ICH in primary or metastatic brain cancer patients with or without anticoagulant treatment: a systematic review and meta-analysis

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Anticoagulant treatment in patients with primary and metastatic brain cancer is a concern due to risk of intracranial hemorrhage (ICH). We performed a systematic review and meta-analysis to evaluate the risk of ICH in patients with primary or metastatic brain cancer treated with or without anticoagulants. Articles on ICH in patients with primary or metastatic brain cancer treated with or without anticoagulants published up to September 2021 were identified by searching PubMed, EMBASE, and Cochrane Library databases. The primary outcome of this analysis was ICH. Thirty studies were included. Rate of ICH was 13.0% in 1009 patients with metastatic brain cancer and 6.4% in 2353 patients with primary brain cancer (relative risk [RR], 3.26; 95% confidence interval [CI], 2.69-3.94; $I^2 = 92.8\%$). In patients with primary brain cancer, ICH occurred in 12.5% and 4.4% of patients treated with or without anticoagulants, respectively (11 studies, 659 treated and 1346 not treated patients; RR, 2.63; 95% CI, 1.48-4.67; $I^2 = 49.6\%$). In patients with metastatic brain cancer, ICH occurred in 14.7% and 15.4% (5 studies, 265 treated and 301 not treated patients; RR, 0.92; 95% CI, 0.43-1.93; $I^2 = 0\%$). ICH occurred in 8.3% of 172 treated with direct oral anticoagulants (DOACs) and in 11.7% of 278 treated with low-molecular weight heparin (LMWH) (5 studies; RR, 0.44; 95% CI, 0.25-0.79; $I^2 = 0$ %). Patients with metastatic brain cancer have a particularly high risk of ICH. Patients with primary brain cancer have an increased risk of ICH during anticoagulation. DOACs are associated with a lower risk of ICH than LMWH.

Introduction

It is estimated that 24 000 new cases of primary brain tumors and 200 000 new cases of brain metastatic cancers occurred in 2020 in the United States. Up to 20% to 25% of metastases spread to the central nervous system, with the highest incidence in patients with breast, melanoma, kidney, and non-small cell lung cancer. Spontaneous intracranial hemorrhage (ICH) is a common complication in patients with primary brain cancer and brain metastases. The clinical presentation of ICH is extremely heterogeneous, ranging from asymptomatic deposition of hemosiderin within the tumor seen on neuroimaging to large bleeds that cause clinical symptoms, either focal or related to intracranial hypertension. The incidence of ICH in patients with primary brain cancer and brain metastases varies according to the imaging diagnostic criteria and the cancer histotype, with rates as high as 50% in patients with brain metastases from melanoma, thyroid carcinoma, renal cell carcinoma, and choriocarcinoma.

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Requests for data sharing may be submitted to the corresponding author (mario. mandala@unipg.it).

The full-text version of this article contains a data supplement.

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When, for any reason, an anticoagulant treatment is required, in patients with primary brain cancer and brain metastases, the conflict between the need for anticoagulation and the risk of bleeding is a challenge. In the context of a timely and debated medical issue, we report the results of a systematic review and meta-analysis in patients with primary brain cancer or brain metastases treated with or without anticoagulant therapy at therapeutic doses to provide summary estimates of ICH across studies, to evaluate these rates in patients with primary or metastatic brain cancer separately, and to investigate the impact of anticoagulation as well as of different anticoagulation strategies on ICH rates.

Materials and methods

This systematic review and meta-analysis was conducted according to the Preferred Reporting Items for Systematic Reviews and Meta-Analyses statement⁶ (http://www.prisma-statement.org). This metaanalysis has been registered in PROSPERO.

Search strategy

We performed an unrestricted search in PubMed, EMBASE, and the Cochrane Library databases from inception to September 2021. The following search terms were used to search clinical trials, registers, and databases: "glioma, glioblastoma, oligodendroglioma, astrocytoma, oligoastrocytoma, brain metastasis" and "anticoagulant, heparin, low-molecular-weight heparin, vitamin K antagonist, oral anticoagulant, direct oral anticoagulant" and "hemorrhage, haemorrhage." Additional studies were identified by hand searching of reference lists of the reviews and retrieved articles. Eligibility assessment was performed independently by two authors (G.P., M.G.), using a data extraction form, in an unblinded standardized approach. Study selection was initially performed by review of titles, and candidate abstracts were then reviewed. A third reviewer (M.M.) resolved disagreements between reviewers and any differences in study selection.

