Research Article

Prospective evaluation of a computerized algorithm for Vitamin K antagonist drug dose calculation

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Summary

Introduction: In an earlier study, we described and validated a VKA dose-finding algorithm (B2A), based on a novel bidirectional factor (BF). We designed a prospective study to evaluate the B2A in a daily care setting.

Methods: In this open-label prospective study, we compared the outcomes of the B2A over the year 2020 with the outcomes of the previous year (2019), using regular algorithms. The outcomes were the duration of Time in the Therapeutic Range (TTR), the percentage of automated dose proposals (PAuP) and the percentage of accepted dose proposals (PAcP). The data were obtained from three anticoagulation centers in the Netherlands, in four locations. The outcomes of this study were based on a non-inferiority level.

Results: The TTR over the year 2020 was at least non-inferior compared with the standard of care treatment. The percentage of automated proposals increased in all centers to approximately 96% of all dosages.

Conclusion: The B2A performs non-inferior compared with the existing algorithms and in some aspects even better.

Introduction

Direct Oral Anticoagulants (DOACs) are nowadays the drugs of choice for the prevention and treatment of most thrombo-embolic diseases [1-4]. Vitamin K Antagonists (VKAs) are prescribed for anticoagulant treatment in selected patients with atrial fibrillation, mechanical heart valves and antiphospholipid syndrome [5,6]. VKAs have a narrow therapeutic window and require monitoring of their anticoagulant activity [7-10]. In addition, frequent dose control and/or adjustments are necessary to optimize safety and efficacy [11-13]. In many countries, VKA dose finding is performed in specialized anticoagulation centers to improve the quality of VKA therapy [14,15]. Despite the expertise available at qualified centers, it is often not possible to increase the quality of VKA therapy. The results of poor VKA therapy are both undertreatments, which may lead to recurrent thrombo-embolic events and overtreatment, with the risk of (major) bleeding. Both risks appeared in the comparisons of VKA treatment with the DOACs. This is the main reason for the preference of DOACs over VKA treatment in several guidelines.

Computerized Decision Support systems (CDS) were designed to improve VKA monitoring and dose-finding [16-20].

In an earlier study, we described and validated a dose-finding algorithm (B2A), based on a novel bidirectional factor (BF) [21]. This BF is a linear transformation of the nonlinear INR. A retrospective study with this B2A, based upon the bidirectional factor (BF), showed similar outcomes compared to VKA dose finding guided by experienced staff in the participating anticoagulation centers.
Improvement of the existing management of VKA treatment could lead to a better quality of control and to decrease in clinical outcomes.

We designed a prospective study to evaluate the B2A in a daily care setting.

**Methods**

**Design**

The algorithm was built within the two major CDS platforms in the Netherlands (Trodis [distributor ASolutions] and Portavita[ CGM]).

In this open-label prospective study, we compared the outcomes of the B2A over the year 2020 with the outcomes of the previous year (2019). In 2019 the medical staff performed dose finding utilizing the existing CDS algorithms in routine situations and manually in difficult cases. The specialized medical staff of the anticoagulation department evaluated all dosage proposals before they were sent to the patients. The quality of VKA therapy is usually expressed as the Time in Therapeutic Range (TTR) [22].

We hypothesized that the algorithm performance was sufficient if the outcomes were at least comparable in both years concerning the TTR and if the percentage of accepted dose proposals would increase.

The Institutional Review Board of the Isala Clinics (Zwolle, the Netherlands) approved the study.

**Outcomes**

The outcomes were the duration of Time in the Therapeutic Range (TTR), the percentage of automated dose proposals (PAuP) and the percentage of accepted proposals (PAcP). The TTR is predictive for thromboembolic and bleeding complications in patients on VKA. The combination of PAuP and PAcP is a key marker of the quality of the algorithm. If both were comparable in the two years, the algorithm would be capable of making a high number of accepted dosage proposals with a comparable TTR as result.

**Data collection**

The data were obtained from three anticoagulation centers in the Netherlands, in four locations (note: one center had two locations). Two of the thrombosis centers used the CDS PortaVita® and one uses both Trodis® (ASolutions) and PortaVita® in two locations.

From the three anticoagulation centers datasets were obtained. Each dataset contained the TTR, the percentage of automated dosage proposals, and the percentage of accepted proposals. The dataset does not generate information on an individual patient level because of privacy restrictions by law.

In The Netherlands, the Dutch Medicines Evaluation Board has approved acenocoumarol (Tablet 1 mg) and phenprocoumon (Tablet 3 mg): Warfarin is not an approved drug in the Netherlands. Therefore, we analyzed the B2A for both acenocoumarol and phenprocoumon.

**Statistical analysis**

We determined the combined TTR values of all patients in the four participating centers over the years 2019 and 2020. Furthermore, we compared the percentage of automated proposals (PAuP) and the percentage of accepted proposals (PAcP) over these years.

The outcomes of this study are based on a non-inferiority level [23,24]. The study has an open-label design and the study population could alter over consecutive years. If the outcomes of the B2A were lower compared to the standard method of treatment, the delta value for the statistical testing would be 4%.

