Update on Direct Oral AntiCoagulants (DOACs)

Perioperative “switching”, drug interactions and persistence

J. Koscielny; C. Rosenthal; C. von Heymann

1 Gerinnungsambulanz mit Hämostaseologie in ambulanten Gesundheitszentrum (AGZ), Charité Campus Mitte (CCM), Universitätsmedizin Berlin; 2 Klinik für Anästhesiologie mit Schwerpunkt operative Intensivmedizin, Charité Campus Virchow Klinikum (CVK), Universitätsmedizin Berlin; 3 Klinik für Anästhesie, Intensivmedizin, Notfallmedizin und Schmerztherapie, Vivantes Klinikum im Friedrichshain, Berlin

Keywords
DOACs, rivaroxaban, apixaban, edoxaban, dabigatran, “switching”, drug interactions, persistence

Summary
Recent findings require an update of previous recommendations for the perioperative use of Direct Oral AntiCoagulants (DOACs). A break in preoperative treatment of 24–96 hours is recommended based on the pharmacokinetic profiles of DOACs and depends on individual patient characteristics, their renal and possibly liver function, and their surgery-related risk of bleeding. In cases of renal or hepatic insufficiency, whether to extend the perioperative interruption of Ila- and Xa-inhibitors is a clinical decision that must be reached on an individual patient basis. In cases of epidural or spinal anaesthesia, more conservative pauses-intervals are recommended due to the risk of persistent neurologic deficits (e.g., paraplegia) following the development of spinal subdural and epidural haematomas. Elective surgery should be postponed according to these recommendations. Preoperative “bridging” with LMWH (more precisely referred to as „switching“) should be omitted due to a significantly increased risk of bleeding. In addition, the incidence of perioperative thromboembolic risks, such as DVT, PE, and stroke, are no different whether interruption or “switching” is undertaken. Postoperatively, the DOACs can be re instituted within the first 24 hours. In cases of major surgery or if there is a higher risk of bleeding, resumption of DOACS should only begin after 24–72 hours. In patients with an elevated thromboembolic risk, transient postoperative LMWH administration can be recommended during this period. Interaction of DOACs with other drugs usually occurs during the absorption, transport and elimination of these drugs. Therefore, substance-specific restrictions and recommendations should be observed during these times. In everyday clinical practice, web-based, independent information portals on drug-interactions are very helpful in providing safe and rapid information about potential interactions when DOACs are used in combination with other drugs, especially during perioperative management. Non-adherence to medications is a worldwide problem that has dangerous and costly consequences. Present data suggest that persistence is the primary factor that supports adherence.

Despite the adherence data presented in the DOACS approval studies (e.g., persistence in the treatment of acute venous thromboembolism has been reported to be between 94–99%), the first registries and meta-analyses provide sobering results regarding the incidence of persistence and the success rate of interventions designed to improve adherence with DOACs in cases of long-term usage.

Schlüsselwörter
DOAKs, Rivaroxaban, Apixaban, Edoxaban, Dabigatran, „Switching”, Medikamenteninteraktionen, Persistenz

Zusammenfassung


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Die Non-Adhärenz von Medikamenten ist weltweit verbreitet, ist gefährlich und teuer in ihren Folgen. Die aktuellen Daten beschrei-ben vorwiegend die Persistenz als ein orientierendes Maß für die Adhärenz. Unabhängig von den Zulassungsstudien der DOAKs (Persistenz bei der Therapie akuter venöser Thromboembolien zwischen 94 – 99%) liefern erste Register und Metaanalysen ermutigende Ergebnisse zur Persistenz und zur Verbesserung der Adhärenz der DOAKs in der Langzeitanwendung.

Introduction

The major medical indications for antico-agulant therapy are prophylaxis for stroke and thromboembolism associated with at-rial fibrillation and for the treatment and prophylaxis of venous thromboembolism and pulmonary embolism. Currently, the established vitamin K antagonists are in-creasingly being replaced by or supple-mented with Direct Oral AntiCoagulants (DOACs) – namely, the thrombin-inhibitor dabigatran and the Xa-inhibitors rivaroxa-ban, apixaban and edoxaban.

