Risk of Death Following Application of Paclitaxel-Coated Balloons and Stents in the Femoropopliteal Artery of the Leg: A Systematic Review and Meta-Analysis of Randomized Controlled Trials

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Background—Several randomized controlled trials (RCTs) have already shown that paclitaxel-coated balloons and stents significantly reduce the rates of vessel restenosis and target lesion revascularization after lower extremity interventions.

Methods and Results—A systematic review and meta-analysis of RCTs investigating paclitaxel-coated devices in the femoral and/or popliteal arteries was performed. The primary safety measure was all-cause patient death. Risk ratios and risk differences were pooled with a random effects model. In all, 28 RCTs with 4663 patients (89% intermittent claudication) were analyzed. All-cause patient death at 1 year (28 RCTs with 4432 cases) was similar between paclitaxel-coated devices and control arms (2.3% versus 2.3% crude risk of death; risk ratio, 1.08; 95% CI, 0.72–1.61). All-cause death at 2 years (12 RCTs with 2316 cases) was significantly increased in the case of paclitaxel versus control (7.2% versus 3.8% crude risk of death; risk ratio, 1.68; 95% CI, 1.15–2.47; number-needed-to-harm, 29 patients [95% CI, 19–59]). All-cause death up to 5 years (3 RCTs with 863 cases) increased further in the case of paclitaxel (14.7% versus 8.1% crude risk of death; risk ratio, 1.93; 95% CI, 1.27–2.93; number-needed-to-harm, 14 patients [95% CI, 9–32]). Meta-regression showed a significant relationship between exposure to paclitaxel (dose-time product) and absolute risk of death (0.4%/C60.1% excess risk of death per paclitaxel mg-year; P<0.001). Trial sequential analysis excluded false-positive findings with 99% certainty (2-sided α, 1.0%).

Conclusions—There is increased risk of death following application of paclitaxel-coated balloons and stents in the femoropopliteal artery of the lower limbs. Further investigations are urgently warranted.

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Key Words: balloon angioplasty • paclitaxel • paclitaxel-coated balloon • paclitaxel-eluting stent

To date, percutaneous transluminal angioplasty and stenting have developed to the mainstream treatment of symptomatic peripheral arterial disease with constantly increasing numbers of procedures worldwide. The femoropopliteal artery is the most common site of involvement in atherosclerosis of the lower limb and is typically characterized by multilevel steno-occlusive disease, often complex calcified morphology, and aggressive postangioplasty neointimal hyperplasia associated with high rates of early vessel restenosis and failure. Drug-eluting stents (DESs) and drug-coated balloons (DCBs) have been extensively investigated as a potential solution to inhibit vessel restenosis and improve clinical outcomes after endovascular revascularization of the femoropopliteal artery.

Following testing in numerous randomized controlled trials (RCTs) and various commercial coating formulations,
paclitaxel has emerged as the single potent and proven antirestenotic agent for the infrainguinal vessels.3–5 Recent meta-analyses of several RCTs with low risk of bias have amassed strong evidence about the clinical effectiveness of paclitaxel DES and DCB in significantly reducing restenosis and thereby reducing the risk of recurrent limb ischemia and target lesion/limb revascularization.4,6,7 Consequently, after extensive preclinical testing and having demonstrated strong clinical efficacy combined with a good safety profile, a number of paclitaxel DES and DCB devices have gradually received the CE mark and Food and Drug Administration approval for use in the femoropopliteal segment of the leg.

However, the INPACT-DEEP (Study of IN.PACT AmphirionTM Drug Eluting Balloon vs. Standard PTA for the Treatment of Below the Knee Critical Limb Ischemia) randomized study has shown higher rates of major amputations in the active paclitaxel arm compared with control.8 In addition, a couple of RCTs with longer-term follow-up have shown hints of increased late patient mortality with the use of paclitaxel DESs9 or DCBs.10,11 In the absence of obvious causal links, these findings have been dismissed by expert review panels as statistical artifacts or anomalies, and both devices are currently under clinical use with extended on label indications. Moreover, coronary drug-eluting stents have been long incriminated for late stent thrombosis with an associated risk of death.12 Hence, we conducted an updated systematic review and quantitative meta-analysis of RCTs investigating paclitaxel-coated balloons and stents in the femoropopliteal artery, in order to analyze the early and late risk of death associated with these novel endovascular technologies that deliver paclitaxel to the vessel wall of the lower limbs.

Material and Methods

Literature Search

The authors declare that all supporting data are available within the tables, figures, and supplemental material of the present article. This systematic review has been registered in the PROSPERO public database (CRD42018099447; http://www.crd.york.ac.uk/PROSPERO). We performed electronic searches of PubMed (Medline), EMBASE (Excerpta Medical Database), AMED (Allied and Complementary medicine Database), Scopus, CENTRAL (Cochrane Central Register of Controlled Trials), archived online content, public filings of regulatory bodies (Food and Drug Administration and European Medicines Agency) and published abstracts from international vascular meetings for eligible RCTs. There were no restrictions on publication language, publication date, or publication status. The literature was screened for randomized studies investigating paclitaxel-coated DESs or DCBs in the femoropopliteal artery of the lower limbs. Search terms included Cochrane, femoral artery, popliteal artery, femoropopliteal artery, late lumen loss, restenosis, target lesion revascularization, peripheral angioplasty, stent, randomized, balloon angioplasty, paclitaxel-eluting balloons, paclitaxel-coated balloons, paclitaxel-eluting stents, paclitaxel-coated stents, paclitaxel-eluting stents, drug-coated balloons, and drug-eluting stents, as well as the corresponding Medical Subjects Headings with Boolean syntax (ie, the logic terms AND and/or OR). The literature search was last updated in August 2018. Trials were considered for inclusion in the present meta–analysis if they fulfilled the following inclusion criteria: (1) Randomized controlled study design, (2) investigation of a paclitaxel-coated/paclitaxel-eluting stent or balloon in the femoropopliteal artery, (3) patient population with peripheral arterial disease of the femoral and/or popliteal artery and symptoms of intermittent claudication and/or critical limb ischemia, (4) clinical follow-up of at least 1 year available.

Evaluation of the quality and risk of bias of the selected RCTs was performed independently by two of the authors (K.K., D.K.) using the Cochrane Collaboration’s tool for assessing risk of bias,13 which evaluates 7 key design items of an RCT: (1) random sequence generation, (2) allocation concealment, (3) blinding of participants and personnel, (4) blinding of outcome assessment, (5) incomplete outcome data, (6) selective outcome reporting, and (7) other potential sources of bias. Each aforementioned domain was evaluated as high, low, or unclear risk of bias according to Cochrane. Disagreements were resolved by consensus.

Data Extraction and Outcome Measures

The trial selection process complied with the Preferred Reporting Items for Systematic Reviews and Meta-Analyses statement.14 The reference lists of all selected articles were
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Results

Included RCTs

The literature search yielded 386 articles eligible for potential inclusion based on their title and content of abstract. Of those, 48 items were found to be relevant and were included in further full-text analysis. Another 20 articles were excluded because they did not meet the predefined inclusion criteria. In all, 28 RCTs with 4663 patients were finally included in the present meta-analysis (Figure S1). Full citation information for all studies, as well as the properties of the 12 different devices tested that were coated with 2.0-, 3.0- or 3.5-µg/mm² paclitaxel, are outlined in Tables S1 and S2. There were 4 RCTs with a paclitaxel DES9,18–22 and 24 RCTs testing different paclitaxel DCB devices. Out of the 24 DCB studies, 16 involved sole application of a paclitaxel-coated balloon (versus balloon PTA)23–48; 4 combined the paclitaxel balloon with bare metal stent (versus PTA and BMS)49–52; and 3 studies investigated use of a DCB for the treatment of in-stent restenosis (versus PTA).53–55 Baseline patient demographics and morphologic lesion variables were largely homogeneously distributed across all studies and in line with previous meta-analyses. The design characteristics of the 28 selected RCTs are shown in Table 1.

