Cilostazol increases patency and reduces adverse outcomes in percutaneous femoropopliteal revascularisation: a meta-analysis of randomised controlled trials

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ABSTRACT

Background: Cilostazol is an oral antiplatelet agent currently indicated for treatment of intermittent claudication. There is evidence that cilostazol may reduce femoropopliteal restenosis after percutaneous endovascular intervention.

Methods: We searched PubMed, Scopus and Cochrane databases from 1966 through September 2013 for randomised controlled trials (RCTs) evaluating the addition of cilostazol to standard care in patients receiving femoropopliteal endovascular treatment. Restenosis, target lesion revascularisation and combined adverse outcomes (death, revascularisation and amputation) within 1–2 years postprocedure were evaluated.

Results: Of 205 articles, three RCTs were included in the analysis. The pooled data provided a total of 396 patients, 195 of whom received cilostazol. When compared to standard medical therapy alone, cilostazol significantly reduced the risk of restenosis (risk difference $-0.20; 95\%\,CI\, -0.29$ to $-0.11;\,p<0.0001;\,number\,needed\,to\,treat\,5$), target lesion revascularisation (risk difference $-0.17; 95\%\,CI\, -0.25$ to $-0.09;\,p<0.0001;\,number\,needed\,to\,treat\,6$). Death and amputation were not different in between groups.

Conclusions and limitation: Cilostazol significantly increases femoropopliteal patency and decreases adverse outcomes in percutaneous endovascular intervention. However, further RCTs are needed because of limited sample size; this meta-analysis represents the best current evidence.

INTRODUCTION

Peripheral arterial disease (PAD) is highly prevalent and has been demonstrated to increase cardiovascular mortality by 15-fold in individuals with severe symptomatic large-vessel disease.1 Nevertheless, it still remains an underdiagnosed and undertreated condition.2 Femoropopliteal lesions account for nearly three quarters of lower extremity PAD.3 4 Revascularisation is required for patients with critical limb ischaemia. The most recent guidelines on the management of femoropopliteal PAD have expanded the indications for endovascular therapy (EVT) to the detriment of surgical revascularisation.5–7

Antiplatelet therapy is recommended after EVT for PAD.5 Cilostazol is a phosphodiesterase-3 inhibitor indicated for the treatment of intermittent claudication.5 8

The potential benefits of cilostazol after EVT for PAD are not addressed in current guidelines. Therefore, we decided to perform a systematic literature review and meta-analysis of the outcomes of randomised controlled trials (RCTs) that compare cilostazol to...
standard therapy alone after EVT for femoropopliteal PAD disease. Cilostazol could potentially improve procedural outcomes at a low cost, as it has been demonstrated to be cost-effective for the treatment of intermittent claudication due to PAD.9 10

MATERIALS AND METHODS
Search strategy
We systematically searched PubMed, Scopus and Cochrane Central Register of Controlled Trials for RCTs from 1966 to September 2013, which compared cilostazol to standard medical therapy in patients with femoropopliteal EVT. The review was performed in accordance with established methods for systematic reviews in cardiovascular medicine.11 The following medical subject heading terms were included for a MEDLINE search and adapted for other databases as needed: “cilostazol” AND (“peripheral arterial disease” OR “endovascular therapy” OR “femoropopliteal”). In addition to searching databases, reference lists of all included studies, meta-analyses and reviews were manually searched, including unpublished data. There was neither language nor patient population size restriction for the search.

Data extraction
Two authors independently completed data extraction after following defined search criteria and quality assessment. They obtained data from tables, text and graphs. When the data were presented in percentage the absolute values were calculated. Disagreements were resolved by consensus after review by the senior author of the study.

Selection criteria
Inclusion criteria were RCTs with (1) a direct comparison between patients treated with cilostazol and a control group who received standard medical therapy alone; (2) patients with femoropopliteal lesions due to PAD; (3) patients who received EVT; and (4) reports of the primary and secondary outcomes. Exclusion criteria were (1) studies that only reported outcomes of cilostazol, without a direct comparison to a control group; (2) observational studies; (3) non-randomised clinical trials; and (4) overlapping patient population, identified by studies developed over the same period of time with common authors or common study centres. In this case, only the study with a greater number of patients was included.