Dabigatran received EU drug approval in September 2011, rivaroxaban in De-cember 2011, apixaban in January 2013, and edoxaban in August 2015 (1–10).

The number of patients taking DOACs has increased in conjunction with the number of surgical procedures in those same patients. For instance, a prespecified analysis of the RE-LY trial showed that approximately 25% of the study participants taking dabigatran underwent an invasive elective or emergency procedure within 24 months after the start of treatment (1).

Thus, an important question that often arises in the perioperative period is when and how long before a procedure should DOACs be discontinued? Appropriate guidelines are particularly needed because antidotes are not available for all DOACs. To address this gap in knowledge, different recommendations and/or guidelines for perioperative management of DOACs have been put forth (11–22). These recommen-dations are not primarily based on data from prospective (randomized) clinical trials, but rather they have been formulated based on pharmacokinetic considerations and registry data derived from clinical practice (23–28). Recent results from these registries show that preoperative „switching“ (e.g., to LMWH) procedures are as-soicated with higher bleeding rates.

Furthermore, approximately seven per-cent of the undesired effects associated with DOACs are due to drug interactions, with the incidence increasing along with the number of concomitant drugs (29, 30).

Nonadherence to medications is a worldwide problem that has dangerous and expensive consequences. According to a WHO report only 50% of patients on aver-age with chronic diseases are adherent to their medications in developed countries (31). The cost of nonadherence to the US health care system is estimated to be ap-proximately 300 billion dollars per year. This represents approximately 13% of the total cost of the US health care system (32).

The main cost drivers are avoidable hos-pitalizations that cost approximately 100 bil-lion dollars. In Germany, these costs are es-timated at 7.5 – 10 billion euros per year (33).

Despite the adherence data in the DOAC approval studies (e.g., adherence to the treatment of acute venous thromboem-bolism was reported to be between 94–99%), the first registries and meta-anal-yses provide sobering results about the level of adherence. Consequently, interven-tions aimed at improving adherence to DOACs in cases of long-term usage are ur-gently needed.

The purpose of this review is to provide a comprehensive review on the perioper-a-tive management of DOACs, the impact of drug interactions on DOAC plasma level-s and the challenge of patient adherence, factors that merit close attention so that medical care for patients using DOACs can be improved in the long run.

Perioperative management of DOACs

The European Heart Rhythm Association (EHRA) has published recommendations for the perinterventional and perioperative management of DOACs (11). These recommendations also take into account the presence of renal insufficiency and are summarized, along with other recommenda-tions, via an „APP“ BridgeAnAcoag (App-Store and Googleplay-Store) from the ACC (American College of Cardiology), as follows: in general, the partial elimination by the kidneys requires timely discontinuation of DOACs in the perioper-a-tive or perinterventional phase, especially in patients with renal insufficiency. In pa-tients with normal renal function, a preop-erative discontinuation of DOACs for 24 hours (2–2.5 half-lives) should be sufficient to reduce plasma-levels to at least 25%, which should be safe for surgical procedur-es with a low risk of bleeding (17, 43).

Only the thrombin inhibitor dabigatran, with a mean half-life of 13.4 hours with a normal glomerular filtration rate (GFR) of > 80 ml / min, should be discontinued 36–48 hours before surgery when there is doubt about the timing of the last dose or concerns about an increased risk of bleeding. In a study by Healey et al. (1) that investigated the perinterventional discontinu-a-tion of dabigatran as a subgroup analysis of the RE-LY study, this time inter-val was associated with a significantly lower rates of bleeding complications com-pared with the interruption of warfarin.
The recommended 24-hour interruption interval comprises roughly 2 half-lives of DOACs, which is comparable to the standard practice of preoperative management of LMWH (17).

