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The risk of deterioration in GCS13-15 patients with traumatic brain injury identified by CT imaging . A systematic review and meta-analysis.

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SCHOLARONE™ Manuscripts Dear Editor of The Journal of Neurotrauma,

Thank you for considering our manuscript and the time that Reviewer 1 has taken to review our manuscript and their useful comments. Our response to their individual points follows below.

Most studies of "mild TBI" currently do not refer to GCS because these truly mild cases almost always have a GCS of 15. Thus, GCS is eliminated as a measure of concussion or the severity of concussion. "Mild TBI" is an undesirable term because we do not know if the authors are referring to the whole range of patients with mild TBI which includes GCS of 13, 14 or 15. For this reason, MTBI is becoming an outmoded term because it encompasses a heterogeneous population ranging from those with focal neurological deficits which are clearly not "mild" and certainly not concussions, and those with no focal brain injuries which are concussions. Currently, the term concussion is preferred for brain injured patients with no focal neurological deficits who are almost always GCS 15. The admixture of GCS 13 and 14 makes this a very heterogeneous group. Since cases with GCS13, 14 or 15 are a heterogeneous group, the data must be looked at separately, as the authors have done in some of their analyses. Those with and without a normal GCS, in other words cases with GCS 13 and 14, should be analysed separately from GCS 15 cases. This paper provides proof that mild TBI is a heterogeneous mixture and should be avoided. They have done this for GCS from 14 to 15, in some of the figures, but why did they exclude GCS of 13? Studies without sufficient data to allow analysis of the effect of GCS should have been excluded.

We agree that the terminology used to categorise traumatic brain injury can be used inconsistently in the literature and in clinical practice. We agree that mild TBI refers to a spectrum of traumatically induced brain dysfunction in GCS13-15 patients, of which only a subset will have injuries identified by CT imaging. We have used the term "mild TBI" to refer to patients with brain injury who present to the Emergency Department with an initial GCS13-15. This is consistent with the definition of mild TBI described in the Reviewer's comments. We tried to make clear that our study population of interest is GCS13-15 patients, who are therefore defined as having mild TBI, with injuries identified by CT imaging. This is outlined in the first 3 lines of page 5. We believe the description of our population of interest as patients with "mild TBI" with injuries identified by CT imaging best defines the group in the absence of a better alternative. We have changed the title, paragraph 6 of the background and paragraph 3 of the section entitled context to try to further clarify that our study population of interest is mTBI patients with injuries identified by CT imaging.

We agree that this population is a heterogenous group with a range of characteristics that mean individual risk for adverse outcomes varies. Our findings suggest that despite being able to identify individual factors that affect risk in this group there currently is no risk model that using these or other factors can reliably identify low-risk patients. Initial GCS certainly represents one important factor that affects the risk of adverse outcomes in this group. We feel stratifying analysis by initial GCS would potentially lose important

information regarding how GCS and other risk factors interact, especially as older patients present with a higher GCS relative to the severity of their injury. We have added to paragraph 3 of the summary section of the discussion to highlight this point. Moreover, the vast majority of the studies that we identified did not stratify their analysis by the initial GCS of the study population and studies that attempted to derive prognostic models included GCS as a prognostic factor. Therefore, it is not possible to assess either outcomes or risk factor effect with only studies that would allow the separate analysis of different initial GCS populations without losing the majority of the study data we have identified. We have assessed the effect that an initial GCS of 15 has on the risk of adverse outcome using both stratification of outcomes by study GCS inclusion criteria (Fig 2 and Fig 5), meta-regression (Fig 4 and Fig 6) and pooling of within study estimates of the effect GCS (supplementary material 6). Figure 2 and Figure 5 include stratification of outcome prevalence by initial GCS 13-15, GCS 14-15 and 15 using study inclusion criteria.

Indeed the abstract indicates that after all their analysis they are saying that the only factors that indicate later deterioration are those with low initial GCS, advancing age and anticoagulation medication. Most clinicians in the field already know this. Then they conclude that research is needed to determine a usable clinical decision rule. In other words as a result of their study they found that there is no useful rule. It is not clear why they did not state a rule that patients with low GCS, advanced age or anticoagulation cannot be discharged from the ED and should be admitted for observation. Wasn't that the purpose of their study?-

We believe that what our study shows is that despite there being a large number of studies that have estimated the risk of adverse outcomes in the population of interest and some studies that have attempted to identify the factors that affect risk in this group, we cannot currently identify individual low risk patients that do not require hospital admission. Until a clinically useable validated multivariable prognostic model with sufficient sensitivity and specificity can accurately identify low-risk patients we believe that the risk of significant adverse outcomes in this group is sufficiently high that all patients in this group should be routinely admitted for observation. This position is outlined in the first 5 lines of page 21.

The exclusions are not clear. For example, did they exclude studies of patients who did not go through ED, and went directly to a hospital ward? Did they exclude patients who went to facilities not connected with a hospital. There was massive exclusion of studies. Case studies were automatically excluded. Why? There was one cohort study included. Why? To the authors are case and cohort studies synonymous? If so, then they should be consistent. They were critical of studies with "bias" and those not seen in emergency departments. Why? Why would head injured patients admitted directly to neurology, neurosurgery or anywhere else be considered a biased sample? Why are those seen in family doctors offices "biased", or remote nursing stations "biased"?

We agree that the explanation of the study exclusion criteria regarding the study setting could be clearer. The section in inclusion criteria entitled participants has now been amended to make it clear that only study participants who attended the ED or were admitted to an inpatient ward were included. The reason we have only included this population is because the study was aimed at informing clinicians evaluating patients in the ED about the potential risk of adverse outcomes in the GCS13-15 patients with brain injuries identified by CT imaging. Patients presenting in a different clinical setting to this may have a different risk profile and therefore conclusions drawn from them may be less applicable to the ED setting.

We feel that the nature of a systematic review means that study exclusion is determined by transparent and a prior defined criteria and that a large number of excluded studies may reflect a sensitive and well conducted search strategy. Our number of studies excluded following title and abstract screening and review of full studies is comparable to that of other systematic reviews including a previous systematic review of prognostic models in TBI that included 53 studies from 3354 studies identified by their search strategy.

Case studies were excluded as it would not be possible to estimate the study prevalence of the adverse outcomes of interest from single case studies or small case series. As indicated in supplementary material 4 all the studies included were cohort studies apart from a single small prospective trial.

The purpose of this study was to identify risk factors which could help clinicians decide whether a patient being evaluated in the ED requires a hospital admission. Therefore, if the patient population was drawn from a context in which patients were likely to have higher acuity injuries, such as patients selected for repeat CT imaging, then outcome estimates may not be as applicable. We agree that bias is not the correct term to describe the effect that different population selection has on outcome measures. The final sentence of the 4th paragraph of the abstract has been changed to reflect this. We do not believe that the use of bias in the rest of the main text refers to study population selection.

What % of cases had MR imaging, and why were they not analysed using normal vs abnormal MRI?

We intended that this study would help clinicians risk stratify patients using the initial CT scan and other patient factors available at presentation. Existing national guidelines including the UK NICE and SIGN guidelines, the Australian New South Wales Guidelines and the Canadian CT Head rule recommend initial CT imaging of head injured patients. We agree that MRI imaging may provide additional useful prognostic information but this may not be available to a clinician in the ED making a decision about whether patients in this group require hospital admission.

It is not clear why some of the focal lesions, especially extradural hematomas fail to make the list of reliable risk factors. We agree that our study indicates the type of focal lesion identified by CT imaging is an important risk factor for deterioration in this group. The 4th paragraph of the abstract and discussion summary section has been amended to highlight the importance the type of focal lesion has on the risk of the adverse outcomes of interest.

I am not sure why IMPACT was mentioned. It would be a completely inappropriate test for this group of patients.

IMPACT and other prognostic models derived in patients with more severe TBI were mentioned to illustrate that it has been possible to develop clinically useful prognostic models for the heterogeneous group of patients with more severe TBI. The 4th paragraph of the background has been amended to make it more explicit that these cannot be applied to the population of interest in this study.

The paper needs some editing for grammar and missing words including the abstract which contains a sentence without a verb.

Paragraph 3 and 4 of the abstract, the section entitled search methods for study identification, paragraph 2 of the section entitled quality assessment and paragraph 4 and 5 of the background have been amended.

For those unfamiliar with the methodology, terms should be more carefully described such as studies "were retrieved". What does this mean in plain language? Jargon such as this should be minimised to improve reader understanding.

We have replaced the term retrieved with selected in the paragraph entitled study selection to improve reader understanding.

The following sentence requires an explanation by the authors: "Factors potentially affecting the risk of adverse outcomes were considered if there were patient characteristics present at admission or available from initial investigations". There are multiple issues that they may have arbitrarily decided to exclude such as drug overdose, alcoholism, diabetes, etc.

This sentence has been amended to make it clear that any factor included in any of the studies providing it was present at admission was included in analysis. This would include drug overdose, alcoholism and diabetes.

"Neurosurgery" as an outcome measure is probably a poor term. Most clinicians regard "neurosurgery as a profession rather than an outcome measure. The performance of a neurosurgical procedure or the requirement for a neurosurgical operation would be better. We have replaced the term neurosurgery with neurosurgical intervention throughout.

, Wentz, R. and Roberts, I. (2006). System.

, BMC Med Inform Decis Mak 6, 38. We hope that we have adequately addressed the feedback and that the paper is now ready to be considered for publication.

Yours sincerely,

Carl Marincowitz

1. Perel, P., Edwards, P., Wentz, R. and Roberts, I. (2006). Systematic review of prognostic models in traumatic brain injury. BMC Med Inform Decis Mak 6, 38.

The risk of deterioration in CT identified GCS13-15 patients with traumatic brain injury identified by

<u>CT imagingmild Traumatic Brain Injur</u>y. A systematic review and meta-analysis.

