Analysis of Recurrent Stroke Volume between VKA (Vitamin K Antagonist) and Three NOACs (Non-Vitamin K Antagonist Oral Anticoagulants) under Oral Anticoagulant Therapy

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Abstract

Objective: We investigated recurrent stroke volume with non-valvular atrial fibrillation (NVAF) patients treated with non-vitamin K antagonist oral anticoagulants (NOACs) about clinical backgrounds and number of recurrent stroke.

Methods: We administered three NOACs, dabigatran, rivaroxaban and apixaban in 84 post cardioembolic stroke with NVAF. In retrospective study, we measured recurrent stroke volume with MRI volumetry and compared them between 7 vitamin K anticoagulant (VKA: warfarin) cases and 10 NOACs cases under anticoagulant therapy.

Results: Of 84 cases, 27 cases were started with VKA and switched to NOACs after 7 recurrent stroke. Other 57 cases were directly started with NOACs and 10 cases with NOACs as first anticoagulants had recurrent stroke. The frequency of recurrent stroke during anticoagulant therapy are not different among VKA group and three NOACs group. Recurrent stroke volume is significantly larger in VKA group (27.4 cm³) than in NOACs group (3.3cm³).

Conclusions: Secondary prevention with NOACs after stroke might be more beneficial by reducing recurrent infarct volume than VKA.

Introduction

Non-vitamin K antagonist oral anticoagulants (NOACs) have gradually changed anticoagulant therapy with non-inferiority to vascular event and decreasing major bleeding in atrial fibrillation and venous thrombosis [1-4]. We have treated post stroke patients with three types of NOACs for 4 years. However, there is no clear consensus for selection of NOACs. A meta-analysis showed that dabigatran 300mg daily (OR 0.66) and apixaban 5mg daily (OR 0.78) were more effective to prevent stroke and systemic embolism than other NOACs. It also showed that edoxaban 30mg daily (OR 0.46) and apixaban 5mg daily (OR 0.69) were more effective to prevent major bleedings than other NOACs [5]. We selected 84 post stroke cases with NOACs for prevention of recurrent ischemic strokes in retrospective study. Although some studies revealed patients taking NOACs appeared to present low risk of intracranial hemorrhage and inhibit increasing of hemorrhage volume, ischemic stroke have not been fully examined [6-9]. So we examined the recurrent stroke volumes and clinical outcome under taking NOACs and compared with recurrent cases under vitamin K anticoagulant (VKA: warfarin).

Subjects and Method

84 post embolic stroke patients with non-valvular atrial fibrillation (NVAF) (mean age 80±9 years) were treated with three types of NOACs for 4 years from October 2011 to September 2015. NVAF was diagnosed by atrial fibrillation within 14 days after admission without rheumatic mitral valve disease, history of prosthetic mitral valve replacement and infectious endocarditis. During the study period, the direct thrombin inhibitor, dabigatran (March 2011), the factor Xa inhibitors, rivaroxaban (April 2012) and apixaban (February 2013) were clinically available in Japan. The eligibility for anticoagulant drugs and selection of NOACs have been freely determined by each stroke neurological physician without any definite intervention. NOACs administration in cardioembolic stroke cases due to NVAF were divided into two arms (Figure 1). Our enrolled 84 patients with NOACs contains 15 cases with dabigatran (DA), 43 cases with rivaroxaban (RI) and 26 cases with apixaban (AP) as using NOACs finally. We divided 84 post stroke patients with three NOACs into two groups which first anticoagulant therapy with VKA or NOACs as prevention therapy of cardioembolic stroke onset.

Basic clinical characteristics included age, gender, body weight (kg) and type of atrial fibrillation (chronic or paroxysmal). To stratify risk factors of stroke event, we calculated CHADS2 score, CHA2DS2-VASC score and HAS-BLED score after the onset of first cardioembolic stroke and before administration of VKA and NOACs. Hypertension was defined as blood pressure levels ≥140/90mmHg or by the use of antihypertensive drugs. Diabetes mellitus was defined as hemoglobin A1c (HbA1c NGSP) level ≥6.5% or by the use of oral antidiabetic drugs or insulin. Dyslipidemia was defined as a serum cholesterol level ≥220mg/dl or by the use of lipid-lowering drugs including statin. CKD was defined as either positive proteinuria or eGFR<60ml/min/1.73m². The data of blood chemistry were noted with the prothrombin time-international normalized ratio (PT-INR), d-dimer, creatinine clearance and brain natriuretic peptide (BNP). The number of cases with recombinit tissue plasminogen activator (rt-PA)on acute stroke therapy was also counted retrospectively.

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The clinical outcomes were estimated with hospitalization period (days) of first stroke event without recurrence and the period of recurrent event. It also includes improvement in National Institute of Health Stroke Scale (NIHSS) between admission and discharge during VKA and NOACs, modified Rankin Scale (mRS) on discharge, tube feeding on discharge. We calculated total stroke volume which was measured with MRI Volumetry soft: MRI cron (http://www.cabiatl.com/mricro/mricron/install.html). Total stroke volume (cm³) is $\Sigma$ all slices of stroke lesions (cm²) × slice thickness (cm) with diffusion weighted image (DWI) axial MR images with the intensity threshold used by our previous reports [10]. Modified Rankin Scale (mRS) as a clinical outcome is estimated correlation with total stroke volume. Cerebral micro bleeds and hemorrhagic infarction were diagnosed with T2*-weighted MRI scans.