The presence of renal insufficiency requires a longer period of discontinuation for all DOACs, as at least 25% of all substances are eliminated renally. Thus, depending on the severity of renal insufficiency, treatment discontinuation of ≥48 hours (approximately 4–5 half-lives) is recommended for the group of factor Xa-inhibitors (Table 1).

For dabigatran, which is predominantly (80%) eliminated through the kidneys, discontinuation for 72–96 hours is recommended depending on the creatinine clearance (Table 2) (11, 17, 44). Because of the diagnostic imprecision of serum creatinine levels in cases of renal insufficiency, renal function in patients with suspected renal impairment should preferably be monitored with a measured or calculated creatinine clearance.

In addition, and as an exception, it may be useful to measure DOAC plasma levels preoperatively using the “diluted thrombin time” (DTT) or a chromogenic assay for dabigatran or a calibrated anti-Xa activity for the FXa-inhibitors (apixaban, edoxaban, rivaroxaban). Even though the plasma levels do not correlate strictly with the risk of bleeding, they may be useful for assessing the risk when a surgical procedure with an increased bleeding risk is undertaken. Despite a lack of evidence from prospective trials, some centres have determined that a cut-off plasma level for DOACs of <30 ng/ml is safe with respect to perioperative bleeding risk.

Placement and removal of epidural catheters as well as administering spinal anaesthesia also require preinterventional pausing of DOACs. The recommended pausing intervals for neuraxial analgesia/anaesthesia have been developed under the auspices of the German Society of Anaesthesiology and Intensive Care Medicine (17) (Table 3). These pausing intervals correlate largely with the abovementioned recommendations (17, 44), though they reside in the more conservative upper limit of pausing-intervals due to the high risk and serious consequences of periprocedural bleeding. In this scenario, the measurement of DOAC plasma levels may be helpful if doubt exists about the last time oral anticoagulants were ingested. In our view, in cases of uncertainty about this timing, one should refrain from neuraxial anaesthesia/anaesthesia to minimize the risk of severe neurological complications resulting from the development of a spinal subdural or epidural haematoma.

No preoperative “switching” of DOACs

The widely used (12–14) practice of “bridging” with LMWH for patients taking vitamin K antagonists (VKA) is strongly discouraged during the preoperative DOAC break (11, 18–22, 45, 46), since this may be associated with a higher risk of perioperative bleeding.

In practice, the different meanings of “bridging” and “switching” should be considered and used correctly.

<table>
<thead>
<tr>
<th>Creatinine clearance (ml/min)</th>
<th>Bleeding risk</th>
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<tr>
<td></td>
<td>low</td>
</tr>
<tr>
<td>≥80</td>
<td>≥24 h</td>
</tr>
<tr>
<td>50–80</td>
<td>≥24 h</td>
</tr>
<tr>
<td>30–50</td>
<td>≥24 h–48 h</td>
</tr>
<tr>
<td>15–30</td>
<td>≥48–72 h</td>
</tr>
<tr>
<td>&lt;15</td>
<td>no approval</td>
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</tbody>
</table>

Because the preoperative pausing intervals for DOACs are usually quite short due to their short half-lives and because there is no evidence for more effective avoidance of thromboembolic complications associated with preoperative switching, DOACs should be stopped preoperatively according to recommended time-intervals (Table 1 and Table 2) before surgery/interventions. Based on recent data analysis, the “switching” method from the registries of individual DOACs showed no reduction in the rate of thromboembolic complications, although there was a significantly increased risk of periinterventional bleeding (Table 4).

