Gestational trophoblastic neoplasia with extrauterine metastasis but lacked uterine primary lesions: a single center experience and literature review

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Abstract

Background: To investigate the clinicopathological characteristics, diagnoses, treatments, and outcomes of a special type of gestational trophoblastic neoplasia (GTN) which only has extrauterine metastases without uterine primary lesions.

Methods: The medical records and pathological sections of the patients who were pathologically diagnosed as GTN, only had extrauterine metastatic lesions but lacked uterine primary lesions, in Women's Hospital of Zhejiang University School of Medicine from February 2014 to March 2021 were collected and reviewed.

Results: Thirteen patients with pathologically confirmed GTN presenting with extrauterine metastases from a missing primary site were included in the past 7 years. The median age was 31.2 years old. 76.9% of patients had a non-hydatidiform pregnancy last time. The intervals between the antecedent pregnancy were > 12 months in 61.5% of patients. Pretreatment serum human chorionic gonadotropin(hCG) levels ranged from 118.7 to 807,270 IU/L. Six patients were misdiagnosed as ectopic pregnancy at initial diagnosis, and 4 as primary tumors at metastatic sites. All of them were diagnosed definitely by surgical pathology including 8 choriocarcinomas (CC), 4 epithelioid trophoblastic tumors (ETTs), and 1 mixed GTN (CC mixed with ETT). All patients achieved complete remission (CR) after treatments. Three patients relapsed; no patient died by the end of follow-up.

Conclusion: GTN presenting with extrauterine metastases from a missing primary site is easily misdiagnosed. Detection of serum hCG in these patients can reduce misdiagnosis. Chemotherapy combined with individualized surgery should be considered for these special GTN patients. Immune checkpoint inhibitors might be potential remedial measures for refractory and recurrent patients.

Keywords: Gestational trophoblastic neoplasia, Neoplasm metastasis, Choriocarcinoma, Diagnosis, Therapy

Introduction

Gestational trophoblastic neoplasia (GTN) is a group of tumors derived from abnormal proliferative placental trophoblastic cells. According to the 2010 World Health Organization (WHO) classification [1], GTN is classified histologically into a series of pregnancy-related malignancies, including, gestational choriocarcinomas (CC)
and placental site trophoblastic tumor (PSTT), and epithelioid trophoblastic tumor (ETT) [2, 3].

GTN can be divided into non-metastatic GTN and metastatic GTN. Non-metastatic GTN refers to lesions confined to the uterus. Metastatic GTN refers to lesions occurring outside the uterus, typically hematomogenous spreading. The lung is the most common site of metastasis, and other metastatic sites include the vagina, tubes, ovaries, liver, spleen, kidneys, bowels, brain, etc. [4] Distinct metastases such as liver and brain are more likely to have a poor prognosis [5]. The majority of metastatic GTN presented as a primary uterine lesion complicated with extraterine metastases. However, a few metastatic GTN only presented with extraterine metastases and the primary lesion was missed. Such metastatic GTN is rare in clinical practice, and most cases are reported as a single case. The first symptoms are usually different due to different metastatic sites and lack of clinical manifestations related to GTN, making diagnosis difficult and easy to misdiagnose.

Thus, it is necessary to conduct retrospective studies aimed at exploring the clinico-pathological data, diagnosis, treatment, and prognosis in patients of this kind of GTN, to provide a reference for diagnosis and treatment of this rare entity.

Materials and methods

Patients and data collection

All patients, who were pathologically diagnosed as GTN presenting with extraterine metastases from a missing primary site in Women’s Hospital of Zhejiang University School of Medicine from February 2014 to March 2021, were collected using computerized databases from the Departments of Gynecologic Oncology and Pathology. Inclusion criteria: 1) New diagnosed GTN patients; 2) Extraterine metastases were diagnosed with GTN by histopathology and immunohistochemistry; 3) No primary uterine lesions were found in the evaluation of gynecological examination combined with transvaginal ultrasound of uterine adnexa, abdominal and pelvic Computed Tomography (CT), with or without pelvic Magnetic Resonance Imaging (MRI). Exclusion criteria: The patients had a history of other malignant tumors. The medical records were reviewed and the following data were collected, including age, symptoms, previous pregnancy, pre- and post-treatment serum human chorionic gonadotropin (hCG) levels, metastasis site, primary clinical diagnosis, pathological diagnosis, stage and prognosis score, treatments, recurrence, and prognosis.

