


BMJ Open Pharmacokinetics and complementary evaluation system-based guidance on prophylaxis of paediatric patients with haemophilia A in China with Kovaltry: protocol of the LEAP study

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To cite: Kun H, Xu W, Zhou M, *et al.* Pharmacokinetics and complementary evaluation system-based guidance on prophylaxis of paediatric patients with haemophilia A in China with Kovaltry: protocol of the LEAP study. *BMJ Open* 2021;**11**:e048432. doi:10.1136/bmjopen-2020-048432

► Prepublication history and additional supplemental material for this paper are available online. To view these files, please visit the journal online (<http://dx.doi.org/10.1136/bmjopen-2020-048432>).

Received 27 December 2020
Accepted 14 June 2021



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ABSTRACT

Introduction Haemophilia A is a rare inherited bleeding disease caused by the deficiency of coagulation factor VIII (FVIII). The main treatment protocol is to administer regular exogenous FVIII concentrate infusions. With the discovery of variability in individualised pharmacokinetics (PK) and bleeding phenotype, the previous weight-based approach needs to be replaced by more advanced PK-tailored prophylaxis with an accurate evaluation system. In this study, we combine individualised PK profiles and a complementary evaluation system to guide prophylaxis in paediatric patients with haemophilia A.

Methods and analysis This is a single-centre, prospective single-arm study. The aim of this study is to assess the effectiveness of a new strategy combining PK and a complementary evaluation system to treat haemophilia A in Chinese paediatric patients. Sixty paediatric patients with haemophilia will be recruited. After PK testing, they will receive a PK-guided stepup prophylaxis in the next 2 years. The dosing regimen will be determined according to individualised PK profiles and complementary evaluation findings. Related indicators at the end of the study will be compared with the values at treatment initiation to examine the effectiveness of this new strategy. The demographic data of the investigated patients will be summarised by descriptive statistics. Quantitative data will be described by summary statistics, including arithmetic median, range, mean and arithmetic SD. Analyses will use t-test to compare indicators such as bleeding rate and imaging score at both ends of the study as well as during follow-up.

Ethics and dissemination The study has been approved by the Ethics Committee of Beijing Children's Hospital (Number 2020-Z-095). The findings will be presented at international meetings such as World Federation of Hemophilia World Congress. Related manuscripts will be submitted to peer-review journals such as *Blood* and *Hemophilia*.

Trial registration number ChiCTR2000037821; Pre-results.

INTRODUCTION

Haemophilia A is an X-linked inherited bleeding disorder due to the deficiency of

Strengths and limitations of this study

- This study will be the first to combine pharmacokinetic and complementary evaluation in haemophilia therapy.
- The complementary evaluation could provide better joint protection by detecting preclinical lesions.
- The small sample size may limit statistical power for further exploratory analyses.

coagulation factor VIII (FVIII). Patients with haemophilia have spontaneous bleeds in muscles or joints, which could cause joint dysfunction or even death. The main treatment option for haemophilia is to administer regular exogenous FVIII infusions. Compared with on-demand therapy, prophylaxis has been considered an optimised therapy regimen to help patients live a normal life, with enhanced ability to decrease bleeds and maintain the function of joints.¹ Prophylaxis is considered the standard treatment of haemophilia in paediatric patients and should be started as soon as possible once prophylaxis is proposed.²

The first prophylaxis regimen was proposed in Sweden and is also known as 'standard prophylaxis'.³ It aims to keep the trough FVIII level of patients with haemophilia A above 1 IU/dL by giving them regular exogenous FVIII concentrate injections (20–40 IU/kg, three times per week or every other day). The experience of the Malmo protocol in more than 50 years has revealed its clinical effectiveness in treating patients with haemophilia.⁴ However, the Malmo protocol has not been widely used in other regions due to the massive consumption of and insufficient access to FVIII concentrate as well as the heavy burden of frequent intravenous

injections. Therefore, other prophylaxis protocols have been developed, including the middle dose prophylaxis in the Netherlands (15–25 IU/kg, 2–3 times per week) and the stepup prophylaxis proposed by Canada.^{5 6} In some developing countries, such as China and India, even low-dose prophylaxis was shown to greatly decrease bleeding compared with on-demand therapy.⁷