Briefly, paclitaxel-coated balloons and stents were used primarily for the treatment of short-distance intermittent claudication (n=4133 of 4663 subjects; 89%) in the majority of the study population and infrequently for a critical limb ischemia indication (n=530). A detailed overview of baseline patient and lesion characteristics is provided in Table S3 for all included studies. Overall, approximately two thirds of the patients were men; the mean age ranged from 67 to 76 years; and there was a high incidence of smoking, hypertension, and hyperlipidemia across all studies. The crude incidence of diabetes mellitus ranged from 21% to 77%. A wide range of intermediate to higher-length lesions was enlisted. With few exceptions, most protocols recommended a short period of 1 to 3 months of dual antiplatelet therapy. The median RCT follow-up period was 2 years (range, 1–5 years). Fifteen studies had 1 year, 10 studies had 2 years, 1 study had 4 years, and 2 studies had 5 years of clinical follow-up available (Table 1). A majority of the RCTs were executed as randomized multicenter studies except for 3 single-center studies and 3 dual-center studies. Randomization and allocation concealment were performed adequately and methodological quality was high for all trials with the exception of an inherently high risk of performance bias in all 28 RCTs because of the universal absence of systematic blindedness of the operators during application of the devices (Figure S2).
| Study and Sources | Year and Study Design | Allocation in Study Arms | Paclitaxel-Coated Device | Primary Study End Point | Maximum Follow-Up Period | Study Registration | Dual Antiplatelet Therapy |
|-------------------|-----------------------|--------------------------|--------------------------|--------------------------|--------------------------|--------------------|--------------------------|
| ZILVER PTX<sup>19,23</sup> | 2011 Multi-center Open label (1:1) | DES (n=241) vs PTA (n=238) | ZILVER-PTX Stent by Cook Medical | Primary patency at 1 y | 5 y | NCT00120406 | >2 mo |
| THUNDER<sup>28,57</sup> | 2008 Multi-center Single-blind (1:1:1) | DCB (n=48) vs PTA (n=54) | Cotavance Balloon by Bavaria Medizin | Late lumen loss at 6 mo | 5 y | NCT00156624 | 1 mo |
| IMPACT SFA<sup>11,25,56,82</sup> | 2015 Multi-center Single-blind (2:1) | DCB (n=220) vs PTA (n=111) | INPACT Admiral by Medtronic | Primary patency at 1 y | 3 y | NCT0175850 NCT01566461 | 1 mo (3 mo if bail-out stenting) |
| FEMPAC<sup>29</sup> | 2008 Multi-center Single-blind (1:1) | DCB (n=45) vs PTA (n=42) | Paccocath Balloon by Bavaria Medizin | Late lumen loss at 6 mo | 2 y | NCT00472472 | Long-term (not specified) |
| LEVANT<sup>17</sup> | 2012 Multi-center Single-blind (1:1) | DCB (n=49) vs PTA (n=52) | Lutonix by CR Bard | Late lumen loss at 6 mo | 2 y | NCT00930813 | 1 mo (3 mo if bail-out stenting) |
| LEVANT<sup>24,26,31,33</sup> | 2015 Multi-center Single-blind (2:1) | DCB (n=316) vs PTA (n=160) | Lutonix by CR Bard | Primary patency at 1 y | 2 y | NCT01412541 | 1 mo |
| ILLUMINATE EU<sup>32,35</sup> | 2017 Multi-center Single-blind (3:1) | DCB (n=222) vs PTA (n=72) | Stellarex by Spectranetics | Primary patency at 1 y | 2 y | NCT01858363 | 1 mo (3 mo if bail-out stenting) |
| CONSEQUENT<sup>30,36</sup> | 2017 Multi-center Single-blind (1:1) | DCB (n=78) vs PTA (n=75) | SeQuent Please by B. Braun | Late lumen loss at 6 mo | 2 y | NCT01970579 | 2 mo |
| ISAR-STATH<sup>51</sup> | 2017 Two-center Open label (1:1:1) | DCB+BMS (n=48) vs PTA+BMS (n=52) | INPACT Admiral by Medtronic | Diameter Stenosis at 6 mo | 2 y | NCT00886752 | 6 mo |
| ISAR-PEBIS<sup>35</sup> | 2017 Two-center Open label (1:1) for ISR | DCB (n=36) vs PTA (n=34) | INPACT Admiral by Medtronic | Diameter stenosis at 6 to 8 mo | 2 y | NCT01083394 | >6 mo |
| INPACT SFA JAPAN<sup>14,41</sup> | 2018 Multi-center Single-blind (2:1) for ISR | DCB (n=68) vs PTA (n=32) | INPACT Admiral by Medtronic | Primary patency at 1 y | 2 y | NCT01947478 | 1 mo (3 mo if bail-out stenting) |
| ACOART<sup>40,42</sup> | 2016 Multi-center Single-blind (1:1) | DCB (n=100) vs PTA (n=100) | Orchid by Acotec Scientific | Late lumen loss at 6 mo | 2 y | Not registered | 6 mo |
| PACIFIER<sup>45</sup> | 2012 Multi-center Single-blind (1:1) | DCB (n=41) vs PTA (n=44) | INPACT Pacific by Medtronic | Late lumen loss at 6 mo | 1 y | NCT01083030 | >2 mo |

Continued
| Study and Sources | Year and Study Design | Allocation in Study Arms | Paclitaxel-Coated Device | Primary Study End Point | Maximum Follow-Up Period | Study Registration | Dual Antiplatelet Therapy |
|-------------------|-----------------------|--------------------------|--------------------------|------------------------|--------------------------|-------------------|--------------------------|
| FAIR54            | 2015 Multicenter Single-blind (1:1) for ISR | DCB (n=62) vs PTA (n=57) | IN.PACT Admiral by Medtronic | 6-mo binary restenosis | 1 y | NCT01305070 | >6 mo |
| BIOLUX P-I44      | 2015 Multicenter Single-blind (1:1) | DCB (n=30) vs PTA (n=30) | Passeo-18 Lux by Biotronik | Late lumen loss at 6 mo | 1 y | NCT01056120 | 1 mo (3 mo if bailout stenting) |
| RANGER-SFA37      | 2018 Multicenter Single-blind (2:1) | DCB (n=71) vs PTA (n=34) | Ranger by Boston Scientific | Primary patency at 1 y | 1 y | NCT02013193 | >1 mo |
| ILLUMENATE pivotal93 | 2017 Multicenter Single-blind (2:1) | DCB (n=200) vs PTA (n=100) | Stellarex by Spectranetics | Primary patency at 1 y | 1 y | NCT01858428 & NCT01912937 | 1 mo |
| DEBATE-SFA50      | 2013 Single-center Open (1:1) | DCB+BMS (n=53) vs PTA+BMS (n=51) | IN.PACT Admiral by Medtronic | 1-y binary restenosis | 1 y | NCT01556542 | 3 mo |
| LUTONIX JAPAN48   | 2018 Multicenter Japan (1:1) | DCB (n=71) vs PTA (n=38) | Lutonix by CR BAND | Primary patency at 1 y | 1 y | Not registered | 1 mo |
| RAPID49           | 2017 Multicenter Double-blind (1:1) | DCB+BMS (n=80) vs PTA+BMS (n=80) | LegFlow by Cardionovum | Primary patency at 1 y | 1 y | ISRCTN47846578 | 3 mo |
| EFFPAC47          | 2018 Multicenter Single-blind (1:1) | DCB (n=85) vs PTA (n=86) | Luminor by iVascular | Late lumen loss at 6 mo | 1 y | NCT02540018 | >1 mo |
| PACUBA53          | 2016 Dual-center Single-blind (1:1) for ISR | DCB (n=85) vs PTA (n=86) | FREEWAY by Eurocor | Primary patency at 1 y | 1 y | NCT01247402 | 3 mo |
| FREEWAY52         | 2017 Multicenter Single-blind (1:1) | DCB+BMS (n=105) vs PTA+BMS (n=99) | FREEWAY by Eurocor | Target lesion revascularization | 1 y | NCT01960647 | Not specified |
| DRECOREST46       | 2018 Single-center Open (1:1) | DCB (n=30) vs PTA (n=30) | IN.PACT by Medtronic for failing bypass | Target lesion revascularization | 1 y | NCT03023098 | 3 mo |

BMS indicates bare metal stent; DCB, drug-coated balloon; DES, drug-eluting stent; ISR, in-stent restenosis; PTA, percutaneous transluminal angioplasty; PTFE, polytetrafluoroethylene.

