Data Descriptor: **Quantitative mapping of RNA-mediated nuclear estrogen receptor β interactome in human breast cancer cells**

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The nuclear receptor estrogen receptor 2 (ESR2, ERβ) modulates cancer cell proliferation and tumor growth, exerting an oncopsuppressive role in breast cancer (BC). Interaction proteomics by tandem affinity purification coupled to mass spectrometry was previously applied in BC cells to identify proteins acting in concert with ERβ to control key cellular functions, including gene transcription, RNA splicing and post-transcriptional mRNA regulation. These studies revealed an involvement of RNA in ERβ interactome assembly and functions. By applying native protein complex purification followed by nano LC-MS/MS before and after in vitro RNA removal, we generated a large dataset of newly identified nuclear ERβ interactors, including a subset associating with the receptor via RNA bridging. These datasets will be useful to investigate further the role of ERβ, nuclear RNAs and the other proteins identified here in BC and other cell types.

**Design Type(s)**
- parallel group design
- cell type comparison design
- protein and RNA interaction identification objective

**Measurement Type(s)**
- protein-protein interaction detection

**Technology Type(s)**
- nanoflow liquid chromatography-tandem mass spectrometry

**Factor Type(s)**
- genotype
- experimental condition
- biological replicate

**Sample Characteristic(s)**
- immortal human breast-derived cell line cell

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Background & Summary

The role of estrogen receptor β (ERβ) in cancer biology is still an open issue, despite its potential use as biomarker and drug target. Extensive in vitro and in vivo evidences point to an oncosuppressive role of this nuclear receptor in breast and other cancers, where it inhibits carcinogenesis, reduces cancer cell proliferation and tumor growth by exerting a specific control on gene transcription and other regulatory functions of the cell. Indeed, ERβ inhibits BC cells proliferation and invasion, both with and without ligand, showing an additive effect with antiestrogen. Furthermore, loss of ERβ expression in invasive BC, in early stages of ductal tumors and in breast tumorigenesis have been reported, together with a positive prognostic value of the presence of ERβ in cancer cells that may also be an indicator of their responsiveness to endocrine therapy. Although some of the results obtained in clinical samples have been challenged, due to the existence of multiple isoform of this protein and uncertainties concerning the specificity of the available antibodies for its detection in tissue specimens, all these evidences points to a key role of ERβ in BC biology. Finally, the functional role of the receptor in the absence of estrogen, a physiological condition during specific phases of the menstrual cycle, before puberty and in post-menopausal women, when the constitutive activities of ERβ might compensate for the absence of circulating hormones, is of great interest but still poorly understood.

Identification and characterization of the multiprotein complexes involved in the functions of ERβ is a critical step to identify the molecular bases of its signaling in BC cells. Interaction proteomics, combining native protein complexes purification and identification by mass spectrometry, is the gold standard to gain such information, and we and others have been mapping ERβ interactomes of human cells under different experimental conditions. By this approach, we recently demonstrated that ERβ can interact with AGO2 in BC cells and that this is mediated by one or more RNAs, suggesting for the first time that RNA plays a role in assembly and/or stabilization of ERβ interactomes, as already shown for other nuclear receptors.

In the present study we generated new ERβ interacting protein datasets by purification of native complexes extracted from C-terminus-tagged expressing ERβ (Ct-ERβ) MCF-7 cell nuclei before and after RNase treatment, followed by label free quantitative proteomics (Fig. 1). Results provide an expanded view of the ERβ nuclear interactome of BC cells, including identification of the protein-protein interactions mediated by RNA, that can now be exploited not only to understand the molecular bases of ERβ activities but also the functions of all other proteins identified.

