

Cilostazol for secondary stroke prevention: systematic review and meta-analysis

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ABSTRACT

Background Stroke is one of the leading causes of death worldwide. Cilostazol, an antiplatelet and phosphodiesterase 3 inhibitor, has not been clearly established for ischaemic stroke use. We aim to determine the efficacy and safety of cilostazol for secondary stroke prevention.

Methods MEDLINE, EMBASE, Cochrane Library, Web of Science and ClinicalTrials.gov were searched from inception to 25 September 2020, for randomised trials comparing the efficacy and safety of cilostazol monotherapy or dual therapy with another antiplatelet regimen or placebo, in patients with ischaemic stroke. Version 2 of the Cochrane risk-of-bias tool for randomised trials (RoB 2) was used to assess study quality. This meta-analysis was reported in line with the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) statement.

Results Eighteen randomised trials comprising 11 429 participants were included in this meta-analysis. Most trials possessed low risk of bias and were of low heterogeneity. Cilostazol significantly reduced the rate of ischaemic stroke recurrence (risk ratio, RR=0.69, 95% CI 0.58 to 0.81), any stroke recurrence (RR=0.64, 95% CI 0.54 to 0.74) and major adverse cardiovascular events (RR=0.67, 95% CI 0.56 to 0.81). Cilostazol did not significantly decrease mortality (RR=0.90, 95% CI 0.64 to 1.25) or increase the rate of good functional outcome (Modified Rankin Scale score of 0–1; RR=1.07, 95% CI 0.95 to 1.19). Cilostazol demonstrated favourable safety profile, significantly reducing the risk of intracranial haemorrhage (RR=0.46, 95% CI 0.31 to 0.68) and major haemorrhagic events (RR=0.49, 95% CI 0.34 to 0.70).

Conclusions Cilostazol demonstrated superior efficacy and safety profiles compared with traditional antiplatelet regimens such as aspirin and clopidogrel for secondary stroke prevention but does not appear to affect functional outcomes. Future randomised trials can be conducted outside East Asia, or compare cilostazol with a wider range of antiplatelet agents.

BACKGROUND

Introduction

The rate of ischaemic stroke recurrence varies from 8%–14% at 1 year to 39% at 10 years,¹² and is associated with poor outcomes.³ To reduce the recurrence of ischaemic strokes,

antiplatelet therapies are widely adopted. Some established regimens include single antiplatelet therapy with aspirin,⁴ monotherapy with clopidogrel⁵ and combination aspirin and extended-release dipyridamole.⁶ Aspirin and clopidogrel combination can be considered for short-term stroke management. However, these antiplatelet therapies, especially long-term combination aspirin and clopidogrel, are known to increase the risk of haemorrhagic conversion and bleeding complications.^{7,8}

Cilostazol is an approved treatment for intermittent claudication and thrombotic complications of coronary angioplasty in combination with aspirin or clopidogrel.⁹ It selectively inhibits phosphodiesterase 3, increasing activation of intracellular cyclic adenosine monophosphate and protein kinase A, thereby inhibiting platelet aggregation. Apart from its antiplatelet properties, cilostazol may prevent recurrent ischaemic stroke by improving endothelial function,¹⁰ reducing triglycerides and increasing high-density lipoproteins.¹¹ The Chinese Guidelines for Secondary Prevention of Ischemic Stroke and Transient Ischemic Attack, Korean Clinical Practice Guidelines for Stroke, Japanese Guidelines for the Management of Stroke and other eastern institutions^{12–15} have accepted cilostazol as a second-line drug to aspirin and/or clopidogrel for secondary stroke prevention. However, cilostazol is neither widely recommended nor approved by most international guidelines.⁴

Objectives

Cilostazol is an accepted monotherapy for secondary stroke prevention in East Asian countries but is not commonly used outside Asia. Previous meta-analyses have also demonstrated the efficacy and safety of cilostazol to this end.^{16–18} However, those reviews do not include a new randomised trial evaluating the short-term use of cilostazol in acute ischaemic

stroke. Since studies performed on these subgroups are relatively scarce, it is useful to perform an updated meta-analysis to (1) confirm the effect of cilostazol as monotherapy; and (2) explore the effect of cilostazol in combination therapy. To investigate the efficacy of cilostazol-based therapies in different situations, we performed subgroup analyses on intracranial arterial stenosis (ICAS), specific antiplatelet regimens, stroke chronicity and duration of treatment. Unlike previous meta-analyses, we graded the quality of evidence for each outcome measure.

METHODS

Search strategy

We report the findings according to the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) statement.¹⁹ The authors screened multiple comprehensive databases (MEDLINE, EMBASE, Cochrane Library, Web of Science and ClinicalTrials.gov) for studies comparing cilostazol and other antiplatelet regimens for secondary prevention of ischaemic stroke.

The literature search was conducted for all publications in the databases from inception to 25 September 2020. We used keywords related to stroke (eg, 'stroke', 'cerebrovascular accident' or 'cerebral infarct') and cilostazol (eg, 'cilostazol', 'Pletal' or 'phosphodiesterase 3 inhibitor'). Medical Subject Headings (MeSH) and EMBASE subject headings (EMTREE) terms were used in conjunction with keywords to refine the search results (online supplemental data). Manual searches were also conducted on the reference lists of included studies and review articles to identify trials that were missed in the electronic search.

Study selection

The inclusion criteria encompassed any randomised trials that compared the efficacy and safety of cilostazol used as monotherapy or combination therapy with other antiplatelet agents, placebo or best medical therapy for the secondary prevention of ischaemic stroke. Non-randomised observational studies and conference proceedings were excluded.

The primary clinical outcome was ischaemic stroke recurrence. Safety outcomes comprised intracranial haemorrhages (ICH) and major haemorrhagic events. Secondary outcomes included any stroke recurrence, all-cause mortality, major adverse cardiovascular events (MACE) and adverse drug events (ADEs) leading to treatment discontinuation. Most studies defined ICH as intracerebral haemorrhage or subarachnoid haemorrhage. The definition for 'any stroke' comprised ischaemic stroke and either haemorrhagic stroke or ICH but did not include transient ischaemic attacks. Good functional outcome was defined as a Modified Rankin Scale score of 0–1 (mRS 0–1) at 3–6 months.

Two reviewers (TCH and WGRA) independently performed title and abstract screening on studies retrieved from the search strategy, followed by full-text screening. Data extraction and quality assessment were

performed independently. Any disagreements were resolved by consensus.

Data collection

The predesigned data extraction form included the following: (1) intervention and control; (2) participant characteristics, for example, age, National Institutes of Health Stroke Scale (NIHSS) on admission and country of study; (3) stroke subtypes; (4) time to randomisation or treatment; (5) duration of treatment and follow-up; and (6) outcome event rates.

Quality assessment

Quality assessment of the included studies was performed using Version 2 of the Cochrane tool for assessing risk of bias in randomised trials (RoB 2).²⁰ For each included study, signalling questions were answered to assess the risk of bias arising from: (1) the randomisation process; (2) deviations from the intended interventions; (3) missing outcome data; (4) the measurement of outcomes; and (5) the selection of reported results. Each component was individually assessed as having low, some concerns or high risk of bias.

The quality of evidence for each outcome was determined using the Grading of Recommendations Assessment, Development and Evaluation (GRADE) system consisting of five components: risk of bias, imprecision, inconsistency, indirectness and publication bias.²¹

Statistical analysis

The random-effects model was used to generate risk ratios (RRs) and associated 95% CIs. Where possible, the intention-to-treat (ITT) population was used. Heterogeneity was evaluated using the chi square (χ^2) and I^2 test. $I^2 < 25%$, $25\%–75%$ and $> 75%$ indicated low, moderate and high degree of inconsistency, respectively. Meta-regression was performed using the mixed-effects model. Publication bias was assessed using funnel plots. A p value ≤ 0.05 was considered statistically significant. All analyses were performed using Review Manager V.5.3 (The Cochrane Collaboration, 2014) and *meta* package V.4.13-0 in R (R Foundation for Statistical Computing, Vienna, Austria).

