Effect of vitamin K\textsubscript{2} on the anticoagulant activity of warfarin during the perioperative period of catheter ablation: Population analysis of retrospective clinical data

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Abstract

Background: Catheter ablation is a non-medication therapy for atrial fibrillation, and during the procedure, warfarin is withdrawn in the preoperative period to prevent the risk of bleeding. In case of emergency, vitamin K\textsubscript{2} can be intravenously administered to antagonize the anticoagulant activity of warfarin. The aims of this study were to conduct population pharmacokinetic/pharmacodynamic modeling for retrospective clinical data and to investigate the effect of vitamin K\textsubscript{2} on the anticoagulant activity of warfarin in the perioperative period of catheter ablation.

Methods: A total of 579 international normalized ratio (INR) values of prothrombin time from 100 patients were analyzed using the nonlinear mixed-effects modeling program NONMEM. A 1-compartment model was adapted to the pharmacokinetics of warfarin and vitamin K\textsubscript{2}, and the indirect response model was used to investigate the relationship between plasma concentration and the pharmacodynamic response of warfarin and vitamin K\textsubscript{2}. Since no plasma concentration data for warfarin and vitamin K\textsubscript{2} were available, 3 literally available pharmacokinetic parameters were used to simultaneously estimate 1 pharmacokinetic parameter and 5 pharmacodynamic parameters.

Results: The population parameters obtained not only successfully explained the observed INR values, but also indicated an increase in sensitivity to warfarin in patients with reduced renal function. Simulations using these parameters indicated that vitamin K\textsubscript{2} administration of more than 20 mg caused a slight dose-dependent decrease in INR on the day of catheter ablation and a delayed INR elevation after warfarin re-initiation.

Conclusions: A pharmacokinetic/pharmacodynamic model was successfully built to explain the retrospective INR data during catheter ablation. Simulation studies suggest that vitamin K\textsubscript{2} should be administered with care and that more than 20 mg is unnecessary in the preoperative period of catheter ablation.

Keywords: Warfarin, Vitamin K\textsubscript{2}, Pharmacodynamics, INR, NONMEM, Catheter ablation

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Background
Atrial fibrillation is the most common sustained cardiac arrhythmia and a major cause of stroke [1, 2]. In order to prevent stroke, an anticoagulant drug, warfarin, is usually used since aspirin was proven ineffective in retrospective analyses [3]. The anticoagulant effect of warfarin does not always correlate with its dose, and polymorphisms in cytochrome P450 (CYP) 2C9 and vitamin K epoxide reductase complex subunit 1 (VKORC1) genes have been proven to influence interindividual variability in the optimal doses, in addition to patients’ primary diseases and characteristics such as age or ethnicity [4, 5]. In Japanese patients, warfarin dose adjustments based on their prothrombin time, an international normalized ratio (INR) of 1.6–2.6 (age ≥70 years) or 2.0–3.0 (age <70 years), are recommended for effective therapy to avoid life-threatening bleeding [6, 7]. When hemorrhagic complications occur, warfarin withdrawal is required and vitamin K₂ or fresh frozen plasma administration is recommended [8–10].

In atrial fibrillation treatment, antiarrhythmic agents are often used, while catheter ablation is also an available option as a non-medication therapy [2]. When catheter ablation, an invasive procedure for complete cure of atrial fibrillation, is selected, anticoagulant therapy with warfarin is withdrawn in the preoperative period to prevent the risk of bleeding, although catheter ablation is sometimes performed in periprocedural therapeutic anticoagulation with warfarin if possible. In some patients, discontinuation of warfarin is not sufficient to lower the INR to the required level before catheter ablation. In such cases, vitamin K₂ is intravenously administered to antagonize the anticoagulant activity of warfarin resulting in prompt recovery of INR to a safe level. Some reports have mentioned the use of pharmacokinetic/pharmacodynamic models for an anticoagulant drug and have conducted population analyses; however, only warfarin was investigated using these models [11, 12]. The effect of vitamin K₂ dose on controlling the anticoagulant activity of warfarin during the perioperative period of catheter ablation has not yet been reported. The aims of this study are to build a population pharmacokinetic/pharmacodynamic model not only for warfarin, but also for vitamin K₂, by using routine clinical data of patients who had been diagnosed with atrial fibrillation and received a catheter ablation, and to obtain information on the optimal vitamin K₂ dose in the preoperative period before catheter ablation.

