reticulocyte lysate kit) and bound protein was detected using autoradiography.

Isothermal Titration Calorimetry. ITC experiments were performed on a VP-ITC Microcal Isothermal Titration Calorimeter (GE Healthcare) following manufacturer's protocol and the resulting data analyzed using the software Origin 7.0 (MicroCal).

Size-Exclusion Chromatography and Multiangle Light Scattering. SEC-MALS experiments were performed using a MALS detector coupled in-line with size-exclusion chromatography and an interferometric refractometer. Data analysis was performed using ASTRA software version 5.2 (Wyatt Technology Corp.).

Sedimentation Velocity Analysis. Sedimentation velocity ultracentrifugation experiments were performed at 25 °C with an XL-A analytical ultracentrifuge

(Beckman) and a TiAn60 rotor. Data were fit using SEDFIT (38) to determine S and f/f_0 and SEDNTERP was used to determine solvent density and viscosity.

HDX Mass Spectrometry. HDX experiments were performed as described previously (39). Briefly, protein solutions consisting either of individual proteins or of relevant complexes were exchanged into an equivalent deuterium buffer solution, exchange quenched, and the peptides subjected to mass spectrometry. Data were processed using in-house software.

Bioinformatic Analyses. For structural variability predictions for PGC1 α 220, the DISOPRED2 algorithm was used, which predicts intrinsic disorder from primary sequence information.

CD Spectroscopy. CD spectra were measured on a J-715 Jasco spectropolarimeter and normalized data was processed using CDFIT and JASCO spectral analysis software.

Fluorescence Spectroscopy. Chemical unfolding of the proteins was characterized using changes in intrinsic tryptophan fluorescence during urea denaturation. Data was collected on a T-format PTI QuantaMaster C-61 spectrofluorimeter following standard protocol and analyzed with Prism 5 (GraphPad).

Small-Angle X-Ray Scattering. X-ray scattering data were measured using synchrotron radiation; specific details are provided in *SI Text*. Forward scattering from the samples examined was recorded on a CCD detector and circularly

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averaged to yield one-dimensional intensity profiles as a function of Q. Scattering data was analyzed using Guinier, Kratky, and Porod–Debye plots, as well as the inverse Fourier transform using the program GNOM (40). Shape analyses using the programs EOM (32), DAMMIN (29), GASBOR (30), and MONSA (29) were performed as described in *SI Text*.

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