Federation International of Gynecology and Obstetrics (FIGO) 2000 stage was used for staging, and World Health Organization (WHO) prognostic index score standard (2014) was used for prognostic score [3].

The study was approved by the Ethics Committee of the Women’s Hospital of Zhejiang University School of Medicine. This study was a retrospective analysis of clinical data, unrelated to human bioethics. Informed consents were obtained for all follow-up contents. the study was performed under the principles of the 2013 Declaration of Helsinki [6]. All methods were performed under the relevant guidelines and regulations.

Immunohistochemistry

Programmed death ligand 1 (PD-L1) expression was detected by Immunohistochemistry in all these patients. Mouse anti-human PD-L1 antibody (ab210931; Abcam, Shanghai, China) diluted at 1:100 was used for detection. Following protocols, the two-step EnVision immunostaining procedure (Dako, Carpentry, CA) was performed. The normal human placenta was regarded as the positive control, and PD-L1 negative cervical squamous cell carcinoma was the negative control. Membrane staining was considered positive. The results were confirmed by pathologists and evaluated by tumor proportion score (TPS) and combined positive score (CPS) with reference to our previous study [7].

Follow-up and outcome

All patients were followed via telephone interviews or at clinics. Complete Remission (CR) was defined as a continuous normalization of serum hCG levels for at least 4 weeks after chemotherapy, while resistance was defined as an increase of hCG levels >10% or stabilization of ±10% within 2 weeks after two courses of chemotherapy. Relapse was defined as hCG levels rising again after three months of CR in the absence of a normal pregnancy. Mixed GTN was defined as the coexistence of choriocarcinoma and/or PSTT and/or ETT components. Undetermined GTN was defined as group serum β-hCG levels elevated in the absence of the pregnancy less than 3 months after completed treatment [8-11].

Statistical analysis

Statistical analysis was performed using SPSS26.0 for Windows (IBM Corporation, Armonk, NY, USA). Continuous data were described as mean ± SD (standard deviation) or median, and categorical data as frequency and percentage. Fisher’s exact test was used to check qualitative variables, and \( P < 0.05 \) was considered statistically significant.

Results

Clinical characteristics

A total of 13 patients with pathologically confirmed GTN presenting with extraterine metastases from a missing primary site were included, and their clinico-pathological
characteristics, diagnosis, treatments, and outcomes were summarized in Table 1. The mean age at diagnosis was 30.5 ± 5.8 years (19–41 years). Imageology ± laparoscopy confirmed that all patients had no uterine lesions throughout the course of the disease. Antecedent pregnancies were molar pregnancy in 3 patients (23.1%), abortion in 8 patients (61.5%), term pregnancy in 1 (7.7%) patient, coexisting pregnancy in 1 (7.7%) patient. In 70% of molar and abortion cases, the nature of the pregnancy substance was confirmed. After an abortion, the serum hCG level was followed up once a week until the value was normal twice in a row. Most cases were reviewed regularly after abortion, except Cases 7, 8, and 11.

The interval between the antecedent pregnancy and treatment was ≤ 12 months for 5 patients including 1 patient during pregnancy and > 12 months in the remaining 8 patients. Pretreatment serum hCG levels ranged from 118.7 to 807,270 IU/L. Notably, 10 of 13 (76.9%) patients had levels of hCG ≤ 2,500 IU/L. Metastases in one or more organs were the most common symptom which was reported in all of the 13 patients including 7 (53.8%) patients presented with pulmonary nodules, 2 (15.4%) patients presented with an adnexal mass, and another 4 (30.8%) patients presented with masses in multiple organs including kidney, liver, brain, pelvic cavity, and ligaments, etc. In addition, all metastases presented as localized hemorrhagic necrosis as shown in Fig. 1. Abnormal vaginal bleeding was the second common symptom which was reported in 8 (61.5%) patients. Other presentations included abdominal pain and amenorrhea.

