Study protocol for SPARED trial: randomised non-inferiority phase III trial comparing dexamethasone on day 1 with dexamethasone on days 1–4, combined with neurokinin-1 receptor antagonist, palonosetron and olanzapine (5 mg) in patients receiving cisplatin-based chemotherapy

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ABSTRACT

Introduction Dexamethasone (DEX) is administered for multiple days to prevent chemotherapy-induced nausea and vomiting for patients receiving highly emetogenic chemotherapy (HEC); however, its notorious side effects have been widely reported. Although our multicentre randomised double-blind comparative study verified non-inferiority of sparing DEX after day 2 of chemotherapy when combined with neurokinin-1 receptor antagonist (NK1-RA) and palonosetron (Palo) for patients receiving cisplatin (CDDP)-based HEC regimens in subgroup analysis. Recently, the efficacy of the addition of olanzapine (OLZ) to standard triple antiemetic therapy on HEC has been demonstrated by several phase III trials. This study aims to confirm non-inferiority of DEX sparing when it is combined with NK1-RA, Palo and OLZ in patients receiving CDDP-based HEC regimens.

Methods and analysis This is a randomised, double-blind, phase III trial. Patients who are scheduled to receive CDDP ≥50mg/m^2 as initial chemotherapy are eligible. Patients are randomly assigned to receive either DEX on days 1–4 or DEX on day 1 combined with NK1-RA, Palo and OLZ (5 mg). The primary endpoint is complete response (CR) rate, defined as no emesis and no rescue medications during the delayed phase (24–120 hours post-CDDP administration). The non-inferiority margin is set at −15.0%. We assume that CR rates would be 75% in both arms. Two hundred and sixty-two patients are required for at least 80% power to confirm non-inferiority at a one-sided significance level of 2.5%. After considering the possibility of attrition, we set our final required sample size of 280.

Strengths and limitations of this study

- This is the first trial to evaluate whether adding olanzapine to neurokinin-1 receptor antagonist, palonosetron and dexamethasone (DEX) can spare DEX administration on day 2–4 for patients receiving cisplatin-based regimens.
- This study is a multicenter, placebo-controlled, double-blind, randomised phase III study.
- A limitation of this study is that it was conducted solely within the Japanese population.

INTRODUCTION

Chemotherapy-induced nausea and vomiting (CINV) is one of the most frequent adverse reactions associated with chemotherapy and considerably reduces patient quality of life (QOL). CINV has been traditionally assessed in overall (0–120 hour postchemotherapy), acute (0–24 hour postchemotherapy) or delayed (24–120 hour postchemotherapy) phases.1 Intravenously administered cytotoxic agents are categorised into four emetic...
risk groups (high, moderate, low and minimal)\(^3\). Highly emetogenic chemotherapy (HEC) including cisplatin (CDDP)-based regimen and anthracycline plus cyclophosphamide (AC) regimen can lead to a >90% incidence of emesis in patients without an adequate antiemetic prophylaxis.\(^5\) When patients were administered prophylactic antiemetics prior to HEC treatment, the incidence of emesis (or requiring additional antiemetics) was found to be 35% according to a recent study.\(^5\)

Dexamethasone (DEX), 5-hydroxytryptamine type 3 receptor antagonists (5-HT3-RA) and neurokinin-1 receptor antagonists (NK1-RA) have been developed to inhibit CINV from HEC.\(^6,7\) DEX is typically administered for multiple days from the start of chemotherapy to care for delayed CINV;\(^7\) whereas frequent administration of corticosteroids has been associated with many adverse effects such as insomnia, hyperglycaemia and reduced bone mineral density.\(^8,9\) Therefore, corticosteroid-minimising (DEX-sparing) strategies, which administer DEX in the acute phase (day 1) and omit DEX in the delayed phase (day 2 and later), have been evaluated.\(^11\)

Our multicentre randomised double-blind comparative study (DEX-1 study) showed the complete response (CR; no emesis, no use of rescue medication) rate in the overall phase of DEX on day 1 provided a non-inferior antiemetic efficacy to a treatment of DEX on days 1–3 when combined with NK1-RA and palonosetron (Palo) for patients receiving HEC including AC and CDDP-based regimen. (44.0% vs 46.9%, p=0.007).\(^12\) In a subgroup analysis of patients receiving CDDP-based regimen, CR rates of acute phase demonstrated a non-inferiority of DEX on day 1 to days 1–3 (95.6% vs 95.6%, p=0.007); whereas CR rates of the overall and delayed phase DEX on day 1 were lower than those of DEX on days 1–3 (57.8% vs 66.7%, p=0.272; 57.8% vs 68.9%, p=0.349, respectively). However, the DEX-1 study was underpowered to multiple days of DEX compared with placebo (79% vs 66%, p<0.001); moreover, no differences were found between two groups in the incidence of sedation.\(^18\) The addition of OLZ in combination with NK1-RA, Palo and DEX has greater benefit and becomes a standard antiemetic therapy in patients receiving CDDP-based regimens.