Study selection

Inclusion criteria of this meta-analysis were: (1) randomized clinical trials or observational studies; (2) patients aged ≥18 years with primary brain cancer and/or brain metastases treated with or without therapeutic doses of anticoagulants; and (3) availability of number of patients who experienced ICH. No language, publication date, or publication status restrictions were imposed. Review articles not reporting original data, case reports and case series with <10 patients, abstracts, editorials/letters, and studies not involving humans were excluded. Studies reporting on the rate of ICH in patients with primary brain cancer or brain metastases who received thromboprophylaxis for venous thromboembolism (VTE) were also excluded. The inter-reviewer agreement for study selection was assessed by the k statistic, which measures agreement beyond chance.7

Study objectives and outcomes

The primary objective of this meta-analysis was to evaluate the overall incidence of ICH in patients with primary brain cancer or brain metastases treated with or without anticoagulants at therapeutic doses. The secondary objectives were to evaluate: (1) the rate of major and fatal ICH; (2) the rate of ICH in relationship to anticoagulant therapy; and (3) the rate of ICH according to the type of anticoagulant treatment. The primary outcome was ICH. Secondary outcomes were major ICH and fatal ICH.

Major ICH was defined as symptomatic, requiring surgical intervention, with a volume ≥10 mL or as fatal.3,4,8-12 Any ICH was defined as the composite of major ICH and the other ICH categories that did not meet the major ICH criteria.

Data extraction

The following data were extracted from each included trial: (1) general data (study design and year of publication); (2) characteristics of trial participants (number, mean age, sex, site of cancer, and cancer histotype); (3) type of anticoagulant (agent, dose, duration, and daily dosing); and (4) study outcomes (primary outcome, secondary outcomes, and length of follow-up). The quality of the studies was evaluated by using the Newcastle-Ottawa quality assessment scale (range, 1-9 [1-3 indicates low quality, 4-6 indicates moderate quality, and 7-9 indicates high quality]).

Statistical analysis

Pooled outcome event rates were calculated by using the logit transformed proportion and corresponding sampling variances. The rates of ICH were pooled by using random effects models and are presented with the corresponding 95% confidence interval (CI). Heterogeneity was assessed by the l^2 test. $l^2 = 0$ was considered to indicate no heterogeneity; $l^2 < 25\%$, 25% to 75%, and >75% were considered to indicate low, moderate, and high degrees of heterogeneity, respectively. 13 To evaluate publication bias, both Egger's test and funnel plots of the logit transformed proportion vs standard error were computed. If the Egger's test confirmed asymmetry, the Duval and Tweedie's trim-and-fill procedure was used to compute an unbiased estimate of the effect size. A mixed effects meta-regression analysis was performed to test differences among subgroups according to the median length of follow-up and the median length of anticoagulation therapy. We also determined pooled relative risk (RR) and 95% Cl using a random effects model for ICH in patients with primary brain cancer or brain metastases treated with or without anticoagulant therapy and according to the anticoagulant type. For studies presenting zero cells, 0.5 was added for a correct estimation of risk measures. The following prespecified sensitivity analyses were performed: (1) rate of ICH in primary brain cancer only; (2) rate of ICH in brain metastases only; (3) rate of major ICH in primary brain cancer or brain metastases; (4) rate of ICH according to the type of anticoagulant treatment; (5) rate of fatal ICH; and (6) rate of ICH in patients with

The statistical analyses, forest plots, and publication bias analyses were produced with Comprehensive Meta-Analysis version 3.0 (Biostat Inc.). P values <.05 were considered statistically significant.

Results

The search of PubMed, EMBASE, and Cochrane Library databases provided a total of 692 articles, while 3 of them were identified through other sources. After removal of duplicates, as well as an additional 530 articles that did not meet the inclusion criteria at abstract review, the full text of the remaining 64 articles was examined in detail. Of these, 34 studies did not meet the inclusion criteria. Thus, 30 studies were included in the systematic review. 3-5,8-12,14-35 No unpublished relevant studies were found.

Table 1. Main clinical features of the studies included according to PICO criteria

