Effectiveness of prothrombin complex concentrate for the treatment of bleeding: A systematic review and meta-analysis

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Abstract
Prothrombin complex concentrate (PCC) is increasingly being used as a treatment for major bleeding in patients who are not taking anticoagulants. The aim of this systematic review and meta-analysis is to evaluate the effectiveness of PCC administration for the treatment of bleeding in patients not taking anticoagulants. Studies investigating the effectiveness of PCC to treat bleeding in adult patients and providing data on either mortality or blood loss were eligible. Data were pooled using Mantel-Haenszel random effects meta-analysis or inverse variance random effects meta-analysis. From 4668 identified studies, 17 observational studies were included. In all patient groups combined, PCC administration was not associated with mortality (odds ratio = 0.83; 95% confidence interval [CI], 0.66-1.06; P = .13; I² = 0%). However, in trauma patients, PCC administration, in addition to fresh frozen plasma, was associated with reduced mortality (odds ratio = 0.64; CI, 0.46-0.88; P = .007; I² = 0%). PCC administration was associated with a reduction in blood loss in cardiac surgery patients (mean difference: −384; CI, −640 to −128, P = .003, I² = 81%) and a decreased need for red blood cell transfusions when compared with standard care across a wide range of bleeding patients not taking anticoagulants (mean difference: −1.80; CI, −3.22 to −0.38; P = .01; I² = 92%). In conclusion, PCC administration was not associated with reduced mortality in the whole cohort but did reduce mortality in trauma patients. In bleeding patients, PCC reduced the need for red blood cell transfusions when compared with treatment strategies not involving PCC. In bleeding cardiac surgery patients, PCC administration reduced blood loss.

Keywords
bleeding, blood coagulation factors, hemostasis, meta-analysis, prothrombin complex concentrate, systematic review
1 | INTRODUCTION

Severe bleeding is a major health problem, occurring in different clinical settings such as in trauma or perioperatively. Excessive bleeding often results in coagulopathy, with incidences ranging from 1% to 45% depending on the severity of trauma or type of surgery, and develops through several mechanisms, such as the consumption of clotting factors. Coagulopathy may aggravate bleeding, increase transfusion requirements, and contribute to adverse outcomes.2

Plasma transfusion is frequently used to correct coagulopathy in bleeding patients because it is believed to replenish a deficit of coagulation factors. European guidelines on the management of bleeding trauma patients recommend transfusion of plasma to correct coagulopathy.6,7 In the operative setting, guidelines state that plasma can be used to treat coagulopathy.8,9 However, the effectiveness of plasma to correct coagulopathy is under debate.10-12 Because the ability of plasma to restore thrombin generation is limited.10 This limited ability may be due to the presence of anticoagulant proteins in plasma, which inhibit thrombin generation.10

An alternative approach for the correction of coagulopathy occurring during bleeding may be the use of factor concentrates, such as prothrombin complex concentrate (PCC). PCC contains either three or four of the vitamin K-dependent coagulation factors (II, IX, X, and sometimes VII). Also, depending on the type of PCC, small amounts of proteins C, S, Z, unfractionated heparin, or antithrombin are present. Because PCCs have gained approval by the US Food and Drug Administration in 2013, they have become the primary treatment for urgent reversal of oral anticoagulation with vitamin K antagonists and congenital vitamin K-dependent coagulation factor deficiencies.13

In addition to these established indications, PCCs are increasingly used to correct coagulopathy in bleeding patients unrelated to coagulopathy.14,15 Guidelines in both trauma and operative settings support the administration of PCC to bleeding patients to correct coagulopathy.7-9 However, this is largely based on expert opinion and is supported by limited evidence. By supplementing coagulation factors, PCCs rapidly correct coagulopathy while having lower risks of complications such as transfusion-associated lung injury, transfusion-associated circulatory overload, bacterial contamination, and allergic reactions when compared with plasma transfusion.16-18 Regarding safety, the risk of thromboembolic complications after administration of 4-factor PCC, when given in the proper dose, seem to be rather low.19

There are limited trials on the effectiveness of PCC. In trauma, a randomized controlled trial showed that PCC is associated with a reduced need for massive transfusion when compared with plasma.13 The trial was terminated early for safety reasons because the plasma group showed more adverse outcome compared with the PCC group. However, as all PCC patients also received fibrinogen, it is not clear from this trial whether PCC, fibrinogen, or the combination contributed most to the observed benefit. To date, a summary of the effectiveness of PCC in bleeding patients not taking anticoagulants is not present. The objective of this review is to systematically evaluate the effectiveness of PCC in bleeding unrelated to anticoagulation in different clinical scenarios.

