Transferrin-functionalized nanographene oxide for delivery of platinum complexes to enhance cancer-cell selectivity and apoptosis-inducing efficacy

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Abstract: Rational design and construction of delivery nanosystems for anticancer metal complexes is a crucial strategy to improve solubility under physiological conditions and permeability and retention behavior in tumor cells. Therefore, in this study, we designed and synthesized a transferrin (Tf)-conjugated nanographene oxide (NGO) nanosystem as a cancer-targeted nanocarrier of Pt complexes (Tf-NGO@Pt). This nanodelivery system exhibited good solubility under physiological conditions. Moreover, Tf-NGO@Pt showed higher anticancer efficacy against MCF human breast cancer cells than the free Pt complex, and effectively inhibited cancer-cell migration and invasion, with involvement of reactive oxygen species overproduction. In addition, nanolization also enhanced the penetration ability and inhibitory effect of the Pt complex toward MCF7 breast cancer-cell spheroids. The enhancement of anticancer efficacy was positively correlated with increased cellular uptake and cellular drug retention. This study provides a new strategy to facilitate the future application of metal complexes in cancer therapy.

Keywords: cancer targeting, nanographene oxide, nanomedicine, transferrin, platinum complex, cell apoptosis

Introduction

Cancer has become the leading threat to human health, and there are currently no effective treatments for most kinds of cancer.1 Chemotherapy is a common treatment for cancer, and Pt complexes have shown potential utility for chemotherapy and photodynamic therapy. Pt-based antineoplastic agents, such as cisplatin, carboplatin, and oxaliplatin, are now in routine clinical practice. These heavy-metal agents possess bioactivity for treatment of various tumors; however, they have limitations, including gastrointestinal symptoms, renal tubular injury, neuromuscular complications, and ototoxicity.2 Cisplatin is a commonly used clinical drug for cancer treatment, but acquired resistance limits its clinical application. As an anticancer agent, platinum has equal or greater antitumor activity and lower toxicity than other metallic cytotoxic compounds to extend the spectrum of activity of metal-based drugs.

In the past 30 years, many Pt complexes have been synthesized and tested for potential antitumor activity. Cisplatin, a typical clinical cancer chemotherapeutic, has been widely prescribed for the treatment of multifarious tumors. Till now, thousands of Pt analogs have been designed and synthesized in order to treat cancer, with higher activity and more compliance. Besides Pt complexes, Ru is also another transition metal of the Pt group that generally has lower toxicity compared to cisplatin, because of specific targeting and accumulation of drugs in cancer tissues. However, clinical application...
of Pt complexes has been hindered by poor solubility under physiological conditions and low permeability retention in tumor cells. Graphene nanoparticles with oxidized form (nanographene oxides [NGOs]) have higher hydrophilicity than NGO, and are sufficiently beneficial to become a drug-delivery system in vivo.\textsuperscript{3–7} NGO has been reported as a metal-complex nanoplatform.\textsuperscript{8–11} Conjugation of cancer-targeted ligands on the surface of a nanosystem is an effective strategy to enhance tumor selectivity by active targeting.\textsuperscript{11–13} This contributes significantly to achieving maximum anticancer effect and significantly reducing side effects.\textsuperscript{14–16}

This study reports on the design of a transferrin (Tf)-conjugated NGO nanosystem, which is a specific targeting molecule for oncotherapy.\textsuperscript{17–19} Tf-NGO@Pt nanoparticles exhibit good water solubility through polyethylenimine (PEI) decoration.\textsuperscript{20} Therefore, they offer a solution to the poor solubility of Pt complexes. Based on this design, we found that Tf-NGO@Pt shows higher anticancer efficacy against MCF human breast cancer cells than the free-Pt complex, and can effectively inhibit cancer-cell migration and invasion by induction of cell apoptosis through upregulation of reactive oxygen species (ROS), important biochemical signaling mediators.\textsuperscript{21–24} In addition, nanolization also enhances the penetration ability and inhibitory effect of the Pt complex toward MCF7 breast cancer-tumor spheroids. The enhancement of anticancer efficacy is positively correlated with the increased amount of cellular uptake and cellular drug retention. This study provides a new strategy to facilitate the future application of metal complexes in cancer therapy.

