Phosphoproteomics Reveals Key Regulatory Kinases and Modulated Pathways Associated with Ovarian Cancer Tumors

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Background: Ovarian cancer (OC) is the seventh most common cancer worldwide for women. However, there are no sufficient diagnostic methods and few treatment options available due to poor understanding of its pathogenic mechanisms.

Methods: To comprehensively analyze the phosphoproteomic characterization for OC, we took advantage of a quantitative global phosphoproteomics method, titanium(IV) immobilized metal affinity chromatography (Ti4+-IMAC) coupled to nanoscale liquid chromatography and quadrupole time-of-flight tandem mass spectrometry (nanoLC/Q-TOF-MS/MS) on ovarian tissue samples obtained from five OC patients and five matched controls.

Results: A total of 722 phosphorylated sites corresponding to 534 proteins were significantly different (fold change ≥ 2, p < 0.01) between OC patients and the controls. Among them, 83 transcription factors mainly consisted of transcription cofactors, zf-C2H2, and chromatin remodeling factors and 29 kinases were included. Further functional analysis suggested significantly biological processes were highly enriched and involved in the pathogenesis of OC, especially fructose and mannose metabolism. Moreover, the regulatory roles of modulated pathways, including MAPK, ErbB, and GnRH signaling pathways were also identified as critical processes involved in OC. The results here highlighted key phosphorylated proteins, particularly kinases, and the corresponding cancer-related metabolic and signal pathways that played important roles in the development of OC. Additionally, the expression levels of two kinases, phosphorylated CDK (T14) and phosphorylated PRKCQ (S695), were validated by Western blot analysis in the other group of ovarian tissue samples.

Conclusion: Altogether, our data not only provided novel insights into the potential biomarkers and therapy options for OC but also extended our knowledge on its pathophysiological mechanism.

Keywords: ovarian cancer, phosphoproteomics, Ti4+-IMAC, kinases, biomarker

Introduction

Ovarian cancer (OC) is the seventh most common cancer worldwide for women.1 Especially in the United States, OC is the leading cause of death among fatal gynecologic malignancies. There were approximately 22,440 new cases of ovarian cancer diagnosed and 14,080 OC deaths in 2018 in the United States.2 Unfortunately, OC is a disease with a poor prognosis, which frequently has no obvious symptoms at its early stages.3 It is usually advanced when it is diagnosed but with low survival rates ranging from 30% to 50%. Nevertheless, risk factors for OC remain largely unknown and there are few promising therapy options for
patients, which motivated researchers to characterize the molecular changes associated with OC development and to identify potential biomarkers for diagnostic purposes.

In the last decade, a series of critical discoveries have clearly demonstrated that OC is a highly heterogeneous disease with a wide spectrum of distinct clinical and molecular entities.\(^4\) Generally, epithelial ovarian neoplasms can be divided into three major groups: borderline, Type I and Type II tumors. Among them, Type I tumors are genetically stable, carrying a distinct set of frequently mutated genes such as BRAF, KRAS, CTNNB1, and PTEN. On the other hand, Type II tumors are the most common type of OC and comprise high-grade serous (HGS) tumors, primarily responsible for the low survival rate.\(^4\) The rapid development of high-throughput omics techniques including genomics, transcriptomics, proteomics, and metabolomics has dramatically extended our understanding on systems biology of OC. The Cancer Genome Atlas (TCGA) performed both genomic and transcriptomic characterization of ovarian high-grade serous carcinoma (HGSOC) and identified the critical role of mutations in the TP53, BRCA1 and BRCA2 genes.\(^5\) Other key findings were also determined including extensive DNA copy alterations and CCNE1 aberrations in HGSC. Moreover, further analysis of genomic data from the TCGA consortium and the integration of microRNA and mRNA expression profiles associated with HGS were conducted to improve the statistical association with patient outcome and identify candidate microRNA targets.\(^6,7\)

