Ticagrelor plus aspirin versus clopidogrel plus aspirin for platelet reactivity in patients with minor stroke or transient ischaemic attack: open label, blinded endpoint, randomised controlled phase II trial

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ABSTRACT

OBJECTIVE
To test the hypothesis that ticagrelor plus aspirin is safe and superior to clopidogrel plus aspirin for reducing high platelet reactivity at 90 days and stroke recurrence in patients with minor stroke or transient ischaemic attack, particularly in carriers of the CYP2C19 loss-of-function allele and patients with large artery atherosclerosis.

MAIN OUTCOME MEASURES
Primary outcome was the proportion of patients with high platelet reactivity at 90 days. High platelet reactivity was defined as P2Y12 reaction units of more than 208. Secondary outcomes included high platelet reactivity at 90 days (7 days either way) in patients carrying genetic variants that would affect clopidogrel metabolism, and any stroke (ischaemic or haemorrhagic) recurrence at 90 days (7 days either way), six months, and one year.

RESULTS
At 90 days, high platelet reactivity occurred in 35 (12.5%) of 280 patients in the ticagrelor/aspirin group and 86 (29.7%) of 290 patients in the clopidogrel/aspirin group (risk ratio 0.40; 95% confidence interval 0.31; 0.49; P<0.001). Patients with large artery atherosclerosis in the ticagrelor/aspirin group had a lower stroke recurrence at 90 days than those in the clopidogrel/aspirin group (6.0% v 0.3%; hazard ratio 0.28; 95% confidence interval 0.18 to 0.49; P=0.001) of patients carrying CYP2C19 loss-of-function alleles. Stroke occurred in 21 (6.3%) of 336 patients in the ticagrelor/aspirin group and 30 (8.8%) of 339 patients in the clopidogrel/aspirin group (hazard ratio 0.70; 95% confidence interval 0.40 to 1.22; P=0.20). Patients with large artery atherosclerosis in the ticagrelor/aspirin group had a lower stroke recurrence at 90 days than those in the clopidogrel/aspirin group (6.0% v 13.1%; hazard ratio 0.45; 95% confidence interval 0.20 to 0.98; P=0.04).

No difference was seen in the rates of major or minor haemorrhagic events between the ticagrelor/aspirin and clopidogrel/aspirin groups (4.8% v 3.5%; P=0.42).

CONCLUSION
Patients with minor stroke or transient ischaemic attack who are treated with ticagrelor plus aspirin have a lower proportion of high platelet reactivity than those who are treated with clopidogrel plus aspirin, particularly for those who are carriers of the CYP2C19 loss-of-function allele. The results of this study should be evaluated further in large scale, phase III trials and in different populations.

TRIAL REGISTRATION
Clinicaltrials.gov NCT02506140.

WHAT IS ALREADY KNOWN ON THIS TOPIC

Studies have shown that patients who are carriers of the cytochrome P450 (CYP) 2C19*2 and *3 loss-of-function alleles do not benefit from dual antiplatelet therapy (aspirin combined with clopidogrel), compared with aspirin alone. Ticagrelor combined with aspirin has been shown to be more efficacious than clopidogrel combined with aspirin for acute coronary syndromes, regardless of CYP2C19 genotypes. However, the safety and efficacy of ticagrelor/aspirin versus clopidogrel/aspirin has not been evaluated in patients with minor stroke or transient ischaemic attack.

WHAT THIS STUDY ADDS

This study suggests the efficacy of ticagrelor/aspirin in reducing high platelet reactivity compared with clopidogrel/aspirin, especially in patients with CYP2C19 loss-of-function alleles at 90 days after symptoms onset. The rate of major or minor haemorrhagic events did not differ between the two groups. As a phase II trial, these results would need to be replicated and investigated further in larger studies and in different populations in the future.
that combined clopidogrel and aspirin treatment is superior to aspirin alone in reducing the risk of stroke, but could increase the risk of non-intracranial haemorrhage. Additionally, about 50% patients with acute ischaemic stroke had a risk of intracranial large artery atherosclerosis (LAA) in Asia, and patients with intracranial arterial stenosis and minor stroke (or a high risk of transient ischaemic attack) had a higher rate of recurrent stroke than those without. The CHANCE genetic substudy showed that patients who were carriers of the cytochrome P450 (CYP) 2C19*2 and *3 loss-of-function alleles benefitted more from using aspirin alone than from using dual antiplatelet therapy.