**Materials and methods**

**Synthesis of Pt complex and preparation of NGO nanosystem**

The Pt complex (Pt [10-methyldipyridophenazine] Cl) was synthesized according to our previous method.\textsuperscript{30} NGOs were prepared according to a slightly modified Hummer protocol with modification.\textsuperscript{25–28} For synthesis of the NGO nanosystem, the Pt-complex solution (1 mg/mL) was dropped in 10 mL NGO solution (1 mg/mL) and ultrasound treated for 15 minutes. The product was allowed to react for 12 hours and filtered with water to remove superfluous Pt complex. PEI solution (2.5 mg/mL) was mixed with the as-prepared solution (NGO-Pt) for 12 hours, and the mixture was dialyzed to remove redundant PEI.

**Tf conjugation and characterization**

For Tf conjugation, 0.2 M 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide hydrochloride (EDC) solution was carefully stirred with Tf (5 mg/mL) and activated the Tf amine group for 2 hours. An added NGO-complex PEI solution then stood at 4°C overnight. Finally, reagents were removed by dialysis for 8 hours. Tf-NGO@Pt was analyzed with atomic-force microscopy (Bioscope Catalyst NanoScope V), transmission electron microscopy (Hitachi H-7650) with 80 kV accelerating voltage, and scanning electron microscopy (EX-250; Horiba). A Zetasizer Nano ZS particle analyzer (Malvern Instruments, Malvern, UK) was used to measured size distribution and ζ-potential of nanoparticles. Chemical construction of the nanosystem was characterized by ultraviolet (UV)-visible spectroscopy (Carry 5000) and Fourier-transform infrared (FTIR) spectroscopy (Equinox 55). A bicinchoninic acid kit (Pierce) was used to measure the concentration of Tf conjugate in the nanosystem. Pt content was determined as previously described by inductively coupled plasma atomic emission spectroscopy (ICP-AES).\textsuperscript{12,29,30}

**Cell lines and cell culture**

Various human cancer-cell lines and normal human cell lines were utilized in this study, including HeLa, CaSki, SiHa, MCF7, Chem5, HepG2 hepatocellular carcinoma, human osteosarcoma MG63, L02 human liver, SV-HUVC1, and A375 cells, obtained from the American Type Culture Collection (Manassas, VA, US). The DMEM media was blended with 50 units/mL of streptomycin (Sigma-Aldrich; St. Louis, MO, USA), 100 units/mL of penicillin (Sigma-Aldrich), and 10% fetal bovine serum. All cell lines were cultured in 5% CO\textsubscript{2} in a fully humidified atmosphere at 37°C.\textsuperscript{31}

**Tf-NGO@Pt cellular uptake**

The concentration of intracellular complexes was determined by ICP-AES. MCF7 and SV-HUVC1 cells were seeded at 3×10\textsuperscript{4} cells per well in six-well plates for 24 hours. The cells were added to Pt complexes and Tf-NGO@Pt (20 μM) for different times. The cells were detached with trypsin EDTA solution (5 minutes, 37°C) and centrifuged to collect the cells. Complex concentration was measured using ICP-AES. The collections were added to mixed acid (concentrated nitric acid:perchloric acid, 3:1) and digested with an infrared (IR) rapid-digestion system (Gerhardt, Königswinter, Germany) at 180°C for 2 hours.

**Cellular retention assay**

Cellular retention of the Tf-NGO@Pt was quantified using a fluorescence microplate reader.\textsuperscript{32} MCF7 and SV-HUVC1 cells were seeded at 2×10\textsuperscript{3} cells per mL (2 mL) and cultured in six-well plates. Then, the medium was replaced by Pt complexes and Tf-NGO@Pt (20 μM) for 8 hours. The medium was then
removed and replaced with medium (2 mL) without phenol. After different periods, medium samples (100 μL) were removed to measure the fluorescence intensity of cell lysis.

**Transferrin competing assay**

Excess Tf receptors (TfRs) were used to bind the TfRs of CMF7 cells to contend with Tf-NGO@Pt. CMF7 cells (100 μL, 7×10^4 cells/mL) were incubated in 96-well plates. After 24 hours, the cells were bound with excess Tf of 0–5 mg/mL and incubated for 1 hour at 37°C, then 20 μM of coumarin-6-loaded Tf-NGO@Pt was added. After 2-hour further incubation, cells were rinsed three times with warm PBS and the fluorescence intensity of cell lysis detected using a fluorescence microplate reader.

**Tf-NGO@Pt intracellular trafficking**

Intracellular localization of Tf-NGO@Pt was characterized by fluorescence microscopy (IX51; Olympus). We incubated MG-63 cells (80 nM) of LysoTracker DND-99 (Sigma-Aldrich) in 2 cm incubator dishes for 2 hours until 70% staining confluence had been reached. Then, 1 mg/mL of DAPI H33258 (Sigma-Aldrich) was added for 30 minutes, cells rinsed with PBS three times and incubated with 20 μM coumarin-6-loaded Tf-NGO@Pt for different periods, and observed by fluorescence microscopy.