2 | METHODS

2.1 | Search strategy

This systematic review and meta-analysis was conducted according to the PRISMA methodology.20 No review protocol is available. To identify all articles investigating the use of PCC for treatment of bleeding, a comprehensive computer-assisted literature search of MEDLINE, EMBASE, and CINAHL electronic databases was performed including articles between 1952 and April 2020. A broad search strategy was used that included multiple synonyms of PCC combined with terms as: “hemorrhage,” “mortality,” and “bleeding.” The full PubMed, EMBASE, and CINAHL search strategies can be found in the supplement (Appendix S1). In addition, the reference list of the most relevant studies and reviews were hand-searched for eligible studies not captured in the initial literature search.

2.2 | Study selection

The selection process was divided in three stages: title, abstract, and full-text selection. The title selection process was done by one author (D.B.). Thereafter, two authors (D.B., N.J.) independently performed the abstract and full-text selection to assess the eligibility of the articles to the predefined criteria. Differences in judgment were resolved by discussion. Inclusion criteria were: (a) patients ≥ 18 years old who received 3- or 4-factor PCC for active bleeding; (b) all patients in the experimental group received PCC; (c) at least one of the following outcomes was reported: mortality, blood product utilization, blood loss or thromboembolic (TE) events; (d) the use of a comparator including: placebo, nothing, usual treatment (eg, fresh frozen plasma [FFP]) or other hemostatic agents (eg, recombinant factor VIIa [rFVIIa]); and (e) language: English. Exclusion criteria were: experimental or preclinical study design; use of activated 4-factor PCC; PCC administration for anticoagulant reversal; use of a different PCC as comparator; and data of outcome could not be extracted. Types of articles eligible for inclusion were randomized controlled trials, nonrandomized controlled trials, and cohort studies. Case reports and congress abstracts were excluded.

2.3 | Data extraction

Using a predefined standard data form, data were collected independently by two authors (D.P., M.W.) for the following data points: study design; number of patients; mean age; study setting; PCC indication; timing of PCC administration; type and dose of PCC; type and dose of comparator; mortality; mortality follow-up days; thromboembolic (TE) events; number of transfused red blood
cell (RBC) units; blood loss (total chest tube output at either 12 or 24 hours postsurgery); other significant outcomes; article authors’ conclusion.

2.4 | Outcomes

All-cause mortality was chosen as the primary outcome. Secondary outcomes were: blood loss, RBC utilization, and TE events. Three separate subgroup analyses were performed, including cause of bleeding (trauma, cardiac surgery, liver surgery, other), comparator (PCC with FFP vs FFP, PCC vs FFP, PCC vs rFVIIa, PCC vs nothing), and PCC dosage (<20, 20-30, >30 IU/kg).

2.5 | Study quality assessment

Study quality assessment of all eligible studies was conducted by two authors (D.P., M.W.) independently. Using the Newcastle-Ottawa quality assessment scale, the risk of bias of cohort and case control studies was assessed. The maximum score of the Newcastle-Ottawa scale is 9 points. All studies were ranked either as poor, moderate, or good quality depending on the total score: studies receiving ≤ 5 points received a poor quality score; studies receiving 6-7 points received a moderate quality score; and studies receiving ≥ 7 points received a good quality score.