The metabolism of ticagrelor is primarily via the CYP3A4 enzyme and does not involve CYP2C19, unlike clopidogrel. A genetic substudy of the Platelet Inhibition and Patient Outcomes (PLATO) trial indicated that ticagrelor is more efficacious than clopidogrel for acute coronary syndromes, regardless of CYP2C19 genotype, but was associated with an increased risk of haemorrhage in patients with a history of stroke. The Acute Stroke or Transient Ischaemic Attack Treated With Aspirin or Ticagrelor and Patient Outcomes (SOCRATES) trial revealed a trend towards better efficacy in reducing the risk of vascular events in the ticagrelor treated group than in the aspirin group in an Asian subpopulation. However, limited data are available on the safety and efficacy of ticagrelor for the treatment of stroke, compared with data for clopidogrel on a background of aspirin in patients with acute stroke.

High platelet reactivity is defined as resistance or non-responsiveness to antiplatelet agents and is a known marker for recurrent ischaemic events in patients with acute coronary syndrome or those patients with percutaneous coronary intervention. Several studies have shown the predictive value of high platelet reactivity for ischaemic and bleeding events after percutaneous coronary intervention or in patients with acute coronary syndrome. Multiple factors can contribute to the variability in platelet function testing results, thus defining the high platelet reactivity status. High platelet reactivity is associated with poor cerebrovascular outcomes, and might be of clinical value for the evaluation of recurrent events in patients with stroke.

We conducted the Platelet Reactivity in Acute Stroke or Transient Ischaemic Attack (PRINCE) II study to compare the efficacy of ticagrelor plus aspirin with clopidogrel plus aspirin in reducing high platelet reactivity at 90 days in patients with minor stroke or transient ischaemic attack. We also compared the clinical outcomes in terms of efficacy and safety before a large scale, phase III study.

Methods

The PRINCE study protocol (NCT02506140) and data collection were approved by the ethics committee of Beijing Tiantan Hospital (ethical approval number KY2014-048-03) and all of the study centres, and conducted in accordance with the Declaration of Helsinki. The trial was a prospective, multicentre, randomised, open label, active controlled, blinded endpoint trial. All participants or their representatives provided written consent before study enrolment.

Study design and participants

From August 2015 to March 2017 in 26 study centres in China, the PRINCE trial enrolled patients aged 40-80 years who had an acute minor ischaemic stroke (National Institutes of Health Stroke Scale score of ≤3 at the time of randomisation) or those with a moderate to high risk of transient ischaemic attack (ABCD² stroke risk score of ≥4 at the time of randomisation or ≥50% stenosis of cervical or intracranial vessels that could account for the presentation) who could be treated with the study drug within 24 hours of symptom onset. Patients were excluded from the trial if they had a diagnosis intracranial haemorrhage, acute coronary syndrome, or other pathology that could account for the neurological symptoms; had a modified Rankin scale score of more than 2 at randomisation; or had a contraindication to ticagrelor, clopidogrel, or aspirin.

The trial included six visits: randomisation (baseline), seven to nine days, 21 days (two days either way), 90 days (seven days either way), six months (14 days either way), and one year (14 days either way). Additional visits at two and 24 hours after the first dose was administered were optional. All visits involved face-to-face interviews, with the exception of the six month follow-up, which was conducted by telephone, with data collected on electronic case report forms. The general schedule of the trial and the collection times for blood and urine samples are listed in the supplementary appendix (blood and urine samples collecting schedule).

Randomisation and procedures

Immediately after signing the written informed consent form, eligible patients were assigned to receive the following within one hour of randomisation, in a 1:1 ratio:

- Intervention (ticagrelor/aspirin): aspirin (a loading dose of 100-300 mg given as one to three 100 mg tablets on day 1, followed by 100 mg once daily until day 21) combined with ticagrelor (180 mg loading dose given as two 90 mg tablets on day 1, followed by 90 mg twice daily until day 90)
- Control (clopidogrel/aspirin): aspirin (a loading dose of 100-300 mg given as one to three 100 mg tablets on day 1, followed by 100 mg once daily until day 21) combined with clopidogrel (300 mg loading dose given as four 75 mg tablets on day 1, followed by 75 mg once daily until day 90).

Patients were allocated via a block randomisation process by investigators at the clinical centres. The block randomisation sequence was provided by an independent statistician using computer generated random numbers with a block size of four. The block size was not listed in the Chinese version of the protocol
(which was provided to the investigators) in order to prevent the investigators from speculating about the group assignment. The loading and maintenance doses of ticagrelor were administered as in the SOCRATES and PLATO studies.\textsuperscript{10,18}

VerifyNow testing was conducted in each study centre by qualified personnel who were blinded to the treatment allocation. The platelet reaction units were measured in each study centre by specially trained and qualified personnel according to a standardised procedure manual. To ensure the validity and reproducibility of the assay, we held two separate training course for all the testing personnel from each centre. Both the investigators and the patients were aware of the study drug assignment, but were blinded to the platelet reactivity data until the end of the trial.

