BMJ Open: Consolidation chemotherapy in postmolar low-risk gestational trophoblastic neoplasia: a systematic review protocol

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ABSTRACT

Introduction: Current evidence remains insufficient to strongly demonstrate the benefits of consolidation chemotherapy to all women with low-risk gestational trophoblastic neoplasia (GTN). This protocol outlines a systematic review to investigate whether consolidation chemotherapy is necessary for all patients with postmolar low-risk GTN after human chorionic gonadotropin normalisation with first-line single-agent chemotherapy.

Methods and analysis: A search string will be used to search the PubMed (MEDLINE), EMBASE, Web of Sciences, Scopus, LILACS and Cochrane Central Register of Controlled Trials databases. Articles will be screened at the title and abstract level, and then at the full article level by two independent reviewers using inclusion/exclusion criteria. Randomised and non-randomised study designs will be included, while case studies, commentaries, editorials, review articles, animal studies, basic science studies and cross-sectional studies, as well as studies not reporting relapse/recurrence rates and/or whether consolidation chemotherapy was delivered will be excluded. There will be no restrictions on date of publication, geographical location, study setting, or language of publication. The primary outcome is rate of recurrence/relapse. The assessments of randomised controlled trials will be performed using the risk of bias tool from the Cochrane Collaboration. Non-randomised studies will be assessed using the Newcastle-Ottawa scale. The quality of evidence will be assessed using the Grading quality of evidence and strength of recommendations (Grades of Recommendations, Assessment, Development and Evaluation) guidelines.

Ethics and dissemination: No formal ethical approval is required as all data collected will be secondary data and analysed anonymously. Results will be disseminated through a peer-reviewed publication and at scientific events.

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Strengths and limitations of this study:

- The number of randomised controlled trials found is expected to be small as this type of study usually requires a large sample size not easily available in the context of rare cancers such as gestational trophoblastic neoplasia (GTN).
- Differences in GTN diagnostic and classification criteria, as well as in the definition of relapse, may confound comparisons.
- The rigorous and transparent study design will possibly reduce the risk of biases.
- Grading the quality of the evidence will provide confidence in the effect estimates.

INTRODUCTION

Gestational trophoblastic neoplasia (GTN) refers to a group of malignant lesions that arise from placental villous and extravillous trophoblast and may follow a hydatidiform mole or a nonmolar pregnancy. Based on GTN classification and stage, treatment consists of either single agent therapy with methotrexate (MTX) or Actinomycin D for low-risk disease, or multiagent therapy for high-risk disease.

Chemotherapy should be continued until complete remission is achieved, with a normalised serum human chorionic gonadotropin (hCG) level. However, tumour resistance or relapse after first-line chemotherapy has been reported in approximately 25% of women with low-risk GTN. To eradicate the remaining tumour cells, consolidation therapy with the last effective agent(s) has been recommended to prevent relapse.

Few studies have addressed consolidation therapy for low-risk GTN. Yang et al identified less than two courses of consolidation chemotherapy as a risk factor for relapsed disease while Sun et al published results suggesting the opposite. This is likely to be due to the fact that besides using different definitions for GTN and relapse, those studies combined patients with postmolar GTN with patients with post-delivery GTN, and patients normalising on MTX with patients who had progressive disease on MTX.
Lybol et al retrospectively compared relapse rates following two (The Netherlands) or three (UK) MTX consolidation courses in women completing MTX therapy for low-risk GTN. They found a 4% relapse rate after three consolidation courses of MTX alternating with folinic acid in contrast to an 8.3% relapse rate following two consolidation courses, suggesting that three courses of consolidation chemotherapy is preferable to two in order to decrease the risk of low-risk GTN relapse. Nonetheless, differences between the Netherlands and the UK regarding the scoring systems employed in defining low-risk disease GTN, and in hCG measurement methods might have influenced the composition of the patient groups studied and the relapse rates observed.8

