Efficacy of different surgical approaches on survival outcomes in patients with early-stage cervical cancer: protocol for a multicentre longitudinal study in China

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ABSTRACT

Introduction Recent studies have revealed that the oncolgical survival outcomes of minimally invasive radical hysterectomy (MIRH) are inferior to those of abdominal radical hysterectomy (ARH) in early-stage cervical cancer, but the potential reasons are unclear. Methods and analysis Each expert from 28 study centres participating in a previously reported randomised controlled trial (NCT03739944) will provide successive eligible records of at least 100 patients who accepted radical hysterectomy for early-stage cervical cancer between 1 January 2009 and 31 December 2015. Inclusion criteria consist of a definite pathological evaluation of stages IA1 (with positive lymphovascular space invasion), IA2 and IB1 according to the International Federation of Gynecology and Obstetrics 2009 staging system and a histological subtype of squamous cell carcinoma, adenocarcinoma or adenosquamous carcinoma. The primary endpoint is 5-year disease-free survival between the MIRH and ARH groups. The secondary endpoints include the MIRH learning curves of participating surgeons, 5-year overall survival between the MIRH and ARH groups, survival outcomes according to surgical chronology, surgical outcomes and sites of recurrence and potential risk factors that affect survival outcomes. A subgroup analysis in patients with tumour diameter less than 2 cm will follow the similar flow diagram. Ethics and dissemination This study has been approved by the Institutional Review Board of Peking Union Medical College Hospital (registration no. JS-1711), and is also filed on record by all other 27 centres. The results will be disseminated through community events and peer-reviewed journals.

Trial registration number NCT03738969

INTRODUCTION

Cervical cancer is the fourth most common malignancy in women worldwide. In addition, 85% of patients with cervical cancer are from developing countries, and cervical cancer is the second leading cause of cancer-related death in women. In China, according to a conservative estimate, the total incidence and mortality of cervical cancer in 2015 were 98 900 and 30 500 cases, respectively, accounting for 28% of the total number of new cases of cervical cancer worldwide. Radical hysterectomy (RH) with pelvic lymphadenectomy is the standard surgical treatment for patients with early-stage cervical cancer. Previously published guidelines indicate that either laparotomy (open surgery) or minimally invasive surgery (MIS) is an acceptable approach to treat RH in patients with early-stage (IA2 to IIA) cervical cancer. In numerous retrospective studies, although MIS and abdominal RH (ARH) have comparable 5-year disease-free survival (DFS) and overall survival (OS) rates, MIS also has better shorter-term quality of life (QoL) and improved surgical outcomes, even in elderly patients. Nevertheless, one prospective randomised study by Ramirez et al and one retrospective epidemiological study by Melamed et al put forward doubts and documented that ARH for...
early-stage cervical cancer is superior to minimally invasive RH (MIRH) regarding DFS and OS. These findings were also confirmed in a recent population-based survey in England. It seems that MIS may have a reputation for this inferior trend, which is significantly different from the situation observed in early-stage uterine, colorectal or gastric cancer.22–24 These studies caused great controversy over the surgical approaches of cervical cancer worldwide, but they certainly changed the idea of the best approach to offer early-stage cervical cancer patients undergoing surgical treatment.25–31 However, the causes of the inferior survival outcomes in the MIS group are still unknown. Many possible reasons for the inferiority of MIS have been debated, such as different radicality, incision methods, ethnic differences, tumour size as a selection criterion, learning curves, data completeness, circulating CO2 levels and the usage of manipulators.35 Hence, we performed a randomised controlled trial (RCT) in 28 Chinese centres (RACC, NCT03739944) to evaluate the survival outcomes of patients who underwent MIRH and ARH and to explore potential risk factors.36 This trial is based on a participating expert-centred design rather than a study site-centred design.36

However, the two most important questions that arose from the RACC study (NCT03739944) await answers. One is the safety of MIRH for patients with early-stage cervical cancer that was inferior, reported in the Laparoscopic Approach to Cervical Cancer (LACC) trial and other studies.13,25,26 The other is how to qualify the experiences of participating surgeons. In one retrospective study on RH procedures performed by one of the participating surgeons, we found that the learning curve probably explains the disadvantage of MIRH, as the survival outcomes of patients who underwent the first dozens of MIRH procedures performed by the surgeon were inferior to those of patients who underwent ARH.37 The current study proposes a multicentre longitudinal study in China to answer these two questions, since the study centre and surgeons who participated in NCT03739944 are the same as those who will participate in the current study.

