Systematic Mapping of Posttranslational Modifications in Human Estrogen Receptor-α with Emphasis on Novel Phosphorylation Sites*†‡§

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A systematic study of posttranslational modifications of the estrogen receptor isolated from the MCF-7 human breast cancer cell line is reported. Proteolysis with multiple enzymes, mass spectrometry, and tandem mass spectrometry achieved very high sequence coverage for the full-length 66-kDa endogenous protein from estradiol-treated cell cultures. Nine phosphorylated serine residues were identified, three of which were previously unreported and none of which were previously observed by mass spectrometry by any other laboratory. Two additional modified serine residues were identified in recombinant protein, one previously reported but not observed here in endogenous protein and the other previously unknown. Although major emphasis was placed on identifying new phosphorylation sites, N-terminal loss of methionine accompanied by amino acetylation and a lysine side chain acetylation (or possibly trimethylation) were also detected. The use of both HPLC-ESI and MALDI interfaced to different mass analyzers gave higher sequence coverage and identified more sites than could be achieved by either method alone. The estrogen receptor is critical in the development and progression of breast cancer. One previously unreported phosphorylation site identified here was shown to be strongly dependent on estradiol, confirming its potential significance to breast cancer. Greater knowledge of this array of posttranslational modifications of estrogen receptor, particularly phosphorylation, will increase our understanding of the processes that lead to estradiol-induced activation of this protein and may aid the development of therapeutic strategies for management of hormone-dependent breast cancer. Molecular & Cellular Proteomics 8:467–480, 2009.

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The α-isofrom of the estrogen receptor (ERα)1 is a 66-kDa nuclear transcription factor that mediates transcriptional regulation of genes involved in cell proliferation and differentiation and plays a pivotal role in the development and progression of breast cancer (1–3). Consequently the development of therapies for management of hormone-dependent breast cancer has targeted signaling pathways based on modulation of ERα activity (4–6). Current knowledge and understanding of ERα activity is derived from over 30 years of accumulated scientific evidence that has sought to delineate ER-mediated mechanisms and signaling pathways under pathological conditions modeled in cultured breast cancer cell lines. Because the function or activity of a protein may depend strongly on the presence of posttranslational modifications (PTMs), significant research has focused on detection and quantitation of such modifications in ERα. The constellation of PTMs on a protein constitutes a molecular code that may dictate protein conformation, localization, and function. Thus biological inference based on ERα structure requires a comprehensive study of all possible PTMs on the constituent amino acids.

Modifications to ERα reported to date are listed in Table I. Two of the most common and important PTMs that modulate the activity of ERα are redox-based modifications of cysteine residues (7–12) and phosphorylation of serine, threonine, and tyrosine residues (13). Biochemical techniques used previously to map ERα phosphorylation sites utilized radioactive labeling with 32P (14–16), Edman degradation, deletion and/or point mutations, and Western blots (17–19). Mutational analysis in combination with estrogen response element-luciferase reporter assays can validate the functional relevance of phosphorylation sites (16, 20). Although these techniques have provided substantial information regarding PTMs in ERα, they have several disadvantages. Autoradiography using radiolabeling of 32P lacks specificity, Edman degradation requires large amounts of purified protein and is not applicable to proteins/peptides with N-terminal acetylation, and

1 The abbreviations used are: ERα, α-isofrom of the estrogen receptor; PTM, posttranslational modification; LTO, linear quadrupole trap; AF, activation factor; DD, DNA-binding domain; MALDI, vac-uum MALDI; ER, estrogen receptor; rER, recombinant ER; MRM, multiple reaction monitoring.
quantitation of PTMs by Western blot analysis relies on prior knowledge of the type and position of specific modifications and on the availability of high quality antibodies. Deletional and mutational analyses on transfected rather than endogenous protein are laborious and time-consuming. Moreover, discrepancies in the literature concerning ERα phosphorylation may arise from the use of different promoters in the reporter constructs (13).

Because of its selectivity and sensitivity, MS/MS has emerged as a powerful tool for the analysis of PTMs such as phosphorylation (21–23). The analysis of phosphopeptides benefits from the use of multiple ionization methods and instrumentation platforms as ESI and MALDI interfaced with different tandem mass analyzers provide complementary information (21). Orthogonal Q-TOF and linear quadrupole trap (LTQ) instruments are widely used for ESI analysis, and several tandem instruments have been developed for or adapted to MALDI methods, including the LTQ (24–26). We previously utilized vacuum (v) MALDI-LTQ to optimize quantitation of cysteine oxidation within the DNA-binding domain (DBD) of recombinant ERα (7). More recently we enriched and immunoaffinity-purified endogenous ERα derived from the human breast cancer cell line MCF-7 in quantities compatible with tandem MS analysis and identified a novel phosphorylation site, Ser154. vMALDI-MS	extsuperscript{+} confirmed Ser154 phosphorylation in human breast cancer cells grown under both ligand-dependent and ligand-independent conditions (27). Coincidentally we also observed known phosphorylation sites Ser118 and Ser167 not previously confirmed by MS methods.

Here we report a comprehensive MS study of ERα to achieve very high sequence coverage for the full-length 66-kDa endogenous protein from estradiol-treated MCF-7 cell cultures with particular emphasis on identifying new phosphorylation sites. Using multiple reaction monitoring (MRM)/MS we demonstrated that one previously unreported phosphorylation sites. Using multiple reaction monitoring (MRM)/MS we previously enriched and immunopurification, and in-gel digestion of ERα from a human breast cancer cell line /H9251 was obtained from the American Type Culture Collection (Manassas, VA). Protocols for cell culture and extraction, immunopurification, and in-gel digestion of ERα from 20 15-cm plates of cells were as described previously (27) except that for each experiment in-gel digestion was with one of the following enzymes: chymotrypsin (300 ng; 37 °C), trypsin (150 ng; 37 °C), Glu-C (250 ng; 25 °C), Lys-C (400 ng; 37 °C), Asp-N (280 ng; 37 °C), or a combination of the last two. The resultant peptides were extracted by vortexing and sonication with 50% ACN, 5% formic acid; vacuum centrifuged to remove most of the solvent; reconstituted in 20 μl of 0.1% formic acid; and stored at −80 °C.