Treatment with OLZ was associated with metabolic effects, including elevated glucose concentrations manifesting as insulin resistance.\(^9\) A phase III study showed that grade 3 hyperglycaemia was observed more frequently in the OLZ versus placebo group.\(^4\) Therefore, there is a concern that the combination of OLZ and multiple-day DEX may worsen glucose intolerance. In another study, Navari et al\(^20\) also demonstrated that OLZ, combined with a single dose of DEX and Palo, was very effective at controlling acute and delayed phase CINV in patients receiving HEC; moreover, this regimen was not associated with significant hyperglycaemia.

Based on these results, we speculate that the antiemetic regimen of OLZ 5mg, NK1-RA, Palo and a single dose of DEX could be effective and safe for delayed phase CINV in patients receiving CDDP-based regimen. We planned this randomised, double-blind, phase III trial to evaluate the non-inferiority of DEX on day 1 compared with DEX on days 1–4 when combined with NK1-RA, Palo and OLZ in patients receiving CDDP-based regimens.

### METHODS AND ANALYSIS

#### Study design

The Standard Protocol Items for Randomized Trials statement and checklist were followed in preparing the protocol. This multicentre, placebo-controlled, double-blinded, randomised, non-inferiority phase III study aims to confirm the non-inferiority of DEX on day 1 compared with DEX on days 1–4 combined with NK1-RA, Palo and OLZ 5mg to prevent CINV in patients with solid malignant tumour receiving CDDP-based regimens.

#### Study setting and participants

Recruiting will be performed in 10 sites across Japan. The inclusion and exclusion criteria are summarised in box 1. The main inclusion criterion is patients who are eligible in the study are 20–74 years old with malignant tumour, excluding haematological malignancies, receiving first-line treatment with CDDP ≥50mg/m\(^2\) (previous use of moderately or low emetogenic chemotherapy is permitted). The main exclusion criteria are as follows: (1) presence of systemic glucocorticoid therapy, (2) patients using antiemetics other than the trial drug, (3) patients receiving moderately emetogenic chemotherapy within 6 days before and after CDDP administration (minimally to low emetogenic agents are allowed), (4) patients receiving radiation therapy to abdomen or pelvis within 6 days prior to enrolment until 6 days after CDDP, (5) patients with symptomatic brain metastasis, diabetes

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Box 1 Inclusion and exclusion criteria

Inclusion criteria
- Patients with malignant tumour, excluding those with haematologic malignancies or those receiving first-line treatment with cisplatin (CDDP) ≥50 mg/m² (previous use of moderate or low emetic chemotherapy is permitted).
- Age: 20–74 years at the time of enrolment.
- Absence of nausea and vomiting within 24 hours prior to registration.
- Eastern Cooperative Oncology Group performance status of 0–1.
- Meeting the following standard values of general clinical tests within 2 weeks prior to enrolment: alanine aminotransferase <100 IU/L, aspartate aminotransferase <100 IU/L, total bilirubin <2.0 mg/dL, serum creatinine <1.5 mg/dL.
- Patients with an expected prognosis of 3 months or more.
- Patients who provided written informed consent.

Exclusion criteria
- Patients undergoing systemic glucocorticoid therapy.
- Patients using antiemetics other than the trial drug.
- Patients receiving moderately emetogenic chemotherapy within 6 days before and after CDDP administration.
- Patients who cannot be hospitalised until after 120 hours of starting CDDP administration as the study requires daily use of an electronic patient-reported outcome system.
- Patients receiving radiation therapy for the abdomen or pelvis within 6 days prior to registration until 6 days after CDDP administration.
- Patients with diabetes mellitus receiving treatment with insulin and/or oral hypoglycemic agents or patients with haemoglobin A1c (National Glycohemoglobin Standardization Program) >6.5% (>6.1% in the event of JDS).
- Patients with symptomatic brain metastasis, convulsive disorder requiring treatment with anticonvulsants and mental illness or psychiatric symptoms that impede activities of daily life.
- Patients who are incapable of taking oral agents.
- Patients with a history of allergy to study drugs or similar compounds.
- Breastfeeding women, pregnant women or patients not willing to use contraception.
- Patients deemed ineligible for the study by the investigator (eg, patients who are unable to maintain medication adherence or who may experience difficulty using electronic devices).