**Cellular uptake mechanisms**

We incubated 7,000 MCF7 cells per well in 96-well plates. After 24 hours, the cells were incubated with endocytosis inhibitors for 1 hour, while nystatin (Sigma-Aldrich) was used for 30 minutes. Subsequently, the cells were cultured with 20 μM of coumarin-6-loaded Tf-NGO@Pt for 3 hours. Specific endocytosis-inhibitor concentrations included nystatin (0.45 M sucrose, 50 mM 2-deoxy-D-glucose (Sigma-Aldrich), and 10 mM sodium azide (NaN₃)). The cells were cultivated with fresh medium at 4°C and treated with coumarin-6-loaded Tf-NGO@Pt for a further 3 hours.

**Tf-NGO@Pt drug release**

To examine the drug release of Tf-NGO@Pt, 10 mg Tf-NGO@Pt was suspended in 10 mL PBS solution to imitate the different conditions of microenvironment, blood (pH 7.4), tumor microenvironment (pH 6.8), lysosomes (pH 5.3, pH 5.3 with lysozyme), and cellular environment (cell-lysis solution). Samples were constantly waggled in tubes at 37°C without illumination. At specific times, known buffer volumes were withdrawn from the tubes and replenished with fresh buffer. The collections were separated by centrifugation and the supernatants measured by fluorescence microplate reader.

**MCF7 tumor spheroids**

MCF7 spheroids were cultured in vitro with a liquid-overlay system. Agarose PBS solution (2% w:v) was prepared at 80°C for 30 minutes, then sterilized using autoclave. The agarose solution (50 μL) was covered in the 96-cell plates, then seeded with MCF7 cells (4,000 cells per well). On the first day, the plates were then gently agitated at set intervals and grown for 5 days.

**Tf-NGO@Pt penetration ability into MCF7 spheroids and inhibitory effects**

To compare the penetration ability of complexes (20 μM) and Tf-NGO@Pt (20 μM), the coumarin-6-labeled nanoparticles were added to the MCF7 spheroids and incubated for 12 hours. The tumor spheroids were subsequently scanned at different layers from the spheroid to the middle using confocal laser-scanning fluorescent microscopy (Zeiss LSM700).

Tumor-spheroid size was used to evaluate the cytotoxicity of complexes (20 μM) and Tf-NGO@Pt (20 μM) using an inverted microscope over 5 days. The major (d_max) and minor (d_min) diameters of each spheroid were determined, and spheroid volume was calculated as V = π × d_max × d_min/6. The volume-change ratio for MCF7 spheroids was calculated as R = (V_day1/V_day0) × 100%. Each assay was repeated three times, with sextuplicate determinations for each dose level.

**Cell-migration assay**

MCF7 cells (30×10^4 cells/mL) were seeded into six-well plates. Then, the medium was removed and cultured with 3% fetal bovine-serum medium for a further 6 hours. Subsequently, the cells were wounded by pipette tips, and drugs were added to the plate. Then, 4’,6-diamidino-2-phenylindole (DAPI) was added to label the cells and fluorescent microscopy was used to photograph the migrated cells.

**Cell-invasion assay**

MCF7 cells (6×10^4) containing the drugs were seeded on the upper side of Matrigel-coated culture insert (8 μM; Corning, NY, USA). After 24-hour cultivation, cotton wipes were used to remove the unmigrated cells on the upper side of the transwell membrane, and methanol was used to fix invaded cells for 10 minutes. After cells had been stained with Giemsa solution, they were observed with the Olympus inverted microscope. As ratio to control samples (%), quantitative data were measured by manual counting and recorded.
MTT assay
As discussed elsewhere, cell viability is an important factor to evaluate the activity of a nanosystem by detecting the ability of cells to transform MTT to a purple formazan dye. Different cells (2x10^4 cells/mL) were incubated in 96-well plates and cultivated, then different concentrations of Tf-NGO@Pt were added at different times. Subsequently, the cells were incubated with 20 mL PBS solution with MTT for 5 hours. Finally, the medium was replaced by 200 μL dimethyl sulfoxide to dissolve purple formazan and UV absorption at 570 nm measured to assess cell viability.

Flow cytometry
Changes in cell-cycle distribution were measured by flow cytometry. Cells were collected and washed three times, then held in precooled 70% ethanol at −20°C overnight. Fixed cells were incubated with propidium iodide. After being dyed for 1 hour, the cell-cycle distribution for treated cells was measured by flow cytometry (Epics XL; Beckman Coulter, Brea, CA, USA) using MultiCycle software to analyze the data.