On-bead Proteolytic Digestion—After immunoprecipitation and PBS washes the immune complex was transferred to a siliconized 1.5-ml Eppendorf tube and washed twice in 100 mM Tris, pH 7, by gentle rotation for 2 min at room temperature and three times in 25 mM NH₄HCO₃ with centrifugation between washes. The pellet beads were suspended in a further 150 μl of NH₄HCO₃ and 170 ng of sequence grade trypsin (10 μl of 17 ng/μl stock solution) and then incubated overnight at 37 °C with shaking at 1,200 rpm. Eppendorf tubes were centrifuged at 12,000 rpm, and the supernatant was transferred to low binding polymer technology 0.65-ml microcentrifuge tubes (PGC Scientifics). Ten microliters of acetonitrile and 1 μl of 10% formic acid were added to the peptide solution, which was then evaporated down to −15 μl. Each sample was divided into three 5-μl aliquots and stored at −80 °C until used for mass spectrometry.

Peptide Mass Fingerprinting by MALDI-TOF—Peptides to be analyzed by MALDI were desalted with C₁₈ ZipTips (Millipore), and then 1 μl of peptide solution and 1 μl of 2.5-dihydroxybenzoic acid or α-cyano-4-hydroxycinnamic acid matrix (50 mg/ml in 4:1 ACN, 0.6% phosphoric acid) were mixed, deposited on a stainless steel sample plate, and air-dried. Alternatively 1 μl of peptide solution was mixed with 1 μl of 50% ACN, 0.5% TFA; spotted on the MALDI plate; and dried. This addition of solvent and drying was repeated three more times, and then 1 μl of matrix solution was added and dried. 1 μl of 0.1% TFA was added and aspirated off by applying a tissue. Mass spectra were recorded on a Voyager-DE STR Plus instrument (Applied Biosystems, Framingham, MA) with 337 nm laser radiation, positive ion delayed extraction, reflector mode, and 20-kV accelerating voltage.

Peptide Identification by Tandem Mass Spectrometry—Peptide mixtures were further analyzed on a vMALDI-LTQ linear ion trap (Thermo Fisher, San Jose, CA) using both manual and automated data acquisition. Both methods used “Tune plus” using the “crystal-positioning system” and low threshold settings for “auto spectrum filter.” MS	extsuperscript{2} spectra were typically collected using threshold values of 500 counts for MS and 150 counts for MS	extsuperscript{2} with “auto gain control” turned on, limiting the number of ions admitted to the trap. A parent mass isolation width of 3 m/z units and fragmentation settings of activation Q = 0.25, activation time of 30 ms, and relative collision energy of 35% were used. For phosphopeptide mapping, potential peptide peaks were interrogated by MS/MS for the neutral loss of phosphoric acid, and (MH − 98) peaks were selected for MS	extsuperscript{3}. Loss of phosphoric acid converted serine residues to dehydroalanine (dS). Reversed phase nano-HPLC-MS/MS (LC-MS/MS) was performed using an Ultimate HPLC instrument (Dionex, Sunnyvale, CA) with a C₁₈ analytical column directly connected to a Q-STAR Pulsar I quadrupole orthogonal TOF mass spectrometer (MDS Sciex, Concord, Canada) as described previously (27).

LC-MS/MS and vMALDI-LTQ fragment ion data were searched against a database constructed for human full-length ERα using Mascot in-house licensed version 2.2 (Matrix Sciences, London, UK) with the “no enzyme” option. LC-MS/MS data were also searched with Paragon (Protein Pilot 2.0, Applied Biosystems).

LC-MRM/MS—Samples were analyzed by nano-LC-MRM/MS on a 4000 QTRAP hybrid triple quadrupole/linear ion trap mass spectrometer (Applied Biosystems). Chromatography was performed using an Eksigent (Dublin, CA) NanoLC-2D LC system with buffer A (0.1% formic acid) and buffer B (90% acetonitrile in 0.1% formic acid). Digests were loaded at 20 μl/min in buffer A onto a 5-mm × 300-μm reversed phase C₁₈ column (5 μm, 100 Å; Dionex) and eluted at 300 n/min with a 75-μm-inner diameter Integrafit column (New Objective, Woburn, MA) packed in house with 10–12 cm of ReproSil-Pur C₁₈ AQ 3-μm reversed phase resin (Dr. Maisch, GmBH) with a gra-
dient of 2–70% B over 32 min. Peptides were ionized using a PicoTip emitter (75/192 m, 15/192 m tip; New Objective). Data acquisition was performed with ion spray voltage at 2450 V, curtain gas pressure at 10 p.s.i., nebulizer gas at 20 p.s.i., and interface heater temperature at 150 °C. Collision energy, declustering potential, and collision cell exit potential were optimized using recombinant ER for maximum sensitivity. MRM transitions were monitored and acquired at unit resolution both in the first and third quadrupoles (Q1 and Q3) to maximize specificity. The y7 transition was monitored at Q1 670.9 m/z and Q3 785.5 m/z for the unmodified peptide and at Q1 710.9 m/z and Q3 865.4 m/z for the phosphorylated peptide with dwell times of 10 and 320 ms, respectively. Furthermore although the y7 transition gave the most intense signal, MRM transitions y8 and y7 were also assayed to confirm the retention time of the Ser294 peptides. A minimum of nine data points were collected per peak.

