Molecular determinants of MED1 interaction with the DNA bound VDR–RXR heterodimer

Anna Y. Belorusova, Maxime Bourguet, Steve Hessmann, Sandra Chalhoub, Bruno Kieffer, Sarah Cianférani and Natacha Rochel

INTRODUCTION

Nuclear receptor (NR) superfamily of ligand-regulated transcription factors activate or repress gene expression by recruiting coactivators, corepressors, chromatin remodelers and the general transcriptional machinery to the target genes (reviewed in (1,2)). Among classical NR coactivators is Mediator, an evolutionary conserved multi-protein complex facilitating multiple stages of gene expression, notably the chromatin remodeling and pre-initiation complex formation (reviewed in (3–5)). It was discovered as a group of factors needed for the yeast RNA polymerase II activity (6,7), and subsequently various mammalian Mediator subcomplexes have been isolated through association with NRs, such as TRAP complex associated with thyroid receptor (TR), (8) and DRIP complex associated with vitamin D receptor (VDR) (9). Other similar complexes included activator-recruited cofactor ARC (10), mammalian mediator (11), mammalian Srb/Mediator complex (12), PC2 (13) and CRSP (14). Mediator is involved in strong ligand-dependent interaction with NRs primarily via its largest subunit 1 (MED1) (15–22), although for some NRs the interaction can include other Mediator subunits and alternative cofactors (23–27).

As a classical NR-binding target, MED1 contains two LXXLL motifs, also called NR-boxes, localized in a central disordered receptor-interacting domain (RID). Binding of the coactivator LXXLL motifs to the activation function 2 (AF-2) of the receptor ligand-binding domain (LBD) has been extensively characterized by structural studies (28–30). Leucines from the coactivator LXXLL motif are buried in the hydrophobic groove of the AF-2 surface formed by hydrophobic residues from helices H3, H4 and H12 of the LBD, and the NR box is locked by a charge clamp formed by a lysine on the NR H3 and a glutamate on H12. The two MED1 LXXLL motifs bind to NRs with different specificity: steroid hormone receptors preferentially bind to the first LXXLL motif, whereas non-steroid hormone receptors, such as TR and VDR, strongly interact with the second LXXLL motif (15,31,32).

ABSTRACT

The MED1 subunit of the Mediator complex is an essential coactivator of nuclear receptor-mediated transcriptional activation. While structural requirements for ligand-dependent binding of classical coactivator motifs of MED1 to numerous nuclear receptor ligand-binding domains have been fully elucidated, the recognition of the full-length or truncated coactivator by full nuclear receptor complexes remain unknown. Here we present structural details of the interaction between a large part of MED1 comprising its structured N-terminal and the flexible receptor-interacting domains and the mutual heterodimer of the vitamin D receptor (VDR) and the retinoid X receptor (RXR) bound to their cognate DNA response element. Using a combination of structural and biophysical methods we show that the ligand-dependent interaction between VDR and the second coactivator motif of MED1 is crucial for complex formation and we identify additional, previously unseen, interaction details. In particular, we identified RXR regions involved in the interaction with the structured N-terminal domain of MED1, as well as VDR regions outside the classical coactivator binding cleft affected by coactivator recruitment. These findings highlight important roles of each receptor within the heterodimer in selective recognition of MED1 and contribute to our understanding of the nuclear receptor-coregulator complexes.

Received June 29, 2020; Revised August 24, 2020; Editorial Decision September 04, 2020; Accepted September 08, 2020

© The Author(s) 2020. Published by Oxford University Press on behalf of Nucleic Acids Research. This is an Open Access article distributed under the terms of the Creative Commons Attribution Non-Commercial License (http://creativecommons.org/licenses/by-nc/4.0/), which permits non-commercial re-use, distribution, and reproduction in any medium, provided the original work is properly cited. For commercial re-use, please contact journals.permissions@oup.com
Mediator-dependent mechanisms of NR regulation by MED1 include looping of enhancers to transcription start sites via an assembly process involving transcription factors, cohesin and non-coding RNAs (33-37); directly linking chromatin remodeling and the pre-initiation complex formation (24,25); or repression of transcription through the core Mediator-associated CDK8 kinase module (reviewed in (38)). At the same time, some regulatory roles of MED1 could be Mediator-independent as it can be recruited to NRs independently from the other Mediator subunits, promoting the association with the Mediator core in a second step. Occupancies of MED1 and NRs on the genome sites are highly correlated (35,39) and MED1 levels are highly elevated on super-enhancers in embryonic stem cells and in cancer cells where NRs act as master regulators (40,41). Recent studies showed that intrinsically disordered regions of MED1 can form phase-separated droplets that compartmentalize and concentrate transcriptional regulators (42).

MED1 has been shown to be essential for various biological functions of large number of NRs (reviewed in (38)). Due to its important role in human physiology, it has been suggested as a possible target for several disorders, such as metabolic syndrome (43), fatty liver (44) and several types of cancer, including breast and prostate cancers (reviewed in (45)).

Despite essential role and high therapeutic potential of MED1, no atomic structural data is available for this protein or its homologs. Furthermore, while much has been discovered about the Mediator complex and its association with transcriptional machinery, the mechanistic details of how MED1 bridges the Polymerase II to NRs are far less understood. Most of the structural investigations on NR-MED1 association, similarly to analogous coregulator complexes, are still limited to the recognition of LXXLL-peptides or short RIDs by NRs (46-48). Recent advances in single particle cryo-electron microscopy (cryoEM) allowed the structural characterization of the full-length estrogen receptor alpha/p160/p300 coactivator complexes (49,50). However, the detailed mechanism of how NRs trigger the formation of big regulatory complexes that directly alter the transcriptional rate has not yet been fully elucidated and remains challenging due to the presence of large intrinsically disordered regions in the coactivators proteins and the associated flexibility of the complexes.

To provide structural insights into the mechanism of the NR-MED1 specific association, in the present study we investigated the complex formed between a large fragment of the coactivator MED1 comprising its structured N-terminal region and the RID encompassing two LXXLL motifs and the full NR heterodimer formed by VDR and the retinoid X receptor (RXR). We combined structural methods including small angle X-ray scattering (SAXS), NMR, hydrogen–deuterium exchange coupled to mass spectrometry (HDX-MS), crosslinking mass spectrometry (XL-MS) as well as biophysical methods to characterize the MED1 recruitment by the receptor heterodimer and to get structural details of this assembly. We show that one molecule of MED1 is recruited by the VDR–RXR heterodimer and confirm primary role of the VDR AF-2 interaction with the second LXXLL motif of MED1 in complex formation. We demonstrate that the RXR AF-2 is not essential for the MED1 recruitment, however is affected upon MED1 binding. We also identify other RXR regions, as well as VDR regions outside the AF-2, which are included in the interaction and could be important for reaching the coactivator selectivity by VDR–RXR. Novel structural information on the NR-MED1 complex presented in this work is essential to understand the molecular organization and the interaction networks between complexes of such type.

**MATERIALS AND METHODS**

### Compounds

1α,25-Dihydroxyvitamin D3 (1,25D3) and 9cis retinoic acid (9cisRA) were purchased from Sigma. The rat Nppa single strands DNAs (5′-AGAGGTCTAGAACAGCATT-3′ and 5′ATAGTCCTTTCATGACCCTT-3′) were purchased from Sigma Aldrich and annealed. The MED1 NR2 peptide (NHPMLMNLLKDN) was synthesized by Pascal Eberling (IGBMC peptide synthesis common facility).