Besides, proteomics and metabolomics have recently emerged as powerful technologies for the identification of cancer-specific proteins and metabolites, which provides potential biomarkers for early diagnosis of OC and expands the insights of their pathophysiological mechanisms. Proteomics allows detecting both qualitatively and quantitatively thousands of proteins and/or peptides in large clinical studies. Coscia et al employed quantitative proteomics on FFPE tumor samples from 25 chemotherapy-naive patients with advanced-stage HGSOC.\(^8\) The results indicated that cancer/testis antigen 45 (CT45) enhances chemosensitivity in metastatic OC, which plays a possible role in the DNA damage response. Importantly, the expression of CT45 may improve the treatment efficacy of immunotherapy or platinum-based chemotherapy for patients with advanced-stage OC. Moreover, Tajmul et al used fluorescence-based 2D-DIGE coupled with MALDI/TOF-MS to profile the salivary protein signatures for non-invasive detection of OC.\(^9\) Forty-four significantly different expressed proteins \((p < 0.05)\) between cancer patients and healthy controls were determined, three of which including lipocalin-2, indoleamine-2, 3-dioxygenase1 and S100A8 were further validated using Western blotting and ELISA. Their work indicated that salivary proteomics might be exploited as a diagnostic biomarker for the screening of OC after further large-scale validation.

Another powerful technology is metabolomics, aiming to simultaneously detection of small molecule metabolites in biological specimens. Likewise, it holds great potential for discovering potential biomarkers for early diagnosis of the disease, and sheds insight into key regulatory metabolic pathways involved in its pathophysiology, thus leading to new therapeutic strategies. For example, Zeleznik et al conducted a prospective analysis of circulating plasma metabolomics and ovarian cancer risk and revealed that an important metabolite, pseudouridine, may be a novel risk factor for OC.\(^10\) And meanwhile, TAGs were also determined to play important roles especially on rapidly fatal tumors.

Recently, phosphoproteomics has been an emerging tool in the study of cancer for the discovery of potential biomarkers and therapy targets.\(^11\) Protein phosphorylation is considered to be one of the most abundant post-translational modifications (PTMs), acting as a key regulator of a series of subcellular processes including proliferation, apoptosis and tumorigenesis. In order to obtain a global view of protein phosphorylation events, here we employed Ti\(^{4+}\)-IMAC coupled with nano-LC-MS/MS technology to comprehensively characterize the phosphoproteomics profiles of OC patients and controls. We aimed to not only find out potential biomarkers and therapy options for OC but also provide novel insights into its pathophysiological mechanism.

### Materials and Methods

#### Subjects

The present study was approved by the local ethics committee of Peking Union Medical College Hospital (PUMCH) and was conducted in accordance with the Declaration of Helsinki, while written informed consent document for each participant was obtained. Totally 20 women were recruited, including 10 patients all with epithelial ovarian cancer (EOC) and 10 matched patients without OC as the controls. Their clinical characteristics were shown in Table S1. All OC patients were defined as serous histological type (type II, G3) without LNM. And meanwhile, their FIGO characterizations were also shown, mostly ranging from stage II to IV. All patients with OC had been diagnosed without any disease of metabolic,
kidney, liver and pelvic inflammatory, as well as any other cancers. Additionally, 10 matched patients in the control group had no history of OC and mainly had hysteromyoma instead. All patients had undergone gynecologic surgery and their diagnosis had been validated by histopathologic examinations before sampling.