Outcomes
The primary outcome studied was 1–2-year incidence of restenosis after endovascular treatment defined by Doppler. Secondary outcomes of interest were 1–2-year target lesion revascularisation and combined major adverse outcomes, which included death, target lesion revascularisation, surgical revascularisation and amputations. When major combined outcomes were not presented as defined it was calculated adding the individual outcomes.

Statistical analysis
Meta-analysis was performed according to recommendations of the Cochrane Collaboration and the Quality of Reporting of Meta-analyses (QUORUM) guidelines and PRISMA statement using Review Manager (RevMan) V.5.2 version (Copenhagen, Nordic Cochrane Centre,
Table 1  Characteristics of studies included in the meta-analysis

<table>
<thead>
<tr>
<th>Author</th>
<th>Follow-up</th>
<th>Inclusion criteria</th>
<th>Exclusion criteria</th>
<th>Control group</th>
<th>Intervention group</th>
</tr>
</thead>
</table>
| Iida et al  | 24 months | 1. De novo FP lesions >50%  
2. Occlusion without inflow lesions  
3. Outflow lesions of below-the-knee arteries of >1 vessel runoff  
4. Symptomatic PAD with claudication (Fontaine 2, 3 or 4) | 1. Acute onset critical limb ischaemia  
2. Previous bypass surgery or  
3. Previous angioplasty for the FP lesions  
4. Presence of untreated pelvic lesions  
5. Intolerance to the medication or contrast agents | 1. Aspirin (100 mg/day)+ticlopidine (200 mg/day)  
2. Patients who received stents were also treated with a thienopyridine | 1. Aspirin (100 mg/day)  
2. Patients who received stents were also treated with a thienopyridine  
3. Aspirin (81–100 mg/day), ticlopidine (200 mg/day) and cilostazol (200 mg/day) |
| Soga et al  | 12 months | 1. Patients with symptomatic PAD greater than Rutherford 1 screened by non-invasive tests to detect limb ischaemia and the presence of de novo FP lesions | 1. Treated with coronary DES  
2. Heart failure symptoms with systolic or diastolic dysfunction evaluated by cardiacechocardiography  
3. Inflow aortoiliac lesions  
4. FP lesions with severe calcification  
5. Poor below-the-knee runoff defined as number of below-the-knee runoff <1 | 1. Aspirin (100 mg/day)  
2. Patients who received stents were also treated with a thienopyridine | 1. Aspirin (81–100 mg/day) and ticlopidine (200 mg/day)  
2. Patients who received stents were also treated with a thienopyridine  
3. Aspirin (81–100 mg/day), ticlopidine (200 mg/day) and cilostazol (200 mg/day) |

Outcomes definition

1. Lesion patency: peak systolic velocity ratio >2.4 by DUS
2. Target lesion revascularisation: reintervention performed for >50% diameter stenosis identified by angiography within 5 mm of the target lesion after documentation of recurrent symptoms of PAD
3. Angiographic restenosis: recurrence of ≥50% diameter stenosis; a peak systolic velocity ratio of >2.0 on Duplex ultrasonography
4. Target lesion: treated segment from 10 mm proximal to 10 mm distal
5. TLR: any repeat EVT for restenosis or other complication of the target lesion with a %DS of >50% in angiography
6. Restenosis: peak systolic velocity ratio of ≥2.4 on Duplex ultrasonography

Endovascular procedure

After balloon inflation for at least 1 min, self-expanding stent was done if:
1. Pressure gradient >10 mm Hg OR
2. >30% residual stenosis OR
3. Flow-limiting dissection
A nitinol stent (Lumixxx, CR Bard, Murray Hill, NJ) or cobalt metallic stent (Wallstenst, Boston Scientific, Natick, Mass) with the diameter 1 mm larger than the reference diameter was used
Stents of 6 mm in diameter were used in most cases

After balloon inflation for at least 1 min, stent was done if:
1. Flow-limiting dissection OR
2. Pressure gradient >10 mm Hg OR
3. >30% residual stenosis
Patients received SMART stents (Cordis Corp, Miami Lakes, Florida, USA) with a diameter 1 mm larger than the reference vessel diameter