Outcomes
The primary outcome of the PRINCE trial was the proportion of patients with high platelet reactivity at 90 days. High platelet reactivity was defined as a P2Y12 reaction unit of more than 208 measured using the VerifyNow P2Y12 assay. Prespecified secondary outcomes included high platelet reactivity at 90 days (seven days either way) in patients carrying genetic variants that would affect clopidogrel metabolism; any stroke (ischaemic or haemorrhagic); and composite clinical vascular events (ischaemic/haemorrhagic stroke, transient ischaemic attack, myocardial infarction, or vascular death) at 90 days (seven days either way), six months, and one year. Each reported composite clinical vascular event and safety outcome was independently adjudicated by two members (KD and Jimei Li) of the clinical event adjudication committee, who were blinded to the treatment group assignments. All discrepancies were reviewed by all five members of the committee and resolved by consensus.

The primary safety outcome was major bleeding, which was defined as that in the PLATO study classification of haemorrhagic events: fatal or life threatening bleed, major bleed, and other (supplementary appendix, PLATO bleeding classification). Secondary safety outcomes included the incidence of intracranial bleeding; dyspnoea events; and mortality at 90 days (seven days either way), six months, and one year.

Genotyping
The prespecified analysis included the single nucleotide polymorphisms (SNPs) CYP2C19*2 (681G >A, rs4244285), CYP2C19*3 (636G >A, rs4986893), and CYP2C19*17 (−806C >A, rs12248560), which were genotyped in all participants with adequate blood samples. Most genotyping of the three SNPs was performed by the Sequenom MassARRAY iPLEX platform (Sequenom, San Diego, CA, USA). We used Sanger sequencing (ABI 3500 Genetic Analyzer, Applied Biosystems) if the results were otherwise inconclusive.

We used star allele nomenclature to categorise patients by the CYP2C19 metaboliser status, based on *2, *3, and *17 genotypes.\textsuperscript{19} Patients with at least one gain-of-function allele (*17) were classified as “gain-of-function allele carriers,” and those with at least one loss-of-function allele (*2 or *3) were classified as “loss-of-function allele carriers.”\textsuperscript{20} Patients who carried at least one *17 allele (*1/*17 or *17/*17) were classified as “ultra-metabolisers,” those without any *2, *3, or *17 allele (*1/*1) were classified as “poor metabolisers,” and those with at least two *2 or *3 alleles (*2/*2, *2/*3, or *3/*3) were classified as “extensive metabolisers.”\textsuperscript{20} Patients with one *17 and a loss-of-function allele (*2/*17 or *3/*17) were classified as “unknown metabolisers,” because the clinical effect of these alleles is uncertain.\textsuperscript{21,22}

Based on our preplanned aims, we compared the recurrence of stroke in patients whose stroke subtype was intracranial large artery atherosclerosis (LAA) with those whose stroke subtype was non-LAA based on the stroke subtype of SSS-TOAST (Stop Stroke Trial of Org 10172 in Acute Stroke Treatment stroke aetiology classification; supplementary appendix, SSS-TOAST classification criteria). Non-LAA included cardioaortic embolism, small artery occlusion, other causes, and undetermined causes.

Statistical analysis
An interim analysis was preplanned in the published protocol.\textsuperscript{17} We calculated that 952 patients (estimated 10% dropout rate) would be required to achieve 90% power with a two sided α=0.05 to detect a relative reduction of 24% in the proportion of the primary outcome in the ticagrelor/aspirin group compared with that in the clopidogrel/aspirin group. The data safety monitoring board opted to terminate the study after the interim analysis based on 476 patients (50% of the projected necessary sample size) who completed 90 days of follow-up, based on achieving a prespecified threshold for efficacy (P<0.005). At the time of this decision, an additional 199 patients had already been recruited and randomised into the trial, and these patients were also followed up to study completion. Therefore, a total of 675 patients were included in the intention-to-treat analyses.