### Biochemistry

In all experiments, human proteins (HsVDR and HsRXRα) were used to form the VDR–RXRα complexes, except for surface plasmon resonance experiment where human VDR and mouse RXRα were used to form the complexes.

cDNAs encoding full-length HsVDR (1–427), HsVDRΔ166–216 (1–427, Δ166–216) and HsVDRΔH12 (1–415) cloned into the pET28b vector were used to generate the N-terminal His-tagged proteins. HsRXRαΔNTD (130–462), MmRXRαΔNTD (132–467), MmRXRαΔNTDΔH12 (132–449), MmRXRαΔNTD AF-2Mut (132–467, K289A,E458A) cloned into pET15b, were expressed as N-terminal His-tagged proteins. Recombinant proteins were produced in Escherichia coli BL21 DE3 after induction with 1 mM IPTG (OD$_{600}$ ∼ 0.7) at 23°C for 4 h. Soluble proteins were purified using chromatography column (HisTrap FF crude, 17-5286-01, GE) followed by size exclusion chromatography (SEC) on HiLoad Superdex 200 (28-9893-35 GE) equilibrated in 20 mM Tris–HCl, pH 8.0, 250 mM NaCl, 5% glycerol, 2 mM CHAPS and 1 mM TCEP. Full-length HsVDR and HsRXRαΔNTD were mixed in stoichiometric amounts and purified by size exclusion chromatography (HiLoad Superdex 200, 28-9893-35, GE) equilibrated in 20 mM Tris–HCl, pH 8.0, 200 mM NaCl, 5% glycerol, 2 mM CHAPS and 1 mM TCEP. Ligands (1,25D3 and 9cisRA) were added to the stoichiometric heterodimer and rat NPPA DNA was mixed in a 1:1 equivalent ratio. The DNA complex was further purified by Size Exclusion Chromatography (SEC) in 20 mM Tris pH 8.0, 75 mM NaCl, 75 mM KCl, 2 mM CHAPS, 5% glycerol, 4 mM MgSO$_4$, 1 mM TCEP. A cDNA encoding truncated human MED1 (50–660) cloned into pBacHGW, pFastBac-1 (InVitrogen) baculovirus transfer vector adapted for Gateway, was used to produce HsMED1 proteins with N-terminal His-tag. SF9 cells were infected with recombinant baculovirus at a multiplicity of infection equal 5 and cultured in TNM–FH supplemented with 10% FCS and 50 mg/ml gentamycin at 27°C for 48 h. Cells were harvested by centrifugation.
(1000 g for 15 min) and cell pellets were stored at −20°C prior purification. Soluble protein was purified using batch/gravity-flow affinity chromatography (cOmplete, Roche). MED1 (50–660) was eluted by 250 mM Imidazole in binding buffer. Following the His-tag removal by Thrombin cleavage, the protein was further purified by SEC on HiLoad Superdex 200 (28-9893-35 GE) equilibrated in 20 mM Tris–HCl, pH 8.0, 250 mM NaCl, 5% glycerol, 2 mM CHAPS and 1 mM TCEP.

The proteins were concentrated to 3–6 mg.mL⁻¹ with an Amicon Ultra 30 kDa MWCO. Purity and homogeneity of all proteins were assessed by SDS and Native PAGE.

Gel retardation in TBE

Six percent polyacrylamide gel was used to examine the migration of DNA-bound complexes and MED1. The samples were loaded onto the polyacrylamide gel, placed in a Bio-Rad chamber for gels and ran with constant current of 6 mA for 3 h at 4°C in TBE migration buffer. Gels were revealed by Coomassie staining.

Small angle X-ray scattering

Synchrotron X-ray data were collected on a Pilatus 1M detector at the ESRF beamline BM29 (51). 100 μL of VDR–RXRΔNTD-DNA, MED1 (50–660) and their complex at concentrations 8.5–10 mg.mL⁻¹ in 25 mM HEPES pH 7.5, 150 mM NaCl, 5% glycerol, 2 mM MgCl₂, 2 mM TCEP were loaded onto a GE Healthcare Superdex 200 10/300 column (equilibrated in the same buffer) at a flow rate of 1 mL.min⁻¹. A scattering profile was integrated every second. Frames were selected based on the examination of the SEC profile together with the calculated Rg and Dmax values. The SAXS data were averaged and processed by standard procedures using PRIMUS (52). The forward scattering I(0) and the radii of gyration Rg were evaluated using the Guinier approximation assuming that at very small angles (s < 1.3/Rg) the intensity is represented as I(s) = I(0) exp(−(sRg)²/3). These parameters were also computed from the entire scattering pattern using the indirect transform package GNOM (53) which also provides the maximum dimension of the particle Dmax and the distance distribution function p(r). The program SASREF (54) was employed for molecular rigid body modeling of the VDR–RXR–DNA complex, based on SAXS and cryoEM structures (48,55). The final fits of the model scattering to the experimental data were computed using CRYSOL (56).

Size exclusion chromatography coupled with multi-angle laser light scattering

The molecular weight and homogeneity of the sample was checked using a SEC column coupled with multi-angle laser light scattering (MALLS) Dawn DSP detector (Wyatt Technology, Santa Barbara, CA, USA). A GE Healthcare Superdex 200 10/300 analytical column was pre-equilibrated with the sample buffer, 25 mM HEPES pH 7.5, 150 mM NaCl, 5% Glycerol, 2 mM MgCl₂, 2 mM TCEP. The system was operated at 20°C, with a flow rate of 0.75 mL.min⁻¹.

Analytical ultracentrifugation

Sedimentation velocity experiments were performed at 4°C in 25 mM Tris–HCl pH 8.0, 100 mM NaCl, 2% glycerol, 1 mM CHAPS, 2 mM MgCl₂, 1 mM TCEP using Beckman Coulter Proteome Lab XL-I analytical ultracentrifuge and the 8-hole Beckman An-50Ti rotor. Sedimentation at 50 000 rpm was monitored by absorbance at 280 nm with boundaries measured each 7 min. MED1 (50–660) at constant concentration (6 μM) was titrated by VDR–RXR–DNA; tested coactivator:heterodimer ratios varied from 1:0.5 to 1:2.7. Density and viscosity of the used buffer were calculated using SEDNTERP software (http://sednterp.unh.edu/) and used for the data correction. Using nonlinear least-squares analysis with SEDPHAT (57), collected datasets were fitted using single site hetero-association model.