Carl Marincowitz¹ MB BChir, MSc, BA

Fiona E. Lecky² MB Ch B, FRCS, DA, MSc, PhD, FCEM

William Townend³ MD FRCS FCEM

William.Townend@hey.nhs.uk

Aditya Borakati⁴ B<u>Sc</u>A

Andrea Fabbri⁵ MD

Trevor A. Sheldon⁶ MSc, MSc, DSc, FMedSci

- 1. **Corresponding Author**. Hull York Medical School, <u>Allam Medical Building</u>Hertford Building, University of Hull, Hull HU6, UK 7RX, Fax: +44 (0) 1482 464705 Tel +44 (0) 870 1245500 Email: carl.marincowitz@hyms.ac.uk
- 2. School of Health and Related Research, University of Sheffield, Regent Court, 30 Regent Street, Sheffield, S1 4DA, UK, Fax: +44 (0)114 222 0749 Tel: (+44) (0)114 222 4345, Email: f.e.lecky@sheffield.ac.uk
- 3. Emergency Department, Hull Royal Infirmary, Hull and East Yorkshire NHS Trust, Anlaby Road, Hull, HU3 2JZ, UK, Fax: (+44) (0) 1482 477857 Tel: (+44) (0) 1482 623065, Email:
- 4. Hull York Medical School, <u>Allam Medical Building</u>Hertford Building, University of Hull, Hull HU6, UK
- 7RX, Fax: +44 (0) 1482 464705 Tel +44 (0) 870 1245500 Email: hyab12@hyms.ac.uk
- 5. Head of Emergency Unit, Presidio Ospedaliero Morgagni-Pierantoni, AUSL della Romagna, via Forlanini 34, 47121 Forlì (FC), Italy Tel +390543735156, email: andrea.fabbri@auslromagna.it,

6. Department of Health Sciences, University of York, Alcuin Research Resource Centre Seebohm

Rowntree Building, Heslington, York, YO10 5DD, Tel +44 (0) 1904 321344, Fax: +44 (0) 1904 32 3433,

e-mail: trevor.sheldon@york.ac.uk

Abstract

The optimal management of mild traumatic brain injury (TBI) patients with injuries identified by CT brain scan is unclear. Some guidelines recommend hospital admission for an observation period of at least 24 hours. Others argue that selected lower-risk patients can be discharged from the Emergency Department (ED).

The objective was to estimate the risk of death, neurosurgical interventionery and clinical deterioration in mild TBI patients with injuries identified by CT brain scan, and assess which patient factors affect the risk of these outcomes.

A systematic review and meta-analysis adhering to PRISMA standards of protocol and reporting.

Study selection was performed by 2 independent reviewers. Meta-analysis using a random effects model was undertaken to estimate pooled risks of: clinical deterioration, neurosurgical interventionery and death. Meta-regression was used to explore between_-study variation in outcome estimates using study population characteristics.

Forty-nine primary studies and 5 reviews were identified that met the inclusion criteria. The estimated pooled risk of the outcomes of interest were: clinical deterioration 11.7% (95% CI: 11.7 to 15.8; neurosurgical interventionery 3.5% (95% CI: 2.2 to 4.9%); death 1.4% (95% CI: 0.8% to 2.2%). Twenty-one studies presented within_-study estimates of the effect of patient factors. Meta_regression of study characteristics and pooling of within_-study estimates of risk factor effect found the following factors significantly affected the risk of adverse outcomes: age; initial GCS; type of injury and anti-coagulation. The generalisability of mMany studies_-wasere limited significantly susceptible to bias due to population selection.

Mild TBI patients with injuries identified by CT brain scan have a small but clinically important risk of serious adverse outcomes. This review has identified <u>severalthe</u> prognostic factors;- rResearch is a Injury; Prognostic modelling; . needed to derive and a validate a usable clinical decision rule so that before low-risk patients can be safely discharged from the ED.

Keywords: Mild Traumatic Brain Injury; Prognostic modelling; Intra-cranial haemorrhage; Minor Head Injury.

Background

There are 1.4 million annual attendances in England and Wales to Emergency Departments (EDs) following a head injury (any trauma to the head), and in 2010 2 .5 million people were treated for traumatic brain injury (TBI- injury to the brain or alteration of brain function due to an external force) in the United States. Approximately 95% of patients have an initial Glasgow Coma Scale (GCS) of 13-15, out of a possible 15, indicating normal or mildly impaired responsiveness and orientation. In this large group with head injury and a high conscious level at presentation research has focused on developing decision rules to identify patients who require computed tomography (CT) imaging due to their risk of life threatening traumatic brain injury (TBI).

In the United Kingdom (UK), National Institute for Health and Care Excellence (NICE) and Scottish Intercollegiate Guidelines Network (SIGN) guidelines are used for this risk assessment, based on the Canadian CT head Rule (CCHR).^{1,3,4} Only 1% of head injured patients have life threatening TBI.^{1,4} However, 7% have TBI identified by CT imaging.⁵

Most TBI patients who require neurosurgical interventionery are identified soon after presentation. The optimal management of the remaining patients in this group remains controversial. A proportion will deteriorate due to the progression of their injuries and so some studies advocate admission to higher dependency levels of care and repeat CT imaging. 6,7

Others studies report that some low risk patients may be safely discharged after a short period of observation in the ED.^{8, 9} Perel et al have previously outlined how prognostic models can aid clinical decision making in TBI.¹⁰ Subsequent prognostic models, including the IMPACT, TARN and CRASH models,—have been useful in predicting adverse outcomes in patients with more severe TBI, but they are not applicable to this patient group are not applicable to this group due to the exclusion of GCS15 patients.¹¹⁻¹³ Equivalent prognostic models for GCS13-15 patients with CT identified TBI may help safely reduce hospital admissions.

This review is the first to give an overview of the risk <u>of adverse outcomes and prognostic factors</u>

<u>inthat</u> patients with mild TBI <u>(-that is-a high or normal conscious level with traumatically induced brain dysfunction) and injuries identified by CT brain scan—and injuries identified by CT brain scan have of adverse outcomes and which patient factors are prognostic. The review specifically:</u>

- (i) Estimates the overall risk of adverse outcomes in patients who are initially GCS13-15 in the ED when traumatic brain injury is identified by CT imaging.
- (ii) Assesses which prognostic factors affect the risk of deterioration and other clinically important outcomes in this population.

Methods

A systematic review was conducted using the PRISMA P protocol and is reported in accordance with PRISMA guidelines. ¹⁴ The review is registered with the PROSPERO prospective register of systematic reviews and the protocol is available at

http://www.crd.york.ac.uk/PROSPERO/display record.asp?ID=CRD42016051585.

Inclusion Criteria:

Participants

Patients aged ≥12 years with an initial GCS of 13-15 with TBI identified by CT imaging. TBI included any traumatic: extradural haemorrhage, subdural haemorrhage, intra-cerebral haemorrhage, subarachnoid haemorrhage, cerebral contusion, or skull fracture. Studies had to be conducted in the context of an emergency hospital attendance including a presentation to the ED or during admission to an inpatient ward.

Prognostic factors

Factors potentially affecting the risk of adverse outcomes were considered if they were included in analysis if they were patient factors present at admission including: demographic characteristics,

comorbidities, medication use, symptoms, other clinical features patient characteristics present at admission or available from initial investigations.

Outcome measures

Primary outcomes: death, neurosurgical interventionery or any other measure of clinical deterioration such that admission to hospital was warranted.

Secondary outcome: progression of TBI on repeat CT imaging.

Types of study design

All studies, other than case studies, were included.

Search methods for study identification:

Studies published before 1996 were excluded due to more liberal use of CT imaging to diagnose TBI after this date.⁵

The following electronic databases were searched with results restricted to English language studies:

- EMBASE (via OVID) searched 24/11/2016 1996 to 2016 Week 47
- MEDLINE (R) (via OVID) searched 24/11/2016 1996 to November Week 3 2016
- CINHAL plus (via EBSCO) searched 24/11/2016 1983 to 2016
- Cochrane Central Register of Controlled Trials (CENTRAL); The Cochrane Library 2016 all available dates. Accessed 24/11/2016

The full search strategy is reported in supplementary material 1.

The reference and citation searches of several national guidelines, reports and reviews included:

NICE, SIGN and Australian New South Wales (NSW) guidelines, National Institute for Health Research
(NIHR) Health Technology Assessment of management strategies for minor head injury, the results
of the World Health Organisation (WHO) Collaboration on prognosis in mild traumatic brain injury,
systematic reviews assessing prognostic factors in traumatic brain injury, and systematic reviews

assessing the utility of repeat CT imaging in minor head injury. ^{1, 3, 10, 15-17} ¹⁸ ^{19, 20} All included studies references and citations were searched.

The Trauma Audit and Research Network (TARN) listed publications were searched via the TARN website: https://www.tarn.ac.uk/Content.aspx?ca=9&c=70 (accessed 10/3/2017).

Data Management and Extraction:

Identified studies were stored in EndNote X8 and duplicates removed.

Study Selection

Two reviewers (CM and AB) independently completed title and abstract screening. Full reports of any studies that potentially met the inclusion were <u>selected and assessed retrieved</u>. These were screened and studies that did not meet the inclusion criteria were discarded with documented reasons. Disagreements were resolved through discussion or arbitration by a 3rd reviewer (TS).

Data Extraction

The following data were extracted using a pre-piloted data extraction tool: study population and demographics, sample size, outcomes assessed, prognostic factors assessed, whether univariable or multivariable modelling had been undertaken and the overall results of the study. The selection criteria of studies were recorded to assess whether sub-populations with different risk profiles had been studied. The data extracted is presented in supplementary material 2.