RESULTS

Study selection and study characteristics

A total of 2415 unique records were retrieved. Of these, the full text of 81 studies were screened (online supplemental data).

We included 18 randomised trials comprising 11 429 patients in the ITT population^{22–39} (tables 1 and 2). Nine studies were conducted in Japan, four in Korea, two in China, two in multiple East Asian countries and one in the UK. Most studies explicitly excluded patients with cardioembolic stroke, but two studies^{23 33} did not specify this. Three studies^{30 31 39} recruited participants with symptomatic ICAS, while two studies^{23 26} included patients with lacunar infarction. Although one study

Table 1 Characteristics of included studies

Study ID	Country	Sample size, ITT		Age (intervention)	Age (control)	Intervention	Control
		(% male)	(% female)				
Aoki 2019 (ADS) ²²	Japan	1201 (66%)	57 (68%)	69 (IQR 60–77)	69 (IQR 61–78)	Cilostazol 200 mg + aspirin 81–200 mg	Aspirin 81–200 mg
Blair 2019 (LACI-1) ²³	UK	57 (68%)	Mean of all participants: 66.1 (11.1)			Cilostazol 100 mg BD + aspirin/clopidogrel	Aspirin/clopidogrel
Gotoh 2000 (CSPS) ²⁴	Japan	1067 (66%)	1067 (66%)	65.2 (8.7)	65.1 (8.8)	Cilostazol 100 mg BD	Placebo
Guo 2009 ²⁵	China	68 (35%)	68 (35%)	59.44 (10.63)	62.02 (11.12)	Cilostazol 100 mg BD	Aspirin 100 mg
Han 2013 (ECLIPse) ²⁶	Korea	182 (25%)	182 (25%)	64.63 (9.07)	65.48 (9.92)	Cilostazol 100 mg BD + aspirin 100 mg	Aspirin 100 mg
Huang 2008 (CASISP) ²⁷	China	719 (69%)	719 (69%)	60.14 (10.05)	60.31 (9.71)	Cilostazol	Aspirin
Johkura 2012 ²⁸	Japan	106 (43%)	106 (43%)	76.7 (9.8)	73.7 (9.4)	Cilostazol 200 mg	Aspirin 100 mg
Kim 2018 (PICASSO) ²⁹	South Korea, Hong Kong, Philippines	1534 (62%)	1534 (62%)	65.5 (10.9)	65.8 (10.7)	Cilostazol 100 mg BD	Aspirin 100 mg
Kwon 2005 (TOSS) ³⁰	South Korea	135 (61%)	135 (61%)	62.28 (10.42)	62.54 (8.97)	Cilostazol 100 mg BD + aspirin 100 mg	Aspirin 100 mg
Kwon 2011 (TOSS-2) ³¹	4 East Asian countries	457 (51%)	457 (51%)	66.42 (11.33)	64.58 (11.11)	Cilostazol 100 mg BD + aspirin 75–150 mg	Clopidogrel 75 mg + aspirin 75–150 mg
Lee 2011 (CAIST) ³²	Korea	458 (61%)	458 (61%)	63 (12)	63 (12)	Cilostazol 200 mg	Aspirin 300 mg
Lee 2017 ³³	Korea	80 (65%)	80 (65%)	57.4 (12.7)	59.5 (11.7)	Cilostazol 100 mg BD	Aspirin 100 mg
Nakamura 2012 ³⁴	Japan	76 (74%)	76 (74%)	66 (12)	67 (10)	Cilostazol 100 mg BD + aspirin 300 mg	Aspirin 300 mg
Ohnuki 2017 ³⁵	Japan	24 (71%)	24 (71%)	60.5 (10)	63.6 (9.1)	Cilostazol 200 mg + aspirin 100 mg	Aspirin 100 mg
Shimizu 2013 ³⁶	Japan	507 (67%)	507 (67%)	66.2 (9.4)	66.6 (8.9)	Cilostazol 200 mg + optimal medical treatment	Optimal medical treatment
Shinohara 2010 (CSPS 2) ³⁷	Japan	2716 (72%)	2716 (72%)	63.5 (9.2)	63.4 (9.0)	Cilostazol 100 mg BD	Aspirin 81 mg
Toyoda 2019 (CSPS.com) ³⁸	Japan	1879 (70%)	1879 (70%)	69.6 (9.2)	69.7 (9.2)	Cilostazol 100 mg BD + aspirin 81–100 mg/clopidogrel 50–75 mg	Aspirin 81–100 mg/clopidogrel 50–75 mg
Uchiyama 2015 (CATHARSIS) ³⁹	Japan	163 (66%)	163 (66%)	68.3 (range 45–84)	68.3 (range 50–82)	Cilostazol 200 mg + aspirin 100 mg	Aspirin 100 mg

All data are in mean (SD) or median (IQR/range).

*After 14 days, both arms were swapped to cilostazol 200 mg until 3 months. Where possible, we analysed data at 14 days instead of 3 months. ITT, intention to treat.

Table 2 Details of included studies

Study ID	Time to randomisation/treatment	Duration of treatment/ follow-up	Type of stroke	
			Cilostazol	Control
Aoki 2019 (ADS) ²²	Within 48 hours Cilostazol: 10.1 hours (IQR 4.6–20.0) Control: 11.5 hours (IQR 4.8–20.9)	14 days, 3 months	Symptomatic ICAS (23%) Large-artery atherosclerosis (13%), LACI (46%), branch atheromatous disease (13%)	Symptomatic ICAS (22%) Large-artery atherosclerosis (15%), LACI (43%), branch atheromatous disease (16%)
Blair 2019 (LACI-1) ²³	Within 4 years Median: 203 days (range 6–920)	Treatment: 6–9 weeks Follow-up: 11 weeks	LACI	LACI
Gotoh 2000 (CSFS) ²⁴	1–6 months Cilostazol: 83.0 days (range 7–1805) Control: 82.4 days (range 8–1079)	Mean follow-up duration: Cilostazol: 651.8 days Control: 569.7 days Treatment duration: Cilostazol: 632.2 (467.7) days Control: 695.1 (456.3) days	Atherothrombotic (14.1%), LACI (75.0%), mixed (9.0%)	Atherothrombotic (13.1%), LACI (73.8%), mixed (11.4%)
Guo 2009 ²⁵	1–6 months	12 months		
Han 2013 (ECLIPse) ²⁶	Within 7 days Median: 5 days (78.3% of patients were randomised within 7 days)	90 days	LACI	LACI
Huang 2008 (CASISP) ²⁷	1–6 months Cilostazol: 77.26 (50.05) days Control: 79.72 (51.96) days	12–18 months (average: 740 person-years)		
Johkura 2012 ²⁸	1–6 months	6 months		
Kim 2018 (PICASSO) ²⁹	Within 180 days Cilostazol: 18 days (IQR 8–40) Control: 17 days (IQR 7–35)	1.9 years (IQR 1.0–3.0)	Ischaemic stroke (95%), TIA (5%)	Ischaemic stroke (94%), TIA (6%)
Kwon 2005 (TOSS) ³⁰	Within 2 weeks	6 months	ICAS	ICAS
Kwon 2011 (TOSS-2) ³¹	Within 2 weeks Cilostazol: 8.03 (3.34) days Control: 7.82 (3.15) days	7 months	ICAS	ICAS
Lee 2011 (CAIST) ³²	Within 48 hours Cilostazol: 33 (12) hours Control: 35 (11) hours	90 days	Large-artery disease (32%), small-vessel disease (55%), cardioembolism (1%), other determined aetiology (<1%), undetermined (12%)	Large-artery disease (25%), small-vessel disease (62%), cardioembolism (1%), other determined aetiology (<1%), undetermined (12%)
Lee 2017 ³³	Within 7 days	3 months	Atherosclerosis (9.4%), small-artery disease (87.5%), TIA (3.1%)	Atherosclerosis (2.9%), small- artery disease (87.5%), TIA (5.9%), unknown (8.8%)