The quality of the participating surgeons can be determined by the MIRH learning curve. Few studies have considered the influencing factors involving surgeons and their learning curves,37,38 and research defining learning curves in gynaecological oncology surgery is limited. Surgeons can be proficient after 20 cases of robotic-assisted RH (RRH) and continually improve between 50 and 70 cases.39–41 Mastery of laparoscopic RH (LRH) requires experience in at least 25 and up to 50 cases.42–44 According to Hwang et al,45 the learning period for LRH and lymph node dissection to reach a turning point was calculated to be 40 cases. The systematic review found a slow learning curve required for a surgeon to gain expertise in laparoscopically assisted vaginal RH.46 These studies support the necessity of evaluating the learning curves in a demanding surgical modality.

In addition, two retrospective reports of a large series showed that ARH and MIRH had different survival outcomes in patients with a tumour diameter >2 cm, while patients with a tumour diameter <2 cm experienced superior outcomes to those with a tumour diameter >2 cm.47 Kim et al.47 also reported that MIRH was not a poor prognostic factor for patients with stage IB1 cervical cancer and cervical masses ≤2 cm in size. Some authors also argued that the reduction in survival in patients receiving MIRH could be due to tumour manipulation and subsequent cancer spread.48

In general, based on the participating surgeons of an RCT,36 we will perform this multicentre longitudinal study to compare the survival outcomes of patients undergoing ARH and MIRH and to qualify the learning curves of participating surgeons. Other essential issues, including tumour diameters and definite surgical procedures which were considered as potential risk factors, will be also evaluated in this study.

Aims and objectives
Primary objectives
1. The primary aim of this longitudinal study is to analyse the 5-year DFS rates of patients with early-stage cervical cancer receiving different surgical routes, including MIS (RRH or LRH) versus ARH, in all enrolled patients and in patients with tumour diameter <2 cm.

Secondary objectives
1. The learning curves of participating surgeons will be presented as their qualification proof.
2. The 5-year OS rates between patients in the MIRH and ARH groups.
3. The 5-year DFS and OS rates according to chronology in general and individually.
4. To compare the time of surgery, estimated blood loss during surgery, volume of transfusion, length of hospital stay, perioperative complications and postoperative complications between the MIRH and ARH groups.
5. To describe patterns of recurrence, including sites, tumour burden and biomarkers.
6. To investigate relevant factors guiding a surgeon’s choice about the best surgical approach for patients who may still benefit from MIS without worsening their survival outcomes. These factors include pathological characteristics, energy devices, uterine manipulators, adjuvant treatment, follow-up protocols, nerve-sparing procedures and expert characteristics. These analysis will be performed in all enrolled patients and in patients with tumour diameter <2 cm.

METHODS AND ANALYSIS
Study design
This is a retrospective, multicentre study comparing the efficacy of different surgical approaches on the surgical and oncological survival outcomes in Chinese patients with early-stage cervical cancer receiving RH. A total of
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28 Chinese centres (online supplementary file 1), same as the RACC trial will collect patients’ medical records. Each expert included in this study will provide the medical records of at least 100 consecutive patients who were hospitalised between 1 January 2009 and 31 December 2015, according to a predefined template. These patients were followed up until 31 December 2019, to guarantee a follow-up of at least 4 years. The study flow chart is illustrated in figure 1.

Recruitment and eligibility
Patients’ medical records will be collected based on the following eligibility criteria.

Inclusion criteria
1. Patients with stage IA1 (with lymphovascular space invasion (LVSI)), IA2 or IB1 cervical cancer according to the International Federation of Gynecology and Obstetrics 2009 staging system and an Eastern Cooperative Oncology Group performance status of 0 to 1.
2. Histological subtypes of squamous cell carcinoma, adenocarcinoma, and adenosquamous carcinoma.
3. Patients older than 18 years at diagnosis.
4. Patients had complete epidemiological and clinicopathological data recorded.
5. All enrolled patients underwent surgery performed by the experts designated in the research centre.