2.6 | Statistical analysis

Data are reported as mean with standard deviation, counts, or percentages. Data retrieved as median with interquartile range were converted to means as reported before to pool data for meta-analysis. Meta-analysis was performed using Review Manager 5.3; statistical heterogeneity across the studies was assessed using the Cochran’s Q test and \( I^2 \) values. Sensitivity analysis to exclude outliers was performed using Review Manager 5.3. The odds ratios (OR) were pooled using the Mantel-Haenszel procedure, which assumes a random effects model. Mean differences were pooled using the Inverse variance procedure, which also assumes a random effects model.

3 | RESULTS

3.1 | Characteristics of included studies

The systematic search identified 4668 studies (Medline: 1554; Embase: 2927; CINAHL: 187; Figure 1). After removing duplicates,
| Reference Number/Study ID | Design of Study/Study Duration | Setting | Sample Size | Intervention Group | Control Group | Conclusion |
|---------------------------|---------------------------------|---------|-------------|--------------------|---------------|------------|
| Zeeshan et al 2019<sup>30</sup> | Retrospective (propensity matched) | Trauma | 468 | 4-factor PCC + FFP | FFP | The use of PCC as an adjunct to FFP is associated with improved survival and reduction in transfusion requirements without increasing the risk of TE |
| Jehan et al 2018<sup>31</sup> | Retrospective (propensity matched) | Trauma | 120 | 4-factor PCC + FFP | FFP | PCC as a component therapy along with FFP is superior to FFP alone in treating coagulopathy |
| Joseph et al 2016<sup>33</sup> | Retrospective (propensity matched) | Trauma | 81 | 3-factor PCC + FFP | FFP | PCC reduced the time to correct INR and time to intervention compared with patients who received FFP |
| Joseph et al 2014<sup>32</sup> | Retrospective (propensity matched) | Trauma | 252 | 3-factor PCC + FFP | FFP | PCC as an adjunct to FFP is associated with reduction of blood product requirement and also lowers overall cost |
| DeLoughery et al 2016<sup>27</sup> | Retrospective | Bleeding | 78 | 4-factor PCC | rFVIIa | PCC was associated with had the shortest LOS among survivors, the rFVIIa group had the lowest mortality |
| Harris et al 2020<sup>20</sup> | Retrospective | Cardiac Surgery | 79 | 4-factor PCC | Nothing | Among Jehovah’s Witness patients undergoing cardiac surgery, 4-PCC was not associated with a difference in Hb change postoperatively, in the event of excessive blood loss 4pcc may provide a viable option |
| Biancari et al 2019<sup>24</sup> | Prospective multicenter trial (propensity matched) | Cardiac surgery | 202 | Both 3- and 4-factor PCC + FFP | FFP | The use of PCC compared with FFP may reduce the need of blood transfusion after CABG; these results should be considered hypothesis generating |
| Zweng et al 2018<sup>39</sup> | Retrospective (propensity matched) | Cardiac surgery | 160 | 3-factor PCC + FFP | FFP | PCC is not associated with an increased risk of TE or unfavorable outcomes compared with conventional treatment. PCC may be acceptable for management of severe perioperative bleeding in open heart surgery |
| Fitzgerald et al 2018<sup>28</sup> | Retrospective (propensity matched) | Cardiac surgery | 234 | 4-factor PCC + FFP | FFP | Use of PCCs as part of a multifaceted coagulation management strategy may have blood-sparing effects |
| Harper et al 2018<sup>29</sup> | Retrospective (propensity matched) | Cardiac surgery | 106 | 3-factor PCC | rFVIIa | Use of rFVIIa vs inactive PCCs was associated with renal failure requiring dialysis and increased postoperative bleeding and transfusions |
| Mehringer et al 2018<sup>35</sup> | Retrospective | Cardiac surgery | 129 | 4-factor PCC | rFVIIa | 4-factor PCC may be an equally efficacious alternative to rFVIIa for patients experiencing significant bleeding during cardiac surgery |