Sample Preparation and Pretreatment for Phosphoproteomics Analysis

All collected tissue samples were ovarian surface epithelium and immediately stored at −80°C prior to further analysis. 5 µL samples were diluted with lysis buffer containing 8M Urea (Sigma, MO, USA), 1 mM EDTA, 1 mM PMSF, 100 mM Tris-HCL (Sigma, MO, USA, pH 8.5); then, centrifuged at 15000 g for 15 min at 4°C to remove the sediment. The concentration of protein was measured according to the BCA protein assay kit (Bi Yuntian, Shanghai, China). For phosphoproteomics analysis, the extracted proteins were further reduced for 60 min at 35°C with 10 mM dithiothreitol and alkylated with 50 mM iodoacetamide for 40 min at room temperature in the dark. Subsequently, protein digestion was conducted by the filter-aided sample preparation (FASP) protocol with trypsin (Promega, Madison, WI) in 100 mM NH₄HCO₃ (Sigma, MO, USA). The concentration of peptide was determined by NanoDrop 2000 instrument (Thermo Scientific, USA) at an absorbance of A280 nm. Moreover, phosphopeptide enrichment was carried out by Ti⁺⁺-IMAC as previously reported. Phosphopeptides were finally diluted from the beads by 10% ammonium hydroxide and dried by vacuum centrifugation.

Nano-HPLC-MS/MS Analysis

Afterwards, the peptides were diluted with 10 µL solvent A (water with 0.1% formic acid) and further analyzed using on-line nanospray LC-MS/MS on an Orbitrap Fusion lumos connected to an EASY-nLC system (Thermo Scientific, MA, USA). For LC-MS/MS analysis, 5 µL peptide sample was loaded onto the trap column (Thermo Scientific Acclaim PepMap C18, 100 µm x 2 cm) at a flow rate of 5 µL/min and subsequently separated on the analytical column (Acclaim PepMap C18, 75 µm x 15 cm) using a 120-min non-linear gradient, ranging from 3% solvent B (ACN with 0.1% formic acid) to 32% solvent B. The column flow rate was 300 nL/min. The electrospray voltage of 2.1 kV relative to the inlet of the mass spectrometer was used. The mass spectrometer was operated under data-dependent acquisition mode, and automatically switched between MS and MS/MS mode in 3.5 s cycle. MS1 mass resolution was set as 120 K with m/z 350–1550, MS/MS isolation window was set 1.6 Da, and MS/MS resolution was set as 30 K under HCD mode with a collision energy was 32%. The dynamic exclusion was set as n=1, and the dynamic exclusion time was 30 s, AGC target was 5e4, max injection time was 120 ms.

Data Analysis

MaxQuant (version 1.5.2.8) was employed here for all data processing. The data were searched against a UniProt human proteome fasta file amended to included common contaminants. A decoy database containing reverse sequences was used to estimate false discovery rates and set the false discovery rate at 1%. Default MaxQuant parameters were used with the following adjustments: Phospho (STY) was added as a variable modification. Match between runs was selected. We obtained the high confidence phosphosite with the following threshold: a PTM site localization probability of 0.75, an Andromeda score of 40 and an Andromeda delta score of 6. The intensity of phosphosite was further pre-processed by metaX. Phosphosites with missing values were imputed with the minimum value to further improve the data quality. Intensity of each phosphosite was normalized with variance stabilizing normalization method. The following criteria were adopted for screening differentially expressed phosphosite: fold change ≥ 1.5 or ≤ 0.67 and p value ≤ 0.05. Here, p values were adjusted using Bonferroni corrections. Functional annotations were further performed by BLAST these proteins against the GO (http://www.geneontology.org/), KEGG (http://www.genome.jp/kegg/pathway.html), COG (http://www.ncbi.nlm.nih.gov/COG/) and eggNOG (http://eggnogdb.embl.de/). Meanwhile, we employed WoLF PSORT to predict protein subcellular localization and used AnimalTFDB for transcription factor analysis.

In the GO and KEGG pathway enrichment analysis of differential phosphoproteins, we compared the significant phosphoproteins to all of the identified proteins, as a background, and the hypergeometric test formula used in these analyses was as follows:

\[ P = 1 - \sum_{i=0}^{m-1} \binom{M}{i} \frac{(N-M)}{n-i} \binom{N}{n} \]

where N is the number of GO/KEGG pathway entries in all identified proteins; n is the number of GO/KEGG
pathway entries that represent significant differentially abundant proteins; M is the number of GO/KEGG pathway entries that can be matched to all identified proteins; and m is the number of GO/KEGG pathway entries that can be matched to a significant differentially expressed protein. If the p-value of the hypergeometric test was less than 0.05, the differential protein was significantly enriched in the GO/KEGG pathway entry.