Proportions were presented for categorical variables, and medians with interquartile ranges or means (standard deviation) were presented for continuous variables. We compared the proportion of high platelet reactivity at the 90 day follow-up (the primary outcome) between the two study groups using genmod models adjusted by the high platelet reactivity status at baseline, reported as a risk ratio with 95% confidence intervals. To evaluate the influence of missing data for the primary outcome, we did sensitivity analyses assuming that all the missing data were high platelet reactivity or not.

The differences in the rates of stroke, composite outcome, death, and bleeding events during the 90 day follow-up were assessed by use of Cox proportional hazards regression, and were reported as hazard ratios with 95% confidence intervals. The proportional hazard assumption for the Cox models...
was examined by including a time dependent covariate with interaction of treatment group, and a logarithmic function of survival time in the model.

We assessed whether the treatment effect differed in certain genotype categories by testing the treatment-by-genotype interaction effect in genmod models for the primary outcome and Cox models for other outcomes, as described earlier. We also tested whether the treatment effect in reducing stroke recurrence differed between patients with LAA and those without LAA by testing the treatment-by-stroke subtype interaction effect in a genmod model. The models also included the main effects of treatment group and genotype or stroke subtype. All statistical analyses were two sided, and differences with a P value of less than 0.05 were considered to be statistically significant. Analyses were performed on the full population.

| Table 1 | Baseline characteristics of patients in the PRINCE trial |
|----------|----------------------------------------------------------|
| Characteristic | Trial group | |
| | Ticagrelor/aspirin (n=336) | Clopidogrel/aspirin (n=339) |
| Age (years) | |
| Mean (standard deviation) | 61.1 (8.5) | 60.5 (9.0) |
| Median (interquartile range) | 62.0 (55.0-67.0) | 61.0 (54.0-67.0) |
| Female sex (No (%)) | 91 (27.1) | 90 (26.5) |
| Systolic blood pressure (mm Hg) | |
| Mean (standard deviation) | 152.3 (22.5) | 154.9 (21.2) |
| Median (interquartile range) | 150.0 (137.5-168.0) | 154.0 (140.0-170.0) |
| Diastolic blood pressure (mm Hg) | |
| Mean (standard deviation) | 87.7 (13.0) | 89.4 (12.8) |
| Median (interquartile range) | 87.5 (80.0-96.0) | 88.0 (80.0-97.0) |
| Body mass index* | |
| Mean (standard deviation) | 25.0 (3.8) | 25.0 (3.8) |
| Median (interquartile range) | 24.6 (22.6-27.0) | 24.8 (22.7-27.3) |
| Pulse rate (beat/min; mean (SD)) | 75.1 (10.1) | 76.3 (11.5) |
| Medical history (No (%)) | |
| Hypertension | 203 (60.4) | 208 (61.4) |
| Glycaemia | 20 (6.0) | 21 (6.2) |
| Diabetes mellitus | 79 (23.5) | 85 (25.1) |
| Ischaemic stroke | 59 (17.6) | 62 (18.3) |
| Transient ischaemic attack | 8 (2.4) | 10 (2.9) |
| Coronary artery disease | 26 (7.7) | 25 (7.4) |
| Known atrial fibrillation | 0 (0.0) | 4 (1.2) |
| Flutter valvular heart disease | 1 (0.3) | 0 (0.0) |
| Pulmonary embolism | 0 (0.0) | 0 (0.0) |
| Smoking status (No (%)) | |
| Non-smoker | 150 (44.6) | 155 (45.7) |
| Current smoker | 160 (47.6) | 159 (46.9) |
| Ex-smoker | 26 (7.7) | 25 (7.4) |
| Drug use before randomisation (No (%)) | |
| Proton pump inhibitor | 2 (0.6) | 3 (0.9) |
| Statin | 36 (10.7) | 30 (8.8) |
| Aspirin | 77 (22.9) | 69 (20.4) |
| Clopidogrel | 5 (1.5) | 10 (2.9) |
| Ticagrelor | 0 (0.0) | 0 (0.0) |
| Time to randomisation after onset of symptoms (h; mean (range)) | 14.0 (8.3-20.6) | 13.8 (8.0-20.8) |
| Qualifying event (No (%)) | |
| Minor stroke | 275 (81.8) | 289 (85.3) |
| Transient ischaemic attack | 61 (18.2) | 50 (14.7) |
| Baseline ABCD² score among patients with transient ischaemic attack as the qualifying event (median (interquartile range))| 5.0 (4.0-5.0) | 4.5 (4.0-5.0) |
| SSS-TOAST stroke subtype (No (%))| |
| Large artery atherosclerosis | 151 (54.9) | 153 (52.9) |
| Cardioaortic embolism | 8 (2.9) | 5 (1.7) |
| Small artery occlusion | 104 (37.8) | 109 (37.7) |
| Other causes | 7 (2.5) | 9 (3.1) |
| Undetermined causes | 5 (1.8) | 13 (4.5) |
| Unknown | 2 (0.7) | 7 (2.4) |
| Unclassified | 3 (1.1) | 6 (2.1) |

*Body mass index is the weight in kilograms divided by the square of the height in metres.