Methods
Patients and data studied
We retrospectively collected data from patients who have had a catheter ablation for atrial fibrillation at the Department of Cardiovascular Medicine, Kyoto University Hospital from January to December in 2008. During this period, 126 Japanese patients underwent catheter ablation, and 111 of these patients were treated with warfarin on the day of admission. A total of 100 patients whose INR values were between 1.0 and 3.0 in the hospitalization period were included in this study. We used 579 INR values obtained from 100 patients during the perioperative period. Clinical laboratory data and medication history for the patients studied were collected from electrical medical records. No patients were taking any medications that may have clinically significantly altered the pharmacokinetics of warfarin, except 4 patients with amiodaron and 1 patient with bucolone [13, 14].

Pharmacokinetic/pharmacodynamic model building
A 1-compartment model was adopted to the pharmacokinetics of warfarin and vitamin K₂ as follows (Fig. 1):

\[ \frac{d(Cp_1 \cdot Vd_1)}{dt} = -k_{10} \cdot (Cp_1 \cdot Vd_1) \]  
\[ \frac{d(Cp_3 \cdot Vd_3)}{dt} = -k_{30} \cdot (Cp_3 \cdot Vd_3) \]

where \( Cp_1 \) and \( Cp_3 \) represent the plasma concentration of warfarin and vitamin K₂, respectively; and \( Vd_1 \) and \( Vd_3 \) represent the distribution volume; and \( k_{10} \) and \( k_{30} \) represent the elimination rate constant for each drug, respectively. Since no plasma concentration data were available for warfarin and vitamin K₂, and INR values were the available data for this study, reported pharmacokinetic parameters for warfarin in Japanese patients [11] and the distribution volume for vitamin K₂ in the product information (Eisai Co., Ltd., Tokyo, Japan) were used in the analysis: \( k_{10} = 0.0129 \) (1/h), \( Vd_1 = 0.183 \) (L/kg) and \( Vd_3 = 0.051 \) (L/kg). Therefore, \( k_{30} \) was the only pharmacokinetic parameter to be estimated in this analysis.

The indirect response model was used to explain the relationship between plasma concentration and pharmacodynamic response of warfarin and vitamin K₂ [11, 14–16]. In this model, the amount of clotting factors was described using a zero-order synthesis rate constant (\( k_s \)) and a first-order degradation rate constant (\( k_d \)) under the hypothesis that coagulant activity was proportional to the amount of clotting factors (Fig. 1). Since both warfarin and vitamin K₂ target the same enzyme that is responsible for clotting factor synthesis [17], the maximum effect models were adopted to describe stimulatory and inhibitory activities of these drugs, respectively, as follows:

\[ \frac{dT}{dt} = k_s \cdot (1 - Cp_1/(Cp_1 + IC_{50})) + E_{max} \cdot Cp_3/(Cp_3 + EC_{50}) - k_d \cdot TT \]

where \( k_s, k_d, IC_{50}, E_{max}, \) and \( EC_{50} \) represent synthesis rate constant (%/h), degradation rate constant (1/h), 50 % inhibitory concentration of warfarin (µg/mL),...
maximum effect of vitamin $K_2$ (no unit), and 50 % effective concentration of vitamin $K_2$ ($\mu$g/mL) were used, respectively (Fig. 1). The Hill coefficient used in the previous study [11] was not included in the present model to simplify the pharmacodynamic model. Since only INR values were collected in this study, thrombotest ($TT$) values were calculated according to Equation 4 using values provided from literature [18]:

$$TT(\%) = 23.77 \times \frac{INR}{(INR-0.8085) - 0.09807 \times INR-23.04} \quad (4)$$

Since the predicted values were outputted by the non-linear mixed-effects modeling program (NONMEM) [19] using $TT$ values, these were then converted into INR values when necessary by solving the quadratic equation obtained from Equation 4.