Currently, prophylactic regimens are determined by a standard weight-based approach, which may cause underdosage or overdosage because of the variability of both FVIII pharmacokinetic (PK) profiles and bleed phenotypes among different patients with haemophilia. According to Chen *et al* FVIII's half-life time varies from 5.52 to 20.02 hour, while in vivo recovery varies from 1.2 IU/kg to 3 IU/kg.⁸ Other studies also confirmed great individual variability in PK profile.^{9 10} Thus, single weight-based prophylaxis may cause either extra bleeds and joint disfunction due to insufficient treatment or unnecessary FVIII concentrate waste with overdosage. Therefore, it has been recommended that individualised PK profiles should be employed for determining the patient's dose and frequency of routine prophylaxis. In the past, it was hard to obtain individualised PK profiles because of the heavy burden of up to 10 time points after a long washout period and single-dose infusion.¹¹ With the application of the Bayes approach to population PK (popPK) in haemophilia, it is currently possible to use blood samples collected at only 2–3 time points to determine individualised PK profiles.¹¹ According to Iorio *et al*, the popPK method is a practical and accurate way to predict individualised PK profiles.¹² Previous studies have revealed the advantages of PK-tailored prophylaxis in haemophilia treatment.¹³ In addition, online PK dosing tools such as Web Accessible Population Pharmacokinetics (WAPPS) have been recommended by official organisations to guide routine prophylaxis according to individualised PK profiles.¹¹

Besides variability in PK profiles, different bleed phenotypes among patients also need to be taken into consideration in routine therapy. Although Collins *et al* clearly demonstrated that break-through bleeds in prophylaxis are correlated to weekly time spent with low FVIII levels, some patients with high-trough FVIII levels in daily prophylaxis still suffer from bleeds, especially those with target joints.¹⁴ According to the sports guidelines for haemophilia, patients with target joints need to keep higher FVIII levels in the same sport compared with those without joint disfunction. In a Dutch study involving more than 400 patients with haemophilia, it seemed that only trough FVIII levels reached 12 IU/dL should the number of target joints decrease to zero.¹⁵ The target trough FVIII level to reach the goal of zero bleed was 15 IU/dL in another study.¹⁶ Besides the joint state, Den Uijl *et al* also suggested that multiple targets should be considered in determining the routine prophylaxis regimen, including physical activity, the quality of life and cost-effectiveness.¹⁶ The haemophilia care team of Beijing Children's Hospital started a study named CHIPS (Chinese Hemophilia

Individualized Prophylaxis Study) in 2016 to explore an evaluation system for paediatric patients with haemophilia, which includes multiple targets such as joint structure assessed by MRI and ultrasound (US) scores, joint function evaluated by Haemophilia Joint Health Score (HJHS) scores, the quality of life assessed by Canadian Hemophilia Outcomes-Kids Life Assessment Tool scores and other aims and scaling scores.¹⁷ In this study, some patients could keep the bleeding rate at 0 in step 1, while others still suffered from frequent bleeds even in step 4, which indicated the variability of bleed phenotype leads to a difference in target trough FVIII levels in routine prophylaxis.

Although some products involving new mechanisms to treat haemophilia are available, most patients around the world are still taking FVIII concentrate for routine prophylaxis, and this situation would not change for a long time.² How to use PK data and a complementary evaluation system to individualise prophylaxis in patients with haemophilia, achieving better clinical outcomes and reduced cost, remains a vital question that needs to be addressed urgently.

STUDY OBJECTIVES

Primary objectives

1. To evaluate the effect of PK-based and complementary evaluation system-based instructions for prophylaxis in paediatric patients with haemophilia A (according to US and/or MRI findings).
2. To establish a popPK model for Kovaltry suitable for paediatric patients with haemophilia A in China.