RESULTS

The overall analytical strategy (Fig. 1A) required optimizing immunoaffinity enrichment of human ERα from estradiol-treated MCF-7 cells for peptide analysis by tandem mass spectrometry. Estradiol treatment of confluent MCF-7 cells for 30 min, overnight incubation of cell lysates with agarose-coupled ERα antibody, and one-dimensional gel electrophoresis separation yielded endogenous ERα protein at 66 kDa (Fig. 1B). Recombinant protein (rERα) was used to confirm this identification and to model in-gel digestions. Comparison with 13 pmol of rERα indicates a yield of ~8–10 pmol of protein/20 plates of cultured MCF-7 cells. Western blots of cell lysates before and after immunoprecipitation indicate the ERα immunocapture to be as much as 98% efficient (Fig. 1C).

PeptideCutter (ExPASy) was used to predict peptide fragments of a size and nature suitable for ionization and sequencing by LC-MS/MS and/or vMALDI-MS, showing that multiple enzymes would be required to achieve comprehensive coverage of the primary sequence. Digests were prepared using trypsin, chymotrypsin, Glu-C, Lys-C, Asp-N, and the combination of Lys-C and Asp-N and analyzed as unseparated mixtures by MALDI-TOF. Fig. 2 shows a typical peptide mass fingerprint derived from a chymotryptic digest of immunopurified MCF-7 ERα. Internally calibrated peak lists were searched against the ERα database on Swiss-Prot using Protein Prospector Version 4.5.1. Peptide sequence information was then obtained by LC-MS/MS and vMALDI-MS. All spectra were interrogated by database searching, and sequence identifications were confirmed by visual inspection. All identified ERα peptides are listed in supplemental Table 1 and summarized in Fig. 3 with combined sequence coverage of 95%. Peptides assigned by peptide mass fingerprinting were not classified as identified unless confirmed by tandem MS. LC-MS/MS analysis yielded more peptides than vMALDI-MS of unseparated mixtures. Chymotrypsin gave the highest sequence coverage by both LC-MS/MS and vMALDI-MS (73% combined) and the highest coverage for detection of critical residues: serine (43 of 45), threonine (20 of 25), tyrosine (19 of 23), and lysine (17 of 29) but not for monitoring another important residue, cysteine. Spectra presented in the text represent previously unreported PTMs in endogenous ERα from MCF-7 cells. Spectra confirming previous reports or observations made only in recombinant protein are presented.
as supplemental data. Data for Ser(P)118, Ser(P)154, and Ser(P)167 have been reported previously and are not repeated here except for a Ser(P)154 peptide that models an unexpected MS/MS fragmentation.

**Monitoring of Cysteine Residues**

To facilitate detection of the 13 ERα cysteine residues, protein samples were reduced and alkylated. For the important DBD, consecutive digestion by Lys-C and Asp-N gave four major peptides, each containing two cysteine residues. Cys381 and Cys417 were not identified in Lys-C/Asp-N digests of MCF-7-derived ERα/H9251, although Cys417 was identified in rERα/H9251 following chymotryptic digestion. Cys530 was observed only in Glu-C digests, and Cys381 was not observed at all. Our studies on the oxidation of cysteine residues in the DBD have been described elsewhere (8, 10, 11).

**Observation of Acetylated Amino Acids**

No identified peptides contained the N-terminal methionine residue predicted by the gene sequence. Broadening the search criteria to allow for loss of this residue together with N-terminal acetylation led to the identification of Ac-2TMTLHTK in tryptic and Lys-C/Asp-N digests. The LC-MS/MS spectrum of the MH2/11001 is illustrated in Fig. 4A. b1 and b2 ions confirm N-terminal acetylation, whereas y2 and y3 exclude the possibility of the acetyl group being located on Lys8. Ac-2TMTLHTKASGMALLHQIQGNE was also identified with a Mascot score of 72 (Glu-C digest; spectrum not shown). Thus there is strong evidence that endogenous MCF-7 ERα undergoes this common modification, which would render it blocked for Edman sequencing.

Two tryptic peptides having similar MS/MS spectra differed in mass by 42 Da, suggesting another acetyl group. Alternatively this mass difference could correspond to trimethylation, which could not be differentiated with the instruments used here. Several ERα acetylation sites have been reported (Table I), but a methylation was also detected recently (28). For the current discussion, it is assumed the modification is an acetylation. vMALDI-LTQ fragmentation of the unmodified MH+/11001 (m/z 1263.4) identified it as 468SLEEKDHIHR with sequence ions y4, y6, y7–NH3, y8, b9, b9–NH3, and b10 (Fig. 4B). The putative acetylated form (m/z 1305.4) showed the same fragments (Fig. 4C), all but y4 (m/z 562.4) were shifted by 42 Da.
consistent with a centrally located modification. MS$^3$ of both $y_4$ ions gave identical spectra for both the non-acetylated and acetylated forms (inset). Thus the acetyl group is within EKD, placing it on the side chain of Lys$^{472}$.

**Phosphopeptide Identification by MS/MS**

Several phosphopeptides were identified in this study (Table II), some confirming previously known sites as well as several that were previously unreported. As a potential aid to phosphopeptide analysis, three Web-based algorithms, Scansite, NetPhos, and Disphos, were used to predict phosphorylation sites, but the correlation between prediction and experiment proved weak (results not shown). MS/MS analysis was performed on digested ER$\alpha$ peptides without any prior phosphopeptide enrichment step. Consequently it was often possible to observe both the phosphorylated and unphosphorylated species. In LC-MS/MS the phosphopeptides were consistently eluted later than the unphosphorylated forms, providing additional chromatographic confirmation of their existence. Phosphopeptides usually resulted in pairs of pep-
tides separated by 80 Da plus $y_n - 98$ ions for the higher mass species. Intact phosphate-containing $y_n$ ions were seen infrequently.

Complementary peptide information was obtained by automated vMALDI-LTQ-MS/MS experiments, initially based on MALDI-TOF-generated inclusion lists. Manual data acquisition was also used to sequence peptides with low ion counts not initially observed by MS and to isolate weak precursor ions and monitor peptides not detected by LC-MS/MS analysis. vMALDI-MS/MS also allowed repeated analysis of the same sample as well as extensive MS$^n$ interrogation of ions, whereas LC analysis was limited to the brief time during which a peptide eluted.

