Objectives: To assess the outcomes and toxic effects of 5-day actinomycin D (Act-D) salvage therapy and to explore the predictors of Act-D resistance in patients with low-risk gestational trophoblastic neoplasia (GTN) who failed 5-day methotrexate (MTX) chemotherapy.

Methods: This retrospective study analyzed patients with low-risk GTN administered Act-D salvage therapy after failing MTX chemotherapy at Women's Hospital, School of Medicine Zhejiang University between January 2000 and December 2015. The clinical parameters of these patients were collected and analyzed.

Results: The final analysis included 89 cases. Of these, 73 cases (82.02%) responded to salvage Act-D. The remaining 16 resistant cases were switched to etoposide, MTX, Act-D/cyclophosphamide, and vincristine chemotherapy and achieved complete remission. Serum human chorionic gonadotrophin levels before Act-D salvage therapy (hCG Act-D) in the Act-D-resistant cases were significantly higher than those in the Act-D responders (median 605 vs. 103 IU/L, p=0.009). However, the range of hCG Act-D values in Act-D responders was wider than that in Act-D-resistant cases (5.76–16,664 IU/L vs. 11.43–6,732 IU/L). Thus, assigning a general cut-off value was difficult considering the individual setting. Except for 2 cases requiring other salvage regimens due to Act-D toxicity, 97.80% of cases (89/91) tolerated the toxicity. During at least 1-year follow-up, the survival rate was 100.00% and no case developed recurrence.

Conclusion: Based on the good therapeutic effect and tolerable toxicity, we recommend Act-D salvage therapy for all patients with low-risk GTN who fail primary MTX chemotherapy. The higher serum hCG levels before Act-D salvage therapy may be associated with resistance to this treatment.

Keywords: Gestational Trophoblastic Neoplasia; Methotrexate; Salvage Therapy; Actinomycin D
INTRODUCTION

Gestational trophoblastic neoplasia (GTN) is a rare malignancy that can be cured by cytotoxic chemotherapy. The International Federation of Gynecology and Obstetrics (FIGO) system recommends single-agent chemotherapy for low-risk GTN patients (FIGO prognostic score ≤6) and multi-agent chemotherapy for high-risk GTN patients (FIGO prognostic score ≥7). Methotrexate (MTX) as a first-line treatment remains the most commonly prescribed drug for low-risk GTN patients [1-3]. However, 4.00%–46.73% of patients develop resistance to MTX [1,3,4-9], while 1.41%–3.69% of patients experience intolerable toxicity to MTX [1,2,10,11], requiring salvage chemotherapy or even surgery [12].

Various salvage regimens for cases of primary MTX failure have been described in the literature, including single-agent pulsed actinomycin D (Act-D) [13,14]; 5-day Act-D [1,3,4,7,15,16]; etoposide and Act-D (EA) [17,18]; etoposide, MTX, Act-D/cyclophosphamide, and vincristine (EMA-CO) [1,3,15]; and bleomycin, etoposide and cisplatin (BEP) [19]. Among these regimens, the use of etoposide in multi-agent chemotherapy (e.g., EMA-CO, EA, and BEP) increases the risk of secondary tumors such as leukemia, melanoma, breast cancer, and colon cancer [20]. EMA-CO treatment also increases the risk of early menopause [21]. Thus, the Act-D regimen is currently preferred due to its simplicity, limited short- and long-term toxicity, and low risk of carcinogenicity [22-24].

However, 5.21%–25.00% of MTX-resistant patients develop Act-D resistance [3,7,14-16] and switch to multi-agent chemotherapy. Some researchers recommended that patients with high levels of serum human chorionic gonadotropin (hCG) at the time of MTX failure be treated directly with multi-agent chemotherapy to reduce drug resistance and shorten the chemotherapy cycle. However, the definition of high serum hCG level is unclear. In the current literature, high serum hCG levels range from 100–1,000 IU/L [3,14-16]. Identification of an accurate threshold is necessary to determine whether to administer a single-agent Act-D or multiple-agent regimen.