In addition, a multi-omics data analysis tool, OmicsBean (http://www.omicsbean.com), was used in the present study for protein–protein interaction network analysis. P-values < 0.05 were considered to be significantly enriched KEGG pathways.

**Western Blot Analysis**

For Western blot analysis, extracted proteins were separated by sodium dodecyl sulfate-polyacrylamide gels (SDS-PAGE) and electroblotted onto polyvinylidene difluoride (PVDF) membranes.14 The antibodies used here were: Rabbit anti-PRKCQ (phosph S695) antibody (Cat. # BS-5584R, Bioss), rabbit anti-CDK2 (phospho T14) antibody (Cat. # ab68265, Abcam), and mouse anti-β-actin antibody (Cat. # GB12001, Servicebio). The final films were photographed and analyzed by using an AlphaEaseFC software (Alpha Innotech). β-actin was used as the loading control.

**Results**

**Phosphoproteomic Profiling of Human Ovarian**

To comprehensively analyze the phosphoproteomic characterization for human ovarian, we took advantage of the label-free method for 10 samples from the patients with OC (Table S1, OC1-OC5) and the controls (Table S1, OV1-OV5). A total of 9619 different phosphopeptides (10,934 unique phosphorylation sites) mapped to 3920 proteins were identified across the test 10 samples, which contained 91.5% phosphoserine sites, 8.2% phosphothreonine sites and 0.3% phosphotyrosine sites (Figure 1). Moreover, 83.8% singly phosphorylated phosphopeptides, 14.3% doubly phosphorylated phosphopeptides and 1.9% phosphopeptides phosphorylated at more than three sites were included in the detected 9619 phosphopeptides. In addition, most of the 3920 proteins had a large number of phosphosites. Especially, more than 10 phosphorylated proteins had nearly four thousands of phosphosites to perform various biological functions.

The identified 3920 proteins were further annotated according to multiple databases including COG, GO, and KEGG database. The publicly available program WolfPsort was also employed here for subcellular localization prediction. As shown in Table S2, the majority of 3920 proteins distributed predominantly in the nucleus and cytoplasm space. Finally, 2140 transcription factors and 193 kinases were identified among those proteins detected in OC tissues (Tables S3 and S4).

**Phosphoproteomic Characterization in Patients with Ovarian Cancer**

To overview the relationship between patients with OC and the controls, we employed pearson correlation analysis and principal component analysis (PCA) based on the intensities of each phosphosite. As shown in Figure 2A, the correlation coefficients between patients with OC ranged from 0.65 to 0.82, the average of which seemed to be higher than those between patients with OC and the controls, but lower than those between the controls. On the other hand, PCA score plots revealed that the samples from the patients with OC were tightly clustered together, but to some extent, separated from the controls (Figure 2B). The results here suggested that patients with OC had a significantly distinct phosphoproteomic profiling. Further statistical analysis indicated 722 out of 10,934 phosphorylated sites that corresponded to 534 proteins were significantly different (fold change ≥ 2, p < 0.01) (Figure 3 and Table S5). Of these phosphosites, 233 sites exhibited significantly greater phosphorylation in patients with ovarian cancer, while 489 phosphorylated sites exhibited less phosphorylation in patients with ovarian cancer than the controls.