Recruitment, randomisation, masking and follow-up

Recruitment
Eligible patients satisfying the screening inclusion and exclusion criteria will be invited to participate in the study by site investigators.

Randomisation
Physicians will introduce the trial to patients. On enrollment and after providing informed consent, eligible patients will be randomly assigned to receive either DEX on day 1–4 or DEX on day 1 with placebo on days 2–4 as part of prophylactic antiemetic therapy. Randomisation is centrally performed by random allocation modules of electronic data captures (EDC) using the minimisation method with balancing prognostic factors for age (<60 years ≥60 years), sex, CDDP dose level (≥70 mg/m² vs <70 mg/m²) and institution.

Masking
Patients and clinicians responsible for treatment will be blinded to administration of DEX or placebo. Only an unblinded pharmacist who prepares the study drug, but is not involved in patient care, will know the assignment and outcome. All study drugs will be prepared by this pharmacist. As a rule, no data will be disclosed until fixed. However, during the trial period, when it is considered necessary to know the details of the trial drug to ensure participant safety, such as for serious adverse events, the study representative and study secretariat will make an inquiry to and discuss the need for disclosure with the Efficacy and Safety Evaluation Committee. When disclosure is deemed necessary as a result of this consultation, the details will be communicated to the study secretariat, and the details of the trial drug will be disclosed.

Data management, central monitoring and auditing
The data centre is located in the Department of Clinical Trial Data Management, Graduate School of Medicine, Tokyo University, Tokyo, Japan. Enrolment, randomisation, data collection and monitoring will be performed using EDC system Viedoc 4 and Viedoc me (Viedoc Technologies). Data entry to the electronic case report form is performed by investigators using EDC at each site. Patient-reported outcome (PRO) data are collected electronically from patients through an electronic tablet device. No personally identifiable information is entered into the EDC, and the data centre does not collect personal information. The central monitoring will be conducted by the data centre, and monthly and semi-annually monitoring reports will be disseminated to investigators to inform about the trial progress and discuss data quality-related issues. The protocol review committee and independent Data Monitoring Committee will assess the protocol amendments, serious adverse events reports and monitoring reports and provide any necessary recommendation to investigators. Auditing will be conducted as necessary in this study.

Harms
Investigators must record all adverse events in the medical records and web systems. The Common Terminology Criteria for Adverse Events (CTCAE, V.4.0) will be used to grade each adverse event. In conjunction with the CTCAE to grade adverse events, the PRO-CTCAE will be administered to patients for their completion to complement information about subjective symptoms. All adverse events are to be followed up continually during the course of treatment. All severe adverse events must be reported to the institutional review board (IRB) and reported to investigators in all sites and discussed through a mail. Patients who are enrolled into the study will be
treated by the healthcare services that are provided by their health insurance.

Treatment
All patients receive Palo (0.75 mg intravenous infusion on day 1 at 30 min before the start of chemotherapy), NKI-RA (aprepitant 125 mg oral administration on day 1 and 80 mg on days 2 and 3 or fosaprepitant 150 mg intravenous infusion on day 1 at 1 hour before the start of chemotherapy) and OLZ (5 mg oral administration on days 1–4 after dinner). DEX is administered as follows: patients in both arms receive DEX 9.9 mg intravenous infusion on day 1; patients receive DEX 6.6 mg or placebo intravenous infusion on days 2–4. When using fosaprepitant, the dose level is increased on days 3 and 4 due to interaction with DEX up to day 2, therefore, patients receive an intravenous DEX 13.2 mg or placebo on days 3 and 4. Patients were allowed to take rescue medication throughout the study period for nausea or vomiting, if necessary. The choice of recommended rescue is determined by each investigator from among prochlorperazine, metoclopramide, domperidone, chlorpheniramine, alprazolam, lorazepam and haloperidol.