After balloon inflation for at least 1 min, self-expanding stent was done if:
1. Angiographic residual stenosis of >30% OR
2. Flow-limiting dissection
A commercially available self-expandable stent was used
Stent type was determined by the operators, and the stent size was chosen to be 1–2 mm larger than the vessel diameter determined

ABI, Ankle Brachial Index; DES, drug-eluting stent; DS, diameter of stenosis; EVT, endovascular therapy; DUS, distal ultrasound; FP, femoropopliteal lesion; MI, myocardial infarction; PAD, peripheral arterial disease; SMART, stent: Cordis Corp, Miami Lakes; TLR, target lesion revascularisation.
The Cochrane Collaboration, 2012). Pooled treatment effects were estimated using risk difference (RD) with the Mantel-Haenszel method. We calculated the number needed to treat (NNT) according to the recommendations of the Cochrane Collaboration. Heterogeneity was assessed using $\chi^2$ tests and $I^2$ statistic; we defined $I^2 < 25\%$ as low heterogeneity according to the Cochrane Handbook of Systematic Reviews. We performed fixed effect analysis when $I^2$ was less than 25% or $p$ value at least 0.10; otherwise we used random effect. We assessed quality for each included trial according to the methods of the Cochrane Collaboration. All included studies were controlled trials and were considered high quality.

RESULTS

The process of study selection is shown in figure 1. Initial MEDLINE search using a systematic approach yielded 205 studies. The search in EMBASE and Cochrane registries did not yield additional studies. Through a review of titles and abstracts, 173 studies were rejected due to lack of relevance to our meta-analysis. The remaining 32 articles were reviewed and assessed for satisfaction of the inclusion or exclusion criteria. Three studies met all criteria and were included in this analysis.

The first study had a follow-up period of 24 months and compared cilostazol/aspirin with ticlopidine/aspirin in 200 patients with femoropopliteal lesions. At 12 and 24 months, the cilostazol/aspirin group reduced restenosis rates (18% vs 43% and 27% vs 52%, respectively). A smaller study in 2009 also found similar results with restenosis rates (43% vs 70.3%). These findings were validated in a larger multicentre study in 2013 with 200 patients that demonstrated an angiographic restenosis rate at 12 months of 20% in cilostazol group versus 49% in the non-cilostazol group.

In order to obtain pooled estimates, a total of 396 patients were included in this analysis. Following EVT for femoropopliteal PAD lesions, 195 individuals were treated with cilostazol and standard medical therapy, whereas 201 received standard medical therapy alone, which included aspirin and a thienopyridine. The main characteristics of individual studies can be found in table 1. Of note, stent restenosis by Doppler was defined as a peak systolic velocity ratio >2.4 in Iida et al and >2 in Iida et al. 16

Table 2 illustrates baseline characteristics of populations in the individual studies. All baseline variables including stenting ratio were similar in all studies with the exception of preprocedural Ankle Brachial Index (ABI) in Iida et al; despite both groups having the ABI on the moderate disease range it was higher on the cilostazol group (0.71 vs 0.66).

For the primary outcome, as seen in figure 2, the follow-up revealed a significantly reduced incidence of restenosis in patients who received cilostazol in addition...
to standard medical (RD $-0.20$; 95% CI $-0.29$ to $-0.11$; $p<0.0001$; NNT 5). This was reflected on a reduced need for target lesion revascularisation (RD $-0.17$; 95% CI $-0.25$ to $-0.09$; $p<0.0001$; NNT 6) with a NNT of 5 for both outcomes (figure 3). We also conducted a sensitivity analysis utilising OR as the primary pooled estimate and the effect size did not change. Death and amputations did not statistically differ between the two groups.

DISCUSSION

The potential benefit of cilostazol following EVT for femoropopliteal PAD was investigated in this study. Our meta-analysis of RCTs revealed that 75 of 201 patients not treated with cilostazol have developed restenosis within 12–24 months and that the chance of restenosis may be mitigated in more than 50% with cilostazol. This result is particularly significant when taken into account that cilostazol is a generic and inexpensive drug that has been shown to be a cost-effective drug in other clinical scenarios, such as treatment of intermittent claudication and secondary prevention of cerebral infarction and that the benefit is obtained with a small NNT.

