



Edinburgh Research Explorer

Sensitivity and specificity of blood-fluid levels for oral anticoagulant-associated intracerebral haemorrhage

Citation for published version:

Almarzouki, A, Wilson, D, Ambler, G, Shakeshaft, C, Cohen, H, Yousry, T, Salman, RA, Lip, GYH, Houlden, H, Brown, MM, Muir, KW, Jäger, HR & Werring, DJ 2020, 'Sensitivity and specificity of blood-fluid levels for oral anticoagulant-associated intracerebral haemorrhage', *Scientific Reports*, vol. 10, no. 1. https://doi.org/10.1038/s41598-020-72504-7

Digital Object Identifier (DOI):

10.1038/s41598-020-72504-7

Link:

Link to publication record in Edinburgh Research Explorer

Document Version:

Publisher's PDF, also known as Version of record

Published In:

Scientific Reports

General rights

Copyright for the publications made accessible via the Edinburgh Research Explorer is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

Take down policy

The University of Edinburgh has made every reasonable effort to ensure that Edinburgh Research Explorer content complies with UK legislation. If you believe that the public display of this file breaches copyright please contact openaccess@ed.ac.uk providing details, and we will remove access to the work immediately and investigate your claim.



Download date: 16. Oct. 2020



natureresearch





OPEN Sensitivity and specificity of blood-fluid levels for oral anticoagulant-associated intracerebral haemorrhage

Abeer Almarzouki^{1,2}, Duncan Wilson², Gareth Ambler³, Clare Shakeshaft², Hannah Cohen⁴, Tarek Yousry⁵, Rustam Al-Shahi Salman⁶, Gregory Y. H. Lip^{7,8}, Henry Houlden¹⁰, Martin M. Brown², Keith W. Muir⁹, Hans Rolf Jäger⁵ & David J. Werring^{2⊠}

Intracerebral haemorrhage (ICH) is a life-threatening emergency, the incidence of which has increased in part due to an increase in the use of oral anticoagulants. A blood-fluid level within the haematoma, as revealed by computed tomography (CT), has been suggested as a marker for oral anticoagulantassociated ICH (OAC-ICH), but the diagnostic specificity and prognostic value of this finding remains unclear. In 855 patients with CT-confirmed acute ICH scanned within 48 h of symptom onset, we investigated the sensitivity and specificity of the presence of a CT-defined blood-fluid level (rated blinded to anticoagulant status) for identifying concomitant anticoagulant use. We also investigated the association of the presence of a blood-fluid level with six-month case fatality. Eighteen patients (2.1%) had a blood-fluid level identified on CT; of those with a blood-fluid level, 15 (83.3%) were taking anticoagulants. The specificity of blood-fluid level for OAC-ICH was 99.4%; the sensitivity was 4.2%. We could not detect an association between the presence of a blood-fluid level and an increased risk of death at six months (OR = 1.21, 95% CI 0.28-3.88, p = 0.769). The presence of a blood-fluid level should alert clinicians to the possibility of OAC-ICH, but absence of a blood-fluid level is not useful in excluding OAC-ICH.

Oral anticoagulant-associated intracerebral hemorrhage (OAC-ICH) is a devastating disease¹, with a reported 90-day case fatality of 42%²⁻⁴. The incidence of OAC-ICH is growing substantially, with increased use of anticoagulant therapy⁵. 5-12% of ICH is related to OAC⁶, and is expected to increase with an ageing population increasingly exposed to oral anticoagulants7. The risk of haematoma growth after OAC-ICH is as high as 54%8. Rapid identification of patients with OAC-ICH is important to allow rapid coagulation reversal, strict blood pressure management and transfer to a higher dependency unit⁹.

The identification of patients with OAC-ICH can be challenging, for example, when patients cannot communicate and there is no clear history from an informant. Computerized tomography (CT) remains the initial neuroimaging tool of choice for identification of acute ICH¹⁰. The presence of a blood-fluid level has been

¹Physiology Department, Faculty of Medicine, King Abdulaziz University, Jeddah, Saudi Arabia. ²Department of Brain Repair and Rehabilitation, UCL Stroke Research Centre, UCL Institute of Neurology and the National Hospital for Neurology and Neurosurgery, Russell Square House, 10-12 Russell Square, London WC1B 5EH, UK. ³Department of Statistical Science, University College London, Gower Street, London, UK. ⁴Haemostasis Research Unit, Department of Haematology, University College London, 51 Chenies Mews, London, UK. 5Lysholm Department of Neuroradiology and the Neuroradiological Academic Unit, Department of Brain Repair and Rehabilitation, UCL Institute of Neurology, Queen Square, London, UK. ⁶Centre for Clinical Brain Sciences, School of Clinical Sciences, University of Edinburgh, Edinburgh, UK. ⁷Liverpool Centre for Cardiovascular Science, University of Liverpool and Liverpool Heart and Chest Hospital, Liverpool, UK. 8Aalborg Thrombosis Research Unit, Department of Clinical Medicine, Aalborg University, Aalborg, Denmark. 9Institute of Neuroscience and Psychology, University of Glasgow, Queen Elizabeth University Hospital, Glasgow, UK. ¹⁰Department of Molecular Neuroscience, UCL Institute of Neurology and the National Hospital for Neurology and Neurosurgery, Queen Square, London WC1N 3BG, UK. [™]email: d.werring@ucl.ac.uk