Firstly, ERβ-containing nuclear protein complexes, purified by affinity chromatography (tandem affinity purification (TAP), partial procedure), were analysed by nano LC-MS/MS, leading to the identification of the largest receptor interactome mapped so far, comprising 1897 specific components, following exclusion of contaminants identified in control samples from ERβ-negative MCF-7 cells processed in the same way, excluding potential contaminants identified in Ct-ERβ samples (e.g. Keratins and Immunoglobulins) (Data Citation 1: 'Identified proteins' table). This ERβ interacting network comprises several sub-networks, comprising proteins involved in cellular functions known to be controlled by this receptor, including transcription, cell death and apoptosis and RNA splicing (Fig. 2).

RNase treatment was then performed in nuclear extracts from Ct-ERβ cells before nuclear complexes purification and mass spectrometry identification. After discarding the contaminants present in the negative control (Data Citation 1: 'Identified proteins' table), and potential contaminants present only in RNase treated Ct-ERβ samples (e.g. keratins and immunoglobulins, see above), 1453 specific ERβ interactors were identified (Data Citation 1: 'RNA-dependent interactors' table). A quantitative approach was then applied, by using MaxQuant tool, to identify proteins whose concentration was significantly increased or decreased between Ct-ERβ and ERβ controls.

Figure 1. Experimental workflow. Summary of the experimental work-flow applied to generate the protein datasets.
reduced by pre-treatment with RNase prior to affinity purification, respect to that in untreated samples. Considering the 1472 proteins unequivocally identified and quantified in both Ct-ERβ datasets (untreated and RNase-treated) according to the statistical analyses reported below, the concentration of about 16% of them was significantly (q-value ≤ 0,05) affected by RNA depletion (Fig. 3a and Data Citation 1: ‘MaxQuant analyses’ table). In detail: the concentration of 149 proteins was decreased up to -189 as difference value in RNase-treated vs untreated samples, indicating that their association with ERβ is mediated, directly or indirectly, by RNA (Fig. 3b). Concerning the ERβ molecular partners showing increased values after RNase treatment, this could be explained by a more efficient in vitro association of these proteins to the receptor moiety following enzymatic treatment, due to a decrease of steric hindrances consequent to removal of RNA and/or associated proteins from the complex.

Interestingly, several of them are known to be involved in the control of gene expression, cell cycle and other functions known to be part of the cellular responses to ERβ (Fig. 3c).

The dataset presented here will be useful to investigate the molecular mechanism of ERβ activity and to design ways to investigate composition and functional roles of macromolecular complexes in BC cell nuclei comprising proteins and RNAs, aiming also at the identification of interactome nodes representing potential drug targets against this, and possibly other, cancers.

Methods

Nuclear proteins extraction and treatments

Ct-ERβ3,14 and ERβ-negative MCF7 cells (which had been steroid deprived by culturing for 5 days in medium without phenol red and with 5% dextran-coated charcoal treated serum), were harvested by scraping in cold PBS and lysed as previously described23. Briefly, cell pellets were resuspended in 3 volumes of hypotonic buffer (20 mM HEPES pH 7.4, 5 mM NaF, 10 μM sodium molybdate, 0.1 mM

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Figure 2. ERβ interaction networks. (a) Global network including all the proteins specifically co-puriﬁed with ERβ by afﬁnity chromatography and mass spectrometry analysis, showing known associations, reported in protein–protein interaction databases. Sub-networks of ERβ-associated proteins involved in (b) transcription and validated nuclear estrogen receptors networks, (c) cell death and apoptosis and (d) splicing, as obtained by FunRich analysis.
Figure 3. ERβ interactome changes upon RNase treatment. (a) Volcano plot summarizing quantitative changes of ERβ-associated proteins upon treatment with RNase. Dotted line (threshold) represents the cut-off (q-value ≤ 0.05) (b) Heatmap of down-represented proteins showing level of LFQ intensities before and after RNase treatment (blu scale). Decrease of protein levels (+ vs – RNase) are shown in green scale. (c) Functional enrichment analysis by IPA of ERβ-associated proteins whose interaction with the receptor was reduced by RNase treatment (upper histogram) and a zoom-in on the Gene Expression functional category (lower histogram).
Two subsequent cleavage reactions were carried out for 2 h and 30 min respectively at 16 °C with gentle 
proteins. Multiple-charged ions were selected for fragmentation on the high-energy collision dissociation (HCD) 
after accumulation to a target of 3,000,000 ions in the quadruple. For MS/MS, the ten most abundant 
acquired from a mass-to-charge ratio (m/z) of 400 to 1,200 with the resolution R