Statistical analysis: We compared clinical factors in two groups of patients with VKA or NOACs after the onset of cardioembolic stroke by Mann Whitney U test (nonparametric data). And we also compared recurrent stroke cases with VKA or NOACs, no recurrent stroke cases with VKA or NOACs and recurrent versus no recurrent stroke in each VKA and NOACs by Mann Whitney U test. We used $\chi$² test between VKA group and NOACs group for evaluating several clinical factors including type of atrial fibrillation, risk factors, concurrent antiplatelet drugs, treatment by t-PA, tube feeding, and recurrent stroke during VKA or NOACs (nonparametric data). Analysis of covariance is used to assess the statistical significance of mean differences in recurrent stroke volumes and mRS between NOACs and VKA. P value of <0.05 was considered statistically significant.

Results

Of 84 cardiac embolic stroke cases due to NVAF, 27 cases were started with VKA and switched to NOACs after 7 recurrent stroke or without 20 recurrent event. Other 57 cases were directly started with three NOACs and 10 of 57 NOACs as first anticoagulant had recurrent stroke. Of 57, 10 cases had recurrent stroke. 6 cases switched to other NOACs. 4 cases kept same NOACs.

In table 1, we showed the clinical background of VKA group and three NOACs group. The NOACs group is significantly lower in CHADS² score, CHA²DS²-VASc score, HAS-BLED score, mRS on discharge and NIHSS score on admission and discharge than VKA group. VKA group is significantly higher rate of CKD with lower Ccr and higher BNP. Although recurrent stroke occurred in 7 of 27 (25.9%) patients in VKA group and in 10 of 57 (17.5%) in NOACs group, we could not find any difference between two groups.

In table 2, we can recognize the baseline characteristics of recurrent or no recurrent case with each anticoagulant, VKA or NOACs. In recurrent case the VKA group had lower body weight and higher risk score with stroke event predicted by CHA²DS²-VASc score (similar to comparison in no recurrent) and HAS-BLED score. The VKA group in recurrent case has significantly higher rate of CKD with lower value of Ccr similar to basic characteristics in table 1. NOACs group had a shorter period of hospitalization, lower mRS on discharge and lower rate of tube feeding. On two columns right end of table 2, we can recognize that low body weight, presence of CKD and tube feeding is characteristic of VKA in recurrent patients. And we could find increased D-dimer in recurrent patients with NOACs (=3.69 µg/ml)
compared to recurrent patients with VKA (=1.43 µg/ml). The mean value of PT-INR on 7 recurrent stroke under VKA was 1.50 with under therapeutic range. Of 7 recurrent stroke under VKA, 5 patients (71%) had been in therapeutic range less than 1.8. Of 27 VKA therapy after second prevention, only 6 (22%) had been in therapeutic range more than 1.8 with PT INR.

In Table 3, 10 recurrent stroke cases (first dosage NOACs of 4 DA, 5 RI and 1 AP) under treatment of NOACs and their clinical backgrounds are presented. They have high age, low adherence, under dose of NOACs, malignancy and systemic atheromatous disease. In Table 4, recurrent stroke patients with 10 patients with NOACs and 7 patients with VKA were presented. Especially case number 10 patient in NOACs group showed small hemorrhage of left thalamus and hematoma had gradually disappeared after 20 days from stroke onset. After adjusting for age, concurrent antiplatelet, NIHSS on admission, microbleeds on MRI and complicated hemorrhage, the recurrent stroke volume is larger in VKA group (27.4 cm³) than in NOACs group (3.3 cm³) (p<0.05, F=5.35). After adjusting for age, concurrent antiplatelet, NIHSS on admission and complicated hemorrhage, the mRS on discharge is lower in NOACs group (1.7) than in VKA group (3.7)(p<0.05, F=3.88).

Discussion

Antithrombotic therapy is recommended on the basis of assessment of risk factor of cerebral infarction with NVAF. Suitable anti-coagulate drugs could be selected by both of CHADS₂ scores and CHA2DS2-VASc...
The incidence of cerebral infarction is ≥ 4% among patients with the score of CHADS2 ≥ 2, so all of four NOACs and VKA are recommended [11, 13]. CHA2DS2-V ASc score considering over 65 years of age, sex category, vascular diseases including prior myocardial infarction of ≥ 1 is also recommended with all of four NOACs and VKA [12, 13]. However we have no definite consensus for selection of NOACs. Our eligibility for anticoagulant drugs and selection of NOACs which had been freely determined by each physician is equal to recent multicenter prospective cohort study (da Vinci study) [14].