(A) The chest CT image of case 4 showed a lobulated mass (black arrowhead) about 2.6×3.1 cm in size in the right upper lung. (B) Abdominal contrast-enhanced CT image of case 11 showed an irregular mass (white arrowhead) near the left ovary, about 4.3×2.7×2.6 cm in size, with uneven internal enhancement on the contrast-enhanced scan. (C) (D) (E) Abdominal enhanced CT image of the case 10 suggested multiple lesions (white arrowhead) in the liver, left kidney, and left quadratus lumbago muscle, which were considered as metastatic tumors. (F) The cranial enhanced MRI image in case 10 showed multiple nodules in the brain parenchyma (the largest one is shown here, about 1.8 cm in length), and metastasis was considered.

**Diagnosis and treatments**

Based on the final operative histopathology results, the 13 patients have confirmed their diagnosis as follows: 8 CC (61.5%), 4 ETTs (30.8%), and 1 mixed GTN (CC mixed with ETT) (7.7%). However, the initial diagnosis of all the 13 patients was incorrect or unclear. Six patients were misdiagnosed as ectopic pregnancy, 2 as the ovarian tumor, 1 as lung cancer, 1 as breast and lung cancer, and the remaining 3 suspected as GTN. During the definitive diagnosis process, 9 patients received multiple surgeries which involved dilation and curettage (D&C) and video-assisted thoracoscopic surgery (VATS) in 1 patient (case 2), D&C combined with laparoscopy and VATS in 3 patients (case 3, 4, 7), percutaneous needle aspiration biopsies of lungs (PTNB) and VATS in 2 patients (case 5, 6), VATS and segmental mastectomy (SM) in 1 patient (case 10), laparoscopy and laparoscopic mass resection (LMR) in 1 patient (case 11), D&C and VATS combined with laparoscopic unilateral adnexectomy (LUA) and laparoscopic mass resection (LMR) in 1 patient (case 12). Other 4 patients were confirmed for GTN by single surgery including case 1 patient by VATS, cases 8 and 9 by laparoscopic adnexal lesions resection, and case 13 by cytoreductive during a cesarean. For related operations other than obstetrics and Gynecology, we invited doctors from all relevant departments to assist us in completing them.

After surgical confirmation and removal of the GTN lesion, all 13 patients underwent further chemotherapy. According to WHO 2000 risk score standard, 2 low-risk CC patients initially received Methotrexate (MTX) regimen, 6 high-risk CC patients initially received TP (Taxol and Carboplatin) or EMA-CO (Etoposide, Methotrexate, Actinomycin-D/Cyclophosphamide, and Vincristine) regimen respectively, 1 ultra-high-risk CC patient initially received EP-EMA (Etoposide, Methotrexate, Actinomycin-D/Etoposide, and Cisplatin) regimen, 4 ETT patients have initially received EP-EMA or EMA-CO regimen. Case 2 presented severe oral ulcers with infection. Bone marrow suppression was found in other patients during chemotherapy, and abnormal liver function was also found in patients 4, 5, 8, 9, and 11.

**Recurrence and prognosis**

All patients had a CR after treatment. The median duration of follow-up was 24.5 months (5–85.2 months), and 3 (case 6, 8, 10) patients had a recurrence within 2 to 5 months, no patients died. Details as shown in Table 2. The case 6 had no vaginal bleeding, cough, abdominal pain, and other discomforts. The patient experienced CR again after a VATS operation by thoracic surgeons to remove the lung lesion and 3 cycles of EMA-CO as consolidation chemotherapy. The case 8 also had no complaints of discomfort. The patient was reexamined with elevated HCG and left ovarian abnormality. She experienced CR again after laparoscopic left adnexectomy and 5 cycles of TP regimens. In case 10, hCG increased again and epilepsy was found after hCG was negative for more than three months. Cranial MRI enhanced scan showed hemorrhage of brain metastasis. She was transferred to a general hospital for...
### Table 1: The clinical characteristics, treatment and outcomes of 13 patients with metastatic GTN without primary lesions

