ORIGINAL RESEARCH

Proteomics and Metabolomics Analysis Reveals the Toxicity of ZnO Quantum Dots on Human SMMC-7721 Cells

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Purpose: ZnO quantum dots (QDs) are composed of less toxic metals than other QDs but have the same interesting photochemical properties. Thus, they have received considerable attention recently. Nevertheless, their toxicity cannot be ignored.

Methods: In this study, we incubated ZnO ODs with human SMMC-7721 cells for 24 h to assess their nanotoxicity through proteomics (Fold change >1.5 and p-value <0.05) and metabolomics (Fold change \ge 1.5; VIP \ge 1; p-value < 0.05) analyses.

Results: Both of 174 and 219 significantly changed metabolites were identified in human SMMC-7721 cells treated with 20 and 50 µg/mL ZnO QDs, respectively. ZnO QDs significantly modified metabolic pathways, including purine metabolism, ferroptosis, morphine addiction, alcoholism, cGMP-PKG signaling, and Cushing syndrome. Moreover, we identified 105 and 8 differentially expressed proteins in cells treated with 20 and 50 µg/mL ZnO QDs, and the pathways of alcoholism and Cushing syndrome were enriched.

Conclusion: ZnO QDs did not affect cell viability in a CCK8 assay, but disturbed the level of intracellular metabolites and proteins at 20 µg/mL. The KEGG analyses of the metabolomics and proteomics data both enriched the alcoholism and Cushing syndrome pathways. These results provide an experimental basis for future research on the safe use of nanomaterials.

Keywords: ZnO quantum dots, cytotoxicity, proteomics, metabolomics

Introduction

Semiconductor quantum dots (QDs) have unique optical properties, including narrow and tunable fluorescence emission spectra, wide excitation wavelength ranges, and good resistance to photochemical degradation and photobleaching.^{1,2} QDs have potential applications in many fields, such as bioimaging, biolabeling, nanomedicine, and optoelectronics.^{1,3,4} Because of this extensive application range, intentional and unintentional environmental contamination is inevitable In this context, ZnO ODs have received considerable attention recently. Indeed, they are composed of less toxic metals than other QDs while keeping interesting photochemical properties. For example, EGCG-modified ZnO QDs potential to be a safe and effective treatment material for diabetic wound.⁵ ZnO QDs with sunlight-driven antibacterial activity can be used for communicable disease protective wearables.⁶ Sarkar et al reported that luminescent defect-engineered ZnO QDs have potential as a new, safe, and economical multifunctional active ingredient for skin UV protection.⁷ ZnO QDs was also used to fabricated gas sensor for NO₂ and methanol detection.^{8,9} Moreover, ZnO QDs was reported significantly promoted tomato (Solanum lycopersicum) and pumpkin (Cucurbita moschata Duch.) growth in comparison with the equivalent concentrations of other sizes of ZnO particles.^{10,11}

Current toxicity studies on ZnO QDs mainly explored aspects such as cytotoxicity, antibacterial activity, reactive oxygen species (ROS) production, oxidative stress, and apoptosis.^{4,12-16} Traditional, single end-point approaches are difficult to achieve toxicity assessment of the growing number of new nanomaterials.¹⁷ However, omics techniques including transcriptomics, proteomics, and metabolomics are promising high throughput methods used in predictive toxicology, which could also detect unsuspected subtle changes before conventional methods.^{18,19} The metabolome is defined as the quantitative collection of low-molecular-weight molecules (metabolites) required for the growth and proper function of a cell.²⁰ Untargeted metabolomic have been an unbiased tool for revealing unforeseen biological effects on cellular or animal models.^{18,21} The effect of ZnO QDs on metabolome has not been reported, except little research of other sized ZnO particles. Meanwhile, proteomics technologies can identify thousands of de-regulated dynamic proteins and their interactions in a cell or organism under different environmental conditions.^{22,23} Furthermore, the proteome and metabolome are directly interconnected, as protein levels influence the metabolic profile, and metabolite affect protein expression.¹⁹ Proteome data affected by ZnO QDs is lacking, and more studies are necessary in these directions.

The liver is an important organ of metabolic clearance and a major site of ZnO QDs accumulation.^{24,25} To investigate the toxicity of ZnO QDs, we selected the human SMMC-7721 cell line, which is commonly used in in vitro models to elucidate cytotoxicity mechanisms on hepatocytes.²⁶ We first utilized a multi-omics (proteomics and metabolomics) approach to gain more information and better understand the nanotoxicity of ZnO QDs. Moreover, we analyzed the differentially expressed proteins and metabolites to identify the metabolic pathways affected by ZnO QDs and elucidate the toxicity mechanisms.

Materials and Methods

Materials

We synthesized and characterized ZnO QDs as described in previous reports.^{16,27} We acquired fetal bovine serum (FBS) from Hyclone (Logan, UT, USA), RPMI-1640, trypsin, penicillin-streptomycin and phosphate-buffered saline (PBS) from Corning (New York, NY, USA), dimethyl sulfoxide from Sigma (St Louis, MO, USA), and a Cell Counting Kit 8 (CCK-8) from Beyotime (Shanghai, China). Other chemicals and reagents used were of analytical grade.