All ER$\alpha$ peptides containing unmodified serine, threonine, or tyrosine residues were interrogated by vMALDI-LTQ-MS/MS at 80 m/z units above the observed molecular ion even if no ions were seen at this mass. Additionally hypothetical phosphopeptide m/z values were generated by in silico proteolysis of ER$\alpha$ and probed for putative phospho-Ser, -Thr, and -Tyr residues in both recombinant and endogenous ER$\alpha$ by multiple stage vMALDI-MS$^n$. The loss of phosphoric acid (98 Da) in the linear ion trap provided a sensitive readout for the presence of a phosphopeptide; thus m/z regions that might correspond to (MH$^+$ − 98)$^+$ ions were examined for indications of phosphorylated residues using MS$^3$ to probe for fragment ions that correlated with the MS$^2$ spectra derived from the corresponding unphosphorylated parent ion.

Systematic Search for Phosphorylation Sites

The N-terminal Domain—Serine phosphorylation within the N-terminal domain containing activation factor-1 (AF-1) contributes to ER$\alpha$ activation and may lead to drug resistance in breast cancer treatment. We recently reported Ser$^{154}$ as a new phosphorylation site within the N-terminal domain of ER$\alpha$ from cultured human breast cancer cells. Here we found no other new sites but observed known sites at Ser$^{102/104/106}$, Ser$^{118}$, and Ser$^{167}$. Data for Ser(P)$^{118}$ and Ser(P)$^{167}$ were reported previously (27) and are not presented here.

We sought to confirm whether each of the previously reported sites Ser$^{102/104/106}$ can be phosphorylated; the data is presented in supplemental Fig. 1. The analysis was on an MCF-7 ER$\alpha$ chymotryptic peptide containing all three residues, $^{90}$GSNGLGFGPPLNSVSPSPL. LC-MS/MS spectra in supplemental Fig. 1, A and B, are from the MH$_2^{2+}$ ions of unphosphorylated peptide (m/z 898.95) eluting at 24.80 min and a monophosphorylated moiety (m/z 938.94) eluting at 26.56 min, i.e. differing by 80 Da. A strong $y_2$ ion of m/z 229.15 is common to both spectra, whereas $y_4$ at m/z 413.26 for the unphosphorylated peptide is replaced by $y_4 - H_2PO_4^-$ (m/z 395.22) in the phosphorylated case, localizing the modification on Ser$^{106}$. All y ions above $y_2$ show the loss of phosphoric acid, but $y_4$ alone definitively locates the site of the modification. Supplemental Fig. 1C shows a weaker MS/MS spectrum for an additional isomer eluting later (28.59 min). The $y_3/y_4$ ions show no phosphorylation, whereas $y_5$ has lost phosphoric acid, suggesting phosphorylation of Ser$^{154}$.

Crystal heterogeneity of unseparated digests can cause some separation of isomers, and different regions of the MALDI spot may yield distinct spectra. Several spectra from phosphorylated $^{90}$GSNGLGFGPPLNSVSPSPL indicated multiple isomeric forms. Supplemental Fig. 1D presents a vMALDI-LTQ-MS/MS spectrum with (MH$^+$ − 98)$^+$ as the base.