Secondary objectives

1. To study the efficacy and safety of the prophylactic regimen under the guidance of PK and complementary evaluation system.
2. To evaluate the PK parameters of paediatric haemophilia A patients in China administered Kovaltry products in China.

METHODS AND ANALYSIS

Ethics and dissemination

The study was approved by the ethics committee of Beijing Children's Hospital (Number 2020-Z-095). Written informed consent was obtained from each enrolled patient and their legally authorised guardians. The SPIRIT list of this study would be available as supplementary files.

Study design

This is a multicentre, prospective single-arm study, including two stages from January 2021 to January 2024.

Stage I is the popPK period lasting for 6 months. The enrolled paediatric patients will be treated with Kovaltry according to current clinical situation, and the therapeutic regimen and bleeding situation will be recorded. PK indicators will be measured comprehensively.

PopPK model building and verification

At this stage, after obtaining the individual PK information of paediatric patients, the PK and personal information of 30 paediatric patients (aged 1–18 years) will be included in WAPPS to generate the Kovaltry popPK model including the data of paediatric patients in China; meanwhile, the individual PK information of another 30 paediatric patients in China not involved in modelling will be included for external verification to ensure the accuracy and availability of the model. Considering the balanced distribution of paediatric patients in various age groups, age distribution for modelling and verification will be 1–6, 6–12 and 12–18 years old. All enrolled paediatric patients should undergo PK testing prior to the trial period. A washout period of at least 72 hours will be retained before PK testing. A single dose of 50 IU/kg coagulation FVIII concentrate (Kovaltry, BAY81-8973) will be infused, with blood samples collected at different time points before and after infusion to determine FVIII concentration. Blood samples will be taken within half an hour before infusion, and at 1 hour, 3 hours, 9 hours, 24 hours, 48 hours and 72 hours after infusion, centrifuged and tested.¹⁸ PK parameters will be obtained through the WAPPS-Hemo team.¹²

Data collection

After enrolment, patients' data on prophylaxis with Kovaltry in the first 6 months will be collected as baseline data in this study.

Stage II is the clinical stage lasting for 2 years. Patients will receive joint assessment and trough FVIII level test every 3 months and PK monitoring every 6 months.

All eligible patients will receive a dose-escalation prophylactic regimen guided by the results of PK and a

complementary evaluation system, including four steps (the first step would be decided according to patients' individualised trough FVIII level in their routine prophylaxis):

Step 1: Maintained trough FVIII concentration=1–2 IU/dL.

Step 2: Maintained trough FVIII concentration=2–3 IU/dL.

Step 3: Maintained trough FVIII concentration=3–4 IU/dL.

Step 4: Maintained trough FVIII concentration=4–5 IU/dL.

Step 5: Maintained trough FVIII concentration >5 IU/dL.

In the above steps, the specific dose and frequency of dosing are not stipulated. The investigators will jointly decide a therapeutic regimen with the subject based on comprehensive assessments and instructions of WAPPS-Hemo PK, the patient's needs for quality of life and other specific conditions.

Prophylactic administration in all eligible patients will be initially (at the seventh month) evaluated as 'insufficient' according to specific criteria (table 1), and the trough FVIII levels of their current prophylaxis will be upgraded to the trough concentrations at the corresponding time. This protocol aims to combine PK and a complementary evaluation system to instruct patients to receive prophylaxis and help them further control bleeding, protecting joint function and improving the quality of life.

