Long non-coding RNA HOTTIP enhances IL-6 expression to potentiate immune escape of ovarian cancer cells by upregulating the expression of PD-L1 in neutrophils

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Abstract

Background: Long non-coding RNA (lncRNA) HOXA transcript at the distal tip (HOTTIP), has been demonstrated to be a vital biomarker when evaluating the prognosis of multiple cancers. Nevertheless, the potential function of HOTTIP in ovarian cancer (OC), a prevalent cancer among women worldwide, remains elusive. Hence, the current study aimed to elucidate the functional relevance of HOTTIP in the development of OC.

Methods: Positive expression of PD-L1 and IL-6 was determined using immunohistochemical staining in the collected OC and normal tissues. The correlation of IL-6 and PD-L1 was analyzed using flow cytometry, Western blot analysis as well as Pearson’s correlation coefficient. The interaction among HOTTIP, c-jun and IL-6 was investigated with the use of RIP, ChIP and dual luciferase reporter gene assays. Finally, the effects of HOTTIP on T cell proliferation and infiltration were identified through gain- and loss-of-function studies in vitro and in vivo.

Results: HOTTIP, IL-6 and PD-L1 were all highly expressed in OC tissues. A positive correlation was observed between IL-6 and PD-L1 and that between HOTTIP and IL-6 in OC tissues. HOTTIP was noted to promote the expression of IL-6 by binding to c-jun, which resulted in a promoted PD-L1 expression in neutrophils and immune escape while inhibiting T cell proliferation as well as tumor immunotherapy.

Conclusion: Taken together, our study unveiled that HOTTIP could promote the secretion of IL-6, and consequently up-regulate the expression of PD-L1 in neutrophils, thus inhibiting the activity of T cells and ultimately accelerating immune escape of OC cells. Our study provides a potential therapeutic strategy by targeting HOTTIP in OC.

Keywords: Ovarian cancer, Neutrophils, Interleukin-6, Programmed death-ligand 1, HOXA transcript at the distal tip, Immune escape

Background

Ovarian cancer (OC) is considered as the most fatal gynecologic malignancy, and the commitments for early detection and novel therapies to lower its mortality have been principally unproductive [1]. Cancer immunotherapy has emerged as a promising therapeutic approach in oncology, and is characterized by the activation of immune system to induce tumor immune surveillance or to reverse the tumor immune escape [2]. Marth et al. have previously highlighted the promising therapeutic effect of immunotherapy in limiting OC [3]. In addition, therapeutic immunotherapy has also been demonstrated to have the ability to improve pancreatic cancer clinical trial outcomes [4]. However, some of the cases in the clinical trials failed as a result of “immune escape” [5]. The host immune system would respond to appearance of tumor cells, while neutrophils signify the first line of host defense against infection [6]. Besides, a previous study conducted on hepatocellular carcinoma (HCC)
revealed that peri-tumoral neutrophils could negatively modulate the adaptive immunity through program death ligand 1 (PD-L1) and PD-1 pathway [7]. Regulatory T (Treg) cells are critical in immune tolerance in patients, and there is an inverse correlation between intertumoral Treg and patients’ survival [8]. Moreover, tumor-related neutrophils can recruit Treg cells, thereby promoting the development of HCC [9].

Long non-coding RNA (lncRNA) is a class of RNA molecules longer than 200 nucleotides that do not have functional protein-coding capacity [10]. The progression of OC has been suggested to be closely associated with lncRNAs [11]. For instance, the clinical significance of HOXA11 has been highlighted in predicting the prognosis of patients suffering from serous OC [12]. Moreover, HOXA transcript at the distal tip (HOTTIP) has been found to have a functional role in the pathogenesis of breast cancer, which is the most common cause of cancer-related death in women via regulation of one of its physical HOXA clusters, HOXA11 [13]. The results from the bioinformatics prediction revealed a binding site between HOTTIP and interleukin-6 (IL-6). IL-6 regulates chemokine generation and leukocyte apoptosis and is therefore a crucial checkpoint regulator of neutrophil trafficking in the inflammatory response [14]. IL-6 is known to have the ability to upregulate PD-L1 expression in tolerogenic antigen-presenting cells by elevating signal transducer and activator of transcription 3 (STAT3) [15]. PD-L1 suppresses host immunity through binding to its receptor PD-1 on lymphocytes, and promotes peritoneal dissemination, thus exacerbating OC [16]. Based on the aforementioned information, we proposed a hypothesis that HOTTIP may affect OC involving the regulation of IL-6 and PD-L1. Thus, the current study aims to investigate the mechanism by which HOTTIP/IL-6/PD-L1 axis influences immune escape of OC cells.

Materials and methods

Ethics statement

This study was conducted with the approval of the Ethics Committee of Tongji Hospital of Tongji University and The Sixth People’s Hospital of Yancheng City. Written informed consents were provided and signed by all participants prior to sample collection. All animal experiments were conducted in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health.

Study subjects

A total of 53 cases of OC tissues were collected from patients with OC who underwent surgical resection at Tongji Hospital of Tongji University and The Sixth People’s Hospital of Yancheng City from May 2017 to October 2018. The average age of OC patients was (54.04 ± 11.88) years, out of which there were 32 patients > 50 years, and 21 patients ≤ 50 years. The clinical characteristics of the patients are listed in Table 1. In addition, normal fallopian tube fimbriae tissues were obtained from 25 patients with benign lesions near the uterus. The average age of normal subjects was (54.24 ± 10.57) years. A part of the specimen was used for the extraction of the total RNA and tumor-related neutrophils. The corresponding information of patients to the paraffin sections was recorded. Histologic diagnosis and grading were performed based on the standards of World Health Organization Classification of Tumors of the Digestive System. The clinical follow-up data included imaging examinations and assessment of survival of patients. The clinical tumor recurrence was defined as tumor recurrence or metastasis examined by clinical imaging. The overall survival (OS) was defined as the intermission between surgery and death.