In the subgroup analysis by Douketis et al. (47), the rate of perioperative/perinterventional major bleeding events in patients receiving dabigatran was 6.5% with “switching” versus 1.8% without “switching” (p <0.001). In contrast, the rates of thromboembolic events and serious embolic complications did not differ between the “switching” and “no switching” groups: 1.2% vs. 0.6%, (p =0.16) and 0.5% vs. 0.3% (p = 0.46), respectively. These results provide evidence that peri-interventional therapy with dabigatran for atrial fibrillation does not require “switching” with heparin prior to minor procedures with a low risk of bleeding. "Real-life" data from a registry in Saxony that investigated the peri-interventional/periprocedural bleeding rate in DOAC-treated patients also supports that preinterventional “switching” is not required (23, 24, 48). The majority of the patients examined were taking rivaroxaban. Of a total of 863 interventions and minor surgical procedures, there were 3 major bleeding events in the group of patients without “switching” to heparin ver-
sus 7 major bleeding events with “switching” (0.5% vs. 2.7%, p = 0.010). In a prospective cohort study of dabigatran with a high medication adherence of 77%, the perioperative bleeding rate without switching was 1.8% (95% confidence interval: 0.7–3.0, 10 of 541 patients) (27). The only thromboembolic event was a transient ischaemic attack (TIA) (0.2%; 95% – confidence interval: 0–0.5). The last dose of dabigatran was given 24 h, 48 h or 96 h (46%, 37% and 6%), preoperatively. A first postoperative dose of dabigatran of 75 mg was administered to 40% of patients at the day of procedure. Another prospective cohort study in patients taking dabigatran, who underwent elective surgery with a standardized break protocol and who were monitored using laboratory analysis, showed the same low perioperative bleeding and low TIA rates of 0.6% (28). Data from the ORBIT-AF registry (25) re-confirmed the significantly increased incidence of bleeding and other complications after “switching” in a group of patients taking dabigatran: 7% of the overall group received oral anticoagulation. Major bleeding was more common in patients with “switching” compared with patients without “switching” (5.0% vs. 1.3%, adjusted OR 3.84, p <0.0001). The incidence of myocardial infarction, ischaemic cerebral infarction or systemic embolism, major bleeding, hospitalization or death within 30 days was significantly more frequent in the group with “switching” (13% vs. 6.3%, adjusted OR 1.94, p = 0.0001). In contrast to “switching”, a subgroup analysis of the pivotal trial of apixaban (ARISTOTLE), investigated the effect of an interruption (> 24 h up to 7 days) of anticoagulation versus no interruption (≤ 24 hours) prior to diagnostic interventions and minor operations. There was no difference in the rate of major bleeding between the groups (1.58% vs. 1.65% [OR 1.023, 95% CI: 0.639 to 1.636]) (43). However, it is noteworthy that these diagnostic interventions and minor surgical procedures were mainly endoscopies of the upper and lower gastrointestinal tract as well as tooth extractions and cataract surgery. Therefore, it is not possible to generalize these findings to the management of other more involved surgical procedures. Interestingly, in a risk-adjusted comparison of groups, thromboembolic complications (stroke, systemic embolism, myocardial infarction) occurred significantly more often without interruption of anticoagulation in the warfarin group (43). This investigation supports the conclusion that for minor surgical procedures and diagnostic interventions, continued anticoagulation with apixaban (interruption less than 24 h) is not associated with more bleeding events compared to a longer interruption of anticoagulation. Periprocedural registry data is not yet available for edoxaban.

Postoperative management of DOACs

The postoperative resumption of anticoagulation with DOACs in the cited studies was usually started in the evening of the day of surgery or on the 1st or 2nd postoperative day. This approach can be recommended safely for the majority of operations, especially those with a low bleeding risk. For operations with an increased risk of bleeding, patients with indwelling epidural catheters for postoperative analgesia or patients with renal or hepatic failure, a later resumption of DOACs seems to be advisable (49). Particularly when using epidural catheters that often remain in place for 3–5 days postoperatively, DOAC treatment should be resumed only after removal of the epidural catheter. Because of this long recommended pausing interval for DOACs prior to catheter removal, patients face an increased risk of thromboembolism.