Hydrogen deuterium exchange coupled to mass spectrometry

HDX experiments of VDR–RXRΔNTD–DNA complex were carried out with and without 2 molar excess of N2R2 motif in 20 mM Tris pH 8.0, 75 mM NaCl, 75 mM KCl, 2 mM CHAPS, 5% glycerol, 4 mM MgSO₄, 1 mM TCEP. The same buffer was used for VDR–RXR–DNA–MED1 (50–660) complex HDX experiment. Preparation and injection of the samples were automatically conducted using a LEAP HDX Automation Manager (Waters), while chromatography was carried out on an Acquity UPLC system with HDX technology (Waters, Manchester, UK). Samples were incubated at different deuteriumation times (0, 0.5, 2, 10 and 30 min) in 95% of deuterated buffer (20 mM Tris pD 8.0, 75 mM NaCl, 75 mM KCl, 4 mM MgSO₄, 1 mM TCEP) before quenching the exchange by adding a 150 mM glycine pH 2.4, 2 M GdHCl, 4 mM MgSO₄, 1 mM TCEP buffer at 1°C during 30s. Digestion of samples (between 20 and 50 pmol injections) was then performed through a pepsin-immobilized cartridge in 0.1% aqueous formic acid solution at a 200 μL/min. Generated Peptides were then trapped on a UPLC pre-column (ACQUITY UPLC BEH C18 VanGuard pre-column, 2.1 mm I.D. × 5 mm, 1.7 μM particle diameter, Waters) and separated on UPLC column (ACQUITY UPLC BEH C18, 1.0 mm I.D. × 100 mm, 1.7 μM particle diameter, Waters) at 0°C. Mass spectrometry analyses were acquired with Synapt G2Si HDMS (Waters) with electrospray ionization, using data-independent acquisition mode (MS²) over an m/z range of 50–2000 and 100 fmol.μL⁻¹ Glu-Fibrinopeptide solution as lock mass correction and calibration. Analyses were performed with the following parameters: capillary voltage, 3 kV; sampling cone voltage, 40 V; source temperature, 80°C; desolation gas, 150°C and 600 L.h⁻¹; scan time, 0.3 s; trap collision energy ramp, 15–40 eV. HDX experiments were realized in triplicate for each time point. Peptide identification was performed using ProteinLynx Global Server 2.5.3 (Waters) with a home-made protein sequence library containing VDR, RXR and MED1(50-660) sequences, with peptide and fragment tolerances set automatically by PLGS, and oxidized methionine set as variable modification. Deuterium uptakes for all identified peptides were then filtered and validated manually using DynamX 3.0 (Waters) as fol-
Crosslinking reactions were conducted with 25 μM protein solutions in 20 mM HEPES pH 8.0, 75 mM NaCl, 75 mM KCl, 4 mM MgSO₄, 1 mM TCEP. Freshly prepared 10 mM stock solution of DSBU and C2-arm version of DSBU in DMSO (CF Plus Chemicals s.r.o., Czech Republic) were added in 50-, 100- and 200-fold molar excess. Crosslinking reactions were conducted during 45 min at room temperature and further quenched during 20 min using NH₄HCO₃ to a final concentration of 20 mM final. Disulfide reduction was next performed by incubating the crosslinked complex solution with 5 mM DTT for 30 min at 60°C, followed by alkylation with 15 mM IAA for 30 min in the dark. Then trypsin (Promega, Madison, USA) was added in 1:50 enzyme: substrate ratio. Samples were incubated overnight at 37°C. Digestion was quenched with 1% formic acid. Peptides were cleaned up using SPE cartridges and samples were concentrated in a SpeedVac concentrator before LC/MS/MS analysis. NanoLC–MS/MS analyses were performed using a nanoAcquity UPLC (Waters, Milford, USA) coupled to the Q-Exactive Plus Orbitrap mass spectrometer (Thermo Scientific, Bremen, Germany) Nanospray Flex™ Ion source. The samples were trapped on a nanoACQUITY UPLC precolumn (C18,180 μm × 20 mm, 5 μm particle size), and the peptides were separated on a nanoACQUITY UPLC column (C18, 75 μm × 250 mm with 1.7 μm particle size, Waters, Milford, USA) maintained at 60°C. The samples were first injected with a 285 min gradient and a flow rate of 450 nL min⁻¹. The Q-Exactive Plus Orbitrap source temperature was set to 250°C and spray voltage to 1.8 kV. Full scan MS spectra (300–1800 m/z) were acquired in positive mode at a resolution of 140 000, a maximum injection time of 50 ms and an AGC target value of 3 × 10⁶ charges, with lock-mass option being enabled (polysiloxane ion from ambient air at 445.12 m/z). The 10 most intense multiply charged ions per full scan (charge states > 2) were isolated using a 2 m/z window and fragmented using higher energy collisional dissociation (30 normalized collision energy, ±3%). MS/MS spectra were acquired with a resolution of 35 000, a maximum injection time of 100 ms, an AGC target value of 1 × 10⁵ and dynamic exclusion was set to 60 s. The system was fully controlled by XCalibur software v3.0.63, 2013 (Thermo Scientific) and NanoAcquity UPLC console v1.51.3347 (Waters). Raw data collected were processed and converted into .mgf format. The MS/MS data were analyzed using MeroX software version 1.6.6 (60). Mass tolerance of 5 ppm for precursor ions and 10 ppm for product ions were applied. A 5% FDR cut-off and a signal-to-noise ≥ 2 were applied. For both crosslinkers, Lys and Arg were considered as protease cleavage sites with a maximum of three missed cleavages. Carbamidomethylation of cysteine was set as fixed and oxidation of methionine as variable modifications (max. mod. 2). Primary amino groups (Lys side chains and N-termini) as well as primary hydroxyl groups (Ser, Thr and Tyr side chains) were considered as crosslinking sites. The cRap database was used in combination with the reporter ion scan event (RISE) mode on. Crosslinks composed of consecutive amino acid sequences were not considered. Each crosslinked product automatically annotated with MeroX was manually validated. Finally, PyMOL software (www.pymol.org) was used to calculate the Cα–Cα distance of each validated linkage sites. The XL-MS data set has been deposited to the ProteomeXchange Consortium via the PRIDE (59) partner repository with the dataset identifier PXD019530.
RESULTS

Binding of MED1 N-terminal domain to VDR–RXR

Previous structural studies uncovered details of the VDR LBD interaction with the second LXXLL motif of MED1 (NR2 motif, residues 645–649) (32,61) (Supplementary Figure S1a), and we have previously shown that the MED1 RID binds to the VDR–RXR heterodimer asymmetrically and remains flexible (47). To gain further insights into specific association between VDR–RXR and MED1, we investigated the binding of the receptor heterodimer to a larger fragment of the coactivator. Based on the disorder prediction for MED1 (Supplementary Figure S1a, b), we selected a protein construct spanning from residues 50–660 encompassing the structured N-terminus and RID, as previous studies suggested that first 570 residues of MED1 are sufficient for its incorporation into the Mediator (62).

We investigated how purified MED1 (50–660) fragment interacts with the heterodimer formed by the full-length VDR and the RXRα lacking the flexible N-terminal domain (NTD), named hereafter VDR–RXR, and associated with a DR3-type vitamin D response element (VDRE). To stabilize the VDR–RXR heterodimer, we selected the VDRE from rat Nppa gene alternatively named rat ANF1 gene (63) displaying a low nanomolar affinity for VDR–RXR complex (Supplementary Figure S2a). In addition, we showed by a transactivation assay in HEK 293 cells that the luciferase gene under control of the rat Nppa promoter is activated by the VDR natural ligand, 1,25D3, at nanomolar concentrations (Supplementary Figure S2b). The complex was formed by an overnight incubation of the purified VDR–RXR bound with their cognate ligands, 1,25D3 and 9cisRA, and the purified MED1 fragment. The presence of the two ligands in the VDR–RXR complex was confirmed by native electrospray ionization mass spectrometry (data not shown). The complex formation with MED1 was observed by gel retardation assay (Supplementary Figure S2c) and size-exclusion chromatography (SEC), where the newly formed high molecular weight complex could immediately be detected as a new specimen appearing on the elution profile before the MED1 (68 kDa) and the VDR–RXR (86 kDa) (Supplementary Figure S2d, e). The molecular weight of the complex as measured by multi-angle light scattering was in close agreement with the respective calculated molecular weight of complex where MED1 binds to VDR–RXR in a 1:1 stoichiometry (Figure 1A). The binding mode of

Figure 1. MED1 forms a complex with VDR–RXR–DNA. (A) Size-exclusion chromatography-coupled multi-angle laser light scattering (SEC-MALLS) of MED1, VDR–RXR–DNA and VDR–RXR–DNA–MED1 complexes showing the elution profile on a SEC S200 10/300 with the direct molar mass measurement of each elution peak. (B) Analytical ultracentrifugation: c(s) distributions. Ratio MED1:VDR–RXR–DNA is indicated and is used for the color representation of the distributions. Left side of the graph corresponds to the top, and right side–to the bottom of the sample cell, which correlates with the direction of migration. (C) Analysis of the interactions of VDR–RXR wild type and VDR–RXRΔH12, VDR–RXRAF2 Mutant and VDRΔ12-RXR mutants with MED1 (50–660) in the presence of 1,25D3 and 9cisRA by surface plasmon resonance and calculated KD.
MED1 to VDR–RXR–DNA was further analyzed by analytical ultracentrifugation in sedimentation velocity mode (Supplementary Figure S3). The distribution of sedimentation coefficients $c(s)$ (Figure 1B) confirmed the formation of the complex between the MED1 and VDR–RXR–DNA and the complex stoichiometry. Interestingly, the $c(s)$ peak shifted in a concentration-dependent way indicating fast kinetics of the interaction ($k_{off} > 10^{-2}$ s$^{-1}$).