Assessment of the risk of bias

The Quality in Prognostic Studies (QUIPS) Tool was used to assess the quality of included studies particularly for the risk of bias.²¹ Six domains were assessed: study participation; study attrition; prognostic factor measurement; outcome measurement; study confounding; and statistical analysis and reporting.

Data Analysis

Three forms of analysis were undertaken: pooling of adverse outcomes reported in studies, identification of risk factors by exploration of between-study variation in outcomes by study characteristics and a synthesis of common risk factors assessed within studies.

A pooled prevalence of the adverse outcomes of interest and confidence intervals for individual studies were estimated using the Metaprop function (STATA-SE 14).²² The Freeman-Tukey double arscine transformation was used to include studies with no adverse outcomes and a random effects model was used due to study heterogeneity.²³

Between-study heterogeneity estimates of outcomes was explored using subgroup analysis. Meta-regression of study characteristics was used to identify factors that affected the risk of the outcomes of interest. Meta-regression of multiple study characteristics' effect on the prevalence of adverse outcomes was assessed using the Metareg function (STATA-SE 14) with weighting incorporating a measure of between study variation (tau2).^{24, 25} The log odds of clinical deterioration, neurosurgeryneurosurgical intervention and death were assessed as dependent variables and the standard error of the log odds was used to approximate the within study standard error. To account for studies with no outcomes, 0.5 was added to both the outcome estimates and the sample size (consequently, in graphic representations of the meta-regression the estimated risk can only tend towards zero).

Where studies had assessed the effect of risk factors on the outcomes of interest using individual data, analysis was categorised as univariable or multivariable. Univariable meta-analysis of prognostic factor effect estimates reported in primary studies was completed using Review Manager 5.3 where possible. A Random Effects model was used due to the heterogeneity of study populations, prognostic factor and outcome measures. Meta-analysis of multivariable models was not possible due to limited numbers and variation in outcome and prognostic factor measurement.

Results

Search Result

The electronic search strategy was completed on the 24/11/2016 and identified 4665 studies. Of these 412 were duplicates, leaving 4253 studies for title and abstract screening (Fig. 1). Following title and abstract screening 69 studies^{6, 9, 27-93} and 2 reviews^{19, 20} were retrieved. A "grey" literature search identified a further 129 studies for title and abstract screening of which 3 were retrieved. Reference and citation searching of included studies and selected reviews and guidelines identified another 46 studies^{7, 8, 39, 97-139} for full retrieval and 3 additional systematic reviews^{17, 18, 140} for reference and citation searches.

In total 118 primary studies and 5 systematic reviews were retrieved.

Study Selection

Forty-nine primary studies met the inclusion criteria. ^{6-9, 27, 28, 30, 32, 37, 41, 42, 52, 54, 55, 57, 59, 60, 62, 63, 65, 66, 69, 71, 73-78, 86, 87, 90, 93, 97-104, 106-109, 114, 125, 130, 139 One review presented new study data. ¹⁸ The 4 remaining reviews formed part of the narrative synthesis. ^{17, 19, 20, 140} The reasons for excluding the remaining 69 studies are presented in supplementary material 3. Anonymised individual patient data were provided by the authors of a cohort study to allow outcomes for initial GCS13-15 patients to be calculated, so this study is included. ¹³⁹}

Study Characteristics

Supplementary material 4 presents the characteristics of included studies. Seven prospective studies were identified ^{28, 66, 74, 75, 90, 114, 139} and 4 studies had a sample size of over 1000. ^{63, 87, 98, 108} Forty-six studies estimated the outcomes of interest and contribute to pooled estimates of risk. ^{6-9, 27, 28, 30, 32, 37, 41, 42, 52, 54, 55, 57, 59, 60, 62, 63, 65, 66, 69, 71, 73-78, 86, 87, 90, 93, 97-104, 106-109, 114, 125, 130, 139} Four studies present data regarding specific injury sub-types. ^{32, 55, 71, 103} One study only contributes to the narrative synthesis

due to the outcome measure it assessed.⁴² Three studies present the Brain Injury Guidelines (BIG) risk stratification tool.^{9, 27, 109} As this tool was applied to all TBI patients and initial GCS forms part of risk stratification, these studies contributed to the narrative synthesis.

Twenty-one studies present either univariate or multivariable analysis assessing prognostic factors' effect on the outcomes of interest. 6, 37, 41, 54, 55, 66, 69, 71, 73-78, 87, 98-101, 130, 139 Sixteen studies present multivariable models using logistic regression or recursive partitioning. 6, 37, 41, 54, 55, 66, 69, 71, 73, 74, 77, 78, 98, 100, 101, 130 Only 2 studies attempted to validate such models by splitting the study data sets. 66, 98

Quality Assessment

QUIPS quality scores are presented in supplementary material 2.²¹ The following common methodological issues were identified.

Study recruitment was often was not representative of all GCS 13-15 patients with TBI identified by CT imaging. Sixteen studies that contribute to the pooled estimates of adverse outcomes only included patients that had undergone repeat CT imaging and so are likely to represent a higher risk population. 7, 18, 54, 74-78, 86, 90, 102, 104, 106, 107, 125, 130 Even when re-imaging was presented as routine practice, it was often indicated that not all patients were re-imaged and included in analysis. Many other studies excluded higher risk anti-coagulated patients or those with more severe injuries.

Prognostic factor measurement was not consistent. Continuous variables were dichotomised at different thresholds or the same risk factor was measured with different methods. For example, the

severity of injury identified by CT imaging was assessed with 10 different measures. Most studies were retrospective and reliant on the accuracy of case notes and radiological reports. The small sample size of many studies prevented multivariable modelling with all variables identified in univariable modelling as affecting deterioration.³⁷

In 32 studies outcomes were assessed during inpatient admission and so patients who were discharged and deteriorated were missed. In other studies, is wasn't clear when outcome measures were assessed. Eight different measures of clinical deterioration were used in 18 studies.

Several studies included patients with extra-cranial injuries and significant comorbidities. Extra-cranial injuries caused clinical interventions, and in studies that measured deterioration in this way this was a potential source of bias. ⁶⁶ Other studies indicated some recorded deaths were related to comorbidities instead of TBI. ^{41, 73}

Risk of Adverse Outcomes and Exploration of Between_-Study Variation

Death

Twenty-seven studies assessed the outcome of death. ^{6, 8, 28, 41, 52, 57, 60, 62, 63, 65, 69, 73-75, 78, 86, 93, 97, 99-102, 104, 114, 125, 130 139} The estimated risk of death for these studies ranged between 0 and 6% (median 1.1%), and with a pooled prevalence of 1.4% (95% CI: 0.8% to 2.2%) (Fig. 2). Studies that selected only initial GCS15 patients had a pooled estimate of mortality of 0.03% (95% CI: 0 to 0.28%). Studies that selected populations for non-ICU admission or other conservative care pathways had an estimated prevalence of death of 0.1% (95% CI: 0 to 0.6%).

The effect on mortality of mean GCS, average age and selection of study population for a lower level of care was explored using meta-regression. Increased age of study population was associated with a higher risk of death (1.05 95% CI: 1.00 to 1.12) (Fig. 3). Whilst higher study population GCS was associated with a lower risk of death (0.12 95% CI: 0.02-0.86) (Fig. 4). The percentage of patients taking anticoagulants in studies was not associated with the prevalence of death (1.05 95% CI: 0.95-1.17), but selection for a lower level of care compared to a higher level of care was (0.27 95% C.I. 0.08-0.94). When average age of the study population and mean study GCS were assessed in a multivariable model they remained statistically significant predictors of mortality (Table 1), with an

adjusted R squared of 38%, indicating that these 2 factors explained over a third of the variation in study estimates.

Neurosurgery Neurosurgical intervention

Thirty-six studies reported neurosurgical outcomes. ^{6-9, 27, 30, 37, 52, 54, 57, 60, 62, 63, 65, 66, 73-78, 86, 90, 93, 97-102, 104, 106, 109, 114, 125, 130, 139} Figure 5 presents the estimates of the proportion of patients that underwent a neurosurgical procedure stratified by the GCS inclusion criteria. Reported neurosurgical intervention prevalence ranged between 0 and 26% (median 3.1%). The high proportion requiring neurosurgeryneurosurgical intervention reported by Beynon et al⁹³ may reflect the greater use of anticoagulants or anti-platelets (33/70 participants).

The pooled estimated neurosurgical intervention risk was 3.5% (95% CI: 2.2 to 4.9%). An I² of 96.4% indicated considerable heterogeneity. Studies conducted on initial GCS 15 patients had a lower prevalence of neurosurgeryneurosurgical intervention: 0.2% (95% CI: 0 to 0.5%). Sensitivity analysis of selection of the study population for reduced care, such as discharge, a non-ICU admission or non-routine repeat CT imaging found the pooled estimate of neurosurgeryneurosurgical intervention in these studies to be 0.1% (95% CI: 0 to 0.5%).

The of result of meta-regression using: mean study population GCS, mean study population age, anticoagulation and selection of study population for non-ICU admission or other reduced care pathways is shown in Figures 6,7,8 and Table 1. Increasing age (1.01 95% CI: 1.02 to 1.11) and increasing percentage of study population taking anti-coagulants (1.1 95% CI: 1.01 to 1.19) was associated with a higher risk, whilst an increasing GCS (0.71 95% CI:0.01 to 0.56) was associated with a lower risk, of neurosurgeryneurosurgical intervention.

Fig. 7 shows a cluster of 4 small studies with low mean ages that appear to have a disproportionately low estimated prevalence of neurosurgeryneurosurgical intervention. 8, 52, 62, 106 This is explained by:

exclusion of anti-coagulated patients, ^{8, 52, 62} selection of patients for non-ICU admission or other reduced other care pathays, ^{8, 52, 62} and exclusion of patients with large injuries⁸.

