Anticoagulant therapy for acute venous thrombo-embolism in cancer patients: A systematic review and network meta-analysis

Anne Rossel1,2, Helia Robert-Ebadi3, Christophe Combescure4, Olivier Grosgeurin1,2, Jérôme Stirnemann1,2, Alfredo Addeo4,5, Nicolas Garin1, Thomas Agoritsas1, Jean-Luc Reny1,2, Christophe Marti1,2*

1 Division of General Internal Medicine, University Hospitals of Geneva, 4, rue Gabrielle Perret-Gentil, CH, Geneva, Switzerland, 2 Faculty of Medicine, University of Geneva, Geneva, Switzerland, 3 Division of Angiology and Haemostasis, University Hospitals of Geneva, 4, rue Gabrielle Perret-Gentil, CH, Geneva, Switzerland, 4 Division of clinical Epidemiology, University Hospitals of Geneva, 4, rue Gabrielle Perret-Gentil, CH, Geneva, Switzerland, 5 Division of Oncology, University Hospitals of Geneva, 4, rue Gabrielle Perret-Gentil, CH, Geneva, Switzerland

* christophe.marti@hcuge.ch

Abstract

Background

Low-molecular-weight heparin (LMWH) is usually recommended for the treatment of cancer-associated thrombosis (CAT) but this treatment requires burdensome daily injections. We did a systematic review to compare the efficacy and safety of direct oral anticoagulants (DOAC), vitamin K antagonists (VKA) and LMWH in patients with CAT.

Methods

We searched Pubmed, Embase and CENTRAL for randomised controlled trials comparing DOAC, VKA and LMWH in patients with CAT. Pairwise and network meta-analyses were computed for venous thromboembolism (VTE) recurrence and bleeding complications.

Results

We identified 14 studies, including 4,661 patients. In pairwise comparison, DOAC were superior to LMWH to prevent VTE recurrence (HR 0.63; 95% CI 0.42–0.96) and LMWH was superior to VKA (HR 0.53; 95% CI 0.40–0.70). The rate of major bleeding was higher with DOAC compared to LMWH (HR 1.78; 95% CI 1.11–2.87). In the network meta-analysis, DOAC had a lower, but non-significant, rate of VTE recurrence compared to LMWH (HR 0.74; 95% CI 0.54–1.01). Both DOAC (HR 0.42; 95% CI 0.29–0.61) and LMWH (HR 0.57; 95% CI 0.44–0.75) were associated with lower rates of recurrence compared to VKA. No significant difference in major bleeding rate was observed in the network meta-analysis. Inconsistency was observed between pairwise and network meta-analysis comparisons for major bleeding.
Introduction

The management of cancer-associated thrombosis (CAT) is challenging. The risk of developing a first venous thromboembolic event (VTE) in cancer patients is seven-fold higher than in the general population [1]. The risk of recurrence after a first episode of VTE is also particularly high in cancer patients after cessation of anticoagulant therapy, reaching a 12-month cumulative incidence of 20% [2]. Prolonged anticoagulant therapy is thus recommended for patients with CAT [3]. However, cancer patients are also at higher risk of bleeding due to cancer itself or to cancer-related interventions such as surgery or chemotherapy [4]. Currently, the standard of care of VTE in patients with cancer consists of subcutaneous low molecular weight heparin (LMWH), for an initial duration of 6 months, which is extended with either LMWH or VKA for an indefinite duration, as long as the cancer is not considered in remission [3]. This recommendation is based on randomised studies showing a reduced risk of VTE recurrence in cancer patients receiving LMWH compared with antivitamin K therapy [5–8]. This benefit was shown to be consistent in several meta-analyses [9–11]. However, LMWH treatment is burdensome, both from an economic and patient perspective as LMWH is expensive and requires daily injections which may further alter quality of life of cancer patients.

Due to the limited tolerance to daily injections in some patients, and to the lack of evidence to recommend LMWH beyond the initial 6 months, VKA are sometimes used in cancer patients [12]. However, the efficacy and safety of VKA may be altered by the difficulties to keep patients with cancer in the therapeutic range. Indeed, VKA therapeutic range is narrow, and VKA are subject to pharmacokinetic and pharmacodynamic interactions which are more numerous in cancer patients than in the general population. Moreover, anorexia or vomiting during chemotherapy can impair regular Vitamin K intake and absorption [13]. The low time in therapeutic range (TTR) obtained even in the setting of clinical trials is an objective reflection of this challenging issue.

DOAC have recently emerged as an alternative to VKA and LMWH for the treatment of VTE in non-cancer patients. Large-scale phase III non-inferiority trials confirmed the efficacy of these molecules compared to VKA to prevent VTE recurrence, with a similar or even more favourable safety profile in terms of bleeding events [14–18]. Previous meta-analyses [10, 11, 19, 20] based on subgroup analyses of these phase III trials concluded that DOAC are effective and safe for the treatment of CAT. However, two main limitations preclude any firm conclusions. First, only a small fraction of the studied population had cancer and they were highly selected and not representative of the general oncologic population. This is well reflected by the surprisingly low VTE recurrence rate for cancer patients, compared to the known high recurrence rate reported in observational studies. Second, these studies compared DOAC with a regimen of 5 days of LMWH followed by VKA, which is not the recommended treatment for CAT.

Recently, two randomised controlled trials (RCT) comparing DOAC to LMWH in cancer patients have been published [21, 22].
In the Hokusai cancer trial [21], edoxaban was shown non-inferior to dalteparin to prevent a composite endpoint including VTE recurrence or major bleeding. Similarly, the SELECT-D trial [22] compared Oral Factor Xa Inhibitor Rivaroxaban to dalteparin. The rate of VTE recurrence was lower in patients allocated to Rivaroxaban, but the rate of clinically relevant non major bleeding (CRNMB) was increased.