*The design of the ZILVER PTX study included a primary randomization (optimal PTA vs primary DES) and a secondary randomization in the case of PTA failure (bailout BMS vs bail-out DES)—results of the 2 randomization levels were pooled for the purposes of the present meta-analysis. For the Thunder trial (3-arm trial), the arm of paclitaxel dissolved in the contrast medium was excluded. For the ISAR-STATH trial (3-arm trial), the arm of directional atherectomy was excluded from the present analysis. For the DEBATE in SFA trial, the 2 BMS arms (with or without cilostazol) were pooled against the ZILVER PTX arm. For the DEBELLUM trial, only femoropopliteal lesions were analyzed. In the LEVANT I trial, randomization between plain BA and PCB was performed after provisional stent placement in a quarter of the cases (26 of 101). In the RAPID study, the Supera biomimetic nitinol stent was used in both arms. Data extraction was supplemented by online archived material from international meetings or regulatory authority filings as cited (US Food and Drug Administration—Japan Pharmaceuticals and Medical Devices Agency).
All-Cause Death at 1 Year

All-cause patient death up to 1 year was reported by all included RCTs for a total of 4432 subjects. There was good evidence that the pooled risk of death did not differ significantly between the active use of paclitaxel-coated balloons or stents versus the control arms. There were 58 deaths out of 2506 patients in the paclitaxel arms (2.3% crude risk of death) compared with 45 deaths in 1926 patients in the control arms (2.3% crude risk of death) with a calculated pooled risk ratio of 1.08 (95% CI, 0.72–1.61; Figure 1). There was no statistically significant heterogeneity between studies ($P=0.98$).

All-Cause Death at 2 Years

In all, 12 studies out of the 28 RCTs reported the incidence of all-cause patient death up to 2 years in a total of 2316 patients. There was good evidence that application of paclitaxel-coated devices in the femoropopliteal artery was related to significantly increased risk of death. There were 101 deaths out of 1397 patients in the paclitaxel arms (7.2% crude risk of death) compared with 35 deaths in 919 patients in the control arms (3.8% crude risk of death) with a calculated risk ratio of 1.68 (95% CI, 1.15–2.47). Absolute risk difference was 3.5% (95% CI, 1.7–5.3%) with a corresponding NNH of 29 patients (95% CI, 19–59). There was no statistically significant heterogeneity between studies ($P=0.80$; Figure 2).

All-Cause Death Up to 5 Years

Long-term analysis of all-cause death up to 5 years was informed by 3 RCTs including 863 cases. One study had 4 years,56 and 2 had 5 years of follow-up.9,57 Some 78 deaths out of 529 cases occurred in the paclitaxel arms (14.7% crude...
risk of death) versus 27 deaths out of 334 cases in the control (8.1% crude risk of death) with a pooled risk ratio of 1.93 (95% CI, 1.27–2.93). Absolute risk difference was 7.2% (95% CI, 3.1–11.3%) and NNH was 14 patients (95% CI, 9–32). There was no statistically significant heterogeneity between studies (P=0.92; Figure 3).

Sensitivity and Subgroup Analyses

There was no visual asymmetry of the respective funnel plots to suggest publication bias at 1 year (Horbold-Egger test, 0.27; P=0.66), 2 years (Horbold-Egger test, 0.47; P=0.55), or 4 to 5 years of follow-up (Horbold-Egger test, −0.43; P=0.11; Figure S3). We opted to report summary estimates from a frequentist random effects model to account for conceptual and study design differences among different RCTs. The randomized studies included here tested numerous designs of paclitaxel-coated devices with variable drug dosages and different drug excipients in slightly different patient populations. Hence, a different, but similar treatment effect was assumed as the basis for the random effects modeling.58 In addition, different methods of analyses (with or without continuity correction and Bayesian methods) were employed to interrogate potential bias and uncertainty arising from meta-analysis of low event rates as recommended elsewhere.59 Bayesian methods generally increased the size of treatment effect (Table S4). The pooled point estimates remained consistent across sensitivity and subgroup analyses with some variation in the magnitude of effect size. There were also differences in the estimated long-term risk of death when examining different paclitaxel doses, although those results are underpowered, with variable follow-up periods, and informed by few studies in each case (Table 2).

Figure 2. Random effects forest plot of all-cause death at 2 years. Pooled point estimate was expressed as risk ratio (RR).

Figure 3. Random effects forest plot of all-cause death at 4 to 5 years. Pooled point estimate was expressed as risk ratio (RR).
Table 2. Sensitivity and Subgroup Analyses of All-Cause Patient Death

| All-cause death at 2 y                      | Risk Ratio (95% CI) |
|--------------------------------------------|---------------------|
| Fixed effects model                        | 1.84 (1.27–2.68)    |
| Random effects model                       | 1.68 (1.15–2.47)    |
| All-cause death at 4 to 5 y                |                     |
| Fixed effects model                        | 1.94 (1.28–2.96)    |
| Random effects model                       | 1.93 (1.27–2.93)    |
| Subgroups (random effects)                 |                     |
| Paclitaxel DES only                        | 1.87 (1.11–3.15)    |
| Paclitaxel DCB only                        | 1.44 (1.04–2.00)    |
| Multicenter studies only                   | 1.48 (1.11–1.97)    |
| Dose subgroups (beyond 1 y)                |                     |
| 3.5 μg/mm² paclitaxel balloon             | 2.31 (1.15–4.63)    |
| 3.0 μg/mm² paclitaxel stent               | 2.10 (1.15–3.83)    |
| 3.0 μg/mm² paclitaxel balloon             | 1.65 (0.95–2.87)    |
| 2.0 μg/mm² paclitaxel balloon             | 1.27 (0.70–2.32)    |
| Trial sequential analysis (TSA; random effects at 2 y) |   |
| TSA diversity adjusted (x=5%, β=20%)      | 1.70 (1.19–2.43)    |
| TSA diversity adjusted (x=5%, β=10%)      | 1.70 (1.24–2.33)    |
| TSA diversity adjusted (x=1%, β=10%)      | 1.70 (1.08–2.69)    |

CI indicates confidence interval; DCB, drug-coated balloon; DES, drug-eluting stent.

Meta-Regression Analysis

In line with Bradford Hill’s criteria for establishing epidemiological evidence for causation, we observed a biological gradient, that is, whether greater exposure leads to greater incidence of the effect. To that end, we performed meta-regression of the absolute risk difference of all-cause death against exposure to paclitaxel in all 28 RCTs. Considering that crystalline paclitaxel delivered by paclitaxel-coated devices has a half-life of weeks to months, we calculated exposure to paclitaxel as the dose-time product after treatment. To account for nominal device dose and treated vessel surface area, the following equation was used for calculation of paclitaxel dose-time product expressed in milligram-years for each individual study (i):

\[
Exposure_i = \text{Dose} \times \left(\pi \times D_i \times \text{Length}_i\right) \times \text{Time}_i
\]

where, Dose, is the nominal paclitaxel dose loaded on the balloon or stent (μg/mm²), D_i is the reference vessel diameter (mm), Length_i is the treated lesion length (mm), and Time_i indicates the available follow-up time period (years). Random effects meta-regression identified a highly significant association between paclitaxel dose-time product and absolute risk of death; there was a 0.4±0.1% excess risk of death for every paclitaxel milligram-year (95% CI, 0.1–0.6%; P<0.001; Figure 5). The result was stable on a resampling permutation test (1000 iterations; P=0.013).

Discussion

Paclitaxel-coated balloon and stents have emerged as the most promising strategies for the inhibition of neointimal hyperplasia following angioplasty of the femoropopliteal artery of the leg. Several randomized controlled studies have already demonstrated strong evidence of clinical effectiveness enrolling predominantly patients suffering from intermittent claudication (≈90% of the sample size). Following the progressive release of longer-term clinical outcomes, a comprehensive updated systematic review and meta-analysis was undertaken to compare the all-cause patient mortality associated with the use of those devices. Overall, there was good evidence that all-cause death at 1 year was equivalent between paclitaxel and control arms. However, the risk of all-cause death appeared to increase dramatically after the first year in the case of paclitaxel arms. Synthesis of study-level outcomes at 2 years documented a significant 68% relative risk increase of all-cause death with a corresponding NNH of 29 patients. Risk of death increased further in the long-term analysis (at 5 years) with a 93% relative risk increase and an NNH of 14. Overall, the present statistical inference appeared to be credible and stable on various sensitivity tests. Furthermore, we found neither any statistical heterogeneity...
nor any major diversity between the included studies. We employed trial sequential analysis to address potential random sampling errors and other unknown bias because of the fact that none of the present studies was designed or adequately powered to explore the outcome of patient mortality. Interestingly, TSA meta-analysis powered at 90% has shown accumulation of adequate information size to exclude false-positive findings (type I error) with 99% certainty.

The authors consider the herein reported findings of particular concern because most of the interrogated devices have already received clearance by regulatory authorities and are currently under routine clinical use. The potential causes of this alarming late increased incidence of death remain unknown. Experience with paclitaxel-coated devices has been previously limited to the coronary TAXUS stents (Boston Scientific, Marlborough, MA), which allow for prolonged release of paclitaxel from a polymer-based stent coating. Of note, long-term results of the safety of the TAXUS paclitaxel stent in the heart (patient-level analysis of 2797 randomized patients receiving TAXUS versus bare stents) have long shown a significant increase of long-term death and myocardial infarction after 1 year following implantation (6.7% versus 4.5%, \( P<0.01 \)).