Cell Culture

The human SMMC-7721 cells were purchased from the Cell Bank of Chinese Academy of Sciences (Shanghai, China). We cultured the cells in RPMI 1640 containing 10% FBS and 1% penicillin-streptomycin. We maintained the cells at 37°C in a humidity- and CO₂-controlled incubator (Thermo Forma, OH, USA). We performed all the cell experiments under a clean atmosphere. The experimental procedures complied with the "Proteomics and metabolomics analysis of the toxicity of ZnO quantum dots on human SMMC-7721 cells" guidelines approved by the Committee of Medical Ethics and Welfare for Experimental Animals of Henan University School of Medicine (no. HUSOM2021-323).

Cell Viability Assay

We assessed cell viability through a CCK-8 assay. Briefly, we seeded human SMMC-7721 cells into 96-well plates $(5.0 \times 10^3 \text{ cells/well})$. We then added ZnO QDs in RPMI 1640 (final concentrations of 0, 10, 20, 30, 40, 50, 60, 70, 80, and 90 µg/mL). After 24 h of incubation, we washed the cells with PBS to remove the excess of QDs. Then, we added 200 µL of fresh medium and 20 µL of CCK-8 reagent per well and incubated the cells at 37°C for 2 h. Finally, we measured the absorbance at 450 nm using a microplate reader. For all the assays, we performed three independent experiments, each in triplicate.

Metabolomics Analysis

Sample Preparation

We incubated human SMMC-7721 cells with 0, 20, or 50 μ g/mL ZnO QDs for 24 h, then collected them. We weighed 25 mg of each sample, then added 800 μ L of a cold methanol, acetonitrile and water solution (2:2:1, v:v:v). We also added the internal standard at this point. Next, we homogenized the mixture and performed ultrasonic extraction. We then centrifugated the mixture, dried it under a vacuum, and re-suspended it before carrying out the ultra-performance liquid

chromatography-tandem mass spectrometry (UPLC-MS/MS) analysis. We prepared quality control (QC) samples by mixing aliquots of all the samples and injected them every 10 samples throughout the analytical run.

UPLC-MS/MS Analysis

We analyzed the samples using a 2D UPLC system (Waters, USA) coupled with a Q Exactive high-resolution mass spectrometer (Thermo Fisher Scientific, USA). We performed the chromatographic separations on a Waters BEH C18 column (2.1×100 mm, 1.7μ m) at 45°C, with a flow rate of 0.35 mL/min and an injection volume of 5 μ L. For positive-mode MS, the mobile phase consisted of 0.1% formic acid (solution A) and 100% methanol with 0.1% formic acid (solution B); for negative-mode MS, it consisted of 10 mM carbamate (solution A) and 95% methanol with 10 mM carbamate (solution B).

The optimized elution conditions were: 0-1 min, 2% B solution; 1-9 min, 2–98% B solution; 9-12 min, 98% B solution; 12-12.1 min, 98–2% B solution; and 12.1-15 min, 2% B solution. We obtained primary mass spectra by scans from 70 to 1050 m/z at a resolution of 70 K, an automatic gain control (AGC) of 3×10^6 , and a maximum injection time of 100 ms. We selected the top 3 based on the parent ion strength, with a resolution of 17.5 K, AGC of 1×10^5 , maximum injection time of 50 ms, and stepped normalized collisional energy of 20, 40 and 60 eV for secondary information. The experimental conditions for the electrospray ionization source were: sheath gas flow rate, 40 arbitrary units; aux gas flow rate, 10 arbitrary units; spray voltage (|KV|), 3.80 for the positive mode and 3.20 for the negative mode; gas temperature, 350° C.

Data Processing and Statistical Analysis

We processed raw UPLC-MS/MS data (peak extraction, peak alignment, and compound identification) using Compound Discoverer 3.1 (Thermo Fisher Scientific, USA). We used the MetaX R software package (BGI, China) for data preprocessing, statistical analysis, metabolite classification and functional annotation. We also conducted unsupervised principal component analysis (PCA) and supervised partial least squares-discriminant analysis (PLS-DA). We selected significantly modified metabolites based on fold change ≥ 1.5 ; VIP values of the first two principal components of the PLS-DA model ≥ 1 ; and *p*-value of Student's *t*-test < 0.05. We achieved the molecular identification of metabolites by automatically matching the purified mass spectra with the BGI Library, mzCloud and ChemSpider database. Finally, we performed an enrichment analysis of the disturbed metabolites using the KEGG (Kyoto Encyclopedia of Genes and Genomes) database and considered that p < 0.05 indicated significantly enriched targets.

Proteomics Analysis

Protein Sample Preparation

We treated human SMMC-7721 cells with 0, 20 or 50 μ g/mL ZnO QDs for 24 h. Next, we added a 1 × Cocktail with ethylenediamine tetraacetic acid and without sodium dodecyl sulfate. We placed the sample on ice for 5 min and added dithiothreitol to a final concentration of 10 mM. We then sonicated the suspension to lyse the cells, centrifuged at 4°C and 25,000 g for 15 min, collected the supernatant, and incubated it at 56°C for 60 min. We then added iodoacetamide to a final concentration of 55 mM and incubated the samples for 45 min in the dark. After centrifugation, we quantified the proteins in the supernatant using the Bradford method. Finally, we analyzed the proteins by sodium dodecyl sulfate-polyacrylamide gel electrophoresis.

Proteolysis and IBT Labeling

We diluted the protein solution (100 μ g) with 0.5 M tetraethylammonium bromide, added 5 μ g of trypsin, and incubated at 37°C for 4 h. After salt removal, we dried the peptides under a vacuum.