Source and author	Study design	No. of pts	Participants	Intervention	Comparator	Median follow-up	Primary outcome	Definition of major ICH
Lee et al ¹⁴ (2021)	ď	Ξ	Pts with primary or secondary brain tumors who received either a DOAC or LMWH for the treatment of VTE	DOAC (55 pts): Rivaroxaban (38) Apixaban (13) Edoxaban (3) Dabigatran (1)	LMWH (56 pts): Enoxaparin	9 mo	Incidence of any ICH within 6 mo	Ψ.
Jo et al ¹⁵ (2021)	œ	220	Pts with high-grade glioma and VTE on LMWH	LMWH (88 pts)	No anticoagulation (22 pts)	12 mo	Incidence of 1-y ICH	NR
Dubinski et al ⁹ (2021)	α	94	Pts who underwent craniotomy for primary tumor resection and PE	DOACs (14 pts): Rivaroxaban (6) Edoxaban (8)	LMWH (32 pts)	15 mo (DOACs) and 9 mo (LMWH)	Clinical course, 6- and 12-mo follow-up and survival	Any hemorrhage that was ≥10 mL in volume, required surgical intervention, or was associated with clinical symptoms, such as nausea and vomiting, or focal neurologic deficit
Burth et al ¹⁶ (2021)	α	172	Pts with glioblastoma and pts with brain metastases with (cases)/without (controls) AF	Full anticoagulation (enoxaparin 40 mg, phenprocoumon, DOAC, or heparin) vs prophylactic anticoagulation/heparin	No anticoagulation	8.6 and 7.2 mo	Incidence of ICH in pts with glioblastoma and brain metastases with/without AF	R.
De Melo Junior et al ¹⁷ (2020)	ш	23	VTE pts on therapeutic anticoagulation started within the first 30 d after intracranial neurosurgical procedure (mostly primary neoplastic lesions)	Total 29 pts: warfarin	Total 21 pts: DOAC (19 rivaroxaban and 2 apixaban) or 4 pts enoxaparin 1 mg/kg twice daily	16 61 d.	Risk of ICH	£
Leader et ai ^{to} (2020)	α	96	Pts with brain metastases and anticoagulation therapy prescribed at therapeutic doses for either VTE or AF	DOAC (41 pts) Apixaban (11) Dabigatran (5) Edoxaban (8) Rivaroxaban (17)	LMWH (55 pts): Enoxaparin (34) Nadroparin (15) Tinzaparin (6)	136 d with DOAC and 175 d with LMWH	Major ICH during 12 mo of follow-up	ICH that measured ≥10 mL in volume, required surgical intervention, or was associated with clinical symptoms, focal neurologic deficits, or changes in cognitive function
Horstman et al ¹⁸ (2018)	α	125	Pts with brain metastasis with or without history of long-term (>1 mo) anticoagulation therapy	67 pts on anticoagulant therapy: once-daily enoxaparin (1.5 mg/kg) or twice-daily enoxaparin (1 mg/kg q12h)	58 pts: Not on anticoagulant therapy	Ä.	Incidence of ICH associated with anticoagulant use	Ϋ́
Carney et al ⁸ (2019)	ď	172	Pts with primary and secondary brain tumors on anticoagulation with a DOAC or LMWH for the treatment of VTE	Primary brain tumors: total 67 pts 20 with DOACs 47 with enoxaparin Brain metastasses: total 105 21 with DOAC 84 with enoxaparin 84 with enoxaparin	Enoxaparin ≥1.5 mg/kg, once daily	£	Major ICH within 12 mo from start of anticoagulation	ICH that measured ≥10 mL in volume, required surgical intervention, or was associated with clinical symptoms, focal neurologic deficits, or changes in cognitive function
AF, atrial fibrillation; IVC, inferior	r vena cava	ι; NR, not	AF, atrial fibrillation; IVC, inferior vena cava; NR, not reported; PE, pulmonary embolism; pts, patients; PICO, Patient/Population, Intervention, Comparison and Outcomes; R, retrospective.	patients; PICO, Patient/Popula	tion, Intervention, Compariso	on and Outcome	s; R, retrospective.	

Table 1. (continued)								
Source and author	Study design	No. of pts	Participants	Intervention	Comparator	Median follow-up	Primary outcome	Definition of major ICH
Gessler et al ¹⁹ (2018)	ď	35	Pts with primary and secondary tumors who underwent craniotomy on anticoagulant treatment for cerebral vein thrombosis	Full therapeutic anticoagulation (25 pts)	Intermediate dosing of LMWH (10 pts)	181 d	Investigate the occurrence, risk factors, and outcomes associated with the development of cerebral vein thrombosis after cranictomy	쫀
Chai-Adisaksopha et al ²⁰ (2017)	α	364	Pts with primary or metastatic brain tumors with VTE (182) treated with anticoagulation and 182 control subjects (pts without brain tumors with VTE on anticoagulant treatment)	Therapeutic dose of LMWH; 5.9% pts received reduced-dose LMWH	162 of 182 pts in the control group received therapeutic dose LMWH; 3.6% pts received-reduced dose LMWH	6.7 то	Incidence of the first major bleeding after starting anticoagulant therapy	
Mantia et al ¹¹ (2017)	œ	133	Pts with primary brain tumors on therapeutic anticoagulation for VTE	Enoxaparin (50 pts) 1 mg/kg, twice daily (76%) or enoxaparin at 1.5 mg/kg (2 pts) or enoxaparin less than standard therapeutic dosing (8 pts)	No anticoagulation (83 pts)	r z	Major ICH from time of diagnosis of primary brain tumor	Any hemorrhage that was ≥10 mL in volume, required surgical intervention, or was associated with clinical symptoms such as nausea and vomiting, focal neurologic deficit, or change in cognitive function
Al Megren et al ²¹ (2017)	œ	152	Glioma pts with VTE or cerebral vein thrombosis	Full anticoagulation with/ without IVC filter with LMWH or unfractionated heparin or fondaparinux (76 pts)	No anticoagulation: glioma pts without VTE (76 pts)	11 0 E	ICH defined as any bleeding into the cranial vault over the follow-up period	NR T
Khoury et al ²² (2016)	œ	173	Pts with glioblastoma and VTE with/without anticoagulation	Total 97 pts: LMWH (69), warfarin (26), heparin (2)	No treatment: 76 pts	6.1 mo	Incidence of ICH	Z.
Donato et al ³ (2015)	α	293	Pts with brain metastasis on the reapeutic enoxaparin for the treatment of VTE	Enoxaparin 1 mg/kg twice daily (76 pts), 1.5 mg/kg once daily (17 pts), and modified dose-reduced therapeutic dosing (11 pts)	189 controls: no anticoagulation	X X	Measurable (>1 mL in volume) ICH from initial diagnosis of brain metastases	Larger volume bleeds (>10 mL), the presence of new symptoms, or the need for surgical intervention
Smith et al ²³ (2015)	œ	69	Pts who underwent surgical resection of primary or metastatic brain tumor	Full anticoagulation with warfarin, enoxaparin, heparin, dalteparin	Prophylactic doses of heparin or enoxaparin	Z Z	Determine the risk factors for VTE in pts who underwent neurosurgical resection of brain tumors	N.
Yust-Katz et al ²⁴ (2015)	α	49	Pts with glioblastoma who developed a VTE during the course of their disease	Anticoagulation alone in 36 pts (8 with Coumadin and 28 with LMWH)	2 pts had IVC filter alone and 21 pts received both an IVC filter and anticoagulation	Ÿ	Estimate the frequency of VTE in glioblastoma pts and identify potential risk factors for the development of VTE during adjuvant chemotherapy	NA N