Study endpoints
The primary endpoint is CR rate (no emesis and no rescue medications) during the delayed phase (24–120 hours post-CDDP administration). Secondary endpoints are as follows: (1) CR rate during the acute phase (24 hours post-CDDP administration) and the overall phase (120 hours post-CDDP administration), (2) complete control (no emesis, no rescue use and no significant nausea) rate, (3) total control (no emesis, no rescue use and no nausea) rate (4) no emesis rate and no nausea rate in the overall, acute and delayed phase, (5) time to treatment failure (ie, time to first emesis or using rescue, whichever occurred first) and (6) severity of nausea during the overall phase. Adverse events are associated with antiemetic therapy (CTCAE V.4.0 Japanese Clinical Oncology Group (JCOG) version and the PRO-CTCAE V.1.0.).

Outcome assessments
Figure 1 provides details of the schedule of enrolment, interventions and assessments.

Presence of emesis and severity of nausea will be assessed by patients using a 2-point categorical scale and 11-point numerical rating scale (NRS), respectively. Significant nausea is defined as 3 points or greater on the NRS. The use of rescue medications will be assessed by pharmacists.

Adverse events will be evaluated according to the CTCAE V.4.0 (JCOG) version, and the PRO-CTCAE V.1.0. The Japanese version of PRO-CTCAE is linguistically and psychometrically validated.21 22 QOL will be assessed by the European Organization for Research and Treatment of Cancer Quality of Life Questionnaire C30 (EORTC QLQ-C30) V.3 that is also validated in the Japanese version.23

Patients are asked to assess QOL before CDDP administration and on day 8, with emesis and nausea assessed the PRO-CTCAE every 24 hours until 120 hours after CDDP administration. The data from the PRO-CTCAE are assessed electronically using a tablet device in the hospital setting, except for QOL on day 8, which is assessed on paper-based questionnaire at home. The PRO-CTCAE data will be not reviewed by the site investigators during the protocol treatment.

Statistical analysis
Sample size calculation is based on an analysis of the primary endpoint. In previous studies with OLZ added to conventional antiemetic treatment for CDDP,20 24 25 the delayed phase CR rate ranged from 75% to 85%, therefore we expect that CR rate in the delayed phase would be 75% in both arms. The non-inferiority margin is set at -15.0%. Two hundred sixty-two patients are required for at least 80% power to confirm non-inferiority at a one-sided significance level of 2.5%. After considering the possibility of attrition, we set our final required sample size of 280. Point estimates and CIs for the CR rate will be calculated and will be compared between groups by using the Mantel-Haenszel test with adjustment for allocation factors. Interim analysis is not planned. We will use a full analysis set. It consists of the registered participant population who received at least a part of the protocol treatment; however, participants who were deemed as ineligible for the study after registration and those who were not administered CDDP-based regimens will be excluded from the analysis set. For the primary analysis, we will impute non-CR for missing primary endpoints.

Patient and public involvement
Patients and/or public were not involved in the design of this study.

Ethics and dissemination
All patients will be required to provide written informed consent (see online supplemental file 1). The study will be performed in accordance with the Declaration of Helsinki and Ethical Guidelines for Medical and Health Research Involving Human Subjects published by Japan’s Ministry of Education, Science and Technology and the Ministry of Health, Labour, and Welfare and the modified Act on the Protection of Personal Information. This protocol was approved by the Ethics Committee (approval ID 4035) of St. Marianna University School of Medicine on July 27, 2018. The protocol was approved by the IRB at each study site (Showa University Northern Yokohama Hospital, Yokohama Rosai Hospital, Nippon Medical School Musashi Kosugi Hospital, Aichi Cancer Center, Gifu University, Kitasato University, and Shizuoka Cancer Center). This trial has been registered at the University Hospital Medical Information Network (UMIN) Clinical Trials Registry. Modifications in the study protocol will be communicated to the IRB at each study site as well as to the protocol review committee. Each Ethics Committee or IRB will revise informed consent
ENROLMENT:
- Eligibility screen
- History and physical
- ECOG PS
- Laboratory studies
- Informed consent

Enrolment

INTERVENTIONS:
- Fosaprepitant (APR) administration
  - X
- PALO administration
  - X
- OLZ administration
  - X X X X
- DEX administration
  - X
- DEX or placebo administration
  - X X X

ASSESSMENTS:
- Risk factor
  - X
- PRO-CTCAE
  - X X X X X X
- QOL
  - X
- Presence of vomiting
  - X X X X X X
- Nausea (NRS)
  - X X X X X X
- Use of rescue medication
  - X X X X X X
- CTCAE v4.0-JCOG
  - X X X X X X

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Figure 1 The schedule of enrolment, interventions and assessments. APR, aprepitant; CTCAE, common terminology criteria for adverse events; DEX, dexamethasone; ECOG, Eastern Cooperative Oncology Group; NRS, numerical rating scale; OLZ, olanzapine; Palo, palonosetron; QOL, quality of life.