When the effect of population selection for reduced clinical management, exclusion of anticoagulated patients (only 23/36 studies reported percentage of anti-coagulated patients), mean age and GCS of the study population were all included in a meta regression, age and GCS were the only statistically significant predictors of neurosurgeryneurosurgical intervention (Table 1). The adjusted R squared of the model was 48%, indicating that these factors accounted for almost half of between study variation.

Clinical Deterioration

Eighteen studies measured prevalence of clinical deterioration. 8, 37, 41, 63, 66, 69, 73, 74, 76-78, 100, 101, 104, 107, 108, 114, 125 The estimated risk of deterioration ranged between 0 and 24.5% (median 12.8%). Figure 9 presents study estimates of the percentage of patients that deteriorated, with 95% confidence intervals and stratified by how the outcome was assessed. A pooled prevalence of 11.7% (95% CI: 8.21 to 5.8%) for some form of clinical deterioration was estimated with an 12 of 95.7%.

Estimates were stratified by: initial GCS of patients, whether the included population were all selected for repeat CT imaging, the inclusion of anticoagulated patients, the follow up period and exclusion of patients with extra-cranial injuries. None of these factors reduced the observed between study heterogeneity.

The effect of: mean GCS study population, mean age study population, study population selection, exclusion of patients with extracranial injuries, and exclusion of anti-coagulated patients was explored using meta-regression. As only 18 studies measured this outcome the model was restricted to 2 variables. No factor assessed individually or in conjunction with another factor was found to statistically affect the risk of clinical deterioration. Higher age and lower GCS were non-statistically associated with a higher risk of clinical deterioration (Table 1).

Progression Repeat CT imaging:

Twenty-six studies assessed the outcome progression of the initial injury on repeat CT imaging. ^{6, 18, 127, 28, 30, 41, 62, 74-78, 87, 90, 97, 99-102, 104, 106-108, 114, 125, 130} The prevalence of this outcome in these studies is presented in Figure 10, stratified by whether studies only included patients that had undergone repeat CT imaging. The pooled estimate for this outcome was 15.6% (95% CI: 11.3 to 20.4%). There is a high degree of heterogeneity with a range in risk of progression between 2% and 48% (median 36.5%) and I²=97%. The non-statistically significant higher pooled risk in studies that included only patients that had undergone repeat CT imaging probably reflects selection of higher risk patients to repeat imaging. Subgroup analysis of study characteristics did not find any factors that accounted for the heterogeneity. This is probably the result of different criteria used to triage patients to repeat CT imaging and definition of progression of injury.

Prognostic Factors Assessed in Primary Studies

Twenty-one studies presented within study estimates of effect of individual risk factors on the outcomes of interest (supplementary material 4) and the factors assessed are presented in supplementary material 5. 6, 37, 41, 54, 55, 66, 69, 71, 73-78, 87, 98-101, 130, 139 The most influential factors were: age; initial GCS; severity of CT finding; type of injury; anti-coagulation; and anti-platelet medication (Table 2). Individual forest plots are presented in supplementary material 6.

Age

Age was evaluated as a factor in prognostic modelling in 18 primary studies. ^{6, 37, 41, 54, 55, 66, 69, 71, 73, 74, 76-78, 98-101, 130} Ten studies ^{37, 41, 54, 66, 73, 74, 76-78, 101} assessed age using 4 different dichotomous cut offs and 11 studies measured age as a continuous factor. ^{6, 55, 69, 71, 73, 76, 77, 98-100, 130} Multivariable models included: logistic regression with age either a dichotomised or continuous variable, or decision tree analysis.

Of these 18 studies: six assessed the outcome of clinical deterioration; 8 assessed the outcome of neurosurgeryneurosurgical intervention; 1 measured death as an outcome; and 8 studies evaluated progression of injury on repeat CT imaging. Despite being the most commonly assessed prognostic factor, due to the variation in measurement and the outcomes assessed, it was not possible to undertake a pooled analysis.

Increased age was associated with an adverse outcome in 9 of the 19 univariable models presented. Age was a significant predictor of an adverse outcome in 2 of 5 multivariable models where it was treated as a continuous variable. ^{69, 71, 98, 130} However, in 4 of 6 multivariable models where it was dichotomised, older age predicted the outcomes of interest. ^{41, 54, 66, 73, 78, 101} This may indicate a non-linear relationship with older age groups having a disproportionately higher associated risk of adverse outcomes.

Initial GCS

Twelve primary studies presented within study estimates of the effect of initial GCS on the risk of the outcomes of interest. 6, 37, 41, 55, 66, 69, 73, 74, 77, 98, 100, 101 Univariable effect estimates of initial GCS 15 were pooled for studies assessing clinical deterioration and neurosurgeryneurosurgical intervention as an outcome with individual patient data provided by Fabbri et al and an initial GCS=15 was protective against clinical deterioration or neurosurgeryneurosurgical intervention (pooled OR 0.35 95% CI: 0.23 to 0.53) (Table 2). 37, 41, 66, 73, 74, 77, 101 Two papers assessed progression of injury on repeat CT imaging and both found initial GCS 15 to be associated with reduced risk of progression. Four studies estimated the effect of an initial GCS of 15 in multivariable models. All 4 multivariable models found initial GCS15 to be associated with a reduced risk of adverse outcomes.

Severity of Injury as assessed by CT findings

Nine studies estimated whether the severity of injury identified by initial CT scan predicted adverse outcomes. ^{6, 41, 54, 55, 66, 73, 76, 78, 100} This was assessed by: the presence of midline shift or mass effect in 5

studies, ^{6, 55, 66, 76, 100} the Marshall classification in 2 studies, ^{41, 73} and measures of haemorrhage thickness or volume in 4 studies. ^{54, 55, 78, 100} The variability in the measures of injury severity and differences in the outcomes assessed prevented pooling.

All studies that assessed presence of midline shift/mass effect found it to be statistically predictive of adverse outcomes. This association remained in the 2 studies that presented multivariable analysis.^{6,}

The Marshall classification was assessed as a continuous⁷³ and dichotomised variable⁴¹ and neither study found a statistically significant association with adverse outcomes.

The 2 studies which assessed the effect of bleed thickness>10mm found this to be statistically predictive of either progression of injury on repeat CT imaging or neurosurgeryneurosurgical intervention in both uni and multivariable analysis. 54, 78

Isolated subarachnoid haemorrhage

Twelve studies presented outcomes for populations with isolated injuries and patients with isolated subarachnoid haemorrhages (iSAH) were the lowest risk for adverse outcomes:

neurosurgeryneurosurgical intervention pooled risk 0.01% (95% CI: 0 to 0.7%) (Fig. 11), and 1.1% (95% CI: 0 to 5.5%) pooled prevalence of clinical deterioration (supplementary material 7). 32, 37, 55, 59, 71, 74, 77, 98, 99, 103, 107, 108

Univariable effect estimates presented in the 2 studies that assessed the effect of the presence of iSAH were pooled with data extracted from 3 additional studies. ^{37, 73,77, 98, 108} The pooled estimate indicated iSAH reduced the risk of neurosurgeryneurosurgical intervention/clinical deterioration (Table 2).

Two multivariable models included iSAH as a prognostic factor. One found iSAH to be associated with a lower risk of clinical deterioration.³⁷The other found iSAH to have no effect on risk.⁹⁸

Isolated extradural haemorrhage

Patients with isolated extradural haemorrhage had the highest risk of neurosurgeryneurosurgical intervention: 13.7% (95% CI: 9.3% to 18.5%) (Fig. 11). 18.5% is estimated from a population of all initial GCS14-15 patients with extradural haemorrhage, whilst the estimates in the other studies are from populations that have been selected for more conservative management. 77, 98, 107, 108

Three studies assessed isolated extradural haemorrhage as a prognostic factor. ^{37, 73, 98} A pooled risk estimate for clinical deterioration or neurosurgeryneurosurgical intervention using these 3 studies and outcome data extracted from a further 2 studies, ^{77, 108} found isolated extradural haemorrhage to be associated with these outcomes (OR 2.26 95% CI: 1.9 to 2.68) (Table 2). Isolated extradural haemorrhage remained statistically associated with neurosurgical outcomes in the only multivariable model that included this factor. ⁹⁸

Anti-coagulation

Twelve studies estimated the prognostic effect of anti-coagulation. ^{6, 37, 41, 55, 74, 76-78, 98, 100, 101, 139}

Measures of anti-coagulation included: any documented coagulopathy, ^{6, 41, 55, 77, 98, 100} pre-injury warfarin use, ^{37, 76, 101} warfarin or antiplatelet therapy as a combined risk factor, ^{78, 100} and continuous laboratory measures of anti-coagulation. ^{6, 74, 101}

Univariable effect estimates of dichotomous measures of anti-coagulation were pooled with individual patient data from Fabbri et al for the composite outcome of clinical deterioration or neurosurgeryneurosurgical intervention (Table 2), pooled estimate: OR 1.45 95% CI: 1.28 to 1.64.

Two studies presented multivariable models that included anti-coagulation and it was not statistically associated with the outcomes of interest in either model. 78, 98

Anti-platelet medication

The effect of anti-platelet use was evaluated by: aspirin use,^{37, 76, 101} clopidogrel use,^{37, 76, 101} and a joint measure of antiplatelet use.^{55, 66, 87} No multivariable models included antiplatelet use. Pooled univariable risk estimates of pre-injury aspirin and clopidogrel use are presented in Table 2. Meta-

analysis indicated a statistical association between clopidogrel with clinical deterioration or neurosurgeryneurosurgical intervention
but no association between aspirin use and this outcome.

Discussion:

Summary

We have completed a thorough systematic review and meta-analysis to identify risk factors for adverse outcomes in this TBI population. This is the first review to provide pooled estimates of clinically important outcomes in this population and identify which factors affect the risk of these outcomes.