Continued

Table 2 Continued

Study ID	Time to randomisation/treatment	Duration of treatment/ follow-up	Type of stroke	
			Cilostazol	Control
Nakamura 2012 ³⁴	Within 48 hours Cilostazol: 25 (12) hours Control: 23 (14) hours	6 months	Large-artery atherosclerosis (21%), small-vessel occlusion (47%), other determined or undetermined aetiology (32%)	Large-artery atherosclerosis (18%), small-vessel occlusion (47%), other determined or undetermined aetiology (34%)
Ohnuki 2017 ³⁵	Within 7 days	4 weeks	Atherothrombosis (15%), LACI (62%), TIA (8%), unknown (15%)	Atherothrombosis (45%), LACI (45%), TIA (9%)
Shimizu 2013 ³⁶	Within 24 hours	3 months	Atherothrombotic (30.7%), LACI (64.1%), others (5.2%)	Atherothrombotic (25.0%), LACI (70.7%), others (4.3%)
Shinohara 2010 (CSPS 2) ³⁷	Within 26 weeks (6 months)	1–5 years Mean: 29 (16) months	Atherothrombotic (33%), LACI (65%), others (3%)	Atherothrombotic (31%), LACI (65%), others (3%)
Toyoda 2019 (CSPS. com) ^{†,‡38}	8–180 days Cilostazol: 27 days (IQR 13–63) Control: 25 days (IQR 13–64)	0.5–3.5 years Median: 1.4 (IQR 0.8–2.2)	ICAS (30%), ECAS (12%) Atherothrombotic (42%), LACI (50%), others/unknown (8%)	ICAS (29%), ECAS (14%) Atherothrombotic (42%), LACI (49%), others/unknown (9%)
Uchiyama 2015 (CATHARSIS) ³⁹	2 weeks to 6 months	2 years Mean: 762 days	ICAS	ICAS

Data are in median (IQR), mean (range) or mean (SD) unless otherwise stated.

*Only patients with asymptomatic or previous intracerebral haemorrhage were included.

†Only patients with high-risk ischaemic stroke were included.

‡ICAS, intracranial arterial stenosis; LACI, lacunar infarction; TIA, transient ischaemic attack.

recruited only patients with asymptomatic or previous intracerebral haemorrhage,²⁹ most studies excluded such patients or participants with a bleeding diathesis. One study included patients with high-risk ischaemic stroke,³⁸ while most others did not include patients with moderate-to-severe disability, for example, NIHSS ≥ 8 ,³⁴ NIHSS ≥ 16 ^{25 30 32} or NIHSS ≥ 20 ^{22 36}, mRS ≥ 4 ^{25 27} or severe cerebral deficits.^{23 24} Some studies explicitly excluded patients with concurrent antiplatelet,^{22 24 26–28 30 32 33 37–39} anticoagulant,^{22–24 26 30 32–34 37–39} or thrombolytic/fibrinolytic^{24 31–34 36 37} use.

Four trials^{22 32 34 36} included patients with acute stroke (onset within 48 hours); three trials^{22 23 35} administered short-term antiplatelet therapy (<3 months). Studies compared cilostazol (CIL) single or dual antiplatelet therapy (SAPT or DAPT) against aspirin (ASA), clopidogrel (CLO), and placebo or best medical therapy (No CIL). For one study,²² we analysed data recorded at 14 days instead of 3 months if available—that is, for most outcomes except for major haemorrhagic events and good functional outcome—to minimise bias from the treatment regimen.

Quality assessment

On the basis of RoB 2, most trials were of high quality and possessed a low overall risk of bias (online supplemental data). One trial²⁸ had a high risk of bias due to missing outcome data from a significant, asymmetric loss to follow-up.

Stroke recurrence and functional outcomes

Ischaemic stroke recurrence

Meta-analysis of all 18 randomised trials (figure 1) revealed cilostazol significantly reduced the risk of recurrent ischaemic stroke (RR=0.69, 95% CI 0.58 to 0.81, $p<0.0001$). There was low heterogeneity across all trials ($I^2=0\%$).

Subgroup analysis based on specific antiplatelet regimens indicated moderate heterogeneity between antiplatelet regimens ($I^2=42.9\%$; $\chi_5^2=8.76$, $p=0.12$). The greatest risk reduction was as follows: (1) CIL+CLO versus CLO (n=1116); followed by (2) CIL versus no CIL (n=1574); (3) CIL+ASA versus ASA (n=2544); and (4) CIL versus ASA (n=5681). Interestingly, CIL+ASA performed worse than ASA+CLO, although this effect was not statistically significant (n=457; RR=1.62, 95% CI 0.60 to 4.37, $p=0.34$). When the specific type of antiplatelet was disregarded, cilostazol DAPT had a greater risk reduction than cilostazol SAPT, both compared with an SAPT.

Any stroke recurrence

Meta-analysis of all 18 trials (figure 2) revealed cilostazol significantly reduced the recurrence of any stroke, both ischaemic and haemorrhagic (RR=0.64, 95% CI 0.54 to 0.74, $p<0.00001$). There was low heterogeneity across all trials ($I^2=0\%$).

Subgroup analysis based on specific antiplatelet regimens indicated moderate heterogeneity between

antiplatelet regimens ($I^2=39.4\%$; $\chi_4^2=6.61$, $p=0.16$). The greatest risk reduction was as follows: (1) CIL+ASA/CLO versus ASA/CLO (n=1936); followed by (2) CIL versus no CIL (n=1574); and (3) CIL versus ASA (n=5681). Similar to ischaemic stroke recurrence, CIL+ASA conferred an insignificant increase in risk of any stroke recurrence compared with ASA+CLO (n=457; RR=1.78, 95% CI 0.67 to 4.73, $p=0.25$). However, unlike ischaemic stroke recurrence, CIL+ASA versus ASA did not entail a significant risk reduction for any stroke recurrence ($p=0.30$), which can be explained by the smaller sample size.

Intracranial haemorrhage

Meta-analysis of all 18 trials (figure 3) revealed cilostazol significantly reduced the risk of ICH (RR=0.46, 95% CI 0.31 to 0.68, $p<0.0001$). There was low heterogeneity across all trials ($I^2=0\%$).

Subgroup analysis of specific antiplatelet therapies indicated a substantial reduction in risk of ICH when cilostazol monotherapy was compared with aspirin (RR=0.36, 95% CI 0.22 to 0.59, $p<0.0001$). The addition of cilostazol to aspirin or clopidogrel did not significantly affect the risk of ICH.

Functional outcome

Meta-analysis of four trials (online supplemental data) showed cilostazol did not significantly increase the probability of a good functional outcome (RR=1.07, 95% CI 0.95 to 1.19, $p=0.28$). There was moderate heterogeneity among trials ($I^2=60\%$; $\chi_3^2=7.49$, $p=0.06$). One trial³⁴ contributed significantly to the heterogeneity (n=76; RR=2.00, 95% CI 1.22 to 3.27); this trial measured patients' mRS scores at 6 months compared with 3 months for the other trials. After removing this trial, I^2 decreased to 0% and $\chi_2^2=0.71$ ($p=0.70$).

Adverse events and mortality

Major haemorrhagic events

Meta-analysis of 14 trials (figure 4) revealed cilostazol reduced the risk of major haemorrhagic events, which included ICH (RR=0.49, 95% CI 0.34 to 0.70, $p<0.0001$). There was low heterogeneity across studies ($I^2=0\%$).