| Patients No | Age  | G/P  | Symptoms       | Signs         | Antecedent pregnancy | Interval (months) | Pretreatment hCG(IU/L) | Initial disposition | Metastasis site                |
|-------------|------|------|----------------|---------------|----------------------|------------------|------------------------|----------------------|------------------------|
| 1           | 28   | 3/1  | Vaginal bleeding | Lung mass     | Mole                 | 17               | 2021                   | GTN                  | Right lung             |
| 2           | 31   | 1/0  | Vaginal bleeding | Lung mass     | Mole                 | 3                | 121                    | GTN                  | Right lung             |
| 3           | 23   | 2/0  | None            | Lung mass     | Abortion             | 11               | 277                    | Ectopic pregnancy    | Right lung             |
| 4           | 32   | 4/2  | Vaginal bleeding, cough | None | Term | 3 | 194 | Lung cancer | Right lung |
| 5           | 30   | 1/0  | Vaginal bleeding | Lung mass     | Mole                 | 14               | 144                    | GTN                  | Right lung             |
| 6           | 27   | 2/0  | Vaginal bleeding | Lung mass     | Abortion             | 36               | 967                    | Ectopic pregnancy    | Right lung             |
| 7           | 36   | 3/1  | Vaginal bleeding | None          | Abortion             | 60               | 1100                   | Ectopic pregnancy    | Left lung              |
| 8           | 31   | 1/0  | Vaginal bleeding | Ovarian mass  | Abortion             | 108              | 612                    | Ectopic pregnancy    | Left ovary             |
| 9           | 19   | 0/0  | Vaginal bleeding | Ovarian mass  | Abortion             | N/A              | 104,400                | Ectopic pregnancy    | Right ovary             |
| 10          | 37   | 2/0  | None            | Lung and breast mass | Abortion | 60 | 21,136 | Lung cancer, breast cancer |
| 11          | 41   | 5/2  | None            | Adnexal mass   | Abortion             | 5                | >2000                   | Ectopic pregnancy    | Right pelvic funnel ligament and meso-colon |
| 12          | 36   | 5/1  | Abdominal pain  | Adnexal mass   | Abortion             | 34               | 119                    | Ovary tumor          | Masses on the right ovary, omentum, and abdominal wall |
| 13          | 26   | 0/0  | None            | Ovarian and liver masses | Pregnancy (29 weeks) | 7 | 807,270 | Ovary tumor | The right lung, pelvic abdomen, liver |

| Patients No | Number of metastases | Initial treatment | Definitive diagnostic method | Pathological diagnosis | FIGO stage | Prognostic score | Chemotherapy | The number of consolidation chemotherapy | Relapse |
|-------------|---------------------|-------------------|------------------------------|------------------------|------------|-----------------|-------------|------------------------------------------|---------|
| 1           | 1                   | 5 cycles of TP    | VATS                         | ETT                    | III        | /               | 2 cycles of EP-EMA + 5 cycles of EP | 5 | No |
| 2           | 1                   | D&C               | VATS                         | CC                     | III        | 5               | 6 cycles of MTX | N/A | No |
| 3           | 1                   | D&C + Laparoscopy | VATS                         | ETT                    | III        | /               | 3 cycles of EP-EMA | 3 | No |
| 4           | 1                   | D&C + Laparoscopy | VATS                         | ETT                    | III        | /               | 3 cycles of EP-EMA | 3 | No |
| 5           | 1                   | PTNB              | VATS                         | ETT                    | III        | /               | 3 cycles of EMA-CO | 3 | No |
| 6           | 1                   | PTNB              | VATS                         | CC                     | III        | 5               | 6 cycles of MTX | 3 | Yes |
| 7           | 1                   | D&C + Laparoscopy + 1 cycle of MTX | VATS | CC | III | 7 | 3 cycles of TP | 3 | No |
| 8           | 1                   | Laparoscopy + 1 cycle of MTX | LOCR | CC | III | 7 | 3 cycles of EMA-CO + 3 cycles of EP-EMA + 4 cycles of FAEV | 5 | Yes |
Table 1 (continued)