The pooled prevalence of adverse outcomes were: 11.7% (95% CI: 8.21 to 5.8%) clinical deterioration, 3.5% (95% CI: 2.2 to 4.9%) neurosurgeryneurosurgical intervention, and 1.4% (95% CI: 0.8% to 2.2%) death. These outcome estimates used a pooled total of 65724 patients and are comparable to the 2.7% craniotomy rate reported for a similar population in a national UK trauma database. The variation in individual study outcomes reflects differences in populations studied and outcome definitions. For the outcomes of neurosurgeryneurosurgical intervention and death heterogeneity could be explained by the age of study populations and different study population GCS scores.

Risk factors for adverse outcomes were identified using both meta-regression of study characteristics and synthesis of prognostic models presented by primary studies. Age, anti-coagulation and initial GCS were found by both methods to affect risk. An increase in mean study population age by 1 year was associated with increased odds of neurosurgeryneurosurgical intervention of 1.09 in multivariable meta-regression (Table 1) and age was a predictor of an adverse outcome in 6/11 multivariable models presented in primary studies. In univariable meta-regression a unit increase in the percentage of the study population taking anti-coagulants was associated with a 1.1 increase in the odds of neurosurgeryneurosurgical intervention (Table 1). Pooling of univariable

models presented in primary studies found anticoagulated patients to have odds 1.45 time greater than patients not anticoagulated for neurosurgeryneurosurgical intervention/clinical deterioration (Table 2). In multivariable meta-regression, a unit increase in mean/median study population GCS was associated with an 0.12 reduction in the odds of neurosurgeryneurosurgical intervention (Table 1). Pooling of univariable models indicated that patients with initial GCS
1). Pooling of univariable models indicated that patients with initial GCS
1). In multivariable meta-regression patients that presented with an initial GCS of 15 lower GCS scores (Table 2). In multivariable meta-regression models including both initial GCS and age, initial GCS had a smaller effect on the risk of either neurosurgical intervention or death than in univariable analysis and this may be due to older patients presenting with higher initial GCS relative to the severity of their injury (Table 1). Patients with extradural haemorrhage had the highest prevalence of adverse outcomes, whilst patients with isolated subarachnoid haemorrhage had the lowest (Fig. 11).

Meta-analysis of multivariable models was not possible due to the small number and variability in how these models were constructed. Therefore, although this review has identified the factors that affect risk, no model that could identify low-risk patients was found or could be reliably constructed.

A thorough search has been conducted, identifying 50 relevant primary studies. Our review fulfils all the AMSTAR systematic review checklist quality domains apart from items 10 and 11, regarding the assessment of publication bias and conflicts of interest. However, the non-interventional nature of the included studies means these domains are less relevant. This review is low-risk for bias in the 5 domains assessed by the Risk of Bias in Systematic reviews (ROBIS) tool. 142

Limitations

Strengths

Many studies identified were small and retrospective with limited follow up of patients after discharge. Instead of attempting to identify low-risk patients through prognostic modelling, several

studies selected patients on study specific characteristics for different care pathways. This variation in study populations contributed to heterogeneity in estimates of outcome prevalence and risk factor effect. The prognostic models that were identified were often derived in cohorts too small to construct multivariable models with all relevant factors. The clinically useful outcome in informing discharge decisions is clinical deterioration, and most prognostic models did not assess this.

Clinical deterioration was defined by 7 different composite outcomes and most commonly by neurological deterioration. This lack of consistency in definition contributed to the heterogeneity in outcome estimates. Neurological deterioration was variably defined and a clinically relevant and consistently used definition or deterioration is required.

No included studies assessed pupillary response and duration of loss of consciousness/amnesia.

These factors are predictive of adverse outcomes in other TBI populations and future research should assess these factors in this population. 13, 143

Context

When the Canadian CT Head Rule was developed, the authors presented a consensus derived list of intra-cranial injuries that would never require neurosurgeryneurosurgical intervention. The implication was that patients with such injuries were safe for discharge. This was rejected by the Society of British Neurological Surgeons. A US group based in Arizona has produced the BIG consensus derived statement that identifies a population with low risk clinical characteristics and intra-cranial injuries similar to those presented by the CCHR authors. They propose such patients are safe for discharge after 6 hours of ED observation. They propose such patients

Kreitzer et al present an alternative policy at a level 1 trauma centre in Cincinnati where the population of interest remain in the ED for observation and undergo repeat CT imaging approximately 6 hours following diagnosis. 86 Neurologically stable patients without progression of injury are discharged. Pruitt et al present a model of care in a Level 1 trauma centre in Chicago in

which all GCS13-15 patients with intra-cranial injuries receive a neurosurgical consultation. Low risk patients identified by the neurosurgeon are left under ED care and discharged after a period of observation. This is similar to the standard of care in the UK NHS.

Others advocate the admission of all GCS13-15 patients_andwith brain injuries mTBI-identified by CT imaging to higher levels of care and routine re-imaging, citing evidence that deterioration in neurological examination may not identify progression of injury that warrants clinical intervention.^{6,}

Multiple reviews have found that this too rare an occurrence to warrant routine re-imaging of all GCS13-15 patients with TBI identified by CT.¹⁷⁻²⁰

Implications

This review supports the view that there are subsets of GCS13-15 patients with injuries identified by CT imaging that may possibly be safely routinely discharged from the ED. However, the current available evidence is insufficient to reliably identify such low-risk patients. The risks of serious adverse outcomes are sufficiently high that, in the absence of evidence to be able to accurately pin point low-risk individual patients, admission for observation probably remains clinically indicated.

No validated model predicting a measure of clinical deterioration that could be used to triage hospital admission was identified. We suggest future research should assess a measure of clinical deterioration that encompasses: neurosurgeryneurosurgical intervention, death, a fall in GCS by 2 or more points, seizure activity, intravenous medical intervention or ICU intervention. These would warrant ongoing inpatient hospital admission.

The BIG criteria, although the best effort at risk stratifying this group in a clinically relevant way, require validation in larger prospective cohorts in different healthcare contexts before being more widely adopted. They were derived by consensus, and empirical prognostic modelling could possibly improve the accuracy of risk stratification.

Decision rules have been employed successfully in the ED to risk stratify patients in a range of conditions, including ankle injuries and suspected pulmonary embolus. ^{144, 145} Equivalent models could be used for patients with mTBI to identify low-risk patients. This review has identified the key factors that are likely to inform such risk stratification, but an adequately powered derivation study with a clinically relevant definition of deterioration and adequate follow up is required.

Conclusion

Mild TBI patients with injuries identified by CT imaging are a heterogenous group. Their overall risk of clinical deterioration and more serious adverse outcomes is small, but clinically significant.

Current research gives an indication to which factors affect the risk of adverse outcomes but is of too low quality to inform clinical decision making. High quality prognostic modelling is needed to help inform discharge decisions.

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Factor	Outcome	Unit Increase Affect Odds Univariable	Unit Increase Affect	
		Model	Multivariable	
Mean Age Study	Death	1.05 (95% C.I. 1.0003-1.12) P= 0.049	1.06 (95% C.I. 1.00	
Population			P= 0.049	
Mean GCS Study	Death	0.12 (95% C.I. 0.02- 0.86) P=0.04	0.09 (95% C.I. 0.01	
Population			P=0.02	
Lower risk study	Death	0.27 (95% C.I. 0.08-0.94) P=0.04		
population versus ICU				
population				
Unselected study	Death	0.81 (95% C.I. 0.22-1.97) P=0.63		
population versus ICU				
population				
Percentage population	Death	1.05 (95% C.I. 0.95-1.17) P=0.32		
Anticoagulated				
	Neuroumgomi	4.04 (050) (0.1.4.02 (4.41) D. 0.04	1.00 (050) 61.1	
Mean Age Study	Neurosurgery	1.01 (95% C.I. 1.02- 1.11) P=0.01	1.09 (95% C.I. 1.0	
Population			P=0.02	
Mean GCS Study	Neurosurgery	0.71 (95% 0.01- 0.56) P=0.01	0.12 (95% C.I. 0.0	
Population			P=0.04	
Population			F=0.04	
Lower risk study	Neurosurgery	0.13 (95% C.I. 0.04- 0.41) P<0.01	0.67 (95% C.I. 0.1	
population versus ICU		•	P=0.66	
population				
Unselected study	Neurosurgery	0.95 (95% C.I. 0.43- 2.12) P=0.90	1.34 (95% C.I. 0.4	
population versus ICU			P=0.58	
population				
1 P: :: 5::				

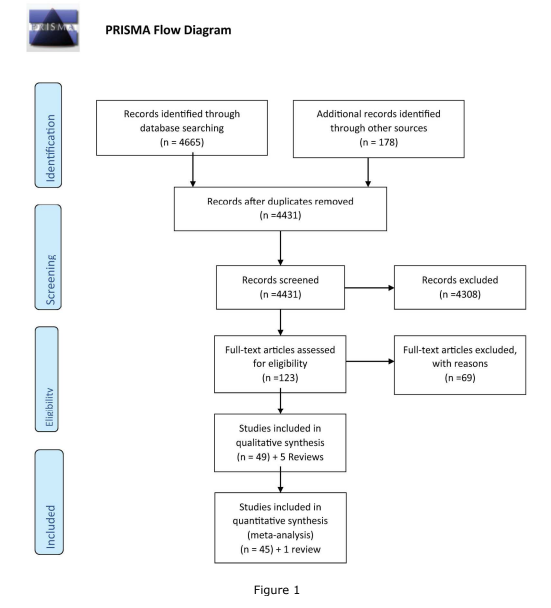
Anticoagulated			
Exclusion of anti-	Neurosurgery	0.63 (95% C.I. 0.27- 1.43) P=0.26	1.33 (95% C.I. 0.51- 3.49)
coagulated patients in			P=0.54
study selection			
Mean Age Study	Clinical	1.01 (95% C.I. 0.95-1.09) P=0.64	1.02 (95% C.I. 0.93-1.12)
Population	Deterioration		P=0.59
Mean GCS Study	Clinical	0.36 (95% C.I. 0.04-3.20) P=0.33	0.26 (95% C.I. 0.02-3.76)
	Deterioration	0.30 (33% C.I. 0.04-3.20) F-0.33	
Population	Deterioration		P=0.29
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Table 2: Summary of effect estimates of risk factors assessed within studies