In order to further assess the evidence on the efficacy and safety of DOAC for the treatment of VTE in cancer patients, we performed a systematic review and network meta-analysis to compare VKA, LMWH and DOAC. In the presence of a limited number of studies directly comparing DOAC to the recommended standard of care (LMWH), network meta-analysis allows to provide direct and indirect comparison between DOAC and LMWH and evaluate their consistency.

Materials and methods

Protocol and registration

Search strategy, study selection, data extraction and analysis were performed according to a pre-defined protocol (available on request) and reported according to the PRISMA guidelines. (S1 and S2 Tables)

Eligibility criteria

We included RCTs comparing the following anticoagulant regimens in adult patients with CAT: LMWH vs VKA, DOAC vs VKA, DOAC vs LMWH. Studies were eligible if patients with cancer were included, even as a subgroup, if results were reported separately. Index VTE events were defined by the presence of an objectively confirmed acute deep vein thrombosis (DVT) and/or pulmonary embolism (PE). DVT had to be diagnosed either by lower limb venous compression ultrasound (CUS) or venography. PE had to be diagnosed either by computed tomography pulmonary angiography, high-probability pulmonary ventilation/perfusion scan, pulmonary angiography, or the presence of a proximal DVT on CUS in a patient with clinically suspected PE.

Thrombotic and haemorrhagic outcomes had to be objectively confirmed and adjudicated, using the same criteria as above for recurrent VTE and the widely accepted ISTH criteria for major bleeding and clinically relevant non-major bleeding.

Studies comparing various regimens of the same anticoagulant, or other classes of anticoagulant therapy such as pentasacharides or unfractionated heparin (UFH), or with a follow-up duration less than 3 months were excluded. As the definition of active cancer may vary in each study, we used the definition applied in each study. When various definitions were reported in the same study, we extracted data from patients with cancer defined as active rather than patients with cancer history or incidental cancers.

Information sources

Two authors (AR, CM) systematically searched Pubmed, Embase and the Cochrane register of controlled Trials (CENTRAL) databases using a comprehensive search strategy without language restriction. The detailed search strategy is available in the supplementary appendix (S2 Table). To ensure a comprehensive literature search, we also examined reference lists from retrieved articles and reference literature (guidelines and systematic reviews).

Study selection and data extraction

Two investigators (AR, CM) independently evaluated studies for possible inclusion. Non-relevant studies were excluded based on title and abstract. For potentially relevant studies, full-text
was obtained and two investigators (AR, CM) independently assessed study eligibility and extracted the data on study design, patient characteristics and outcomes. Disagreement about study inclusion or data extraction was resolved by consensus or by discussion with a third author (HRE).

Outcomes
The main outcome was the rate of objectively confirmed new VTE event. VTE events had to be diagnosed as described in the eligibility criteria.

Safety outcomes were the rate of major bleeding, clinically relevant non-major bleedings (CRNMB), all-cause mortality and gastro-intestinal (GI) bleeding.

According to the International Society of Thrombosis and Haemostasis, major bleeding is defined as a decrease in haemoglobin level of ≥2g/dl, transfusion of ≥2 units of packed red blood cells, bleeding that occurs in a critical site (intracranial, intra-spinal, intraocular, pericardial, intra-articular, intramuscular with compartment syndrome, or retroperitoneal) or fatal bleeding [23]. CRNMB is defined as overt bleeding that does not meet the criteria for major bleeding but is associated with medical intervention, unscheduled contact with a physician, interruption or discontinuation of study drug, or discomfort or impairment of activities of daily living [24].

The longest follow-up duration was used. According to inclusion criteria, minimum follow-up was 3 months.

Risk of bias
We used the quality criteria of the Cochrane Collaboration’s tool [25] to assess risk of bias in the following domains: selection bias (random sequence generation, allocation concealment), performance bias (blinding of participants and personnel), detection bias (blinding of outcome assessment), attrition bias (incomplete outcome data), reporting bias (selective reporting), and other biases.

Two investigators assessed study quality independently. Disagreements were resolved by consensus or by a third reviewer if no consensus was found.

Data analysis
For all outcomes, pooled risks were obtained for the different treatment arms using random effect models. Hazard ratios (HR) for comparison between treatment arms were pooled across studies using network meta-analyses with random effects [26]. Estimates were obtained by using a frequentist method with the package netmeta for R program version 3.5.0 [27]. In addition, to further assess the inconsistency of the network, the direct evidence was obtained for each pair of compared treatments by combining only studies directly comparing the two treatments. For this purpose, a model with random effects (DerSimonian and Laird approach) was used.

When HRs were not reported in original publications, they were estimated by the ratio of logarithms of event-free proportions calculated from the reported sample sizes and the number of events [28]. In order to assess the reliability of this approximation method, we computed estimates also in studies providing HRs and compared the calculated results to the actual reported HRs.

To facilitate the interpretation of pooled HRs, the effect sizes were expressed as absolute one-year risk difference (RD) between treatment arms. For this purpose, we used the formula of the ratio of logarithms of event-free proportions with the risks in LMWH arm of the
Hokusai-cancer VTE study as baseline risks as this study was the largest and most recent RCT dedicated to VTE treatment in patients with cancer [21].