Figure 1. Typical blood-fluid level on an acute CT scan of a patient with an intracerebral hemorrhage in the left parietal lobe.

suggested as a marker for OAC-ICH^{11,12}, but with the exception of one study (a sub-study of INTERACT-2) most published studies are case reports or were done on small samples¹¹⁻¹⁵. In a study of 2065 patients from the INTERACT-2 study, blood-fluid levels on baseline CT (found in 19 patients in the sample) were associated with the use of warfarin as well as poor outcome 90 days after ICH¹⁶.

In this study, we aimed to determine the prevalence, sensitivity and specificity of blood-fluid levels as a marker for OAC-ICH and its prognostic significance in a large multicentre prospective cohort of patients with ICH.

Methods

Participants. We included participants with CT-confirmed acute ICH (scanned within 48 h of symptom onset) recruited in the observational Clinical Relevance of Microbleeds in Stroke Study (CROMIS-2) conducted at 79 hospitals throughout the UK (and one in the Netherlands) between 2012 and 2015. The protocol has been published elsewhere¹⁷. Briefly, patients were eligible if they were 18 years or older and had a spontaneous ICH not secondary to major trauma or a macrovascular cause, as previously described¹⁸.

The CROMIS-2 study was approved by the UK National Health Service Research Ethics Committee and was carried out in accordance with the Declaration of Helsinki. Patients with capacity gave informed written consent. When patients could not consent, it was obtained in written form from a proxy (as defined by relevant local legislation).

Imaging. CT imaging was performed within 48 h of ICH onset in all patients. Digital CT images were collected in uncompressed Digital Imaging and Communication in Medicine (DICOM) format and analysed centrally. CT scans were examined for the presence of a blood-fluid level by two researchers (blind to anticoagulant status and clinical outcome). The raters were a neuroscience graduate student (A.A.) and a vascular neurologist (D.W.) with experience in neuroimaging research—A.A. performed the initial ratings, which were then checked and discussed with D.W.. As described previously¹¹, a blood-fluid level was defined according to the following features: (1) upper compartment hypodense to the brain, (2) lower compartment hyperdense to the brain and (3) a sharply defined horizontal interface between the upper and lower compartments (Fig. 1). ICH location was classified as infratentorial, deep, or lobar (cortical or cortical–subcortical) using a validated rating instrument¹⁹.

Baseline and follow up data. Data collected at baseline included age, sex, ethnicity, pre-morbid function assessed by the modified Rankin scale, clinical information, history of comorbidities (e.g., hypertension, atrial fibrillation), history of previous ischaemic stroke, ICH or TIA), and use of antithrombotic drugs prior to ICH. We recorded examination findings on admission, systolic and diastolic blood pressure, Glasgow coma scale score, the international normalized ratio (INR), and the use of surgical intervention. Mortality was assessed at six months post-ICH using information provided by the National Health Service digital data (Health and Social Care Information Centre)¹⁷.

Statistical analyses. We performed statistical analyses using SPSS version 24 and R version 3.4.3. We visually inspected the distribution of the data using histograms for continuous variables. We analysed data that were not normally distributed using appropriate non-parametric tests. The results of this study were expressed as

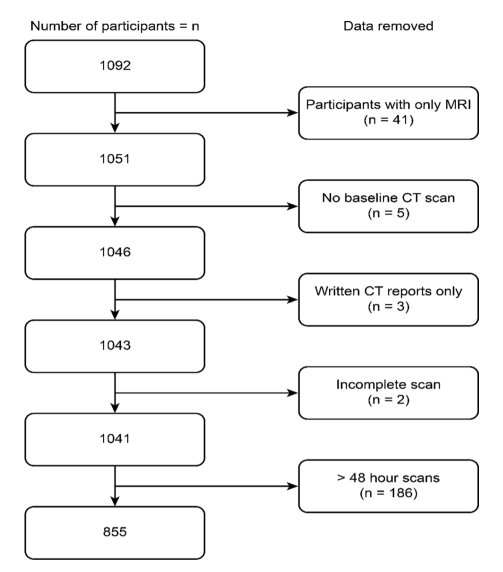


Figure 2. Flowchart showing the recruitment of patients for this study, including reasons for exclusion.