Likewise, long-term results from the SYNTAX (TAXUS Drug-Eluting Stent Versus Coronary Artery Bypass Surgery for the Treatment of Narrowed Arteries) study (1800 patients with complex coronary artery disease randomized between TAXUS paclitaxel stent and coronary artery bypass surgery) have shown a significantly higher cardiovascular mortality up to 5 years in the case of paclitaxel coronary interventions compared with bypass grafting (9.6% versus 5.6%; \( P=0.008 \)) to be explained mostly by late myocardial infarctions. The latter was further confirmed in the nested nonrandomized SYNTAX registries extending to involve both a cardiovascular (12.1% versus 4.7%) and an inexplicable noncardiovascular (14.9% versus 5.3%) mortality difference. Consequently, the coronary field has gradually moved away from paclitaxel DES also because of the well-recognized issues of vessel wall tissue inflammation, aneurysm formation, and late stent thrombosis.

In addition, most drug-coated balloons and stents for the femoropopliteal artery contain at least an order of magnitude higher payload of paclitaxel in comparison with paclitaxel-eluting coronary stents (a 3.5\( \times \)32 mm coronary TAXUS stent contains around 200 \( \mu \)g paclitaxel compared with around 1.2 mg for the ZILVER-PTX 6.0\( \times \)120 stent, 4.5 mg for the LUTONIX 6.0\( \times \)120 drug-coated balloon and 8.5 mg for the IN.PACT 6.0\( \times \)120 balloon). From a pharmacological viewpoint, paclitaxel-coated balloons and stents have been engineered to deliver prolonged levels of paclitaxel into the vessel tissues so as to inhibit smooth muscle cell proliferation and avert vessel restenosis. Paclitaxel is a lipophilic cytotoxic agent that blocks the cell cycle during mitosis by interfering with the spindle disassembly. To enable sustained tissue bioavailability without a polymer, most modern balloons and stents (Table S2) are coated with a mostly solid crystalline form of paclitaxel combined with a unique paclitaxel spacer or

**Figure 4.** Trial sequential analysis of all-cause death. External red lines denote the O’Brien-Fleming alpha spending trial sequential monitoring boundaries. Internal red wedge lines denote the futility O’Brien-Fleming beta spending lines. Cumulative Z curve (blue line) crossed the alpha monitoring boundaries and the required information size (patient sample) has been reached in both illustrative scenarios; (A) \( \alpha=5\% \), \( \beta=10\% \); and (B) \( \alpha=1\% \), \( \beta=10\% \). Vertical red line denotes the calculated required sample size, whereas the Z value is the test statistic (\( |Z|=1.96 \) corresponds to a \( P \) value of 0.05; the higher the Z value, the lower the \( P \) value).
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Figure 5. Meta-regression (mixed effects model) of all-cause death against paclitaxel exposure (dose-time product calculated in milligram-years). The size of the blue symbols is inversely proportional to the variance of the estimated treatment effect for each study. Solid and dotted red lines indicate the regression line with its corresponding 95% confidence bands. Intercept is \(-0.8\pm0.9\%\) and coefficient of the regression line is \(0.4\pm0.1\%\) (95% confidence interval, 0.1–0.6%; \(P<0.001\)). The equation of the regression line is \(Y=(−0.008)+0.004X\). The “metareg” function of the “meta” library was employed in R language.

The authors postulate that late paclitaxel toxicity may be the reason for the observed increased death rate. Contrary to solvent-based (eg, cremophor) intravenous paclitaxel used in cancer chemotherapy that has a half-life of around 6 hours,72 paclitaxel crystals loaded on DCB or DES for the peripheral arteries have a half-life of weeks to months, depending on the exact chemical properties of the applied paclitaxel formulation.62–64 Increased paclitaxel crystallinity helps achieve higher tissue uptake and retention and improved biologic effect, however, at the expense of microparticle formation that may embolize in the downstream systemic circulation.63 Worrisome rates of potential downstream embolization of the skeletal muscles of the lower limbs have been confirmed on the bench and animal studies62,73 and are postulated to be responsible for the significantly higher rates of major amputations noted in the active paclitaxel arm of the randomized INPACT-DEEP study.8,71 Preclinical follow-up studies have shown that in the case of paclitaxel-coated balloons, \(\approx1\%\) to \(10\%\) of the paclitaxel dose gets transferred into the target vessel wall, and as much as \(90\%\) (or as much as 8.5 mg of paclitaxel on a 6.0 × 120 mm 3.5 \(\mu\)g/mm\(^2\) device) gets lost into the systemic circulation with unknown consequences.64,74 Rates of distal paclitaxel embolization, if any, in the case of paclitaxel DES remain unknown.70

Within the modern epidemiologic framework of structural causal modeling developed by Judea Pearl,75,76 the present work shows strong signals of biomedical causality between paclitaxel and mortality within multiple controlled randomized trials. According to the more traditional Bradford Hill criteria61 for establishing a causal relationship between a presumed cause and an observed health effect, the present work has shown evidence of strength, consistency, temporality, and biological gradient. Risk of death was identical at 1 year but more than doubled during the second year following intervention. Twelve of 13 studies showed increased mortality between 1 and 2 years after intervention. In addition, a significant relationship between dose-time exposure to paclitaxel and incidence of deaths was identified; risk of death increased by \(0.4\pm0.1\%\) per paclitaxel milligram-year. Interestingly, the risk of death beyond 1 year also seemed to vary among different paclitaxel dosages, being significantly higher in the 3.5 \(\mu\)g/mm\(^2\) devices compared with the lower-dose devices (Table 2). Still, the present meta-analysis is underpowered to discern outcome differences between the different paclitaxel devices as some devices are supported by a single trial and follow-up beyond 2 years is missing in most cases. The authors would therefore encourage collection of longer-term follow-up (beyond 1 year) in case of all studies to help confirm or refute the present findings.

Two-year clinical outcomes from large-scale phase IV registries have been recently released for 2 DCB and 1 DES device in the femoropopliteal artery. The ZILVER-PTX postmarket single-arm DES surveillance registry reported a 2.6% annualized risk of death (41 of 787 subjects at 2 years),19 the Lutonix Global SFA registry stated a 3.0% risk (38 of 637 died at 2 years),77 and the IN.PACT Global postmarket DCB study a 3.5% annualized risk of death (89 of 1269 at 2 years).78 The

Table 3. Causes of Death

| Cause                  | Paclitaxel-Coated Balloon (INPACT SFA) at 3 Years\(^{10,82}\) | Paclitaxel-Coated Stent (ZILVER PTX) at 2 Years\(^{19,23}\) |
|------------------------|---------------------------------------------------------------|----------------------------------------------------------|
|                        | Paclitaxel Control | Paclitaxel Control | Paclitaxel Control | Paclitaxel Control |
| Cardiovascular         | 9                  | 0                  | 18                 | 8                  |
| Cancer                 | 2                  | 2                  |                    |                    |
| Infectious             | 5                  | 0                  |                    |                    |
| Pulmonary              | 3                  | 0                  |                    |                    |
| Other                  | 3                  | 0                  | NA                 | NA                 |

NA indicates not applicable.
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The aforementioned rates of patient death appear to be consistent with an annualized 3.1% incidence of death in the case of paclitaxel DES and DCB documented in the present meta-analysis (compared with 1.9% in the control angioplasty arms). Population-based contemporary series of intermittent claudication that constituted 89% of the patient population included here have documented a 1.0% to 2.0% annual mortality rate,\(^6\) which is consistent with the 1.9% incidence of all-cause death in the control arms of the current analysis.