All together, the biophysical data indicates that only one MED1 molecule binds to the liganded VDR–RXR heterodimer and we show that the process of the protein association–dissociation is dynamic and the complex is rather transient.

**Topology of the MED1-VDR–RXR–DNA complex**

To gain better insights into the overall structures of the multi-modular VDR–RXR–DNA and VDR–RXR–DNA–MED1 complexes, we determined their solution structures using SAXS. SAXS data were collected from samples of MED1, liganded VDR–RXR–DNA and liganded VDR–RXR–DNA–MED1 using on-line SEC (SEC-SAXS) (Supplementary Figure S4). The SAXS profiles are shown in Figure 2A, B and Supplementary Figure S5 and the structural parameters including the radius of gyration, $R_g$, and the maximum particle dimension, $D_{max}$, are reported in Table 1. For MED1 alone, the Kratky plot representation of MED1 scattering data (Supplementary Figure S5c) suggests that the fragment is globular and has one core with flexible parts/linkers. The structural parameters values for VDR–RXR–DNA are slightly larger than those previously determined for the related VDR$\Delta$-RXR–DNA complex where VDR-specific insertion localized between H1 and H3 was truncated (47). Binding of MED1 to VDR–RXR–DNA increases the average size of the complex but does not induce a large change in the overall SAXS profile: similar distinctive ‘humps’ around 1 nm$^{-1}$ are observed in the SAXS curves for VDR–RXR–DNA alone and in complex with MED1 (Figure 2A, C). The probability distribution of real-space scattering pair distances, or $p(r)$ profiles, reveals a similar shoulder around 60–80 Å for both complexes (Figure 2B). This indicates that the shape of the VDR–RXR–DNA in complex with MED1 remains similar to that of the DNA-bound heterodimer, where the DNA binding domains (DBDs) are spatially separated from the LBDS. Comparison of the most representative ab initio models of VDR–RXR–DNA and VDR–RXR–DNA–MED1 complexes obtained with DAMMIN (64) reveals overall shape similarities with two distinguishable domains (Figure 2C) indicating no major conformational change of the DNA-bound heterodimer upon MED1 interaction. For the coactivator complex, an additional electron density at the region occupied by both LBDS is visible indicating that the globular domain of MED1 is likely located on ‘top’ of the heterodimer and is interacting with the VDR–RXR LBD heterodimer via an extensive area.

**VDR insertion domain modulates MED1 interaction with the VDR–RXR heterodimer**

Whereas the ab initio SAXS envelope visualizes the overall shapes, the rigid-body refinement provides a model that reflects the overall distribution of conformers in solution and is not restricted to a particular low-energy conformation of the macromolecules. For the VDR–RXR–DNA complex, we built an ensemble of SAXS compatible models (Figure 2D) using the crystal structures of the DNA and ligand binding domains of VDR and RXR (65–67) with missing regions (VDR NTD, hinges, VDR's insertion) modeled as dummy residues. The insertion region in VDR LBD is a 50 amino acid domain specific for VDR and poorly conserved between VDR family members and disordered in context of the isolated LBD (68). In the obtained refined models of VDR–RXR–DNA (Figure 2A), it occupies a defined region of space similar in all refined models suggesting that the insertion domain although possibly flexible is not totally disordered. Modelling also suggests that the VDR NTD (residues 1–23) is rather flexible, adopting various extended conformation in solution and not interacting with the DNA.

The best refined model docked into the SAXS envelope of VDR–RXR–DNA–MED1 (Figure 2E) reveals a proximity and possible overlap of the areas occupied by the VDR insertion domain and MED1, thus suggesting that this VDR region may be interacting with MED1. To further characterize the involvement of the disordered VDR insertion domain in MED1 association, a NMR analysis of N$^{15}$-labeled VDR complexes was performed. As VDR exhibits two disordered regions, the short NTD and the insertion domain, we compared the $^1$H, $^{15}$N HSQCs recorded for DNA bound heterodimer where VDR was either full-length or truncated of its insertion domain, with or without addition of MED1. The size of the complexes filters out signals from folded regions leaving only amide resonances from disordered regions belonging to the NTD and insertion in full-length LBD (Figure 2F, G). The addition of MED1 to VDR–RXR–DNA complex led to the specific disappearance of a small number of cross peaks specifically found in the full-length VDR indicating that the disordered insertion is involved, either directly or indirectly, in the interaction with MED1.

**Roles of VDR and RXR AF-2 in association with MED1**

To analyze the roles of the VDR and RXR AF-2 in interaction with MED1 (50–660) fragment, we mutated VDR and RXR and determined the impact of mutations on the association with MED1 by surface plasmon resonance. MED1 fails to interact to VDR–RXR when VDR H12 is deleted, or in presence of VDR antagonist ligand (ZK168281) (69) (Figure 1C and Supplementary Figure S8), thus confirming that the main anchoring interaction are agonist-dependent and mediated through VDR AF-2, in agreement with previous studies showing that a synthetic peptide comprising classical NR-interacting LXXLL motif competes with MED1 for interaction with the VDR (70). In contrast, deletion of RXR H12 does not prevent MED1 binding to VDR–RXR–DNA. A slightly increased MED1 binding is observed for VDR–RXRDeltaH12. Mutations of RXR residues forming a charge clamp for proper orientation and binding of the LXXLL motif of the coactivator (71,72) has no effect on MED1 interaction.
Figure 2. Solution structure of MED1 complex with VDR–RXR–DNA. (A) Averaged SEC-SAXS data for VDR–RXR–DNA (blue) and VDR–RXR–DNA–MED1 (green) complexes. (B) Corresponding p(r) profiles calculated from the SAXS data. (C) Representative ab initio molecular envelopes for VDR–RXR–DNA–MED1 complex (green mesh) with superimposed ab initio molecular envelope of VDR–RXR–DNA (blue mesh). (D) SAXS-based ensemble models (4 conformations) of the VDR–RXR–DNA complex. (E) Best SAXS model of VDR–RXR–DNA fitted into the SAXS envelope of VDR–RXR–DNA–MED1. (F) 1H–15N HSQC spectra of the VDR–RXR–DNA complex lacking the VDR LBD insertion before (blue) and after addition of stoichiometric amount of MED1 (red). (G) 1H–15N HSQC spectra of the VDR–RXR–DNA complex harboring the full-length VDR before (blue) and after addition of stoichiometric amount of MED1 (red). The red stars indicate correlation peaks whose intensities are affected by the addition of MED1.
Table 1. SAXS parameters, $R_g$ and $D_{max}$ as determined from Guinier plot or $p(r)$ distribution

| Sample                  | $R_g$, Å (from GNOM) | $R_g$, Å (from GNOM) | $D_{max}$, Å (from GNOM) |
|-------------------------|----------------------|----------------------|--------------------------|
| VDR–RXR–DNA–MED1        | 63.8 ± 2.3           | 63.6 ± 0.13          | 202                      |
| VDR–RXR–DNA             | 42.8 ± 0.14          | 43.3 ± 0.1           | 135                      |
| MEDI                    | 41.5 ± 0.27          | 41.6 ± 0.12          | 140                      |

In addition, we investigated the impact of the binding of the peptide bearing the MED1 NR2 motif (residues 645–649) on the VDR–RXR–DNA complex using HDX-MS. We observed that two regions of VDR corresponding to the NTD and the insertion domain exhibit fast H/D exchange rates (Supplementary Figure S6a) which is a typical phenomenon for highly flexible regions (73), thus supporting SAXS results. Similar fast HDX rates were also observed for several regions of RXR (Supplementary Figure S6b), encompassing the hinge region, helix H1, H2, the C-terminus of H3, H11, the C-terminus of H12, and the region after H12. The comparison of relative fractional uptakes (RFU) of VDR–RXR–DNA and VDR–RXR–DNA–MED1 NR2 motif revealed that mainly VDR region 411–419 spanning the C-terminus of H11n and the N-terminus of the helix H12 was protected from H/D exchange upon NR2 motif binding (Figure 3A and Supplementary Figure S7a). These results are in agreement with the crystal structures of VLD LBD complexes with coactivator peptides (74,75), highlighting the role of VDR AF-2 in its interaction with MED1 NR2 through H3, H4 and H12 helices. On the contrary, no significant differences were detected for RXR upon MED1 NR2 motif binding (Figure 3 and Supplementary Figure S7b), supported by the low affinity of MED1 NR2 for RXR (72,76). Taken together, our data show that the AF-2 of VDR but not RXR is directly involved in agonist-induced binding of MED1.