(i0% FA for mass spectrometry analysis, that was carried out as previously described 28. In brief, peptide 
mixtures were analyzed with an Easy nLC1000 nano-LC system connected to a quadrupole Orbitrap mass 
spectrometer (QExactive Plus, ThermoElectron, Bremen, Germany) equipped with a nanoelectrospray 
ion source (EasySpray/Thermo). Liquid chromatography separation of the peptides was performed with 
an EasySpray column capillary of 50 cm bed length (C18, 2 μm beads, 100 A, 75 μm inner diameter, 
Thermo) at 300 nL/min flow rate. Peptides were eluted with a 2-30% gradient of solvent B in 60 min 
(solvent A: aqueous 0.1% formic acid, solvent B: 100% acetonitrile / 0.1% formic acid). Data-dependent 
acquisition automatically switched between MS and MSMS mode and survey full scan MS spectra were 
acquired from a mass-to-charge ratio (m/z) of 400 to 1,200 with the resolution R = 70,000 at m/z 200, 
after accumulation to a target of 3,000,000 ions in the quadruple. For MS/MS, the ten most abundant 
multiple-charged ions were selected for fragmentation on the high-energy collision dissociation (HCD) 
cell at a target value of 100,000 charges or maximum acquisition time of 100 ms. The MS/MS scans were

| Sample name (cell line) | Protocol 1 | Protocol 2 | Protocol 3 | Treatment | Data |
|------------------------|------------|------------|------------|-----------|------|
| MCF7 (CTRL)_1          | Nuclear protein extracts | Tandem Affinity Purification | Nano LC-MS/MS | - | PRIDE PXD006720 (P4883-P4887) |
| MCF7 (CTRL)_2          | Nuclear protein extracts | Tandem Affinity Purification | Nano LC-MS/MS | - | PRIDE PXD006720 (P4889-P4894) |
| MCF7 (CTRL)_3          | Nuclear protein extracts | Tandem Affinity Purification | Nano LC-MS/MS | - | PRIDE PXD006720 (P4896-P4901) |
| Ct-ERβ (sample)_1      | Nuclear protein extracts | Tandem Affinity Purification | Nano LC-MS/MS | - | PRIDE PXD006720 (P4834-4839) |
| Ct-ERβ (sample)_2      | Nuclear protein extracts | Tandem Affinity Purification | Nano LC-MS/MS | - | PRIDE PXD006720 (P4841-P4846) |
| Ct-ERβ (sample)_3      | Nuclear protein extracts | Tandem Affinity Purification | Nano LC-MS/MS | - | PRIDE PXD006720 (P4848-P4853) |
| Ct-ERβ (sample)_4      | Nuclear protein extracts | Tandem Affinity Purification | Nano LC-MS/MS | Rnase A | PRIDE PXD006720 (P4866-P4867) |
| Ct-ERβ (sample)_5      | Nuclear protein extracts | Tandem Affinity Purification | Nano LC-MS/MS | Rnase A | PRIDE PXD006720 (P4869-P4874) |
| Ct-ERβ (sample)_6      | Nuclear protein extracts | Tandem Affinity Purification | Nano LC-MS/MS | NT | PRIDE PXD006820 (P5164) |
| Ct-ERβ (sample)_7      | Nuclear protein extracts | Tandem Affinity Purification | Nano LC-MS/MS | NT | PRIDE PXD006820 (P5166) |
| Ct-ERβ (sample)_8      | Nuclear protein extracts | Tandem Affinity Purification | Nano LC-MS/MS | NT | PRIDE PXD006820 (P5168) |
| Ct-ERβ (sample)_9      | Nuclear protein extracts | Tandem Affinity Purification | Nano LC-MS/MS | AGO2. Kd | PRIDE PXD006820 (P5170) |
| Ct-ERβ (sample)_10     | Nuclear protein extracts | Tandem Affinity Purification | Nano LC-MS/MS | AGO2 Kd | PRIDE PXD006820 (P5172) |
| Ct-ERβ (sample)_11     | Nuclear protein extracts | Tandem Affinity Purification | Nano LC-MS/MS | AGO2 Kd | PRIDE PXD006820 (P5174) |