The present meta-analysis has several limitations. First, we excluded studies with DCB or DES in the below-knee infrapopliteal arteries as they pertain to a distinctively different patient population mostly with critical limb ischemia associated with high morbidity and mortality rates. Second, some study protocols did not include an independent blinded clinical events committee for event adjudication, and nearly universally a single-blind or open-label study design was applied that may have introduced detection or performance bias, respectively. Third, unfortunately and with few exceptions,\(^9,10,19,82\) most studies did not report the actual causes of death to help infer potential causal links with paclitaxel use. An association of paclitaxel with more cardiovascular deaths, but also of infectious, gastrointestinal, and pulmonary origin, was noted in the INPACT SFA (Randomized Trial of IN.PACT Admiral\(^\text{TM}\) Drug Coated Balloon vs Standard PTA for the Treatment of SFA and Proximal Popliteal Arterial Disease) and ZILVER PTX (Evaluation of the Zilver PTX Drug-Eluting Peripheral Stent) studies (Table 3). Although baseline demographics were generally well balanced, in a few studies a numerically greater incidence of patient comorbidities—eg, smoking, hyperlipidemia, hypertension, or diabetes mellitus—were noted in the paclitaxel arms, for example, in the ZILVER PTX study. Hence, undetected sources of heterogeneity could not be explored in depth in the absence of individual patient data. Finally, we could not establish a plausible mechanism between paclitaxel and deaths, but as Sir Bradford Hill noted, knowledge of the mechanism may be limited by current knowledge.\(^6\)

In conclusion, there seems to be an increased long-term risk of death beyond the first year following femoropopliteal application of paclitaxel-coated balloons and stents in the lower limbs. Actual causes for this serious late side effect remain unknown, and further investigations with longer-term follow-up are urgently warranted.

Acknowledgments

Data sharing: All authors had unrestricted access to the data sets and can take responsibility for the integrity of the data and the accuracy of the data analysis. The lead and corresponding author (Katsanos) performed all statistical analyses and has final overall responsibility for the submitted version of the manuscript (study guarantor). The lead and corresponding author (Katsanos) affirms that the 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. Raw data are presented in the submitted tables and figures.

Disclosures

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SUPPLEMENTAL MATERIAL
### Table S1. Design characteristics of the included randomized controlled trials*

| Study and sources | Year and study design | Allocation in study arms | Paclitaxel coated device | Primary study endpoint | Maximum follow-up period | Study registration | Dual antiplatelet therapy |
|-------------------|-----------------------|--------------------------|--------------------------|------------------------|--------------------------|-------------------|--------------------------|
| ZILVER-PTX 1-3    | 2011 Multi-center Open label (1:1) | DES (n=241) vs PTA (n=238) | ZILVER-PTX Stent by COOK Medical | Primary Patency at 1 year | 5 years | NCT00120406 | >2 months |
| THUNDER 4-5       | 2008 Multi-center Single-blind (1:1:1) | DCB (n=48) vs PTA (n=54) | Cotavance Balloon by Bavaria Medizin | Late Lumen Loss at 6 months | 5 years | NCT00156624 | 1 month |
| IN.PACT SFA 6-10  | 2015 Multi-center Single-blind (2:1) | DCB (n=220) vs PTA (n=111) | IN.PACT Admiral by Medtronic | Primary Patency at 1 year | 3 years | NCT01175850 NCT01666461 | 1 month (3 months if bail-out stenting) |
| FEMPAC 11         | 2008 Multi-center Single-blind (1:1) | DCB (n=45) vs PTA (n=42) | Paccocath Balloon by Bavaria Medizin | Late Lumen Loss at 6 months | 2 years | NCT00472472 | Long-term (not specified) |
| LEVANT I 12       | 2012 Multi-center Single-blind (1:1) | DCB (n=49) vs PTA (n=52) | Lutonix by CR BARD | Late Lumen Loss at 6 months | 2 years | NCT00930813 | 1 month (3 months if bail-out stenting) |
| LEVANT II 13-16   | 2015 Multi-center Single-blind (2:1) | DCB (n=316) vs PTA (n=160) | Lutonix by CR BARD | Primary Patency at 1 year | 2 years | NCT01412541 | 1 month |
| ILLUMENATE EU 17  | 2017 Multi-center Single-blind (2:1) | DCB (n=222) vs PTA (n=72) | Stellarex by Spectranetics | Primary Patency at 1 year | 2 years | NCT01858363 | 1 month (3 months if bail-out stenting) |
| CONSEQUENT 18     | 2017 Multi-center Single-blind (1:1) | DCB (n=78) vs PTA (n=75) | SeQuent Please by B.Braun | Late Lumen Loss at 6 months | 2 years | NCT01970579 | 2 months |
| ISAR-STATH 19     | 2017 Two-center Open label (1:1:1) | DCB+BMS (n=48) vs PTA+BMS (n=52) | IN.PACT Admiral by Medtronic | Diameter Stenosis at 6 months | 2 years | NCT00986752 | 6 months |
| ISAR-PEBIS 20     | 2017 Two-center Open label (1:1) for ISR | DCB (n=36) vs PTA (n=34) | IN.PACT Admiral by Medtronic | Diameter Stenosis at 6-8 months | 2 years | NCT01083394 | >6 months |
| IN.PACT SFA JAPAN 21 | 2018 Multi-center Single-blind (2:1) | DCB (n=68) vs PTA (n=32) | IN.PACT Admiral by Medtronic | Primary Patency at 1 year | 2 years | NCT01947478 | 1 month (3 months if bail-out stenting) |
| Study | Year | Center | Randomization | Intervention 1 | Intervention 2 | Endpoint | Duration | Registration | Follow-up |
|-------|------|--------|---------------|----------------|----------------|----------|----------|-------------|-----------|
| ACOART | 2016 | Multi-center | Single-blind (1:1) | DCB (n=100) vs PTA (n=100) | Orchid by Acotec Scientific | Late Lumen Loss at 6 months | 2 years | Not registered | 6 months |
| FINN-PTX | 2018 | Multi-center | Open label (2:1) | DES (n=28) vs PTFE (n=18) | ZILVER-PTX Stent by COOK Medical | Secondary Patency at 2 years | 2 years | NCT01450722 | 3 months (Aspirin in control group) |
| BATTLE | 2018 | Multi-center | Open label (1:1) | DES (n=86) vs BMS (n=85) | ZILVER-PTX Stent by COOK Medical | In-stent binary restenosis at 1 year | 1 year | NCT02004951 | >2 months (clopidogrel to continue for 2 years) |
| DEBATE-IN-SFA | 2018 | Multi-center | Open label (1:1:1) | DES (n=85) vs BMS (n=170) | ZILVER-PTX Stent by COOK Medical | In-stent binary restenosis at 1 year | 1 year | UMIN000010071 | >2 months (Aspirin to continue lifelong) |
| DEBELLUM | 2014 | Single-center | Open (1:1) | DCB (n=25) vs PTA (n=25) | IN.PACT Admiral by Medtronic | Late Lumen Loss at 6 months | 1 year | Not registered | 1 month |
| PACIFIER | 2012 | Multi-center | Single-blind (1:1) | DCB (n=41) vs PTA (n=44) | IN.PACT Pacific by Medtronic | Late Lumen Loss at 6 months | 1 year | NCT01083030 | >2 months |
| FAIR | 2015 | Multi-center | Single-blind (1:1) for ISR | DCB (n=62) vs PTA (n=67) | IN.PACT Admiral by Medtronic | 6-month binary restenosis | 1 year | NCT01305070 | >6 months |
| BIOLUX P-I | 2015 | Multi-center | Single-blind (1:1) | DCB (n=30) vs PTA (n=30) | Passeo-18 Lux by Biotronik | Late Lumen Loss at 6 months | 1 year | NCT01056120 | 1 month (3 months if bail-out stenting) |
| RANGER SFA | 2018 | Multi-center | Single-blind (2:1) | DCB (n=71) vs PTA (n=34) | Ranger by Boston Scientific | Primary Patency at 1 year | 1 year | NCT02013193 | >1 month |
| ILLUMENATE pivotal | 2017 | Multi-center | Single-blind (2:1) | DCB (n=200) vs PTA (n=100) | Stellarex by Spectranetics | Primary Patency at 1 year | 1 year | NCT01858428 & NCT01912937 | 1 month |
| DEBATE-SFA | 2013 | Single-center | Open (1:1) | DCB+BMS (n=53) vs PTA+BMS (n=51) | IN.PACT Admiral by Medtronic | 1-year binary restenosis | 1 year | NCT01556542 | 3 months |
| LUTONIX JAPAN | 2018 | Multi-center | Japan (1:1) | DCB (n=71) vs PTA (n=38) | Lutonix by CR BARD | Primary Patency at 1 year | 1 year | Not registered | 1 month |
| Study | Year   | Setting | Randomization | Intervention 1 | Intervention 2 | Primary Endpoint | Follow-up | Study ID | Time Frame |
|-------|--------|---------|---------------|----------------|----------------|------------------|-----------|----------|------------|
| RAPID | 2017   | Multi-center | Double-blind (1:1) | DCB+BMS (n=80) vs PTA+BMS (n=80) | LegFlow by Cardionovum | Primary Patency at 1 year | 1 year | ISRCTN7846578 | 3 months |
| EFFPAC| 2018   | Multi-center | Single-blind (1:1) | DCB (n=85) vs PTA (n=86) | Luminor by iVascular | Late Lumen Loss at 6 months | 1 year | NCT02540018 | >1 month |
| PACUBA| 2016   | Dual-center | Single-blind (1:1) for ISR | DCB (n=85) vs PTA (n=86) | FREEWAY by Eurocor | Primary Patency at 1 year | 1 year | NCT01247402 | 3 months |
| FREEWAY| 2017   | Multi-center | Single-blind (1:1) | DCB+BMS (n=105) vs PTA+BMS (n=99) | FREEWAY by Eurocor | Target lesion revascularization | 1 year | NCT01960647 | Not specified |
| DRECOREST| 2018 | Single-center | Open (1:1) | DCB (n=30) vs PTA (n=30) | IN.PACT by Medtronic for failing bypass | Target lesion revascularization | 1 year | NCT03023098 | 3 months |
Table S2. Design characteristics of the tested paclitaxel DES and DCB devices.