RE-LY trial showed that the rate of hemorrhagic stroke was 0.38% per year in the VKA group, as compared with 0.12% per year with 110 mg of dabigatran (P<0.001) and 0.10% per year with 150 mg of dabigatran (P<0.001) [1]. ROCKET-AF trial showed that rivaroxaban group showed significant reduction in intracranial hemorrhage as compared with VKA group (0.5% vs. 0.7%, P=0.02) [4]. ARISTOTLE trial showed that the rate of major bleeding was 2.13% per year in the apixaban group, as compared with 3.09% per year in the VKA group (hazard ratio, 0.69; 95% CI, 0.60 to 0.80; P<0.001)[3].

Table 3: Stroke occurrence during NOACs.
DA: Dabigatran, RI: Rivaroxaban, AP: Apixaban,
Type of stroke: CE: Cardioembolic Stroke, AT: Atherothrombotic Brain Infarction, TIA: Transient Ischemic Attack, CH: Cerebral Hemorrhage

Table 4: 10 recurrent stroke cases with NOACs and 7 recurrent stroke cases with VKA.
Type of stroke: CE: Cardioembolic Stroke, AT: Atherothrombotic Brain Infarction, TIA: Transient Ischemic Attack, CH: Cerebral Hemorrhage
*p<0.05
All of three NOACs have lower rate in major bleeding and intracranial hemorrhage in these major studies and superior to VKA in prevention of bleeding events. Patients with HAS-BLED score ≥ 3 have high risk for major bleeding, so all of our three NOACs groups with mean HAS-BLED score ≥ 3 might been adequate to use NOACs [15].

Our NOACs group is significantly lower in CHADS2, CHA2DS2-VASc, HAS-BLED, NIHSS on admission and discharge than VKA group. And our VKA group in recurrent case has significantly higher rate of CKD with lower value of Ccr. Toyoda et al. and SAMURAI Study Investigators reported that post stroke patients taking NOACs due to NVAF had more frequently lower scores for CHADS2, CHA2DS2-VASc, HAS-BLED, admission NIHSS and discharge mRS [16]. Fujimoto et al reported that NOACs were preferentially chosen for the patients with lower severity and better renal function in current status of choice of oral anticoagulants [17]. Our results equal to these reports in Japan. Our results showing the rate of stroke recurrence with 25.9% in VKA group and 17.5% in NOACs group without significant difference between two groups also equals to mega trial in ROCKET-AF and sub analysis in ARISTOTLE trial [3,4,18]. The reason for the characteristic of dabigatran could not be not crushed in ROCKET-AF and sub analysis in ARISTOTLE trial [3,4,18].

The reason for the characteristic of dabigatran could not be not crushed and were not fitted with a nasogastric tube in severe stroke patients with dysphagia. So our group might be tend to choice dabigatran for patients with milder neurological symptom[6,14]. Following oral administration of apixaban, urinary excretion is approximately 27% of the total clearance in healthy subjects, and it is lowest elimination in kidney among four NOACs [19-21]. Relative low renal excretion could make us choice apixaban in subjects with renal dysfunction. Exactly our patients treated with apixaban was 54.5 (mL/min) in creatinine clearance and is not significantly lower than two other NOACs.

We can see several studies about stroke volume and anti-coagulant therapy. Hagii et al. reported that rivaroxaban-associated intracranial hemorrhage (ICH) had relatively small hematoma and no expansion of hematoma compared with VKA-associated ICH [7]. Matsumoto M et al reported that therapeutic anticoagulation by VKA (PT-INR=1.6) reduces infarct volume and improves neurological outcome after ischemic stroke in patients with NVAF [22]. Hakan et al. reported that patients who were on therapeutic PT-INR (>2.0) had smaller infarcts compared with patients without preadmission V ka use (p<0.001). Preadmission V ka use associated with therapeutic level of anticoagulation can offer a benefit in limiting the extent of ischemic injury and volume in an event of acute stroke [23]. NOACs could act as neuro-protectors by inhibiting the activation of matrix metalloproteinase-9 which significantly reduces brain infarct size in early inhibition [24,25]. We hypothesized that NOACs might contribute stroke volume at minimum because of decreased ischemic stroke events [5]. Coagulation Factor X is an important amplifier of both the intrinsic and extrinsic activation pathways. Under-dose of V KA with sub-therapeutic PT-INR values were found in 74% of first ischemic stroke with NVAF and 68% of those with recurrent stroke [26,27]. Of our 7 recurrent ischemic stroke with VKA 5 patients (71%) were under sub-therapeutic PT-INR value, and its percentage is near the previous reports. Direct Factor Xa inhibitor, rivaroxaban cause more permeable to flow in fibrin network and decreased degradation of plasma clots being formed with a looser structure in the presence of ribaroxaban [28]. The efficacy of stable blood concentration, the direct thrombin inhibitor or the factor Xa inhibitors as specific and narrow blockers in coagulating pathways with NOACs might contribute to less stroke event and smaller volume in ischemic stroke.

Study Limitations

Our study may have been influenced by relatively small sample size and retrospective study. Randomized control trial will be needed to make anticoagulant treatment definitively.

Conclusion

Secondary prevention with NOACs after stroke might be more beneficial by reducing infarct volume and modified Rankin Scale in recurrent stroke compared to VKA, warfarin.

Competing Interests

The authors declare that they have no conflict of interest in this work.

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