| Patients No | Number of metastases | Initial treatment | Definitive diagnostic method | Pathological diagnosis | FIGO stage | Prognostic score | Chemotherapy | The number of consolidation chemotherapy | Relapse |
|-------------|----------------------|------------------|-------------------------------|------------------------|------------|-----------------|-------------|-------------------------------------------|---------|
| 9           | 1                    | Laparoscopy      | LOCR                          | CC                     | II         | >7              | 7 cycles of EMA-CO  | 3                     | No      |
| 10          | Multiple             | VATS + SM        | VATS + SM                     | CC                     | IV         | 17              | 2 cycles of EMA-CO + 8 cycles of EP-EMA + 2 cycles of MTX intrathecal injection | 5                     | Yes     |
| 11          | Multiple             | Laparoscopy + 2 cycles of MTX | LMR                          | CC                     | II         | 9               | 3 cycles of EMA-CO + 2 cycles of EMA | 3                     | No      |
| 12          | Multiple             | D&C + Laparoscopy | LUA + LMR                    | CC/ETT                 | IV         | 12              | 4 cycles of EP-EMA | 3                     | No      |
| 13          | Multiple             | Cesarean         | Cytoreductive surgery         | CC                     | IV         | 7               | 5 cycles of EMA-CO | N/A                    | No      |

G/P, Gravida/para; hCG, Human chorionic gonadotropin; IU/L, International units per liter; FIGO, International Federation of Obstetrics and Gynecology; GTN, Gestational trophoblastic neoplasm; D&C, Dilation and curettage; PTNB, Percutaneous needle biopsy of the chest; VATS, Video-assisted thoracoscopic surgery; SM, Segmental mastectomy; LOCR, Laparoscopic ovarian cyst resection; LUA, Laparoscopic unilateral adnexectomy; LMR, Laparoscopic mass resection; ETT, Epithelioid trophoblastic tumor; CC, Choriocarcinoma; TP, Paclitaxel; Cisplatin; MTX, Methotrexate; EP, Etoposide; Methotrexate; Actinomycin-D; Etoposide, Cisplatin; EP, Etoposide; Cisplatin; EMA, Etoposide; Methotrexate; Actinomycin-D; Cyclophosphamide; Vincristine; FAEV, Floxuridine; Actinomycin-D. None; no positive symptoms and signs; N/A, not available.

(The patients returned to the local hospital for treatment after hCG values returned to normal levels, and the regimen and cycles of consolidation chemotherapy are uncertain. Follow-up showed that hCG was reduced to normal.) /, ETT does not apply to the FIGO scoring system and is not rated.
gamma-knife, and postoperative pathology considered CC with hemorrhage and necrosis of brain metastases. Then she was given weekly treatment with Albumin Paclitaxel combined with PD-1 immunotherapy (Tirelizumab) for 5 cycles and achieved CR again, then Tirelizumab maintenance therapy. Well controlled by follow-up.

Only one patient (case 9) had reproductive requirements who had delivered 2 healthy boys in 2017 and 2019 respectively. No recurrence of the GTN was observed.

Univariate analysis showed that age, antecedent pregnancy, interval months from index pregnancy, pretreatment serum hCG level, site and number of metastases, prognosis score, and pathological type were not associated with recurrence (supplyment1).
PD-L1 expression

All available GTN tissue samples from patients were tested for PD-L1 expression, showing strong positive staining. Positive immunohistochemistry staining images were shown in Fig. 2. (A) (B) Pelvic mass tissue, choriocarcinoma, Immunohistochemistry showed diffuse membrane staining, PD-L1 expression > 50% (magnification, 100 × and 200x). (C) Lung tissue, PD-L1 positive, TPS > 50%, CPS > 50% (magnification, 400x). (D) Breast tissue, PD-L1 positive, TPS > 50%, CPS > 50% (magnification, 400x).

Discussion

The 13 cases accounted for 0.04% of the total cases of gynecological oncology in our hospital, and 6.5% of the total cases of gestational trophoblastic tumors from February 2014 to March 2021. Metastatic GTN without primary lesion is rare in clinical practice, and mostly
described in case reports, little is known about its clinical characteristics, which is easy to be misdiagnosed and mistreated. At present, the occurrence of this special GTN has been speculated as the primary uterine lesion is too small or grows slowly after tumor metastasis, which cannot be detected by existing imaging techniques, or the primary lesion is cleared by the autoimmune system after metastasis. For this purpose, we performed a retrospective study and literature review to better understand these special GTNs.