We dissolved the peptides in 200 mM tetraethylammonium bromide to obtain a final concentration of 40 μ g/ μ L. Next, we placed 100 μ g of peptides in tubes and added 2 mg of IBT reagent (BGI, Shenzhen, China) dissolved in 80 μ L of isopropanol in each tube. We rapidly mixed, shook, and centrifuged the tubes, then checked that the pH was between 7.0 and 8.0. We then incubated the mixture was at room temperature for 2 h to obtain sufficient labeling. We labeled control samples as "C", 20 μ g/mL ZnO QDs-treated samples as Z20, and 50 μ g/mL ZnO QDs-treated samples as Z50. The

isobaric tagging of samples was performed as follows: C1:118C; C2:119; C3:115N; Z20-1:115C; Z20-2: 116N; Z20-3: 116C; Z50-1: 117N; Z50-2: 117C; and Z50-3: 118N.

Peptide Fractionation and UHPLC-MS/MS

We dissolved 20 μ g samples of dried peptides in mobile phase A (5% acetonitrile, pH = 9.8) and fractionated them on a Shimadzu LC-20AD system, using a 5 μ m × 20 cm × 180 μ m with a 5%–35% gradient of buffer B (95% acetonitrile, pH = 9.8). According to the chromatographic elution peaks at 214 nm, we obtained 20 components, which we freeze-dried.

We dissolved the dried peptide samples in mobile phase A (2% acetonitrile, 0.1% formic acid), then centrifuged the solutions at 20,000 g for 10 min. We separated the obtained supernatant on a Thermo UltiMate 3000 UHPLC system. The sample was enriched in trap column and desalted, then placed on a self-packed C18 column (75 μ m × 3 μ m × 25 cm). The liquid-phase gradient was: 0–5 min, 5% mobile phase B (98% acetonitrile, 0.1% formic acid); 5–45 min, 5–25% B; 45–50 min, 25–35% B; 50–52 min, 35%–80% B; 52–54 min, 80% B; 54–60 min, 5% B; flow rate, 300 nL/min.

The UHPLC system was connected to the mass spectrometer; peptides were ionized by a nano-electrospray ionization source and passed to a Q-Exactive HF X tandem mass spectrometer (Thermo Fisher Scientific, San Jose, CA, USA) for data-dependent acquisition. The MS parameters were: ion source voltage, 1.9 kV; MS1 scanning range, 350–1500 m/z; the resolution for MS1, 60,000; MS2 starting m/z, 100; resolution for MS2, 30,000; AGC for MS1, 3×10^6 ; AGC for MS2, 1×10^5 ; dynamic exclusion time, 30 s. The ion screening conditions for MS2 fragmentation were: charge 2+ to 6+, and the top 20 parent ions with a peak intensity exceeding 20,000. The ion fragmentation mode was higher-energy collisional dissociation, and the fragment ions were detected in Orbitrap.

Proteomics Data Analysis and Bioinformatics

We converted raw MS data to the MGF format, and identified proteins using the Mascot search engine (version 2.3.02). We quantified the proteins using IQuant software (BGI, Shenzhen, China).²⁸ We set the false discovery rate to $\leq 1\%$ for both protein and peptide identification. We set fold change > 1.5 and *p*-value < 0.05 as significance thresholds for differentially expressed proteins (DEPs). Finally, we functionally annotated DEPs by KEGG pathway enrichment analysis (<u>http://www.genome.jp/kegg/pathway.html</u>) and considered that *p*-value < 0.05 indicated significant enrichment.

Statistical Analysis

The CCK-8 assay results are presented as mean \pm SEM. We compared the means of multiple groups by one-way analysis of variance (ANOVA) followed by Dunnett's test using SPSS version 16.0. We compared pairs of groups by using an unpaired *t*-test. We considered that p < 0.05 indicated statistical significance.

Results and Discussion

As shown in <u>Figure S1A</u>, transmission electron microscopy (JEM2100Plus, JEOL, Japan) revealed that ZnO QDs had an average diameter of 7.98 ± 0.31 nm. They also had strong fluorescence centered at 566 nm and an excitation maximum located at 368 nm (<u>Figure S1B</u>), as recorded on a Fluorolog-3 spectrofluorometer (HORIBA Scientific, USA).

Cytotoxicity of ZnO QDs in Human SMMC-7721 Cells

We assessed the viability of human SMMC-7721 cells incubated with ZnO QDs for 24 h using a CCK-8 assay (Figure 1). We observed significant dose-dependent cytotoxicity starting at 30 μ g/mL. This result is consistent with our previous experiment, which showed that 25 μ g/mL ZnO QDs did not affect the cell viability of Hela and HEK-293T, while 50 μ g/mL QDs killed approximately 50% of the cells.¹⁶ Roshini et al reported that 10 μ g/mL ZnO QDs killed approximately 40% of MCF-7 and MDA-MB-231 breast cancer cells.¹² Based on the cytotoxicity of ZnO QDs on SMMC-7721 cells, we selected the 20 (no toxicity dose) and 50 (dose of about 50% inhibition of cell activity) μ g/mL concentrations for the following metabolomics and proteomic experiments.



Figure 1 Cytotoxicity of ZnO QDs against SMMC-7721 cells after 24 h exposure. Cell viability was determined by CCK-8 assay and calculated relative to negative controls. All data are presented as mean \pm SEM (n = 3). *p < 0.05, and ***p < 0.001 versus control according to ANOVA followed by Dunnett's test.