Similar to ICH, cilostazol monotherapy substantially reduced the risk of major haemorrhagic events compared with aspirin (RR=0.39, 95% CI 0.25 to 0.62, $p<0.0001$), and the addition of cilostazol to aspirin or clopidogrel did not affect the risk of major haemorrhagic events.

Mortality

Meta-analysis of 15 trials (figure 5) revealed cilostazol did not significantly reduce the risk of mortality (RR=0.90, 95% CI 0.64 to 1.25, $p=0.53$). Two trials^{31 32} presented data on vascular deaths instead of all-cause mortality. There was low heterogeneity across studies ($I^2=0\%$). There was no significant difference between antiplatelet regimens ($I^2=0\%$; $\chi_4^2=0.32$, $p=0.99$); there was no statistically significant reduction in mortality for both cilostazol combination therapy and monotherapy.

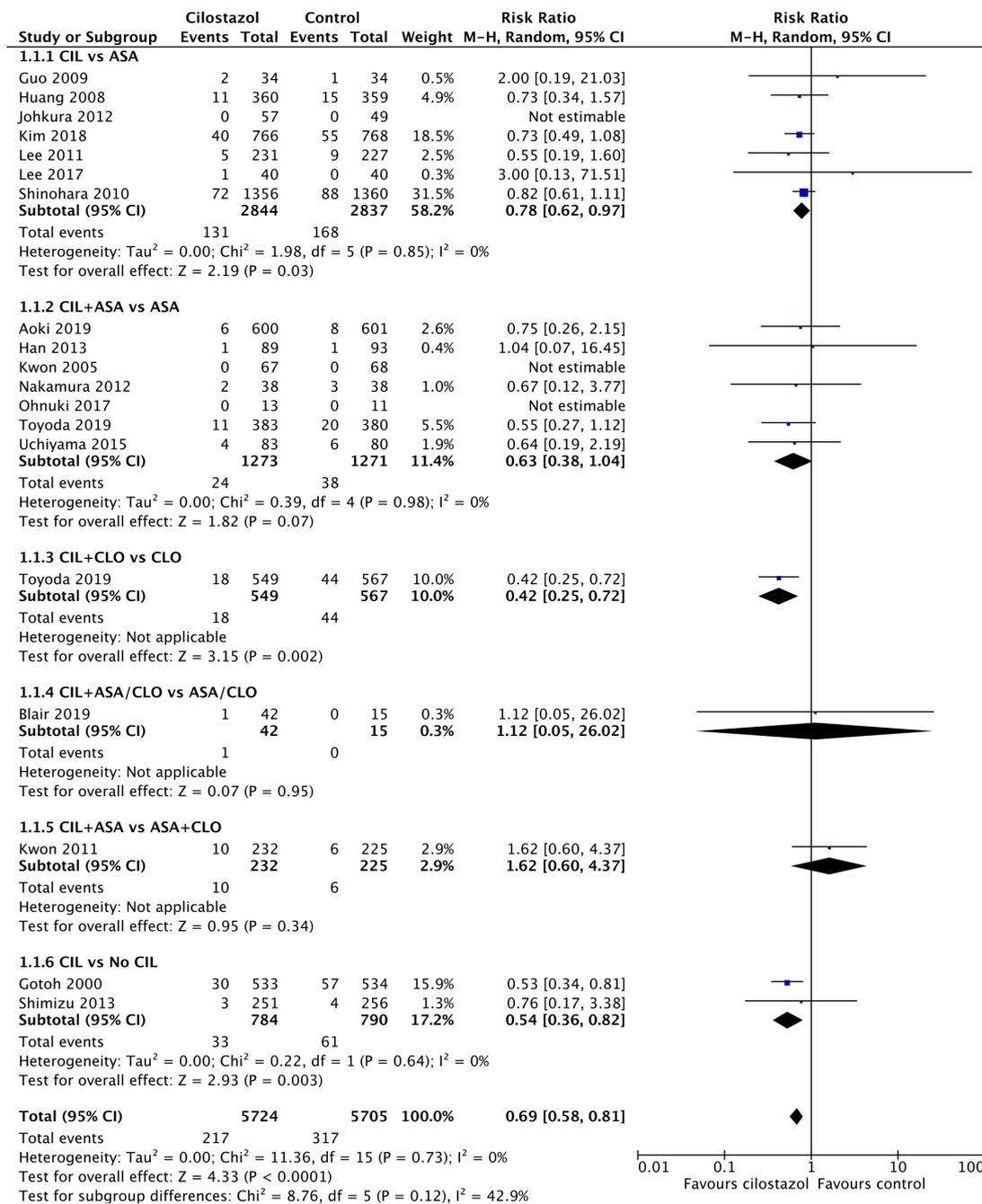


Figure 1 Forest plot depicting risk of ischaemic stroke recurrence. ASA, aspirin; CIL, cilostazol; CLO, clopidogrel; No CIL, placebo or best medical therapy.

Major adverse cardiovascular events

Most studies defined MACE as a composite of strokes, ICH, myocardial infarctions and vascular deaths. Meta-analysis of 13 trials (online supplemental data) revealed cilostazol significantly reduced the risk of MACE (RR=0.67, 95% CI 0.56 to 0.81, $p < 0.0001$). There was low heterogeneity across studies ($I^2 = 0\%$).

Subgroup analysis based on specific antiplatelet regimens indicated moderate heterogeneity between antiplatelet regimens ($I^2 = 54.6\%$; $\chi^2_4 = 8.82$, $p = 0.07$). The greatest risk reduction was (1) CIL+ASA/CLO versus ASA/CLO ($n = 1879$, from one trial)³⁸; followed by (2) CIL versus no CIL ($n = 1574$). However, CIL versus ASA

and CIL+ASA versus ASA did not significantly reduce the risk of MACE ($p = 0.08$ and 0.75 , respectively). Similar to ischaemic stroke recurrence, CIL+ASA performed worse than ASA+CLO, but this effect was not statistically significant ($n = 457$; RR=1.45, 95% CI 0.67 to 3.17, $p = 0.35$). When the specific type of antiplatelet was disregarded, cilostazol DAPT had a greater risk reduction ($n = 3497$; RR=0.58, 95% CI 0.42 to 0.80, $p = 0.0009$) than cilostazol SAPT ($n = 2140$; RR=0.77, 95% CI 0.57 to 1.03, $p = 0.08$), both compared with an SAPT. Since one trial was able to considerably skew the results of CIL DAPT versus ASA SAPT, it suggests that the other studies are underpowered for MACE.