Risk Factor	Number of Studies Assessed in	Pooled Univariable Effect*	Effect Multi-variable Models**	Likely Effect on Risk	
Age	18 ^{6, 37, 41, 54, 55, 66, 69,} 71, 73, 74, 76-78, 98-101, 130		+6/11	+	
Initial GCS 15	7 ^{37, 41, 66, 73, 74, 77, 101}	OR 0.35 95% CI: 0.23 to 0.52	- 4/4	-	
Severity CT brain	9 ^{6, 41, 54, 55, 66, 73, 76, 78,} 100		+7/8	+	
Isolated SAH	5 ^{37, 73, 77, 98, 108}	OR 0.19 95% CI: 0.07 to 0.5	-1/2	-	
Isolated EDH	5 ^{37, 73, 77, 98, 108}	OR 2.26 95% CI: 1.9 to 2.68	+1/1	+	
Isolated SDH	5 ^{37, 73, 77, 98, 108}	OR 1.82 95% CI: 0.69 to 4.77	+2/2		
Isolated Contusion	3 ^{37, 98, 108}	OR 0.24 95% CI: 0.2-0.28	0/1		
Anti-coagulation	12 ^{6, 37, 41, 55, 74, 76-78,} 98, 100, 101, 139	OR 1.45 95% CI: 1.28-1.64	0/2	+	
Aspirin	6 ^{37, 55, 66, 76, 87, 101}	OR 1.30 95% CI: 0.95-1.78			
Clopidogrel	6 ^{37, 55, 66, 76, 87, 101}	OR 1.79 95% CI:1.17-2.72		+	
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^{*}Pooled estimate of effect on risk of neurosurgery or clinical deterioration

^{**}Indicates number of multivariable models where factor was found to be a significant predictor and direction of effect on risk



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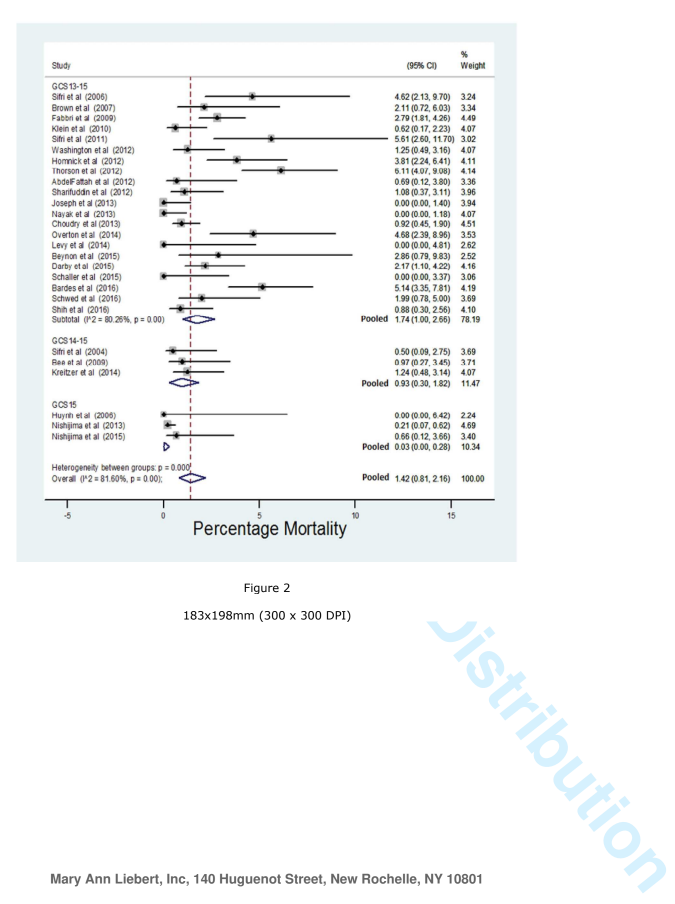


Figure 2 183x198mm (300 x 300 DPI)

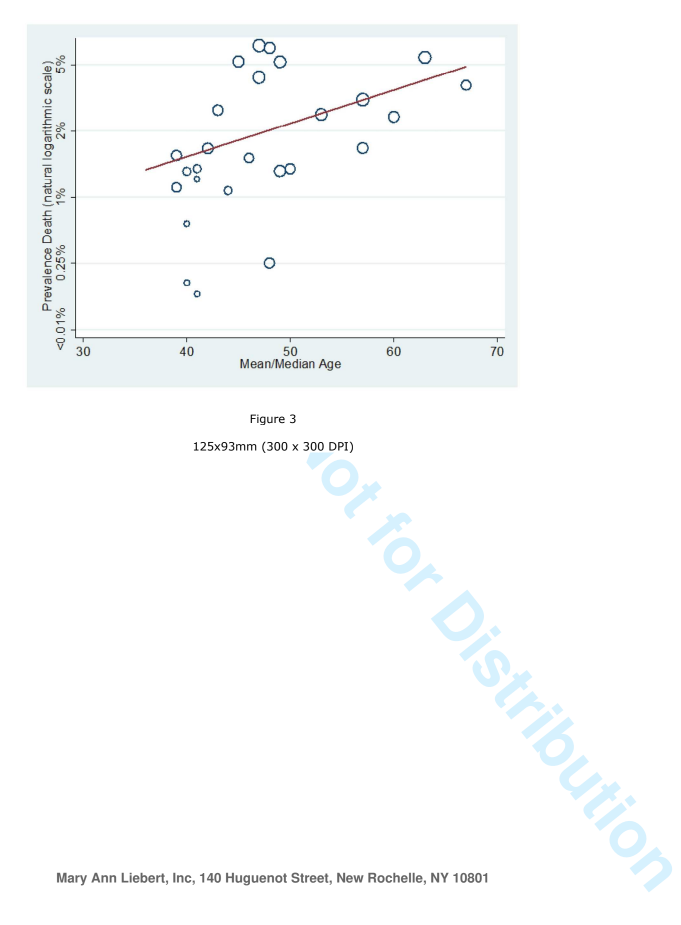


Figure 3 125x93mm (300 x 300 DPI)

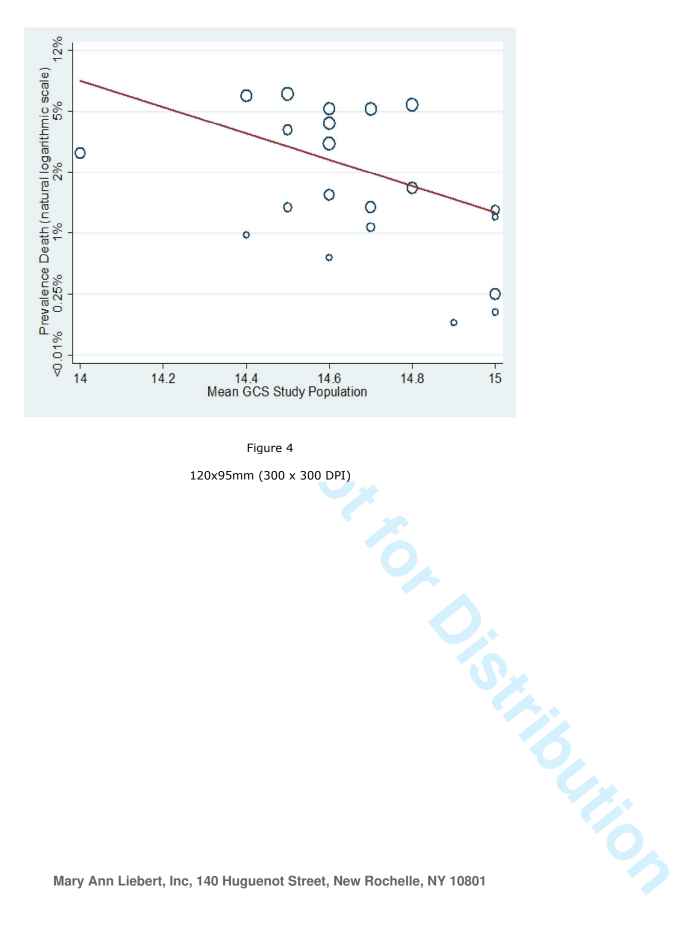


Figure 4 120x95mm (300 x 300 DPI)

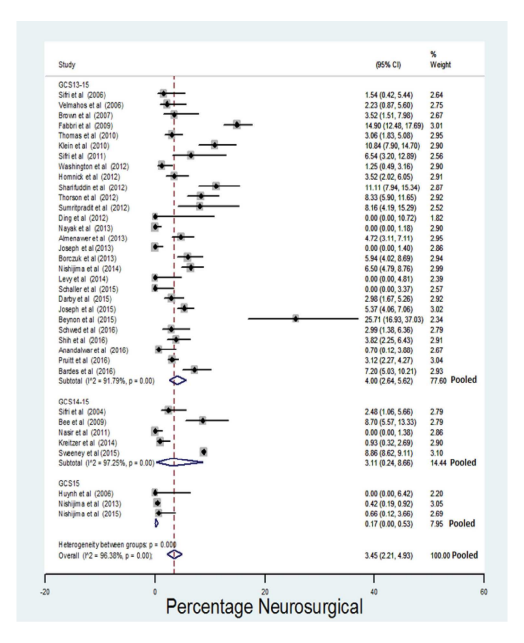


Figure 5 190x233mm (300 x 300 DPI)