Effects of ZnO QDs Exposure on the Metabolome of Human SMMC-7721 Cells

We analyzed the metabolites of human SMMC-7721 cells incubated with 0, 20, or 50 μ g/mL ZnO QDs for 24 h using UPLC-MS/MS. Figure S2 shows representative base peak chromatograms in positive and negative mode for untreated SMMC-7721 cells. The overlay of the base peak of all QC samples (Figure S3) suggested that the stability of the UPLC-MS/MS system was acceptable, while QC samples together in PCA score plots indicated the sufficiently reproducibility of the method (Figure S4). We identified a total of 1258 metabolites in positive model (538 with identification information) and 1080 in negative model (428 with identification information) by automatically matching the mass spectra of purified compounds with the BGI Library, mzCloud and ChemSpider databases. Furthermore, we compared the metabolites of control and ZnO QD-treated cells (20 and 50 μ g/mL) using unsupervised PCA and supervised PLS-DA. We observed marked differences in the PCA score plots both in positive and negative modes (Figure 2A and B). Ellipses represents the 95% confidence intervals, and we used all of the samples in the following analysis to obtain maximum information.



Figure 2 Discrimination plots of control and ZnO QDs-treated cells from the PCA of UPLC-MS/MS data: (A) positive mode, (B) negative mode.



Figure 3 Metabolic profiles of the different ZnO QDs doses and controls at 24 h. (A–D) score plots of the PCA model, (E–H) score plots of the PLS-DA model, (I–L) plot of the permutation test (200 times) of the PLS-DA model.

Table I Overlap of Significantly Changed Metabolites Between Different Dosage of ZnO QDs and Controls

Name	Low Dose vs Control		High Dose vs Control			
	Ratio	p value	VIP	Ratio	p value	VIP
(±)-pantetheine	4.0394	0.0106	1.3888	74.1569	0.0001	2.5884
(2-hydroxy-2-oxido-1,3,2-dioxaphospholan-4-yl)methyl (9z)-9-octadecenoate	2.5656	0.0176	1.2296	4.9084	0.0018	1.197
(2-hydroxy-2-oxido-1,3,2-dioxaphospholan-4-yl)methyl palmitate	2.9252	0.0337	1.1397	7.2363	0.0005	1.3161
(2-hydroxy-2-oxido-1,3,2-dioxaphospholan-4-yl)methyl stearate	2.6216	0.0141	1.1177	6.7481	0.0006	1.3001
(2r)-1-{[(2-aminoethoxy)(hydroxy)phosphory]]oxy}-3-hydroxy-2-propanyl (11z)-	2.7337	0.0039	1.3256	7.8905	0.0001	1.8935
I l-icosenoate						
(2r)-1-{[(2-aminoethoxy)(hydroxy)phosphoryl]oxy}-3-hydroxy-2-propanyl (13z)-	2.6115	0.0067	1.3248	7.183	0.0001	1.8556
I3-docosenoate						
(2r)-2-acetoxy-3-[(9z)-9-octadecen-1-yloxy]propyl 2-(trimethylammonio)ethyl	2.4033	0.0171	1.221	5.0328	0.0398	1.2604
phosphate						
(3s,5z,7e)-26,26,26,27,27,27-hexafluoro-9,10-secocholesta-5,7,10-triene-3,25-	5.5868	0.0013	1.706	8.6811	0.0003	1.3916
diol						
I-(Iz-hexadecenyl)-sn-glycero-3-phosphocholine	3.1404	0.0036	1.3458	15.5062	0.0001	2.2395
I-(Iz-octadecenyl)-2-hexadecanoyl-sn-glycero-3-phosphoethanolamine	2.4605	0.0403	1.5421	2.3114	0.0425	1.0755
I-(9z-octadecenoyl)-sn-glycero-3-phospho-(1'-sn-glycerol)	6.536	0.0003	1.7791	9.6255	0	1.407
I.4-d-xylobiose	4.2433	0.0006	1.5741	15.2279	0.0002	2.1008
16-heptadecyne-1.2.4-triol	2.5507	0.0085	1.2532	3.6556	0.0016	1.0263
I-arachidonoyl-sn-glycero-3-phosphocholine	2.6867	0.0222	1.0055	10.2014	0.0012	1.8414
I-hentadecanovI-sn-glycero-3-phosphocholine	2.6347	0.0461	1,2939	4.1147	0.0472	1.1691
	5.5149	0.0007	1.4387	60.8509	0.0013	2.4636
I-methylinosine	4 3693	0.0009	1 5063	18 9429	0.0005	2 1553
LoctadecanovI-2-(7z 10z 13z 16z)-docosatetraenovI-sn-glycero	2 525	0.0361	16117	2 9513	0.0238	1 29
-3-phosphoethanolamine	2.025	0.0501	1.0117	2.7515	0.0250	1.27
I-o-bexadecyl-lyso-sn-glycero-3-phosphocholine	2 3629	0.0082	1 2088	6 592	0.0021	1 6224
Loleovi-sn-giveero-3-phospho-d-myo-inositol	2.5027	0.0278	1.1969	4 6675	0.0026	1.1603
Loleovi-sn-giveero-3-phosphoethanolamine	31168	0.0148	1 4456	5 3647	0.0001	1.1005
Lo-oleovi-sn-givcero-3-phosphoserine	3 3124	0.0011	1 4702	5.5012	0.0003	1.2327
I-stearoyl-sn-glycero-3-phosphoethanolamine	2 6408	0.0099	1.1702	62159	0	1.2105
Leteradecanovi-2-[(5z 8z 1 / z 1 4z)-eicosatetraenovi]-sn-givcero	2.6100	0.0272	13128	6 3644	0.0095	1.2720
-3-phosphocholine		0.0272	1.5120	0.5011	0.0075	1.1371
2-/2 6-dihydroxy-4-[6-hydroxy-7-(3-methyl-2-huten-1-yl)-1-henzofuran-2-yl]		0.0459	1 0973	69 0412	0.0001	1 9905
phenyl}-6-(2 4-dihydroxyphenyl)-5-bydroxy-4-methyl-3-cyclohexene-		0.0107	1.0775	07.0112	0.0001	1.7705
L-carboxylic acid						
2-2minoethyl (2r)-3-[(1z)-1-hevadecen-1-vloxy]-2-hydroxypropyl hydrogen	2 7578	0	12137	5 8933	0	1 2602
	2.7570	Ŭ	1.2157	5.0755	Ŭ	1.2002
2-linoleovl-sn-glycero-3-phosphoethanolamine	28176	0.001	1 2804	5 486	0.0002	1 2505
2-molecyl-singlycelo-3-phosphoeenanolamine	2.0170	0.0231	1.2001	3 2029	0.0002	1.2005
3-({[(2s)-2 3-dihydroxypropoxy](hydroxy)phosphory]}oxy)-2-hydroxypropy	5 2378	0.0001	1.3958	29 161	0	1.7831
	5.2570	0.0001	1.5750	27.101	Ū	1.7051
3-/[/2 3-dihydroxypropoxy)(hydroxy)phosphoryl]oxy}-2-hydroxypropyl stearate	3 3566	0.0002	1 0842	21.646	0	1 703
	4 8535	0.0002	2 0926	9 2487	0.0037	1.705
4 (alpha d ducosaminyl) I d myo inositol	4 4 9 5 9	0.0074	1 3 2 3 3	22 0401	0.0037	1.295
- (alpha-u-glucosalilling)-iu-inositol		0.0003	1.3233	4 1 4 9 4	0.0013	1.075
l vl 4 (1 azonanyl)butanosto	2.0105	0.0077	1.5057	4.1070	0.0015	1.5752
8 hydroxy dooxyguanosing	6 7107	0.0032	1 9924	22 9453	0.0011	2 1927
Abaditaral	2 3935	0.0032	1.7020	3 6734	0.0042	1 0403
Adenine	2.3733	0.0019	1.1317	64 4745	0.0072	1.0703
Adenosine	8 6697	0.0012		76 299	0	1.00/7
	23121	0.0000	1.0204	4 2222		1.7372
	2.3121	0.002	1.0300	0 4220	0 0001	1.112/
	3.0312	0.0037	1.2013	0.7220	0.0001	1.2000