In these cases, postoperative temporary “switching” to LMWH or UF-heparin for anticoagulation is a favourable solution. When comparing the recommendations for the perioperative management of anticoagulation for patients receiving VKA or DOACs, it appears that “bridging” of VKA is indicated in certain circumstances. This should be considered at least for the period prior to surgery or the intervention in patients with an increased thromboembolic risk. “Switching” is probably not required in most patients with DOACs, due to the

<table>
<thead>
<tr>
<th>Drug</th>
<th>Half-time in plasma</th>
<th>Before neuroaxial process</th>
<th>After neuroaxial process</th>
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<tbody>
<tr>
<td>dabigatran (max. 1 x 150–220mg/d)</td>
<td>14–17h</td>
<td>28–34h</td>
<td>6h</td>
</tr>
<tr>
<td>dabigatran (max. 2 x 150mg/d)</td>
<td>14–17h</td>
<td>56–85h</td>
<td>6h</td>
</tr>
<tr>
<td>rivaroxaban (1 x 10mg/d)</td>
<td>11–13h</td>
<td>22–26h</td>
<td>4–5.5h</td>
</tr>
<tr>
<td>rivaroxaban (2 x 15mg/d, 1 x 20mg/d)</td>
<td>11–13h</td>
<td>44–65h</td>
<td>4–5.5h</td>
</tr>
<tr>
<td>apixaban (2 x 2.5mg/d)</td>
<td>10–15h</td>
<td>26–30h</td>
<td>5–7h</td>
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<tr>
<td>apixaban (2 x 5mg/d)</td>
<td>10–15h</td>
<td>40–75h</td>
<td>5–7h</td>
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<tr>
<td>edoxaban (1 x 30mg/d)</td>
<td>10–14h</td>
<td>20–28h</td>
<td>6–7h</td>
</tr>
<tr>
<td>edoxaban (1 x 60mg/d)</td>
<td>10–14h</td>
<td>40–60h</td>
<td>6–7h</td>
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</table>

1 under ASA-medication (100 mg); free intervals of additional anticoagulants 4–5 half-times in plasma
2 CAFE: half-time in plasma depending on renal function
3 CAFE: half-time in plasma depending on liver function
4 individual risk calculation

Tab. 3 The recommended time intervals for the individual substances before or after neuroaxial anesthesia – from the German Society of Anaesthesiology and Intensive Care Medicine (DGAI) guidelines (17)
Most patients are usually taking several drugs simultaneously, resulting in drug-drug interactions. The definition of a drug interaction is one that results in a negative impact on the therapeutic use (synergistic or antagonistic drug effects). These interactions may occur during absorption, transportation, elimination and distribution of the drugs. For DOACs, the first three routes are of great importance (50–56). To varying degrees, DOACs are substrates of CYP3A4 and/or the P-glycoprotein-transporter, which are also inhibitors of transport processes in various cell types. Usually, a co-medication will influence these proteins, placing the DOACs in a „victim“ role and unveiling a new dimension of possible interactions (50–56). Approximately seven percent of unwanted drug effects are due to drug interactions, and the incidence increases exponentially with the number of co-administered medications (29–30). To receive fast and reliable information in clinical practice, the following web-based, independent providers are helpful: www.dosing.de or www.wechselwirkungskontrolle.de

From a pharmacological view, the exposure of the co-administered drug at steady state is crucial. Here, the degree of accumulation of the “victim” drug, i.e., how much higher the concentration at steady state is compared with the first dose, will be determined (50–56). ▶ Figure 1 provides an overview for orientation.