| Brand name       | Paclitaxel dose (μg/mm²) | Excipient/spacer                              | Manufacturer                                      |
|------------------|--------------------------|------------------------------------------------|---------------------------------------------------|
| IN.PACT          | 3.5 μg/mm² (3.7 μg/mm²) | Urea (FreePac)                                 | Medtronic (dose based on FDA submission)          |
| ZILVER-PTX       | 3.0 μg/mm² (0.37 μg/mm²) | None (polymer-free stent)                      | COOK Medical (area adjusted dose in case of stents) |
| Cotavance        | 3.0 μg/mm²               | Paccocath (iodinated contrast)                 | Bavaria Medizin Technologie MedRad, later Bayer  |
| Passeo-18 Lux    | 3.0 μg/mm²               | Butyryl-tri-n-hexyl citrate (BTHC)             | Biotronik                                        |
| SeQuent Please   | 3.0 μg/mm²               | Resveratrol                                    | B.Braun                                          |
| FREEWAY          | 3.0 μg/mm²               | Shellac (shellolic and aleuritic acid resin)   | Eurocor                                          |
| LegFlow          | 3.0 μg/mm²               | Nanocrystalline 0.1-μm paclitaxel in ammonium salt | Cardionovum                                    |
| Orchid           | 3.0 μg/mm²               | Magnesium stearate                             | Acotec Scientific                                |
| Lutonix          | 2.0 μg/mm²               | Polysorbate and sorbitol                       | C.R. BARD                                       |
| Luminor          | 3.0 μg/mm²               | Organic ester                                  | iVascular                                       |
| Stellarex        | 2.0 μg/mm²               | Polyethylene glycol                            | Spectranetics                                   |
| Ranger           | 2.0 μg/mm²               | acetyl tributyl citrate – ATBC (Transpax)      | Boston Scientific                               |

Nominal paclitaxel dose is expressed in micrograms/mm² (μg/mm²). Based on the relevant FDA submission, dose is around (3.7μg/mm²) for the IN.PACT drug-coated balloon. In case of the ZILVER-PTX drug-coated stent, nominal paclitaxel dose adjusted for corresponding vessel surface area is actually 0.37μg/mm² based on corresponding FDA filing data. The latter doses were used for calculation of paclitaxel dose for the purposes of meta-regression analysis.
Table S3. Baseline patient characteristics of included randomized clinical trials.

| Study allocation | ZILVER-PTX | THUNDER | IN.PACT SFA | FEMPAC |
|------------------|------------|---------|-------------|-------|
| Patients (limbs) | DES        | PTA     | DCB         | PTA   |
| n                | 241        | 238     | 220         | 45    |
| Age (years)      | 68±10      | 68±11   | 68±10       | 67±6  |
| Male gender      | 155 (66%)  | 152 (64%) | 31 (65%) | 143 (65%) |
| Smoking          | 204 (86%)  | 200 (84%) | 11 (23%) | 85 (39%) |
| Hypertension     | 210 (89%)  | 194 (82%) | 38 (79%) | 201 (91%) |
| Hyperlipidemia   | 180 (76%)  | 166 (70%) | 33 (69%) | 186 (85%) |
| Diabetes mellitus| 116 (49%)  | 100 (42%) | 24 (50%) | 89 (41%) |
| Coronary artery disease | 50 (21%) | 41 (17%) | NR | 122 (57%) |
| Renal insufficiency | 24 (10%) | 25 (11%) | NR | NR |
| Intermittent claudication | 217 (90%) | 216 (91%) | 35 (73%) | 209 (95%) |
| Critical limb ischemia | 24 (10%) | 22 (9%) | 13 (27%) | 11 (5%) |
| Lesions treated | n=247      | n=251   | n=86        | n=221 |
| Lesion Length (cm) | 6.6±3.9  | 6.3±4.1 | 7.5±6.2     | 8.9±4.9 |
| Vessel Diameter (mm) | NA      | NA      | 5.0±0.7     | 4.7±0.8 |
| Total occlusions | 73 (30%)  | 62 (25%) | 13 (27%)    | 57 (26%) |
| Bail-out stenting | NA      | (2-level random*) | 2 (4%) | 16 (7%) |
Table S3. Baseline patient characteristics of included randomized clinical trials (continued).

| Study allocation | LEVANT I | LEVANT II | ILLUMENATE EU | CONSEQUENT |
|------------------|----------|-----------|---------------|------------|
| Patients (limbs) | DCB      | PTA       | DCB           | PTA        |
| n=49             | n=52     | n=316     | n=160         | n=222      |
| Age (years)      | 67±8     | 70±10     | 68±10         | 69±9       |
| Male gender      | 34 (69%) | 30 (58%)  | 193 (61%)     | 107 (67%)  |
| Smoking          | 15 (31%) | 20 (39%)  | 111 (35%)     | 54 (34%)   |
| Hypertension     | 47 (96%) | 45 (87%)  | 282 (89%)     | 140 (88%)  |
| Hyperlipidemia   | 29 (59%) | 36 (69%)  | 283 (90%)     | 138 (86%)  |
| Diabetes mellitus| 22 (45%) | 26 (50%)  | 137 (43%)     | 67 (42%)   |
| Coronary artery disease | 19 (39%) | 23 (44%) | 157 (50%) | 77 (48%) |
| Renal insufficiency | NR      | NR        | 11 (4%)       | 7 (4%)     |
| Intermittent claudication | 46 (94%) | 48 (92%) | 291 (92%) | 147 (92%) |
| Critical limb ischemia | 3 (6%) | 4 (8%) | 25 (8%) | 13 (8%) |
| Lesions treated | n=49 | n=52 | n=322 | n=165 |
| Lesion Length (cm) | 8.1±3.7 | 8.0±3.8 | 6.3±4.0 | 6.3±4.0 |
| Vessel Diameter (mm) | 4.1±0.6 | 4.2±0.7 | 4.8±0.8 | 4.8±0.8 |
| Total occlusions | 20 (41%) | 22 (42%) | 65 (21%) | 35 (22%) |
| Bail-out stenting | 12 (24%) | 14 (27%) | 8 (2.5%) | 11 (6.9%) |
Table S3. Baseline patient characteristics of included randomized clinical trials (continued).