Name	Low Dose vs Control		High Dose vs Control			
	Ratio	p value	VIP	Ratio	p value	VIP
Anhydrotetracycline	0.3822	0.0275	1.096	0.129	0.0015	2.074
Asp-trp	0.1814	0.0343	1.3815	0.0155	0.0003	1.7167
Buprenorphine	2.2176	0.0212	1.1428	5.0672	0.0012	1.5218
Cholest-5-en-3-ol	2.4496	0.0224	1.4487	2.62	0.0165	1.1653
Cilastatin	0.3879	0.0127	1.0941	0.1044	0.0005	2.1675
Dicoumarol	3.1582	0.0028	1.388	5.8572	0.0004	1.2756
Diflucortolone	0.1109	0.0233	1.6751	0.0327	0.0019	1.4772
Docosatetraenoylethanolamide	2.2087	0.0477	1.6544	2.6528	0.0357	1.2972
Epsilon-(gamma-glutamyl)-lysine	0.3374	0.0356	1.5185	0.0911	0.0025	1.2232
Eptapirone	0.0261	0	2.5248	0.0096	0	1.9218
Ethyl docosahexaenoate	2.3676	0.0155	1.2751	29.4454	0.0005	2.1416
Fructoselysine	0.4289	0.0085	1.4639	0.1309	0	1.3256
Glycerophospho-n-palmitoyl ethanolamine	2.2521	0.0153	1.1756	4.5658	0.0013	1.1933
Guanine	3.3957	0.0116	1.3202	12.2321	0.0044	1.8658
Guanosine	6.5347	0.0004	1.4879	23.7246	0	1.637
Ho-dpeg8-oh	0.2014	0.0024	1.3861	0.0433	0	1.6574
Inosine	5.003	0.0089	1.9071	11.2705	0.0059	1.862
L-alpha-aspartyl-l-phenylalanine	0.2282	0.0194	1.7209	0.0544	0.0001	1.4543
Leucylasparagine	4.419	0.0073	2.078	0.0167	0.0007	1.6411
Leucylproline	0.014	0.0124	1.9507	0.0022	0.0014	1.9162
Leu-gly-pro	0.3133	0.0028	1.2626	0.1637	0.0002	1.2025
Leu-leu	0.1242	0.0353	1.5045	0.0815	0.0154	1.1168
Leu-val	0.0782	0.043	1.5628	0.1873	0.0025	1.0058
Lysopc	2.5419	0.0051	1.0913	10.4208	0.0101	1.7301
Lysopc a c28:1	2.9685	0.0172	1.5536	3.8575	0.0073	1.3844
Lysophosphatidylcholine 14:1(9z)/0:0		0.0018	1.6601	15.5481	0.0001	1.6107
Lysophosphatidylinositol	5.7105	0.0044	1.4258	19.4967	0	1.6291
Lys-pro	0.0903	0.013	2.326	0.0801	0.008	1.7408
Marimastat	0.0897	0.0172	1.4972	0.0073	0.0002	1.9425
Menatetrenone	6.1628	0.0003	2.1968	7.1793	8000.0	1.8274
Methohexital	0.028	0.0119	1.8357	0.0068	0.0021	1.6979
Mfcd00037235	0.1898	0.0027	1.3533	0.0401	0	1.6755
Mfcd00059633	2.3441	0.0039	1.1576	3.5991	0.0003	1.0341
Militarinone a	2.0566	0.0195	1.0908	3.9369	0.0023	1.4119
Miltefosine	2.4969	0.0071	1.3794	3.4773	0.0296	1.1826
Mono(2-ethylhexyl) phthalate (mehp)	2.815	0.0048	1.2443	3.9364	0.0001	1.047
Myxochelin a	2.638	0.0204	1.0137	10.7227	0.0001	1.4674
N,n-dimethyladenosine	5.989	0.0147	1.8854	28.0035	0.0173	2.0285
N2-dimethylguanosine	3.8202	0.0077	1.7025	9.7792	0.0044	1.8046
N-tridecanoylglycine	7.1757	0.009	1.169	102.7646	0	1.9833
O-[{(2r)-3-[(4z,7z,10z,13z,16z,19z)-4,7,10.13.16.19-docosahexaenovloxv]		0.0186	1.2507	11.4219	0.0001	2.0721
-2-[(9z, 2z, 5z)-9, 2, 5-octadecatrienovloxy]propoxy}(hvdroxy)phosphoryl]-						
I-serine						
Pc	2.2032	0.0473	1.2059	3.1576	0.0337	1.3232
Porphyra-334	2.5718	0.0041	1.1974	3.7278	0.0001	1.0442
Progesterone	2.2609	0.0101	1.3353	2.7313	0.0303	1.0556
Promegestone	1.8318	0.0334	1.1505	2.4483	0.0084	1.1819
Riboprine	2.9004	0.0145	1.3823	10.6261	0.0022	1.897
Ro 20–1724	0.3362	0.0335	1.0439	0.1636	0.0021	1.0928