Immunohistochemical staining

The sections were dewaxed with dimethylbenzene, hydrated with gradient alcohol, repaired in the citrate repair solution under high pressure and high temperature for 1.5 min, and cooled down at room temperature. After receiving a wash with phosphate-buffered saline (PBS), each section was incubated with 50 μL 3% H2O2 at room temperature for 20 min to eliminate the activity of endogenous peroxidase. Each section was then probed overnight with rabbit antibodies against PD-L1 (ab228415), IL-6 (ab9324) or CD3 (ab16669) in a 4°C freezer, with the normal rabbit serum serving as the negative control (NC) instead. All of the above antibodies were purchased from Abcam Inc. (Cambridge,

| Table 1 Clinical characteristics of the patients with OC |
|----------------|-------------------|
|                | Normal           | Ovarian tumor |
| Case           | 25               | 53            |
| Age (years)    | 54.24 ± 10.57    | 54.04 ± 11.88 |
| > 50           | 16               | 32            |
| ≤ 50           | 9                | 21            |
| Tumor stage (n)|                  |               |
| I              | 20               |               |
| II             | 23               |               |
| III            | 7                |               |
| IV             | 3                |               |
| Pathological classification (n) | | |
| Serous         | 27               |               |
| Mucinous       | 15               |               |
| Endometrioid   | 9                |               |
| Clear cell carcinoma | 2 | |

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MA, USA). Next, each section was incubated with 50 μL polymer enhancer at 37 °C for 20 min and 50 μL enzyme-labeled goat anti-rabbit immunoglobulin G (IgG) at 37 °C for 30 min. Each section was added with 2 drops or 100 μL freshly prepared diaminobenzidine (DAB) for development and observed under a microscope for 3–10 min. A positive result was indicated by the presence of brown staining. After being washed with distilled water, the sections were counterstained with hematoxylin, dehydrated with gradient alcohol, mounted and observed under the microscope.

The expression of PD-L1 and IL-6 was characterized by the presence of fine brown granules in tumor cells. Grading was performed in accordance with the percentage of positive cells and the staining intensity: 0 point, the rate of positive cells ≤10%; 2 points, 11–51% positive cells; 3 points, 51–81% positive cells; 4 points: the rate of positive cells ≥81%; 1 point, weak intensity; 2 points, medium intensity; 3 points, high intensity. The total score was calculated by adding the two scores above: 0–7 points.

Collection and preservation of blood samples from patients

The patients with OC confirmed by pathologists after surgery were subjected to peripheral venous blood collection using anticoagulant tubes. The blood was then transferred to the laboratory in an immediate manner and stored at 4 °C for subsequent experiments.

Cell culture

The 293 T cells were obtained from the Cell bank of the Chinese Academy of Sciences (Shanghai, China), and SKOV3, OVCAr3, and Hy-A8 cells were obtained from the American Type Culture Collection (ATCC, Manassas, VA, USA). These cell lines were cultured in Dulbecco’s modified Eagle’s medium (DMEM; 31,800,022, Gibco, Carlsbad, CA, USA) containing 1.5 g/L NaHCO₃, 10% fetal bovine serum (FBS) and 1% penicillin and streptomycin. Firstly, 5 mL medium was transferred into 15 mL sterile centrifuge tubes with sterile pipettes and water-bathed at 37 °C to melt the cells quickly. The cells then underwent centrifugation at 800 rpm for 5 min and were re-suspended in 1 mL culture solution. The cell suspension was subsequently cultured in 6 cm culture dish added with 2 mL solution in an incubator with 5% CO₂ at 37 °C. The cell growth was observed the following day and the culture solution was changed every 1–2 days.

Neutrophil isolation and identification

Extraction and identification of neutrophils from peripheral blood were conducted. Briefly, a total of 5 mL fresh heparin anticoagulant was collected from healthy adults. The serum was diluted with aseptic PBS at the ratio of 1:1. Afterwards, 3 mL of lymphocyte separation medium was added into a 15-mL centrifuge tube, followed by careful addition into the diluted serum. The tube was then centrifuged at 2000 rpm for 20 min. The intermediate layer of peripheral blood mononuclear cells (PBMCs) was transferred to a new centrifuge tube, washed twice with PBS and then centrifuged at 1500 rpm for 10 min. After resuspension with Roswell Park Memorial Institute (RPMI)-1640 medium containing penicillin and streptomycin, a flow cytometer was employed for the sorting of the CD66b⁺ cells (130–104-413, 1:10; Miltenyi biotec, Bergisch Gladbach, Germany).

Tumor infiltrated neutrophil separation was conducted. In short, the tissues were cut into 0.5 cm × 0.5 cm blocks, transferred into a tube with 10 mL DMEM medium, 100 μL collagenase IV, 100 μL DNase I, 100 μL MgCl₂ as well as 50 μL CaCl₂, and then mixed evenly. The tissue blocks were centrifuged and ground, and filtered with a 200-mesh steel mesh. The single cell suspension was harvested in a centrifuge tube (50 mL) and centrifugation was carried out for 10 min at 1700 rpm. The cells were re-suspended with PBS and filtered again. Next, the cells were collected in two centrifuge tubes (1.5 mL) and centrifuged for 5 min at 2000 rpm. Finally, the CD66b⁺ cells were sorted with the use of flow cytometry.

T cell isolation and identification

Heparin anticoagulant was added and the cells centrifuged, after which the serum was discarded. The erythrocytes were lysed with 3 volumes of erythrocyte lysis buffer at room temperature for 10 min. After centrifugation for 5 min, the cells were washed with 5 mL PBS, centrifuged, and re-suspended with sterile PBS followed by cell counting. The CD3⁺ cells were sorted by flow cytometry with the PE-labelled CD3 antibody (130–113-129: 1:50; Miltenyi biotec, Bergisch Gladbach, Germany).

Flow cytometric analysis of surface marker and intracellular proteins

The cells were dispersed into single cell suspension and re-suspended by the staining buffer (BD Biosciences, San Jose, CA, USA). The neutrophils were identified using FITC-labeled anti-CD66b (130–104-413, 1:10; Miltenyi biotec, Bergisch Gladbach, Germany). Allophycocyanin (APC)-labeled PD-L1 antibody (130–117-694, 1:10; Miltenyi biotec, Bergisch Gladbach, Germany) was used to determine PD-L1 expression. PE-labeled CD3 antibody (130–117-139, 1:50; Miltenyi biotec, Bergisch Gladbach, Germany) was used for
the separation of T cells and Pacific blue-labeled interferon (IFN)-γ antibody (#505817, 1:50; BioLegend, San Diego, CA, USA) was used for the measurement of IFN-γ content. The detection of intracellular proteins required a membrane-breaking perforation of the cells. Flow cytometry was performed on the FACS Canto II analyzer (Becton Dickinson, Franklin Lakes, NJ) and the data were analyzed by the FlowJo software.