**Functional Annotation of the Differentially Phosphorylated Proteins**

Gene ontology-based functional enrichment analysis of 534 differentially phosphorylated proteins was firstly performed to identify their possible functional significance. The results showed that these phosphorylated proteins were significantly enriched in 99 terms from the Component Ontology, 63 terms from the Function Ontology, and 326 terms from the Process Ontology (Tables S6–S8). According to RichFactor values, the top 20 most significantly enriched GO terms were shown in Figure S1, including cellular component organization, cellular component organization or biogenesis, protein
binding and binding. Moreover, the left 16 GO terms of the cellular component were also included, such as non-membrane-bounded organelle, intracellular non-membrane-bounded organelle, nuclear lumen and so on. To better understand metabolic pathways activated in the pathogenesis of OC, pathway enrichment analysis were further carried out. On one hand, all 534 differentially phosphorylated proteins were involved in 41 pathways, including signal transduction, cancers: overview, an infectious disease: viral (Figure S2). On the other hand, six pathways were finally determined to be significantly enriched: fructose & mannose metabolism, systemic lupus erythematosus, arrhythmogenic right ventricular cardiomyopathy (ARVC), spliceosome, RNA transport, and tight junction (Figure 4). Regarding the subcellular localization of 534 differentially phosphorylated proteins, 59.6% were localized in the nucleus, and 21.9% were localized in the cytoplasm (Figure S3). Moreover, among those differentially phosphorylated proteins, 83 transcription factors were included, most of which were transcription cofactors, zf-C2H2, and chromatin remodeling factors (Figure S4). Additionally, 29 kinases were also included in the differentially phosphorylated proteins, belonging to eight groups including AGC, CMGC, APK, and STE (Table S9). The top two groups were AGC, which mainly consists of protein kinases A, C and N, and CMGC, which mostly contains CDK kinases and DYRK families. Moreover, 12 kinases were with up-regulated sites, while the left 17 kinases were with down-regulated sites.

Figure 1 Summary of the phosphoproteomic profiling. (A) The percentages of phosphoserines, phosphothreonines and phosphotyrosines. (B) The distribution of phosphopeptides. (C) The number of phosphosite for phosphorylated proteins. (D) The number of phosphorylated protein annotated according to multiple databases including COG, GO, and KEGG database.
Protein–Protein Interaction Network Analysis of Significantly Changed Kinases

Kinases play central roles in various cellular signaling processes, and dysregulated kinase signaling has been observed in cancer.\(^\text{15}\) Therefore, we accordingly employed protein–protein interaction network analysis of significantly changed kinases to provide insights into the regulatory pathways involved in the developmental and physiological processes underlying OC. Fifty-three significantly enriched pathways were determined (Table S10), the top 10 of which are shown in Figure 5, involving 16 associated kinases. Obviously, kinases including PRKCA, CDK2, and EGFR belonging to several important families such as PKC, CDK, and EGFR were connected to these cancer-related pathways such as prostate cancer, bladder cancer, and non-small cell lung cancer. Signaling pathways including MAPK, ErbB, and GnRH were also included in the interaction network. Additionally, different kinases belonging to the same family may change in the opposite direction. For instance, in the PKC family, PRKCA and PRKACB were significantly down-regulated in OC patients, while PRKCD and PRKCQ were significantly up-regulated. The two kinases EGFR and ERBB2, all belonging to the EGFR family, also displayed conversely regulatory roles in OC pathogenesis.

Validation of the Expression Levels of p-CDK and p-PRKCQ

To confirm the expression levels of those significantly changed proteins, two important kinases, p-CDK2 (T14) and p-PRKCQ (S695) were chosen for Western blot analysis in the other group of ovarian tissue samples (Table S1) using commercially available antibodies. As shown in Figure 6, p-CDK2 (T14) and p-PRKCQ (S695) had significantly higher levels in OC patients than those in the controls ($p = 0.015$ and $p = 0.016$, respectively). The results here were obviously consistent with the findings in our phosphoproteomic study.
Discussion

OC remains one of the top 10 cancers worldwide with a high mortality rate, highlighting the importance of understanding its pathogenesis given no sufficient diagnostic methods and few treatment options available. Therefore, to comprehensively reveal the molecular characterizations of OC, herein we employed Ti^{4+}-IMAC coupled with nano-LC-MS/MS technology to perform phosphoproteomics analysis on tumor tissue samples and matched samples that served as the controls. The data revealed key phosphorylated sites, phosphorylated proteins and regulatory signal/metabolic pathways that are involved in OC pathogenesis, provided potential biomarkers for early diagnosis, and expanded the current knowledge of its physiopathology.