median and interquartile range (IQR) for continuous variables and as numbers and percentages (%) for categorical variables. We divided the sample into a blood-fluid level group and a non-blood-fluid level group. We tested the differences between the two groups using independent-sample Mann–Whitney U tests for continuous variables, while Fisher's exact test was used for categorical variables. The assumptions of these tests were checked and the corresponding effect sizes are reported. The effect sizes used were r for the Mann–Whitney U test (the Z-value divided by the square root of the total number of observations)²⁰ and the odds ratio for the chi-squared test

We calculated the sensitivity, specificity, and positive and negative predictive values of fluid level for anticoagulation (as well as their 95% confidence intervals; 95% CI). We tested the association between the anticoagulant status and CT-defined blood-fluid level using a chi-squared test. The association between the presence of a blood-fluid level and various clinical and imaging parameters was assessed using logistic regression. The association between blood-fluid level and six-month mortality was assessed using logistic regression with adjustment for prespecified clinically important variables known to be associated with outcomes: age, sex, premorbid mRS, ICH location, hematoma volume, intraventricular extension of the hemorrhage, and oral anticoagulant use. All model assumptions were checked. For all analyses, p < 0.05 (two-tailed) was considered significant.

Results

We included 857 patients with a median age of 75.5 years (IQR = 65.8–82.4), of whom 361 (42.1%) were female (Fig. 2). Anticoagulant status was unknown in six patients in the study sample, none of whom had a blood-fluid level. Of the 360 patients on anticoagulant therapy in the study sample, 20 were on a Factor Xa inhibitor, two were on Dabigatran, and the rest were on warfarin.

A blood-fluid level was demonstrated in eighteen patients (2.1%) by acute CT imaging (within 48 h of the ICH). Table 1 summarises the characteristics of the participants between the two groups. Compared to the

| aracteristic Total number n = 857 Fluid level | | Fluid level n = 18 | el n = 18 Non fluid level n = 839 | | |
|---|------------------|--------------------------------|-----------------------------------|--------|--|
| Age (years) (median, IQR) | 75.5 (65.8–82.4) | 79.5 (70.5–83.0) | 75.5 (65.6-82.3) | 0.2588 | |
| Male n, (%)* | 496 (57.9%) | 14 (77.8%) | 482 (57.4%) | 0.0957 | |
| White background n, (%) | 743 (86.7%) | 15 (88.2%) | 728 (86.8%) | 0.5657 | |
| Anticoagulants n, (%) | 360 (42.0%) | 15 (83.3%) | 345 (41.1%) | 0.0004 | |
| Comorbidities | | | | | |
| Atrial fibrillation n, (%) | 303 (35.4%) | 14 (77.8%) 289 (34.4%) | | 0.0008 | |
| Hypertension n, (%) | 566 (66.0%) | 12 (66.7%) | 554 (66.0%) | 1.0000 | |
| History of previous stroke | 1 | | • | | |
| IS n, (%) | 118 (13.8%) | 4 (25.0%) | 114 (13.6%) | 0.2647 | |
| ICH n, (%) | 35 (4.1%) | 2 (12.5%) | 33 (3.9%) | 0.1430 | |
| TIA n, (%) | 98 (11.4%) | 1 (6.7%) | 97 (11.6%) | 1.000 | |
| Medication on admission | 1 | | | | |
| Aspirin n, (%) | 189 (22.1%) | 1 (5.9%) | 188 (22.4%) | 0.1403 | |
| Warfarin n, (%) | 323 (37.7%) | 14 (77.8%) | 309 (36.8%) | 0.0008 | |
| Clinical measurements | | 1 | • | | |
| SBP (median, IQR) | 166 (147-190) | 160 (140-172) | 167 (147-190) | 0.1475 | |
| DBP (median, IQR) | 90 (79–105) | 83 (78.5-91.5) | 90 (79–105) | 0.2745 | |
| GCS (median, IQR) | 15 (14–15) | 15 (14–15) | 15 (14–15) | 0.8221 | |
| Blood results on admission | 1 | 1 | | | |
| INR (median, IQR) | 1.11 (1.00-2.40) | 3.05 (2.23-3.90) | 1.10 (1.00-2.30) | 0.0001 | |
| Management received | | 1 | | | |
| Surgery n, (%) | 29 (3.4%) | 2 (11.1%) 27 (3.2%) | | 0.1242 | |
| AC reversal n, (%) | 269 (31.4%) | 13 (72.2%) 256 (31.1%) | | 0.0005 | |
| Haematoma location** | | | | 0.0524 | |
| Lobar n, (%) | 296 (34.5%) | 10 (55.6%) 286 (34.1%) | | | |
| Deep n, (%) | 459 (53.6%) | 5 (27.8%) 454 (54.1%) | | | |
| Infratentorial n, (%) | 79 (9.2%) | 3 (16.7%) 76 (9.1%) | | | |
| Haematoma volume (median, IQR) | 7.1 (2.4–17.5) | 14.1 (5.5–23.7) 7.0 (2.3–17.2) | | 0.0451 | |
| Intraventricular extension n, (%) | 249 (29.1%) | 2 (11.1%) | 247 (29.4%) | 0.1345 | |
| Deceased n, (%) | 127 (14.8%) | 3 (20.0%) | 124 (14.8%) | 0.7312 | |