Effect of MED1 (50–660) binding on the VDR–RXR heterodimer

We next performed HDX-MS experiments with the larger construct of MED1 to determine the effect of binding on both the coactivator and the VDR–RXR heterodimer. HDX-MS of MED1 (50–660) fragment revealed that the protein is mostly structured (Supplementary Figure S9a), in agreement with the disorder prediction (Supplementary Figure S9b). Several MED1 regions exhibit fast H/D exchange rates and can be characterized as highly flexible, including residues 61–74, 161–174, 191–201, 231–255, 276–296, 327–339 and 480–487 which could correspond to flexible loops between the secondary structure elements (Supplementary Figure S9c). MED1 RID is also highly flexible, in agreement with our previous data (47).

We compared HDX-MS RFU values of VDR and RXR in VDR–RXR–DNA complex with those in VDR–RXR–DNA–MED1 (50–660) state to characterize their conformational dynamics upon MED1 binding. As expected, C-terminus of H11n and N-terminus of H12 of VDR (protected upon MED1 NR2 binding) were similarly affected upon MED1 (50–660) binding (Figure 3B, C and Supplementary Figure S10a). Additional regions of VDR were protected, including helices H3, the C-terminus of H5, H6, H7, H10 and H11. These regions are spatially close to the VDR AF-2 domain (Figure 3B), revealing a higher protection effect on this region upon MED1 (50–660) binding compared to the binding of MED1 NR2 motif.

Interestingly, several protected regions of RXR LBD were identified upon MED1 (50–660) binding, corresponding to helices H3, H5 and the β-strand, H11 and the N-terminus of H12 (Figure 3B, D and Supplementary Figure S10b). These regions are spatially close to each other and are located on ‘top’ of the heterodimer LBDs supporting the SAXS models indicating that this region creates a large MED1 interaction surface. Of note, region 419–429 of RXR, covering helix H10 and comprising the heterodimerization interface, shows deprotection upon MED1 binding at shorter time points suggesting a conformational change leading to its higher flexibility (Figure 3B and Supplementary Figure S9b).

MED1 (50–660) is also protected upon formation of the complex with VDR–RXR–DNA (Supplementary Figure S11). Residues 243–255 of the structured N-terminal domain of MED1 are particularly protected upon VDR–RXR–DNA binding (Figure 3E). Other affected regions are 74–107, 123–150, 191–201, 509–527 as well as regions 560–604, 636–645 and 649–657 situated within the unstructured part of MED1 comprising RID. Residues 560–604 and 636–645 encompass the first leucine residues of both NR boxes 1 and 2 of MED1 RID domain respectively (Leu604 and Leu645), suggesting a stabilization of both motifs upon NR binding.

All together, these results show that MED1 (50–660) binding affects extended regions within the liganded VDR–RXR LBD heterodimer, in contrast to the MED1 NR2 motif stabilizing solely the C-terminus of VDR. In addition to the VDR coactivator cleft and H12 strongly stabilized upon MED1 binding, several RXR regions including AF-2 and H5/β-strand are affected indicating role of RXR in establishing a specific association with the coactivator.

Investigation of the interaction surface between MED1 and VDR–RXR heterodimer

To determine molecular constraints and amino acids of VDR–RXR and MED1 located in close proximity, we next performed XL-MS on VDR–RXR–DNA–MED1 (50–660) complex using DSBU (spacer arm 12.5 Å) (77) and a C2-arm version of the DSBU (spacer arm 6.2 Å) (see Material and Methods). Both crosslinkers are MS-cleavable, allowing more confident MS/MS identification and validation thanks to the detection of characteristic doublet peaks along with the peptide backbone fragments (78). Both DSBU crosslinker versions target primary amines as well as hydroxyl groups and can bridge residues with Cα–Ca distances up to 26–30 Å and 20–24 Å for DSBU and the C2-arm, respectively (79). We identified 42 intra- and inter-crosslinked peptides: 11 crosslinks intra-RXR, 12 intra-VDR, 6 intra-MED1, 12 inter-VDR–RXR and 1 inter-MED1–RXR (Figure 4 and Supplementary Figure S12a). Twelve crosslinks involving VDR and RXR were found in the proximity to or within the DBDs when mapped on the...
Figure 3. HDX-MS analysis of the VDR–RXR–DNA interaction with MED1. (A) Relative fractional uptake (RFU) differences between VDR–RXR–DNA and VDR–RXR–DNA–MED1 NR Box 2 complexes mapped on the representative SAXS model of the VDR–RXR–DNA after 0.5 min of deuteration. (B) Relative fractional uptake (RFU) differences between VDR–RXR–DNA and VDR–RXR–DNA–MED1 (50–660) complexes mapped on the representative SAXS model of the VDR–RXR–DNA after 0.5 min of deuteration. (C) Deuterium uptake of the peptides 224–230 (H3), 385–389 (H10) and 413–419 (H11–12) from VDR perturbed upon MED1 (50–660) binding plotted as a function of deuteration time. (D) Deuterium uptake of the peptides 271–278 (H3), 326–330 (β) and 450–454 (H12) from RXR perturbed upon MED1 (50–660) binding plotted as a function of deuteration time. (E) Deuterium uptake of selected MED1 peptides perturbed upon binding to VDR–RXR–DNA: 243–255 from the structured N-terminal part, 595–604 comprising Leu604 of the NR Box 1 and 636–645 comprising Leu645 from the NR Box 2.
Figure 4. Crosslinked sites observed for the VDR–RXR–DNA–MED1(50–660) complex. (A) Identified VDR–RXR-MED1 (50–660) crosslinks representation. Red: NR boxes 1 and 2 of MED1; purple: LBD region; grey: hinge region; pink: DBD region for each nuclear receptor, VDR and RXR. Blue lines represent inter-protein crosslinks and purple curved lines intra-protein crosslinks. (B) Representation of Lys321 of RXR, identifies as crosslinked to Thr236 of MED1, within the VDR–RXR LBD heterodimer.
VDR–RXR heterodimer model (55) and with a Co–Co distance below the cut-off mentioned, increasing confidence in our XL-MS approach (Supplementary Figure S12b). Interestingly, an inter-protein crosslink involving Thr236 of MED1 and Lys321 of RXR was identified. Lys321 is located in the β-strand between H5 and H6 of RXR LBD (Figure 4B) on ‘top’ of the LBD heterodimer and within the region protected from the H/D exchange upon MED1 binding correlating with the SAXS and HDX-MS results. Taken together, our data suggests that although RXR H12 is not required for MED1 recruitment, RXR LBD possesses an extensive surface directly interacting with MED1.