Our hospital is a tertiary medical institute and one of the GTN centers in China. We have previously reported on the treatment and management of GTN [4,25-30], including the reasons for treatment failure in low-risk GTN patients administered initial single-agent MTX chemotherapy [4,25,26]. The present study describes our 16-year experience in treating low-risk GTN patients who failed MTX chemotherapy. Our purpose was to assess the outcomes and side effects of the 5-day Act-D salvage therapy after MTX failure and to explore predictors of Act-D resistance.

MATERIALS AND METHODS

This retrospective cohort study included low-risk GTN patients who failed primary MTX chemotherapy at the Women’s Hospital, School of Medicine Zhejiang University between January 2000 and December 2015. This study was approved by the Ethics Committee of Women’s Hospital, School of Medicine Zhejiang University (09/27/2018, No.20180106). All patients included in this study were anonymized.

We searched the clinical records in the Medical Record Review System. Our recruitment criteria were: 1) FIGO prognostic score ≤6. 2) First-line regimen of MTX (0.4 mg/kg per day,
maximum 25 mg) injected intramuscularly or intravenously injection for 5 days every other week. 3) Patients switched to 5-day Act-D salvage regimen (10 μg/kg per day, intravenous injection, for 5 days every 2 weeks) for MTX resistance or intolerable toxicity. 4) Patients completing the entire treatment in our hospital. FIGO prognostic score and stage were assessed according to the FIGO 2000 statement [26].

The following clinical information was recorded for each subject: 1) Clinical characteristics: age, antecedent pregnancy, interval months from antecedent pregnancy, stage, and FIGO prognostic score. 2) Imaging exams including pelvic ultrasound, chest X-ray, chest computed tomography (CT), abdominal and brain CT, or magnetic resonance imaging: detectable lesions, maximum lesion diameters, and number of metastases. 3) Therapeutic regimen: number of MTX cycles, the reasons switching to second-line Act-D or/and other salvage regimens, and treatment outcome. 4) Serum hCG monitoring before each chemotherapy cycle. 5) Chemotherapy-related toxicity: oral mucositis, alopecia, nausea, vomiting, complete blood cell count, platelet count, and renal and liver function tests (serum creatinine, serum alanine aminotransferase, aspartate aminotransferase, and bilirubin).

The drug resistance criteria were: serum hCG levels not showing a logarithmical decline, remaining at a plateau level, or increasing during 2 successive chemotherapy cycles or imaging files indicating that the tumor size did not decrease or increase or that new lesions appeared [31]. The trends in hCG levels were classified as elevation and declination. An elevation in hCG level was defined as an increase in serum hCG level during 2 successive MTX chemotherapy cycles, while the opposite situation was defined as a declination. Drug toxicity was assessed in each cycle according to World Health Organization criteria [32]. First-line MTX chemotherapy was stopped in the cases of drug resistance, severe side effects, or intolerable toxicity. Complete remission was defined as cases in which the hCG level remained normal within 3 months after treatment. To determine recurrence and survival, we performed a follow-up at least 1 year.

The normality of all continuous data was tested by 1-sample Kolmogorov–Smirnov tests. If the data were normally distributed, t-tests were used to compare means between the 2 groups. Otherwise, nonparametric Wilcoxon–Mann–Whitney tests were used. The heterogeneity of all distributions was assessed by χ² or Fisher’s exact tests. Receiver operating characteristic (ROC) curves were used to test the discriminating potential of important factors identified in the above-described comparative analysis. An area under the curve (AUC) with a lower 95% confidence interval (CI) of >0.5 was considered significantly discriminant. The p<0.05 (2-tailed) was considered statistically significant. All analyses were performed using PASW Statistics for Windows, version 18.0 (SPSS, Inc., Chicago, IL, USA) on Windows XP (Microsoft Corp., Redmond, WA, USA).