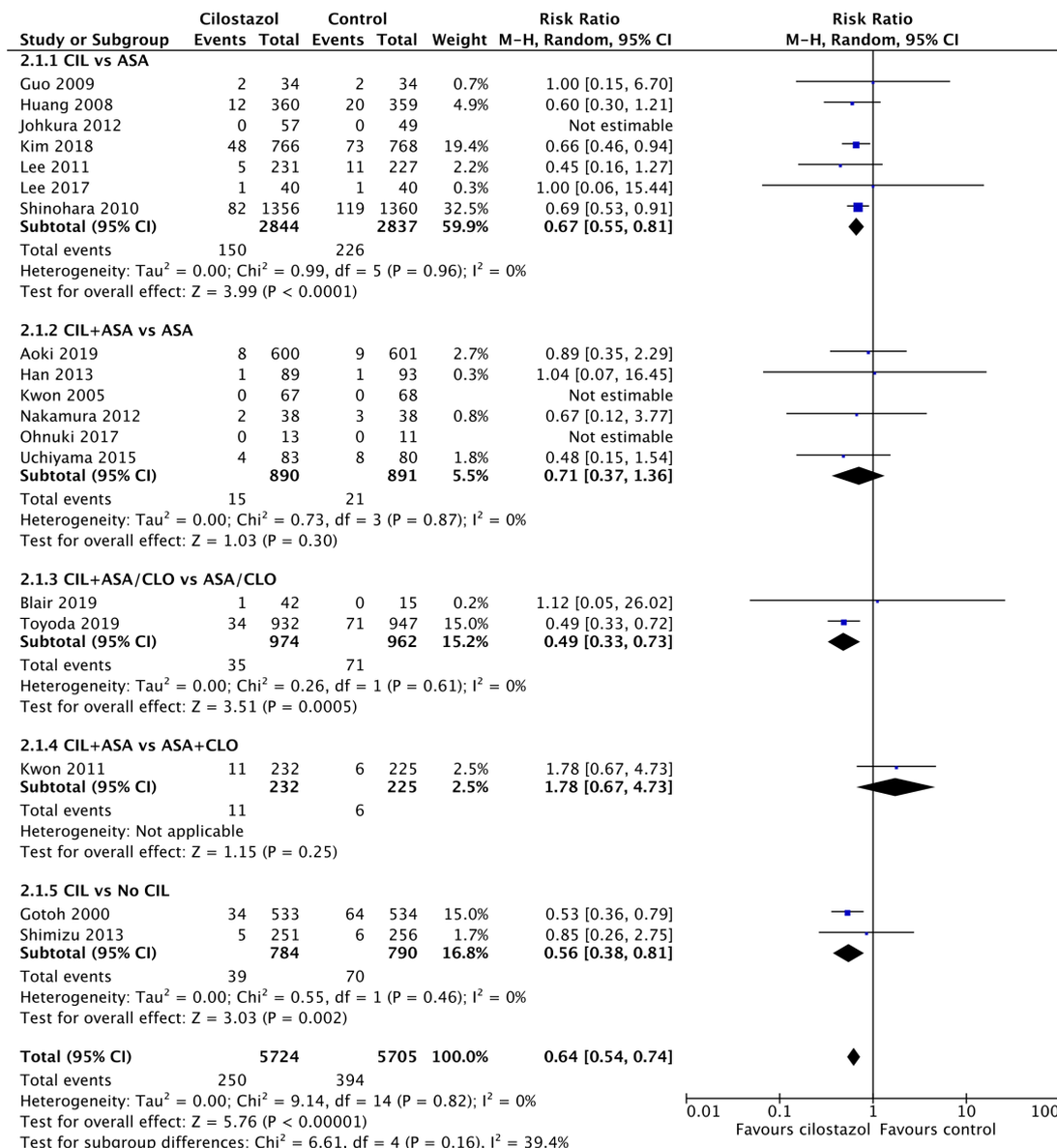


Figure 2 Forest plot depicting risk of any stroke recurrence. ASA, aspirin; CIL, cilostazol; CLO, clopidogrel; No CIL, placebo or best medical therapy.

ADEs leading to treatment discontinuation

Meta-analysis of 13 trials (online supplemental data) revealed cilostazol increased the risk of ADEs leading to treatment discontinuation (RR=1.83, 95% CI 1.30 to 2.59, $p=0.0006$). There was high heterogeneity across studies ($I^2=81\%$). In three studies,^{26 29 32} the safety population comprising all participants who consumed the drug was used in place of the ITT population.

Subgroup analysis of the antiplatelet treatments demonstrated high heterogeneity among different regimens ($I^2=82.9\%$; $\chi^2=17.58$, $p=0.0005$). The greatest statistically significant increase in relative risk occurred in the CIL+ASA/CLO versus ASA/CLO subgroup ($n=1936$; RR=5.59, 95% CI 3.04 to 10.27), followed by the CIL versus No CIL subgroup ($n=1067$; RR=2.13, 95% CI 1.43 to 3.16). There was no significant increase in ADEs leading to drug

discontinuation in the CIL versus ASA and CIL+ASA versus ASA subgroup ($n=5476$ and 1639 , $p=0.11$ and 0.10 , respectively).

Different administrative strategies

Acute versus delayed administration for stroke

Acute/subacute stroke was defined as stroke within 72 hours of symptom onset. Based on the time from onset to randomisation or treatment, 4 studies comprising 2242 patients presented data on cilostazol administration during acute/subacute stroke, while 14 studies comprising 9187 patients provided data for administration during chronic stroke (table 3). Cilostazol administration during the acute and chronic stroke phase demonstrated no significant difference in outcomes ($I^2\leq 4.8\%$; $p>0.05$).

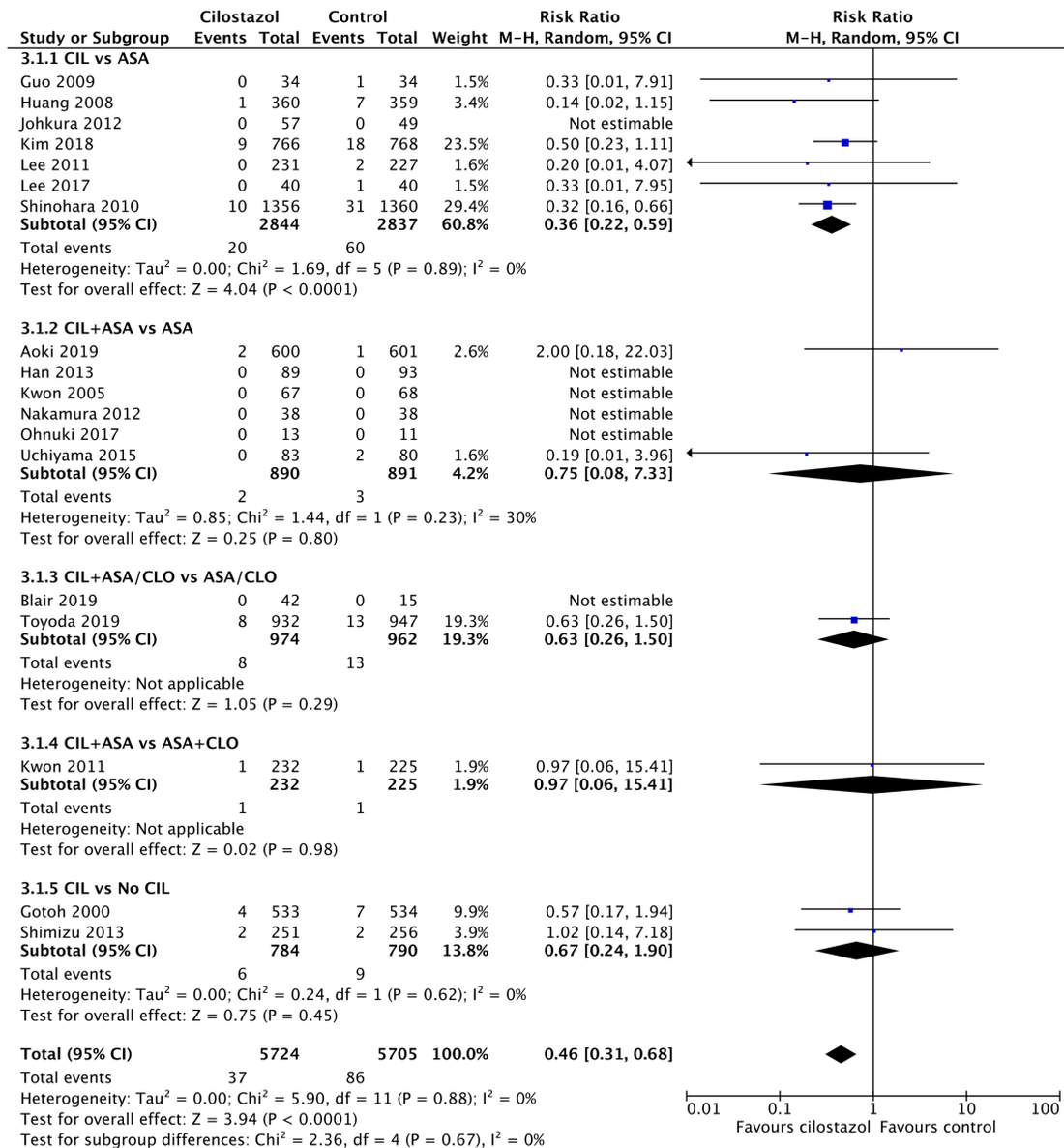


Figure 3 Forest plot depicting risk of intracranial haemorrhage. ASA, aspirin; CIL, cilostazol; CLO, clopidogrel; No CIL, placebo or best medical therapy.