TUNEL–DAPI costaining assay
DNA fragmentation in MCF7 and SV-HUVC1 cells caused by Tf-NGO@Pt was confirmed by terminal deoxynucleotidyl transferase dUTP nick end labeling (TUNEL) assay. Tf-NGO@Pt was added to MCF7 and SV-HUVC1 cells and incubated for 24 hours. Cells were blended with formaldehyde (3.7%) for 10 minutes and incubated with Triton X-100 (0.1%). Cells were then incubated with TUNEL reaction mixture for 1 hour, then 1 mg/mL DAPI was added at 37℃. After 15 minutes, cells were rinsed with PBS and measured by fluorescence microscopy (Nikon Eclipse 80i).

Intracellular reactive oxygen species generation
Production of intracellular ROS in cells induced by Tf-NGO@Pt was measured by fluorescence intensity in cells, which were stained with dihydroethidium (DHE; Beyotime, Jiangsu, People’s Republic of China). The collected cells were rinsed three times and resuspended in PBS, and next the cells were incubated with DHE (10 μM) at 37°C. After 30 minutes, the various concentrations of Tf-NGO@Pt were added to cells for different periods. Finally, the level of intracellular ROS production was monitored with a microplate reader at 300 and 610 nm.

Statistical analysis
All experiments were carried out at least in triplicate, and results are expressed as means ± SD. SPSS was used for statistical analysis. Statistical significance between two groups was analyzed using P<0.05 or P<0.01. Bars with different characters show statistical significance at P<0.05.
Overall, the Tf-NGO@Pt remained stable in conditions of aqueous solution and human plasma. In Figure S2B, changes in \( \zeta \)-potential for Tf-NGO@Pt in two kinds of solutions are verified over 40 days. The \( \zeta \)-potential for Tf-NGO@Pt was 40.7 mV, and only slightly reduced to average 38.4 mV after 30 days and change to 37.6 mV in 40 days.

**In vitro anticancer activity of Tf-NGO@Pt**

To determine the selectivity for Tf-NGO@Pt, MG63, HeLa, CaSki, SiHa, MCF7, A375, and HepG2 malignant cells were used as cancer-cell models, and Chem5, NIH3T3, HK2, and SV-HUVC as normal-cell models. Table 1 shows that the free nanocarrier of NGO and NGO-PEI was not cytotoxic against any cell lines at 2 mg/mL NGO and NGO-PEI. Tf-NGO@Pt significantly enhanced anticancer activity for Pt and simultaneously reduced cytotoxicity toward normal cells. For example, Tf-NGO@Pt showed a 15-fold greater anticancer effect on MCF7 cells, and SV-HUVC cells were even more cytotoxic. These results indicated that the IC\(_{50}\) of Tf-NGO@Pt was 6.5 \( \mu \)M and most sensitive to MCF7 cells. Tf-NGO@Pt considerably inhibited MG63-cell growth in a dose-dependent manner. Moreover, Figure S3 shows that Tf-NGO@Pt had higher anticancer effects than free-Pt complexes or NGO-Pt-PEI in MCF7 cells at 24 and 48 hours. Pt complexes and NGO-Pt-PEI expressed negligible cytotoxicity in 24 hours. Furthermore, Tf-NGO@Pt was noncytotoxic toward SV-HUVC cells (IC\(_{50}\) 95.8 \( \mu \)M). In brief, Tf-NGO@Pt showed outstanding selectivity for cancer cells, ensuring its effective application with safe doses in vivo. It is possible that, the advantage of Tf-NGO@Pt could be due to Tf protein-surface decoration, which could effectively improve the selectivity of Tf-NGO@Pt between cancer and normal cells.

**Tf-NGO@Pt cellular uptake**

Cellular uptake assays were conducted to verify the selectivity of Tf-NGO@Pt. In brief, we quantified Tf-NGO@Pt-uptake efficacy in MCF7 and SV-HUVC cells by the standard-curve method from internalized fluorescein isothiocyanate-loaded nanoparticles to assess Tf-NGO@Pt selectivity. Figure 2A and B shows that this was time dependent in the continuous process of Tf-NGO@Pt intracellular uptake in MCF7 and SV-HUVC cells, and cellular uptake of Tf-NGO@Pt in MCF7 cells was considerably higher (Figure 2A). However, Tf-NGO@Pt compared with Pt complexes showed lower fluorescence intensity in SV-HUVC cells (Figure 2B). These results were expected, and indicate the feasibility of Tf-guided selectivity in Tf-NGO@Pt between cancer cells and normal cells.
The efficiency of drug retention is a crucial parameter in biological application. Therefore, we examined the fluorescence intensity of Pt complexes and Tf-NGO@Pt in cancer cells, as shown in Figure 2C. Tf-NGO@Pt compared with Pt complexes had significantly higher intensity in MCF7 cells in a time-dependent manner. However, Figure 2D shows that retention efficacy was decreased in SV-HUVC cells. These outcomes suggest that Tf-NGO@Pt-enhanced retention overwhelmed Pt complexes, which ensured effective drug accumulation in cancer cells, dramatically improving its anticancer ability in MCF7 cells.