(Continues)
| Reference Number/Study ID | Design of Study/Study Duration | Setting | Sample Size | Intervention Group | Control Group | Conclusion |
|---------------------------|--------------------------------|---------|-------------|--------------------|---------------|------------|
| Cappabianca et al 2015\textsuperscript{23} | Prospective observational (propensity matched) | Cardiac surgery | 450 | 3-factor PCC | FFP | The use of PCC compared with FFP was associated with decreased postoperative blood loss and RBC transfusion requirements. However, PCC may be associated with a higher risk of AKI |
| Bradford et al 2015\textsuperscript{25} | Retrospective | Cardiac surgery | 68 | 3-factor PCC | Nothing | PCC in LVAD insertion does not appear to be associated with a significant increase in thromboembolic events |
| Ortmann et al 2014\textsuperscript{36} | Retrospective (propensity matched) | Cardiac surgery | 100 | 4-factor PCC | FFP 15 | PCC may be an alternative to FFP in patients who are coagulopathic and bleeding after cardiac surgery |
| Tanaka et al 2013\textsuperscript{37} | Retrospective | Cardiac surgery | 150 | 3-factor PCC | rFVIIa | 3-factor PCC could be hemostatically effective in dilutional coagulopathy because of its high prothrombin content despite the lower FVII content |
| Colavecchia et al 2017\textsuperscript{26} | Retrospective (propensity matched) | Liver surgery | 117 | 4-factor PCC + FFP | FFP | Use of PCC and fibrinogen concentrate during liver transplantation did not reduce intraoperative blood product requirements |
| Kirchner et al 2014\textsuperscript{34} | Retrospective | Liver surgery | 266 | 4-factor PCC + FFP | FFP | In liver transplantation, ROTEM-guided treatment with fibrinogen and/or PCC did not increase the occurrence of thrombosis and ischemic events |

Abbreviations: AKI, acute kidney injury; CABG, coronary artery bypass graft; Hb, hemoglobin; INR, International Normalized Ratio; LOS, length of stay; LVAD, left ventricular assist device; ROTEM, rotational thromboelastometry; TE, thromboembolic events.
3423 studies remained. Of these, 3297 were excluded based on exclusion criteria, yielding 126 studies. Of these, full text was assessed, leading to the exclusion of another 109 studies, resulting in 17 studies that were included in the analysis. Studies included two prospective studies and 15 retrospective studies, yielding a total of 3060 patients (Figure 1). Characteristics of the included studies are shown in Table 1. Studies were conducted in patients undergoing cardiothoracic surgery (10 studies with a total of 1678 patients), where a 3-factor PCC was evaluated in seven studies. PCC had an increase use of 2.1 RBC units in comparison to patients not receiving PCC. Of importance, all analyses showed high heterogeneity. In contrast, patients undergoing liver transplant surgery did not report any data on mortality. Subgroup analyses on the dose-dependent effectiveness of PCC on mortality is available in the supplement (Figure S5). There were no significant differences between lower and higher PCC dosages. A sensitivity analysis excluding outliers did not change results (OR 0.79; 95% CI, 0.62-1.01; P = .06; I² = 0%; P for heterogeneity = .78; Figure S7).

3.4 | Blood loss outcome

Data on blood loss were available in five studies with a total of 875 patients, all of which were performed in patients undergoing cardiac surgery. The timeframe in which blood loss was recorded was either 12 or 24 hours. Four studies reported blood loss in median values, which were converted into mean values. Mean blood loss ranged from 353 to 1159 mL in patients receiving PCC and 480 to 1644 mL in patients not receiving PCC. Total blood loss was significantly lower in the PCC group (mean difference −293 mL; 95% CI, −546 to −41; P = .02; I² = 86%; Figure 3).
not receiving PCC (95% CI, 1.2 to −2.8; P < .00001; I² = 0%; P for heterogeneity = .64; Figure 4). Subgroup analyses on the dose-dependent effectiveness of PCC on RBC utilization is available in the supplement (Figure S6). Dosages < 30 IU/kg showed effects in favor of PCC whereas dosages > 30 IU/kg showed no differences between groups.