**Dabigatran**

Dabigatran has a low potential for drug interactions (19, 53, 54). It is neither metabolized through CYP-450 enzymes nor does it affect them. However, it is a substrate of the efflux transporter P-glycoprotein, resulting in increased dabigatran plasma concentrations when P-glycoprotein inhibitors are used simultaneously. Therefore, concomitant use of dabigatran with P-glycoprotein inhibitors such as amiodarone, verapamil, ketoconazole, cyclosporine,itraconazole and dronedarone is contraindicated due to an increased risk of bleeding. Dabigatran administration combined with other anticoagulants, antiplatelet drugs, NSAIDs (nonsteroidal anti-inflammatory drugs), SSRIs (selective serotonin reuptake inhibitors), SSNRI (selective serotonin-norepinephrine-reuptake-inhibitor) and other drugs that may affect haemostasis is also contraindicated (54). Exceptions include when there is a change of the anticoagulant therapy or when unfractionated heparin is given in doses that are necessary to prevent clot formation within central venous or arterial lines (19, 53, 54). When dabigatran and the aforementioned drugs have to be administered concomitantly, it is recommended to adjust the dose of dabigatran, for example, from 2 x 150 mg to 2 x 110 mg daily.

Concomitant use of dabigatran with non-retarded P-glycoprotein-inducers, such as carbamazepine, phenytoin, St. John’s wort, as well as drugs with P-glycoprotein inhibitors, such as posaconazole, tacrolimus, protease inhibitors, including ritonavir, which reduce the bioavailability and thus the plasma levels of dabigatran, is not recommended (53).

**Rivaroxaban**

Rivaroxaban is subject to the metabolism of the CYP3A4 system and is a substrate of P-glycoprotein transporters. When administered concomitantly with inhibitors of these enzyme systems, the plasma levels...
rise. Use of rivaroxaban is therefore not recommended in patients receiving concomitant systemic treatment with azole antifungals (such as ketoconazole, itraconazole, voriconazole and posaconazole) or HIV protease inhibitors (e.g., ritonavir), as this may lead to an increased risk of bleeding (20, 51, 52). Concomitant administration of rivaroxaban with strong CYP3A4 inducers (e.g., rifampicin, phenytoin, carbamazepine, phenobarbital or “St. John’s wort” (Hypericum perforatum)) may reduce plasma concentrations of rivaroxaban and therefore should be avoided (51). However, if non-recommended co-medications are necessary, the patient should be closely monitored for signs and symptoms of adverse effects such as thrombosis. Concomitant administration of rivaroxaban and 500 mg naproxen has not been shown to lead to clinically relevant prolongation of bleeding time. Although no clinically significant pharmacokinetic or pharmacodynamic interactions were observed with concomitant administration of rivaroxaban and ASA (500 mg) or clopidogrel (300 mg initial dose followed by 75 mg maintenance dose), the EMA recommends careful use of rivaroxaban and NSAIDs (including ASA) plus antiplatelet agents due to the increased risk of bleeding (20, 51, 52).

**Apixaban**

The use of apixaban is not recommended in patients receiving concomitant systemic treatment with strong CYP3A4 and P-glycoprotein inhibitors, such as azole antifungals (e.g., ketoconazole, itraconazole, voriconazole and posaconazole) and HIV protease inhibitors (e.g., ritonavir) (21, 56). These drugs may increase plasma levels of apixaban by a factor of 2 or more in the presence of additional factors that reduce apixaban elimination (e.g., severe renal impairment with GFR 15–30 ml/min). In contrast, concomitant use of apixaban with strong inducers of CYP3A4 and P-glycoprotein (for example: rifampicin, phenytoin, carbamazepine, phenobarbital or St. John’s wort) may lead to a reduction in apixaban plasma levels of up to approximately 50%. In patients who are concomitantly treated with those drugs, apixaban is not recommended for use in the treatment of venous thromboembolisms (56). Concomitant use with other anticoagulants is contraindicated due to the increased risk of bleeding risk. This risk is significantly increased particularly in patients under so-called „triple“ anticoagulation regimens that include ASA and ADP-receptor blockers after percutaneous coronary intervention (PCI) with stent placement (21, 56).