Short-term versus long-term treatment

Long-term antiplatelet treatment was defined as drug administration for ≥ 3 months. Three studies comprising 1282 patients administered short-term antiplatelet therapy, while 15 studies comprising 10 147 patients administered treatment for ≥ 3 months. For most outcomes (ischaemic stroke recurrence, any stroke recurrence, major haemorrhagic events and MACE), there was no significant difference in the efficacy/safety of cilostazol for short-term and long-term administration ($I^2 \leq 3.0\%$). For mortality, there was insufficient events for short-term cilostazol administration to make a comparison. However, for ICH, there was moderate heterogeneity between short-term and long-term cilostazol administration ($I^2 = 32.3\%$; $\chi_1^2 = 1.48$, $p = 0.22$). Long-term cilostazol administration reduces the risk of ICH (RR=0.44, 95% CI 0.30 to 0.66, $p < 0.0001$), while short-term cilostazol administration was underpowered

and seemed to confer no benefit (RR=2.00, 95% CI 0.18 to 22.03, $p = 0.57$).

Symptomatic ICAS

Three studies comprising 755 patients included only patients with symptomatic ICAS. Two studies^{22 25} that did not specify whether ICAS was symptomatic or was associated with the infarct territory were excluded. For most outcomes, there was no significant difference between the symptomatic ICAS and mixed/other stroke subgroup. Most strikingly, in the ICAS subgroup, cilostazol administration is associated with an insignificant increase in risk of MACE (RR=1.38, 95% CI 0.67 to 2.85, $p = 0.38$), compared with a statistically significant decrease in MACE risk in studies which did not exclusively recruit patients with symptomatic ICAS. Furthermore, there was no significant decrease in risk of ICAS progression or worsening

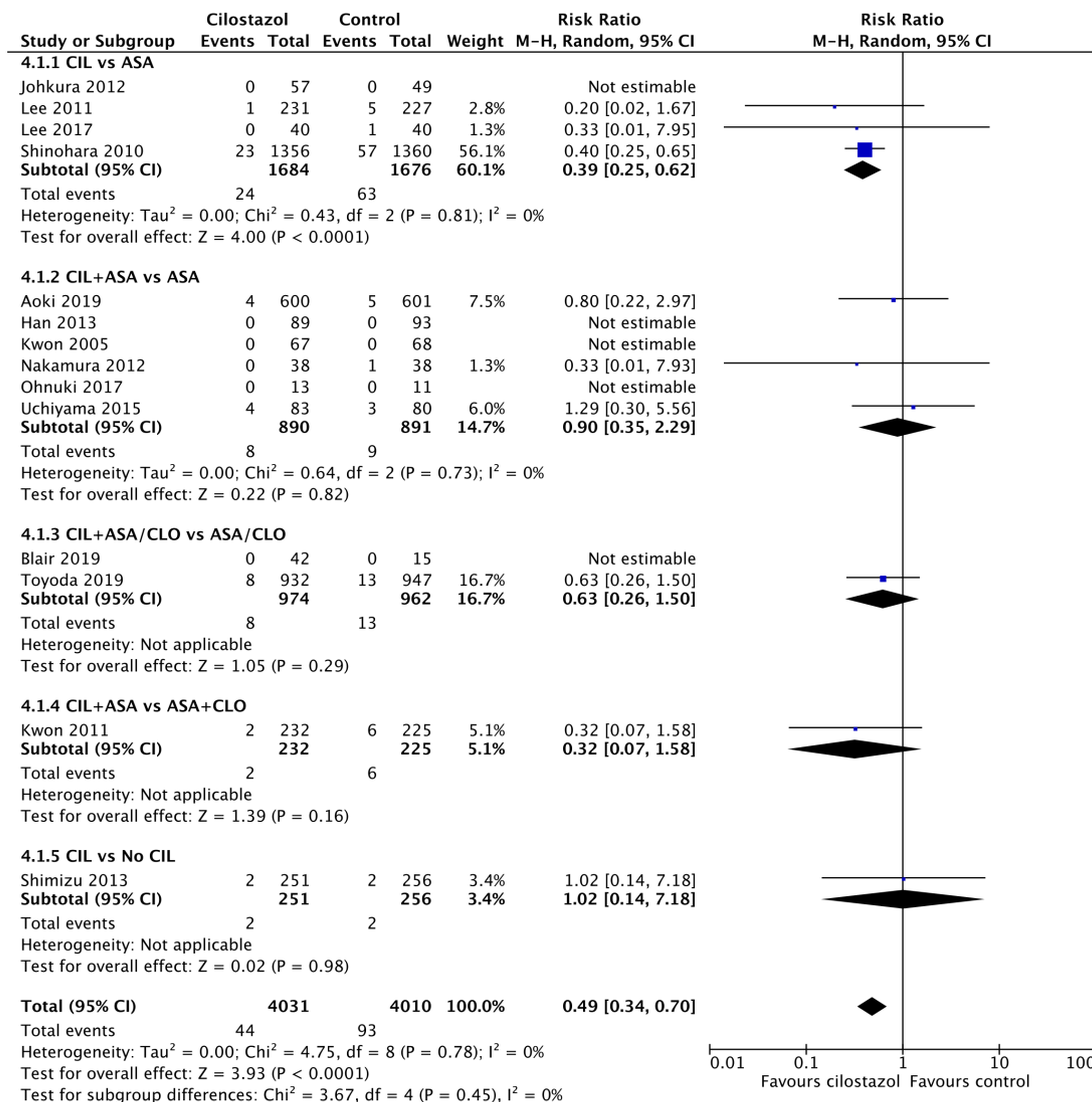


Figure 4 Forest plot depicting risk of major haemorrhagic events. ASA, aspirin; CIL, cilostazol; CLO, clopidogrel; No CIL, placebo or best medical therapy.

(RR=0.63, 95% CI 0.25 to 1.58, p=0.33) after cilostazol administration (online supplemental data).

Further analyses

Meta-regression of the mean or median: (1) duration of treatment or follow-up; (2) time from stroke onset to randomisation or treatment; or (3) proportion of patients with lacunar infarction did not reveal any significant effect on ischaemic stroke recurrence (online supplemental data). Funnel plots did not suggest publication bias for any outcome.

Using the GRADE tool, the quality of evidence was high for ischaemic stroke, any stroke, ICH, major haemorrhagic events and MACE; moderate for mortality and ADEs leading to treatment discontinuations; and low for good functional outcome.

DISCUSSION

Overall, cilostazol demonstrated good efficacy in secondary stroke prevention, significantly reducing the

rates of ischaemic stroke recurrence, any stroke recurrence and MACE.