Aside from the 13 cases in our retrospective study, other 12 cases of extrauterine metastatic GTN without primary lesion were obtained by the literature reviewed [12–20]. The characteristics, initial diagnoses, treatments, and outcomes of the 12 patients in the literature were summarized in Table 3. The mean age at diagnosis was 39.4 years (25–56 years). Antecedent pregnancies were completed to term in 3 (25%) patients, ended in mole pregnancy in 3 (25%) patients, ended in abortion in 1 (8.3%) patient, and unknown in other 5 (41.7%) patients. The interval between the antecedent pregnancy and treatment was from 2 weeks to 16 years. Pretreatment serum hCG levels ranged from 6.9 to 610,000 IU/L. However, only 2 of 12 patients had high levels of hCG > 2,500 IU/L. Hemorrhagic necrotic metastases in one or more organs were the most common symptoms, usually in the lung (83.3%). Other symptoms were atypical and varied with different sites of metastasis, such as vaginal bleeding, abdominal pain, hemoptysis, etc. The vast majority of patients were misdiagnosed at the initial diagnosis, 1 patient was misdiagnosed as ectopic pregnancy and 8 patients were misdiagnosed as a primary tumor at the metastatic site. Most patients were diagnosed surgically, 5 patients were diagnosed by thoracoscopy, 1 by vaginal mass resection, 1 by laminectomy, 1 by external iliac lymph node biopsy, and 1 by total hysterectomy. Among the 12 patients, 8 patients were histopathological confirmed ETT, 4 were CC. Among the 8 patients whose data were collected, 6 were alive without disease, 1 relapsed after remission and yet was alive after salvage treatment, one died. Most patients received EMA-CO or EP-EMA regimen; 1 patient received an immune checkpoint inhibitor (pembrolizumab) for treatment.

Based on our cohort and cases reported in the literature, it showed that this special GTN was a rare particular type of GTN which different from ordinary GTN in clinical presentation, histopathology, diagnosis, and treatment. This particular type of GTN had the following clinical presentation: 1) Most antecedent pregnancies were non-hydadiform mole; 2) It always presented as hemorrhagic necrotic metastases in one or more organs rather than abnormal vaginal bleeding; 3) Serum hCG level can be slightly elevated in the majority patients and exceed 2,500 IU/L in a very few patients. Due to the lack of typical clinical presentations, this particular type of GTN was easily misdiagnosed as the primary tumor at the metastatic site at the first diagnosis, and surgical diagnosis is often required. Thus, for women with the above manifestations, serum hCG should be tested to avoid misdiagnosis.

ETT and CC were the main pathological types in this special GTN which were confirmed by surgical histology. Patients with single lung metastases and low hCG levels were more likely to have ETT, accounting for 20% of the 25 patients. As we know, the ordinary GTN is a solid tumor diagnosed and treated based on clinical evidence without a histological diagnosis. The treatment principle is chemotherapy, supplemented by surgery [21]. Chemotherapy regimens depend on stage and score, with low-risk patients (GTN < 7) receiving single-agent chemotherapy first, high-risk patients (GTN ≥ 7) receiving combination regimens, and surgery being used only for removal of uncontrolled bleeding or drug-resistant lesions. However, surgery was of great value in this particular type of GTN patient, which could not only confirm the diagnosis but also play an important role in the treatment. In this group, 10 patients were cured by surgery, of which 30% were drug-resistant lesions removed and 70% were ETT whose hCG was reduced to normal range after confirmed surgery. EMA-CO and EP-EMA were the first choices for chemotherapy. 41.7% (5/12) of the cases with available data in the literature received remission after treatment with EMA-CO or EP-EMA, and 3 of the 13 (23.1%) patients in this group were resistant to both MTX single-agent chemotherapy and TP for the first time, but still achieved complete remission after treatment with EMA-CO or EP-EMA. Thus, we suggested that obtaining pathology was an important way to guide the treatment, and EMA-CO or EP-EMA chemotherapy combined with individualized surgical treatment was the first choice for this special GTN.

Of the 12 patients reported in the literature, 1 patient recurred and 1 patient died; of the 13 patients in our cohort, 3 patients relapsed and none died. As the literature reported that the recurrence rate of conventional low-risk and high-risk GTN was 1.6–3.1% and 6.9% respectively [22–24]. In our study, the recurrence rate of this special GTN was 16% which was slightly higher than that of ordinary GTN, which may be related to the prolonged treatment time, the metastasis of important organs, and the insensitive to chemotherapy for ETT. Fortunately, the overall prognosis of these patients was generally good, most patients with recurrence could still achieve CR after surgery and chemotherapy.