Name	Low Dose vs Control		High Dose vs Control			
	Ratio	p value	VIP	Ratio	p value	VIP
S-adenosylhomocysteine	13.1582	0	2.2135	23.8809	0	1.6587
Sm(d18:0/14:0)	2.4895	0.0373	1.4481	2.4578	0.0411	1.0274
St2975000	4.9473	0.0011	1.7002	17.2064	0.0007	2.1077
Thymidine	9.2648	0.0022	2.2155	11.6621	0.0063	1.2082
Thymine	5.2527	0.0004	1.986	11.146	0	2.027
Transfluthrin	4.0575	0.0026	1.0373	21.1084	0	1.5297
Vitamin e nicotinate	2.1556	0.0308	1.2546	2.7571	0.0128	1.1491
Xanthosine	4.9458	0.0022	1.3315	13.2779	0.001	1.2717

We performed a multivariate analysis comparing the metabolites of control and ZnO QD-treated cells (20 or 50 μ g/mL). The PCA score plots of ZnO QD-treated cells (at both concentrations) and those of control cells were notably different in positive and negative mode (Figure 3A-D). PLS-DA, a supervised statistical method, can better reflect the differences between experimental and control samples. Indeed, we observed a better separation in the PLS-DA model, with a seven-fold cross-validation (Figure 3E-H), suggesting a significant metabolic difference between ZnO QD-treated and control cells. Next, we performed response permutation testing 200 times to confirm that the PLS-DA model was not random and overfitting (Figure 3I-L). The criteria for identifying significantly modified metabolites were: fold change ≥ 1.5 ; VIP values of the first two principal components of the PLS-DA model ≥ 1 ; and *p*-value of Student's *t*-test < 0.05. Thus, we identified 174 significantly changed metabolites in cells treated with 20 μ g/mL ZnO QDs (96 in positive mode and 102 in negative mode), and 219 in cells treated with 50 μ g/mL ZnO QDs (98 in positive mode and 159 in negative mode). A total of 95 metabolites were changed in both treated groups (Table 1). According to the CCK8 assay results, 20 μ g/mL ZnO QDs did not induce cytotoxicity in SMMC-7721 cells but affected intracellular metabolite levels.

We analyzed the differential metabolites in positive and negative modes to find ZnO QD-modified metabolic pathways. In positive mode, cells treated with 20 and 50 µg/mL ZnO QDs had almost the same pathways significantly enriched (Figures 4 and S5A), namely purine metabolism, ferroptosis, morphine addiction, alcoholism, cGMP-PKG signaling pathway, and Cushing syndrome. Purines are important components of DNA replication and RNA synthesis, and play a key role in neurotransmission and neuromodulation.^{29,30} Zinc oxide nanoparticles can reduce the level of purine metabolites in Saccharomyces cerevisiae.³¹ Ferroptosis is a newly identified programmed cell death driven by iron-dependent lipid peroxidation.³² Zhang et al demonstrated that "iron free" zinc oxide nanoparticles triggered ferroptosis by increasing ROS production and lipid peroxidation in vitro.³³ Wu et al also reported that graphene QDs caused ferroptosis via mitochondrial oxidative stress in microglia.³⁴ Adenosine is a major component of adenine nucleotides and ribonucleic acids, as well as a signal molecule mediating signal transduction.³⁵ ZnO ODs disturbed adenosine in many signaling pathways, including morphine addiction, alcoholism, and cGMP-PKG signaling. In negative mode, cells treated with 20 and 50 µg/mL displayed different enriched pathways (Figure S5B and C). The 20 µg/mL ZnO QDs treatment significantly enriched pathways such as biosynthesis of unsaturated fatty acids, linoleic acid metabolism, purine metabolism, fatty acid biosynthesis, regulation of lipolysis in adipocytes, while the 50 µg/mL ZnO QDs treatment affected protein digestion and absorption, biosynthesis and metabolism of amino acids, aminoacyl-tRNA biosynthesis, mineral absorption, purine metabolism, ABC transporters, glyoxylate and dicarboxylate metabolism, and pyrimidine metabolism.