Heterogeneity was detected using the test based on Cochran’s Q statistic with a 10% level of statistical significance (P < 0.1) and I² > 50%. In the network meta-analysis, the generalized Cochran’s Q statistic was used to test heterogeneity and inconsistency with a 10% level of statistical significance due to the lack of power of this type of tests [29]. A sensitivity analysis was performed to evaluate the strength of the pooled HRs, by the leave-one out principle.

The two-sided significance level was 0.05 for all analyses (except for the heterogeneity and inconsistency tests). In order to determinate treatment rankings for efficacy and safety outcomes, P-scores were derived from the network meta-analysis. The P-score can be interpreted as the mean extent of certainty that treatment is better than another treatment, averaged over all competing treatments and can be seen as the frequentist equivalent of surface under the cumulative ranking (SUCRA) value [30].

Results

Study selection and characteristics
The search retrieved 5054 references, among which 756 were duplicates (Fig 1). Fourteen RCTs including 4,661 patients were included in the meta-analysis. Seven studies [5–8, 31–33] compared LMWH to VKA (2,095 patients), 5 studies compared DOAC to VKA [34–38] (1,114 patients) and the 2 remaining [21, 22] (1,452 patients) compared DOAC to LMWH. Four studies comparing DOAC to VKA, consisted of subgroup analyses of patients with cancer, included in studies for VTE treatment in a general population. Characteristics of included studies are provided in Table 1. The diagram of the network meta-analysis is provided in Fig 2.

Venous thromboembolism recurrence
Fourteen studies including 4,661 patients reported VTE recurrence (Table 1) [5–8, 21, 22, 31–38]. Recurrence occurred in 4.1% (66/1310) of patients allocated to DOAC, 8.3% (151/1,797) allocated to LMWH and in 9.5% (156/1554) of those allocated to VKA. (S3 Table)

In direct comparison (Table 2), treatment by DOAC was associated with a decreased rate of recurrence compared to LMWH (HR 0.63; 95% CI 0.42–0.96, RD -3.8; 95% CI -6.0 to -0.9: Table 2, Fig 3 and S4 Table). There was no significant difference between DOAC and VKA (HR 0.62; 95% CI 0.34 to 1.15). The rate of recurrence was lower in patients treated by LMWH compared to patients treated by VKA (HR 0.53; 95% CI 0.40–0.70, RD -8.9; 95% CI -14.6 to -4.4. In the network meta-analysis patients treated by DOAC had a lower, though no longer statistically significant, rate of VTE recurrence compared to LMWH (HR 0.74; 95% CI 0.54–1.01). Both DOAC (HR 0.42; 95% CI 0.29–0.61) and LMWH (HR 0.57; 95% CI 0.44–0.75) were associated with a lower rate of recurrence compared to VKA. (Table 2)

Major bleeding
Twelve studies including 4546 patients reported major bleeding [5–8, 21, 22, 31, 33–37].

The pooled risk of major bleeding was 4.4% (63/1283) among patients receiving DOAC, 4.7% (76/1769) among patients treated with LMWH and 4.9% (63/1494) among patients treated with VKA. (S3 Table).

Direct comparisons showed a higher rate of major bleeding in patients treated by DOAC compared to LMWH (HR 1.78, 95% CI 1.11 to 2.87, RD 3.0; 95% CI 0.4 to 7.1). (Fig 4 and S4 Table). There was no significant difference in the rate of major bleeding between DOAC and VKA (HR 0.76, 95% CI 0.38 to 1.53) or between LMWH and VKA (HR 0.96, 95% CI 0.57 to
1.63; Fig 4). The network meta-analysis showed no significant difference in major bleeding between treatment groups (Table 2).

**Clinically relevant non-major bleeding**

CRNMB was reported in 11 studies including 4400 patients [5, 6, 8, 21, 22, 31, 33–37]. The pooled risk of CRNMB was 12.6% (161/1283) among patients receiving DOAC, 13.8% (203/1698) among those receiving LMWH and 16.5% (217/1419) among those treated with VKA.
Table 1. Characteristics of included studies.