Table 1. Patient baseline and follow up characteristics in each group. *ICH* intracerebral haemorrhage, *TIA* transient ischemic attack, SBP systolic blood pressure, DBP diastolic blood pressure, GCS Glasgow coma scale, INR International normalized ratio, AC anticoagulant. *Note that data on sex was missing from one patient in the sample. *Note that data on hematoma location was missing from 23 patients in the sample. The differences between the groups were statistically significant (p<0.05) for the variables shown in bold. IS: ischemic stroke.

non-blood-fluid level group, the blood-fluid level group had a significantly higher INR (3.05 (2.23–3.9) versus 1.10 (1.00–2.30), p = 0.0001, r = 0.13). The blood-fluid level group were more often taking an anticoagulant (warfarin in all cases) at ICH onset than the non-blood-fluid level group (15 [83.3%] versus 345 [41%]; p = 0.0004). The blood-fluid level group also underwent OAC reversal more frequently, in 13 (72.2%) versus 256 (30.5%) (p = 0.0005) and had significantly more atrial fibrillation (14 [77.8%] compared to 289 [34.4%], p = 0.0008) in the non-blood-fluid level group. Overall, there was no statistically significant difference in death at six months post-ICH between the blood-fluid level group and the non-blood-fluid level group (3 [20.0%] vs. 124 [14.8%], p = 0.7312).

Blood-fluid level sensitivity and specificity to OAC-ICH. Of the patients in whom a blood-fluid level was seen (n = 18), the majority (15/18 (83.3%) were on anticoagulants (see Supplementary Table 1). The non-blood-fluid level group consisted of 839 patients. Less than half of this group was on anticoagulants (345/839, 41%). The presence of a blood-fluid level was significantly associated with the use of anticoagulants (χ^2 = 12.6, p < 0.001, odds ratio [OR] 7.1, 95% CI 2.0–38.6; p < 0.001).

The presence of a blood-fluid level identifies just 4% (95% CI 2–7%) of patients on anticoagulants (sensitivity). However, the absence of a blood-fluid level identifies 99% (95% CI 98–100%) of patients who are not on anticoagulants (specificity). Of all the patients with a blood-fluid level, 83% were on anticoagulants (positive predictive value, 95% CI 59–96%). The negative predictive value, the probability that patients with no blood-fluid level were not on anticoagulants, was 59% (95% CI 55–62%).

Haematoma characteristics of OAC-ICH associated with blood-fluid level on CT. Haematomas of the patients in the blood-fluid level group (median = 14.1, IQR = 5.5-23.7) were significantly larger than those of patients in the non-blood-fluid level group (median = 7.0, IQR = 2.3-17.2) (p = 0.045). Blood-fluid level-asso-

| | OR | 95% CI | p value | | |
|------------------------------|------|-----------|----------|--|--|
| Unadjusted model | | | | | |
| Blood-fluid level (yes) | 1.21 | 0.28-3.88 | 0.7690 | | |
| Fully adjusted model | | | | | |
| Blood-fluid level (yes) | 0.82 | 0.67-3.08 | 0.7852 | | |
| Age | 1.06 | 1.03-1.08 | < 0.0001 | | |
| Sex (female) | 0.97 | 0.61-1.54 | 0.8903 | | |
| Location (lobar) | 0.99 | 0.66-1.47 | 0.9549 | | |
| Pre-morbid MRS | 1.52 | 1.28-1.81 | < 0.0001 | | |
| Hematoma volume | 1.02 | 1.01-1.03 | < 0.0001 | | |
| Intraventricular extension | 2.13 | 1.32-3.43 | 0.0018 | | |
| Oral anticoagulant use (yes) | 1.84 | 1.18-2.91 | 0.0080 | | |

Table 2. Association between blood-fluid level and mortality. Reference categories are shown in parentheses for the categorical variables in the models.

ciated haematomas tended to have a lobar location (55.6%), but there were no statistically significant differences in haematoma location between the groups (p = 0.0524).