ABCD² stroke risk scores range from 0 to 7, with higher scores indicating higher risk; data provided in the table are only for the group of 111 patients whose qualifying event was transient ischaemic attack for inclusion in the trial.

SSS-TOAST stroke subtype=Stop Stroke Study Trial of Org 10172 in Acute Stroke Treatment stroke aetiology classification (supplementary appendix, SSS-TOAST classification criteria); data provided in the table are only for the group of 564 patients whose qualifying event was minor stroke for inclusion in the trial.
performed by use of SAS software, version 9.4 (SAS Institute, Cary, NC, USA).

**Patient and public involvement**
The design, outcome measurement, recruiting plans, or implementation of the study were independent of any patient. The gene and platelet reactivity testing results of every patient will be delivered to the patient himself or his appointed relatives after the primary results of the results published through email or telephone. The results of the research will be broadcasted to all the participants and general public through internet news, popular science articles, newspapers, and social media.

**Results**
Between August 2015 and March 2017, 5644 patients with stroke or transient ischaemic attack were screened at 26 hospitals, and 675 patients (mean age...
Among the 675 patients enrolled, 650 (mean age 60.8 [standard deviation 8.7]) had complete genotype data for all three SNPs, of whom 321 were randomly assigned to the ticagrelor/aspirin group and 329 were assigned to the clopidogrel/aspirin group (supplementary tables B, C, and D show details of the baseline characteristics of patients enrolled in the genetic substudy). Of the 650 participants, 374 (57.5%) were classified as CYP2C19 loss-of-function carriers.

We obtained valid measurements in 627 (92.9%) and 570 (84.4%) patients for the VerifyNow P2Y12 assay at the seven day and 90 day follow-up periods, respectively. The P2Y12 reaction units before receiving the study drugs were similar in the ticagrelor/aspirin and clopidogrel/aspirin groups (mean 256.4 [standard deviation 61.3] v 246.9 [53.7]; P=0.13). The ticagrelor/aspirin group had significantly fewer P2Y12 reaction units than the clopidogrel/aspirin group (69.3 [87.0] v 173.5 [67.6]; P<0.001) at 90 day follow-up (fig 2).

The primary outcome (high platelet reactivity) was observed in 35 of 280 patients (12.5%) in the ticagrelor/aspirin group, and in 86 of the 290 patients (29.7%) in the clopidogrel/aspirin group at 90 days (risk ratio 0.40; 95% confidence interval 0.28 to 0.56; P<0.001; table 2). Similar results were observed for the primary outcome in sensitivity analyses assuming that all missing data for the primary outcome showed high platelet reactivity or not (supplementary table E). The primary safety outcome (PLATO major hemorrhagic event) occurred in five patients (1.5%) in the ticagrelor/aspirin group and two (0.6%) in the clopidogrel/aspirin group, and in 8 (2.6%) and 1 (0.3%) of the 336 patients in the ticagrelor/aspirin and clopidogrel/aspirin groups, respectively. The P2Y12 reaction units before receiving the study drugs were similar in the ticagrelor/aspirin and clopidogrel/aspirin groups (mean 256.4 [standard deviation 61.3] v 246.9 [53.7]; P=0.13). The ticagrelor/aspirin group had significantly fewer P2Y12 reaction units than the clopidogrel/aspirin group (69.3 [87.0] v 173.5 [67.6]; P<0.001) at 90 day follow-up (fig 2).

Among the patients with LAA, stroke recurrence at 90 days was markedly lower in the ticagrelor/aspirin group (10.8% v 35.4%; risk ratio 0.31, 95% confidence interval 0.18 to 0.49; P<0.001; supplementary table F). The event rates of stroke, composite events, and haemorrhagic events varied by treatment assignment and phenotype (fig 3). Other primary and secondary outcomes in patients with available genotype data are listed in supplementary table G.