| Author (Study name) year | Study drug | Dose | Comparator | Dose | Treatment duration | TTR (%) | Included patients | Recurrence study drug vs comparator (%) | Major Bleeding study drug vs comparator (%) | CRNMB study drug vs Comparator (%) | Death study drug vs Comparator (%) |
|--------------------------|------------|------|------------|------|-------------------|---------|-----------------|------------------------------------------|---------------------------------------------|----------------------------------------|----------------------------------------|
| Raskob (Hokusai-VTE Cancer) 2018 [21] | LMWH 5–7 days, then Edoxaban | 60mg or 30mg/d | Dalteparin | 200U/kg/d during 30days, then 150U/kg/d | Min. 6months | NA | 1046 | 7.9 vs 11.3 | 6.9 vs 4.0 | 14.6 vs 11.1 | 39.5 vs 36.6 |
| Young (Select-D) 2018[22] | Rivaroxaban | 15mg bd 3 weeks, then 20mg/d | Dalteparin | 200U/kg/d during 30days, then 150U/kg/d | Min. 6months | NA | 406 | 3.9 vs 8.9 | 5.4 vs 3.0 | 12.3 vs 3.4 | 23.6 vs 27.6 |
| Agnelli (Amplify) 2015[34] | Apixaban | 10mg bd 7d, then 5mg bd | Enoxaparin-Warfarin* | 1mg/kg bd min 5d, then for INR 2–3 | 6months | NA | 159 | 3.7 vs 6.4 | 2.3 vs 5.0 | 10.3 vs 17.5 | 0.0 vs 7.7 |
| Mazilu 2014[38] | Dabigatran | 150 mg bd | LMWH-Acenocoumarol* | Target INR 2–3 | 6months | NA | 46 | 0 vs 0 | NA | NA | 0 vs 4 |
| Prins (EINSTEIN-DVT/PE) 2014[35] | Rivaroxaban | 15mg bd 3 weeks, then 20mg/d | Enoxaparin-Warfarin* | 1mg/kg bd min 5d, then for INR 2–3 | 3, 6 or 12 months | 57% | 462 | 2.3 vs 3.9 | 1.9 vs 3.9 | 11.6 vs 13.2 | 14.7 vs 11.8 |
| Raskob (Hokusai-VTE) 2016 [36] | Edoxaban | 60mg/d or 30mg/d | LMWH-Warfarin* | Target INR 2–3 | Min. 3 months | 63% | 208 | 3.6 vs 7.0 | 4.6 vs 3.0 | 14.7 vs 23.2 | 28.4 vs 26.3 |
| Schulman (RE-COVER) 2015 [37] | Dabigatran | 150mg bd | LMWH-Warfarin* | Target INR 2–3 | 6 months | 48% | 221 | 3.5 vs 4.7 | 3.8 vs 3.0 | 9.5 vs 9.0 | 14.0 vs 15.0 |
| Deitcher (ONCENOX) 2006 [31] | Enoxaparin | 1mg/kg bd 5d then 1 or 1.5mg/kg/d | LMWH-Warfarin* | Target INR 2–3 | 6 months | NA | 91 | 6.6 vs 10 | 9.0 vs 2.9 | 58.2 vs 50.0 | 32.8 vs 32 |
| Hull (LITE) 2006 [6] | Tinzaparin | 175U/kg/d | LMWH-Warfarin* | Target INR 2–3 | 3 months | NA | 200 | 7.0 vs 16.0 | 7.0 vs 7.0 | 20.0 vs 17.0 | 47.0 vs 47.0 |
| Lee (CLOT) 2003 [5] | Dalteparin | 200U/kg/d | LMWH-VKA* | Target INR 2–3 | 6 months | 46% | 673 | 8.0 vs 15.8 | 5.6 vs 3.6 | 14.0 vs 19.0 | 38.7 vs 40.5 |
| Lee (CATCH) 2015 [8] | Tinzaparin | 175U/kg/d | LMWH-Warfarin* | Target INR 2–3 | 6 months | 47% | 900 | 6.9 vs 10.0 | 2.7 vs 2.4 | 10.9 vs 15.3 | 33.4 vs 30.6 |
| Lopéz-Beret 2001 [33] | Nadroparin | 1.025U/10kg bd | LMWH-Acenocoumarol* | Target INR 2–3 | 6 months | NA | 35 | 2.5 vs 9.0 | 0 vs 5.2 | 4.9 vs 0 | 11.1 vs 7.8 |
| Meyer (CANTHANOX) 2002[7] | Enoxaparin | 1mg/kg once daily | LMWH-Warfarin* | Target INR 2–3 | 3 months | 41% | 146 | 3 vs 4 | 6.7 vs 16 | NA | 11.3 vs 22.7 |
| Romero 2009[32] | Tinzaparin | 175U/kg once daily | LMWH-Acenocoumarol* | Target INR 2–3 | 6 months | NA | 69 | 6 vs 21 | NA | NA | NA |

VKA: Vitamin K antagonists, d: day, bd: bi-daily, TTR: Time in the therapeutic range
* LMWH for min 5 days overlapped and followed by VKA
NA Not applicable.

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Gastro-intestinal bleeding

GI bleeding was reported in 2 studies comparing DOAC to LMWH (1,452 patients) [21, 22] and in 2 studies comparing VKA to LMWH (181 patients) [32, 33]. The pooled risk of GI bleeding was 5.7% (37/725) among patients receiving DOAC, 2.5% (16/815) among patients receiving LMWH, and 8.7% (8/93) among those receiving VKA (S3 Table). In direct comparison, the rate of GI bleeding was increased in patients treated with DOAC compared to LMWH (HR 2.88, 95%CI 1.53–5.44, RD 8.7, 95%CI 2.5 to 19.3) and was similar in patients treated with LMWH or VKA (HR 0.42; 95% CI 0.12 to 1.48). The network meta-analysis showed no significant difference of GI bleeding when DOAC were compared to VKA (HR 1.22, 95%CI 0.30 to 4.96) or LMWH to VKA (HR 0.42, 95% CI 0.12 to 1.48) (Table 2 and S5 Table).

Mortality

Twelve studies including 4610 patients reported mortality [5–8, 21, 22, 31, 33–37]. The pooled risk of death was 18.2% (344/1816) among patients receiving DOAC, 33.8% (612/1767) among patients receiving LMWH and 24.9% (439/1527) among patients receiving VKA (S3 Table). There were no significant differences in mortality between treatments in direct comparison, as in the network meta-analysis (Table 2 and S5 Table).
Study quality and risk of bias

The method of randomisation, allocation concealment and reporting were considered adequate in most studies. Participants were not blinded in most studies [5–8, 21, 22, 31–33, 35]. The risk of bias of individual studies is provided on S6 Table.

Sources of heterogeneity and sensitivity analyses

A low level of heterogeneity (I² < 25%) was observed in direct comparisons for VTE recurrence, major bleeding and GI bleeding. Moderate heterogeneity (I² < 50%) was observed among studies comparing DOAC to LMWH for mortality and high heterogeneity (I² > 50%) was observed among studies comparing LMWH to DOAC or VKA for CRNMB (Table 2). Exploration of heterogeneity, publication bias and sensitivity analysis were hindered by the small number of studies in the different comparison arms. The leave-one out meta-analysis did not significantly alter comparison estimates. There were some differences in treatment estimates between direct and indirect evidences (Table 2 and S7 Table) although p-values for comparison were not statistically significant. The proportion of direct evidence in the network
analysis varied from 36 to 100%. P-value for inconsistency was statistically significant for major bleeding (S8 Table).