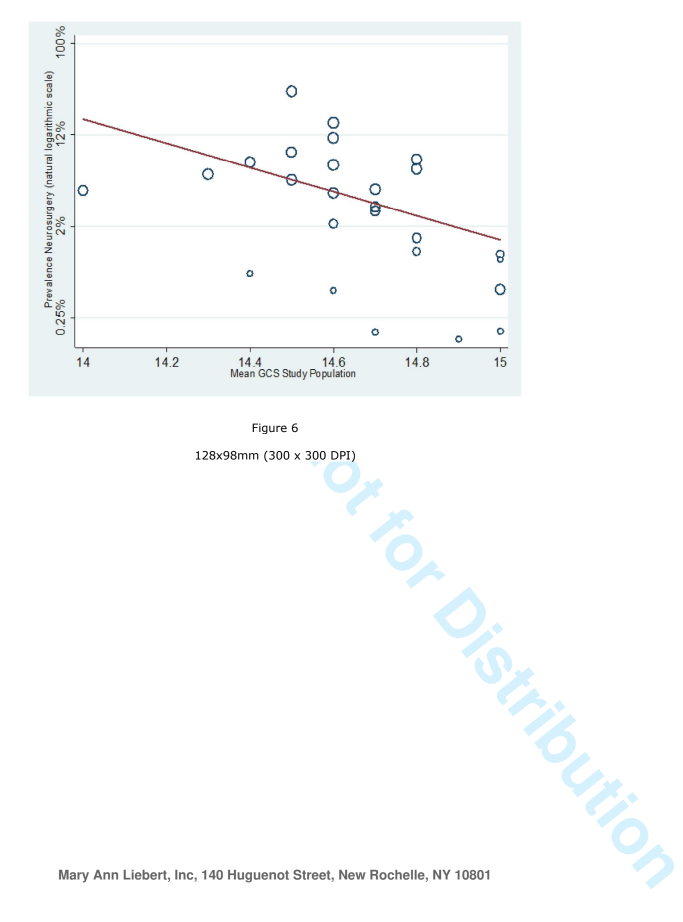


Figure 6 128x98mm (300 x 300 DPI)

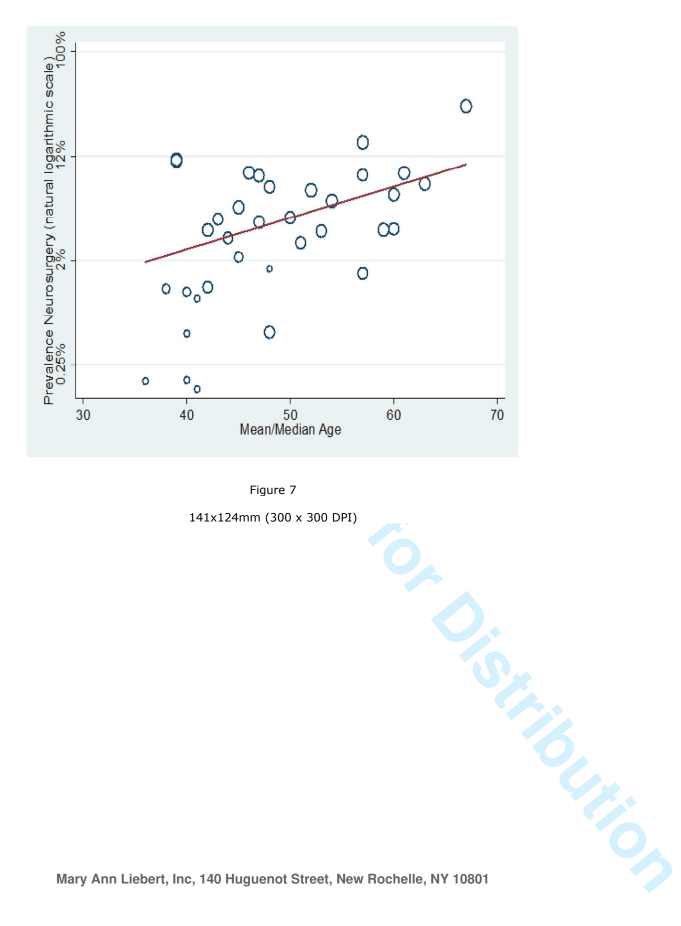
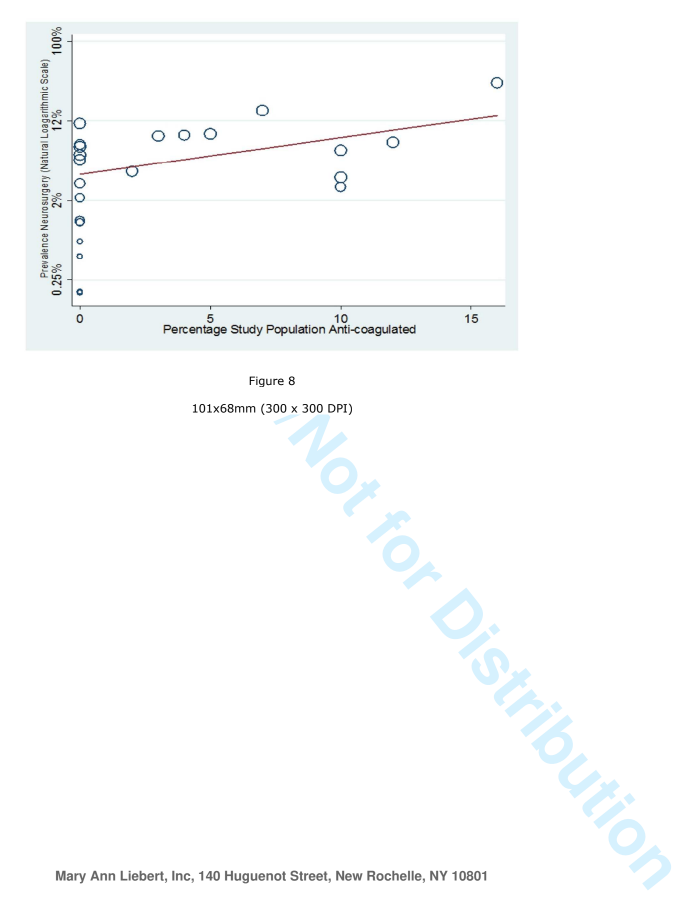


Figure 7 141x124mm (300 x 300 DPI)



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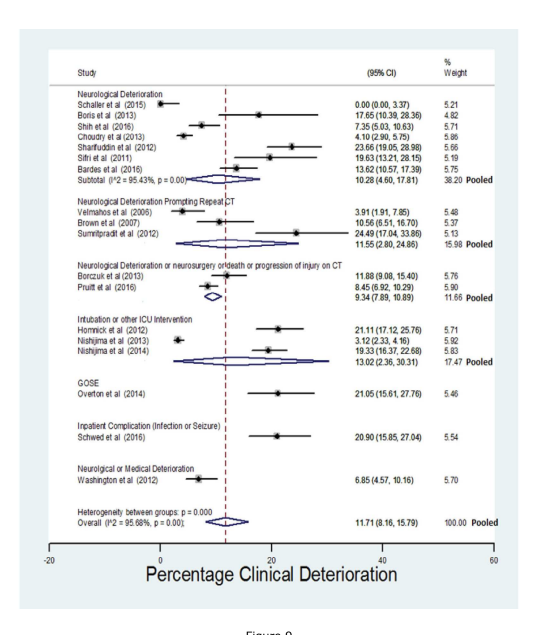


Figure 9 166x192mm (300 x 300 DPI)

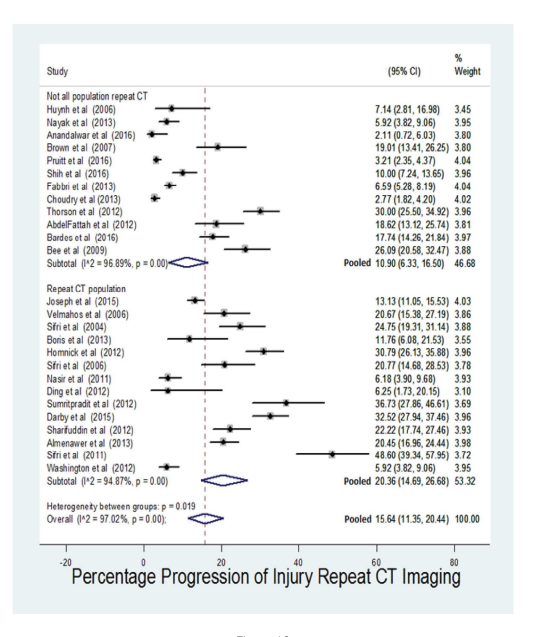
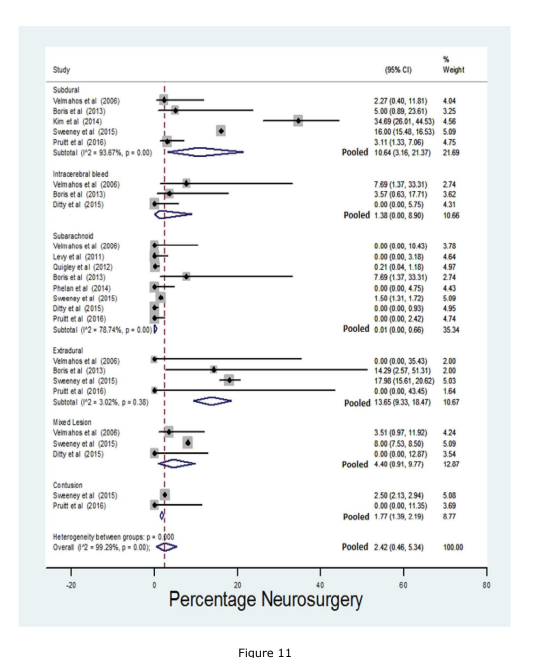