Couder et al retrospectively analysed the predictive factors of relapse in a homogeneous population with FIGO-2000-defined low risk GTN that received two consolidation courses after hCG normalisation with MTX alone. They found a relapse rate of 5.7%, consistent with the rate reported in previous series where the FIGO scoring system was not uniformly used. Additionally, they demonstrated that postpartum low-risk GTN and patients who need more than four courses of MTX for normalisation are at a higher risk of relapse than other low-risk patients (8.66-fold and 6.7-fold higher relapse risk, respectively). According to these authors, these findings suggest different biological behaviours among low-risk GTN patients and support previously reported relapse rates as low as 3% without any consolidation.9

Taken together, these data indicate that current evidence on the benefits of consolidation chemotherapy to all women with low-risk GTN remains insufficiently strong. Thus, this review aims at investigating whether consolidation chemotherapy is necessary for all patients with postmolar low-risk GTN after hCG normalisation with first-line single-agent chemotherapy. To this end, the proposed review will determine patient, disease and treatment characteristics of the women with postmolar lowrisk GTN included in each of the studies of interest; and assess the relationship of relapse rate with these characteristics, use of consolidation chemotherapy, and number of consolidation chemotherapy cycles used (if any).

Review question
The components of population, intervention, comparator, outcome and time frame are as follows:

Population: women with postmolar low-risk GTN who achieved remission (hCG normalisation) with single-agent chemotherapy.

Intervention: Consolidation chemotherapy after hCG normalisation with first-line single-agent chemotherapy.

Comparator: No consolidation treatment or placebo after hCG normalisation with first-line single-agent chemotherapy.

Outcome: The primary outcome is rate of recurrence/relapse. Due to possible variation in disease definitions over time, definitions of outcomes will be extracted as reported in individual studies. Secondary outcomes will include consolidation chemotherapy toxicity and time to relapse recurrence.

Time frame: up to 1 year of follow-up after hCG normalisation with first-line single-agent chemotherapy.

This protocol outlines the procedures for a systematic literature review intended to answer the question: Is consolidation chemotherapy necessary for all patients with postmolar low-risk GTN to prevent recurrence after hCG normalisation with first-line single-agent chemotherapy?

METHODS AND ANALYSIS
This protocol was designed according to PRISMA (Preferred Reporting Items for Systematic Reviews and Meta-Analysis Protocols) guidelines10 (checklist in online supplemental file 1).

Eligibility criteria
As information on consolidation therapy for postmolar low-risk GTN is limited, the following randomised and non-randomised study designs will be included: randomised and quasi-randomised trials (including cluster and cross-over trials), non-randomised controlled trials (RCTs), prospective and retrospective cohort studies, case-control studies, controlled before-and-after studies, historically controlled studies, case series, and interrupted time-series studies. Abstracts and conference proceedings will also be considered.

Case studies, commentaries, editorials, review articles, animal studies and basic science studies, cross-sectional studies, as well as studies not reporting relapse/recurrence rates and/or whether consolidation chemotherapy was delivered will be excluded. In studies also including women with other types of GTN (eg, nonmolar, high risk), only data for the postmolar low-risk subgroup will be extracted. To ensure literature saturation, the reference lists of the studies included will be hand searched.

There will be no restrictions on date of publication, geographical location, study setting or language of publication.

Search strategy
The specific search strategies will be created by a Health Sciences Librarian with expertise in systematic review searching with input from the project team. Searches will be conducted in PubMed (MEDLINE), EMBASE, Web of Sciences, Scopus, LILACS and Cochrane Central Register of Controlled Trials (CENTRAL). A draft search strategy for all databases to be used is presented in online supplemental file 2. After the PubMed strategy is finalised, it will be adapted to the syntax and subject headings of the other databases. Reference lists of included studies identified through the search will be scanned to identify any potentially eligible studies. The search will be updated towards the end of the review to capture recently published literature.
Data management

Two independent reviewers will perform data abstraction and quality assessment. Disagreements will be resolved through discussion and/or with the involvement of third-party arbitration. Using standardised forms, the reviewers will extract data independently and in duplicate from each eligible study. To ensure consistency across reviewers, calibration exercises will be conducted before starting the review. The Grading of Recommendations, Assessment, Development and Evaluation Profiler (GRADEpro) software will be used to create tables for summary of findings and quality assessment.