**Cell treatment**

The pLKO.1 plasmids (containing Puromycin resistance gene) knocking down HOTTIP, IL-6 or c-jun, the pBABE plasmids (containing Puromycin resistance gene) overexpressing HOTTIP or c-jun and the lentiviral package plasmids (pVSVG-G, pREV, pMDL, PIK) were purchased from Cyagen (Suzhou, Jiangsu, China). The plasmids were transduced to the E.coli DH5α competent cells. The corresponding plasmids were extracted using plasmid extraction kit (DP103–03, TIANGEN Biotechnology Co., Ltd., Beijing, China). The plasmid was transduced into the HRK 293 T cells in strict accordance with the instructions of the Turbofect transfection reagent (R0531, Thermo Fisher Scientific Inc., Waltham, MA, USA). After 12 h, the culture medium was replaced, and collected at 24th and 48th h, respectively. The viruses were obtained after being filtered with the 0.45 μm filter, after which the OC cell line was infected. The corresponding cell line stably silencing HOTTIP, IL-6 or c-jun or that overexpressing HOTTIP or c-jun was screened out using Puromycin (2 μg/mL).

**Detection of the proliferation of T cells induced by neutrophils**

Anti-CD3/anti-CD28 tetramer antibody (STEMCELL Technologies, Vancouver, BC, Canada) was used to coat the 96-well plates. IL-2 (20 IU/mL) was added to the collected CD3⁺ T cells. CD3⁺ T cells labeled by Carboxyfluorescein succinimidyl ester (CFSE; S1076, Beijing Solabio Life Sciences Co., Ltd., Beijing, China) were co-cultured with the isolated neutrophils in RPMI-1640 with 5% CO₂ standing on ice for 5 min. Following 5 min of centrifugation at 4 °C, 500 g, the supernatant (i.e. cytoplasm) was removed into 2 mL aseptic and enzyme-free tube, centrifuged (500 g) 5 min at 4 °C. The cell precipitate was transferred into a fresh eppendorf (EP) tube, and then centrifuged at 8000 rpm for 8 min. The content of CFSE⁻ T cells was measured at 450 nm using a microplate reader [18].

**Determination of IL-6 secretion**

The serum or OC culture medium supernatant was transferred into a fresh eppendorf (EP) tube, and then centrifuged at 8000 rpm for 8 min. The content of IL-6 was determined with the application of the IL-6 enzyme-linked immunosorbent assay (ELISA) kit (ab178013, Abcam Inc., Cambridge, MA, USA). The 96-well plate was added with 100 μL of coating solution containing IL-6 and sealed at 4 °C overnight. Afterwards, the wells in each well were incubated with 200 μL ELISA diluent for 1 h at room temperature. Next, 50 μL ELISA diluent was used to dilute 50 μL standard products, a total of 8 wells. The remaining wells were sealed with 100 μL isolated serum at room temperature for 2 h, with 3 parallel wells set up. Then, the sample was probed with the secondary antibody (100 μL) for 1 h, then reacted with ELISA diluent for 40 min, and developed with 100 μL chromogenic solution for 5–10 min devoid of light at room temperature. Finally, 50 μL of termination solution was added into each well. The optical density (OD) value was measured at 450 nm using a microplate reader.

**RNA isolation and quantification**

The total RNA of tissues or cells was extracted using the Trizol kit (15596–018, Beijing Solabio Life Sciences Co., Ltd., Beijing, China), and the concentration of RNA was determined. The primers used in this study were synthesized by Takara Biotechnology (Dalian, Liaoaning, China) (Table 2). The reverse transcription was performed with 25 μL assay system based on the instructions of the complementary DNA (cDNA) reverse transcription kit (K1622, Reanta Co., Ltd., Beijing, China). The cDNA obtained was diluted to 50 ng/μL for the following RT-qPCR, which was carried out on the fluorescence quantitative polymerase chain reaction instrument (ViiA 7, DAAN Gene Co., Ltd. of Sun Yat-sen University, Guangzhou, Guangdong, China). Glyceraldehyde-3-phosphate dehydrogenase (GAPDH) was used as an internal reference. The fold changes were calculated by means of the relative quantification (2−ΔΔCt method).

**Fractionation of nuclear/cytoplasmic RNA**

The experiment was conducted in accordance with the PARIS™ Kit Protein and RNA Isolation System (Life Technologies, Carlsbad, CA, USA). In brief, OC cells were collected, washed with PBS, trypsinized and centrifuged (500 g) 5 min at 4 °C. The cell precipitate was rinsed with PBS, after which the supernatant was discarded. Next, 500 μL of cell fractionation Buffer was added to the cells and dissociated gently, followed by standing on ice for 5–10 min. Following 5 min of centrifugation at 4 °C, 500 g, the supernatant (i.e. cytoplasm) was removed into 2 mL aseptic and enzyme-free tube, followed by additional centrifugation at 4 °C, 500 g for 5 min. The precipitate obtained (i.e. nucleus) was added with 500 μL Cell Fractionation Buffer and mixed by

**Table 2 Primer sequences for RT-qPCR**

| Primer sequence | Forward (5′-3′) | Reverse (5′-3′) |
|-----------------|----------------|----------------|
| HOTTIP          | CCTAAAGCCAGGTCCTTTG | TGCAAGGCTGAGATCCCTA |
| GAPDH           | ACAACAGGCTCAAGATCA | TGCCACACCTGACAGST |