The population pharmacokinetic and pharmacodynamic analysis was performed using the NONMEM (version VI), using the first-order conditional estimation method [19]. In this study, exponential error models for both inter- and intraindividual variability were chosen as follows:

$$P_{ij} = P_{pop.i} \times \exp(\eta_{ij}) \quad (5)$$

$$TT_{jk} = TT_{\text{pop}.i} \times \exp(\epsilon_{jk}) \quad (6)$$

where $P_{ij}$ is the $i$-th individual pharmacokinetic or pharmacodynamic parameter for patient $j$; $P_{pop.i}$ is the $i$-th population mean parameter; and $\eta_{ij}$ is the individual random perturbation from the population mean parameter that is distributed with a mean of zero and variance $\omega^2$. $TT_{jk}$ is the observed $TT$ value at time $k$ for patient $j$; $TT_{\text{pop}.i}$ is the corresponding predicted $TT$ value; and $\epsilon_{jk}$ represents the independent identically distributed error with a mean of zero and variance of $\sigma^2$ for the $TT$ value.

The number of $\eta$ used in the model was determined by the method of minimum Akaike information criterion (AIC) estimation [20].

$$AIC = OBJ + 2M \quad (7)$$

where $OBJ$ is the objective function values calculated using the NONMEM and $M$ is the number of independently adjusted parameters within the model.

Next, the influence of renal function on each parameter was examined using Equation 8, by the forward selection method.

$$P_{pop.i} = P_{\text{pop}.i} \times \theta^{RF} \quad (8)$$

where $RF = 1$ if serum creatinine was higher than our in-hospital reference value, namely 1.1 mg/dL or higher for men, and 0.8 mg/dL or higher for women, otherwise $RF = 0$. $P_{\text{pop}.i}$ is the $i$-th population mean parameter in the patient whose serum creatinine is within our in-hospital reference value. The parameter set that had the smallest objective function value was selected, and the null hypothesis that $\theta$ was not statistically different from unity was examined using the likelihood ratio test. A difference of 7.88 in $OBJ$ with 1 degree of freedom was used to measure statistical significance ($P<0.005$ by the chi-squared distribution).

**Simulation for INR transition**

**A) Effect of vitamin $K_2$ dose**

Simulations were carried out using the obtained population mean parameters based on a typical patient whose
body weight was 50 kg with/without renal failure. The maintenance dose of warfarin was set to 3 mg/day (7 PM) and was stopped on day −1 (the day prior to the operation), and 5 mg/day was administered for 2 days after the operation as a loading dose, followed by a maintenance dose of 3 mg/day. Vitamin K₂ was administered at 20 mg 0, 1, 2, or 3-times every 4 hours after 4 PM on day −1 with the total dose administered ranging from 0 mg to 60 mg.

For quantitative evaluation, we obtained 4 parameters, namely ΔINR, 1st loading, 95% recovery, and INR/day. The ΔINR represents the difference in INR values between before warfarin withdrawal and before the loading dose; the 1st loading represents an INR increase after the first warfarin loading dose; and the 95% recovery represents the time needed for INR elevation in the postoperative period up to 95% of the preoperative steady state INR value. In addition, INR/day was calculated by dividing ΔINR by 95% recovery (day).

(B) Effect of warfarin dose
Simulations with various warfarin maintenance doses were conducted. As a maintenance dose, 3 to 6 mg of warfarin was administered and it was stopped on day −1 without vitamin K₂ administration. Warfarin (2 mg) was added to each maintenance dose as a loading dose, and it was administered for 2 days after the operation, followed by each maintenance dose. Cases where 20 to 60 mg of vitamin K₂ was administered were also simulated.

(C) Effect of interindividual variability
Simulations were also conducted using several parameter sets in which 1 of the mean parameters was altered using the interindividual variability (+ or − ω) from the population mean value. Warfarin and vitamin K₂ doses were set to 3 and 20 mg, respectively, in each simulation.

Results

Patients’ characteristics and INR transitions
Table 1 shows the characteristics of patients used in this study. Each patient received anticoagulant therapy of 1 to 7 mg/day of warfarin to prevent thromboembolic events. The median initial INR value on the day of admission was 1.76, and the median maintenance dose before hospitalization and the median loading doses of warfarin after the operation were 3 and 5 mg, respectively. To antagonize warfarin after its withdrawal in the preoperative period, a total of 20 to 70 mg of vitamin K₂, determined by the physician responsible, was intravenously administered to 76 patients before the operation. There were 4 patients with a total bilirubin concentration greater than our in-hospital reference value, but not substantially higher. Eight patients had an albumin concentration lower than our in-hospital reference values. Twenty-two patients had a serum creatinine concentration greater than our in-hospital reference value. Twenty-six patients had an estimated glomerular filtration rate from 0 mg to 60 mL/min/1.73 m². Figure 2 shows the INR transitions of each patient from day −5 to day 10, where the day of operation was day 0. The INR values decreased during the preoperative period and gradually increased again during the postoperative period.