AF, atrial fibrillation; VC, inferior vena cava; NR, not reported; PE, pulmonary embolism; pts, patients; PICO, Patient/Population, Intervention, Comparison and Outcomes; R, retrospective.

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and of treatment-related

complication

The incidence of perioperative VTE

Ä

Total: 17 pts who had vena cava filters placed

Total: 109 81 treated with heparin, 28 with enoxaparin

Pts who underwent surgery for primary or metastatic brain tumors

126

œ

Chaichana et al²⁵ (2013)

Table 1. (continued)								
Source and author	Study No. design of pts	No. of pts	Participants	Intervention	Comparator	Median follow-up	Primary outcome	Definition of major ICH
Aishima and Yoshimoto ²⁶ (2013)	<u>~</u>	23	23 Pts who underwent surgery for	Screening cohort	Nonscreening cohort	2	The effectiveness and	<u>د</u> ک

Source and author	Study design	No. of pts	Participants	Intervention	Comparator	Median follow-up	Primary outcome	Definition of major ICH
Aishima and Yoshimoto ²⁶ (2013)	œ	23	Pts who underwent surgery for primary or metastatic brain tumors	Screening cohort with serum D-dimer level	Nonscreening cohort	Σ Σ	The effectiveness and safety of screening strategy for the detection and prevention of VTE	NR
Alvarado et al ²⁷ (2012)	œ	74	Pts with melanoma with brain metastasis and VTE	Total: 57 pts Anticoagulation alone in 26 pts, 31 also IVC filter placement	No coagulation in 17 pts: 3.4 mo 13 had IVC filter, 4 only supportive measures	o Œ	Risk and benefits of systemic anticoagulation in these pts	X.
Norden et al ⁴ (2011)	ď	282	Pts with gloma treated with bevacizumab and anticoagulants for VTE	Total 64: Enoxaparin (49) Dalteparin (1) Fondaparinux (1) Warfarin (13)	218 pts treated without anticoagulation	χ α	Hemorrhagic risk of concurrent use of bevacizumab and anticoagulants in glioma pts	Any hemorrhage of grade 3 or greater
Pan et al ²⁸ (2009)	œ	6 8	Pts with glioblastoma with VTE or ICH	Total: 25 pts Fourteen with IVC filter and anticoagulation, 11 with anticoagulation without IVC filter	Total: 14 pts Pts with IVC filter without anticoagulation	χ α	Incidence of initial and recurrent compared with the incidence of ICH in pts with glioblastoma	Ψ Z
Nghiemphu et al ¹² (2008)	œ	265	Pts with gliomas who were treated concurrently with bevacizumab and anticoagulation for VTE	Total: 21 pts Nine on LMWH and 12 on warfarin	Total: 244 pts No anticoagulation	184 d	Incidence of major ICH	ICH with severe neurologic deficits
Ghanim et al ²⁹ (2007)	œ	175	Patients with VTE and primary or metastatic brain tumors and/or brain hemorrhage	Anticoagulants (total 39 pts): Vena cava filter Prophylactic dose (7 pts) (136 pts) or therapeutic dose (32 pts)	Vena cava filter (136 pts)	92 d	Mortality risk for VTE between pts treated with IVF and anticoagulants	X.
Schiff and DeAngelis ⁵ (1994)	œ	24	Pts with brain metastases who experienced VTE	Total: 42 29 on IV heparin followed by warfarin 2 on IV heparin 2 with warfarin alone 7 with IV heparin and subcutaneous heparin 2 with IV heparin, subcutaneous heparin, and warfarin	IVC filters	p 88	The efficacy and complications of filters and anticoagulation	NA NA
Quevedo et ai ^{so} (1994)	œ	16	Pts with primary brain tumors	Heparin followed by warfarin sodium	No anticoagulation	œ Z	The associations between VTE and factors related to the risk of occurrence of VTE.	X.
Altschuler et al ³¹ (1990)	œ	23	Pts with malignant glioma treated with anticoagulant therapy for VTE	Continuous IV heparin and an oral dose of 10 mg of warfarin	NR	R R	Safety and effectiveness of anticoagulation treatment of VTE	NR
Olin et al ⁹² (1987)	α	49	Olin et al ³² (1987) R 49 Pts with primary and secondary Total 25 pts: Total: 24 pts treated NR The complication mortality betw continuous infusion of 500 with IVC filter treated with IVC filter treated with IVC once daily or and anticoagu	Total 25 pts: IV heparin sodium at continuous infusion of 500 U/kg once daily or warfarin (17 pts)	Total: 24 pts treated with IVC filter	Z Z	The complications and mortality between pts treated with IVC filter and anticoagulants	NA NA