RESULTS

At our center, a total of 299 low-risk GTN patients received a 5-day MTX regimen as first-line chemotherapy between January 2000 and December 2015. Among these patients, 101 cases switched to Act-D as second-line chemotherapy, including 83 cases developing MTX resistance and 18 cases with intolerable toxicity. Except for 10 cases administered Act-D as consolidation therapy and 2 cases that stopped Act-D due to intolerable toxicity, 89 patients were included in the final analysis. Fig. 1 shows the flow chart of subject inclusion.
Table 1 shows the clinical characters of 89 patients administered Act-D salvage therapy after MTX failure. The median patient age was 28 years, ranging from 16 to 50 years. The response rate for Act-D was 82.02% (73/89). Sixteen cases with Act-D resistance were switched to multiple-agent EMA-CO chemotherapy. All patients continued to receive at least 2 cycles of consolidation chemotherapy after hCG levels decreased to normal (<5.3 IU/L) and were defined as achieving complete remission. Sixty-six cases were monitored for 1 – 16 years after complete remission. No case of recurrence was observed and the overall survival rate was 100%.

We compared the clinical features of Act-D-resistant cases to those of Act-D responders (Table 1). The results showed that the serum hCG levels before Act-D salvage therapy (hCG_{Act-D}) were significantly higher in the Act-D-resistant cases than those in the Act-D responders (median 605 vs. 103 IU/L, p=0.009). However, other factors, such as age, FIGO prognostic score, and serum hCG level before primary MTX therapy, did not show significant differences.

The trends in serum hCG levels during primary MTX therapy did not differ significantly between the 2 groups, although the FIGO statement indicated that elevation in hCG level might be associated with Act-D resistance [33]. In our 16-year data, while 13/89 patients (14.61%) had elevated serum hCG levels during primary MTX chemotherapy, most (84.62%, 11/13) responded to Act-D. Even the case with the largest increase in serum hCG level (from 1,517 to 12,630 IU/L) in successive MTX cycles still responded to Act-D salvage therapy. Furthermore, the trend in serum hCG level during primary MTX therapy was not significantly associated with age, FIGO prognostic score, number of Act-D cycles, or other clinical factors (Table 2).

Thus, hCG_{Act-D} was the only factor associated with Act-D resistance in our study. ROC curve analysis showed an AUC of hCG_{Act-D} of 0.71 (95% CI, 0.58–0.84), indicating its discriminatory potential (Fig. 2). The optimal hCG_{Act-D} cut-off value for Act-D resistance was 330 IU/L, with
68.80% sensitivity and 69.90% specificity. Patients with hCGAct-D ≥330 IU/L had a 5-fold higher risk of Act-D resistance compared to that in cases with hCGAct-D <330 IU/L (risk ratio [RR]=5.10; 95% CI=1.58–16.42; p=0.004).

However, the scatter diagram showed that the range of hCGAct-D values in Act-D responders was wider than that in the Act-D-resistant cases (5.76– 16,664 IU/L vs. 11.43–6,732 IU/L, Fig. 3). Most (71.43%, 10/14) cases with high hCGAct-D level (>1,000 IU/L) responded to Act-D. Therefore, hCGAct-D values did not provide overwhelming evidence to predict whether individual cases would develop Act-D resistance.

Except for 2 cases requiring other salvage regimens due to Act-D toxicity (Fig. 1), nearly all patients (97.80%, 89/91) tolerated the toxicity. Assessment of Act-D side effects for each cycle revealed that 59.55% of patients (53/89) experienced mild (grade I or II) chemotherapy-related toxicity. Neutropenia occurred at the highest frequency, developing at grade IV and III.
in 6.74% (6/89) and 23.60% (21/89) of patients. All patients were administered granulocyte colony-stimulating factor instead of stopping Act-D salvage therapy (Table 3). Patients with mild to moderate stomatitis were successfully treated with various “stomatitis cocktails.” No grade III hepatotoxicity related to Act-D salvage therapy was observed. Grade III nausea/vomiting was not a common side effect.