Table 1 Patient characteristics

| Characteristics                        | Number or median (min-max) |
|----------------------------------------|----------------------------|
| Total number of patients (M/F)         | 100 (70/30)                |
| Age (years)                            | 64 (31–80)                 |
| Body weight (kg)                       | 63.8 (34.9–92.6)           |
| Initial INR                            | 1.76 (1.03–2.64)           |
| Warfarin maintenance dose (mg)         | 3.0 (1.0–7.0)              |
| Warfarin loading dose (mg)             | 5.0 (1.0–9.0)              |
| Number of patients treated with vitamin K₂ | 76                         |
| Total dosage of vitamin K₂ (mg)        | 40 (20–70)                 |
| 20 mg                                  | 19                         |
| 30 mg                                  | 2                          |
| 40 mg                                  | 35                         |
| 60 mg                                  | 19                         |
| 70 mg                                  | 1                          |
| Total bilirubin concentration (mg/dL)  | 0.7 (0.3–1.7)              |
| Serum albumin (g/dL)                   | 4.3 (3.6–5.0)              |
| Serum creatinine concentration (mg/dL) | 0.8 (0.5–9.6)              |
| Estimated glomerular filtration rate (mL/min/1.73 m²) | 67.7 (5–120) |

INR, prothrombin internationalized ratio

Model development
When interindividual variability was considered for all population pharmacokinetic/pharmacodynamic mean parameters (η = 6), AIC was 3398. To simplify the model in which only ηₖₐ and η₅₀ were included (η = 2), AIC was 3394, and was decreased to 3393 when another η for k₃₀ was included in the model (η = 3). Thus, the model with the minimum AIC value was adopted, which reflected the interindividual variability of kₚ, IC₅₀, and k₃₀.

Next, a search for covariates of population mean parameters was conducted using the forward selection method. When the effect of serum creatinine on each population mean parameter was examined, significant effects of renal function on kₚ, k₃₀, and IC₅₀ were observed (P < 0.005). Since the effect on IC₅₀ showed
the largest $-2$ log likelihood difference ($-2LLD$) of 27.2, this effect was incorporated into the second step. At the second step, additional effects of renal function on other parameters were examined, but no significant differences were observed ($-2LLD < 0.61$). We also examined the effect of renal function on the $IC_{50}$ using the value of estimated glomerular filtration rate, but the model fitting was better in the model using serum creatinine. Therefore, we chose the model in which only $IC_{50}$ was affected by renal function as follows:

$$d(TT)/dt = k_s \cdot \left(1 - Cp_1/(Cp_1 + IC_{50} \cdot \theta^{RF})\right) + E_{max} \cdot Cp_3/(Cp_3 + EC_{50}) - k_d \cdot TT$$

(9)

There were only 4 patients out of 100 patients whose total bilirubin concentration exceeded our in-hospital reference value, and those values were not remarkably high. Therefore, the effect of hepatic function on population mean parameters was not further examined. The anticoagulant effect of warfarin is generally considered to be associated with its unbound plasma concentration [10]. We examined the effect of serum albumin concentration on the $IC_{50}$ or $k_{10}$, but we could not obtain any significant effects.

Table 2 shows the final population mean parameters obtained and inter- and intraindividual variability. The interindividual variability for $k_s$, $IC_{50}$, and $k_{30}$ were 26.5 %, 37.9 %, and 41.4 %, respectively, and intraindividual variability was 28.2 % as a coefficient of variation (CV). In patients with decreased renal function, the $IC_{50}$ value was reduced to 61.4 % of those with normal renal function, suggesting enhanced sensitivity to warfarin.

Validity of population mean parameters

Figure 3 shows the plot of population or individual (post-hoc Bayesian) predicted versus observed $TT$. Although there was significant variability between the population predicted and observed $TT$, each plot individually predicted by the Bayesian method was closer to the unit line. The validation of the final population parameters was further confirmed by comparing the predicted $INR$ values versus observed $INR$ values (Fig. 4). Three typical patients were randomly selected each from 3 different groups classified by $INR$ values on the day of admission (high, median and low), and their predicted values were compared to the observed values. The time course profiles predicted by the Bayesian method were closer to the observed values than those predicted by the population mean parameters were, although there was still some discrepancy between these plots.