RT-qPCR reverse transcription quantitative polymerase chain reaction, HOTTIP HOXA transcript at the distal tip, GAPDH glyceraldehyde-3-phosphate dehydrogenase
flicking. Next, 500 µL 2 × Lysis/binding solution was dropped to the mixture, mixed well following gentle dissociation, and subsequently allowed to stand on ice. Following the removal of the supernatant, precooled 500 µL Cell Disruption Buffer was added and whirled violently for complete mixture. Thereafter, 500 µL absolute ethanol was added and mixed by gentle dissociation. The absorption column was put into collection tubes, after which 700 µL Wash solution 1 was added followed by 30-s centrifugation at 12000 g and the removal of the supernatant. The liquid in the collection tubes was harvested. Similarly, 500 µL Wash solution 2/3 was added and centrifugation was carried out at 12000 g for 30 s, after which the supernatant was withdrawn and the liquid in the tubes was collected. Next, the empty column was rotated at a maximum speed for 1 min, and the collection tubes were discarded. The adsorption column was loaded into a new collection tube, and was eluted by the addition of 40 µL Elution solution (water bath to 95 °C in advance) and 30 s of centrifugation at 12000 g, finally, reverse transcription quantitative polymerase chain reaction (RT-qPCR) was carried out to determine the expression of HOTTTIP in cytoplasm (with 12S rRNA regarded as the loading control) and nucleus (with 45S rRNA regarded as the loading control) (Table 3).

**Dual luciferase reporter gene assay**

The National Center for Biotechnology Information (NCBI) database (http://www.ncbi.nlm.nih.gov/gene) was adopted for the retrieval of the promoter sequence and gene sequence of IL-6. The promoter region of IL-6 gene was cloned into the pmirGLO vector (Promega Corporation, Madison, WI, USA), and the wild type (wt) pmirGLO-IL-6 prom vector (IL-6-prom-wt vector). The plasmids were co-transfected with the renilla luciferase expression vector pRL-TK (TaKaRa Biotechnology Co. Ltd., Liaoning, China) in accordance with the instructions provided on the lip2000. The cells were then transfected with plasmids of IL-6-prom-wt + NC, IL-6 prom wt + shRNA (sh)-c-jun + NC, IL-6 prom wt + pLV-EGFP-N-HOTTIP, IL-6 prom wt + c-jun, IL-6 prom wt + pLV-EGFP-N-HOTTIP + c-jun and IL-6 prom wt + sh-c-jun + pLV-EGFP-N-HOTTIP. Following transfection, the cells were cultured for 24 h, and Dual-Luciferase Reporter Assay System (Promega Corporation, Madison, WI, USA) was used to determine the dual luciferase activity in each group, with renilla luciferase pRL-TK as loading control (TaKaRa, Biotechnology Co. Ltd., Liaoning, China).

**Chromatin immunoprecipitation (ChIP) assay**

The binding sites between c-jun and IL-6 were predicted using the JASPAR website (http://jaspar.genereg.net/) (site 1: GCTATGATGCAATT; site 2: TGGATGACCTCAC). The cells were fixed by formaldehyde for 10 min to produce DNA-protein crosslinking. The DNA was sonicated for 10 s (15 times at an interval of 10 s). Following centrifugation at 12000×g at 4 °C for 10 min, the supernatant was collected and divided into two tubes, both of which underwent incubation overnight with rabbit antibody against c-jun (#9165, Cell Signaling Technologies, Beverly, MA, USA) or IgG (NC, ab109489, 1:300, Abcam Inc., Cambridge, MA, USA) at 4 °C. Protein Agarose/Seaphorase was used to precipitate DNA-protein complex. The non-specific complex was then washed, and the crosslinking was reversed at 65 °C overnight. The DNA fragment was purified and extracted by phenol/chloroform. The site 1 primers [forward (F) sequence: 5′-TGCACTCCCATTTGCTG-G-3′; reverse (R) sequence: 5′-TGATACCTGACCAGAAAGGCT-3′] and the sequences of site 2 primers (F: 5′-ACACAAAAAGATCCCCAGCAG-3′; R: 5′-ATTTCA-GGACCCGCCTTGG-3′) were designed. The length of the amplified product was 130 bp and 100 bp, respectively, and the transcription start sites (TSS) were 1846 bp and 27 bp. The sequence far from the promoter region of IL-6 gene was also amplified as the NC (F: 5′-ACACATTGTACATCCAGCAG-3′; R: 5′-GGGTAC-GCCAGAACATGGGAC-3′) with the length of 89 bp, and the TSS of 3867 bp. RT-qPCR was carried out using the primers. The purified DNA obtained by overexpression of HOTTTIP was the same as the above described with the site 2 primer used as the primers for RT-qPCR.

**RNA-binding protein immunoprecipitation (RIP)**

The cells were treated with radioimmunoprecipitation assay (RIPA) lysis buffer and then the lysate was isolated. The magnetic beads were incubated with rabbit anti-human c-jun (#9165, 1:100, Cell Signaling Technologies, Beverly, MA, USA) to construct the magnetic bead-c-jun antibody complex. The complex was then mixed with the lysate. Subsequently, isolation and purification of RNA from the complex were conducted.

**Western blot analysis**

After the treatment with the RIPA lysis buffer containing phenylmethanesulfonyl fluoride (PMSF) and phosphatase

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**Table 3** Primer sequences for fractionation of nuclear/cytoplasmic RNA

| Gene     | Sequence                      |
|----------|-------------------------------|
| 45S rRNA | F: 5′-GTGCCCTACAGTTTACTTT-3′ |
|          | R: 5′-TAGGAGACAAACCCTGAGCCT-3′ |
| 12S rRNA | F: 5′-TGATAAACCCCCCGCTACCT-3′ |
|          | R: 5′-TGCTACACCTTGGACTAACGT-3′ |

F: forward, R: reverse
inhibitor in an ice-bath for 30 min, the proteins were isolated by centrifugation at 12000 rpm at 4°C for 30 min. The protein concentration was determined with the use of the bicinchoninic acid (BCA) kit. Following separation by 10% SDS-polyacrylamide gel electrophoresis (PAGE), the proteins were transferred onto the polyvinylidene fluoride (PVDF) membrane by a wet transfer method. The membrane was then blocked with 5% skimmed milk at room temperature for 1 h and incubation was carried out with diluted rabbit primary antibodies: IL-6 antibody (ab9324), PD-L1 antibody (ab228415), STAT3 antibody (ab68153), p-STAT3 antibody (ab76315), and mouse anti-β-actin antibody (ab8226, 1:100). All of the aforementioned antibodies were purchased from Abcam Inc. (Cambridge, MA, USA). After being washed, the membrane was incubated with secondary antibody horseradish peroxidase (HRP)-conjugated goat anti-mouse IgG or goat anti-rabbit IgG (Transgene Biotech, Beijing, China) for 1 h. The immunocomplexes on the membrane were visualized using enhanced chemiluminescence (ECL) fluorescent detection kit (BB-3501, Amersham Pharmacia, Piscataway, NJ, USA) and band intensities were quantified using Bio-Rad image analysis system (Bio-Rad Laboratories, Hercules, CA, USA) and Quantity One v4.6.2 software. The ratio of the gray value of the target protein band to that of β-actin band was regarded as the relative protein expression.