AF, atrial fibrillation; IVC, inferior vena cava; NR, not reported; PE, pulmonary embolism; pts, patients; PICO, Patient/Population, Intervention, Comparison and Outcomes; R, retrospective.

	Study	, o				Median	Primary	Definition
Source and author	design	of pts	Participants	Intervention	Comparator	follow-up	outcome	of major ICH
Choucair et al ³³ (1987)	ш	98	Pts with malignant gliomas with a score of ≥60% on the Kamofsky performance scale who underwent intracranial surgery	Total: 22 pts IV heparin for 7-10 d followed by subcutaneous heparin (5000-8000 U twice daily) for at least 3 mo or oral warfarin	Total: 14 pts No anticoagulation	α Σ	Risk of ICH in pts with malignant gliomas, treated with anticoagulant for late postoperative thromboembolism	۳
Ruff and Posner ³⁴ (1983)	ď	266	Pts with malignant astrocytoma or glioblastoma multiforme	Total: 95 pts IV heparin for 7-14 d, followed by warfarin for 6 to 14 wk	Total: 171 pts No anticoagulant	96 wk and group 2 for 36 wk	Incidence, prevention, Ni and treatment of VTE	Z.
Ruff and Posner ³⁵ (1981)	œ	375	Pts with malignant astrocytomas	Total: 103 pts Heparin followed by warfarin for 6 to 14 wk	Total: 272 pts No anticoagulation	NR	Incidence of VTE and the risk of systemic anticoagulation	ď

not reported; PE, pulmonary embolism; pts, patients; PICO, Patient/Population, Intervention, Comparison and Outcomes; R, retrospective.

atrial fibrillation; IVC, inferior vena cava; NR,

ĄF,

The flow diagram of the literature search is shown in supplemental Figure 1, and the main features of the studies are reported in Table 1 and supplemental Tables 1 and 3. The inter-reviewer agreement for study selection was very good (κ statistic, 0.85).

The studies selected for the review included 3893 patients (range for individual studies, 16-364 patients), and all were retrospective. Fifteen studies included patients with primary brain cancer only. 4,9,11,12,15,17,21,22,24,28,30,31,33-35 six studies included patients with brain metastases only, 3,5,10,16,18,27 and nine studies included both patients with primary brain cancer and patients with brain metastases.^{8,14,19,20,23,25,26,29,32} Overall, this analysis includes 2353 patients with primary brain cancer and 1009 with brain metastases. Seven studies (531 patients) did not separately report ICH occurring in primary cancer or brain metastases. 17,19,23,25,26,29,32

Mean age varied from 51 to 72 years, and men were slightly more represented than women. Seventeen studies (2896 patients) included both patients treated (1072 patients) and not treated (1824 patients) with anticoagulants, whereas 13 studies (997 patients) included only those treated with anticoagulants. The main indication for anticoagulant treatment was acute VTE (25 studies, 3313 patients), followed by atrial fibrillation (2 studies, 268 patients), cerebral vein thrombosis (2 studies, 187 patients), or any indication for anticoagulant treatment (1 study, 125 patients) (Table 1).

The anticoagulant agent was heparin in 21 studies (1338 patients),3-5,8-12,14-25 warfarin in 13 studies (475 tients), 4,5,12,17,22-24,30-35 and direct oral anticoagulants (DOACs) in 5 studies (172 patients).8-10,14,16

The median patient's observation period was 125 days, ranging from to 27 days and 240 days. The median duration of anticoagulant treatment at the time of ICH was 8.1 months.

The quality of studies assessed according to the Newcastle-Ottawa quality assessment scale was poor in 22 studies and good in 8 studies. The quality assessment is reported in supplemental Table 2.