**Treatment ranking**

According to the network meta-analysis, DOAC were associated with the highest certainty to be superior to other treatments for VTE recurrence (P-score 0.98) followed by LMWH (P-score 0.51) and VKA (P-score 0.01). LMWH was associated with the highest certainty to be superior to other treatments for safety outcomes, followed by VKA and DOAC (Table 3).

**Discussion**

This meta-analysis suggests that DOAC are an efficacious alternative to LMWH to prevent VTE recurrence in cancer patients. In the present study, DOAC were associated with a lower rate of VTE recurrence compared to LMWH in direct comparison studies.
In the network meta-analysis, treatment effect estimates were also in favour of DOAC and the upper limit of the confidence interval was close to statistical significance. Additionally, the present study confirms the superiority of LMWH over VKA to prevent VTE recurrence. These results are in accordance with previous meta-analyses and observational studies comparing anticoagulant strategies in cancer patients. [9, 39–41].

Table 3. Ranking of treatments according to network meta-analysis.

| P-score for ranking of treatments | DOACs | LMWH | VKA |
|----------------------------------|-------|------|-----|
| Recurrence                       | 0.986 | 0.515| 0   |
| Major bleeding                   | 0.214 | 0.857| 0.429|
| CRNMB                            | 0.261 | 0.950| 0.290|
| GI bleeding                      | 0.196 | 0.955| 0.349|
| Mortality                        | 0.178 | 0.839| 0.482|

Fig 4. Direct meta-analysis major bleeding forest plot.
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While DOAC were shown superior to other anticoagulant strategies to prevent VTE recurrence, we also found an increased rate of major bleeding in patients receiving DOAC compared with those receiving LMWH. Interestingly, estimates from direct and network comparisons were inconsistent as this increased rate of bleeding was observed only in the direct comparison analysis. Similarly, previous meta-analyses reported contrasting conclusions according to their design: direct comparison meta-analyses \[9, 40\] concluded to an increased rate of major bleeding in patients allocated to DOAC compared to LMWH, while network meta-analyses did not \[39, 42\].

We believe that network meta-analyses including indirect evidence from studies comparing DOAC to VKA possibly failed to demonstrate this increased rate of major bleeding because these studies included patients with less active or advanced cancer and thus at lower bleeding risk. The rate of major bleeding in the DOAC arm was two to three times lower in studies comparing DOAC to VKA than in studies comparing DOAC to LMWH (S3 Table). Including indirect evidence from studies comparing DOAC to VKA in the network meta-analysis underestimates the overall rate of bleeding for DOAC compared to LMWH. This may artificially favour DOAC in the indirect comparison with LMWH. Therefore, we consider that estimates from direct comparisons studies dedicated to cancer patients should be considered as more appropriate and applicable.

The increased rate of major bleeding in patients allocated to DOAC in studies comparing DOAC to LMWH was mainly due to the higher rate of upper gastro-intestinal bleeding. These events occurred mainly among patients who entered the trials with GI cancer. A safety analysis of the SELECT-D trial after enrolment of 220 patients showed an increased rate of bleeding among patients with oesophageal or gastric cancer and these cancers were subsequently excluded from enrolment. In the Hokusai-cancer study, subgroup analysis showed a significant interaction between GI cancer and increased risk of major bleeding in patients receiving edoxaban.

Mechanisms of DOAC-related gastro-intestinal bleeding could be due to an incomplete intestinal absorption, which can increase the amount of active molecule inside the gastrointestinal lumen and exacerbate existing lesions \[43\]. Nevertheless, DOAC seem to increase bleeding risk of other mucosa, such as vaginal and upper airways \[44, 45\]. Direct factor Xa inhibitors also decrease local thrombin generation and platelet activation which may contribute to mucosal bleeding \[46\]. As the number of studies comparing DOAC to LMWH was limited, we were not able to evaluate if this increased risk of bleeding was class specific or differed between molecules. A population-based study with a propensity-score matched cohort showed that risk of GI bleeding may vary between different Factor Xa inhibitors \[47\].

Another explanation to the superior efficacy of DOAC to prevent recurrence, and its increased rate of bleeding might be a lower degree of anticoagulation in the LMWH arm. The two studies comparing DOAC to LMWH used a conventional regimen of dalteparin as a comparator. This regimen includes a 25% dose reduction after the first month of anticoagulation, which might contribute to a lower efficacy and improved safety. Finally, the time under treatment was longer among patients allocated to DOAC than those allocated to LMWH, which reflects the better adherence to oral treatment and may also contribute to the lower rate of recurrence and increased rate of bleeding. In real world studies, the differences in adherence to oral or parenteral treatment are even more marked \[12, 48, 49\] which may further increase safety and efficacy differences.

Overall, DOAC appeared as the most probably superior treatment to prevent recurrence while the LMWH strategy was likely safer for bleeding complications. In terms of absolute differences, the reduction of VTE recurrence with DOAC compared to LMWH and the increase in major bleeding were of similar magnitude precluding to conclude to the superiority of any
class of treatment but rather suggesting to tailor the choice to each patient’s thrombotic/bleed-
ing risk profile.