Many studies have shown that ligand/receptor-mediated targeting is an important pathway in drug selectivity between cancer and normal cells. Therefore, we investigated Tf-NGO@Pt selective cytotoxicity mechanisms. A Tf competing assay was conducted to verify the role of Tf/TfR-mediated targeting in the process of Tf-NGO@Pt causing cell death. Figure 3 shows that excess Tf dose dependently blocked Tf-NGO@Pt-enhanced retention.

Table 1 In vitro cytotoxicity of Tf-NGO@Pt (72 hours)

| Samples          | IC<sub>50</sub> (μM) |
|------------------|----------------------|
|                  | MG63     | HeLa     | CaSki    | SiHa     | MCF7     | A375     | HepG2    | Chem5*   | NIH3T3*  | HK2*     | SV-HUVC1* |
| NGO (mg/mL)      | 2.0      | 2.0      | 2.0      | 2.0      | 2.0      | 2.0      | 2.0      | 2.0      | 2.0      | 2.0      | 2.0      |
| NGO-PEI (mg/mL)  | 132.7    | 34.4     | 35.3     | 35.8     | 29.3     | 59.8     | 55.8     | >150     | >150     | >150     | >150     |
| Pt complex       | 126.3    | 30.6     | 31.2     | 29.6     | 10.2     | 54.7     | 50.2     | >150     | >150     | >150     | >150     |
| NGO-Pt-PEI       | 110.6    | 25.2     | 25.6     | 23.4     | 6.5      | 50.2     | 46.7     | >150     | >150     | >150     | >150     |
| Tf-NGO@Pt        | 110.6    | 25.2     | 25.6     | 23.4     | 6.5      | 50.2     | 46.7     | >150     | >150     | >150     | >150     |

Notes: *Chem5, NIH3T3, HK2, and SV-HUVC1 are normal cells.
Abbreviations: Tf, transferrin; NGO, nanographene oxide; PEI, polyethylenimine.

Figure 2 Selective cellular uptake and intracellular retention of Tf-NGO@Pt.

Notes: Quantitative analysis of 6-coumarin-loaded Pt complex (12 mM) and Tf-NGO@Pt (12 mM) by cellular uptake in MCF7 (A) and SV-HUVC1 (B). Quantitative analysis of 6-coumarin-loaded Pt complex (12 mM) and Tf-NGO@Pt (12 mM) by retention efficacy in MCF7 (C) and SV-HUVC1 (D).

Abbreviations: Tf, transferrin; NGO, nanographene oxide.
Pt uptake into MCF7 cells. Therefore, the Tf-NGO@Pt achieved upregulated cellular uptake by specifically binding to TfRs on MCF7 cells. As such, the excess Tf treatment dramatically declined Tf-NGO@Pt toxicity in MCF7 cells: 69.24% vs 23.35% from a single Tf-NGO@Pt treatment by reduced amount of internalized Tf-NGO@Pt. Taken together, these outcomes show that Tf-NGO@Pt obviously targeted cancer cells and reduced normal-cell uptake, on account of TfR-mediated endocytosis. Tf-NGO@Pt has superior selectivity uptake between cancer and normal cells.

**Tf-NGO@Pt intracellular localization, uptake pathways, and pH-responsive drug release**

Figure 4 shows the process of intracellular location of Tf-NGO@Pt in MCF7 cells. Imaging was performed to visualize the subcellular localization of Tf-NGO@Pt (Figure 4A). DAPI (blue) and LysoTracker (red) were specific probes used to label the nucleus and lysosome, respectively. Figure 4A shows that Tf-NGO@Pt localized in the lysosomes within 1 hour after being internalized, with some further gradual accumulation in the lysosome over 8 hours, shown as the high coincidence of green and red fluorescence in the cells, and shifted to the nucleus in MCF7 cells. These outcomes suggest that the lysosome is the main target of Tf-NGO@Pt after being internalized.