**DISCUSSION**

This study reported our 16-year experience in managing low-risk GTN patients who failed primary MTX chemotherapy. These patients were administered an Act-D salvage regimen and 82.02% responded to 5-day Act-D. The response rate was consistent with that in our previous paper, which showed 81.25% response rate in 16 patients from 1999 to 2004 [25]. Both datasets stated the appropriate therapeutic effect of the 5-day Act-D salvage therapy on management of low-risk GTN with failed primary MTX.
the low-risk GTN patients with MTX failure. To our knowledge, the number of subjects in our study was larger than that in many existing studies on 5-day Act-D salvage therapy in low-risk GTN [3,6,7,15,16,34].
Meanwhile, around 20% of cases in the present study developed Act-D resistance after MTX resistance. Repeated drug resistance prolongs the treatment period, delays pregnancy, and increases psychological stress. Several studies have explored predictors of resistance to Act-D salvage therapy. Kang et al. [35] analyzed 38 cases, reporting response rates of 89.29% (25/28) for cases with hCG_{Act-D} <1,000 IU/L, 42.86% (3/7) for 1,000–10,000 IU/L, and 100% (0/3) for ≥10,000 IU/L. In our study, hCG_{Act-D} levels in the Act-D-resistant cases were significantly higher than those in Act-D responders (p=0.009). Thus, the possibility of resistance to Act-D salvage therapy increased with increasing serum hCG_{Act-D} level. Gynecological oncologists agree with prescribing Act-D as salvage therapy for patients with low hCG_{Act-D} and EMA-CO for those with high hCG_{Act-D} [3,14-16].

However, the definition of high hCG_{Act-D} remains controversial, varying from 100 to 1,000 IU/L in the literature [3,14-16]. In our study, the optimal cut-off value for Act-D resistance was 330 IU/L with 68.80% sensitivity, 69.90% specificity, and RR of 5.1. However, the range of hCG_{Act-D} values in Act-D responders was wider than that in the Act-D-resistant cases. Only 33.33% of patients (11/33) with hCG_{Act-D} levels over the cut-off value showed Act-D resistance, while 28.57% of patients (4/14) showed Act-D resistance with hCG_{Act-D} ≥1,000 IU/L, the threshold proposed by Prouvot et al. [16]. Although the risk of drug resistance increased with increasing hCG_{Act-D} level, resistance developed in a small proportion of the population. Thus, it is reckless to use multiple agent regimens for the small number of potentially drug-resistant cases, which might sacrifice more patients to tolerate the side effects of multiple drugs. Therefore, the predictive power of hCG_{Act-D} to identify individuals with Act-D resistance was limited.

The 2018 FIGO cancer report recommended the initiation of multiple agents in cases with significant elevation in hCG level during the first single-agent treatment [33]. However, this report did not give a precise numerical definition of “significant”. Furthermore, other literature also failed to define this term in this context [33,36]. In the 16 years of data of our institute, 14.61% of cases (13/89) administered Act-D salvage treatment experienced elevated hCG levels during MTX chemotherapy. While we have identified an association between elevation in hCG level during MTX chemotherapy and Act-D resistance, additional studies with more patients are required to reach clear conclusions regarding this relationship.

In terms of drug side effects, most of the patients in this study experienced mild chemotherapy-related toxicity. In comparison, half of the ultra-high-risk GTN patients (50.00%, 12/24) administered multiple-agent EMA-CO chemotherapy in our center experienced severe neutropenia (grade III or IV) during the same period [29]. Among these patients, 37.50% (9/24) stopped cyclophosphamide and vincristine (CO) chemotherapy for severe adverse side effects [29]. These results indicate that the toxicity of Act-D was much lower than that of EMA-CO [16,21]. No cases of relapse were observed in follow-ups performed at least 1 year. The overall survival rate was 100.00%. Because only a few cases developed resistance to Act-D salvage therapy, unless strongly indicated, we do not suggest the initial use of EMA-CO protocols due to the risk of serious side effects.

One of the limitations of this study was that we did not compare cases directly treated by EMA-CO to those subsequently treated by Act-D and EMA-CO after MTX failure. This comparison is difficult to perform in clinical studies. Moreover, data on long-term response were unavailable for the 23 patients (23/89, 25.84%) lost to follow-up. Finally, we retrieved data retrospectively, resulting in a potential bias that may have underestimated some of the toxicities.
In conclusion, the predictive value of hCG_{Act-D} for Act-D salvage therapy could not be determined, although the hCG_{Act-D} level was higher in resistant cases than that in responders. Based on the good therapeutic effect and tolerable toxicity, we recommended 5-day Act-D salvage therapy for all patients who fail primary MTX chemotherapy. Additional studies are required to determine the proper criteria for regimen selection.

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