Factors associated with blood-fluid level in patients on anticoagulants. In a multivariate logistic regression including age (OR 1.00, 95% CI 0.95–1.97, p=0.867), sex (OR 2.68, 95% CI 0.70–13.38 , p=0.176), haematoma location (OR 0.93 , 95% CI 0.391–2.04 , p=0.858), haematoma volume (OR 0.99, 95% CI 0.96–1.02, p=0.775), intraventricular extension (OR 0.15, 95% CI 0.007–0.93, p=0.097), hypertension (OR 1.05 , 95% CI 0.31–4.34, p=0.944), and platelet count (OR 1.00, 95% CI 0.99–1.01, p=0.559), only the INR was significantly associated with the presence of a blood-fluid level in patients on anticoagulants (OR 1.58 , 95% CI 1.08–2.34 , p=0.0174).

Prognostic significance of blood-fluid level on mortality. Six hundred and ninety-nine patients (81.6%) had follow up at 6 months and complete data for the logistic regression analysis. In this analysis, the presence of a blood-fluid level was not significantly associated with higher odds of death at six months (OR 1.21, 95% CI 0.28–3.88, p=0.7690). In a multivariable analysis, older age, higher pre-morbid mRS, larger haematoma volume, intraventricular extension, and oral anticoagulant use were all associated with higher mortality (Table 2).

Discussion

In this study, we found a blood-fluid level in 18 out of 855 (2%) patients with acute ICH using CT imaging. A blood-fluid level was associated with the use of anticoagulants with a high specificity (99%) but low sensitivity (4%). Patients with a blood-fluid level also had larger ICH volumes but no evidence of higher mortality.

Our reported prevalence of a blood-fluid level is similar to that in a recent large ICH study that reported a prevalence rate of 1%¹⁶; however, no sensitivity level was reported in their study. In terms of diagnostic agreement, our results differed somewhat from those reported in previous studies. The sensitivity of the blood-fluid level for coagulopathy was found to be 59.4% in the study by Pfleger et al.¹¹ and 41.6% in the study by Gökce et al.¹², while it was considerably lower (4%) in our study. However, Pfleger et al.'s study did not group patients based on use of OACs, but based on the presence of coagulopathy in general. Pfleger et al.'s study included patients on warfarin, patients with liver failure, diffuse intravascular coagulation, and other causes of coagulopathy, the pathophysiology of which may be different to OAC-associated ICH. In addition, our study had a substantially larger sample size than previous studies and included patients on non-vitamin K antagonist oral anticoagulants, which may have influenced the sensitivity estimates.

Although there have been a few case reports of a blood-fluid level in patients with no underlying coagulopathy^{21–23}, our findings suggest that in acute ICH a blood-fluid level is highly specific for OAC-ICH (99%), consistent with previous clinical studies^{11,12,16}. We have also confirmed previous observations that a blood-fluid level is associated with larger haematoma volume^{11,12,16}, which is associated with worse clinical outcome²⁴. The presence of a blood-fluid level was not associated with mortality at six months in our study. This result suggests that the value of this sign is in diagnosing intracerebral hemorrhage associated with anticoagulants, but that it has no prognostic significance. This is at odds with the results from the INTERACT-2 data¹⁶, however, the relatively low prevalence of patients with blood-fluid level in both these studies may have led to spurious results. As previously suggested^{16,25}, a possible association between blood-fluid levels and increased mortality may be related to faster haematoma growth, but this needs to be further investigated in larger, longitudinal imaging studies.

Our study has strengths. We recruited a large cohort of ICH patients from 80 hospitals, so our findings should be widely generalisable. We used a strict definition of blood-fluid level, included only acute (<2 days) CT scans, and performed the blood-fluid level rating blinded to anticoagulant use and outcome. The main limitation of our study is the small number of participants in the blood-fluid level group, which limits its statistical power. Although our study suggests that higher INR is associated with a higher prevalence of blood-fluid level in patients

on anticoagulants, larger future cohorts are needed to investigate the relationship between INR and blood-fluid level in more detail, to determine whether the presence of a blood-fluid level occurs mainly with INRs outside the therapeutic range. One mechanism proposed to explain this is that, in the absence of normal blood clotting, red blood cells fall to the bottom of the haematoma, leaving plasma at the top, causing a sedimentation level. If this mechanism underlies blood-fluid levels, clotting might take time to eventually occur, meaning that the detection of this sign would be dependent on the time from ICH onset to imaging. This could not be investigated in the current study because the exact time from onset to imaging was not available, but future studies may shed light on this possibility. Finally, our study required consent from either the patient or a proxy, effectively excluding patients with large, clinically devastating ICH. Overall, this led to our included cohort being less mildly affected than a general ICH cohort.