Two patients (case 10 and Bell etc. [17]) were finally treated with PD-1 antibody which could control the
Table 3 The clinical data of patients with metastatic GTN without primary lesions reported in the literature

| Author       | Age | G/P | Symptoms                              | Signs               | Antecedent pregnancy | Intervals times | Pretreatment hCG(IU/L) | Initial diagnosis | Metastasis site   | Number of metastases | Diagnostic method | Pathological diagnosis | FIGO stage and Prognostic score | Subsequent treatment | Outcomes |
|--------------|-----|-----|---------------------------------------|--------------------|----------------------|-------------------|-----------------------|-------------------|-------------------|---------------------|-------------------|-----------------------|-------------------------------|---------------------|----------|
| Lei et al    | 49  | 1/1 | None                                  | Lung mass          | Mole                 | N/A               | N/A                   | N/A               | Lung              | 1                   | VATS              | ETT                   | /                            | N/A                 | NED      |
| Ru et al     | 33  | 3/1 | None                                  | Vaginal mass       | Abortion             | 6 months          | 70                    | N/A               | Vaginal wall, lung| N/A                | Mass excision | ETT                   | /                            | 2 cycles of EMA-CO     | NED      |
| Chohan et al | 36  | 4/4 | Severe low back pain                  | Term               | 2 weeks              | 16                | Infusing tumor        | Spine, liver, lung| Multiple          | Laminctomy          | ETT               | /                    |                               | EMA-CO              | DOD      |
| Uraie et al  | 38  | 5/4 | Poor physical condition              | Lung mass          | Term                 | N/A               | 80.1                  | CC                | Lung              | Multiple            | VATS              | ETT                   | /                            | 6 cycles of EMA-CO, thoracoscopy | Relapse  |
| Hamazaki et al | 32 | 3/1 | None                                  | Lung mass          | Mole                 | 5 years           | N/A                   | Lung cancer       | Lung              | 1                   | VATS              | ETT                   | /                            | N/A                 | NED      |
| Hamazaki et al | 42 | 2/2 | Hemoptysis and cough                 | Small nodules in lung and liver | N/A | N/A | N/A | Lung cancer | Lung, liver | Multiple | VATS | ETT | / | Chemotherapy | NED |
| Bell et al   | 47  | 2/2 | Recurrent respiratory infections     | Pulmonary emboli, pulmonary nodules, and infarcts | N/A | 12 years | 9 | N/A | Lung, the infrarilar mass, recto abdominis muscles, a right external iliac lymph node | Multiple | External iliac lymph node biopsy | ETT | / | 7 cycles of EP-EMA, 29 times of pembrolizumab | NED |
| Davis et al  | 49  | N/A | None                                  | Term               | 16 years             | 69                | N/A                   | Intra-abdominal   | N/A               | TAH, BSO, tumor debulking | ETT | / | EP-EMA, prior to surgery | NED |
| Whaley et al | 56  | N/A | Metastatic cervical carcinoma with liver hemorrhage | None               | N/A                 | N/A | 5365.1 | N/A | Liver | N/A | CC | IV | N/A | N/A |
| Whaley et al | 40  | N/A | Splenic injury with liver hemorrhage | None               | N/A                 | 13.4 | N/A | N/A | Liver | N/A | CC | IV | N/A | N/A |
| Whaley et al | 25  | N/A | Retroperitoneal bleed                 | None               | N/A                 | 610,000 | N/A | Lung | N/A | N/A | CC | III | N/A | N/A |
| Maruoka et al | 26 | N/A | High level of hCG, a pulmonary nodule | Lung mass          | Mole                 | 6 years           | 233.8                  | Eclectic pregnancy | Lung | 1       | VATS | CC | III,6 | N/A | N/A |