Our study has limitations. First, systematic reviews and meta-analyses rely on the quality of
included studies. Although most included studies were considered at low risk of bias, patients
were not blinded in most studies. Moreover, the number of included studies comparing
DOAC to LMWH and the number of events in some comparison arms were small, which lim-
its the precision of treatment effect estimates and precludes analysing additional sources of
heterogeneity.

We compared different classes of anticoagulants but molecules and treatment regimens var-
ied across studies. Unfortunately, the limited number of available studies was insufficient to
evaluate a molecule effect within a given class. Nevertheless we pooled the results of these dif-
ferent molecules and regimens without observing a large amount of heterogeneity.

Our study has also several strengths. First, we performed a thorough literature search to
provide an exhaustive summary of the current best evidence. Second, in contrast to previous
meta-analyses, we reported combined hazard ratios which are measures of association better
suited to survival data than risk ratios or odds ratios, especially when lengths of follow-up vary
across studies [50]. Third, we aimed to support clinical decision by providing relative and
absolute risk differences and treatment ranking for efficacy and safety outcomes. Finally, we
provided both estimates for direct and network comparisons in order to take into account
direct and indirect evidence and contribute to explain contrasted data observed in previously
published systematic reviews on this topic.

Conclusion
Our meta-analysis shows that DOAC may be more effective than LMWH to prevent VTE
recurrence in cancer patients but are associated with an increased risk of major bleeding, espe-
cially in patients with GI cancer. Therefore, the choice of the best anticoagulant strategy should
be personalised, taking into account recurrence and bleeding risks, including cancer site, and
patients’ values and preferences. Results of ongoing studies may help clarify which patients
may benefit most from DOAC and potential differences between available Xa inhibitors.

Supporting information
S1 Table. PRISMA checklist.
(DOCX)
S2 Table. Detailed search strategy.
(DOCX)
S3 Table. Pooled risks in different treatment arms.
(DOCX)
S4 Table. Risk differences extrapolated from baseline risk in the Hokusai cancer VTE
study.
(DOCX)
S5 Table. Forrest plots for each outcome.
(DOCX)
S6 Table. Risk of bias of included studies.
(DOCX)
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Author Contributions

Conceptualization: Anne Rossel, Helia Robert-Ebadi, Christophe Combescure, Alfredo Addeo, Jean-Luc Reny, Christophe Marti.

Data curation: Helia Robert-Ebadi, Christophe Combescure, Jean-Luc Reny.

Formal analysis: Anne Rossel, Christophe Combescure, Christophe Marti.

Methodology: Anne Rossel, Helia Robert-Ebadi, Christophe Combescure, Jean-Luc Reny, Christophe Marti.

Supervision: Helia Robert-Ebadi, Jean-Luc Reny, Christophe Marti.

Validation: Anne Rossel, Helia Robert-Ebadi, Christophe Combescure, Olivier Grosgeurin, Jérôme Stirnemann, Alfredo Addeo, Nicolas Garin, Christophe Marti.

Visualization: Christophe Marti.

Writing – original draft: Anne Rossel, Helia Robert-Ebadi, Christophe Combescure, Jean-Luc Reny, Christophe Marti.

Writing – review & editing: Anne Rossel, Helia Robert-Ebadi, Christophe Combescure, Olivier Grosgeurin, Jérôme Stirnemann, Alfredo Addeo, Nicolas Garin, Thomas Agoritsas, Jean-Luc Reny, Christophe Marti.

References

1. Heit JA, O’Fallon WM, Petterson TM, Lohse CM, Silverstein MD, Mohr DN, et al. Relative impact of risk factors for deep vein thrombosis and pulmonary embolism: a population-based study. Arch Intern Med. 2002; 162(11):1245–8. Epub 2002/06/01. PMID: 12038942.

2. Blom JW, Doggen CJ, Osanto S, Rosendaal FR. Malignancies, prothrombotic mutations, and the risk of venous thrombosis. JAMA. 2005; 293(6):715–22. Epub 2005/02/11. https://doi.org/10.1001/jama.293.6.715 PMID: 15701913.

3. Kearon C, Akl EA, Ornelas J, Blaivas A, Jimenez D, Bounameaux H, et al. Antithrombotic Therapy for VTE Disease: CHEST Guideline and Expert Panel Report. Chest. 2016; 149(2):315–52. Epub 2016/02/13. https://doi.org/10.1016/j.chest.2015.11.026 PMID: 26867832.

4. Prandoni P, Lensing AW, Piccioli A, Bernardi E, Simioni P, Girolami B, et al. Recurrent venous thromboembolism and bleeding complications during anticoagulant treatment in patients with cancer and venous thrombosis. Blood. 2002; 100(10):3484–8. Epub 2002/10/24. https://doi.org/10.1182/blood-2002-01-0168 PMID: 12393647.

5. Lee AYY, Levine MN, Baker RL, Bowden C, Kaakkar AK, Prins M, et al. Low-molecular-weight heparin versus a coumarin for the prevention of recurrent venous thromboembolism in patients with cancer. New England journal of medicine. 2003; 349(2):146–53. https://doi.org/10.1056/NEJMoa025313 PubMed Central PMCID: PMCPHarmacia. PMID: 12853587.