Rates of ICH in primary or metastatic brain cancer

Overall, the weighted incidence rates in patients with primary brain cancer or brain metastases estimated by using a random effects model was 7.7% (95% Cl. 5.1-11.5; $l^2 = 92.8\%$; 445 events in 3893 patients) (Table 2). Egger's tests revealed the presence of publication bias (t = 5.90; P < .001). After using the trim-and-fill procedure, 5 studies were trimmed, and the ICH adjusted rate was 9.1% (95% Cl, 6.2-13.2). Bias assessment plots are reported in supplemental Figure 2A.

Rates of ICH in patients with primary brain cancer were reported in 18 studies and ranged from 1.1% 25.4%. ^{4,9-12,16,18-21,23,27,28,30,31,33-35} The weighted incidence rate was 6.4% (95% Cl, 4.1-9.9; $l^2 = 84.4\%$; 156 events in 2353 patients) (supplemental Figure 3A), and the risk of publication bias was significant (t = 2.39; P = .03) (supplemental Figure 2B). Adjusted value after the trim-and-fill procedure was 7.5% (95% CI, 4.9-11.3).

Rate of ICH in patients with metastatic brain cancer was reported in 9 studies and ranged from 2.7% to 47.6%. 3,5,9,12,14-16,18,26 In these studies, the weighted incidence rate of ICH was 13.0% (95% CI, 6.5-24.2; $l^2 = 93.7\%$; 218 events in 1009 patients) (supplemental Figure 3B). The risk of publication bias was significant (t = 5.10; P = .001); after the trim-and-fill procedure, 2 studies were trimmed,

Table 2. Rates of ICH, major ICH, and fatal ICH in patients with primary brain cancer or brain metastases

Outcome	No. of studies	No. of ICH/no. of patients	Rate	95% CI	/ ²
Overall ICH	30	445/3893	7.7%	5.1-11.5	92.8%
Major ICH	7	117/1287	6.2%	2.8-13.0	91.5%
Fatal ICH	11	13/764	2.9%	1.7-4.7	0%
ICH in PBC patients	18	156/2353	6.4%	4.1-9.9	84.4%
ICH in MBC patients	9	218/1009	13.0%	6.5-24.2	93.7%
Major ICH in PBC patients	4	30/793	3.9%	1.3-11.6	87.6%
Major ICH in MBC patients	3	87/494	15.4%	9.4-24.2	74.6%
ICH in patients with VTE	25	384/3313	7.1%	4.4-11.5	93.7%

MBC, metastatic brain cancer; PBC, primary brain cancer.

and the adjusted rate of ICH was 17.5% (95% CI, 9.6-29.8) (supplemental Figure 2C).

Risk of ICH was significantly higher in patients with metastatic brain cancer than in patients with primary brain cancer (RR, 3.26; 95% Cl, 2.69-3.94; $\dot{l}^2 = 92.8\%$).

At meta-regression analysis, the median length of a patient's observation period (P = .99; $I^2 = 87.7\%$) and the median length of anticoagulant therapy before ICH (P = .763; $I^2 = 92.4\%$) did not influence the rate of ICH.

For patients with VTE, the overall weighted incidence rate of ICH was 7.1% (95% Cl. 4.4-11.5: $l^2 = 93.7\%$: 25 studies, 384 events in 3313 patients)^{3-5,8,9,11,12,14,15,20-35} (Table 2). Specifically, the weighted incidence rate of ICH was 6.1% (95% Cl, 3.7-9.7%; $l^2 = 85.3\%$; 17 studies) and 13.6% (95% CI, 5.6-29.2; $l^2 = 94.1\%$; 6 studies) in patients with primary and metastatic brain cancer, respectively. Rates of major and fatal ICH in patients with primary or metastatic brain cancer are reported in Table 2.

Rates of ICH in patients treated with or without anticoagulant therapy

Overall, rates of ICH in patients treated with anticoagulants was 11.5% (95% Cl. 7.4-17.6: $l^2 = 83.7\%$: 152 events in 1072

patients) and 6.0% in those not treated with anticoagulants (95% Cl. 3.0-11.5: $f^2 = 92.2\%$: 177 events in 1824 patients) (RR. 1.81: 95% Cl, 1.15-2.84; P = .001; $I^2 = 60.3\%$) (Table 3). No publication bias was observed (t = 1.41; P = 0.17). In patients with primary brain cancer, anticoagulant therapy was associated with an increased risk of ICH and of major ICH (Figure 1; Table 3). In patients with metastatic brain cancer, anticoagulant therapy was not associated with an increased rate of ICH. Fatal ICH while on anticoagulant therapy was reported in 11 studies, and the weighted incidence rate was 2.7% (95% CI, 1.6-4.5; $l^2 = 0\%$; 13 events in 764 patients). 4,13,14,18,21,25,26,30,32,34,35