Figure 4 Pathway enrichment analysis of significantly different phosphorylated sites in OC patients.

Figure 5 The top 10 significantly enriched pathways according to protein–protein interaction network analysis of 29 significantly different kinases.
Phosphoproteomics has been demonstrated to be a very promising technology for the identification of diagnostic biomarkers and novel therapeutic targets for cancer. The success of phosphoproteomics especially relies on efficient phosphopeptide enrichment prior to mass spectrometric (MS) analysis. Among those selected methods, TiO$_2$ and Ti$^{4+}$-IMAC are typically applied to enrich phosphorylated peptides for phosphoproteomics analysis. Pinkse et al once revealed that TiO$_2$ has a strong affinity for phosphopeptides and thus indicated its high potential in the field of phosphoproteomics. Moreover, several studies indicated that Ti$^{4+}$-IMAC for enriching phosphopeptides could achieve a greater depth of coverage, with high specificity and very large dynamic ranges in large-scale phosphoproteome analysis. Particularly, Ti$^{4+}$-IMAC was determined to be the most efficient when systematically compared with some developed isolation methods such as TiO$_2$, ZrO$_2$, Fe$^{3+}$-IMAC, and Zr$^{4+}$-IMAC for their specificity and efficiency. In the present study, we took advantage of the powerful tool, Ti$^{4+}$-IMAC coupled with nano-LC-MS/MS, for an in-depth understanding of the biological functions of phosphoproteomes involved in OC development. To the best of our knowledge, there have been limited phosphoproteomics studies on OC by using the Ti$^{4+}$-IMAC method. Francavilla et al recently employed TiO$_2$-based chromatography for phosphoproteomics to report epithelial ovarian cancer signatures and 7658 distinct phosphorylation sites were determined from 13 independent patient samples. Here we identified 10,934 unique phosphorylation sites across the 10 tissue samples, much more than that in the previous study by Francavilla et al. Obviously, our phosphoproteomics study represented a larger number of phosphorylation sites in human OC tissues, corroborating the power of this method to reveal the molecular characteristics of the syndrome.

Altered metabolism in cancer cells, especially the Warburg effect, is well established as a hallmark of tumors. It is characterized by an increase in glucose uptake and lactate production, which is often accompanied by a decrease in oxidative phosphorylation. The aerobic glycolysis in tumor supports high energetic and biosynthetic demands to promote cell growth, proliferation, and metastasis. Although the pathway of glycolysis/glucogenesis was not significantly enriched in our study, another glycolysis-related pathway, fructose and mannose metabolism, was found to be significantly enriched based on 534 differentially phosphorylated proteins. Four glycolytic enzymes (PFKFB2, PFKL, ALDOA, and TPI1) mapped to fructose and mannose metabolism were significantly changed in OC, which may play important roles in glycolysis and further promote ovarian tumor growth. For example, PFKFB-2 is the cardiac isoform of the most critical glycolytic regulator, phosphofructokinase-2, while the PFKFB2 gene has determined to be overexpressed, ranking in the top 33% according to the TCGA database with more than 300 ovarian cancers. Meanwhile, knockdown of PFKFB2 could induce ovarian cancer cell growth inhibition and may enhance sensitivity to paclitaxel-based chemotherapy in OC patients. The results here revealed that the phosphorylation level of PFKFB2 on the Ser466 residue was much higher in OC patients than the controls, and its high activity probably further promotes glycolysis in ovarian cancer cells. Likewise, ALDOA plays an important role in glycolysis and keeping the maintenance of glucose homeostasis. We found a down-regulation of phosphorylated ALDOA protein in OC, though a large number of studies have reported high ALDOA expression
in other various types of tumors including osteosarcoma, lung cancer, oral squamous cell carcinoma, and hepatocellular cell carcinoma. Additionally, other two significantly down-regulated enzymes, PFKL and TPI1, were also reported to be involved in targeting glucose metabolism of human OC cells by ABT737.26