Among the patients with LAA, stroke recurrence at 90 days was markedly lower in the ticagrelor/aspirin group than in the clopidogrel/aspirin group (6.0% v 13.1%; hazard ratio 0.45, 95% confidence interval 0.20 to 0.98; P=0.04; table 3 and fig 4). However, among patients with non-LAA, the risk of stroke recurrence was similar in the ticagrelor/aspirin group and clopidogrel/aspirin groups (8.1% v 7.4%; 1.1, 0.46 to 2.63; P=0.84; table 3 and fig 4). The proportional hazard assumption was met (P=0.93).

The primary safety outcome (PLATO major hemorrhagic event) occurred in five patients (1.5%) in the ticagrelor/aspirin group and four (1.2%) in the clopidogrel/aspirin group (hazard ratio 1.27; 95% confidence interval 0.34 to 4.72; table 2). Three patients (0.9%) in the ticagrelor/aspirin group and two (0.6%)
in the clopidogrel/aspirin group had intracranial haemorrhage. However, the rate of any haemorrhagic events occurring was higher in the ticagrelor/aspirin group (22.3%) than in the clopidogrel/aspirin group (14.2%; 1.65, 1.15 to 2.37; table 2). All of the proportional hazard assumptions were met (P=0.99 for major haemorrhagic event and P=0.82 for any haemorrhagic events). The rate of major bleeding did not vary significantly between the ticagrelor/aspirin and clopidogrel/aspirin groups amongst the carriers of the CYP2C19 loss-of-function allele (rate 0.0% v 0.04 to 3.33; P=0.43) and non-carriers (0.0% v 0.35, 95% confidence interval 0.04 to 3.33; P=0.43) and non-carriers (0.0% v 0.7%; supplementary table G).

Dyspnoea was more common in the ticagrelor/aspirin group (n=54, 16.1%) than in the clopidogrel/aspirin group (n=11, 3.2%; supplementary table H). A total of 69 (20.5%) patients in the ticagrelor/aspirin group and 47 (13.9%) in the clopidogrel/aspirin group stopped receiving the study drug before 90 days (fig 1); the most common reasons were dyspnoea and epistaxis. The rate of permanent discontinuation caused by dyspnoea was 4.2% (14/336) in the ticagrelor/aspirin group and 0.0% in the clopidogrel/aspirin group, and the rate of permanent discontinuation caused by epistaxis was 1.8% (n=6) in the ticagrelor/aspirin group and 0.0% in the clopidogrel/aspirin group (supplementary table 1). Serious adverse events and adverse events leading to permanent drug discontinuation within 90 days are listed in supplementary table 1.

### Discussion

#### Principal findings

This PRINCE trial results indicated that the proportion of high platelet reactivity at 90 days reduced with ticagrelor compared with clopidogrel, in patients with acute minor stroke and those at moderate-to-high risk of transient ischaemic attack treated with aspirin. This trial was not powered to study clinical events. However, we observed fewer strokes and composite outcomes at 90 days in patients who were treated with dual antiplatelet therapy using ticagrelor/aspirin compared with clopidogrel/aspirin within 24 hours of the onset of minor stroke or transient ischaemic attack, without increasing the risk of major, minor, or intracranial haemorrhage. Treatment discontinuation was, however, higher with ticagrelor due to an increased rate of dyspnoea and minimal haemorrhagic events (that is, epistaxis).

#### Comparison with other studies

The major haemorrhage rate in the present study was lower than that found in the PLATO study (ticagrelor/aspirin group 1.5% v 11.6%; clopidogrel/aspirin group

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**Table 2 | Effect of ticagrelor/aspirin versus clopidogrel/aspirin on efficacy and safety outcomes in PRINCE trial**