### 3.6 | Thromboembolic events

Data on TE events was available in 15 of the 17 included studies with a total of 2745 evaluable patients. The TE rate ranged between 0% and 41% in the PCC group and between 4% and 26% in the no-PCC group. Overall, PCC administration was not
This systematic review analyzes the effectiveness of PCC to control bleeding in various patient populations who were no on anticoagulants. Overall, we found that PCC did not reduce mortality in comparison to plasma or to other hemostatic agents. However, in the subgroup analysis, PCC administration when added to FFP was associated with reduced mortality in trauma patients.

An explanation for the difference of an effect between patient groups could be that PCC could play a more important role in patients that are bleeding more excessively, which is more probable in trauma patients compared with the cardiac and liver surgery patients, which may experience more "controlled" blood loss.

Most of the included studies compared PCC in conjunction with FFP to solely FFP. Whereas PCC added to FFP was associated with a reduction in mortality, this effect was not noted in the other comparisons, such as PCC versus FFP or PCC versus no therapy. This implies that PCC is beneficial as an adjunct to standard treatment including FFP rather than a replacement of FFP. This benefit could lie in a more rapid correction of coagulopathy and coagulation factor replacement while not losing the benefit of FFP.

In cardiac surgery patients, a significantly reduced blood loss in favor of the PCC group was found. However, substantial statistical heterogeneity may hamper credibility of results. This heterogeneity is explained by two reasons. First, blood loss was recorded in different time frames (either within 12 or 24 hours postsurgery), yielding a wide range. Second, we extrapolated means from medians in four studies. However, results on amount of blood loss are congruent with the finding of a reduction in RBC utilization in the PCC-treated patients when compared with the patients not receiving PCC. In most patient categories, results point in the same direction, suggesting that PCC is associated with reduced blood loss.

Of note, liver surgery patients showed opposite results, with more RBC products needed in patients treated with PCC versus a comparator. This may be due to baseline imbalances between groups, with more severe liver injury and coagulopathy in the PCC group. Also, two studies with small sample sizes were included. Therefore, we feel that results in the subgroup of liver failure patients are uncertain.

Regarding safety, the use of PCC did not result in an increase in TE events when compared with patients not receiving PCC. Of note, the CIs of the ORs were very wide, suggesting differential results across studies. However, the large number of patients included in this analysis is a strength. Taken together, PCCs do not seem to come at a cost of increased venous thromboembolic events.

This review has certain limitations because a small number of eligible studies were available with substantial heterogeneity. This heterogeneity is caused by a number of reasons. First, most

| Study or subgroup | Mean difference IV, Random, 95% CI | Year | Mean (95% CI) | Study or subgroup | Mean difference IV, Random, 95% CI | Year | Mean (95% CI) |
|-------------------|----------------------------------|------|--------------|-------------------|----------------------------------|------|--------------|
| 1.1.1 Trauma      |                                  |      |              |                   |                                  |      |              |
| Joseph 2014       | 6.6                              | 4.1  | 63           | 10                | 8.3                             | 189  | 9.2%         |
| Joseph 2016       | 3.2                              | 1.9  | 27           | 5.4               | 4.1                             | 54   | 9.6%         |
| Zeehan 2019       | 6                                | 4    | 234          | 10                | 4                              | 234  | 10.2%        |
| Subtotal          | 364                              |      | 384          |                     | 357                             |      | 38.4%        |
| Heterogeneity:    | Tau² = 0.80; Chi² = 9.45, df = 3 | P < 0.0001; I² = 68% |                     | Test for overall effect: Z = 5.44 (P < 0.0001) |

**FIGURE 4** Forest plot comparison of overall and subgroup RBC utilization in patients treated with PCC vs patients not treated with PCC associated with the occurrence of TE (OR 1.11; 95% CI, 0.82-1.50; P = .49; I² = 0%; Figure 5). In subgroup analyses, similar results were found (Figure 5).

4 | DISCUSSION

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This systematic review analyzes the effectiveness of PCC to control bleeding in various patient populations who were no on anticoagulants. Overall, we found that PCC did not reduce mortality in comparison to plasma or to other hemostatic agents. However, in the subgroup analysis, PCC administration when added to FFP was associated with reduced mortality in trauma patients.