6. Hull RD, Pineo GF, Brant RF, Mah AF, Burke N, Dear R, et al. Long-term Low-Molecular-Weight Heparin versus Usual Care in Proximal-Vein Thrombosis Patients with Cancer. American journal of medicine. 2006; 119(12):1062–72. https://doi.org/10.1016/j.amjmed.2006.02.022 PubMed Central PMCID: PMCPLeo(D enmark). PMID: 17145251.
16. Büllner HR, Prins MH, Lensing AWA, Decousus H, Jacobson BF, Minar E, et al. Oral rivaroxaban for the treatment of acute venous thromboembolism in patients with active cancer: A randomized controlled trial. JAMA—Journal of the American Medical Association. 2015; 314(7):677–86. https://doi.org/10.1001/jama.2015.9243 PubMed Central PMCID: PMCPMC6. PMID: 26284719

17. Lee AYY, Kamphuisen PW, Meyer G, Bauersachs R, Janas MS, Janer MF, et al. Tinzaparin vs warfarin for treatment of acute venous thromboembolism in patients with active cancer: A randomized clinical trial. JAMA—Journal of the American Medical Association. 2015; 314(7):677–86. https://doi.org/10.1001/jama.2015.9243 PubMed Central PMCID: PMCPMC6. PMID: 26284719

18. Schulman S, Kearon C, Subcommittee on Control of Anticoagulation of the S, Standardization Committee of the International Society on T, Haemostasis. Definition of major bleeding in clinical investigations of antithrombotic medicinal products in non-surgical patients. J Thromb Haemost. 2005; 3(4):692–4. https://doi.org/10.1111/j.1538-7836.2005.01204.x PMID: 15842354.
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24. Kaatz S, Ahmad D, Spyropoulos AC, Schulman S, Subcommittee on Control of A. Definition of clinically relevant non-major bleeding in studies of anticoagulants in atrial fibrillation and venous thromboembolic disease in non-surgical patients: communication from the SSC of the ISTH. J Thromb Haemost. 2015; 13(11):2119–26. PMID: 26764429.

25. Higgins JP, Altman DG, Gotzsche PC, Juni P, Moher D, Oxman AD, et al. The Cochrane Collaboration’s tool for assessing risk of bias in randomised trials. BMJ. 2011; 343:d5928. https://doi.org/10.1136/bmj.d5928 PMID: 22008217; PubMed Central PMCID: PMC3916245.

26. Mills EJ, Thorlund K, Ioannidis JP. Demystifying trial networks and network meta-analysis. BMJ. 2013; 346:i2914. Epub 2013/05/16. https://doi.org/10.1136/bmj.i2914 PMID: 23674332.

27. Rucker G. Network meta-analysis, electrical networks and graph theory. Res Synth Methods. 2012; 3 (4):312–24. Epub 2012/12/01. https://doi.org/10.1002/jrsm.1058 PMID: 26053424.

28. Perneger TV. Estimating the relative hazard by the ratio of logarithms of event-free proportions. Contemp Clin Trials. 2008; 29(5):762–6. Epub 2008/07/22. https://doi.org/10.1016/j.cct.2008.06.002 PMID: 18639651.

29. Riley RD, Jackson D, Salanti G, Burke DL, Kirkham J, et al. Multivariate and network meta-analysis of multiple outcomes and multiple treatments: rationale, concepts, and examples. BMJ. 2017; 358:j3932. Epub 2017/09/15. https://doi.org/10.1136/bmj.j3932 PMID: 28903924; PubMed Central PMCID: PMC5596393 interests and declare: none.

30. Rucker G, Schwarzer G. Ranking treatments in frequentist network meta-analysis works without resampling methods. BMC Med Res Methodol. 2015; 15:58. Epub 2015/08/01. https://doi.org/10.1186/s12874-015-0060-8 PMID: 26227148; PubMed Central PMCID: PMC4521472.

31. Deitcher SR, Kessler CM, Merli G, Rigas JR, Lyons RM, Fareed J. Secondary prevention of venous thromboembolism in patients with active cancer: Enoxaparin alone versus initial enoxaparin followed by warfarin for a 180-day period. Clinical and applied thrombosis/hemostasis. 2006; 12(4):389–96. https://doi.org/10.1177/1076029606235922 PMID: 17000884.

32. Romero A, Cairols MA, Vila-Coll R, Martí X, Colomé E, Bonell A, et al. A Randomised Open-Label Trial Comparing Long-term Sub-Cutaneous Low-Molecular-weight Heparin Compared with Oral-Anticoagulant Therapy in the Treatment of Deep Venous Thrombosis. European journal of vascular and endovascular surgery. 2009; 37(3):349–56. https://doi.org/10.1016/j.ejvs.2008.11.030 PubMed Central PMCID: PMC3924489.

33. López-Beret P, Orgaz A, Fontcuberta J, Doblas M, Martínez A, Lozano G, et al. Low molecular weight heparin versus oral anticoagulants in the long-term treatment of deep vein thrombosis. Journal of vascular surgery. 2001; 33(1):77–90. https://doi.org/10.1067/mva.2001.109336 PubMed Central PMCID: PMSanofi Winthrop(France). PMID: 11137927.

34. Agnelli G, Buller HR, Cohen A, Gallus AS, Lee TC, Pak R, et al. Oral apixaban for the treatment of venous thromboembolism in cancer patients: Results from the AMPLIFY trial. Journal of Thrombosis and Haemostasis. 2015; 13(12):2187–91. https://doi.org/10.1111/jth.13153 PMID: 26407753.

35. Prins MH, Lensing AW, Brighton TA, Lyons RM, Rehm J, Trajanovic M, et al. Oral rivaroxaban versus enoxaparin with vitamin K antagonist for the treatment of symptomatic venous thromboembolism in patients with cancer (EINSTEIN-DVT and EINSTEIN-PE): a pooled subgroup analysis of two randomised controlled trials. Lancet Haematol. 2014; 1(1):e37–46. https://doi.org/10.1016/S2352-3026(14)70018-3 PMID: 27030066.