**Edoxaban**

Edoxaban is absorbed primarily in the upper gastrointestinal tract (22, 50, 55). In contrast to rivaroxaban, edoxaban is not substrate of P-glycoprotein and therefore has not been shown to lead to clinically relevant prolongation of bleeding time. Although no clinically significant pharmacokinetic or pharmacodynamic interactions were observed with concomitant administration of edoxaban and ASA (500 mg) or clopidogrel (300 mg initial dose followed by 75 mg maintenance dose), the EMA recommends careful use of edoxaban and NSAIDs (including ASA) plus antiplatelet agents due to the increased risk of bleeding (20, 51, 52).
sulted in elevated plasma concentrations of edoxaban. With such a combination, the daily dose of edoxaban should be reduced to 30 mg (50). According to clinical data on concomitant use of edoxaban and quinidine, verapamil or amiodarone, no dose reduction is recommended. The use of edoxaban with other P-glycoprotein-inhibitors, including HIV protease inhibitors, has not been studied. In the ENGAGE AF-TIMI 48 study, concomitant use of thienopyridines (e.g., clopidogrel) as monotherapy was allowed and was associated with a higher rate of clinically relevant bleeding (9, 22). However, the risk of bleeding with edoxaban was lower than with warfarin. There is very limited experience with the combined use of edoxaban with a dual platelet aggregation inhibitor or with fibrinolytics. According to the authors’ opinion, this should only be attempted with caution and close clinical monitoring. In clinical trials, the concomitant use of NSAIDs increasingly led to clinically relevant bleeding (9, 22, 50). The long-term use of NSAIDs together with edoxaban is not recommended. The concomitant use of edoxaban and other anticoagulants is contraindicated due to the increased risk of bleeding. However, the concomitant use of low-dose ASA (≤ 100 mg) had no impact on the edoxaban peak levels or total exposure (55), indicating that edoxaban may safely be used together with low-dose ASA (≤ 100 mg / day).

Persistence of DOACs

Non-adherence to medications is a worldwide phenomenon that has dangerous and expensive consequences. According to a WHO report, in developed countries, only 50% of patients on average with chronic diseases are adherent (31). The costs to the US health care system of drug nonadherence are estimated to be approximately 300 billion dollars per year. This represents approximately 13% of the total cost of the US health care system (32). The main cost drivers are avoidable hospitalizations that cost approximately 100 billion dollars. In Germany, these costs are estimated at 7.5 – 10 billion euro per year (33). Despite the results from DOAC registration studies (e.g., DOACs adherence in the treatment of acute venous thromboembolism reported to be between 94–99%), the first registry and meta-analyses provide sobering results regarding the level of adherence. Approaches to improving the adherence of DOACs in cases of long-term usage are urgently needed. Current data suggest that persistence is the primary guiding force that improves adherence.

In a large meta-analysis (18 randomized controlled trials) that included 101,801 patients (34), the rate of discontinuous intake of DOACs did not differ significantly from that of the comparable therapies for venous thrombosis (risk ratio [RR], 0.91; 95% CI, 0.74–1.13; P = 0.40) as well as the comparable therapies (warfarin, ASA) for prophylaxis of ischaemic cerebral infarct in patients with non-valvular atrial fibrillation (RR, 1.01; 95% CI, 0.87–1.17; P = 0.92). The studies found no detectable improvement in persistence with oral anticoagulation with DOACs compared with vitamin K antagonists. For example, only 50% of patients prescribed DOACs for atrial fibrillation took the oral anticoagulants as prescribed (35). Persistence has traditionally been a major challenge for patients taking oral anticoagulants (37, 38). The question is whether the situation is any better with the new oral anticoagulants (DOACs) under real-life conditions. In a registry analysis, the Dresden research team analysed data on persistence from 1,775 patients with atrial fibrillation treated with rivaroxaban for stroke prevention (n = 1,200) and patients with venous thromboembolism (n = 575) (36). According to this study, which was carried out with a high degree of medical oversight, only 13.6 discontinuations per 100 patients per year occurred – a value well below of 23.7% value in the pivotal study (2). This difference was even more distinct with everyday use of vitamin K antagonists (VKA). Approximately 30% of the patients discontinued the treatment in the first year and another 10% in the second year of treatment (37, 38). For 4,863 US veterans with atrial fibrillation, the adherence rate to anticoagulant therapy with dabigatran was only 72.2% (39). However, this result has to be interpreted in light of substantial variability between the 67 medical institutions included in this study. Factors such as a good selection of the patients, the availability of training, and in particular, structured aftercare were associated with improved persistence. In facilities without such factors, adherence was below 50%, which represents a sustained safety risk for the patients.