An explanation for the difference of an effect between patient groups could be that PCC could play a more important role in patients that are bleeding more excessively, which is more probable in trauma patients compared with the cardiac and liver surgery patients, which may experience more "controlled" blood loss.

Most of the included studies compared PCC in conjunction with FFP to solely FFP. Whereas PCC added to FFP was associated with a reduction in mortality, this effect was not noted in the other comparisons, such as PCC versus FFP or PCC versus no therapy. This implies that PCC is beneficial as an adjunct to standard treatment including FFP rather than a replacement of FFP. This benefit could lie in a more rapid correction of coagulopathy and coagulation factor replacement while not losing the benefit of FFP.

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Of note, liver surgery patients showed opposite results, with more RBC products needed in patients treated with PCC versus a comparator. This may be due to baseline imbalances between groups, with more severe liver injury and coagulopathy in the PCC group. Also, two studies with small sample sizes were included. Therefore, we feel that results in the subgroup of liver failure patients are uncertain.

Regarding safety, the use of PCC did not result in an increase in TE events when compared with patients not receiving PCC. Of note, the CIs of the ORs were very wide, suggesting differential results across studies. However, the large number of patients included in this analysis is a strength. Taken together, PCCs do not seem to come at a cost of increased venous thromboembolic events.

This review has certain limitations because a small number of eligible studies were available with substantial heterogeneity. This heterogeneity is caused by a number of reasons. First, most
studies had a retrospective study design, in which PCC was administered based on clinical judgment. This may introduce confounding and bias because PCC may potentially be administered to more severely ill and hemodynamically unstable patients, resulting in underrepresentation of the actual effect. Second, because there is no optimal dose for PCC, there is considerable variety in the type and dose of PCC, as well as the trigger for administration. This could lead to both an under- or overrepresentation of the actual effects as certain PCCs might have better effectiveness and safety profiles compared with others. Third, not all included studies had strict inclusion or exclusion criteria of their study population, leading to a heterogeneous study population. Furthermore, we only included articles published in English, which might lead to language bias.

Of interest, our search also yielded multiple studies, including a randomized controlled trial,\textsuperscript{40} that used a point-of-care coagulation testing-based protocol in which both PCC and fibrinogen concentrate as were administered as an adjunct to FFP in comparison to a FFP-based protocol.\textsuperscript{15,40-43} Unfortunately, these studies were not included because outcome data from patients receiving PCC could not be separated from those not receiving PCC. Innerhofer et al found that patients receiving PCC and fibrinogen had a decreased need for massive transfusion and rescue therapy when compared with the FFP group.\textsuperscript{40} Schöchl et al and Nienaber et al also showed reduced exposure of trauma patients to blood products in the PCC/fibrinogen group when compared with the FFP group.\textsuperscript{41,43} Görlinger et al also showed reduced blood product utilization after implementing the PCC/fibrinogen based protocol.

| Study or Subgroup | PCC | no PCC | Weight | Odds Ratio |
|------------------|-----|--------|--------|------------|
| **1.4.1 Trauma** |     |        |        |            |
| Joseph 2014      | 2   | 63     | 3      | 2.8%       |
| Joseph 2016      | 4   | 27     | 5      | 4.6%       |
| Jehan 2018       | 1   | 40     | 2      | 1.5%       |
| Zeeshan 2019     | 11  | 234    | 17     | 14.9%      |
| **Subtotal (95% CI)** | 364 | 557     | 23.8% | 0.90 [0.49, 1.67] |
| Total events     | 18  |        |        |            |

Heterogeneity: Tau\textsuperscript{2} = 0.00; Chi\textsuperscript{2} = 2.38, df = 3 (P < 0.50); I\textsuperscript{2} = 0%
Test for overall effect: Z = 0.33 (P = 0.74)

| **1.4.2 Cardiac surgery** |     |        |        |            |
| Ortmann 2014            | 1   | 45     | 1      | 1.2%       |
| Bradford 2015           | 12  | 29     | 6      | 6.6%       |
| Cappabianca 2016        | 26  | 225    | 22     | 25.2%      |
| Mehringer 2018          | 2   | 56     | 4      | 3.0%       |
| Zweng 2018              | 8   | 80     | 4      | 5.9%       |
| Fitzgerald 2018         | 8   | 117    | 9      | 9.3%       |
| Harper 2018             | 8   | 53     | 14     | 9.7%       |
| Biancari 2019           | 5   | 101    | 3      | 4.3%       |
| Harris 2020             | 2   | 19     | 2      | 2.2%       |
| **Subtotal (95% CI)**   | 725 | 791    | 67.5% | 1.19 [0.82, 1.70] |
| Total events            | 72  |        |        |            |