As a monotherapy, our updated meta-analysis confirmed that cilostazol was superior to aspirin. However, cilostazol monotherapy was not compared with clopidogrel. As combination therapy, the performance of cilostazol is promising. While bleeding outcomes did not increase when cilostazol was added to aspirin or clopidogrel, its efficacy in secondary ischaemic stroke prevention requires confirmation. Compared with aspirin monotherapy, the risk reduction of cilostazol DAPT was greater but just missed significance; we were not able to comment on its efficacy against clopidogrel monotherapy as it was compared with clopidogrel monotherapy in only one trial.³⁸ In contrast to ASA+CLO combination therapy whose use is typically limited to 3 months or less,⁴⁰ our results suggest that cilostazol combination therapy is a promising option for long-term secondary stroke prevention, similar to aspirin plus dipyridamole.⁴¹ Finally, when

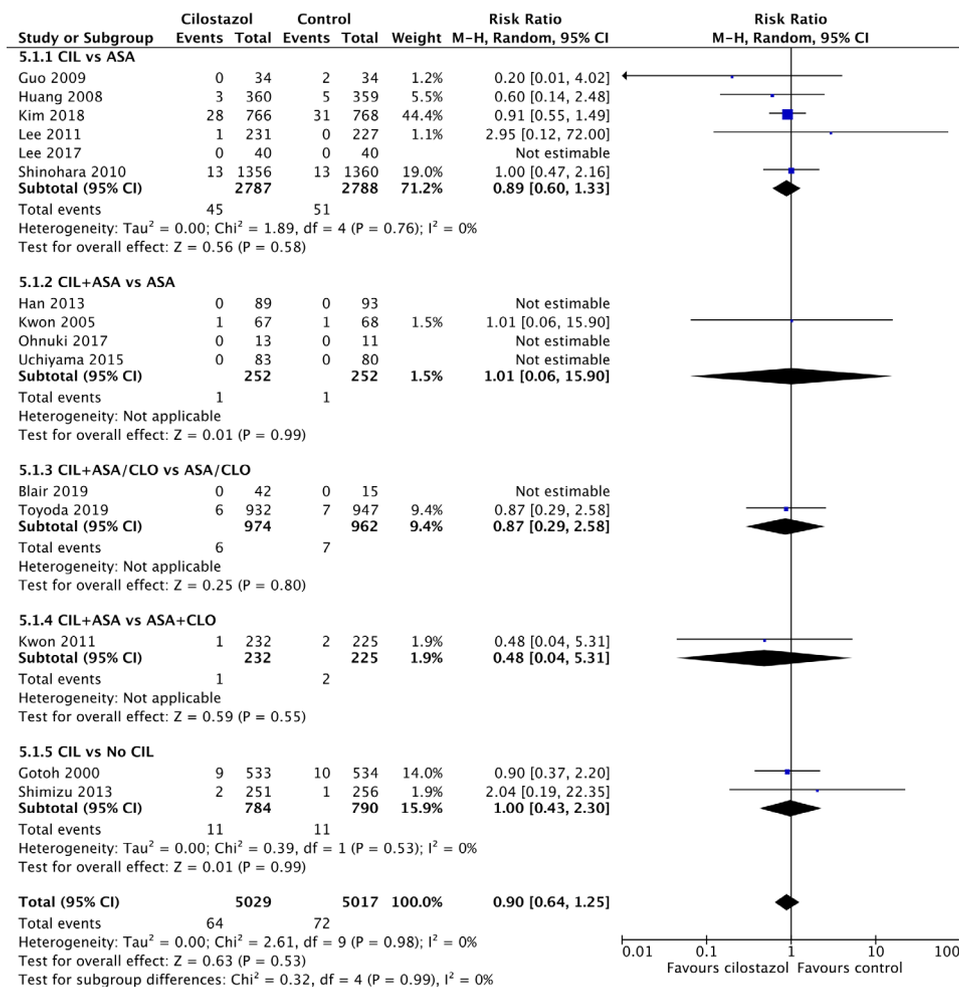


Figure 5 Forest plot depicting risk of mortality. ASA, aspirin; CIL, cilostazol; CLO, clopidogrel; No CIL, placebo or best medical therapy.

compared with other DAPT, CIL combination versus ASA+CLO was limited by wide CI and the availability of only one study³¹ (which demonstrated possible inferiority in stroke recurrence); the CIL+CLO option was not tested.

The ‘No CIL’ subgroup is a heterogeneous classification. One study²⁴ compared cilostazol with placebo. Another study³⁶ compared cilostazol with best medical therapy—both arms were subjected to a variety of both oral and intravenous antiplatelets, antithrombin, anti-coagulants and/or free radical scavengers—without a placebo. Resultantly, it is difficult to comment on this subgroup.

Previous studies have demonstrated high mortality rates after a secondary stroke.⁴² Interestingly, despite a reduction in stroke recurrence and haemorrhagic complications, cilostazol did not significantly reduce mortality rates. One potential explanation is that studies were underpowered to assess the benefits of cilostazol administration on all-cause mortality. Although there were 644 ‘any stroke’ recurrences, there were only 136 cases of mortality; given that the control mortality rate was only 1.44%, the optimal information size was not met for an assumed 25% relative risk reduction.⁴³

Additionally, the rate of good functional outcomes did not increase with cilostazol use. Apart from the fact that only four trials reported mRS scores, treatment was only administered for 3 or 6 months, with the latter achieving better functional outcomes. Future studies could assess the rates of mRS 0–1 with longer duration of antiplatelet administration.

Cilostazol use was associated with an increase in drug discontinuations due to ADEs, especially when administered as combination therapy. Common adverse effects included headaches, dizziness, palpitations, arrhythmias and gastrointestinal disturbances. However, this analysis was complicated by high heterogeneity, which may be attributed to the wide variation in definition among trials. Some trials included vascular events and deaths, while others included medication non-compliance. Nevertheless, existing literature suggests that most of these adverse events are mild and temporary.¹⁰

Unexpectedly, cilostazol did not further reduce the rates of efficacy and safety outcomes in patients with symptomatic ICAS. This does not corroborate with existing literature, which suggests that the antiplatelet and vasodilating effect⁴⁴ of cilostazol could prevent ICAS