Study population

Inclusion criteria

1. Severe haemophilia A (FVIII: C<1%), aged 1–18 years.

Table 1 Escalation criteria determined by the complementary evaluation system

	Parameter	Time	Description	Score
Bleeding (1–18 years)	Bleeding	Every 3 months	No bleeding	0
			1 bleed	+1
			≥2 bleeds	+2
Clinical imaging (1–18 years)	HEAD-US	Every 3 months	No change or improved	0
			HEAD-US scores+1 or new significant haematoma/ joint haematoma/haemosiderosis	+1
			HEAD-US scores+2 or new severe haematoma/ joint haematoma/haemosiderosis	+2
Joint function (4–18 years)	HJHS	Every 3 months	No change or improved	0
			Single joint score increased by ≥1	+1
			Single joint score increased by ≥2	+2
Motion (7–18 years)	FISH	Every 6 months	Total score decreased by <2	0
			Total score decreased by 2–4	+1
			Total score decreased by ≥4	+2

Evaluation:<2 points:maintain the prophylactic dose;≥2 points: increase the prophylactic dose into the next step.

FISH, functional independence score in haemophilia; HEAD-US, Hemophilia Early Arthropathy Dection with Ultrasound; HJHS, Haemophilia Joint Health Score.

2. A history of knee, elbow or ankle bleeding.
3. >50 exposure days (calculated from previous treatments with FVIII products).
4. No FVIII inhibitors at enrolment.
5. Regular clinical visits and medical records available.
6. Informed consent from the legal guardians of patients before enrolment.
7. Prophylaxis with Kovaltry being administered at the time of enrolment and baseline data on prophylaxis with Kovaltry for at least 6 months prior to phase I available.

Exclusion criteria

1. Other haemorrhagic diseases such as von Willebrand disease (VWD).
2. Generation of FVIII inhibitors: >0.6BU (confirmed by two separate tests).
3. A previous history of inhibitors and presence of FVIII inhibitor at any time in the study period.
4. Planning to participate (or previous involvement) in other Kovaltry-related studies, other interventional studies or any studies expected to affect the study protocol.
5. Using other FVIII concentrates for routine prophylaxis.

Sample size

According to the guidance of the WAPPS team and the number of potential patients available in our centre, the PK data of 60 paediatric patients with haemophilia A would be sufficient for this study. In addition, due to the novelty of this study, the sample size could not be estimated through previous studies.

Study endpoints and outcomes measures

Primary endpoints

1. Percentages of MRI/US scores of joints improved/unchanged from baseline.
2. A valid Kovaltry popPK model established for patients with paediatric haemophilia A in China

Second endpoints

1. Annual bleeding rate, annual joint bleeding rate and annual target joint bleeding rate.
2. Bleeding rates will be calculated according to the routine electronic record of patients.
3. Joint function (HJHS).
4. Joint structure (X-ray Pettersson score).
5. Motion (functional independence score in haemophilia >7 years of age).
6. Quality of life.
7. Consumption and therapeutic dose regimen of Kovaltry.
8. Assessment of family disease burden.
9. Treatment compliance and reasons for noncompliance of patients in various age groups.

Escalation criteria

According to the criteria detailed in [table 1](#), the prophylactic dose would increase into the next step with a score ≥ 2 points. Detailed variables and evaluation methods are described in [table 2](#).

The study flowchart is depicted in [figure 1](#).

Statistical analysis plan

Descriptive analysis of all variables will be performed by appropriate statistical methods. Categorical variables will be analysed using frequency distribution tables (absolute and relative frequencies). Continuous variables will be analysed using sample statistics (ie, mean, SD, minimum,

Table 2 Variables and evaluation methods

Study outcomes	Variables and methods
Percentages of MRI/ultrasound scores of representative joints improved/unchanged from baseline at the end of the study.	MRI: IPSPG MRI score Ultrasound: HEAD-US score
ABR, AJBR and ATJBR	Annual bleeding and joint bleeding rates
X-ray outcome	Pettersson score
Joint function	HJHS
Motion	FISH (>7 years old)
Quality of life	CHO-KLAT score, China V.2.0
Percentage of patients remaining at each step of administration at the end of this study	Percentage of patients
FVIII consumption	Frequency and volume of FVIII infusion
Inhibitors	Incidence of inhibitors
Treatment compliance	Comparison of the actual infusion volume received with the individualised prevention protocol prescribed by physicians

ABR, annual bleeding rate; AJBR, annual joint bleeding rate; ATJBR, annual target joint bleeding rate; CHOKLAT, Canadian Hemophilia Outcomes-Kids Life Assessment Tool; FISH, functional independence score in haemophilia; HEAD-US, Hemophilia Early Arthropathy Decision with Ultrasound; HJHS, Haemophilia Joint Health Score; IPSPG, International Prophylaxis Study Group.