36. Raskob GE, van Es N, Segers A, Angchaisuksri P, Oh D, Boda Z, et al. Edoxaban for venous thromboembolism in patients with cancer: results from a non-inferiority subgroup analysis of the Hokusai-VTE randomised, double-blind, double-dummy trial. Lancet Haematol. 2016; 3(8):e379–87. Epub 2016/08/02. S2352-3026(16)30057-6 [pii] https://doi.org/10.1016/S2352-3026(16)30057-6 PMID: 27476789.

37. Schulman S, Goldhaber SZ, Kearon C, Kakkar AK, Schellong S, Eriksson H, et al. Treatment with dabigatran or warfarin in patients with venous thromboembolism and cancer. Thromb Haemost. 2015; 114 (1):150–7. Epub 2015/03/06. 14-11-0977 [pii] https://doi.org/10.1160/TH14-11-0977 PMID: 25739680.

38. Mazzi P, Suceveanu A, Suceveanu A, Baz R, Catrinoiu D. Venous thromboembolism: secondary prevention with dabigatran vs. acenocumarol in patients with paraneoplastic deep vein thrombosis. Results from a small prospective study in Romania. Cardiovascular Research. 2014; 103(suppl_1): S39. https://doi.org/10.1093/cvr/cvu082.154.

39. Vedovati MC, Giustozzi M, Bonitta G, Agnelli G, Becattini C. Efficacy and safety of anticoagulant agents in patients with venous thromboembolism and cancer: A network meta-analysis. Thromb Res. 2018; 170:175–80. https://doi.org/10.1016/j.thromres.2018.08.023 PMID: 30196195.

40. Li A, Garcia DA, Lyman GH, Carrier M. Direct oral anticoagulant (DOAC) versus low-molecular-weight heparin (LMWH) for treatment of cancer associated thrombosis (CAT): A systematic review and meta-analysis. Thromb Res. 2018. https://doi.org/10.1016/j.thromres.2018.02.144 PMID: 29506866; PubMed Central PMCID: PMC619655.
41. Khorana A.A. MK, Milentijevic D., McCormick N., Laliberté F., Crivera C., Lefebvre P., Lejeune D., Rozjabek H., Schein J., Streiff M.B. VTE recurrence and safety of Anticoagulants among patients with cancer treated for venous thromboembolism. Blood. 2017;(130 (Suppl 1)):4631.

42. Sobieraj DM, Baker WL, Smith E, Sasiela K, Trexler SE, Kim O, et al. Anticoagulation for the Treatment of Cancer-Associated Thrombosis: A Systematic Review and Network Meta-Analysis of Randomized Trials. Clin Appl Thromb Hemost. 2018:1076029618800792. https://doi.org/10.1177/1076029618800792 PMID: 30244595.

43. Abraham NS, Horsley-Silva JL. Gastrointestinal bleeding secondary to the new anticoagulants. Curr Opin Gastroenterol. 2016; 32(6):474–80. https://doi.org/10.1097/MOG.0000000000000310 PMID: 27636303.

44. Brekelmans MP, Scheres LJ, Bleker SM, Hutten BA, Timmermans A, Buller HR, et al. Abnormal vaginal bleeding in women with venous thromboembolism treated with apixaban or warfarin. Thromb Haemost. 2017; 117(4):809–15. Epub 2017/02/10. 16-11-0874 [pii] https://doi.org/10.1160/TH16-11-0874 PMID: 28180233.

45. Becattini C, Franco L, Beyer-Westendorf J, Masotti L, Nitti C, Vanni S, et al. Major bleeding with vitamin K antagonists or direct oral anticoagulants in real-life. Int J Cardiol. 2017; 227:261–6. https://doi.org/10.1016/j.ijcard.2016.11.117 PMID: 28043907.

46. Honda Y, Kamisato C, Morishima Y. Edoxaban, a direct factor Xa inhibitor, suppresses tissue-factor induced human platelet aggregation and clot-bound factor Xa in vitro: Comparison with an antithrombin-dependent factor Xa inhibitor, fondaparinux. Thromb Res. 2016; 141:17–21. https://doi.org/10.1016/j.thromres.2016.02.026 PMID: 26962981.

47. Abraham NS, Noseworthy PA, Yao X, Sangaralingham LR, Shah ND. Gastrointestinal Safety of Direct Oral Anticoagulants: A Large Population-Based Study. Gastroenterology. 2017; 152(5):1014–22 e1. https://doi.org/10.1053/j.gastro.2016.12.018 PMID: 28043907.

48. Khorana AA, McCrae KR, Milentijevic D, Fortier J, Nelson WW, Laliberte F, et al. Current practice patterns and patient persistence with anticoagulant treatments for cancer-associated thrombosis. Res Pract Thromb Haemost. 2017; 1(1):14–22. https://doi.org/10.1002/rth2.12002 PMID: 30046670; PubMed Central PMCID: PMCPMC6058198.

49. Streiff MB, Milentijevic D, McCrae K, Yannicelli D, Fortier J, Nelson WW, et al. Effectiveness and safety of anticoagulants for the treatment of venous thromboembolism in patients with cancer. Am J Hematol. 2018; 93(5):664–71. https://doi.org/10.1002/ajh.25059 PMID: 29396864; PubMed Central PMCID: PMCPMC5947542.

50. Combescure C, Courvoisier DS, Haller G, Perneger TV. Meta-analysis of binary outcomes from two-by-two tables when the length of follow-up varies and hazards are proportional. Stat Methods Med Res. 2011; 20(5):531–40. Epub 2010/08/19. https://doi.org/10.1177/0962280210379172 PMID: 20716589.