To examine Tf-NGO@Pt-endocytosis mechanisms, cells were pretreated with four series of endocytosis inhibitors at the initial stage. Figure 4B shows that with the combination of 2-deoxy-D-glucose and Na$_3$H$_2$PO$_4$ treatments and low-temperature (4°C), cellular internalization of Tf-NGO@Pt was reduced to 38.68% and 72.22%, which indicates that Tf-NGO@Pt is internalized through endocytosis by energy dependent pathways to enter MCF7 cells. There were three main mechanisms of endocytosis: clathrin, caveolae/lipid raft-mediated apoptosis, and phagocytosis/macropinocytosis. Since MCF7 cells are not phagocytic, other pathways were investigated. The specific receptor TfR was conjugated to Tf through lipid raft-coated vesicles to bind ferric ions. Tf-NGO@Pt cellular uptake was estimated using serial treatment of sucrose, an inhibitor of clathrin-mediated endocytosis. Sucrose decreased Tf-NGO@Pt uptake to 35.37% of control, demonstrating that clathrin-mediated endocytosis was the primary pathway. Dynamin, a GTP-binding protein, is required for receptor-mediated endocytosis. Dynasore, a specific inhibitor, was necessary for dynamin-mediated lipid-raft endocytosis-restricted Tf-NGO@Pt internalization to 53.9% of control, which indicates that the dynamin-mediated pathway was the primary pattern of lipid raft-dependent endocytosis of Tf-NGO@Pt in MCF7 cells. Nystatin, a specific inhibitor for lipid raft-dependent endocytosis, reduced Tf-NGO@Pt uptake to 61.36%, indicating that lipid raft-dependent endocytosis still caused Tf-NGO@Pt endocytosis. On the whole, Tf-NGO@Pt cellular uptake in MCF7 cells involved clathrin-mediated endocytic and dynamin-dependent lipid raft-mediated pathways.

To further identify clathrin-mediated endocytosis as the primary endocytic pathway, the cellular viability of MCF7 cells was determined after treatment with sucrose and Tf-NGO@Pt, as shown in Figure 4C. Pretreatment with sucrose significantly reduced Tf-NGO@Pt-caused cytotoxicity in MCF7 cells. Therefore, clathrin mediated endocytosis was confirmed as the primary endocytic pathway. Sucrose decreased Tf-NGO@Pt uptake to 35.37% of...
solution, the NGO shell was significantly dissolved, leading to rapid drug release, owing to enzymes or intracellular environments. Therefore, Tf-NGO@Pt is a bioresponsive nanomedicine within the intracellular environment.

In summary, Tf-NGO@Pt cellular uptake in MCF7 cells involves dynamin-dependent lipid raft- and clathrin-mediated endocytic pathways. For drug release, Tf-NGO@Pt powder was dispersed in PBS solution as buffer to simulate the physiological environment in cancer cells, as shown in Figure 5. Pt-complex release was 2.19%, 4.19%, 6.62%, and 7.85% for pH 7.4, 6.8, 5.3, and 5.3+ buffer, respectively, within 1 hour, and increased to 26.82%, 47%, 63.38%, and 72.46%, respectively, within 72 hours.

Generally, the NGO layer was negatively charged (Figure S2B) and conjugated with positively charged complexes. However, the complexes may compete with protons in the solution under acidic environments for interaction with the nanoparticle composition, which could allow free complexes to be released into the solution from the nanoparticle. This was verified by the nanosystem possessing pH-responsive drug release, as indicated. In cell-lysis

Figure 4 Intracellular localization and endocytosis mechanism of Tf-NGO@Pt.

Notes: (A) Intracellular location of Tf-NGO@Pt in MCF7 cells. Magnification ×200. (B) Intracellular uptake of Tf-NGO@Pt in MCF7 cells. *P<0.05; **P<0.01. (C) Cytotoxicity of Tf-NGO@Pt in MCF7 cells after sucrose pretreatment.

Abbreviations: Tf, transferrin; NGO, nanographene oxide; DAPI, 4′,6-diamidino-2-phenylindole; DG, 2-deoxy-D-glucose.

Figure 5 Drug release of complex in Tf-NGO@Pt and in vitro release of Pt complex from nanosystem in different PBS solutions.