Because of their important roles in cell growth and proliferation, these four glycolytic enzymes and the corresponding pathway may be promising targets for treating ovarian cancer. Besides, 29 differentially expressed kinases were also identified in OC patients, which are key phosphoproteins involved in various cellular processes including signal transduction.27 Importantly, dysregulated kinase signaling usually occurs in many diseases including cancer and thus targeting specific kinases is expected of particular interest in cancer therapy. Castillo et al performed human testis phosphoproteome and indicated kinases as potential targets in spermatogenesis and testicular cancer.28 As respect for OC, certain kinases including CDK7 have been identified to be potential players in EOC development and highly associated with poor survival.21,29 In the present study, 29 kinases belonged to 22 kinase families were found to be associated with OC development, some of which were previously reported including CDK2, EGFR, MAP3K5, BRAF, and PKCa.21,29 Four well-known kinase families such as PKC, DYRK, CDK and EGFR were also included and were expected to play crucial roles in ovarian tumor growth. For example, Zhang et al revealed the PKC family plays a key regulatory role in various cancer-associated signal transduction pathways and indicated PKC\textsubscript{1} as a potential oncogene in ovarian carcinoma.30 Protein–protein interaction network analysis of those 29 kinases also suggested that 16 key kinases could modulate key signaling pathways and regulate tumor proliferation. Notably, different kinases belonging to the same family changed in the opposite direction, indicating their different roles in the physiological processes underlying OC.

There were still some deficiencies in this study. Firstly, fallopian tube epithelium (FTE) and ovarian surface epithelium (OSE) were both considered candidates for the cell-of-origin of HGSOC.31 Here, no samples of FTE were used for the comparison to high-grade serous carcinoma. It would be more accurate to parallelly use both FTE and OSE as the controls. Secondary, Ti\textsuperscript{4+}-IMAC beads do have high specificity and large dynamic ranges for the enrichment of various types of phosphopeptides without bias. However, because of its lower endogenous frequency of occurrence, the analysis of tyrosine phosphorylation still poses a larger challenge than threonine and serine phosphorylation.32 Therefore, other complementary methods such as immunoprecipitation (IP) should be employed to selectively probe tyrosine phosphorylation levels. Thirdly, the number and types of samples were limited. In the next steps, extensive validation of our findings on phosphoproteomics changes would be performed in a larger scale study that features adequate amounts of ovarian samples. Targeted strategies including parallel reaction monitoring assay are supposed to be used to demonstrate the changes of those important proteins and their phosphorylated forms. In addition, we would deeply reveal the biological roles of those important phosphorylated proteins involved in tumorigenesis of ovarian cancer according to gain and loss of function experiments in future work.

Conclusion
Altogether, the present study comprehensively revealed the molecular characteristics in relation to OC and identified significantly distinct phosphorylated sites and phosphorylated proteins underlying the syndrome. The results here dissected regulatory elements (especially kinases) and pathways involved in cancer-related pathways including fructose & mannose metabolism, MAPK, ErbB, and GnRH signaling pathway. These findings may not only provide potential biomarkers and novel therapeutic targets for OC but also bolstered our knowledge on the physiopathology of this syndrome.

Data Sharing Statement
All the raw mass spectrometry data in the present study have been deposited to the ProteomeXchange Consortium (http://proteomecentral.proteomexchange.org) via the iProX partner repository with the dataset identifier PXD016183.

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Disclosure
The authors declared no conflicts of interest in this work.
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