Figure 4 Scatterplot of significantly enriched KEGG pathways based on the differential metabolites induced by 20 µg/mL ZnO QDs (data obtained in positive mode UPLC-MS/MS). The size of the dots represents the number of metabolites.

Effects of ZnO QDs Exposure on the Proteome of Human SMMC-7721 Cells

Overall, we identified 6469 proteins under the 1% false discovery rate filter. The DEP selection criteria were: fold change > 1.5 and *p*-value < 0.05. Compared with the control cells, cells treated with 20 μ g/mL ZnO QDs had 105 DEPs, including 64 upregulated and 41 downregulated proteins (Table 2). Next, we carried out a KEGG pathways

Accession	Protein Name	Abbreviation	Regulation			
Low Dose vs Control						
Q9Y4G2	Pleckstrin homology domain-containing family M member I	РКНМІ	Up			
P24386	Rab proteins geranylgeranyltransferase component A I	RAEI	Up			
075570	Peptide chain release factor I, mitochondrial	RFIM	Up			
O15049	NEDD4-binding protein 3	N4BP3	Up			
Q96RU7	Tribbles homolog 3	TRIB3	Up			
Q5T4F4	Protrudin	ZFY27	Up			
Q8N0Z6	Tetratricopeptide repeat protein 5	TTC5	Up			

 Table 2 Significantly Differentially Expressed Proteins Induced by ZnO QDs

Table 2 (Continued).

Accession	Protein Name	Abbreviation	Regulation
No.			
P61088	Ubiquitin_conjugating enzyme F2 N	LIBE2N	Ць
092954	Proteoglycan 4	PRG4	Up
P13929	Beta-enolase	FNOB	Up
014522	Recentor-type tyrosine-protein phosphatase T	PTPRT	Un
O9NPR9	Protein GPR 108	GP108	Un
Q16514	Transcription initiation factor TEIID subunit 12	TAFI2	Un
08N876	Discoidin CLIB and LCCL domain-containing protein L		Un
P36894	Bone morphogenetic protein receptor type-14	BMRIA	Up
		NEBI	Up
	Protein FAM71F1	F7IFI	Up
Q16650	They brain protein I	TBRI	Up
P04264		K2C1	Up
	F3 ubiguitin-protein ligase LIBR2	LIBR2	Un
	Zinc finger HIT domain-containing protein 2		Un
Q96AB3	Isochorismatase domain-containing protein 2		Up
Q05BO5	MBT domain-containing protein 1	MBTDI	Up
P49593	Protein phosphatase IF	PPMIE	Up
09H117	Protein Wrt-Sh	WNT5B	Up
014061	Cytochrome c oxidase concer chaperone		Up
	WD and tetratricopentide repeats protein L	WDTCI	Up
014795	Protain unc-13 homolog B		Up
OPBVG8		KIEC3	Up
P04259		K2C6B	Up
P55789	FAD_linked sulfbydryl oxidase ALR	AL R	Up
P07602	Prosposin	SAP	Up
P02790	Hemopexin		Up
P62487	DNA-directed RNA polymerose II subunit RPB7	RPB7	Up
013907	Isopentenyl-dinboshate Delta-isomerase I		Up
P46020	Phosphorylase h kinase regulatory subunit alpha skeletal muscle isoform	KPRI	Up
099757	Thioradoxin mitochondrial	THIOM	Up
09H3D4		P63	Up
P21817	Ryanodine receptor I	RYRI	Up
P13473	l vsosome-associated membrane glycoprotein 2		Up
015145	Actin-related protein 2/3 complex subunit 3	ARPC3	Up
086Y37	CDK2-associated and cullin domain-containing protein 1		Up
Q00137	Conserved aligometric Galgi complex subunit 3	COG3	Up
Q96FK7	Constitutive coactivator of perovisome proliferator-activated receptor gamma	EL20B	Up
096674	OTU domain-containing protein 5		Up
Q9UB79	DNA repair protein REVI	REVI	Un
P14174	Macrophage migration inhibitory factor	MIF	Un
P35527	Keratin type I cytoskeletal 9	кіся	Un
015055	Period circadian protein homolog 2	PER2	Up
095835	Serine/threonine-protein kinase ATS1		Up
015243	lentin recentor gene-related protein	OBRG	Up
015245 096CP7			Up
013404	Le containe protein i		Up
P56277	Cy9C motif.containing protein 4	CMC4	Up
065558	Capping protein Arp2/3 and myosin-I linker protein 2		Up
P14209	CD99 antigen	CD99	Up
04VC31	Coiled-coil domain-containing protein 58	CCD58	Ср Цр
211031	Concercon domain-containing protein 50		Ч С

Table 2 (Continued).