Exclusion criteria
1. Patients with a histological subtype of neuroendocrine carcinoma, clear cell carcinoma, serous cell type carcinoma or metastatic carcinoma.
2. Patients with stage IA1 disease without LVSI or clinically advanced disease (stages IB2–IV).
3. Pregnant women.
4. Patients receiving neoadjuvant chemotherapy or radiotherapy before RH.
5. Patients with a history of pelvic radiotherapy, pelvic reconstruction, and brain/spinal cord diseases.
6. Patients with a positive HIV status, autoimmune disorders and systemic disease (such as hormone treatment diseases, severe liver and kidney dysfunction) or a severe mental illness and pre-existing cancer diagnosis.
7. Patients without integrated medical records, such as details on epidemiology, surgery and pathology.

Interventions
Study centre and surgeon selection
The selection and determination of study centres and surgeons were described previously. All the surgeons (ie, the principal investigators) from all study centres listed in online supplementary file 1 approved the study protocol and signed a research agreement form.

Surgical treatment
The surgical procedures performed have been described previously: these consisted of RH, bilateral salpingo-oophorectomy and pelvic lymphadenectomy (the resection of sentinel lymph nodes also needed clear notification), with/without concomitant para-aortic lymph node dissection. For young patients requiring the preservation of ovaries, bilateral salpingectomy was performed along with ovary suspension to the peritoneum above the level of the anterior superior spine. All the major procedures, including lymphadenectomy and parametrial resection with or without a nerve-sparing procedure, were primarily performed by one surgeon according to the Piver-Rutledge-Smith classification or to class B or C of the Q-M classification. Patients who converted to laparotomy due to intraoperative complications were allocated to the ARH group. During the review of medical records, the following surgical issues will receive special attention and recording: energy devices, artificial pneumoperitoneum, manipulator usage, vaginal incision methods used during the surgeries and nerve-sparing procedures.

Postoperative adjuvant treatment
Postoperative adjuvant therapies, including systematic chemotherapy, radiotherapy and chemoradiotherapy, will be recorded. Therapy regimens, dosages, durations and adverse events will also be addressed.

Sample size calculation
The primary objective of this study is to explore whether there are differences between MIRH and ARH with respect to DFS. We assume that the 5-year DFS rate will...
be approximately 85%, and the non-inferiority threshold of 5% is clinically acceptable. The corresponding HR is set at 1.50 based on the significance level, and the power is set at 0.8. A total of 1140 patients are needed for the MIRH and ARH groups. Considering the possible 20% loss to follow-up rate, 2850 patients are needed to accomplish the study goal.

In this study, since each participating hospital will be asked to provide the successive eligible records of at least 100 patients, approximately 3000 or more patients could be examined in total to achieve the estimated sample size. However, such estimation does not guarantee a sufficient sample size in the MIRH or ARH group.

Measurement
The patients’ detailed epidemiology, surgical details, pathological characteristics, perioperative complications and follow-up data, as well as postoperative adjuvant therapy data, will be collected by medical staff. All data obtained from domestic research centers will be input into the database by trained medical staff. Details are as follows:

1. Surgical outcomes include estimated blood loss, transfusion, surgical duration and hospital stay after RH.
2. Pathological outcomes include the measurement of critical parameters from available records, consisting of the width or length of the resected parametrium, vagina and uterosacral ligaments under their natural conditions; numbers and locations of harvested lymph nodes; and feasibility of sentinel lymph node biopsy. Due to the results and concerns from LACC trial and other studies, patients with tumour diameter <2 cm will be included a subgroup analysis as to guarantee the safety of patients in future trials.
3. Complications will be recorded as intraoperative, perioperative, early (<4 weeks) postoperative and delayed postoperative (4 weeks to 6 months after RH) according to the protocol of the LACC trial, and severity will be judged by the Common Terminology Criteria for Adverse Events (CTCAE) V.4.03. Definite complications include intraoperative complications (bladder injury, ureteral injury, bowel injury, vascular injury and obturator nerve injury), postoperative complications (urinary retention with catheterisation, cerebrovascular, pulmonary and renal diseases, ileus, abdominal wound and vault complications, septicemia, thromboembolic complications, pelvic haematoma, lymphocyst formation and/or secondary infection, postoperative vesicovaginal fistula, ureterovaginal fistula, rectovaginal fistula, chylous leakage and lymphedema).
4. Survival outcomes: the time to recurrence will be calculated from the date of surgery to the time the patient is alive.