| Study allocation | ISAR-STATH | IN.PACT SFA JAPAN | DEBELLUM | PACIFIER |
|------------------|------------|-------------------|----------|----------|
| Patients (limbs) | n=48       | n=68              | n=25     | n=41     |
|                  | n=52       | n=32              | n=25     | n=44     |
| Age (years)      | 70±9       | 69±8              | 73±7     | 67±7     |
|                  |            | 74±6              | 67±6     | 71±7     |
| Male gender      | 33 (69%)   | 37 (71%)          | 50 (74%) | 19 (76%) |
|                  |            | 26 (81%)          | 18 (72%) | 26 (59%) |
|                  |            |                   | 18 (72%) | 30 (64%) |
| Smoking          | 36 (75%)   | 34 (65%)          | 18 (26%) | 17 (68%) |
|                  |            | 10 (31%)          | 14 (56%) | 21 (49%) |
|                  |            |                   | 28 (60%) |          |
| Hypertension     | 40 (83%)   | 40 (77%)          | NR       | 19 (76%) |
|                  |            |                   | 15 (60%) | 29 (66%) |
|                  |            |                   | 31 (66%) |          |
| Hyperlipidemia   | 45 (94%)   | 45 (87%)          | NR       | 12 (48%) |
|                  |            |                   | 17 (68%) | 22 (50%) |
|                  |            |                   | 22 (47%) |          |
| Diabetes mellitus| 10 (21%)   | 15 (29%)          | 40 (59%) | 13 (52%) |
|                  |            |                   | 9 (36%)  | 19 (43%) |
|                  |            |                   | 13 (28%) |          |
| Coronary artery disease | 25 (52%) | 24 (46%)          | 34 (50%) | 16 (50%) |
|                  |            |                   | NR       | 14 (32%) |
|                  |            |                   | 15 (32%) |          |
| Renal insufficiency | NR       | NR                | 6 (9%)   | 4 (13%)  |
|                  |            |                   | NR       | 12 (48%) |
|                  |            |                   | 17 (68%) | 22 (50%) |
|                  |            |                   | 22 (47%) |          |
| Intermittent claudication | 45 (94%) | 48 (92%)          | 65 (96%) | 31 (97%) |
|                  |            |                   | 23 (92%) | 22 (88%) |
|                  |            |                   | 42 (95%) | 45 (96%) |
| Critical limb ischemia | 3 (6%)  | 4 (8%)            | 3 (4%)   | 1 (3%)   |
|                  |            |                   | 2 (8%)   | 3 (12%)  |
|                  |            |                   | 2 (5%)   | 2 (4%)   |
| Lesions treated  | n=48       | n=52              | n=68     | n=44     |
|                  | n=32       | n=32              | n=44     | n=48     |
|                  | n=44 (62)  | n=47 (55)         |          |          |
| Lesion Length (cm) | 6.8±4.4  | 7.4±5.6           | 13.4±5.1 | 13.7±5.6 |
|                  |            | 13.7±5.6          | 7.6±0.6  | 7.8±0.7  |
|                  |            | 7.0±5.3           | 6.6±5.5  |          |
| Vessel Diameter (mm) | 5.0±1.0  | 5.0±0.9           | 4.8±0.8  | 4.7±0.7  |
|                  |            |                   | NA       | NA       |
|                  |            |                   | 4.9±1.3  | 4.9±1.3  |
| Total occlusions | 28 (58%)   | 35 (67%)          | 11 (16%) | 5 (16%)  |
|                  |            |                   | 5 (11%)  | 9 (19%)  |
|                  |            |                   | 10 (23%) | 18 (38%) |
| Bail-out stenting | 48 (100%) | 52 (100%)         | 3 (4%)   | 1 (3%)   |
|                  |            |                   | 20 (45%) | 21 (44%) |
|                  |            |                   | 9 (21%)  | 16 (34%) |
Table S3. Baseline patient characteristics of included randomized clinical trials (continued).

| Study allocation | ISAR-PEBIS | FAIR | BIOLUX P-I | RANGER-SFA |
|------------------|------------|------|------------|------------|
| Patients (limbs) | n=36       | n=62 | n=30       | n=71       |
| Age (years)      | 70±10      | 69±8 | 70±10      | 68±8       |
| Male gender      | 24 (67%)   | 33 (53%) | 17 (57%) | 53 (75%) |
| Smoking          | 21 (58%)   | 18 (29%) | 19 (63%) | 29 (41%) |
| Hypertension     | 33 (92%)   | 52 (84%) | 23 (77%) | 58 (82%) |
| Hyperlipidemia   | 35 (97%)   | 48 (78%) | 18 (60%) | 49 (69%) |
| Diabetes mellitus| 12 (33%)   | 28 (45%) | 11 (37%) | 28 (39%) |
| Coronary artery disease | 17 (47%) | 26 (42%) | 8 (27%) | 24 (34%) |
| Renal insufficiency | NR | 8 (13%) | NR | 8 (11%) |
| Intermittent claudication | 35 (97%) | 59 (95%) | 24 (80%) | 71 (100%) |
| Critical limb ischemia | 1 (3%) | 3 (5%) | 6 (20%) | 0 (0%) |
| Lesions treated  | n=36       | n=62 | n=33       | n=70       |
| Lesion Length (cm) | 13.2±6.5 | 8.2±7.1 | 5.1±4.7 | 6.8±4.6 |
| Vessel Diameter (mm) | 5.0±1.1 | 5.1±0.9 | 4.6±0.8 | 5.0±0.9 |
| Total occlusions | 13 (36%)   | 15 (24%) | NR | 24 (34%) |
| Bail-out stenting | NA | NA | 2 (7%) | 15 (21%) |
Table S3. Baseline patient characteristics of included randomized clinical trials (continued).

| Study allocation | ACOART-I | DEBATE-SFA | ILLUMENATE pivotal | LUTONIX JAPAN |
|------------------|----------|------------|--------------------|---------------|
| | DCB  | PTA  | DCB+BMS  | PTA+BMS  | DCB  | PTA  | DCB  | PTA  |
| Patients (limbs) | n=100  | n=100  | n=53  | n=51  | n=200  | n=100  | n=71  | n=38  |
| Age (years) | 66±9  | 66±9  | 74±9  | 76±8  | 68±10  | 70±10  | 72±10  | 78±8  |
| Male gender | 73 (73%)  | 74 (74%)  | 50 (75%)  | 42 (63%)  | 112 (56%)  | 64 (64%)  | 45 (63%)  | 46 (68%)  |
| Smoking | 29 (29%)  | 33 (33%)  | 25 (47%)  | 28 (55%)  | 168 (84%)  | 75 (75%)  | 53 (75%)  | 26 (68%)  |
| Hypertension | 62 (62%)  | 72 (72%)  | 47 (89%)  | 45 (88%)  | 187 (94%)  | 94 (94%)  | 60 (85%)  | 35 (92%)  |
| Hyperlipidemia | 27 (27%)  | 29 (29%)  | 33 (62%)  | 27 (53%)  | 176 (88%)  | 90 (90%)  | 47 (66%)  | 26 (68%)  |
| Diabetes mellitus | 54 (54%)  | 57 (57%)  | 41 (77%)  | 36 (71%)  | 99 (50%)  | 52 (52%)  | 33 (47%)  | 18 (47%)  |
| Coronary artery disease | NR  | NR  | 21 (40%)  | 18 (35%)  | 90 (45%)  | 48 (48%)  | 31 (44%)  | 14 (37%)  |
| Renal insufficiency | NR  | NR  | NR  | NR  | 36 (18%)  | 16 (16%)  | 5 (7%)  | 2 (5%)  |
| Intermittent claudication | 60 (60%)  | 66 (66%)  | 11 (21%)  | 16 (31%)  | 192 (96%)  | 95 (95%)  | 71 (100%)  | 37 (97%)  |
| Critical limb ischemia | 40 (40%)  | 34 (34%)  | 42 (79%)  | 35 (69%)  | 8 (4%)  | 5 (5%)  | 0 (0%)  | 1 (3%)  |
| Lesions treated | n=100  | n=100  | n=55  | n=55  | n=200  | n=100  | n=72  | n=40  |
| Lesion Length (cm) | 14.7±11.0  | 15.2±10.9  | 9.4±6.0  | 9.6±6.9  | 8.0±4.5  | 8.9±4.6  | 6.8±4.3  | 5.7±5.1  |
| Vessel Diameter (mm) | 3.8±0.6  | 3.7±0.8  | 5.0±0.5  | 5.1±0.5  | 4.9±0.9  | 5.2±1.1  | 4.9±0.7  | 4.7±0.7  |
| Total occlusions | 57 (57%)  | 52 (52%)  | 30 (55%)  | 38 (69%)  | 38 (19%)  | 18 (18%)  | 13 (18%)  | 2 (5%)  |
| Bail-out stenting | 19 (19%)  | 21 (21%)  | NA  | NA  | 12 (6%)  | 6 (6%)  | 1 (2%)  | 3 (8%)  |
Table S3. Baseline patient characteristics of included randomized clinical trials (continued).