### Table I

| Residue | Modification | Ref. |
|---------|--------------|------|
| Serine 10 | O-Linked N-acetylglucosamine | Cheng and Hart (33) |
| Serine 104 | Phosphorylation | Le Goff et al. (29) |
| Serine 106 | Phosphorylation | Le Goff et al. (29) |
| Serine 118 | Phosphorylation | All et al. (34) |
| Serine 154 | Phosphorylation | Britton et al. (27) |
| Serine 167 | Phosphorylation | Arnold et al. (14) |
| Serine 236 | Phosphorylation | Chen et al. (15) |
| Various cysteine residues | S-Nitrosylation | Garban et al. (35) |
| Cysteines 221, 227, 237, and 240 | Thiol oxidation | Whittal et al. (11) |
| Lysines between 251 and 305 | Sumoylation | Sentis et al. (36) |
| Lysine 266 and lysine 268 | Acetylation | Kim et al. (37) |
| Lysine 299, 302, and 303 | Acetylation | Wang et al. (16) |
| Lysine 302 | Methylation | Subramanian et al. (28) |
| Serine 305 | Phosphorylation | Wang et al. (30) |
| Threonine 311 | Phosphorylation | Lee and Bai (31) |
| Cysteine 447 | S-Palmitoylation (C$_{16}$) | Marino et al. (38) |
| Tyrosine 537 | Phosphorylation | Arnold et al. (32) |
| Threonine 575 | O-Linked N-acetylglucosamine | Jiang and Hart (39) |
| Lysines between 302 and 534 | Ligand-independent ubiquitination | Tateishi et al. (40) |
| Residues 535–595 (lysine 581) | Ligand-dependent ubiquitination | Lonard et al. (41) |
| Mod. site | Protein source | Enzyme | Peptide | Sequence | vMALDI | LC-MS/MS |
|----------|----------------|--------|---------|----------|---------|----------|
|          |                |        |         |          | MH⁺ | Precursor | Error | Msc score | Modified | t | Unmodified |
| Thr(Asp)² | Both Trypsin | 2–8 M | | AcTMTLHTK | A | 437.23⁺² | 33 | 25 |         |       |
| Both Glu-C | 2–22 M | | AcTMTLHTKASGMAHHQIQGNE | L | 775.07⁺³ | 13 | 72 |         |       |
| Ser(P)¹⁰² | MCF7 Chymo. | 90–108 F | | GSGNLGFPPVLPSVSPSPL | M | 1876.9 27 | 938.97⁻¹² | 21 | 9 | 28.6 | 24.8 |
| Ser(P)¹⁰⁴ | Both Chymo. | 90–108 F | | GSGNLGFPPVLPSVSPSPL | M | 1876.9 25 | 938.94⁻¹² | 1 | 17 | 26.6 | 24.8 |
| Ser(P)¹⁰⁶ | Both Chymo. | 90–108 F | | GSGNLGFPPVLPSVSPSPL | M | 1876.9 24 | 0 | 17 | 26.6 | 24.8 |
| Ser(P)¹¹⁸ | Both Chymo. | 109–120 L | | mLHPPPQLpSFP | L | 1472.7 10 | 736.85⁻¹³ | 9 | 16 | 37.6 | 36.72 |
| Both Chymo. | 109–120 L | | mLHPPPQLpSFP | L | 1456.4 11 | 728.88⁻¹³ | 39 | 18 | 48.5 | 47.06 |
| rER Chymo. | 109–130 L | | mLHPPPQLpSFP | L | 1474.7 10 | 868.82⁻¹³ | 67 | 15 | 49.8 |       |
| rER Glu-C | 109–130 L | | mLHPPPQLpSFP | L | 1463.6 10 | 868.85⁻¹³ | 39 | 9 | 50.1 |       |
| Both Chymo. | 111–120 L | | mLHPPPQLpSFP | L | 1212.4 7 | 606.77⁻¹² | 27 | 35 | 39.8 | 37.59 |
| MCF7 Asp-N | 118–132 L | | pSPFQLPHQQVYPY | L | 927.54⁻¹² | 119 | 61 | 33.1 |       |
| Ser(P)¹⁰⁵ | Both Trypsin | 143–158 R | | EAGPAFYRNNpSDNRR | Q | 1927.0 6 | 642.97⁻¹³ | 33 | 10 | 27.8 | 26.97 |
| Both Asp-N | 143–154 R | | EAGPAFYRNNpS | D | 1385.7 |       |       |       |       |       |
| Ser(P)¹⁶⁷ | MCF7 Asp-N | 155–169 S | | DNRRQGQRERApSTN | D | 453.25⁻¹⁴ | 81 | 17.3 | 16.9 |       |
| MCF7 Glu-C | 164–177 E | | RLPSTNDKGSMAmE | S | 541.61⁻¹³ | 105 | 36 | 17.9 | 17.83 |
| MCF7 Trypsin | 165–180 R | | LAPSTNDKGGMAMESAK | E | 574.29⁻¹³ | 81 | 35 | 15.2 | 13.93 |
| MCF7 Trypsin | 165–180 R | | LAPSTNDKGGMAMESAK | E | 579.57⁻¹³ | 3 | 14 | 13.5 | 12.94 |
| MCF7 Trypsin | 165–180 R | | LAPSTNDKGGMAMESAK | E | 584.90⁻¹³ | 11 | 33 | 12.7 | 12.37 |
| MCF7 Chymo. | 165–177 T | | LAPSTNDKGGMAMESAK | S | 484.22⁻¹³ | 50 | 12 | 25.7 |       |
| Ser(P)²¹² | Both Asp-N | 203–217 C | | EGKAKAFPRpSDGHN | D | 1778.9 |       |       |       |       |
| Ser(P)²³⁶ | rER Lys-C | 236–244 K | | pSQCACRLR | C | 420.19 | 6.2 | 5 | 16.1 | 15.5 |
| Ser(P)²⁹⁴ | Both Lys-C/Asp-N | 285–298 E | | DMRAANLWPpSPLMIK | R | 608.15⁻¹³ | 80 | 22 | 40.1 | 38.93 |
| MCF7 Chymo. | 287–296 M | | RAANLWPpSPLMIK | M | 1204.8 |       |       |       |       |
| MCF7 Trypsin | 288–299 R | | AANLWPpSPLMIK | R | 1420.7 |       |       |       |       |
| MCF7 Trypsin | 288–299 R | | AANLWPpSPLMIK | R | 1436.7 | 718.88⁻¹² | 35 | 20 | 23.4 | 22.44 |
| Lys(Asp)²⁷² | Both Trypsin | 468–477 K | | SLEAAcDKDIHR | V | 1305.7 |       |       |       |       |
| Ser(P)²⁶⁴ | rER Chymo. | 549–555 R | | LHAPTSP | G | 861.4 5 | 431.21⁻¹² | 23 | 21 | 18.1 | 14.83 |
| rER Chymo. | 550–568 L | | HAAPTSP | A | 686.96⁻¹³ | 0.38 | 15 | 13.6 | 12.69 |
| Ser(P)²⁵⁹ | Both Trypsin | 556–577 G | | GGApSVEETDQSHLATAGSTSSHSLQ | Y | 2654.8 8 | 667.10⁻¹⁴ | 77 | 52 | 27.1 | 26.4 |
| rER Chymo. | 556–577 R | | GGApSVEETDQSHLATAGSTSSHSLQ | S | 552.9⁻¹⁴ | 148 | 12 | 25 |       |
peak. N-terminal fragments b₁₂ and below show no evidence of phosphate, whereas b₁₃ and above are all b

/₁₁₀⁰² sug-

suggesting phosphorylation at Ser 102. Another spectrum (not shown) gave similar fragments, but a distinctive b₁₃ ion (m/z 1198.2) indicated some molecules not phosphorylated at Ser102, a n day6/₁₁₀⁰² ion (m/z 581.2) implied phosphorylation of Ser104.

Peptides possessing both two and three phosphate groups prove that multiple serine residues can be phosphorylated. Supplemental Fig. 1E shows vMALDI-MS/MS of a doubly phosphorylated peptide of m/z 1956.5. The y₁₁ and y₁₁₁/₁₁₀⁰² ions confirm two phosphates C-terminal to Phe₉⁷ not involv-

ing Ser₉₁, whereas b₁₅ - 98 and b₁₆ - 98 - H₂O ions suggest that Ser₁₀⁶ is not phosphorylated. Supplemental Fig. 1F shows a triply phosphorylated species of m/z 2036.6. The stoichiometry for triple phosphorylation is clearly low, but three consecutive neutral losses of 98 Da confirm all three serine residues can be phosphorylated simultaneously.