In vivo xenograft experiments
Twenty-five non-obese diabetic/severe combined immunodeficient (NOD/SCID) mice (5–7 weeks old; weight 18–22 g; purchased from Shanghai Lingchang, Shanghai, China) were divided into 5 groups with 5 mice in each group. All animals were allowed to acclimatize for 1 week in the specific-pathogen-free (SPF) animal facility in Tongji Hospital of Tongji University and The Sixth People’s Hospital of Yancheng City before any intervention was initiated. The mice were provided with sterile feeding and drinking water with alternated day and night for 12 h. The SKOV3 single cell suspension was prepared with sterile and drinking water with alternated day and night for 12 h. The SKOV3 single cell suspension was prepared with the concentration of 2 × 10⁶ cells/mL and 0.2 mL of cell suspension was subcutaneously injected into the left axillary of the mice using 1 mL syringes. The mice (5 in each group) were intraperitoneally injected respectively with 100 μL PBS, neutrophils + T cells (5 × 10⁶ cells for either; 1:1, a total of 100 μL), SKOV3-neutrophils + T cells (5 × 10⁶ cells for either; 1:1, a total of 100 μL), SKOV3-neutrophils overexpressing HOTTIP + T cells (5 × 10⁶ cells for either; 1:1, a total of 100 μL), or PD-L1 antibody (20 μg/mL)-treated SKOV3-neutrophils overexpressing HOTTIP and T cells (5 × 10⁶ cells for either; 1:1, a total of 100 μL). Following inoculation, all the nude mice were kept in the SPF animal room.

The short diameter (a) and long diameter (b) of tumors were measured every 3 days using the calipers. The tumor volume was estimated using the following formula: \( \pi \left(\frac{a^2 \times b}{6}\right) \). Once the mice were euthanized, the tumors were weighed. A portion of the tissue was further excised for immunohistochemical staining, and another portion was used to prepare tumor-infiltrating lymphocyte (TIL) using the tumor dissociation kit (Miltenyi biotec, Bergisch Gladbach, Germany) and TIL gentleMACS dissociator (Miltenyi biotec, Bergisch Gladbach, Germany). TILs were isolated from tumor suspensions through density gradient centrifugation with gradient Percoll II solution (GE Healthcare, Pittsburgh, PA, USA). Flow cytometry was used for the analysis of IFN-γ expression in TILs [17].

Statistical analysis
All analyses were conducted using SPSS 21.0 statistical software (IBM Corp, Armonk, NY, USA). Measurement data were expressed as mean ± standard deviation. Data with normal distribution and homogeneity of variance between two groups were compared using paired t-test (paired data) or unpaired t-test (unpaired data). One-way analysis of variance was performed for comparisons among multiple groups, followed by Tukey’s post hoc test. The repeated measures analysis of variance with Bonferroni post hoc test was applied for the comparison of data at different time points. Correlation between parameters was evaluated by Pearson’s correlation coefficient. OS was assessed using the Kaplan-Meier method and the log-rank test. \( p < 0.05 \) was considered a statistically significant value.

Results
IL-6 is highly expressed in OC tissues and positively associated with PD-L1 in neutrophils
To evaluate the potential role of neutrophils in the immune system of OC patients, the number of neutrophils in peripheral blood collected from clinical samples was assessed by flow cytometry. Notably, OC patients presented with a greater number of neutrophils relative to normal controls (Fig. 1a, \( p < 0.05 \)). Next, infiltrated neutrophils in OC were isolated from peripheral blood using flow cytometry and then the expression of PD-L1 in neutrophils from peripheral blood and infiltrated neutrophils was evaluated. The results revealed that PD-L1 expression in infiltrated neutrophils was elevated in comparison to that of the neutrophils from peripheral blood (Fig. 1b, \( p < 0.05 \)), suggesting that tumor cells might be engaged in the PD-L1 expression in infiltrated neutrophils in OC.

Furthermore, in order to explore the association between the PD-L1 expression in infiltrated neutrophils and the IL-6 expression in peripheral blood of OC
patients, flow cytometry and RT-qPCR were conducted to detect the expression of PD-L1 in infiltrated neutrophils and IL-6 in OC patients and normal controls. The results illustrated a significant enhancement in IL-6 expression in peripheral blood of OC patients relative to normal controls, and the higher expression IL-6 in peripheral blood of OC patients was closely associated with the higher expression of PD-L1 in infiltrated neutrophils in OC (Fig. 1c, $p < 0.05$). The results inferred that the abnormal expression of IL-6 in OC patients resulted in higher expression of PD-L1 in infiltrated neutrophils in OC. In consistent, the immunohistochemical staining exhibited that neutrophils were accumulated in the OC tissues compared to the normal fallopian tube fimbriae tissues (Fig. 1d, $p < 0.05$). Following extraction of the infiltrated neutrophils in OC, we conducted flow cytometry to measure the expression of PD-L1. Combined with the immunohistochemical staining results of the upregulated PD-L1 expression in OC tissues, a positive relationship was noted between IL-6 expression in OC tissues and PD-L1 expression in infiltrated neutrophils (Fig. 1e, $p < 0.05$). Collectively, these data suggested upregulated IL-6 in OC tissues, which might secrete to the surrounding infiltrated neutrophils, thereby contributing to PD-L1 overexpression.