The implications of this study are that the presence of a blood-fluid level on the CT scan of a patient with ICH should alert clinicians that the patient is likely to be taking an OAC. This is associated with a high risk of hematoma growth and mortality²⁶. In such patients, higher-level care, intensive blood pressure control, rapid coagulation assays, and reversal of anticoagulation might be appropriate^{27–30}. However, physicians must consider the small chance that, in rare cases, the presence of blood-fluid level may not be associated with OAC use. This is particularly important because blind reversal could lead to adverse events such as thrombosis³⁰.

In conclusion, the presence of a blood-fluid level on an acute CT scan is specific for anticoagulant treatment and is not associated with higher mortality at 6 months. Clinicians should be aware that patients with this sign are likely to be taking an OAC, which may help guide the acute management of such patients.

Data availability

Analyses for the CROMIS-2 study are ongoing; once all of these analyses are completed, the CROMIS-2 Steering Committee will consider applications from other researchers for access to anonymised source data.

Received: 16 February 2020; Accepted: 31 July 2020

Published online: 23 September 2020

References

- 1. Caceres, J. A. & Goldstein, J. N. Intracranial hemorrhage. Emerg. Med. Clin. N. Am. 30, 771 (2012).
- Dowlatshahi, D. et al. Poor prognosis in warfarin-associated intracranial hemorrhage despite anticoagulation reversal. Stroke J. Cerebral Circul. 43(7), 1812–1817 (2012).
- 3. Wilson, D. *et al.* Outcome of intracerebral hemorrhage associated with different oral anticoagulants. *Neurology* **88**, 1693–1700 (2017).
- 4. Inohara, T. et al. Association of intracerebral hemorrhage among patients taking non-Vitamin K antagonist vs vitamin K antagonist oral anticoagulants with in-hospital mortality. JAMA J. Am. Med. Assoc. 319, 463–473 (2018).
- 5. Flaherty, M. L. et al. The increasing incidence of anticoagulant-associated intracerebral hemorrhage. Neurology 68, 116-121 (2007).
- 6. Cervera, Á, Amaro, S. & Chamorro, Á. Oral anticoagulant-associated intracerebral hemorrhage. J. Neurol. 259, 212-224 (2012).
- 7. Veltkamp, R., Rizos, T. & Horstmann, S. Intracerebral bleeding in patients on antithrombotic agents. Semin. Thromb. Hemost. 39, 963–971 (2013).
- 8. Huttner, H. B. et al. Hematoma growth and outcome in treated neurocritical care patients with intracerebral hemorrhage related to oral anticoagulant therapy. Stroke 37, 1465–1470 (2006).
- 9. Kuramatsu, J. B. et al. Anticoagulant reversal, blood pressure levels, and anticoagulant resumption in patients with anticoagulation-related intracerebral hemorrhage. JAMA J. Am. Med. Assoc. 313, 824–836 (2015).
- 10. Heit, J. J., Iv, M. & Wintermark, M. Imaging of intracranial hemorrhage. J. Stroke Cerebrovasc. Dis. 19, 11 (2017).
- 11. Pfleger, M. J., Hardee, E. P., Contant, C. F. & Hayman, L. A. Sensitivity and specificity of fluid-blood levels for coagulopathy in acute intracerebral hematomas. *AJNR Am. J. Neuroradiol.* 15, 217–223 (1994).
- 12. Gökçe, E., Beyhan, M. & Acu, B. Evaluation of oral anticoagulant-associated intracranial parenchymal hematomas using CT findings. Clin. Neuroradiol. 25, 151–159 (2015).
- 13. Kase, C. S. et al. Anticoagulant-related intracerebral hemorrhage. Neurology 35, 943–943 (1985).
- 14. Weisberg, L. A. Significance of the fluid-blood interface in intracranial hematomas in anticoagulated patients. *Comput. Radiol.* 11, 175–179 (1987).
- 15. Qureshi, A. I. et al. Intensive blood-pressure lowering in patients with acute cerebral hemorrhage. N. Engl. J. Med. 375, 1033–1043 (2016).
- Sato, S. et al. Determinants and prognostic significance of hematoma sedimentation levels in acute intracerebral hemorrhage. Cerebrovasc. Dis. 41, 80–86 (2016).
- 17. Charidimou, A. *et al.* The clinical relevance of microbleeds in stroke study (CROMIS-2): Rationale, design, and methods. *Int. J. Stroke* 10, 155–161 (2015).
- 18. Wilson, D. *et al.* Volume and functional outcome of intracerebral hemorrhage according to oral anticoagulant type. *Neurology* **86**, 360–366 (2016).
- Charidimou, A. et al. The Cerebral Haemorrhage Anatomical RaTing inStrument (CHARTS): Development and assessment of reliability. J. Neurol. Sci. 372, 178–183 (2017).
- 20. Two-sample Mann-Whitney U Test. in R Handbook (ed. Mangiafico, S.) (2016).
- 21. Dublin, A. & Norman, D. Fluid-fluid level in cystic cerebral metastatic melanoma. J. Comput. Assist. Tomogr. 3, 650-652 (1979).
- 22. Iplikçioğlu, A. C. et al. A fluid level in an acute extradural haematoma. Neuroradiology 36, 31-32 (1994).
- 23. Jung, Y.-J., Kim, M.-S., Chang, C.-H. & Choi, B.-Y. The fluid-blood level in a spontaneous intracerebral hematoma case report. Korean J. Cerebrovasc. Surg. 13, 80–83 (2011).
- Lauer, A., Pfeilschifter, W., Schaffer, C. B., Lo, E. H. & Foerch, C. Intracerebral haemorrhage associated with antithrombotic treatment: Translational insights from experimental studies. *Lancet Neurol.* 12, 394–405 (2013).
- 25. Sato, S. et al. The speed of ultraearly hematoma growth in acute intracerebral hemorrhage of the intensive blood pressure reduction in acute cerebral. *Neurology* 83, 2232–2238 (2014).
- Cucchiara, B., Messe, S., Sansing, L., Kasner, S. & Lyden, P. Hematoma growth in oral anticoagulant related intracerebral hemorrhage. Stroke 39, 2993–2996 (2008).
- Aguilar, M. I. et al. Treatment of warfarin-associated intracerebral hemorrhage: Literature review and expert opinion. Mayo Clin. Proc. 82, 82–92 (2007).