N/A, Not available; None, no positive symptoms and signs; G/P, Gravida/para; hCG, Human chorionic gonadotropin; IU/L, International units per liter; FIGO, International Federation of Obstetrics and Gynecology; CC, choriocarcinoma; ETT, epithelioid trophoblastic tumor; VATS, video-assisted thoracoscopic surgery; TAH, total abdominal hysterectomy; BSO, bilateral salpingo-oophorectomy; CC, choriocarcinoma; /, ETT does not apply to the FIGO scoring system and is not rated. EMA-CO, Etoposide, Methotrexate, Actinomycin-D/Cyclophosphamide; Vincristine; EP-EMA, Etoposide, Methotrexate, Actinomycin-D/Etoposide, Cisplatin. NED, no evidence of disease; DOD, dead of disease.
disease well. The refractory and recurrent cases achieved CR with chemotherapy combined with immune checkpoint inhibitors (pembrolizumab) based on the high expression of PD-1/PD-L1 [6, 25]. Studies showed that about 30%-40% of high-risk patients responded poorly to first-line therapy or relapsed after remission. Remedial chemotherapy with Etoposide and Platinum drugs, which can be combined with surgical excision of drug-resistant lesions, has a good cure rate. The chemotherapy regimen included TP/TE (Paclitaxel and Cisplatin interchanged weekly with Paclitaxel and Etoposide), BEP (Bleomycin, Etoposide; HDC: High dose chemotherapy.

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Conclusion
In conclusion, improving the understanding of this special GTN, avoiding misdiagnosis and mistreatment, chemotherapy combined with individualized surgical treatment was the key to improving the prognosis of patients. Immune checkpoint inhibitors might be potential remedial measures for refractory and recurrent patients.

Abbreviations
GTN: Gestational trophoblastic neoplasia; WHO: World Health Organization; IHH: Invasive hydatidiform mole; CC: Chorioncarcinomas; PSTT: Placental site trophoblastic tumor; ETT: Epithelioid trophoblastic tumor; CT: Computed Tomography; MRI: Magnetic Resonance Imaging; hCG: Serum human chorionic gonadotropin; FIGO: Federation International of Gynecology and Obstetrics; PD-L1: Programmed death-ligand 1; TPS: Tumor proportion score; CPS: Combined positive score; CR: Complete remission; D&C: Dilatation and curettage; VATS: Video-assisted thoracoscopic surgery; PTNBL: Percutaneous needle aspiration biopsies of lungs; SM: Segmental mastectomy; LMR: Laparoscopic mass resection; LUA: Laparoscopic unilateral adnexitomy; MTX: Methotrexate; TP: Taxol, Carboplatin; EMA: CO Etoposide, Methotrexate, Actinomycin-D/Cyclophosphamide, Vincristine; EP-EMA: Etoposide, Methotrexate, Actinomycin-D/Etoposide, Cisplatin; TP/TE: Paclitaxel and Cisplatin interchanged weekly with Paclitaxel and Etoposide; BEP: Bleomycin, Etoposide, Cisplatin; VIP: Etoposide, Ifosfamide, Cisplatin; ICE: Ifosfamide, Carboplatin, Etoposide; HDC: High dose chemotherapy.

Supplementary Information
The online version contains supplementary material available at https://doi.org/10.1186/s12885-022-09620-2.

Additional file 1. Supplement 1: The univariate correlation analysis of metastatic GTN without primary tumor and the outcome of recurrence.

Acknowledgements
The authors are thankful to personnel of the medical record library of Women’s Hospital of Zhejiang University School of Medicine for data acquisition.

Authors’ contributions
YS, XX and WL contributed to the conception of this study. JL contributed to data acquisition • data analysis and presentation • writing the initial draft. YW contributed to data acquisition and data presentation. BL contributed to pathological images and pathological analysis. YS contributed to the design of methodology • manuscript editing and review.

Funding
Not applicable.

Availability of data and materials
All data generated or analyzed during this study are included in this published article (and supplementary information files).

Declarations
Ethics approval and consent to participate
The study was approved by the Ethics Committee of the Women’s Hospital of Zhejiang University School of Medicine. Informed consents were obtained before data collection.

Consent for publication
Not applicable.

Competing interests
The authors declare that the research was conducted in the absence of any potential conflict of interest.

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Received: 26 December 2021 Accepted: 28 April 2022

Published online: 06 May 2022

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Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

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