Table 1. Summary of the protocols and datasets used.

EDTA, 1 mM PMSF and 1X protease inhibitors cocktail (Sigma Aldrich) and incubated on ice for 15 min. 
Cytosolic fraction was discarded after adding 0.5% Triton X-100 and spinning for 30 s at 15000 X g at 
4 °C. Nuclear pellets were then suspended in 1 volume of nuclear lysis buffer (20 mM HEPEs pH 7.4, 25% 
glycerol, 420 mM NaCl, 1.5 mM MgCl2, 0.2 mM EDTA, 1 mM PMSF and 1X protease inhibitors cocktail (Sigma Aldrich), incubated for 30 min at 4 °C rotating and centrifuged for 30 min at 15000 X g at 4 °C. Supernatants were finally recovered, diluted 1:3 with nuclear lysis buffer without NaCl to restore the 
physiological saline concentration and quantified.

Purification of ERβ nuclear complexes

IgG-Sepharose beads (GE Healthcare), pre-treated according to the manufacturer’s instructions and 
equilibrated in TEV buffer (50 mM Tris- HCl pH 8.0, 0.5 mM EDTA, 0.1% Triton X-100, 150 mM NaCl, 
1 mM DTT), were added to nuclear protein extracts and incubated for 3 h at 4 °C with gentle rotation, as 
described earlier14,16,23,25,26. Where indicated (see Table 1), 100 μg/ml RNaseA were added to the samples 
before binding as previously described19,27. After incubation, unbound proteins were discarded following 
centrifugation and the beads were thoroughly washed with 100XVol of IPP150 buffer (20 mM HEPEs pH 
7.5, 8% glycerol, 150 mM NaCl, 0.5 mM MgCl2, 0.1 mM EDTA, 0.1% Triton X-100) and equilibrated in 30XVol of TEV Buffer in Poly-Prep Chromatography columns (0.8x4 cm, Bio-Rad) at 4 °C. Then 4xBeads 
Vol of Cleavage Buffer (TEV Buffer containing 1U/µl beads of TEV protease, Invitrogen) were added and 
two subsequent cleavage reactions were carried out for 2 h and 30 min respectively at 16 °C with gentle 
shaking. The eluates were then collected, after sedimentation of beads still binding uncut and non-specific 
proteins.