DISCUSSION

While structural requirements for ligand-dependent binding of classical coactivator motifs to NRs have been fully elucidated (reviewed in (80)), the molecular mechanism of co-activation remains poorly understood. In particular, studies providing structural insights into recognition of full-length or truncated coactivators by full NR complexes remain extremely scarce (47,49,50,81). Here, we describe novel structural data (Figure 5) on the complex formed between the agonist- and DNA-bound VDR–RXR heterodimer and a large fragment of the classical nuclear receptor coactivator MED1, an important regulator of VDR activation (85). Our results confirm that MED1 NR2-VDR interaction is driving the complex formation. However, by using a larger fragment of MED1, MED1 (50–660), we have also demonstrated that other VDR–RXR regions outside the VDR AF-2 as well as MED1 regions other than RID modulate the association and form an extended interaction surface.

Both LXXLL motifs of MED1, as well as proper spacing between the motifs, were reported to be required for optimal MED1 binding to DNA-bound VDR–RXR heterodimer (32). It was suggested that each LXXLL motif is recognized by VDR and RXR coactivator binding clefts simultaneously when the complex with MED1 is formed. In this case the 35 amino acid spacer between the two LXXLL motifs has to clasp around the LBDs as the coactivator binding sites are located on the opposite sites of the heterodimer. However, we have previously shown that the MED1 RID binds to the VDR–RXR heterodimer asymmetrically and remains flexible (47). Interestingly, in the present study, we observed the perturbation of both LXXLL motifs of MED1 (50–660) upon formation of the complex with VDR–RXR, suggesting that while the NR1 box is not accommodated within the classical coactivator binding site, it could be either interacting with an alternative site of the receptors or stabilized allosterically. Similar observations have previously been made for the three NR boxes of the SRC-2 RID binding to PPARG-RXR (86). Alternative interaction with androgen receptor AF-1 was previously described for two noncanonical α-helical motifs of MED1 located between residues 505 and 537 and in proximity to the NR box 1 (87).

Here we demonstrate that, in addition to RID, the structured N-terminal domain of MED1 is also affected upon binding to VDR–RXR and is likely interacting with both VDR and RXR LBDs. In particular, MED1 region 243–255 is largely stabilized in the complex with the receptor heterodimer. Neighboring Thr236 of MED1 was identified within inter-MED1-RXR crosslink, suggesting that this MED1 region is in physical proximity to the RXR β-strand which, in turn, is also perturbed upon the interaction. As N-terminal region of MED1 is involved in important downstream interactions such as incorporation into Mediator (62) and recruitment of alternative cofactors to enhancer-bound NRs (26), it is tempting to speculate that by creating an extended surface for MED1 accommodation and altering its conformation, RXR could be important for achieving optimal MED1-mediated transcription activation.

Among other novel MED1-interacting regions within the VDR–RXR heterodimer is the flexible insertion domain in the VDR LBD located between H1 and H3. By using 15N NMR, we show that it undergoes conformational changes upon interaction with MED1. This effect is not seen in the HDX-MS experiment; however the observed difference could be attributed to a different temporal resolution of the two methods. While the VDR insertion domain does not play a major role in receptor selectivity for 1,25D3 (70) nor bile acids (88), previously published data suggests that it has a functional role in regulation of VDR signaling. Several phosphorylations modulating VDR transcriptional activity, S182 and S208, have been identified within
this region (89). Interestingly, S208 phosphorylation has previously been reported to enhance VDR interaction with MED1 but not with the SRC-1 coactivator (90,91).

In this study we used the full VDR–RXR heterodimer in agonist-bound form, where both VDR and RXR ligands were present. Hypothetically, each receptor can recruit a coactivator, however, we demonstrate that only one molecule of MED1 is recruited by the VDR–RXR heterodimer, suggesting that H12 of RXR is not essential for the complex formation. Indeed, truncation of RXR H12 and mutations in RXR charge clamp do not prevent MED1 interaction, and binding of a peptide comprising MED1 NR2 does not induce any change in RXR as seen by HDX-MS. However, using HDX-MS, we identified H3, H11 and the N-terminus of H12 comprising the AF-2 among RXR regions largely stabilized upon MED1 binding. The observed stabilization event could originate from direct non-canonical interaction with MED1 as well as from a distal allosteric effect.

As MED1 is a classical NR coactivator, it is recognized by NRs similarly to other coregulators, e.g. SRCs, and their binding sites overlap. Interestingly, significant differences could be observed between the VDR–RXR complex with MED1 described here and the analogue complexes with the SRCs. In presence of 1,25D3 only RXR AF-2 within the full VDR–RXR complex was insensitive to the binding of SRC-1 RID, indicating that it is primarily associated to VDR AF-2 (92). However, RXR and its ligand modulate the interaction and in the presence of both VDR and RXR agonists SRC-1 RID binding has been shown to stabilize VDR AF-2 as well as RXR H3 (227–273) and H10-H11 (433–451) (81). Differences in binding mode between SRCs and MED1 were previously suggested for ER where a single mutation differently affects MED1 and SRC-2 RID interaction (93), or for PPAR-RXR where RXR ligand only induces SRCs but not MED1 binding to RXR (83). Such differences in the binding modes could serve as molecular determinants of how the NRs discriminate between coactivators and sequentially recruit them.

This work contributes to a growing number of studies revealing the complexity of coactivator binding to NRs. Extended characterization of allosteric mechanism within large NR coregulator complexes should increase the potential of novel targets for drug design and discovery programs.

DATA AVAILABILITY
HDX-MS data have been deposited to the ProteomeXchange Consortium via the PRIDE partner repository with the dataset identifier PXD019530.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

ACKNOWLEDGEMENTS
We thank the staff of BM29 at ESRF for assistance in using the beamline. We thank the IGBMC cell culture service, Isabelle Kolb-Channel and Nathalie Troffer-Charlier (IGBMC) for insect cell production, Pascal Eberling (IGBMC) for peptide synthesis, Pierre Poussin-Courmontagne (IGBMC) for assistance with SEC-MALLS, Camille Costmann (IGBMC) for assistance with surface plasmon resonance, Catherine Birck (IGBMC) for help with analytical ultracentrifugation data collection, as well as Carole Peluso-Ilits (IGBMC) for excellent technical assistance.

Author contributions: A.Y.B. and S.C. purified the proteins and performed biochemical and structural analyses. M.B., S.H. and S.C. performed the structural mass-spectrometry experiments and analyzed the data. B.K. performed the NMR analysis. N.R. designed and supervised the study. A.Y.B. and N.R. wrote the manuscript with contribution from all the authors. All authors have given approval to the final version of the paper.

FUNDING
Agence Nationale de la Recherche (ANR) [ANR-13-BSV8-0024-01]; French Infrastructure for Integrated Structural Biology [FRIISBI ANR-10-INBS-05, Instruct-ERIC, ANR-10-LABXX-0030-INRT]; French State fund managed by the Agence Nationale de la Recherche under the program Investissements d’Avenir [ANR-10-IDEX-0002]; French Proteomic Infrastructure [ProFI ANR-10-INBS-08-03]; GIS IBiSA and Région Alsace for financial support in purchasing a Synapt G2SI HDMS instrument; M.B. was supported by a fellowship from the Région Alsace; A.Y.B. was supported by a FRM fellowship [FDT20140930978]. Funding for open access charge: Agence Nationale de la Recherche.

Conflict of interest statement. None declared.