This view is supported by results from a Danish registry of 2,960 patients in which similar factors were found to influence better adherence to anticoagulant therapy with dabigatran (40): more than 75% of the patients showed a persistence of over 80% in the first year. Patients with a higher morbidity and more frequent physician contact had the highest persistence rate. A recent US health care research study has examined a database of a large US insurance carrier more closely (41) involving nearly 65,000 records. All the patients had atrial fibrillation and started oral anticoagulation for the first time between 2010 and 2014. Nine out of ten patients had a CHA2DS2-VASc score of more than one. The percentage of patients using vitamin K antagonists was 59%, rivaroxaban 19%, dabigatran 16% and apixaban 6%. The median follow-up period was 1.1 years. Within this period, 47.5% of patients treated with DOACs demonstrated good persistence, defined as taking the treatment according to the package insert for at least eight out of ten days. This result was, indeed, significantly better than the result for treatment with vitamin K antagonists (40.2%, p < 0.001), but confirmed a disappointing low adherence for DOACs. Moreover, the investigation yielded results on the medical consequences of suboptimal persistence with oral anticoagulation. Particularly problematic was poor persistence, as expected, in patients at high risk, specifically a CHA2DS2-VASc score of four or above. Patients in this group with low persistence for more than one month during the follow-up period had an incidence of stroke at least twice as high as those who only showed non-adherence for less than seven days. The risk of stroke increased with the degree of nonpersistence. The risk of stroke in patients with lower CHA2DS2-VASc score (2 or 3) was significantly increased only when they did not take their medi-
cation as prescribed for more than six months in total.

Taking together, the data confirm that persistence in anticoagulated patients with atrial fibrillation, deep vein thrombosis / pulmonary embolism is more important for the prognosis than the type of anticoagulation. This suggests that new approaches are urgently needed to improve drug adherence, e.g., via mobile communications through social networks, e-mails, electronic messaging, offers for training, and in particular, structured aftercare, whose effect must be examined in the future in structured care studies.

In this context, the quality of the available care studies remains problematic. In a meta-analysis of 182 studies that examined various approaches to improve persistence, only 17 studies (9.3%) fulfilled quality standards for analysis (42). In only 5 of these studies was an improvement in patients' health (outcome criterion) detectable. The situation with current studies is sobering, and so far, no general recommendations for improving persistence can be provided.

Conflicts of Interest

Adj. Prof. Dr. J. Koscielny declares the following conflicts of interest: speaker honoraria from Aspen, Bayer Health Care Pharmaceuticals, Daichi Sankyo, Boehringer Ingelheim, CSL Behring, Sanofi-Aventis, Pfizer, BMS, Mitsubishi Pharma, Ferring GmbH, Mylan Healthcare GmbH and Novo Nordisk. Adj. Prof. Dr. J. Koscielny is also a medical advisor for CSL. Behring International, Bayer HealthCare Pharmaceuticals (national and international) and Novo Nordisk (national) for the last three years.

Dr. C. Rosenthal received honoraria and/or travel reimbursements over the last three years for lectures and consultancy work related to the topic of this article from Bayer AG, Boehringer Ingelheim, CSL Behring, Daiichi Sankyo, NovoNordisk, Pfizer GmbH and TEM International.

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