Heterogeneity: Tau\textsuperscript{2} = 0.00; Chi\textsuperscript{2} = 7.57, df = 8 (P = 0.48); I\textsuperscript{2} = 0%
Test for overall effect: Z = 0.84 (P = 0.38)

| **1.4.3 Liver surgery** |     |        |        |            |
| Kirchner 2014           | 11  | 156    | 5      | 7.7%       |
| **Subtotal (95% CI)**   | 156 | 110    | 7.7%  | 1.59 [0.54, 4.72] |
| Total events            | 11  |        |        |            |

Heterogeneity: Not applicable
Test for overall effect: Z = 0.84 (P = 0.40)

| **1.4.4 Bleeding various reasons** |     |        |        |            |
| Dellaoughery 2016       | 0   | 16     | 3      | 1.0%       |
| **Subtotal (95% CI)**   | 16  | 26     | 1.0%  | 0.20 [0.01, 4.21] |
| Total events            | 0   |        |        |            |

Heterogeneity: Not applicable
Test for overall effect: Z = 1.03 (P = 0.30)

| Total (95% CI)           | 1261 | 1484 | 100.0% | 1.11 [0.82, 1.50] |
| Total events             | 101  | 100  |        |            |

Heterogeneity: Tau\textsuperscript{2} = 0.00; Chi\textsuperscript{2} = 12.12, df = 14 (P= 0.60); I\textsuperscript{2} = 0%
Test for overall effect: Z = 0.68 (P = 0.49)
Test for subgroup differences: Chi\textsuperscript{2} = 2.16; df = 3 (P= 0.54); I\textsuperscript{2} = 0%

**FIGURE 5** Forest plot comparison of overall and subgroup thromboembolic events in patients treated with PCC vs patients not treated with PCC
in cardiac surgery patients. Inclusion of these studies probably would have strengthened the suggestion that a PCC and/or fibrinogen-based approach may be a reasonable strategy to treat bleeding patients.

5 | CONCLUSION

PCC administration in bleeding patients not using anticoagulants had no effect on mortality in the whole cohort of patients. However, in trauma patients, a resuscitation strategy using both PCC and FFP transfusion was associated with reduced mortality when compared to a resuscitation strategy involving solely FFP. Also, PCC reduced the need for RBC transfusions when compared with treatment strategies not involving PCC. In bleeding cardiac surgery patients, PCC administration reduced perioperative blood loss. Risk of TE events was not increased. However, results are subject to considerable heterogeneity and should be interpreted with caution. These data, derived from observational studies, can be used to design trials to further explore the effectiveness of PCC in different clinical scenarios of bleeding.

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Not applicable.

CONFLICT OF INTEREST

None.

AUTHOR CONTRIBUTIONS

Daan P. van den Brink contributed substantially to the conception, design, acquisition, analysis, and interpretation of data for the work; and drafted and revised the work. Mathijs R Wirtz contributed substantially to the acquisition, analysis, interpretation of data, and revising the intellectual content. A. Serpa Neto made substantial contributions to the interpretation of data and revising the intellectual content. Herbert Schöchl made substantial contributions to the interpretation of data and revising the intellectual content. Victor Viersen made substantial contributions to the interpretation of data and revising the intellectual content. J Binnekade made substantial contributions to the statistical analyses of the data for the work. Nicole P. Juffermans contributed substantially to the conception, design, acquisition, analysis, and interpretation of data for the work; and drafted and revised the work.

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**SUPPORTING INFORMATION**

Additional supporting information may be found online in the Supporting Information section.

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