Nano LC-MS/MS and Data Analysis

Three biological replicates of partially purified samples from control MCF-7 and from Ct-ERβ cells before 
and after RNase treatment were analyzed. The protein extracts were precipitated with 10% TCA in 
Acetone solution, resuspended in Laemmli sample buffer, separated by SDS-PAGE and visualized with 
silver-staining. The SDS-PAGE lanes were sliced into 6 pieces/each and the proteins were in-gel digested 
with trypsin (Promega) overnight at 37 °C and eluted as previously described14,16,23,25,26. Peptides were 
desalted and concentrated before mass spectrometry by the STAGE-TIP method using a C18 resin disk 
(3M Empore). The peptides were eluted twice with 0.1% TFA / 50% ACN, dried and solubilized in 7 μl 
0.1% FA for mass spectrometry analysis, that was carried out as previously described28. In brief, peptide 
mixtures were analyzed with an Easy nLC1000 nano-LC system connected to a quadrupole Orbitrap mass 
spectrometer (QExactive Plus, ThermoElectron, Bremen, Germany) equipped with a nanoelectrospray 
ion source (EasySpray/Thermo). Liquid chromatography separation of the peptides was performed with 
an EasySpray column capillary of 50 cm bed length (C18, 2 μm beads, 100 A, 75 μm inner diameter, 
Thermo) at 300 nL/min flow rate. Peptides were eluted with a 2-30% gradient of solvent B in 60 min 
(solvent A: aqueous 0.1% formic acid, solvent B: 100% acetonitrile / 0.1% formic acid). Data-dependent 
acquisition automatically switched between MS and MSMS mode and survey full scan MS spectra were 
acquired from a mass-to-charge ratio (m/z) of 400 to 1,200 with the resolution R = 70,000 at m/z 200, 
after accumulation to a target of 3,000,000 ions in the quadruple. For MS/MS, the ten most abundant 
multiple-charged ions were selected for fragmentation on the high-energy collision dissociation (HCD) 
cell at a target value of 100,000 charges or maximum acquisition time of 100 ms. The MS/MS scans were
collected at a resolution of 17,500. Target ions already selected for MS/MS were dynamically excluded for 30 s. The resulting MS raw files were submitted for protein identification to Proteome Discoverer 2.1 interface using the Mascot 2.5 search engine. The search criteria for Mascot searches were: trypsin digestion with two missed cleavage allowed, Carbamidomethyl (C) as fixed modification and Acetyl (N-term), Gln->pyro-Glu (N-term Q), Oxidation (M) as variable modifications. The parent mass tolerance was 10 ppm and MS/MS tolerance 0.1 Da. The SwissProt human database was used (August 2016, with 154,660 entries) for the database searches. All of the reported protein identifications were statistically significant ($p < 0.05$) in Mascot and filtered in ProteomeDiscoverer for high confidence peptides. Only proteins identified in the Ct-ER$\beta$ samples and not present in ER$\beta$- control samples prepared in the same way were considered as receptor specific interactors. Additionally, potential contaminants identified only in the Ct-ER$\beta$ samples (e.g. Keratins, Immunoglobulins) were discarded and excluded from further analysis. Label-free quantitation (LFQ) was employed with default settings. The SwissProt human database was used (August 2016, with 154,660 entries) for the database searches. Statistical analysis was performed with the Perseus software (version 1.5.6.0). Proteins identified in the decoy reverse database, or only by site, were not considered for data analysis. Also, known and potential contaminants provided by MaxQuant and present in the samples were excluded together with all proteins identified by ProteomeDiscoverer/Mascot searches also in the control, ER$\beta$- samples were discarded. The label-free quantification (LFQ) data were log10 transformed, filtered to include only proteins identified and quantified in at least two out of three replicates in at least one experimental group, and missing values were imputed with values representing a normal distribution with default settings in Perseus (generated at 1.8 standard deviations of the total intensity distribution, subtracted from the mean, and a width of 0.3 standard deviations).
deposited to the ProteomeXchange Consortium via the PRIDE\textsuperscript{30} partner repository with the dataset identifier PXD006720 (Data Citation 2) for interaction and quantitative proteomics datasets comprising CTRLs and Ct-ER$\beta$ samples before and after RNase treatment. The list of proteins whose association with the receptor changed after depletion of RNA (Data Citation 1: 'RNA-dependent interactors' and 'MaxQuant analyses' tables) was compared with that comprising the proteins whose association with the receptor changed upon AGO2 silencing\textsuperscript{19} (Data Citation 3). Results can be found in the figshare repository (Data Citation 1: 'Comparison of ER beta interactors after either RNA or AGO2 depletion' table).