Abbreviations: Tf, transferrin; NGO, nanographene oxide.
Selective induction of cancer-cell apoptosis

Apoptosis has been identified as a main action mechanism in cancer treatment with metal complexes. This study verified Tf-NGO@Pt-action modes by flow cytometry. Figures 6 and S4 show that Pt complexes and Tf-NGO@Pt induced large-scale apoptosis in MCF7 cells (see the sub-G1 populations in the graphs). Tf-NGO@Pt possessed a particularly higher level of apoptosis than Pt complexes. For example, MCF7 cells treated with 3, 6, and 12 μM Tf-NGO@Pt increased from 0.2% (control) to 17%, 31.7%, and 46.4%, significantly larger than those of Pt complexes (7.5%, 10.1%, and 16.3%, respectively) at equivalent concentrations. In contrast, Pt complexes and Tf-NGO@Pt had negligible apoptosis in SV-HUVC cells. Furthermore, as shown in Figure 7, TUNEL-assay results showed that Tf-NGO@Pt caused a dose-dependent increase in DNA fragmentation (green fluorescence) and nucleus condensation (blue fluorescence) in MCF7 cells. These results demonstrate that cell death induced by Tf-NGO@Pt nanoparticles was caused mainly by apoptosis. In sum, Tf-NGO@Pt was verified to induce MCF7 apoptosis, but showed lower effects on normal cells.

Tf-NGO@Pt triggers ROS overproduction in cancer cells

ROS plays an important role in cell apoptosis, which involves hydrogen peroxide, superoxide, and hydroxyl radicals. High ROS levels induce macromolecular damage and ROS-mediated apoptosis. To investigate intracellular ROS induced by Tf-NGO@Pt, we examined intracellular ROS levels by measuring the DHE-fluorescence intensity. Figure 8A and B shows that Tf-NGO@Pt overwhelmed Pt complexes by triggering ROS overproduction in MCF7 cells after 5 minutes of treatment, then reached 232.78% at 50 minutes, with a gradual reduction to 147.97%. In contrast, Pt complexes and Tf-NGO@Pt caused lower ROS levels in SV-HUVC cells after 2 hours. Figure 8C shows that Tf-NGO@Pt induced strong fluorescence intensity in MCF7 cells but only slight fluorescence in SV-HUVC cells, which verifies selective ROS induction by Tf-NGO@Pt. In contrast, Tf-NGO@Pt caused lower ROS levels in SV-HUVC cells compared to Pt after 2-hour incubation, which can be attributed to limited Tf-NGO@Pt uptake, as discussed earlier. Therefore, Tf-NGO@Pt exerted significantly higher anticancer effects in MCF7 cells via ROS-activated apoptosis.

Figure 6 Flow-cytometry analysis of MCF7 cells after incubation with Pt complex and Tf-NGO@Pt.
Abbreviations: Tf, transferrin; NGO, nanographene oxide.
Figure 7 Representative images of DNA fragmentation and nuclear condensation.

Note: Fragmentation and condensation caused by 24-hour treatment of Pt complex (12 μM) and Tf-NGO@Pt (6 and 12 μM), as detected by TUNEL assay and DAPI staining. Magnification ×200.

Abbreviations: Tf, transferrin; NGO, nanographene oxide; TUNEL, terminal deoxynucleotidyl transferase dUTP nick end labeling; DaPI, 4′,6-diamidino-2-phenylindole.

Figure 8 Changes of ROS generation induced by Tf-NGO@Pt.

Notes: (A) MCF7 cells were treated with Pt complex and Tf-NGO@Pt for 120 minutes; (B) SV-HUVC1 cells were exposed to different concentrations of Pt complex and Tf-NGO@Pt for 120 minutes; (C) MCF7 and SV-HUVC1 cells were dyed by 10 μM DHE at 0, 10, 20, 40, 80, 120 minutes. Magnification ×200.

Abbreviations: ROS, reactive oxygen species; Tf, transferrin; NGO, nanographene oxide; DHE, dihydroethidium.
Tf-NGO@Pt penetration ability and inhibitory effects on MCF7 spheroids

Less sensitivity to drugs has been reported for cancer cells in tumors than in cultured cells. Recently, tumor-spheroid cell-culture systems have received increasing interest, because they can restore in vivo environments. In particular, tumor spheroids with diameters up to 200 μm sufficiently reflect in vivo like cell–cell and cell–matrix interactions. This study successfully cultivated MCF7 spheroids following an in vitro formation process (Figure 9A).

To examine the penetration ability of different Tf-NGO@Pt nanosystems, MCF7 spheroids were treated with fluorescein isothiocyanate-labeled nanoparticles for 12 hours. The resultant tumor spheroids were scanned at different layers from the top of the spheroid to the middle using confocal laser-scanning fluorescent microscopy, as shown in Figure 9C. Tumor spheroids treated with the nanoparticles displayed varied fluorescence intensity. Among them, Tf-NGO@Pt showed excellent advantages in penetrating ability of the MCF7 spheroids, reaching the core.