**OC cells secret IL-6 to activate the STAT3/PD-L1 pathway in neutrophils and repress T cell proliferation**

To discover the probable functions of IL-6 on neutrophils, IL-6 (3 μg/mL) was supplemented to neutrophils, and the expression of PD-L1 and STAT3 in addition to the extent of STAT3 phosphorylation in neutrophils were measured as well. As shown in Fig. 2a, content of PD-L1 and the extent of STAT3 phosphorylation increased ($p < 0.05$), suggesting that IL-6 could promote the extent of STAT3 phosphorylation and increase expression of PD-L1 in neutrophils. In order to verify whether IL-6 promoted expression of PD-L1 through STAT3 phosphorylation, the STAT3 phosphorylation inhibitor Stattic was applied. The results found that the content PD-L1 was not up-regulated following the addition of IL-6 ($p < 0.05$). The results showed that IL-6 stimulated the up-regulation of PD-L1 mainly by promoting phosphorylation of STAT3.

The expression of IL-6 in OC cell lines was determined. It was validated that the expression of IL-6 in SKOV3 and OVCAR3 cells were higher than that in Hey-A8 cells (Fig. 2b, $p < 0.05$). Next, cell culture solution of SKOV3, Hey-A8 and OVCAR3 was added to neutrophils and the expression of PD-L1 was evaluated.
using RT-qPCR and Western blot analysis. As shown in Fig. 2c, PD-L1 expression was enhanced in neutrophils cultured with SKOV3 or OVCAR3 cells versus that with Hey-A8 cells \((p < 0.05)\), suggesting that OC cells with higher expression of IL-6 induced PD-L1 expression in neutrophils. For further verification, IL-6 gene was knocked out in SKOV3 cells, and IL-6 was overexpressed in Hey-A8 cells. The PD-L1 expression in neutrophils was found to be decreased following culture with IL-6-knockout SKOV3 cells and Hey-A8 cells silencing IL-6 (on the left) and added with Hey-A8 cells treated with overexpressed IL-6 and SKOV3 cells silencing IL-6 (on the right). d The proliferation of CFSE-labeled T cells co-cultured with neutrophils treated with Hey-A8/IL-6 and SKOV3/si-IL-6, as examined by flow cytometry. e Western blot analysis of PD-L1 protein in neutrophils added with Hey-A8 and SKOV3 culture solution. * \(p < 0.05\) vs. controls; # \(p < 0.05\) vs. SKOV3 cell culture solution; The measurement data were summarized as mean ± standard deviation. The unpaired \(t\) test was used to compare the data between two groups. The one-way analysis of variance was adopted to compare the data among multiple groups, followed by Tukey’s post hoc test. The experiment was repeated 3 times independently.

**Fig. 2** IL-6 secreted by OC cells inhibits T cell proliferation by activating the STAT3/PD-L1 pathway in neutrophils. a Western blot analysis of PD-L1 and STAT3 proteins as well as the extent of STAT3 phosphorylation in neutrophils added with IL-6 (3 \(\mu\)g/mL) and STAT3 phosphorylation inhibitor Stattic (10 \(\mu\)M). b IL-6 expression in OC cell lines: SKOV3, Hey-A8 and OVCAR3 as determined by Western blot analysis and ELISA. c Western blot analysis of PD-L1 protein in neutrophils added with three kinds of OC cell culture medium (on the left) and added with Hey-A8 cells treated with overexpressed IL-6 and SKOV3 cells silencing IL-6 (on the right). d The proliferation of CFSE-labeled T cells co-cultured with neutrophils treated with Hey-A8/IL-6 and SKOV3/si-IL-6, as examined by flow cytometry. e Western blot analysis of PD-L1 protein in neutrophils added with Hey-A8 and SKOV3 culture solution. * \(p < 0.05\) vs. controls; # \(p < 0.05\) vs. SKOV3 cell culture solution; The measurement data were summarized as mean ± standard deviation. The unpaired \(t\) test was used to compare the data between two groups. The one-way analysis of variance was adopted to compare the data among multiple groups, followed by Tukey’s post hoc test. The experiment was repeated 3 times independently.
expression of PD-L1 in neutrophils was significantly reduced. Therefore, we may conclude that the OC cells up-regulated PD-L1 expression in neutrophils by secreting IL-6, thus inhibiting T cell proliferation.

**HOTTIP promotes IL-6 transcription by regulating transcription factor c-jun**

Next, the mechanism of IL-6 overexpression in OC was studied. The bioinformatics prediction website indicated that HOTTIP may promote IL-6 transcription by binding to c-jun (Fig. 3a). Fractionation of nuclear/cytoplasmic RNA displayed that HOTTIP was predominantly localized in nucleus (Additional file 1: Figure S1). Dual luciferase reporter gene assay revealed that both c-jun and HOTTIP could increase the activity of IL-6 promoter (Fig. 3b, $p < 0.05$). Moreover, two binding sites between c-jun and IL-6 promoter with the highest score were predicted, which were further conformed by the ChIP assay. In addition, the effect of HOTTIP on the binding of c-jun and IL-6 promoter was evaluated. The results noted that HOTTIP enhanced the binding of c-jun and IL-6 promoter (Fig. 3c, $p < 0.05$). Moreover, the interaction between c-jun and HOTTIP was verified in SKOV3 cells by RIP assay (Fig. 3d, $p < 0.05$). When HOTTIP was overexpressed in SKOV3 cells, the expression and secretion levels of IL-6 were enhanced; however, the expression and secretion levels of IL-6 were reduced following HOTTIP silencing (Fig. 3e & f, $p < 0.05$).

**Fig. 3** HOTTIP stimulates IL-6 transcription via regulating c-jun. **a** The relationship between HOTTIP, IL-6 and c-jun predicted by LNCmap. **b** The binding relationship between HOTTIP and IL-6, as well as between c-jun and IL-6 confirmed by dual luciferase reporter gene assay. **c** The binding sites between c-jun and IL-6 promoter predicted and verified by ChIP assay. **d** The interaction between c-jun and HOTTIP validated by RIP assay. **e** Western blot analysis of IL-6 protein in SKOV3 cells overexpressing or silencing HOTTIP. **f** The secretion of IL-6 in SKOV3 cells overexpressing or silencing HOTTIP as measured by ELISA. *$p < 0.05$ vs. cells treated with IL-6-prom-wt plus NC; $^\#p < 0.05$ vs. cells without treatment; $^\&p < 0.05$ vs. cells treated with IgG; The measurement data were summarized as mean ± standard deviation. The unpaired t test was used to compare the data in two groups. The one-way analysis of variance was adopted to compare the data among multiple groups, followed by Tukey’s post hoc test. The experiment was repeated 3 times independently.
suggesting HOTTIP can promote expression of IL-6. Altogether, HOTTIP could augment transcription of IL-6 via transcription factor c-jun.