| Outcomes                                      | Trial participants (No with event/total No (%)) | Hazard ratio or risk ratio (95% CI)* | P   |
|-----------------------------------------------|-----------------------------------------------|-------------------------------------|-----|
| **Primary efficacy outcomes†**                |                                               |                                     |     |
| Baseline                                      | 268/333 (80.5)                                | 260/336 (77.4)                      | 1.04 (0.96 to 1.13) | 0.33 |
| 7 ± 2 days                                    | 12/306 (3.9)                                  | 89/321 (27.7)                       | 0.14 (0.07 to 0.23) | <0.001|
| 90 ± 7 days                                   | 35/280 (12.5)                                 | 86/290 (29.5)                       | 0.40 (0.28 to 0.56) | <0.001|
| **Secondary efficacy outcomes**               |                                               |                                     |     |
| Stroke                                        | 21/336 (6.3)                                  | 30/339 (9.0)                        | 0.70 (0.40 to 1.22) | 0.20 |
| Composite events‡                            | 22/336 (6.5)                                  | 12/339 (3.6)                        | 0.68 (0.40 to 1.18) | 0.17 |
| Ischaemic stroke                              | 33/336 (9.9)                                  | 27/339 (8.0)                        | 0.64 (0.35 to 1.16) | 0.14 |
| Haemorrhagic stroke                           | 5/336 (1.5)                                   | 2/339 (0.6)                         | 1.52 (0.25 to 9.08) | 0.65 |
| Myocardial infarction                         | 0/336 (0.0)                                   | 1/339 (0.3)                         | —    | —    |
| Death from cardiovascular causes              | 1/336 (0.3)                                   | 2/339 (0.6)                         | 0.50 (0.05 to 5.55) | 0.58 |
| Death from any cause                          | 3/336 (0.9)                                   | 2/339 (0.6)                         | 1.50 (0.25 to 9.00) | 0.65 |
| Transient ischaemic attack                    | 1/336 (0.3)                                   | 2/339 (0.6)                         | 0.50 (0.05 to 5.53) | 0.57 |
| **Primary safety outcomes§**                 |                                               |                                     |     |
| Major bleeding                                | 5/336 (1.5)                                   | 4/339 (1.2)                         | 1.27 (0.34 to 4.72) | 0.72 |
| Major, fatal, life threatening bleeding       | 4/336 (1.2)                                   | 3/339 (0.9)                         | 1.35 (0.30 to 6.03) | 0.69 |
| Fatal bleeding                                | 1/336 (0.3)                                   | 1/339 (0.3)                         | 1.01 (0.06 to 16.13) | 1.00 |
| Intracranial haemorrhage                      | 33/336 (9.9)                                  | 27/339 (8.0)                        | 1.27 (0.34 to 4.72) | 0.72 |
| Major, other                                  | 1/336 (0.3)                                   | 1/339 (0.3)                         | 1.01 (0.06 to 16.18) | 0.99 |
| Minor bleeding                                | 11/336 (3.3)                                  | 8/339 (2.4)                         | 1.40 (0.56 to 3.47) | 0.47 |
| Major or minor bleeding                       | 16/336 (4.8)                                  | 12/339 (3.5)                        | 1.36 (0.64 to 2.88) | 0.42 |
| Minimal bleeding                              | 64/336 (19.0)                                 | 36/339 (10.6)                       | 1.86 (1.24 to 2.80) | 0.003|
| Any bleeding                                  | 75/336 (22.3)                                 | 48/339 (14.2)                       | 1.65 (1.15 to 2.37) | 0.007|
| **Other safety outcomes**                     |                                               |                                     |     |
| Respiratory, thoracic, and mediastinal disorders | 22/336 (6.5)                               | 0/339 (0.0)                         | —    | <0.001|
| Dyspnoea                                      | 14/336 (4.2)                                  | 0/339 (0.0)                         | —    | <0.001|
| Epistaxis                                     | 6/336 (1.8)                                   | 0/339 (0.0)                         | —    | 0.04 |

| *Risk ratios used for the primary efficacy outcome and hazard ratios used for secondary efficacy outcome. |
| †Primary outcome indicates high platelet reactivity, which was defined as a P2Y12 reaction unit of more than 208, as measured by a VerifyNow P2Y12 assay. |
| ‡A composite event was defined as a new clinical vascular event, including stroke, transient ischaemic attack, myocardial infarction, or death from cardiovascular causes. |
| §Primary safety outcomes were defined according to the PLATO criteria (supplementary appendix, PLATO bleeding classification). All 675 patients were included in the analysis of safety outcomes. Other safety outcomes included those leading to permanent drug discontinuation. |
1.2% v 11.2%). This difference could be partly due to the short term use of dual antiplatelet therapy in our study compared with the PLATO study (21 v 277 days).

By contrast, a slightly higher major haemorrhagic rate was found in PRINCE than in the Asian subgroup in the SOCRATES trial (1.5% in the ticagrelor/aspirin group in PRINCE v 0.6% in the ticagrelor used as monotherapy in SOCRATES). This difference could be related to the combined use of two drugs in our study. Similar to that reported in the PLATO and SOCRATES trials, dyspnoea occurred more commonly with ticagrelor than with clopidogrel treatment.