Materials given to participants and adapt the informed consent according to their own institution’s guidelines. The main result will be presented at an international conference and published in an English journal. Authorship will be ascribed in accordance with the International Committee of Medical Journal Editors guidance.

Access to data
Only clinical data managers at the central data centre have access to collected data through the EDC system.
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Contributors  HM, NI and TEN contributed to the trial conception and are the principal investigators. HM, NI, TK, TM, TY and TEN participated in the design of the study. TY played a primary role in designing statistical analysis. TK and TM played a primary role in designing the data management approach. HM, NI, KS, KH, HI, YO, YI, HA, HM, NH, MS, CK, SN, HI, AT and TT have carried out recruitment and collected the data. Data analysis and interpretation will be conducted by HM, NI, TK, TM, TY and TEN. HM and NI wrote the first draft of the manuscript. All authors have read, approved the paper and meet the criteria for authorship as established by the International Committee of Medical Journal Editors.

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Competing interests  NI has received honoraria from Takeda Pharma Co., Ltd, Eli Lilly, Ono, Pharma Co., Ltd, and Daiichi Sankyo Company. Author HA has received grant from Taiho, Chugai and Nippon Kayaku; personal fees from Novartis, Sanofi, Ono, Kyowa Kirin and Takeda. TY has received grant and personal fees from Ono Pharmaceutical Co., Ltd. TEN has received grant and personal fee from Taiho Pharmaceutical Co., Chugai Pharmaceutical Co., Takeda Pharmaceutical Co., Sanofi K.K., Daiichi Sankyo Co., Eli Lilly Japan K.K., Nippon Kayaku Co., Ono Pharmaceutical Co. and MSD K.K.; personal fees from Mochida Pharmaceutical, Celltrion Healthcare Japan, Mercck Serono Co., Sawai Pharmaceutical Co., Bayer Yakuhin, Bristol-Myers Squibb, Teijin Pharma, Pfizer Japan Inc., Novartis Japan, Yakult Honsha Co. and Nipco Co; grant from Astellas Pharma Inc., Sumitomo Dainippon Pharma Co., Eisai Co and Solasia Pharma K.K. The other authors have declared no conflicts of interest.

Patient consent for publication  Not required.

Provenance and peer review  Not commissioned; externally peer reviewed.

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REFERENCES  
Consent Form

St. Marianna University School of Medicine Hospital

Dear Hospital Director:

Title of the study: Randomized noninferiority phase III trial comparing dexamethasone on day 1 with dexamethasone on day 1-4 combined with neurokinin-1 receptor antagonist, palonosetron, and olanzapine (5 mg) in patients receiving cisplatin-based chemotherapy

Description
1. Introduction about clinical trials.
2. The purpose of this study.
3. The method of this study.
4. The expected duration of participation in this study.
5. The expected number of participants in this study.
6. The expected effects of the medication under investigation and the possible side effects.
7. If you do not use this medication, other treatment options are available.
8. Participation in this study is voluntary.
9. If you agree to participate in the study, it requires observation of your first 5 days of chemotherapy.
10. Potential harmful effects to your health, which may occur during this study.
11. The chance that we may stop using this medication.
12. If you participate in this study, your medical records and other information may be examined during and after this study.
13. Your identity will not be revealed if the results of this study are made public.
14. We will keep you informed regarding this medication.
15. Your cost burden.
16. Information regarding the bioethics committee.
17. The institution participating in the study.
18. Information about your physician and consultation.

[Patient’s signature line]
I have been fully informed of the above information, have received the letter of consent, and fully understand the details of this study. I voluntarily consent to participate in this study.

Date of consent: ________________________________

Patient’s name: (Signed) ______________________________________

[Physician’s signature line]
I have fully briefed the patient on this clinical trial.

Date of presentation: ____________________________________________

Affiliation: ____________________________________________________

Name: (Signed) ________________________________________________