Statistical analysis
Continuous variables conforming to normal distribution will be described as the means and SDs and compared with parametric methods. Non-parametric tests will be used to assess the categorical data, and discrete variables that do not conform to normal distribution will be summarised as the medians, ranges and IQRs. Survival curves will be generated by the Kaplan-Meier method, and Cox proportional hazards models will be used to estimate the HRs and 95% CIs for the effect of surgical routes on DFS, progression-free survival and OS. A multivariable analysis of DFS will be performed after adjusting for important pathological risk factors. Statistical analysis will be performed using SPSS V.22.0 (SPSS Inc, Chicago, Illinois, USA). Significance will be set at a p value of 0.05.

Safety and adverse events
This trial will be conducted in compliance with this study protocol. Although it is a retrospective study, the Data Safety Monitoring Committee (online supplementary file 2) in the RACC trial will still receive study data regularly, including complications and survival outcomes. The Committee will review the final data of individual patients with MIRH to find out the potential plateau of DFS. ARH, abdominal radical hysterectomy. DFS, disease-free survival. MIRH, minimally invasive radical hysterectomy.
surgeon’s learning curve, as to make recommendation about further investigation. The concern about the tumour diameter (less or $\geq$2 cm) will receive specific attention.

**Limitations**
The retrospective nature of the current study is the most important limitation. Disease-specific and treatment-specific issues affecting QoL cannot be adequately assessed. The emphasis on the individual surgeon’s experience and skill probably limits generalisation to all surgeons. How to eliminate or decrease the restriction of the learning curve for young surgeons is critical in oncology education.

**Patient and public involvement**
Epidemiological data and clinicopathological characteristics were retrospectively collected by trained medical staff from a third party by reviewing the medical records. Complications related to RH within 6 months were collected as adverse events according to the CTCAE V.4.03.55 Data on recurrence and mortality were obtained from the medical records or follow-up data by the certificated third party. Sites of recurrence were verified by surgeries and/or imaging evaluations. A detailed follow-up by telephone was also provided to patients with unknown or uncertain survival outcomes. In the current trial, no patients were involved in the design of the study or in the selection of outcome measures.

**Ethics and dissemination**
The study will be practiced in compliance with this study protocol. The patient’s consent to accept the treatment and the approval of the trial by each study centre will be reviewed and collected by the principle investigator (LL and MW). All procedures performed in the study involving human participants were in accordance with the ethical standards of the institutional and/or national research committee and with the 1964 Declaration of Helsinki and its later amendments or comparable ethical standards.

This is the first version of protocol (Identifier 1.0, 16 February 2019). The protocol will be public on the website of the principle study centre (http://www.pumch.cn). Any important protocol modifications must be communicated to all study centres and investigators, and must be reported to and be approved by the Institutional Review Board of all study centres.

Personal information about potential and enrolled participants will be masked by systemic sequence number so as to be collected, shared and maintained in order to protect confidentiality before, during and after the trial. The final data will be available on the online dataset (http://pi.kangruihiealth.com) for data sharing after anonymization processing.

**CONCLUSION**
Due to the complex and complicated characteristics of RH, the learning time needed to achieve comparable survival outcomes is probably long and is equally essential for laparotomy and laparoscopy. Data from large, well-designed observational studies are urgently needed. After adjusting the bias of surgeons, this study may provide essential evidence of whether MIRH has inferior oncological outcomes to ARH and the potential reasons involved despite of several assumptions, especially in patients with tumour diameter less or $\geq$2 cm. Other important surgical approaches associated with controversial issues, including definite surgical procedures, will also be discussed in this study.

**Contributors**
LL conceived of the original idea for the study, carried out the statistical analysis, edited the paper and was overall guarantor. XC obtained ethical approval and contributed to drafts of the paper. MW, SZ, SM, XT and XS contributed to the study design and commented on drafts of the paper.

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**Competing interests**
None declared.

**Patient and public involvement**
Patients and/or the public were not involved in the design, or conduct, or reporting or disseminating plans of this research.

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Obtained.

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