Safety of study drug

The premature discontinuation of the study drug was more frequent in the ticagrelor/aspirin group than in the clopidogrel/aspirin group, although the rate of serious adverse events did not differ between the two groups. The treating physician or patient had the option of temporarily or permanently discontinuing

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**Table 3 | Stroke recurrence at 90 days, by cause**

| Cause of stroke* | Ticagrelor/aspirin (n=336) | Clopidogrel/aspirin (n=339) | Hazard ratio (95% CI)* | P | P for interaction |
|------------------|---------------------------|-----------------------------|-----------------------|---|------------------|
| Large artery atherosclerosis | 9/151 (6.0) | 20/153 (13.1) | 0.45 (0.20 to 0.98) | 0.04 | 0.13 |
| Non-large artery atherosclerosis | 10/124 (8.1) | 10/136 (7.4) | 1.10 (0.46 to 2.63) | 0.84 | — |

*Cause of stroke classified by the SSS-TOAST stroke subtype (SSS-TOAST=Stop Stroke Trial of Org 10172 in Acute Stroke Treatment stroke aetiology classification (supplementary appendix, SSS-TOAST classification criteria)). Non-large artery atherosclerosis included patients with cardioaortic embolism, small artery occlusion, other causes, and undetermined causes.
the drug in case of an adverse event, and could have taken this opportunity more frequently because of the open label design, when using an unapproved drug (that is, ticagrelor) in this indication. Furthermore, because early discontinuation and non-adherence are likely to contribute to the increased platelet reactivity observed at 90 days (compared with that observed at seven days), the efficacy of ticagrelor on high platelet reactivity could have been even higher in patients in the case of a blinded study design.

Clinical efficacy

Although we tested the clinical efficacy (recurrent stroke) between the two groups, the sample size was small and this phase II trial was not powered to find a clinical effect. Our finding of fewer recurrent stroke and composite events in patients treated with ticagrelor/aspirin than in those treated with clopidogrel/aspirin would need to be replicated, because the current study provided only a 25.5% power to show a 30% relative risk difference with a two sided test at a 5% significance level. Given a 90% power and a significance level of 5% (two sided), a total of 4690 patients would need to be included in a phase III trial to detect the relative risk difference between the ticagrelor/aspirin and clopidogrel/aspirin groups, based on the event rates in our study.

The stroke recurrence rate in LAA group was higher than that in the non-LAA group in our study. Patients with minor stroke or transient ischaemic attack with LAA might benefit more from ticagrelor plus aspirin than from clopidogrel plus aspirin treatment. Considering the large numbers required for the stroke subtyping analyses, our results were exploratory and hypothesis generating, so future studies are needed. We also found a relatively high (7.7%) stroke recurrence rate in patients with non-LAA (table 3), compared with studies conducted in other developed countries. This might be relevant with the overall recurrent stroke rate after minor stroke or transient ischaemic attack also being relatively high, because the recurrent stroke included a new stroke and also rapid worsening (National Institutes of Health Stroke Score ≥4) of an existing focal neurological deficit in our study.

Among patients randomised to clopidogrel, we saw a dose-response association between the number of CYP2C19 loss-of-function alleles and the proportion of high platelet reactivity, with patients carrying more loss-of-function alleles having a higher proportion of platelet aggregation. A larger benefit with ticagrelor compared with clopidogrel was observed in the proportion of high platelet reactivity in patients carrying more loss-of-function alleles. While a similar trend in clinical outcomes was seen among patients randomly allocated to clopidogrel, those allocated to ticagrelor also had higher rates of clinical events if they carried a greater number of loss-of-function allele. However, these results should be interpreted with caution, owing to the very low number of patients and events.

Limitations

This study had several limitations. Firstly, the primary outcome used high platelet reactivity as a marker of risk for events but not yet proved as a causal risk...
factor of thrombotic events. Further studies are needed to evaluate the clinical efficacy of dual antiplatelet therapy as the primary endpoint in this target population. Secondly, about 15% of patients were lost to follow-up for the evaluation of high platelet reactivity at 90 days. However, similar results were observed after assuming all the missing data were high platelet reactivity or not. Thirdly, potential selection bias could exist because we enrolled patients from sites that were mostly urban hospitals and that had more experts and medical resources. Fourthly, the cause of stroke and the genetic differences in the CYP2C19 gene differ between Chinese patients with stroke and European patients with stroke. The results of our study should be evaluated in different populations in the future. Finally, the open label design could have led to a placebo effect, 23 which might have caused potential bias in adverse events assessment, drug continuation, and even the physicians’ or patients’ decisions.

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Data sharing: The technical appendix, dataset, and statistical code are available from the corresponding author at yongjunwang@ncrcnd.org.cn.

The lead author affirms that this manuscript is an honest, accurate, and transparent account of the study being reported; that no important aspects of the study have been omitted; and that any discrepancies from the study as planned (and, if relevant, registered) have been explained.

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