Accession	Protein Name	Abbreviation	Regulation
No.			-
	Gynsy retrotransposon integrase-like protein l	GINI	
Q75N03	E3 ubiguitin-protein ligase Hakai		Up
Q751105	Protain PATI homolog I		Up
	Puridina nucleatida disulfida avidaradustasa domain containing protain L		Up
Q877010			Ор
015495	Putative LI2 small puclear ribonucleoprotein auxiliary factor 35 kDa subunit related protein L		Up
	Venetropic and polytropic retrovirus receptor I		Ор
	E2 ubiguitin protein ligase PNEL69		Ор
Q011 VV 3	ES ubiquiun-protein ligase KINF166	RINIGO	Down
Q6V107	Phospholurin acidic cluster sorting protein 1	FACSI	Down
			Down
015504	Nucleoporin NUP42	NUP42	Down
095429	BAG family molecular chaperone regulator 4	BAG4	Down
076039	Cyclin-dependent kinase-like 5	CDKLS	Down
Q92733	Proline-rich protein PRCC	PRCC	Down
Q6ZS81	WD repeat- and FYVE domain-containing protein 4	WDFY4	Down
Q9BTM1	Histone H2A.J	H2AJ	Down
Q9Y2E4	Disco-interacting protein 2 homolog C	DIP2C	Down
P16455	Methylated-DNA–protein-cysteine methyltransferase	MGMT	Down
P16401	Histone H1.5	HI5	Down
A6NDN3	Golgin subfamily A member 6B	GOG6B	Down
Q8WWT9	Solute carrier family 13 member 3	S13A3	Down
Q9NZRI	Tropomodulin-2	TMOD2	Down
Q9UPM8	AP-4 complex subunit epsilon-1	AP4E1	Down
P04908	Histone H2A type I-B/E	H2A1B	Down
Q8N302	Angiogenic factor with G patch and FHA domains I	AGGFI	Down
Q14934	Nuclear factor of activated T-cells, cytoplasmic 4	NFAC4	Down
Q9H0G5	Nuclear speckle splicing regulatory protein I	NSRPI	Down
Q15004	PCNA-associated factor	PAF15	Down
Q9BXW9	Fanconi anemia group D2 protein	FACD2	Down
Q15532	Protein SSXT	SSXT	Down
Q9Y6J0	Calcineurin-binding protein cabin-l	CABIN	Down
Q9BSM1	Polycomb group RING finger protein I	PCGFI	Down
Q8IWW6	Rho GTPase-activating protein 12	RHG12	Down
Q9BT25	HAUS augmin-like complex subunit 8	HAUS8	Down
O6P2P2	Protein arginine N-methyltransferase 9	ANM9	Down
O5VTL8	Pre-mRNA-splicing factor 38B	PR38B	Down
P15056	Serine/threonine-protein kinase B-raf	BRAF	Down
P63173	60S ribosomal protein 138	RI 38	Down
015072	Zinc finger protein QZE	OZE	Down
	Protein FAM186A	5186A	Down
096823	Uncharacterized protein C18orf25	CR025	Down
015555	Microtubule-associated protein RP/ER family member 2	MARE2	Down
Q15555	Histone H2A type 2 C		Down
	Polyhomootis like protoin 3		Down
			Down
			Down
		UBAFZ	Down
	Innibitor of growth protein I		Down
P22/36	Nuclear receptor subfamily 4 group A member 1	NK4AT	Down

Table 2 (Continued).

Accession	Protein Name	Abbreviation	Regulation				
110.							
High Dose vs Con	High Dose vs Control						
Q9H4H8	Protein FAM83D	FA83D	Up				
Q8IXZ2	Zinc finger CCCH domain-containing protein 3	ZC3H3	Up				
P26447	Protein S100-A4	S10A4	Up				
Q9BSE4	Homocysteine-responsive endoplasmic reticulum-resident ubiquitin-like domain member 2	HERP2	Up				
	protein						
P62328	Thymosin beta-4	TYB4	Up				
O94966	Ubiquitin carboxyl-terminal hydrolase 19	UBP19	Down				
076039	Cyclin-dependent kinase-like 5	CDKL5	Down				
A6NE01	Protein FAM186A	F186A	Down				

analysis to determine the function of the DEPs. The significantly enriched pathways included the hippo signaling pathway, signaling pathways regulating pluripotency of stem cells, Toll and Imd signaling pathway, systemic lupus erythematosus, alcoholism, Cushing syndrome, transcriptional misregulation in cancer, and basal cell carcinoma (Figure 5). Interestingly, the metabolomics and proteomics analysis both identified the alcoholism and Cushing syndrome pathways as enriched. Compared with the control cells, cells treated with 50 μ g/mL ZnO QDs only five upregulated and three downregulated proteins (Table 2). According to the KEGG pathway analysis, this dose notably affected the interleukin-17 signaling pathway. Interleukin-17, an inflammatory cytokine, is key to the host-protective capacity, while unrestrained interleukin-17 signaling is related to autoimmune disease, immunopathology, and cancer progression.³⁶

Conclusion

ZnO QDs significantly disturbed the metabolism of human SMMC-7721 cells, and changed the level of 174 and 219 metabolites at 20 and 50 μ g/mL, respectively. The proteomics analysis revealed 105 and 8 DEPs. The KEGG



Figure 5 Scatterplot of significantly enriched KEGG pathways based on differential proteins induced by 20 µg/ mL ZnO QDs. The size of the dots represents the numbers of proteins.

analyses of the metabolomics and proteomics data both identified the alcoholism and Cushing syndrome pathways as enriched. ZnO QDs had no effect on the cell viability in the CCK8 assay but affected the intracellular levels of metabolites and proteins at 20 μ g/mL. These findings will be helpful for future research on ZnO QDs and their applications.

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Disclosure

The authors report no conflicts of interest in this work.

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