| Study allocation | RAPID | EFFPAC | FINNPTX | BATTLE |
|------------------|-------|--------|---------|--------|
| Patients (limbs) | n=80  | n=85   | n=23    | n=18   |
| Age (years)      | 68±8  | 67±8   | 68±10   | 71±12  |
| Male gender      | 52 (65%) | 50 (63%) | 60 (70%) | 62 (72%) |
| Smoking          | 40 (50%) | 39 (49%) | NR      | 9 (39%) |
| Hypertension     | NR    | NR     | 74 (87%) | 15 (65%) |
| Hyperlipidemia   | NR    | NR     | 60 (71%) | 13 (57%) |
| Diabetes mellitus| 23 (29%) | 24 (30%) | 31 (37%) | 9 (39%) |
| Coronary artery disease | NR | NR | NR | NR |
| Renal insufficiency | NR | NR | NR | NR |
| Intermittent claudication | 66 (82%) | 67 (84%) | 82 (96%) | 68 (79%) |
| Critical limb ischemia | 14 (18%) | 13 (16%) | 3 (4%) | 14 (18%) |
| Lesion treated   | n=80  | n=80   | n=85    | n=23   |
| Lesion Length (cm) | 15.8±7.4 | 15.8±7.6 | 5.6±3.9 | 11.3±4.0 |
| Vessel Diameter (mm) | 5.1±0.7 | 5.2±0.8 | 5.4±0.7 | 5.8±0.6 |
| Total occlusions | 61 (76%) | 56 (70%) | 17 (20%) | 18 (100%) |
| Bail-out stenting | NA    | NA     | 13 (15%) | 86 (100%) |

|     | DCE+BMS | PTA+BMS | PTA  | DES  | PTFE | DES  | BMS  |
|-----|---------|---------|------|------|------|------|------|
| Patients (limbs) | n=80 | n=80 | n=85 | n=86 | n=18 | n=86 | n=85 |
| Age (years) | 68±8 | 67±8 | 68±9 | 68±10 | 67±9 | 71±12 | 68±12 |
| Male gender | 52 (65%) | 50 (63%) | 60 (70%) | 17 (74%) | 12 (67%) | 62 (72%) | 62 (73%) |
| Smoking | 40 (50%) | 39 (49%) | NR | NR | 9 (39%) | 6 (33%) | 20 (23%) | 28 (33%) |
| Hypertension | NR | NR | 74 (87%) | 73 (85%) | 15 (65%) | 15 (83%) | 59 (69%) | 52 (61%) |
| Hyperlipidemia | NR | NR | 60 (71%) | 59 (69%) | 13 (57%) | 15 (83%) | 55 (65%) | 61 (73%) |
| Diabetes mellitus | 23 (29%) | 24 (30%) | 31 (37%) | 35 (41%) | 9 (39%) | 6 (33%) | 41 (48%) | 22 (26%) |
| Coronary artery disease | NR | NR | NR | NR | 6 (26%) | 5 (28%) | 27 (31%) | 34 (40%) |
| Renal insufficiency | NR | NR | NR | NR | NR | NR | 8 (9%) | 6 (7%) |
| Intermittent claudication | 66 (82%) | 67 (84%) | 82 (96%) | 85 (99%) | 17 (74%) | 17 (94%) | 68 (79%) | 70 (82%) |
| Critical limb ischemia | 14 (18%) | 13 (16%) | 3 (4%) | 1 (1%) | 6 (26%) | 1 (6%) | 18 (21%) | 15 (18%) |
| Lesion treated | n=80 | n=80 | n=85 | n=86 | n=23 | n=18 | n=86 | n=85 |
| Lesion Length (cm) | 15.8±7.4 | 15.8±7.6 | 5.6±3.9 | 13.2±6.2 | 11.3±4.0 | 7.3±3.2 | 7.3±3.9 |
| Vessel Diameter (mm) | 5.1±0.7 | 5.2±0.8 | 5.4±0.7 | NA | NA | 5.8±0.6 | 5.8±0.5 |
| Total occlusions | 61 (76%) | 56 (70%) | 17 (20%) | 22 (26%) | 23 (100%) | 18 (100%) | NR | NR |
| Bail-out stenting | NA | NA | 13 (15%) | 16 (19%) | 23 (100%) | NA | 86 (100%) | 85 (100%) |
Table S3. Baseline patient characteristics of included randomized clinical trials (continued).

| Study allocation | DEBATE in SFA | PACUBA | DRECOREST | FREEWAY |
|------------------|--------------|--------|-----------|---------|
| Patients (limbs) | n=85         | n=170  | n=35      | n=105   |
| Age (years)      | 73±8         | 73±9   | 68±9      | 68±10   |
| Male gender      | 60 (71%)     | 112 (66%) | 20 (57%) | 15 (52%) |
| Smoking          | 24 (28%)     | 41 (24%) | 17 (52%) | 17 (59%) |
| Hypertension     | 68 (80%)     | 142 (84%) | 26 (79%) | 23 (79%) |
| Hyperlipidemia   | 52 (61%)     | 101 (59%) | 18 (55%) | 25 (74%) |
| Diabetes mellitus| 50 (59%)     | 90 (53%) | 17 (52%) | 13 (38%) |
| Coronary artery disease | 44 (52%) | 66 (39%) | NR | NR |
| Renal insufficiency | 17 (20%)     | 36 (21%) | 6 (19%) | 6 (21%) |
| Intermittent claudication | 79 (93%) | 146 (86%) | 35 (100%) | 38 (100%) |
| Critical limb ischemia | 6 (7%)       | 24 (14%) | 0 (0%) | 16 (55%) |
| Lesions treated | n=85         | n=170  | n=35      | n=105   |
| Lesion Length (cm) | 11.1±5.0    | 9.8±4.0 | 17.3±11.3 | 18.4±8.8 |
| Vessel Diameter (mm) | 5.3±1.0   | 5.2±1.1 | 5.7±1.0 | 5.4±0.9 |
| Total occlusions | 43 (51%)     | 67 (37%) | 11 (31%) | 11 (28%) |
| Bail-out stenting | 86 (100%)   | 85 (100%) | NA | NA |
Table S4. Sensitivity analyses of rare events (Risk Ratio; 95%CI or CrI) 41

(R ‘meta’ package (version 4.9-2 – Bayesian with https://gemtc.drugis.org)

|                                | Fixed effects | Random effects |
|--------------------------------|---------------|----------------|
| **All-cause death at 1 year**  |               |                |
| Continuity correction 0.5      | 1.06 (0.73-1.55) | 1.08 (0.72-1.61) |
| Continuity correction 0.01     | 1.07 (0.72-1.58) | 1.05 (0.69-1.62) |
| Treatment arm continuity correction (TACC) | 1.06 (0.73-1.55) | 1.07 (0.71-1.60) |
| Mantel-Haenszel exact method   | 1.07 (0.72-1.58) | 1.05 (0.68-1.61) |
| (no continuity correction)     |               |                |
| Bayesian binomial/log model (risk ratio) | 1.59 (1.12-2.31) | 1.64 (1.12-2.46) |
| (burn-in 50000 and inference 200000 iterations) |         |                |
| **All-cause death at 2 years** |               |                |
| Continuity correction =0.5     | 1.84 (1.27-2.68) | 1.68 (1.15-2.47) |
| Continuity correction =0.01    | 1.87 (1.28-2.72) | 1.64 (1.11-2.42) |
| Treatment arm continuity correction | 1.85 (1.27-2.69) | 1.69 (1.15-2.48) |
| Mantel-Haenszel exact method   | 1.87 (1.28-2.73) | 1.63 (1.11-2.41) |
| (no continuity correction)     |               |                |
| Bayesian binomial/log model (risk ratio) | 2.12 (1.48-3.13) | 2.28 (1.45-4.27) |
| (burn-in 50000 and inference 200000 iterations) |         |                |
| **All-cause death at 4-5 years** |               |                |
| Continuity correction =0.5     | 1.94 (1.28-2.96) | 1.93 (1.27-2.93) |
| Continuity correction =0.01    | 1.94 (1.28-2.96) | 1.93 (1.27-2.93) |
| Treatment arm continuity correction | 1.94 (1.28-2.96) | 1.93 (1.27-2.93) |
| Mantel-Haenszel exact method   | 1.94 (1.28-2.96) | 1.93 (1.27-2.93) |
Bayesian binomial/log model (risk ratio)
(burn-in 50000 and inference 200000 iterations) | 2.00 (1.35-3.11) | 2.01 (1.15-3.61)
Figure S1. Literature search and study selection process following the PRISMA statement.

Records identified through database searches (n = 355) → Additional records identified through online material (n = 33) → Records after duplicates removed (n = 386) → Records screened (n = 386) → Records excluded (n = 342) → Full-text items assessed for eligibility (n = 48) → Studies included in qualitative and quantitative synthesis (meta-analysis) (n = 28) → Articles excluded not meeting inclusion criteria
- Non-randomized
- Follow-up <1 year
- Interim results
- Not paclitaxel
- Below-knee vessels
- Unclear - others (n = 20)
Figure S2. Evaluation of risk of bias of each RCT according to the Cochrane Collaboration Tool.
Figure S3. Funnel plots of all-cause death analyses at (A) 1 year, (B) 2 years, and (C) 4-5 years of follow-up.

The SE of the logRR was plotted against the RR for each trial.
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