**Upregulation of HOTTIP restrains T cell proliferation by promoting IL-6**

To investigate the effects of HOTTIP on the expression of PD-L1 in neutrophils and the proliferation of T cells in OC, HOTTIP expression was enforced in SKOV3 cells and then the cell culture medium was supplemented to neutrophils. Next, Western blot analysis was conducted to detect the protein expression of STAT3 and PD-L1 as well as the extent of STAT3 phosphorylation in neutrophils, which suggested that the protein expression of PD-L1 and the extent of STAT3 phosphorylation were elevated in neutrophils treated with SKOV3 cell culture medium (Fig. 4a, \( p < 0.05 \)). Next, the neutrophils added with SKOV3 cells both overexpressing HOTTIP and silencing IL-6 were co-cultured with T cells for a time, after which the T cell proliferation was assessed using flow cytometry. It was implied that the proliferation of T cells was repressed by neutrophils cultured with SKOV3 cell culture medium (Fig. 4b, \( p < 0.05 \)).

Afterwards, the mechanism of suppression of T cell proliferation by HOTTIP was explored. In OC cells treated with overexpressed HOTTIP, IL-6 was silenced. After a period of culture, the culture medium was obtained and added to the neutrophils. The protein expression of PD-L1 and STAT3 as well as the extent of STAT3 phosphorylation in neutrophils added with SKOV3 cells both overexpressing HOTTIP and silencing IL-6 (Fig. 4a, \( p > 0.05 \)), indicating that HOTTIP augmented PD-L1 expression in neutrophils mainly through IL-6. Meanwhile, the neutrophils added with SKOV3 cells both overexpressing HOTTIP and silencing IL-6 were dropped into the T cells labeled by CFSE and then co-cultured. Flow cytometry was followed to assess the proliferation ability of T cells. The results demonstrated no remarkable difference in the proliferation ability of T cells (Fig. 4b, \( p < 0.05 \)). The aforementioned results suggest that HOTTIP exerted a promoting role in neutrophils induced inhibition of T cell proliferation by promoting IL-6 in OC cells.

**Upregulation of HOTTIP increases PD-L1 expression and inhibits T cell-mediated tumor immunotherapy**

The effect of HOTTIP on T cell immunity was evaluated in vivo. At the 10th d of tumorigenesis, the neutrophils treated with the medium from the normal SKOV3 cells or SKOV3 cells overexpressing HOTTIP were intraperitoneally injected with PBS and intraperitoneally injected with CD3\(^+\) T cells into the established NOD/SCID mice with or without PD-L1 antibody. Subsequently, the body weight of the mice and the tumor volume were recorded. As revealed in Fig. 5a, no significant difference was visible in body weight upon different treatments (\( p > 0.05 \)). However, the tumor volume of mice injected with neutrophils treated with SKOV3 cells overexpressing HOTTIP was much heavier than that in neutrophils treated with normal SKOV3 cells (Fig. 5b, \( p < 0.05 \)). Then all the tumors were excised and the percentage of CD3\(^+\) T cells was measured by immunohistochemical staining. A decline of the infiltrated CD3\(^+\) T cells was observed in mice injected with neutrophils added with SKOV3 cells overexpressing HOTTIP while CD3\(^+\) T cells increased following the addition of PD-L1 antibody (Fig. 5c, \( p < 0.05 \)). Meanwhile, the IFN-\(\gamma\) expression in CD3\(^+\) T cells was detected through flow cytometry to

![Fig. 4](image-url)
determine the immune activity of T cells. The results showed that the expression of IFN-γ in CD3+ T cells was relatively lower in neutrophils added with SKOV3 cells overexpressing HOTTIP, while the results was restored by adding PD-L1 antibody (Fig. 5d, *p < 0.05*).

These results indicated that HOTTIP could promote the expression of PD-L1 in neutrophils, while suppressing T cell immunity in vivo.

**HOTTIP is highly expressed in OC tissues and positively correlates with PD-L1 in neutrophils**

To confirm the HOTTIP expression in OC tissues as well as the association with prognosis of OC patients, initially, the expression of HOTTIP was evaluated in OC tissues and normal fallopian tube fimbriae tissues using RT-qPCR. The results found that HOTTIP expression was elevated in OC tissue relative to that in normal fallopian tube fimbriae tissues (Fig. 6a, *p < 0.05*). Next, we conducted clinical follow-up to analyze the association of HOTTIP expression with prognosis of OC. It was revealed that the OS of patients with higher expression of HOTTIP (cut off value was 3.22) was far shorter than that of OC patients with lower HOTTIP expression (Fig. 6b, *p < 0.05*).

Subsequent correlation analysis revealed a positive correlation between expression of IL-6 and expression of HOTTIP (Fig. 6c, *p < 0.05*). Moreover, HOTTIP could also positively associate with the PD-L1 expression in neutrophils (Fig. 6d, *p < 0.05*). The aforementioned data supported a strong and positive correlation between HOTTIP expression and PD-L1 expression in neutrophils.

**Discussion**

Recent studies have reported that HOTTIP plays critical roles in the progression of several types of human malignancies, such as nasopharyngeal cancer [19], oral tongue squamous cell carcinoma [20] as well as breast cancer [21]. In this expanding field, much remains unknown regarding the function of HOTTIP in the development of OC and the underlying mechanism. Hence, the present study aims at demonstrating the possible mechanism by which HOTTIP was implicated in OC development. Collectively, HOTTIP could elicit the secretion of IL-6 and up-regulate PD-L1 expression in neutrophils, so as to suppress the activity of T cells, ultimately accelerating immune escape of OC cells.

Our findings indicated a robust expression of HOTTIP and IL-6 in OC tissues. Similarly, HOTTIP was up-regulated in gastric cancer (GC) cell lines [22]. Also, HOTTIP expression was notably higher in OC tissue samples in comparison to adjacent normal tissue samples [23]. In addition, OC patients presented with consistently elevated expression of IL-6 [24]. Furthermore,
IL-6 has been noted to be in a positive correlation with PD-L1, which was expressed intensely in OC cells [25], which was consistent with our findings. 