REFERENCES
1. Lonard,D.M. and O’Malley,B.W. (2012) Nuclear receptor coregulators: modulators of pathology and therapeutic targets. Nat. Rev. Endocrinol., 8, 598–604.
2. Giudici,M., Goni,S., Fan,R. and Treuter,E. (2015) Nuclear Receptor Coregulators in Metabolism and Disease. In: Herzog, S. (ed). Metabolic Control. Handbook of Experimental Pharmacology. Springer, Vol. 233.
3. Jeronimo,C. and Robert,F. (2017) The mediator complex: at the nexus of RNA polymerase II transcription. Trends Cell Biol., 27, 765–783.
4. Soutourina,J. (2018) Transcription regulation by the mediator complex. Nat. Rev. Mol. Cell Biol., 19, 262–274.
5. Verger,A., Monté,D. and Villeret,V. (2019) Twenty years of mediator complex structural studies. Biochem Soc Trans., 47, 399–410.
6. Kelleher,R.J., Flanagan,P.M. and Kornberg,R.D. (1990) A novel mediator between activator proteins and the RNA polymerase II transcription apparatus. Cell., 61, 1209–1215.
7. Flanagan,P.M., Kelleher,R.J., Sayre,M.H., Tschochner,H. and Kornberg,R.D. (1991) A mediator required for activation of RNA polymerase II transcription in vitro. Nature, 350, 436–438.
8. Fondell,J.D., Ge,H. and Roeder,R.G. (1996) Ligand induction of a transcriptionally active thyroid hormone receptor coactivator complex. Proc. Natl. Acad. Sci. U.S.A., 93, 8329–8333.
9. Rauchez,C., Suldan,Z., Ward,J., Chang,C.P., Burakov,D., Erdjument-Bromage,H., Tempst,P. and Freedman,L.P. (1998) A novel protein complex that interacts with the vitamin D3 receptor in a ligand-dependent manner and enhances VDR transactivation in a cell-free system. Genes Dev., 12, 1787–1800.
10. Niäär,A.M., Beaurang,P.A., Zhou,S., Abraham,S., Solomon,W. and Tjian,R. (1999) Composite co-activator ARC mediates chromatin-directed transcriptional activation. Nature, 398, 828–832.
11. Jiang,Y.W., Veschambre,P., Erdjument-Bromage,H., Tempst,P., Conaway,J.W., Conaway,R.C. and Kornberg,R.D. (1998)
Mammalian mediator of transcriptional regulation and its possible role as an end-point of signal transduction pathways. *Proc. Natl. Acad. Sci. U.S.A.*, *95*, 8538–8543.

12. Boyer,T.G., Martin,M.E., Lees,E., Ricciardi,R.P. and Berk,A.J. (1999) Mammalian Srb/Mediator complex is targeted by adenovirus E1A protein. *Nature*, *399*, 276–279.

13. Malik,S., Gu,W., Wu,W., Qin,J. and Roeder,R.G. (2000) The USA-derived transcriptional coactivator PC2 is a submodule of TRAP/SMCC and acts synergistically with other PC2s. *Mol. Cell.*, *5*, 753–760.

14. Ryu,S. and Tjian,R. (1999) Purification of transcription cofactor complex CRSP. *Proc. Natl. Acad. Sci. U.S.A.*, *96*, 7137–7142.

15. Yuan,C.X., Ito,M., Fondell,J.D., Fu,Z.Y. and Roeder,R.G. (1998) Mammalian Srb complex CRSP. *Mol. Cell.*, *3*, 510–519.

16. Nolte,R.T., Wisely,G.B., Westin,S., Cobb,J.E., Lambert,M.H., Spiegelman,B.M. and Roeder,R.G. (2002) Transcription coactivator TRAP220 is required for PPAR gamma 2-stimulated adipogenesis. *Proc. Natl. Acad. Sci. U.S.A.*, *99*, 7399–7404.

17. Raceczek,C., Lemon,B.D., Suldan,Z., Bromleigh,V., Gamble,M., Nänä,A.M., Erjument-Bromage,H., Tempst,P. and Freedman,L.P. (1999) Differential regulation of glucocorticoid receptor function. *Nature*, *398*, 824–828.

18. Kagey,M.H., Guermah,M., Yuan,C.X., Ito,M., Fondell,J.D., Fu,Z.Y. and Roeder,R.G. (1998) The TRAP220 component of a thyroid hormone receptor-associated protein (TRAP) coactivator complex interacts directly with nuclear receptors in a ligand-dependent fashion. *Proc. Natl. Acad. Sci. U.S.A.*, *95*, 7399–7404.

19. Ge,K., Guermah,M., Yuan,C.X., Ito,M., Fondell,J.D., Fu,Z.Y. and Roeder,R.G. (1999) Mammalian Srb/Mediator of transcriptional regulation and its possible role as an end-point of signal transduction pathways. *Proc. Natl. Acad. Sci. U.S.A.*, *96*, 7137–7142.

20. Malik,S., Gu,W., Wu,W., Qin,J. and Roeder,R.G. (2000) The USA-derived transcriptional coactivator PC2 is a submodule of TRAP/SMCC and acts synergistically with other PC2s. *Mol. Cell.*, *5*, 753–760.

21. Boyer,T.G., Martin,M.E., Lees,E., Ricciardi,R.P. and Berk,A.J. (1999) Mammalian Srb/Mediator complex is targeted by adenovirus E1A protein. *Nature*, *399*, 276–279.

22. Wang,Q., Sharma,D., Ren,Y. and Fondell,J.D. (2002) A coregulatory complex CRSP. *Mol. Cell.*, *3*, 510–519.

23. Kim,J.H., Yang,C.K., Heo,K., Roeder,R.G., An,W. and Stallcup,M.R. (2008) CCAR1, a key regulator of mediator complex super-enhancers links phase separation and gene control. *Science*, *323*, 6400.

24. Chen,W., Zhang,X., Birsoy,K. and Roeder,R.G. (2010) A muscle-specific knockout implicates nuclear receptor coactivator MED1 in the regulation of glucose and energy metabolism. *Proc. Natl. Acad. Sci. U.S.A.*, *107*, 10196–10201.

25. Bai,L., Jia,Y., Viswakarma,N., Huang,J., Viuggens,A., Wolins,N.E., Jafari,N., Rao,M.S., Borensztajn,J., Yang,G. et al. (2011) Transcriptional coactivator mediator subunit MED1 is required for the development of fatty liver in the mouse. *Hepatology*, *53*, 1164–1174.

26. Weber,H. and Garabedian,M.J. (2018) The mediator complex in genomic and non-genomic signaling in cancer. *Steroids*, *133*, 8–14.

27. Osz,J., Breilhet,Y., Peluso-Ilitsi,C., Cura,V., Eiler,S., Ruff,M., Bourguet,W., Rochel,N. and Moras,D. (2012) Structural basis for a molecular allosteric control mechanism of cofactor binding to nuclear receptors. *Proc. Natl. Acad. Sci. U.S.A.*, *109*, E588–E594.

28. Rochel,N., Ciesielski,F., Godejot,J., Moman,E., Roessle,M., Peluso-Ilitsi,C., Moulin,M., Haertlein,M., Callow,P., Mély,Y. et al. (2011) Common architecture of nuclear receptor heterodimers on DNA direct repeat elements with different spacings. *Nat. Struct. Mol. Biol.*, *18*, 564–570.

29. Pavlin,M.R., Brunzelle,J.S. and Fernandez,E.J. (2014) Agonist ligands mediate the transcriptional response of nuclear receptor heterodimers through distinct stoichiometric assemblies with coactivators. *J. Biol. Chem.*, *289*, 24771–24778.
49. Yi, P., Wang, Z., Feng, Q., Pintilie, G.D., Foulds, C.E., Lanz, R.B., Ludtke, S.J., Schmid, M.F., Chiu, W. and O’Malley, B.W. (2015) Structure of a biologically active estrogen receptor-coactivator complex on DNA. *Nucleic Acids Research*, 2015.

50. Yi, P., Wang, Z., Feng, Q., Chou, C.K., Pintilie, G.D., Shen, H., Foulds, C.E., Fan, G., Serysheva, I., Ludtke, S.J. et al. (2017) Structural and Functional Impacts of ER Coactivator Sequential Recruitment. *Nucleic Acids Research*, 67, 733–743.

51. Pernot, P., Theveneau, P., Giraudt, T., Nogueira Fernandes, R., Nurizzo, D., Spruce, D., Serry, A., McSweeney, S., Round, A., Felisaz, F. et al. (2010) New beamline dedicated to solution scattering from biological macromolecules at the ESRF. *J. Phys.: Conf. Ser.*, 2010.