Table 3 Summary of outcomes

Subgroup	N	Cilostazol	Control	RR (95% CI)	P value	I ² (%)	Test for subgroup differences
Ischaemic stroke recurrence							
A Overall	18	217/5724	317/5705	0.69 (0.58–0.81)	<0.0001	0	NA
B CIL SAPT vs SAPT	7	131/2844	168/2837	0.78 (0.62–0.97)	0.03	0	$\chi^2=7.50$, $p=0.06$, $I^2=60.0\%$
CIL DAPT vs SAPT	8	43/1864	82/1853	0.52 (0.36–0.75)	0.0005	0	
CIL vs No CIL	2	33/784	61/790	0.54 (0.36–0.82)	0.003	0	
C Stroke onset <3 days	4	16/1120	24/1122	0.67 (0.36–1.25)	0.20	0	
Stroke onset >3 days	14	201/4604	293/4583	0.69 (0.56–0.84)	0.0002	8	
D Short term (<3 months)	3	7/655	8/627	0.78 (0.29–2.12)	0.63	0	$\chi^2=0.07$, $p=0.80$, $I^2=0\%$
Long term (≥3 months)	15	210/5069	309/5078	0.68 (0.58–0.81)	<0.0001	0	
E Symptomatic ICAS	3	14/382	12/373	1.10 (0.45–2.68)	0.84	24	$\chi^2=1.33$, $p=0.52$, $I^2=0\%$
LACI	2	2/131	1/108	1.08 (0.14–8.56)	0.95	0	
Others/mixed/unknown	13	201/5211	304/5224	0.67 (0.56–0.79)	<0.00001	0	
Any stroke recurrence							
A Overall	18	250/5724	394/5705	0.64 (0.54–0.74)	<0.00001	0	NA
B Stroke onset <3 days	4	20/1120	29/1122	0.70 (0.39–1.23)	0.21	0	$\chi^2=0.11$, $p=0.75$, $I^2=0\%$
Stroke onset >3 days	14	230/4604	365/4583	0.63 (0.54–0.74)	<0.00001	0	
C Short term (<3 months)	3	9/655	9/627	0.91 (0.37–2.24)	0.83	0	$\chi^2=0.61$, $p=0.43$, $I^2=0\%$
Long term (≥3 months)	15	241/5069	385/5078	0.63 (0.54–0.74)	<0.00001	0	
D Symptomatic ICAS	3	15/382	14/373	0.96 (0.27–3.45)	0.95	65	$\chi^2=0.71$, $p=0.70$, $I^2=0\%$
LACI	2	2/131	1/108	1.08 (0.14–8.56)	0.95	0	
Others/mixed/unknown	13	233/5211	379/5224	0.62 (0.53–0.73)	<0.00001	0	
Intracranial haemorrhage							
A Overall	18	37/5724	86/5705	0.46 (0.31–0.68)	<0.0001	0	NA
B Stroke onset <3 days	4	4/1120	5/1122	0.91 (0.24–3.53)	0.89	0	$\chi^2=1.05$, $p=0.31$, $I^2=4.8\%$
Stroke onset >3 days	14	33/4604	81/4583	0.43 (0.29–0.65)	<0.0001	0	
C Short term (<3 months)	3	2/655	1/627	2.00 (0.18–22.03)	0.57	NA	$\chi^2=1.48$, $p=0.22$, $I^2=32.3\%$
Long term (≥3 months)	15	35/5069	85/5078	0.44 (0.30–0.66)	<0.0001	0	
D Symptomatic ICAS	3	1/382	3/373	0.46 (0.06–3.57)	0.46	0	$\chi^2=0.00$, $p=1.00$, $I^2=0\%$
LACI	2	0/131	0/108	NA	NA	NA	
Others/mixed/unknown	13	36/5211	83/5224	0.46 (0.31–0.68)	0.0001	0	
Major haemorrhagic events							
A Overall	14	44/4031	93/4010	0.49 (0.34–0.70)	<0.0001	0	NA
B Stroke onset <3 days	4	7/1120	13/1122	0.60 (0.24–1.52)	0.28	0	$\chi^2=0.23$, $p=0.63$, $I^2=0\%$
Stroke onset >3 days	10	37/2911	80/2888	0.47 (0.32–0.69)	0.0001	0	
C Short term (<3 months)	3	4/655	5/627	0.80 (0.22–2.97)	0.74	NA	$\chi^2=0.59$, $p=0.44$, $I^2=0\%$
Long term (≥3 months)	11	40/3376	88/3383	0.47 (0.32–0.68)	<0.0001	0	
D Symptomatic ICAS	3	6/382	9/373	0.67 (0.17–2.59)	0.56	37	$\chi^2=0.24$, $p=0.62$, $I^2=0\%$
LACI	2	0/131	0/108	NA	NA	NA	
Others/mixed/unknown	9	38/3518	84/3529	0.47 (0.32–0.68)	<0.0001	0	
Mortality							
A Overall	15	64/5029	72/5017	0.90 (0.64–1.25)	0.53	0	NA
B Stroke onset <3 days	2	3/482	1/483	2.33 (0.34–15.82)	0.39	0	$\chi^2=0.98$, $p=0.32$, $I^2=0\%$
Stroke onset >3 days	13	61/4547	71/4534	0.87 (0.62–1.22)	0.43	0	

Continued

Table 3 Continued

Subgroup		N	Cilostazol	Control	RR (95% CI)	P value	I ² (%)	Test for subgroup differences
C	Short term (<3 months)	2	0/55	0/26	NA	NA	NA	NA
	Long term (≥3 months)	13	64/4974	72/4991	0.90 (0.64–1.25)	0.53	0	
D	Symptomatic ICAS	3	2/382	3/373	0.67 (0.11–4.06)	0.66	0	$\chi^2_{21}=0.11$, $p=0.74$, $I^2=0\%$
	LACI	2	0/131	0/108	NA	NA	NA	
	Others/mixed/unknown	10	62/4516	69/4536	0.91 (0.65–1.27)	0.58	0	
MACE								
A	Overall	13	184/3826	276/3842	0.67 (0.56–0.81)	<0.0001	0	NA
B	Stroke onset <3 days	4	25/1120	33/1122	0.76 (0.45–1.27)	0.30	0	$\chi^2_{21}=0.18$, $p=0.67$, $I^2=0\%$
	Stroke onset >3 days	9	159/2706	243/2720	0.67 (0.52–0.86)	0.002	20	
C	Short term (<3 months)	2	12/613	12/612	1.00 (0.45–2.21)	1.00	NA	$\chi^2_{21}=1.03$, $p=0.31$, $I^2=3.0\%$
	Long term (≥3 months)	11	172/3213	264/3230	0.66 (0.55–0.79)	<0.0001	0	
D	Symptomatic ICAS	2	17/299	12/293	1.38 (0.67–2.85)	0.38	0	$\chi^2_{22}=4.22$, $p=0.12$, $I^2=52.6\%$
	LACI	1	1/89	1/93	1.04 (0.07–16.45)	0.98	NA	
	Others/mixed/unknown	10	166/3438	263/3456	0.64 (0.53–0.77)	<0.00001	0	

CIL, cilostazol; DAPT, dual antiplatelet therapy; ICAS, intracranial arterial stenosis; LACI, lacunar infarction; No CIL, placebo or best medical therapy; RR, risk ratio; SAPT, single antiplatelet therapy.

progression, which, unlike extracranial artery stenosis, cannot be equilibrated by collateral circulation.⁴⁵

Subgroup analysis did not reveal any significant effect of stroke chronicity, or duration of treatment or follow-up, on efficacy and safety outcomes. However, the risk reduction of ICH is greater when cilostazol is administered long term, which could be due to the increase in incidence of ICH over time when aspirin is employed. Surprisingly, there was a lower overall rate of ischaemic stroke recurrence when cilostazol is administered during the acute/subacute phase (1.78%) versus chronic phase (5.38%) of ischaemic stroke. This may be attributed to a shorter duration of follow-up (14 days to 6 months) in the former subgroup; alternatively, these four trials excluded patients with NIHSS ≥8, 16 or 20, which may reduce the risk of ischaemic stroke recurrence.

Limitations

All but one trial were performed in the East Asian population, thus the results of this meta-analysis may not be fully generalisable to other ethnicities. Additionally, the following analyses were underpowered: (1) mortality; (2) subgroup analysis for symptomatic ICAS; and (3) subgroup analysis for the duration of treatment. Moreover, there was moderate to high heterogeneity in the meta-analysis of (1) good functional outcome; and (2) ADEs leading to treatment discontinuation. Furthermore, due to the limited trials available, we were unable to compare the efficacy and safety of cilostazol with antiplatelet agents other than aspirin and clopidogrel. Finally, for one study,²² the time points at which the different outcomes were measured varied due to the limited information available.

Implications for future research

We identified knowledge gaps and areas for further research. For example, studies could perform trials on non-Asian populations. Furthermore, studies could compare the safety and efficacy profile of cilostazol with other antiplatelet regimens, in particular: (1) CIL versus CLO monotherapy; (2) CIL combination versus ASA+CLO, especially in the short term; and (3) newer antiplatelet agents, for instance, CIL versus ticagrelor (TIG) or CIL versus short-term ASA+TIG. While network meta-analyses may be performed to evaluate indirect evidence, further randomised controlled trials for direct comparisons are necessary.

CONCLUSIONS

Cilostazol SAPT and DAPT significantly reduced the risk of recurrent ischaemic stroke, any stroke and MACE but did not have a significant effect on functional outcomes. Cilostazol reduced ICH and major haemorrhagic complications compared with aspirin.

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