A pull-down assay confirmed putative binding sites of lncRNA DILC within the IL-6 promoter in HCC [26]. Knockdown of MALAT1 suppresses the phosphorylation of c-jun N-terminal kinase 1 (JNK1) in OC cells [27]. Consistently, our results revealed that HOTTIP could induce the release of IL-6 via c-jun. In addition, IL-6 abrogates T cell proliferation by promoting the extent of STAT3 phosphorylation through the enhancement of PD-L1 expression in neutrophils. Downregulation of IL-6/JAK/STAT3 pathway activation subsequently reduces the expression of PD-L1 in non-small cell lung cancer [15]. Cancer-associated fibroblasts induce PD-L1 in neutrophils by promoting the IL-6-STAT3 pathway that fosters immune suppression in HCC [28]. IL-6 mediates STAT3 activation in tumor and tumor progression, and IL-6 neutralization reduces STAT3 activity in vivo [29]. The aforementioned findings were in line with our findings that the addition of STAT3 inhibitor Stattic eliminated the promoting effects of IL-6 on PD-L1 expression. Notably, the IL-6/STAT3 axis can simultaneously result in the elevation of the growth of immunosuppressive cells or modify the balance of T-cell subsets [30]. The decline in IL-6 may be suggestive of the contrasting role of effector T cells and oppressive myeloid cells [31], and our study demonstrated that neutrophils treated with OC cells with the up-regulation of HOTTIP plus IL-6 silencing did not result in a significant change in T-cell proliferation.

Moreover, enhanced HOTTIP has been observed to elevate PD-L1 expression and result in increased immunosuppression, as demonstrated by suppressed T cell proliferation, IFN-γ expression as well as elevated tumor weigh in nude mice. Previous data revealed that the expression of HOTTIP and PD-L1 was simultaneously dysregulated in non-small cell lung cancer [32]. LncRNA MALAT1 promotes tumorigenesis and immune escape of diffuse large B cell lymphoma by promoting PD-L1 capable of enhancing the tumorigenesis and immune escape abilities of cancers [33]. In GC, PD-L1 restoration in infiltrated neutrophils played an inhibitory role on IFN-γ expression and T cell proliferation ability, further highlighting the significance of PD-L1 in tumor-related immunosuppression [17]. Furthermore, PD-L1 is elevated on the CD44+ subpopulation, and preferentially expressed PD-L1 can, at least partially, induce the suppression of T cell proliferation through these cells [34]. Although PD-L1 is not expressed in the majority of the OC cell lines, PD-L1 expression in response to IFN-γ is detected in OC cells in patient ascites, and PD-L1 knockdown in tumor cells can diminish the tumor-promoting effect of IFN-γ [16]. In recent years, the immunotherapy based on PD-1 impairment has profoundly improved the survival of melanoma patients in advanced stage that had
restricted therapeutic options [35]. Some partial response patients with GC exhibited diminished effector regulatory T cells and PD-1 expression through CD8+ T cells in TILs, which may reflect reduced tumor burdens [36].

Conclusion
Altogether, our study identified that HOTTIP can potentially accelerate immune escape of GC cells via enhancement of IL-6-dependent PD-L1 expression (Fig. 7). This presents a novel mechanism for the role of HOTTIP in regulating immune escape of OC cells and provides a potential target for OC therapy. However, if experimental funds are permitted, further studies will be required based on the syngeneic model that is more ideal to demonstrate anti-tumor immune response to explain the specific underlying mechanism by which HOTTIP/IL-6/PD-L1 network influences OS.

Additional file

Additional file 1: Figure S1. The localization and transfection efficiency of HOTTIP. A, Subcellular localization of HOTTIP in cells as measured using Fractionation of nuclear/cytoplasmic RNA. B and C, Silencing efficiency of HOTTIP in the nucleus as detected by RT-qPCR. (EPS 505 kb)

Abbreviations
ATCC: American Type Culture Collection; cDNA: Complementary DNA; ChIP: Chromatin immunoprecipitation; DAB: Diaminobenzidine; DMEM: Dulbecco’s modified Eagle’s medium; ECL: Enhanced chemiluminescence; ELISA: Enzyme-linked immunosorbent assay; EP: Eppendorf; F: Forward; FBS: Fetal bovine serum; GAPDH: Glyceraldehyde-3-phosphate dehydrogenase; HCC: Hepatocellular carcinoma; HOTTIP: HOXA transcript at the distal tip; HRP: Horseradish peroxidase; IFN: Interferon; IgG: Immunoglobulin G; IL-6: Interleukin-6; LncRNA: Long non-coding RNA; NC: Negative control; NCBI: National Center for Biotechnology Information; NOD/SCID: Nonobese diabetic/severe combined immunodeficient; OC: Ovarian cancer; OD: Optical density; OS: The overall survival; PAGE: Polyacrylamide gel electrophoresis; PBMCs: Peripheral blood mononuclear cells; PBS: Phosphate-buffered saline; PD-L1: Program death ligand 1; PMSF: Phenylmethanesulfonyl fluoride; PVDF: Polyvinylidene fluoride; R: Reverse; RIPA: Radioimmunoprecipitation assay; RPMI: Roswell Park Memorial Institute; SDS: Sodium dodecyl sulfate; sh: Short hairpin RNA; si: Small interfering RNA; SPF: Specific-pathogen-free; TIL: Tumor-infiltrating lymphocyte; Treg: Regulatory T; TSS: Transcription start sites; wt: Wild type

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Authors’ contributions
WW, CG, CC, BZ and YY designed the study. PJ, JS, JW, and WL collated the data, carried out data analyses and produced the initial draft of the manuscript. AS, ZS and DL contributed to drafting the manuscript. All authors have read and approved the final submitted manuscript.

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Availability of data and materials
The datasets generated/analysed during the current study are available.

Ethics approval and consent to participate
This study was conducted with the approval of the Ethics Committee of Tongji Hospital of Tongji University and The Sixth People's Hospital of Yancheng City. Written informed consents were provided and signed by all participants prior to sample collection. All animal experiments were conducted in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

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