**Code availability**

The following software and versions were used for quality control and data analysis:

1. For protein identification from raw MS data, Proteome Discoverer software (Thermo) version 2.1 was used: https://tools.thermofisher.com/ with the Mascot 2.5 search engine.
2. For quantitative data analysis, MaxQuant software version 1.5.7.4 with the Andromeda search engine was used: http://www.coxdocs.org/doku.php?id=maxquant:start
3. The SwissProt human database was used (August 2016, with 154,660 entries) as database for protein searches: http://web.expasy.org/
4. For functional interaction networks visualization FunRich v3.0 was used: http://www.funrich.org
5. Functional analysis were performed with IPA software version 01-07: http://www.qiagenbioinformatics.com/products/ingenuity-pathway-analysis
6. Statistical analyses were performed using R www.r-project.org

**Data Records**

Data records can be downloaded from our Figshare archive (Data Citation 1). The newly described mass spectrometry proteomics data has been deposited to the ProteomeXchange Consortium via the PRIDE\textsuperscript{30} partner repository with the following dataset identifier: PXD006720 for interaction and quantitative proteomics datasets comprising CTRLs and Ct-ER$\beta$ samples before and after RNase treatment (Data Citation 2).

**Technical Validation**

Data shown were obtained from three, independent biological replicates (cell cultures processed independently, each comprising 30 × 150 mm$^2$ dishes at 80% confluency), for both controls and samples, analysed in parallel. The best practice applied in the experimental workflow and technical validations are described below.

Firstly, authenticated cell lines with no more than 5 passages after thawing and mycoplasma-free were used. Affinity purification was performed independently for each replicate processing each control and sample simultaneously in parallel. To quantify the effects of the RNase treatment, RNA was purified from an aliquot of each nuclear extract before TAP and analysed by microfluidic electrophoresis as described\textsuperscript{19} previously (Fig. 4a). Furthermore, aliquots of each sample were kept from each passage and analysed by western blotting to assess presence and integrity of the bait protein ER$\beta$ and its known partner AGO2 (ref. 19) in ER$\beta$+ samples and the background in control samples (Fig. 4b–d). Moreover, to exclude that the observed proteome changes were not due to a reduction of ER$\beta$ concentration in the starting materials or of its purification efficiency following RNase treatment, ER$\beta$ LFQ intensities were compared before and after RNase treatment in Ct-ER$\beta$ expressing cells (Figure 4e). The correlation among the triplicate purifications was verified by analysing log2 of LFQ intensities with R version 3.4.2 (see code availability 6), applying the function cor from R-package sats v3.4.2 with Pearson correlation method.

Label-free quantitative analysis with MaxQuant was performed to obtain information concerning the impact of RNase on ER$\beta$ interactome (differentially purified proteins). Finally, we compared data from the newly obtained datasets with those previously obtained following AGO2 silencing\textsuperscript{19}. Data analysis, considering as reference the dataset generated here (PXD006720), confirmed unliganded ER$\beta$ interactome composition, with an 85% overlapping between the two datasets. We then compared the proteins whose concentration changed in the purified samples after depletion of either RNA or AGO2, by RNase or siRNA treatment respectively\textsuperscript{19} (Data Citation 1: 'Comparison of ER beta interactors after either RNA or AGO2 depletion' table and Data Citation 3). A summary of all data records related to the samples analysed here is reported in Table 1.

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Data Citations
1. Weiss, A. et al. fgshare http://doi.org/10.6084/m9.fgshare.c.3952927 (2018).
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Author Contributions
All authors participated in conception and design of the study. R.T., G.N., A.S., E.A. and F.R. performed in vitro experimental work, T.N. and G.N. performed the proteomics analyses, G.G. performed the statistical and bioinformatics analyses, R.T., G.G., G.N. and A.W. coordinated and finalized figure preparation, manuscript drafting and revision. All authors read and approved the final manuscript.
Additional Information

Competing interests: The authors declare no competing financial interests.

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