Rate of ICH according to the type of anticoagulant treatment

In patients with primary brain cancer or brain metastases, DOACs were associated with a lower risk of ICH than low-molecular weight heparin (LMWH) (8.3% vs 11.7%; RR, 0.44; 95% Cl, 0.25-0.79; I² 0%; 5 studies in 450 patients) (supplemental Figure 4A).8-10,14,17 In patients with primary brain cancer, DOACs were associated with a reduced risk of ICH (RR, 0.19; 95% Cl, 0.04-0.99; $l^2 = 0\%$; 0 events in 69 DOAC-treated patients and 19 events in 95 LMWH-treated patients, 4 studies). In patients with brain metastases, 12 events were observed in 103 DOAC-treated

Table 3. Rates of ICH in patients with primary brain cancer or brain metastases treated with or without anticoagulants

Setting	No. of studies	No. of ICH/no. of patients treated	Anticoagulant therapy	No. of ICH/no. of patients not treated	No anticoagulant therapy	RR	95% CI	P	l²
Overall patients	17	152/1072	11.5% (95% CI, 7.4-17.6)	177/1824	6.0% (95% CI, 3.0-11.5)	1.81	1.15-2.84	.001	60.3%
Patients with PBC	11	80/659	12.5% (95% CI, 8.0-18.8)	50/1346	4.4% (95% CI, 2.5-7.7)	2.58	1.59-4.19	<.001	45.5%
Patients with MBC	4	61/265	14.7% (95% Cl, 4.4-39.2)	81/301	15.4% (95% CI, 5.3-37.2)	0.86	0.45-1.65	.287	0%
Patients treated with DOACs vs LMWH	5	12/172	8.3% (95% Cl, 4.4-15.3)	71/278	11.7% (95% CI, 2.9-37.0)	0.44	0.25-0.79	.007	0%
Patients treated with LMWH vs warfarin	4	15/211	5.9% (95% Cl, 1.5-20.5)	8/198	5.4% (95% Cl, 1.5-17.3)	1.45	0.56-3.79	.185	0%
Overall major ICH	4	33/239	10.4% (95% Cl, 4.0-24.5)	47/734	3.4% (95% CI, 0.6-17.6)	1.93	0.79-4.73	.001	38.7%
Major ICH in patients with PBC	3	9/135	6.3% (95% Cl, 1.7-20.3)	9/545	1.8% (95% CI, 0.9-3.4)	3.75	1.6-4.5	.003	0%

MBC, metastatic brain cancer; PBC, primary brain cancer.

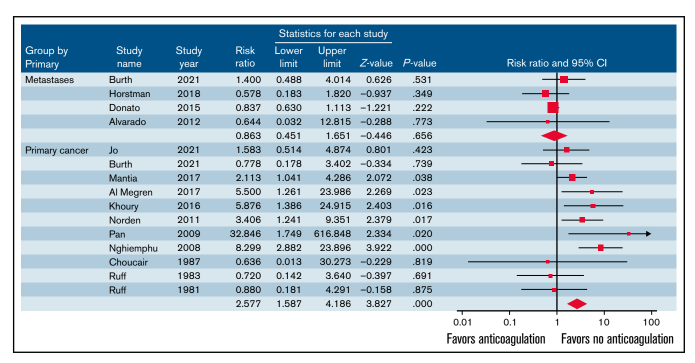


Figure 1. Risk of ICH in patients with primary or metastatic brain cancer treated with or without anticoagulants.

patients and 52 events in 183 LMWH-treated patients (RR, 0.65; 95% Cl, 0.36-1.16; $l^2 = 0\%$; 3 studies). When considering only studies in patients receiving anticoagulants for VTE, the RR for ICH was 0.36 (95% CI, 0.18-0.71; $l^2 = 0\%$; 4 studies) in patients treated with DOACs (8 events in 131 patients) compared with LMWH (64 events in 223 patients). 9,11,14,16 Risk of ICH was not significantly different with LMWH vs warfarin (4 studies, 15 events in 211 LMWH-treated patients vs 8 events in 198 warfarin-treated patients) (RR, 1.45; 95% Cl, 0.56-3.79; $I^2 = 0\%$) (supplemental Figure 4B). 17,22,23,27

Discussion

The current meta-analysis provides the following findings: (1) the rate of ICH and major ICH is higher in patients with metastatic brain cancer compared with those with primary brain cancer; (2) anticoagulant therapy is associated with an increase in ICH and major ICH in patients with primary brain cancer but not in those with brain metastases; and (3) the risk of ICH is lower in patients with primary or metastatic brain cancer treated with DOACs compared with those treated with LMWH.

According to international guidelines, presence of an intracranial primary or metastatic brain cancer is not an absolute contraindication for anticoagulation.³⁶ Nevertheless, limited data support the use of anticoagulant therapy in these patients. In this context of uncertainty, our results may be relevant and timely for clinical decision-making and design of future clinical trials. Of potential clinical interest, our analysis provides data on brain cancer overall as well as separately in patients with primary or metastatic brain cancer. Data are also provided concerning the rate of ICH in patients treated with or without anticoagulants.