Figure 9B shows MCF7 spheroid-volume ratios measured after treatment with free complexes and Tf-NGO@Pt nanoparticles. Tf-NGO@Pt significantly inhibited growth of the MCF7 spheroids. For example, Tf-NGO@Pt reduced MCF7 spheroid-volume ratios by 71% of control after 120 hours, significantly more effective than free-Pt complexes (31.4%). Therefore, although formation of a protein corona was observed, Tf-NGO@Pt showed excellent advantages in penetration ability and inhibitory effects for future clinical use.

Tf-NGO@Pt suppresses MCF7-cell migration and invasion in vitro

Cell migration and invasion are also pivotal for tumor growth. To investigate the antimetastasis and invasion ability of Tf-NGO@Pt, we conducted wound-healing and transwell assays, as shown in Figure 10A and D. Migration and invasion of MCF7 cells were effectively suppressed by free complexes, but Tf-NGO@Pt exhibited significantly higher inhibition than the free complexes. In particular, Tf-NGO@Pt exhibited higher activity than the free complexes. Figures 9C and 10B show the migratory and invaded ratios. These results further confirm that functionalized Tf-NGO@Pt nanosystems possess remarkable anticancer activity and application potential in cancer therapy.

Figure 9 Penetration ability and inhibitory effect toward MCF7 spheroids.
Notes: (A) The formation process of MCF7 tumor spheroids. (B) Inhibitory effect of Pt complex and Tf-NGO@Pt toward MCF7 spheroids. Magnification ×200. Values presented as mean ± SD of triplicate results. (C) Penetration ability of Tf-NGO@Pt to MCF7 tumor spheroids. The spheroids were treated with coumarin-6-labeled nanoparticles for 12 hours. Drug concentration was 60 mM.

Abbreviations: Tf, transferrin; NGO, nanographene oxide.
Figure 10 Wound-healing assay of Pt complex and Tf-NGO@Pt on MCF7 cells (4×10^5 cells/L).
Notes: (A) Migratory assay in MCF7 cells induced by Pt complex and Tf-NGO@Pt. (B) Migratory ratio of Tf-NGO@Pt. Quantitative data were analyzed by manual counting (% of control). (C) Invasion assay in MCF7 cells induced by Pt complex and Tf-NGO@Pt. (D) Invasion ratio of Pt complex and Tf-NGO@Pt. Magnification ×200. Quantitative data were analyzed by manual counting (% of control). Values expressed as mean ± SD of triplicate results.
Abbreviations: Tf, transferrin; NGO, nanographene oxide.

Conclusion
In summary, we have design and synthesized a cancer-targeted Tf-NGO@Pt nanosystem, in order to overcome the poor solution of metal complexes and to enhance their anticancer efficacy. The Tf-NGO@Pt nanosystem through transferrin decorated effectively enhances the cellular uptake of drugs with increased discrimination between cancer and normal cells. The efficacy and safety index of Tf-NGO@Pt were greater than that of the free-metal complexes. This effect was closely related to the enhancement of cellular uptake and cellular drug retention. Moreover, Tf-NGO@Pt nanoparticles have shown excellent advantages in penetration ability and inhibitory effect toward MCF7 spheroids compared to free complexes. This study highlights the usefulness of cancer-targeted nanoparticles for delivery of anticancer metal complexes to achieve enhanced anticancer efficacy and to expand the usefulness of metal complexes.
Disclosure

The authors report no conflicts of interest in this work.

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Supplementary materials

Figure S1 Structural characterization of Tf-NGO@Pt complex.
Notes: (A) Fourier-transform infrared spectra of Tf-NGO@Pt; (B) ultraviolet-visible spectra of Tf-NGO@Pt.
Abbreviations: Tf, transferrin; NGO, nanographene oxide.

Figure S2 Structural characterization of Tf-NGO@Pt complex.
Notes: (A) Time course of size distribution of Tf-NGO@Pt (2 mg/mL); (B) time course of ζ-potential of Tf-NGO@Pt (2 mg/mL).
Abbreviations: Tf, transferrin; NGO, nanographene oxide.
**Figure S3** Cytotoxicity of Pt complex, NGO-Pt-PEI, and Tf-NGO@Pt in MCF7 cells at 24, 48, and 72 hours.

**Abbreviations:** Tf, transferrin; NGO, nanographene oxide.

**Figure S4** Induction of cancer-cell apoptosis by Tf-NGO@Pt and flow-cytometry analysis of Pt complex and Tf-NGO@Pt-treated SV-HUVC1 cells for 72 hours.

**Abbreviations:** Tf, transferrin; NGO, nanographene oxide.