The DBD—A new phosphorylation site was detected at Ser₂¹² within the loop between the two zinc fingers of this domain. Two Asp-N peptides separated by 80 Da gave similar fragments in vMALDI-LTQ-MS/MS. The unphosphorylated MH⁻ (m/z 1778.9) showed losses of water and ammonia and weak backbone fragments even at increased collision energy (Fig. 5A). Fragment ions b₉, b₁₃, b₁₄, y₇, y₉ - NH₃, y₁₃, and y₁₃ - NH₃ identified the peptide as ³⁰³EGCKAFFKRSQGHN with cysteine modified by iodoacetamide. The heavier mass MH⁻ (1858.8) gave a weaker spectrum (Fig. 5B) but showed loss of phosphoric acid (-98 Da) and additional losses of water and ammonia. An N-terminal fragment common to both spectra was identified as b₁₄ - H₂O for the unmodified peptide and b₁₄ - H₃PO₄ for the phosphopeptide. The heavier molecular ion also retained phosphate but lost combinations of water, ammonia, and carbon dioxide. The peptide is clearly phosphorylated, and the only possible location for a phos-

phate group in this peptide is Ser₂¹².

It was hypothesized that such a phosphopeptide with N-terminal glutamate might undergo the combined loss of phosphoric acid and water to form a new ionic moiety containing dehydroalanine and pyroglutamate. Identical fragmentation was observed for an analogous peptide incorporating the known Ser₁⁵⁴ site. Fig. 5C shows MS/MS of the unmodified peptide of m/z 1305.6 with (MH - NH₃ - H₂O)⁻ as the base peak plus b₁₁, b₁₂, b₁₃ - H₂O, b₁₄ - H₂O, b₁₅ - H₂O, y₁₁, y₁₁₀, and y₁₀ that confirm this peptide as ¹⁴³EAGPPAFYRPNS. Fig. 5D shows the spectrum of the phosphopeptide at m/z 1385.6.

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**Estrogen Receptor Posttranslational Modifications**

**Fig. 5. Identification of novel phosphorylation site Ser²¹² and comparison with a structurally similar phosphopeptide containing Ser(P)¹⁵⁴.** A, vMALDI-LTQ-MS/MS of MH⁻ for the Asp-N peptide ³⁰³EGCKAFFKRSQGHN (m/z 1778.9) from endogenous ERα with expansion of weak fragment peaks at m/z 800–1600. B, the corresponding MH⁻ spectrum from the phosphorylated peptide (m/z 1858.8). C, vMALDI-LTQ-MS/MS of MH⁻ for the Asp-N peptide ¹⁴³EAGPPAFYRPNS (m/z 1305.6) from endogenous ERα. D, the corresponding MH⁻ spectrum from the phosphorylated peptide (m/z 1385.6). pS, phosphoserine.
Again the base peak corresponds to combined losses of phosphoric acid and water, and other prominent peaks combine losses of phosphoric acid, H₂O, NH₃, and CO₂. The only significant backbone fragment (b₉) also shows water loss; thus such fragmentation appears to characterize phosphoserine peptides with N-terminal glutamate.

A previously reported phosphorylation site falling within the second zinc finger of the DBD at Ser²³⁶ was confirmed here in rER/H₉₂₅₁ only. LC-MS/MS of a Lys-C digest of rER/H₉₂₅₁ identified the triply charged peptide²³⁶SCQACRLRK (with iodoacetamide-derivatized cysteines) with and without serine phosphorylation; the modified form eluted half a minute later. Supplemental Fig. 2A represents the unmodified MH⁺³⁺ (m/z 393.5) with fragment ions MH₃⁻⁻, NH₃⁺, y₆⁺²⁺, y₇⁺²⁺, and y₈⁺²⁺. Supplemental Fig. 2B shows the corresponding spectrum of the phosphorylated MH₃⁺⁺ (m/z 420.2) with base peak corresponding to the loss of phosphoric acid (triply charged) and y₆⁺²⁺, y₇⁺²⁺, and y₈⁺²⁺ ions that locate the modification at Ser²³⁶.

The Hinge Region between the DBD and AF-2—Although no phosphorylation has been reported for this region, residues 263–301, we identified Ser²⁹⁴ as a phosphorylation site in MCF-7 cells. Fig. 6A shows the vMALDI-LTQ-MS/MS of MH₁₁₀₀₁ from chymotryptic peptide ²⁸⁷RAANLWSPL (m/z 1124.8). B, the corresponding MH⁺ spectrum from the phosphorylated peptide (m/z 1204.5). C, MS³ spectrum of (MH – 98)⁺ (m/z 1106.5) from the phosphopeptide MH⁺. D, LC-MS/MS spectrum of MH₃⁺⁺ of phosphorylated Lys-C/Asp-N peptide ²⁸⁵DMRAANLWSPMLIK (m/z 608.15). pS, phosphoserine; dS, dehydroalanine.
NH\textsubscript{3}, b\textsubscript{9}, b\textsubscript{g} + H\textsubscript{2}O, and MH\textsuperscript{+} \(-2\text{NH}_{3}\) identify \textsuperscript{287}RAANLWPSPL. The N-terminal arginine is consistent with predominant b ions. Fig. 6B for the analogous species 80 Da heavier (m/z 1204.5) shows (MH – 98\textsuperscript{+}) at m/z 1106.5, which was selected for MS\textsuperscript{3} analysis (Fig. 6C). This gives identical fragments to MS\textsuperscript{2} of the unmodified peptide for ions b\textsubscript{g} and below, but b\textsubscript{9} – NH\textsubscript{3}, b\textsubscript{g}, b\textsubscript{h} + H\textsubscript{2}O, and (MH – 2NH\textsubscript{3})\textsuperscript{+} are all 18 Da lower, confirming that the phosphate is situated between the b\textsubscript{g} and b\textsubscript{h} positions, i.e. Ser\textsuperscript{294}. Further evidence for this new site came from a 15-residue Asp-N/Lys-C peptide, \textsuperscript{285}DMR-LWPSPLMIK. The LC-MS/MS spectrum of MH\textsubscript{3} \textsuperscript{3+} (m/z 608.15) is illustrated in Fig. 6D; Ser(P)\textsuperscript{294} is confirmed by the y\textsubscript{9}, y\textsubscript{7}, and y\textsubscript{7} ions.