- 28. Steiner, T., Rosand, J. & Diringer, M. Intracerebral hemorrhage associated with oral anticoagulant therapy. *Stroke* 37, 256–262 (2006).
- Siegal, D. M. & Crowther, M. A. Acute management of bleeding in patients on novel oral anticoagulants. Eur. Heart J. 34, 489–498 (2013).
- 30. Kahn, M. R., Amara, R. S. & Halperin, J. L. Risks to reversal of anticoagulation: The cardiology perspective. *Am. J. Gastroenterol. Suppl.* **3**, 22–28 (2016).

Acknowledgements

The CROMIS-2 study was funded by the Stroke Association and British Heart Foundation. This work was undertaken at UCLH/UCL who received a proportion of funding from the Department of Health's NIHR Biomedical Research Centres funding scheme. The authors thank the CROMIS-2 collaborators:

| John Aeron-Thomas |
|----------------------|
| Prasanna Aghoram |
| Elaine Amis |
| Peter Anderton |
| Sreeman Andole |
| Ijaz Anwar |
| John Bamford |
| Azra Banaras |
| Aian Barry |
| Ruth Bellfield |
| Aienne Benford |
| Ajay Bhalla |
| Maneesh Bhargava |
| Biju Bhaskaran |
| Neelima Bhupathiraju |
| Ekaterina Biggs |
| Jonathan Birns |
| Adrian Blight |
| Angie Bowring |
| Ellen Brown |
| David Bruce |
| Amanda Buck |
| Kerry Bunworth |
| Ilse Burger |
| Laura Burgess |
| Mathew Burn |
| Evelyn Burssens |
| Maudrian Burton |
| Nicola Butler |
| Denise Button |
| Sarah Caine |
| Michael Carpenter |
| Dinesh Chadha |
| Kausik Chatterjee |
| Lillian Choy |
| David Cohen |
| Bridget Colam |
| Lynne Connell |
| Martin Cooper |
| John Corrigan |
| Donna Cotterill |
| Gillian Courtauld |
| Linda Cowie |
| John Coyle |
| Susan Crawford |
| Claire Cullen |
| Krishna Dani |
| Amelia Daniel |
| Khaled Darawil |
| Kliaicu Dalawii |