Effect of renal function on $INR$ transition

Figure 5 shows the simulation curves for the effect of renal function. In a patient with decreased renal function, the $INR$ value at a steady state rose from 1.65 in a patient with normal renal function to 1.99 with a maintenance dose of 3 mg/day (Fig. 5a). The $INR$ transitions of a patient with decreased renal function showed more dynamic changes with variable vitamin $K_2$ doses than those with normal renal function in the perioperative period. Table 3A shows the values from quantitative evaluation of Fig. 5. The $\Delta INR$ increased depending on the total dose of vitamin $K_2$, while 95 % recovery was remarkably prolonged by the increased dose of vitamin $K_2$. Specifically, without the administration of vitamin $K_2$ to a patient with normal renal function, the 95 % $INR$
recovery was 8 h, while it increased to 100 h when 20 mg of vitamin K$_2$ was administered. The calculated INR/day also decreased from 0.39 (0 mg of vitamin K$_2$) to 0.072 (20 mg of vitamin K$_2$) in patients with normal renal function. In patients with decreased renal function, a similar but greater INR change compared with those with normal renal function is shown in Fig. 5 and Table 3A.

**Effect of warfarin dose on INR transition**

Figure 6a shows the simulation curves with various warfarin maintenance doses. The INR values increased almost directly according to the increase in warfarin dose. In Table 3B, each quantitative index in Fig. 6a is shown, and the cases where 20 to 60 mg of vitamin K$_2$ was administered are indicated. The ΔINR increased, ranging from 0.130 to 0.754 depending on both the warfarin maintenance dose and the vitamin K$_2$ total dose. The 95% recovery depended both on the warfarin maintenance dose and on the vitamin K$_2$ total dose.

**Effect of interindividual variability on INR transition**

Figure 6b shows the effects of interindividual variability on INR transition. The simulated curves suggested that the interindividual variability of $k_{30}$ had a relatively small effect on INR variability, while $k_s$ and $IC_{50}$ had greater effects although they varied by 26.5 % or 37.9 %, respectively, from each population mean value. The INR values under a warfarin maintenance dose of 3 mg ranged from 1.47 to 1.98, and INR values after warfarin withdrawal ranged from 1.23 to 1.55, depending on $k_s$, $IC_{50}$, and $k_{30}$ values. Table 3C shows quantitative indices of the results of Fig. 6b. The ΔINR values ranged from 0.237 to 0.416, from 0.220 to 0.504, and from 0.294 to 0.310, when $k_s$, $IC_{50}$, and $k_{30}$ were increased or decreased by the interindividual variability from the population mean value, respectively. The interindividual variability of $k_s$, $IC_{50}$, and $k_{30}$ had similar effects on the 1st loading. Unlike the ΔINR values, the interindividual variability of $k_s$ and $IC_{50}$ had a small effect on the 95% recovery, while the $k_{30}$ value strongly affected the 95% recovery.

**Discussion**

It is widely known that the warfarin dose suitable for a patient varies among individuals and that careful monitoring of its anticoagulant activity is necessary for preventing excessive anticoagulation or hemorrhagic events [6, 7]. Vitamin K$_2$ can effectively antagonize warfarin, for example, in the preoperative period and when life-

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**Table 2** Final population pharmacokinetic and pharmacodynamic parameters

| Mean parameters | Estimate | RSE |
|-----------------|----------|-----|
| $k_s$ (%/h)     | 3.97     | 17.5|
| $k_d$ (1/h)     | 0.0611   | 9.90|
| $IC_{50}$ (μg/mL) | 0.604   | 24.5|
| $E_{max}$      | 0.324    | 15.9|
| $EC_{50}$ (μg/mL) | 5.30    | 17.6|
| $k_{30}$ (1/h) | 0.0194   | 19.2|
| θ               | 0.614    | 13.9|

| Interindividual variability | Estimate (CV%) | RSE |
|-----------------------------|----------------|-----|
| $ω_{ks}^2$                  | 0.0704 (26.5)  | 25.6|
| $ω_{IC_{50}}^2$             | 0.144 (37.9)   | 43.3|
| $ω_{k_{30}}^2$              | 0.171 (41.4)   | 85.3|

| Residual variability (%) | Estimate (CV%) | RSE |
|--------------------------|----------------|-----|
| σ$^2$                     | 0.0798 (28.2)  | 11.8|