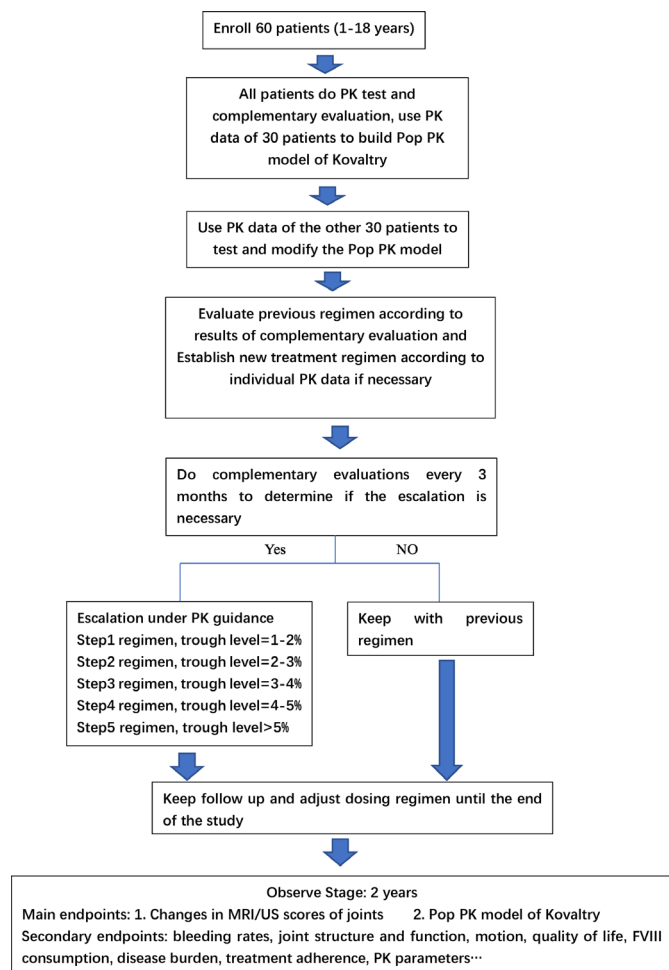


Figure 1 The study flowchart. FVIII, factorVIII; PK, pharmacokinetics.

median, quartile and maximum). Continuous variables will be described using absolute values from each time point and represented as changes from baseline (if applicable). The statistical package for social sciences (SPSS) software V.13.0 will be used for statistical processing. Student's t-test will be performed for the analysis of normally distributed data. The χ^2 test will be carried out for enumeration data. Non-normally distributed data will be analysed by the rank sum test. $p < 0.05$ will be considered statistically significant.

Ethics and dissemination

This study has been approved by the Ethics Committee of Beijing Children's Hospital (BCH). Informed consent will be obtained from all boys with severe haemophilia A and their legally authorised guardians. The results will be organised into manuscripts and submitted to peer-review journals as well as international academic meetings. The original data will be stored at Beijing Children's Hospital, and disclosure will only be available on reasonable request by e-mail to the corresponding authors. The main

finding will be open to all participants and the Haemophilia Society.

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Contributors HK proposed the study, defended during ethical review and wrote the manuscript. RW and ZC designed the study, applied for funding and reviewed the manuscript. WX, MZ, XL, ZX, YF and CL discussed on the planning, conduct and reporting of this study and reviewed the manuscript.

Funding The current work was in part supported by grants from Research on the application of clinical characteristics of the Beijing Municipal Science and Technology Commission (code Z181100001718182) and Bayer Health Company (grant number 20006429).

Competing interests None declared.

Patient consent for publication Not required.

Provenance and peer review Not commissioned; externally peer reviewed.

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