One-year incidence of restenosis after balloon angioplasty of femoropopliteal lesions has been reported as high as 64%. Sirolimus-eluting stents and endovascular brachytherapy have failed to demonstrate a long-term decrease in the restenosis incidence following lower extremity EVT for PAD, but paclitaxel-eluting stents have shown increased patency and a lower rate of events. Nevertheless, patency has greatly improved since the introduction of nitinol stents. When compared to percutaneous transluminal angioplasty, the use of nitinol stenting decreased the 12-month restenosis incidence from 81.3% to 36.7% in a RCT. Nevertheless, the results of this meta-analysis suggest that patency rates can be further increased with cilostazol in addition to nitinol stents, given that stent use did not differ among cilostazol and control groups, as illustrated in table 2.

Cilostazol has been shown to decrease restenosis and repeat revascularisation after percutaneous coronary intervention in patients with coronary artery disease, without an increased bleeding risk. Similarly, in this meta-analysis, target lesion revascularisation occurred in about every third patient on the control group over a 12–24 month follow-up. The chance of requiring a repeat target lesion revascularisation was reduced in more than 60% with the use of cilostazol. The combined incidence of death, revascularisation and amputation was also significantly reduced in the cilostazol group but it was mostly driven by the reduced need for revascularisation.

The main mechanism for cilostazol-mediated decrease in restenosis and target lesion revascularisation after EVT is likely inhibition of intimal hyperplasia. Cilostazol has been shown to suppress neointimal hyperplasia in animal models. Furthermore, RCTs have demonstrated that triple antiplatelet therapy with aspirin, clopidogrel and cilostazol was more effective than dual therapy alone in suppressing intimal hyperplasia. A second mechanism for improved outcomes in the cilostazol group is vasodilation. Studies have shown that cilostazol increases walking distance in patients with PAD and improves ankle-brachial index due to vasodilation. Moreover, cilostazol mediates an in-vitro inhibition of smooth muscle cell proliferation. Whether this effect contributes to improved EVT procedural outcomes is unknown. Cilostazol also inhibits platelet aggregation.
This meta-analysis has some limitations. Owing to the limited number of studies the sample size is small. However, despite the limited sample size, our meta-analysis represents the best available evidence for the use of cilostazol to increase femoropopliteal patency after endovascular intervention. As with any meta-analysis, our study is subject to publication bias that is potentially mitigated by the exclusive use of RCTs. Although the time frame of outcomes varied from 12 to 24 months, most cases of restenosis following EVT for femoropopliteal lesions occurred from 6 to 12 months, a time frame that was included in this study. The available data are exclusively from Asian populations and it is possible that our results apply exclusively to Asian populations, but that is unlikely due to prior positive cilostazol studies in other populations. Finally, regarding the antiplatelet properties of cilostazol, bleeding complications could not be assessed given that this outcome was not reported in all included studies and this could be considered to be an important limitation of our analysis. Only Soga’s study attempted comparing major bleeding in dual versus triple therapy but no patients had events. Although not reported in these studies, there is robust evidence that when cilostazol is used in combination with dual antiplatelet therapy for other indications, there is no increased risk of bleeding compared to dual antiplatelet therapy alone. For example, in an RCT with patients with acute coronary syndrome undergoing stenting, triple therapy had similar bleeding events compared with dual antiplatelet therapy.

In summary, in our meta-analysis of RCTs, cilostazol following endovascular treatment for femoropopliteal PAD was shown to significantly reduce restenosis, target lesion revascularisation and combined adverse outcomes when compared to a standard therapy control group. Large RCTs are urged to confirm our findings.

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Contributors AMB, DCG, JSJ and TC were involved in the conception and study design. AMB, DCG, RMNC and TPM were involved in the literature review. GEE-H, GNN, EFA and JJD were involved in the data collection. GNN, GEE-H, EFA and TPM were involved in the statistical analysis. AMB, DCG, GNN and TC were involved in the revising manuscript critically for important intellectual content. AMB, DCG, RMNC, TPM, GEE-H, GNN, EFA, JJD and TC were involved in the final approval of the manuscript submitted.

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