52. Svergun, D.I. (1992) Determination of the regularization parameter in small-angle scattering data analysis. *J. Appl. Cryst.*, 36, 1277–1282.

53. Svergun, D.I. (1992) Determination of the regularization parameter in indirect-transform methods using perceptual criteria. *J. Appl. Cryst.*, 25, 495–503.

54. Petrov, M. V. and Svergun, D. I. (2005) Global rigid body modeling of macromolecular complexes against small-angle scattering data. *Biophys. J.*, 89, 1237–1250.

55. Orlov, I., Rochel, N., Moras, D. and Klaholz, B.P. (2012) Structure of the full human RXR/VDR nuclear receptor heterodimer complex with its DR3 target DNA. *EMBO J.*, 31, 291–300.

56. Svergun, D.I., Barberato, C. and Koch, M.H.J. (1995) CRYSOL - a program to evaluate X-ray solution scattering of biological macromolecules from atomic coordinates. *J. Appl. Cryst.*, 28, 768–773.

57. Schuck, P. (2000). Size-distribution analysis of macromolecules from solution scattering using simulated annealing. *J. Biol. Chem.*, 275, 1237–1250.

58. Svergun, D.I., Barberato, C. and Koch, M.H.J. (1995) CRYSOL - a program to evaluate X-ray solution scattering of biological macromolecules from atomic coordinates. *J. Appl. Cryst.*, 28, 768–773.

59. Perez-Riverol, Y., Csordas, A., Bai, J., Bernal-Llinares, M., Eisenacher, M. and Lepretre, J.M. (2019) The PRIDE database and related tools and resources in 2019: improving support for quantification data. *Nucleic Acids Res.*, 47, 442–D450.

60. Götz, M., Pettekau, J., Frischke, R., Ihling, C., Schäfer, M. and Sinz, A. (2015) Automated assignment of MS/MS cleavable cross-links in protein 3D-structure analysis. *J. Am. Soc. Mass. Spec.*, 26, 83–97.

61. Teichert, A., Arnold, L.A., Oteino, S., Oda, Y., Augustinaita, I., Geistlinger, T.R., Kriwacki, R.W., Guy, R.K. and Bille, D.D. (2009) Quantification of the vitamin D receptor-coregulator interaction. *Biochemistry*, 48, 1454–1461.

62. Ge, K., Cho, Y.W., Guo, H., Hong, T.B., Guermah, M., Ito, M., Yu, H., Kalkum, M. and Roeder, R.G. (2008) Alternative mechanisms by which mediator subunit MED1 regulates peroxisome proliferator-activated receptor gamma-stimulated adipogenesis and target gene expression. *Mol. Cell. Biol.*, 28, 1081–1091.

63. Atil, C., Goetze, M., Ihling, C.H., Hage, C., Schaefer, M. and Sinz, A. (2010) Cleavable cross-linker for protein structure analysis: reliable identification of cross-linking products by tandem MS. *Anal. Chem.*, 82, 6958–6968.

64. Atil, C., Goetze, M., Ihling, C.H., Hage, C., Schaefer, M. and Sinz, A. (2016) An integrated workflow for structural proteomics studies based on cross-linking/mass spectrometry with an MS/MS cleavable cross-linker. *Anal. Chem.*, 88, 7930–7937.

65. Merkley, E.D., Rysavy, S., Kahraman, A., Hafen, R.P., Daggett, V. and Adkins, J.N. (2014) Distance restraints from crosslinking mass spectrometry: Mining a molecular dynamics simulation database to evaluate lysine-lysine distance. *Protein Sci.*, 23, 747–759.

66. Weikum, E.R., Liu, X. and Ortlund, E.A. (2018) The nuclear receptor superfamily: a structural perspective. *Protein Sci.*, 27, 1876–1892.

67. Zhang, J., Chalmers, M.J., Stayrook, K.R., Burris, L.L., Wang, Y., Busby, S.A., Pascal, B.D., Garcia-Ordonez, R.D., Bruning, J.B., Istrate, M.A. et al. (2011) DNA binding alters coactivator interaction surfaces of the intact VDR–RXR complex. *Nat. Struct. Mol. Biol.*, 18, 556–563.

68. Oda, Y., Hu, L., Bul, V., Elalieh, H., Reddy, J.K. and Bille, D.D. (2012) Coactivator MED1 ablation in keratinocytes results in hair-cycling defects and epidermal alterations. *J. Invest. Dermatol.*, 132, 1075–1083.

69. Yang, W., Rachez, C. and Freedman, L.P. (2000) Discrete roles for peroxisome proliferator-activated receptor gamma and retinoid X receptor in recruiting nuclear receptor coactivators. *Cell. Biol.*, 20, 8008–8017.

70. Malik, S., Guermah, M., Yuan, C.X., Wu, W., Yamamura, S. and Roeder, R.G. (2004) Structural and functional organization of TRAP220, the TRAP/mediator subunit that is targeted by nuclear receptors. *Cell. Biol.*, 26, 9244–9254.

71. Jimenez-Lara, A.M. and Aranda, A. (1999) Lysine 246 of the vitamin D receptor is crucial for ligand-dependent interaction with coactivators and transcriptional activity. *J. Biol. Chem.*, 274, 13503–13510.
86. de Vera, I. M. S., Zheng, J., Novick, S., Shang, J., Hughes, T. S., Brust, R., Munoz-Tello, P., Gardner, W. J. Jr, Marciano, D. P., Kong, X. et al. (2017) Synergistic regulation of coregulator/nuclear receptor interaction by ligand and DNA. *Structure*, 25, 1506–1518.

87. Jin, F., Claessens, F. and Fondell, J. D. (2012) Regulation of androgen receptor-dependent transcription by coactivator MED1 is mediated through a newly discovered noncanonical binding motif. *J. Biol. Chem.*, 287, 858–870.

88. Krasowski, M. D., Ai, N., Hagey, L. R., Kollitz, E. M., Kullman, S. W., Reschly, E. J. and Ekins, S. (2011) The evolution of farnesoid X, vitamin D, and pregnane X receptors: insights from the green-spotted pufferfish (Tetraodon nigroviridis) and other non-mammalian species. *BMC Biochem.*, 12, 5.

89. Zenata, O. and Vrzal, R. (2017) Fine tuning of vitamin D receptor (VDR) activity by post-transcriptional and post-translational modifications. *Oncotarget*, 8, 35390–35402.

90. Barletta, F., Freedman, L. P. and Christakos, S. (2002) Enhancement of VDR-mediated transcription by phosphorylation: correlation with increased interaction between the VDR and DRIP205, a subunit of the VDR-interacting protein coactivator complex. *Mol. Endocrinol.*, 16, 301–314.

91. Arriagada, G., Paredes, R., Olate, J., van Wijnen, A., Lian, J. B., Stein, G. S., Stein, J. L., Onate, S. and Montecino, M. (2007) Phosphorylation at serine 208 of the 1alpha,25-dihydroxy Vitamin D3 receptor modulates the interaction with transcriptional coactivators. *J. Steroid Biochem. Mol. Biol.*, 103, 425–429.

92. Zheng, J., Chang, M. R., Stites, R. E., Wang, Y., Bruning, J. B., Pascal, B. D., Novick, S. J., Garcia-Ordonez, R. D., Stayrook, K. R., Chalmers, M. J. et al. (2017) HDX reveals the conformational dynamics of DNA sequence specific VDR co-activator interactions. *Nat. Commun.*, 8, 923.

93. Acevedo, M. L., Lee, K. C., Stender, J. D., Katzenellenbogen, B. S. and Kraus, W. L. (2004) Selective recognition of distinct classes of coactivators by a ligand-inducible activation domain. *Mol. Cell.*, 13, 725–738.