The Ligand-binding Domain—The largest domain of ER\textsubscript{\alpha} extends from residue 302 to the C terminus at residue 595 and contains AF-2. The extreme C-terminal region 552–595 (the F domain) may have a specific modulatory function that affects the agonist/antagonist effectiveness of antiestrogens and the transcriptional activity of ligand-bound ER in cells. No phosphorylation sites were identified within AF-2, but we found two novel phosphorylation sites in the F-domain at Ser\textsuperscript{294} and Ser\textsuperscript{559}. The assignment of Ser\textsuperscript{559} was based on observations made with recombinant protein, not with MCF-7 protein, so it is illustrated in supplemental Fig. 3. Panel A shows a vMALDI-MS/MS spectrum, and panel B shows an LC-MS/MS spectrum for a tryptic phosphopeptide of sequence \textsuperscript{549}LHAPTSR phosphorylated at either Thr\textsuperscript{553} or Ser\textsuperscript{554}. Both spectra show strong fragment ions, including the loss of phosphoric acid. Only the vMALDI spectrum contains a strong b\textsubscript{8} fragment ion that confirms the phosphate group to be on Ser\textsuperscript{554} rather than Thr\textsuperscript{553}.

Fig. 7 shows MS/MS spectra that identify phospho-Ser\textsuperscript{559} in MCF-7-derived ER\textsubscript{\alpha} digested with trypsin. Fig. 7A shows the LC-MS/MS spectrum of the unmodified peptide MH\textsubscript{3} \textsuperscript{3+} (m/z 862.4) eluting at 26.4 min and identified as \textsuperscript{556}GGAS-VEETDQSHLATAGSTSSHSLQK. Fig. 7B shows the corresponding spectrum of the modified form (m/z 889.1) eluting at 27.1 min. This 26-residue peptide contains six serines and three threonines, any one of which could potentially be the phosphorylation site. Comparing the two spectra, the similarity of the y-series ions indicates that the modification is not in the C-terminal region, whereas strong b\textsubscript{9} – b\textsubscript{7} ions show the loss of phosphoric acid, which unambiguously establishes Ser\textsuperscript{559} as the site of the modification. The same peptide was also observed by vMALDI. Fig. 7C shows the vMALDI-MS\textsuperscript{2} spectrum of the unmodified peptide at m/z 2585.3. Fig. 7D shows the species 80 Da heavier at m/z 2665.3, and Fig. 7E represents the MS\textsuperscript{3} spectrum of the (M – 98\textsuperscript{+}) at m/z 2567.2. Although these spectra show numerous fragment ions that clearly define the sequence, no unique ions identify the specific location of the phosphate group, unlike the LC-MS/MS data.

Quantitation of Ser\textsuperscript{294} Phosphorylation by LC-MRM/MS

To determine the biological relevance of these previously unreported phosphorylation sites an MRM/MS assay was developed to monitor any change in Ser\textsuperscript{294} phosphorylation following estradiol treatment of MCF-7 cells. MRM is the most sensitive and robust quantitative mass spectrometric method because of (i) coincident detection of both the precursor ion (Q1) and a designated fragment ion (Q3) and (ii) rapid cycling that allows each transition to be sampled multiple times during chromatographic elution of the selected species. Fragment ions with m/z values greater than that of the precursor ion lead to lower noise levels, which requires that the fragment ion has a lower charge state than the precursor. The LC-MS/MS spectra in Fig. 8, A and B, of tryptic peptide \textsuperscript{288}AANLWPSPLMIK with and without phosphorylation of Ser\textsuperscript{294} show that the most abundant fragment ions that fit this criterion are y\textsubscript{7} – phosphoric acid (m/z 767.48) from the doubly charged Ser\textsuperscript{294} phosphopeptide (m/z 710.9) and y\textsubscript{7} (m/z 785.47) from the doubly charged unmodified peptide (m/z 670.9).

Because of the high specificity of MRM, gel isolation of the immunoprecipitated ER\textsubscript{\alpha} from background antibody proteins, protein A, and other contaminants was not required before trypsin digestion. This eliminated the sample losses typical of in-gel digestion and also minimized methionine oxidation, significantly increasing the peptide signal strength as this was not split between the oxidized and non-oxidized forms. Fig. 8C illustrates MRM data for the doubly protonated molecular ion fragmenting to y\textsubscript{7} for unmodified and phosphorylated Ser\textsuperscript{294} peptides from immunoprecipitated ER\textsubscript{\alpha} obtained from control conditions and following 30-min incubation with 10 nM estradiol. As expected, the phosphopeptide eluted 1–2 min later than the unmodified peptide. The increased peak heights following estradiol treatment relative to control conditions clearly indicate significant induction of phospho-Ser\textsuperscript{294}. The fold induction shown in Fig. 8D was obtained by dividing the phospho-Ser\textsuperscript{294} integrated peak areas by the corresponding Ser\textsuperscript{294} integrated peak areas normalized to the value obtained for the control sample. Three biological replicates demonstrated a mean 27.8-fold induction of phosphorylation following estradiol treatment relative to control with a standard deviation of 9.8.