In this meta-analysis, the rates of ICH and major ICH were higher in patients with metastatic brain cancer than in patients with primary

brain cancer. The safety profile of anticoagulant therapy seems to be different in patients with primary or metastatic brain cancer. Of clinical relevance, in patients with primary brain cancer, therapeutic anticoagulation was associated with an increased risk of ICH and major ICH. In contrast, in patients with metastatic brain cancer, the administration of anticoagulant therapy was not associated with an increased rate of ICH. Although ICH from metastatic brain cancers is a relatively common clinical observation, its pathogenesis has not been fully elucidated. Several biomarkers involved in angiogenesis reportedly contribute to the vascular instability of brain metastases.37-39 Our meta-analysis was not able to elucidate the risk of bleeding associated with specific tumor histotypes. Future studies are needed to definitively establish rates of bleeding and the safety of anticoagulation in malignancies associated with high ICH rates, including metastatic melanoma, choriocarcinoma, and renal cell carcinoma.

Due to the high bleeding risk, patients with brain cancer should be, as a priority, the target for future studies with new, potentially safer anticoagulant agents, including the anti-factor XI inhibitors. These agents have been recently shown to be associated with a lower risk of bleeding than LMWH^{40,41} when given for thromboprophylaxis in patients who underwent major orthopedic surgery.

The data regarding the relative safety of anticoagulation in patients with brain metastases seem to be reassuring. Thus, exclusion of these patients from clinical trials on the treatment of cancerassociated VTE should no longer be supported, although caution should remain for patients with primary brain cancer. It should be recommended that patients with primary or metastatic cancer, respectively, should be subjected to a priori stratification before study randomization.

Interestingly, although the number of patients on DOACs in our analysis was relatively low, we found that treatment with DOACs was associated with a lower risk of ICH than treatment with LMWH in patients with primary or metastatic brain cancer, with no heterogeneity across studies. These results are in agreement with recent studies suggesting the safety profile of DOACs in terms of ICH rates in patients with brain cancer 42,43 and make these agents an attractive strategy in particular for VTE treatment. Indeed, in our study, anticoagulants were mostly used for the treatment of VTE, and DOACs were associated with one-third the risk of ICH compared with that of LMWH.

The overall rates of ICH varied considerably, ranging from 1.4% to 47.6%. Specifically, among patients receiving anticoagulant therapy, the rate of ICH in patients with primary and metastatic brain cancer ranged from 1.4% to 29.0% and from 2.7% to 47.6%, respectively. Several reasons can justify these large ranges: the retrospective design of the included studies, the heterogeneity in ICH monitoring, the definition of ICH and imaging modalities used, cancer histotype, and the type of the adopted anticoagulant therapeutic strategy.

Our meta-analysis had several limitations, including the fact that none of the studies in the analysis was prospective. Furthermore, for different subgroup analyses, there was a significant heterogeneity due to differences in study target populations or targeted effects, protocol-scheduled imaging, types and duration of anticoagulant therapy, and/or analytical methods, including covariate adjustments. Moreover, the definition of major and nonmajor was not formally standardized, although the same definition was consistently reported in different studies. 3,4,8-12 Finally, due to the paucity of data, we were not able to distinguish rates of ICH in patients with brain cancer treated with or without chemotherapy or radiotherapy.

Nevertheless, our meta-analysis has several strengths, including: (1) an extensive search of available data that makes this meta-analysis the most updated reported so far; (2) the punctual estimation of the rates of ICH and major ICH in large series of patients with primary or metastatic brain cancer; and (3) the assessment of RRs in patients with primary or metastatic brain cancer treated with or without anticoagulants.

In summary, our study confirms the not-negligible risk of ICH in patients with primary brain cancer or brain metastases. Anticoagulation is associated with an increase in the risk of ICH and major ICH in patients with primary brain cancer. This increase in risk does not seem to occur in patients with brain metastases. DOACs seem to be associated with a lower risk of ICH than LMWH. Prospective controlled studies on new anticoagulant strategies, potentially associated with a reduced risk of bleeding, are needed in patients with brain cancer, with these patients and clinical settings as a priority in the unmet clinical need-based clinical research.

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Authorship

Contribution: M.M. and M.G. were responsible for study conception and design; G.P., M.G., and M.M. were responsible for data acquisition; and M.G. and C.B. performed the statistical analysis; and all authors contributed to interpretation of the data, drafting of the manuscript, critical revision of the manuscript for important intellectual content, and gave final approval of the manuscript.

Conflict-of-interest disclosure: C.B. reports lecture fees and consultancies for Bayer Health Care, Bristol Myers Squibb, and Daiichi Sankyo outside the submitted paper. G.A. reports lecture fees from Bayer, Boehringer Ingelheim, Bristol Myers Squibb, and Daiichi Sankyo outside the submitted paper. M.M. received honoraria for participation at advisory boards from Novartis, BMS, MSD, Pierre Fabre, and Sanofi; reports lecture fees from Novartis, BMS, MSD, Pierre Fabre, and Sanofi; and reports grants from Roche and Novartis. The remaining authors declare no competing financial interests.

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