Screening

All screening steps will be conducted independently by two reviewers. Duplicate references will be removed, and two review authors will independently screen the titles and abstracts yielded by the search. Full-text articles that meet the inclusion criteria will then be retrieved and screened. The reasons for excluding trials will be recorded. Neither of the review authors will be blind to the journal titles or to the study authors or institutions.

To facilitate collaboration among reviewers, search results will be uploaded to a web-based systematic review tool (eg, Rayyan** or COVIDENCE) during the study selection. Training will be provided to members of the review team not familiar with such tools. Prior to the formal screening process, screening questions and forms will be developed and tested for assessments based on the inclusion and exclusion criteria.

The reviewers will meet after each stage to assess agreement. Disagreements will be resolved through discussion. If the conflict persists, a third independent reviewer will be consulted to reach unanimity.

The PRISMA flow diagram10 will be used to report the screening process.

Data abstraction

A data abstraction form will be created and piloted by two reviewers on a sample subset of publications for this review. Data to be collected include study title, authors, publication date, number of patients with postmolar low-risk GTN, low-risk GTN definition, type of antecedent pregnancy, pretreatment hCG, single-agent chemotherapy regimen used, number of single-agent chemotherapy cycles needed to achieve hCG normalisation, interval between chemotherapy cycles until hCG remission, number of consolidation chemotherapy cycles administered, consolidation chemotherapy adverse events, relapse/recurrence rate, relapse/recurrence definition, time to relapse/recurrence, relapse/recurrence prognostic factors identified.

Quality assessment

The assessments of RCTs will be performed using the risk of bias tool from the Cochrane Collaboration,11 which evaluates potential bias for seven items across six domains: selection bias (random sequence generation; allocation concealment), performance bias (blinding of participants and personnel), detection bias (blinding of outcome assessment), attrition bias (incomplete outcome data), reporting bias (selective reporting), and other sources of bias. Each study will be rated as 'high', 'unclear' or 'low' risk of bias.

Non-randomised studies will be assessed using the Newcastle-Ottawa scale12 that evaluates each study on eight items falling into three categories: (1) selection of study groups, (2) comparability of groups and (3) ascertainment of exposure or outcome of interest.

The quality of evidence will be assessed using the Grading Quality of Evidence and Strength of Recommendations (GRADE) guidelines,13 which covers risk of bias, inconsistency, indirectness, imprecision and publication bias. The overall quality of evidence is rated as high, moderate, low or very low by each outcome measure.

The GRADEpro software will be used to create tables for summary of findings and quality assessment.14

Data analysis

Aggregate data will be used, and quantitative synthesis planned where possible if the studies included are sufficiently homogeneous in terms of design and disease type or outcome definitions (homogeneous outcomes in at least two studies). Continuous data will be expressed as means and SD. For dichotomous data, the OR will be calculated, and a log-rank approach will be used to estimate a HR, both with 95% CI. Random effects is the model chosen for meta-analysis as treatment effect is assumed to vary among studies. For quantitative synthesis, Review Manager V5.3 software will be used to pool the results of trials for each outcome.

Inconsistency among the results of the included studies will be ascertained by visual inspection of a forest plot (absence of overlap of CIs around the effect estimates of individual studies), as well as by the Higgins inconsistency test or I², where I²>50% indicates moderate likelihood of heterogeneity.

The quality of evidence of intervention effect estimation for outcomes that can be plotted in meta-analysis will be generated according to GRADE.

Qualitative synthesis is planned for outcomes in which quantitative synthesis is not feasible or appropriate.

Any amendments to this protocol will be documented with their corresponding rationale in the full review.

Patient and public involvement

There is no patient and public involvement in this study.

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Contributors MB-S will serve as the first author of the protocol and review paper. She led all stages of protocol development, including development of the research question and objectives, search strategy, and extraction and analysis plans. IM, KE,
REFERENCES