DISCUSSION

Previous comprehensive mapping of ER\textsubscript{\alpha} and its PTMs by mass spectrometry had been hampered by the inability to purify/enrich sufficient endogenous protein. The optimization of ER\textsubscript{\alpha} immunoaffinity enrichment from MCF-7 cells for MS analysis reported here is a significant achievement in ER\textsubscript{\alpha} biochemical studies as demonstrated by our recent observation of the novel phosphorylation site Ser\textsuperscript{154} in the N-terminal domain of ER\textsubscript{\alpha} (27).
Several PTMs of ERα summarized in Table II and Fig. 9 were identified in this tandem mass spectrometry study, which focused on detection and characterization of phosphorylation sites. The majority of phosphorylation sites identified here occur in endogenous protein isolated from the estradiol-stimulated MCF-7 human breast cancer cell line (Ser102, Ser104, Ser106, Ser118, Ser154, Ser167, Ser212, Ser294, and Ser559), although two were observed only in recombinant protein (Ser236 and Ser554). Ser154 was a novel phosphorylation site previously detected by us using the same methods, but several other sites had not been reported previously: namely Ser212, Ser294, Ser554, and Ser559. Most or all of these serine phosphorylations are likely estradiol-dependent, although evaluation of the induction level and functional consequences for these various residues are not yet completed. However, in regard to Ser294, Fig. 8 demonstrates that estra-

Fig. 7. Identification of novel phosphorylation site Ser559 in endogenous MCF-7 ERα. A, LC-MS/MS spectrum of MH + of tryptic peptide 556GGASVEETDQLATAGSTSSSHSLQK (m/z 889.1) eluting at 26.4 min. B, LC-MS/MS spectrum of MH + of the analogous phosphorylated peptide (m/z 889.1) eluting at 27.1 min. C, vMALDI-LTQ-MS/MS spectrum of MH + of the unmodified peptide (m/z 2585.3). D, the corresponding spectrum of MH + of the phosphorylated peptide (m/z 2665.3). E, MS3 spectrum of (MH – 98)+ (m/z 2567.2) from the phosphopeptide. pS, phosphoserine; dS, dehydroalanine.
diol stimulation of MCF-7 cells produced an approximate 28-fold phosphorylation enhancement of this residue, strongly implicating its functional relevance to ER activation. It is interesting to note that an early study examining the phosphorylation of ER expressed in COS-1 cells failed to detect Ser\(^{294}\) phosphorylation (29). We note that our studies differ from these initial studies in several key aspects including our inclusion of the potent phosphatase inhibitor okadaic acid in the immunoprecipitation buffer; the use of MCF-7 cells, which endogenously express ER instead of the more artificial system.

**FIG. 8.** LC-MS/MS of tryptic peptides from endogenous MCF-7 ER\(\alpha\).

A, spectrum of MH\(^{3+}\) (m/z 710.9) of peptide 28AANLWPpsPLMK (where pS is phosphoserine). B, spectrum of MH\(^{3+}\) (m/z 670.9) of the analogous unphosphorylated peptide. C, MRM/MS analysis of Ser\(^{294}\) induction following estradiol (E2) treatment showing chromatograms of the \(y_7\) Q1/Q3 MRM transitions for the unmodified (670.9/785.5) and phosphorylated (710.9/865.4) peptide from control and estradiol-treated (10 nM; 30 min) cells. D, comparing peak areas for the phosphorylated with unmodified peptides demonstrates that estradiol induces Ser\(^{294}\) oxidation 28-fold. The error bar represents the standard deviation for three biological replicates. XIC, extracted ion chromatogram; cps, counts/s.

**FIG. 9.** Summary of ER\(\alpha\) modifications observed in this study. ** signifies previously unreported modifications identified in ER\(\alpha\) from MCF-7 cells. * signifies a previously unreported modification identified only in rER\(\alpha\). In parentheses previously known modification identified here only in rER\(\alpha\). All others are previously known modifications identified in ER\(\alpha\) from MCF-7 cells. LBD, ligand-binding domain; NTD, N-terminal domain; pSer, phosphoserine.
of ER expressed in COS-1 cells; and an estradiol exposure time of 30 min in our experiments versus the 4 h used in the earlier study.

Previously reported phosphorylation sites Ser\textsuperscript{305}, Thr\textsuperscript{311}, and Tyr\textsuperscript{537} were not confirmed here, and Ser\textsuperscript{336} phosphorylation was observed only in recombinant protein. However, Ser\textsuperscript{336} is phosphorylated by cAMP-dependent protein kinase (ligand-independent) so 30-min treatment with estradiol may not induce phosphorylation in MCF-7 cells, although 1–5-min exposure to estradiol might activate the cAMP-dependent protein kinase pathway through membrane ER activation (15). Residue 305 is in a region for which no peptide was observed so no data were obtained to support or refute its phosphorylation. However, it is phosphorylated by PAK1 (growth factors heregulin and epidermal growth factor induce PAK1 activation), which again is ligand-independent (30). Unmodified Thr\textsuperscript{311} was identified by LC-MS/MS in chymotryptic peptide\textsuperscript{306}SLTADQMVSSALL with good mass accuracy (an error of 19 ppm) and a significant Mascot score of 35 and probably would be observed as phosphorylated if that actually occurred. This residue was reported phosphorylated by p38 kinase and was estradiol-dependent in cells overexpressing ER and tagged p38 using \textit{in vitro} kinase assays (31). We observed unmodified Tyr\textsuperscript{537} peptides with trypsin (532–548), Glu-C (524–538), and Asp-N (527–537), so any phosphorylation of this residue would probably be observed, but phosphorylation may not be induced by 30-min estradiol treatment as it may be Src-dependent (32).

This study demonstrates the sensitivity of tandem mass spectrometric methods for detection of phosphorylation sites in low level proteins such as ER\textalpha, and it reiterates the advantages of using complementary methods of ionization and analysis as no single method was able to detect all the modifications reported here. Additional modifications detected in ER\textalpha from MCF-7 cells were N-terminal cleavage of methionine accompanied by acetylation of Thr\textsuperscript{2} and side chain acetylation or possibly trimethylation of Lys\textsuperscript{472}.

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