$k_s$, synthesis rate constant; $k_d$, degradation rate constant; $IC_{50}$, 50 % inhibitory concentration of warfarin; $E_{max}$, maximum effect of vitamin K$_2$; $EC_{50}$, 50 % effective concentration of vitamin K$_2$; $k_{30}$, elimination rate constant of vitamin K$_2$; θ, a factor for the effect of decreased renal function on $IC_{50}$; RSE, relative standard error

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**Fig. 3** Population predicted thrombotest ($TT$, A) or individual predicted $TT$ (B) versus observed $TT$ in each patient. The unit line is shown.
threatening bleeding occurs [9]. Although the recommended dose of vitamin K$_2$ was under 5 mg [9], 20–70 mg of vitamin K$_2$ was administered to decrease the INR value in the preoperative period (Table 1). Thus, caution must be exercised to find a balance between over- and under-coagulation. The pharmacokinetics and pharmacodynamics of warfarin have been studied since 1960’s [11, 12, 14, 21, 22], while combined pharmacokinetic/pharmacodynamic analyses of both warfarin and vitamin K formulations have not yet been reported. In the present study, we built a model that describes the pharmacokinetics/pharmacodynamics of these drugs for the first time. However, because this is a retrospective study wherein only patients’ pharmacodynamic data were used and because we converted the INR values to TT values while calculating the pharmacokinetic/pharmacodynamic parameters, special attention should be paid when drawing conclusions from the results obtained herein. Additionally, the obtained pharmacokinetic and pharmacodynamic parameters should be carefully treated, since these values greatly depended on the fixed pharmacokinetic parameters of warfarin and vitamin K$_2$ in the model.

Final population pharmacokinetic/pharmacodynamic parameters had reasonably small relative standard errors except $\omega^2_{k30}$ (Table 3), and both individual predicted TT and INR values were well correlated with the observed values (Figs. 3 and 4), indicating that reliable population mean parameters were obtained in this study. Some patients had the INR values between 1.0-1.5 on the day of admission (Fig. 2). We could not check drug
compliance in the patients before the hospitalization, but good compliance was expected in the hospital. Since the prediction bias of the TT was not observed against time (data not shown), effects of non-compliance on the present results were considered to be small. Since coadministration of amiodarone or bucolome was reported to inhibit the warfarin metabolism mediated by CYP2C9 [13, 14], we examined the effect of these drugs on the $k_{10}$. Although the coadministration of these drugs decreased $k_{10}$, this effect did not reach a statistical significance level ($-2\text{LLD} = 7.61 < 7.88$). Therefore, we did not include the effect of amiodarone and bucolome in the final model. The estimated population mean parameters for $k_s$, $IC_{50}$, and $k_{30}$ were similar to those in a previous report [11], and interindividual variability for $k_s$, $IC_{50}$, and $k_{30}$ was minimal, although the intra-individual variability was quite significant.

The several simulations of INR transition by the obtained population pharmacokinetic/pharmacodynamic parameters showed that vitamin K$_2$ could antagonize the anticoagulant activity of warfarin in a dose-dependent manner. While more than 20 mg of vitamin K$_2$ showed only a small effect on the extent of INR decreases in the preoperative period, the time required for warfarin to exert its anticoagulation activity again in the postoperative period depended on the total dose of vitamin K$_2$. An inability to anticoagulate promptly after the operation may lead to prolonged hospitalization and consequently decrease patients’ quality of life, as well as increase medical costs. Although it is important to examine the effect of less than 20 mg vitamin

### Table 3

Quantitative evaluation of the simulated INR transitions corresponding to Figs. 5 and 6

(A) Effect of renal function on INR transitions.

| Renal Function | Normal | Decreased renal function |
|----------------|--------|--------------------------|
| Vitamin K$_2$ (mg) | 0 20 | 40 60 0 20 40 60 |
| $\triangle$INR ($\times 10^{-1}$) | 1.30 3.02 | 3.41 3.58 2.15 5.11 5.73 5.98 |
| 1st Loading ($\times 10^{-1}$) | 0.72 0.40 | 0.23 0.15 1.12 0.55 0.30 0.17 |
| 95 % Recovery (h) | 8 100 | 148 172 16 126 174 200 |
| INR/day ($\times 10^{-1}$) | 3.90 0.72 | 0.55 0.50 3.23 0.97 0.79 0.72 |