| Prabel Datta Michelle Davis Nicola Day Sharon Dealing Mandy Doherty Deborah Dellafera Catherine Douglas Karen Dunne Collette Edwards Charlotte Edinton |
|--|
| Nicola Day Sharon Dealing Mandy Doherty Deborah Dellafera Catherine Douglas Karen Dunne Collette Edwards |
| Sharon Dealing Mandy Doherty Deborah Dellafera Catherine Douglas Karen Dunne Collette Edwards |
| Mandy Doherty Deborah Dellafera Catherine Douglas Karen Dunne Collette Edwards |
| Deborah Dellafera Catherine Douglas Karen Dunne Collette Edwards |
| Catherine Douglas Karen Dunne Collette Edwards |
| Karen Dunne Collette Edwards |
| Collette Edwards |
| |
| Charlotte Eglinton |
| Charlotte Eglinton |
| Abduelbaset Elmarimi |
| Hedley Emsley |
| Timothy England |
| Daniel Epstein |
| Renuka Erande |
| Bernard Esisi |
| Rachel Evans |
| Pamela arren |
| Pauline Fitzell |
| Glyn Fletcher |
| Rachel Gallifent |
| Rachel Gascoyne |
| Elio Giallombardo |
| |
| Bindu Gregary |
| Gunaratam Gunathilagan |
| Paul Guyler |
| Brigid Hairsine |
| Michael Haley |
| Anne Hardwick |
| David Hargroves |
| Frances Harrington Ahamad Hassan |
| Amanda Hedstrom |
| Melinda Holden |
| Clare Holmes |
| Senussi Hussein |
| Tanya Ingram |
| Sissi Ispoglou |
| Liz Iveson |
| Venetia Johnson |
| · · · · · · · · · · · · · · · · · · · |
| Frances Justin Shahid Kausar |
| Shahid Kausar Karen Kee |
| |
| Michael Keeling |
| Deborah Kelly |
| Shagufta Khan |
| Agnieszka Kieliszkowska |
| Hayley Kingwell |
| Vinodh Krishnamurthy |
| Sagal Kullane |
| Balakrishna Kumar |
| Geraldine Landers |
| Enas Lawrence |
| Simon Leach |
| Sana Leason |
| Paula Lopez |
| Caroline Lovelock |
| Robert Luder |

| Barbara Madigan |
|--------------------------------|
| Stuart Maguire |
| Holly Maguire |
| Karim Mahawish |
| Linetty Makawa |
| Maam Mamun |
| Dulka Manawadu |
| David Mangion |
| Aravindakshan Manoj |
| Syed Mansoor |
| Tracy Marsden |
| Rachel Marsh |
| Sheila Mashate |
| Michael McCormick |
| Clare McGoldrick |
| Madeleine McKee |
| |
| Emma Mckenzie |
| Sanjeevikumar Meenakishundaram |
| Zoe Mellor |
| Amulya Misra |
| Amit Mistri |
| Azlisham Mohd Nor |
| Mushiya Mpelembue |
| Peter Murphy |
| Arumug Nallasivam |
| Ann Needle |
| Vinh Nguyen |
| Janice O'Connell |
| Paul O'Mahony |
| James Okwera |
| Chukwuka Orefo |
| Peter Owusu-Agyei |
| Anthea Parry |
| Adrian Parry-Jones |
| Kath Pasco |
| Chris Patterson |
| Cassilda Peixoto |
| Jane Perez |
| Nicola Persad |
| Mia Porteous |
| Michael Power |
| Mathuri Prabhakaran |
| Christopher Price |
| Harald Proschel |
| Shuja Punekar |
| Janet Putterill |
| Marc Randall |
| Ozlem Redjep |
| * - |
| Habib Rehman |
| Emma Richards |
| Victoria Riddell |
| Christine Roffe |
| Gill Rogers |
| Anthony Rudd |
| Kari Saastamoinen |
| Mahmud Sajid |
| Banher Sandhu |
| Christine Schofield |
| |

| Jon Scott |
|----------------------|
| Lakshmanan Sekaran |
| Pankaj Sharma |
| Jagdish Sharma |
| Simon Sharpe |
| Matthew Smith |
| Andrew Smith |
| Nikola Sprigg |
| Julie Staals |
| Amy Steele |
| Gail Storey |
| Kelley Storey |
| Santhosh Subramonian |
| Appu Suman |
| Jane Sword |
| Grainne Tallon |
| Garryck Tan |
| Margaret Tate |
| Jennifer Teke |
| Natalie Temple |
| Teresa Thompson |
| Sharon Tysoe |
| Djamil Vahidassr |
| Anouk van der Kwaak |
| Roland Veltkamp |
| Deborah Walstow |
| Caroline Watchurst |
| Fran Watson |
| Dean Waugh |
| Peter Wilkinson |
| David Wilson |
| Jennifer Wilson |
| Sarah Wilson-Owen |
| Belinda Wroath |
| Inez Wynter |
| Emma Young |

Author contributions

A.A., D.W., T.Y., M.M.B., and D.J.W. conceived and designed the study. D.W., C.S., H.C., T.Y., H.H., M.M.B., K.W.M., H.R.J., and D.J.W. collected the data. A.A., D.W., and D.J.W. analyzed the data. A.A., D.W., G.A., R.A.S.S., G.Y.H.L., and D.J.W. interpreted the data. A.A., D.W., G.A., T.Y., R.A.S.S., G.Y.H.L., K.W.M., and D.J.W. wrote the manuscript. All authors critically revised the manuscript.

Competing interests

The authors declare no competing interests.

Additional information

Supplementary information is available for this paper at https://doi.org/10.1038/s41598-020-72504-7.

Correspondence and requests for materials should be addressed to D.J.W.

Reprints and permissions information is available at www.nature.com/reprints.

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2020