We used MedCalc® software for statistical analysis. If the results of the B2A were lower than the results of the existing software we would do several statistical tests in order to analyze this finding. If the outcomes were equal or better, then the outcome would be that the B2A was non-inferior.

**Results**

Datasets were obtained from the three anticoagulation centers (four locations) over the years 2019 and 2020. In 2019 there were 268,214 INR values and in 2020 257,641. This decrease was mainly due to the ongoing transfer of patients from VKA to DOAC. At the end of 2020, there was a small increase in INR measurements on account of the COVID-19 vaccinations.

**TTR**

Table 1 shows the TTR results over the years 2019 and 2020.

The TTR over the year 2020 was at least non-inferior compared with the standard of care treatment Figure 1.

The percentage of automated proposals (PAuP) and the percentage of accepted proposals (PAcP).

The percentage of automated proposals increased in all centers to approximately 96% of all dosages (Table 2, Figure 2).

The percentage is never 100% because the B2A only proposes when a former dosage is available. This is not the case with new patients.

<table>
<thead>
<tr>
<th>Center</th>
<th>2019</th>
<th>2020</th>
</tr>
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<tbody>
<tr>
<td>1</td>
<td>82.1</td>
<td>83.3</td>
</tr>
<tr>
<td>2</td>
<td>79.2</td>
<td>81.8</td>
</tr>
<tr>
<td>3</td>
<td>76.1</td>
<td>79.3</td>
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The percentage of accepted proposals increased in all centers in subsequent years (Table 3, Figure 3).

There are several possible reasons for the alternation of a dosage proposal, for instance, a planned medical intervention or a change of medication intake in the previous period. The percentage of accepted dose advice differs between the three centers, but all centers showed an increase in the percentage of acceptance over the two years.

**Discussion**

VKA treatment is difficult and hazardous because it needs several skills and experience to achieve an acceptable quality of care. The hazards are both a lack of effectiveness in the sense of recurrence of thrombo-embolic events and problems with safety on account of a high incidence of major bleeding. Several studies have shown that DOACs perform better regarding both issues. In these studies, the quality of VKA control was low, when regarding the TTR.

A CDS can be used in order to assist in VKA dose-finding. A CDS needs a dose-finding algorithm to calculate the VKA dosage of a patient for the following period of time after the last INR measurement. Most CDSs have an algorithm, but the outcomes are often dissatisfying. Sometimes constructing a new algorithm or algorithm provides no proposal and often the proposal needs to be adjusted.

We have made a new algorithm based on a mathematical concept of transforming the exponential INR value into a linear value. With this value, a new dosage proposal can be calculated easily. We have proven the correctness of this concept in a previous retrospective study.

We designed a new prospective study to compare our dose-finding algorithm with regular software. The open, non-randomized design of the study was chosen because of an expected change in population characteristics over the years 2019 and 2020 as a result of an ongoing transition of patients using VKAs to DOAC [25,26]. This fact and the COVID-19 pandemic in 2020 made the population characteristics not entirely compatible [27].

The TTR over the year 2020 was not inferior compared with the TTR over the year 2019. The percentage of automated dosage proposals increased from 87% in 2019 to 96% in 2020, which means that the B2A proposes in almost all cases. Of the computer-generated proposals, 76% - 88% were considered to be correct by the thrombosis center staff.

Every working day the CDSs produce three different lists of dosage proposals. The first list contains proposals that are probably correct, the second list contains proposals that need some attention and the third list appears the difficult cases. The percentage of acceptance decreases from almost 100% for list 1% to 60% for list 3. This implies that list 1 is almost an expert system because the algorithm provides a
correct dosage proposal for almost every case in this list. List 3 contains patients that are ill, patients with new interacting medication, or patients with a planned medical intervention in the nearby future. You may explain the lower percentage of acceptance of this list.

This prospective study has shown that the B2A performed well in two existing CDSs.

The performance of our B2A proved to be comparable and in some aspects even better than the CDSs used in the anticoagulant centers. We confirmed that the results from our retrospective study are also achievable in a daily care setting.

The B2A can support the VKA dose finding of the medical staff. It is not an expert system, so medical evaluation of the outcomes is always necessary. In the future, B2A may further evolve into an expert system, that can perform dosage advice in non-complicated situations.

Building the B2A into existing CDSs can improve the TTR of a patient and hopefully improve the issues of effectiveness and safety of VKA therapy.

Conclusion

The B2A performs non-inferior compared to the existing algorithms and in some aspects even better.

Acknowledgment

The study was supported by a Grant from the Dutch Federation of Thrombosis Centers.

Conflict of interest: Maarten Beinema is a member of the Scientific Board of Portavita and a member of the Medical Committee of Portavita. Henk Adriaansen is a member of the Medical Committee of Portavita.

Addendum

The role and contribution of the authors: MB: Study design and founder of the B2A; JB: Study design and input manuscript; HA: Input manuscript, a leading physician of one of the centers; FJ: Study design, input manuscript.

References