(B) Effects of combinations of various warfarin maintenance doses and vitamin K$_2$ doses on INR transitions.

| Warfarin (mg) | 3 | 4 |
| Vitamin K$_2$ (mg) | 0 20 | 40 60 0 20 40 60 |
| $\triangle$INR ($\times 10^{-1}$) | 1.30 3.02 | 3.41 3.58 1.75 4.10 4.62 4.83 |
| 1st Loading ($\times 10^{-1}$) | 0.72 0.40 | 0.23 0.15 0.77 0.38 0.20 0.11 |
| 95 % Recovery (h) | 8 100 | 148 172 26 | 126 172 196 |
| INR/day ($\times 10^{-1}$) | 3.90 0.72 | 0.55 0.50 1.62 0.78 0.64 0.59 |
| Warfarin (mg) | 5 | 6 |
| Vitamin K$_2$ (mg) | 0 20 | 40 60 0 20 40 60 |
| $\triangle$INR ($\times 10^{-1}$) | 1.30 3.02 | 3.41 3.58 3.58 4.10 4.62 4.83 |
| 1st Loading ($\times 10^{-1}$) | 0.72 0.40 | 0.23 0.15 0.77 0.38 0.20 0.11 |
| 95 % Recovery (h) | 28 150 | 196 206 32 | 158 200 224 |
| INR/day ($\times 10^{-1}$) | 1.89 0.84 | 0.72 0.72 2.00 0.99 0.87 0.81 |

(C) Effect of interindividual variability on INR transitions.

$\Delta$INR, a decrease before and after warfarin withdrawal; 1st Loading, an INR increase by the first warfarin loading; 95 % Recovery, a time to elevate to the 95 % of the INR value before warfarin withdrawal; INR/day, $\triangle$INR divided by 95 % Recovery
K₂ on INR, we could not obtain clinical data using less than 20 mg vitamin K₂. Effects of lower dose of vitamin K₂ on INR remains to be examined in a future study.

In this study, we clarified the enhanced anticoagulant activity of warfarin in patients with decreased renal function. Warfarin is well known to inhibit the vitamin K-dependent synthesis pathway of coagulation factors in the liver and to be degraded in the liver [10]. Thus, great caution is required while using warfarin in patients with hepatic disorders [10]. According to the package insert of warfarin, caution is also required while use in those with renal dysfunction. Recent studies reported that renal function influences warfarin responsiveness and hemorrhagic complications [23, 24]. The maintenance warfarin dose was positively correlated with kidney function in Japanese patients [25]. Precise mechanisms for the enhanced sensitivity to warfarin in patients with decreased renal function should be investigated further in future studies.

Conclusions
We built and analyzed a pharmacokinetic/pharmacodynamic model of both warfarin and vitamin K₂ by using retrospective clinical data during the catheter ablation. Simulations using the obtained population pharmacokinetic/pharmacodynamic parameters indicated that vitamin K₂ should be administered with care and that more than 20 mg is unnecessary in the preoperative period of catheter ablation. Low-dose (5 mg or less) of vitamin K is recommended in the guideline [9].

Abbreviations
AIC, Akaike information criterion; Cp, plasma concentration; CYP, cytochrome P450; EC₅₀, 50% effective concentration; Eₘₐₓ, maximum effect; IC₅₀, 50% inhibitory concentration; INR, international normalized ratio; kₛ, synthesis rate constant; kₐ₀, degradation rate constant; LLD, log likelihood difference; OBJ, objective function; TT, thrombostest; Vd, distribution volume; kₐ₀, elimination rate constant; VKORC1, vitamin K epoxide reductase complex subunit 1.

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Authors’ contributions
ZZ, IY, SO, and SS conceived the study, designed the protocol. ZZ, IV, SO, and YM carried out the study and drafted the manuscript. SS, MH, TK, AA, KI, and KM participated in interpretation of the data and contributed the discussions. All authors read and approved the final manuscript.

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Competing interests
The authors have no competing interests to declare for this study.
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Not applicable.

Ethics approval and consent to participate
This study was conducted in accordance with the Declaration of Helsinki and its amendments. The study protocol was approved by the Ethics Committee of the Kyoto University Graduate School of Medicine and Kyoto University Hospital (R0264).

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