Figure 10 181×213mm (300 × 300 DPI)



194x237mm (300 x 300 DPI)

- Figure 1: PRISMA flow-diagram showing selection of studies for inclusion in the systematic review
- Figure 2: Risk of Death stratified by initial GCS
- Figure 3: Meta-regression risk of death by mean age study population (Coefficient odds 1.05 (95% CI: 1.00 to 1.12) P=0.049)
- Figure 4: Meta-regression risk of death by mean GCS study population (Coefficient odds 0.12 (95% CI: 0.02 to 0.86) P=0.04)
- Figure 5: Risk of neurosurgery stratified by the initial GCS of the study population
- Figure 6: Meta-regression of risk of neurosurgery by mean GCS study population (Coefficient odds 0.71 (95% 0.01- 0.56) P=0.01)
- Figure 7: Meta-regression of risk of neurosurgery by mean age study population (Coefficient odds 1.01 (95% C.I. 1.02- 1.11) p=0.01)
- Figure 8: Meta-regression of risk of neurosurgery by percentage of study population taking anticoagulants (Coefficient odds 1.1 (95% C.I. 1.01-1.19) p=0.04)
- Figure 9: Estimates of clinical deterioration stratified by the outcome measure
- Figure 10 Risk on repeat CT imaging of progression of injury stratified by whether entire population selected for repeat imaging
- by the out.
 of injury stratifie.
 d by isolated injury type ide. Figure 11: Pooled risk of neurosurgery stratified by isolated injury type identified by initial CT imaging

Supplementary material 1: Full Search Strategy

Embase search 24/11/2016 1996 to 2016 Week 47:

	12	1 and 10 and 11	3167					
	11	2 or 3 or 4 or 5 or 6 or 9	104649					
	10	7 or 8	2298555					
	9	"cerebral contusion".mp. or exp brain contusion/	2627					
	8	exp outcome variable/ or outcome.mp. or exp critical care outcome/ or exp adverse outcome/	1787765					
	7	exp prognosis/ or prognos*.mp.	704898					
	6	exp subarachnoid hemorrhage/ or "traumatic subarachnoid h#em*".mp.	28977					
	5	"extradural h#em*".mp.	225					
	4	exp epidural hematoma/ or "epidural h#em*".mp.	4775					
	3	exp subdural hematoma/ or "subdural h#em*".mp.	10281					
	2	exp Intracranial Hemorrhages/ or "intracranial h#em*".mp.	92720					
	1	"traumatic brain injury".mp. or traumatic brain injury/ or head injury/	69888					
M	EDLINE	Ovid MEDLINE(R) without Revisions 1996 to November \	Week 3 2016					
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24/11/2016

9	1 and 7 and 8	1143	
8	2 or 3 or 4 or 5 or 6	34984	
7	exp Risk Factors/ or risk.mp. or exp Risk/ or exp Risk Assessment/	1502469	
6	"traumatic subarachnoid h#emorrhage".mp. or exp Subarachnoid Hemorrhage Traumatic/	e,231	
5	exp Cerebral Hemorrhage, Traumatic/ or exp Hematoma, Epidural, Cranial/ or "extradural haemorrhage".mp.	1434	
4	exp Hematoma, Subdural/ or "subdural h#em*".mp.	3712	
3	exp Intracranial Hemorrhages/ or "intracranial h#em*".mp.	34253	
2	exp Cerebral Hemorrhage/ or "intracerebral h#em*".mp.	14418	
1	"head injury".mp. or exp Craniocerebral Trauma/	75438	
		NY 10801	

CINHAL plus access through EBSCO 24/11/2016 1983-2016:

		1
Search Terms	Search Options	
S11	((S3 OR S4 OR S5 OR S6) AND (S3 OR S4 OR S5 OR S6 OR S7)) AND (S8 AND S9 AND S10)	View Results (292)
S10	(S3 OR S4 OR S5 OR S6) AND (S3 OR S4 OR S5 OR S6 OR S7)	View Results (6,995)
S9	S1 OR S2	View Results (17,827)
S8	prognosis or outcome	View Results (592,464)
S7	brain contusion OR cerebral contusion	<u>View Results</u> (106)
S6	extradural haematoma OR extradural hematoma OR (epidural hematoma or epidural hemorrhage)	<u>View Results</u> (753)
S5	intracerebral hemorrhage OR intracerebral haemorrhage OR intracerebral bleed	View Results (2,456)
S`4	intracranial hemorrhage OR intracranial haemorrhage OR intracranial hematoma OR intracranial haematoma	View Results (3,176)
S3	subdural hematoma OR subdural hemorrhage OR subdural haematoma OR subdural haemorrhage	View Results (1,246)
S2	traumatic brain injury	View Results (10,081)
S1	head injury	View Results (7,746)

Cochrane CENTRAL:

Search Name: Prognostic systematic Review

Date Run: 24/11/16 11:33:55.251

- ID Search Hits
- #1 Craniocerebral Trauma 417
- #2 head injury 2563
- #3 #1 or #2 2704
- #4 Hematoma, Subdural 228
- #5 Hematoma, Epidural, Cranial 20
- #6 Cerebral Hemorrhage 2609
- #7 Skull Fracture 130
- #8 Skull Fracture, Basilar 6
- #9 Skull Fracture, Depressed 13
- #10 brain contusion 131
- #11 #4 or #5 or #6 or #7 or #8 or #9 or #10 2969
- #12 #3 and #11 211

All Results (211)

- Cochrane Reviews (138)
- All Review Protocol
- Other Reviews (4) Trials (63) Methods Studies (0) Technology Assessments (0)
- Economic Evaluations (1) Cochrane Groups (5)

Only trials retrieved.

		Studies Onl	y Included in Meta	-Analysis of	Prevalence of Outcomes N=26	
Reference	Population	Study Design	Outcome Measures	Prognostic factors assessed	Results	Quality Appraisal
Nishijima et al	Multicenter-8 sites	Retrospective Cohort	Initial ICU admission from ED	Age	11240 patients coded as bleeds	Study Recruitment: Mod risk bias
2013	Western USA. All Level 1	Study		Initial GCS	771 excluded due to missing data	Dependent on accuracy on recording of
Sacromento			Proportion of patients	Initial BP		trauma registry. Does have some qualit
USA	Trauma registries	Objective:	receiving crit care	LOS hosp	1412 remaining met inclusion criteria.	assessment of data imputation
	searched for ICD-9 codes	1) assess the variability of	intervention defined as:	ICU stay		
	intra-cranial	ICU use in a cohort of	Neurosurgical intervention	Procedures as	888/1412 admitted ICU, significant variation between sites	Note initial GCS 15- lower risk group
Variability of	haemorrhage 2005-2010	patients with minor	Mechanical ventilation	coded in trauma		
ICU Use in		traumatic intra-cranial	Vasopressor/ionotropic use	registry	44/1412 (3.1%) had critical care intervention	Attrition: Low risk
adult patients	Inclusion Criteria:	haemorrhages across	Transfusion blood product	AIS	6/1412 neurosurgical intervention	Follow up only during hospital admission
with minor	 Age ≥ 18 years 	multiple trauma centres.	Invasive monitoring			
traumatic intra-	Traumatic ICH				847/888 patients admitted ICU no crit care intervention	Prognostic factor measurement: Low risk
cranial	 Initial ED GCS 15 	2)Estimate the proportion				Doesn't really apply as testing disposition
haemorrhages	ISS less than 16	of minor traumatic			Mean/median GCS=15	not outcomes
		intracranial haemorrhages			Mean/median age= 48	
		patients admitted to ICU				Outcome measures: Low risk
		that do not receive an ICU				No measure of outcomes after discharge
		intervention				but study primarily about disposition.
						Does not report deaths.
						Confounding Factors:
						States IIS increases ICU admission- will b
						related to other injuries
						Statistical techniques: low risk
						N/A
						N/A
						Overall
						Only GCS15 patients with low ISS.
Nishijima et al	Level1 trauma centre	Retrospective Cohort	Prospective long term	age	188 met inclusion criteria	Study Recruitment: Mod risk bias
2015		Study	outcome measure at 6	sex,		Dependent on accuracy on recording or
Sacromento	2008-2013	•	months	mechanism of	151/188 complete data= cohort	trauma registry and accuracy of case notes.
USA	Inclusion Criteria:	Aim	Either GOS-E 8 fully	injury initial ED	106 admitted ICU (70%)	5 , , , , , , , , , , , , , , , , , , ,
	 Age ≥ 18 years 	compare long-term	recovered or GOS-E 1-7 not	GCS score, initial	45 admitted ED (30%)	Low risk group- GCS 15 and benign CT
	Identified ICH ICD9	neurological outcomes in	fully recovered	(SBP)		
	code trauma	low- risk patients with		heart rate,	1/151 patients neurosurgical intervention as inpatient	Attrition: Low risk
	registry	traumatic intracranial		respiratory rate,	1/151 patient died as inpatient	Loss of 37 patients to follow up
Long-term	Initial ED GCS 15	hemorrhage (tICH)		blood alcohol	78 (52%) GOS-E 8 at 6 months	
Neurological	Isolated Head	admitted to the ICU		level, AIS score		Prognostic factor measurement: Low risk
Outcomes in	Injury based on AIS	(intensive care unit)		ISS score	Does present analysis for outcome at 6 months GOSE but	As recorded in case notes so dependent or
Adults with	score	versus patients admitted		INR	no inpatient measures of deterioration.	accuracy

				Journ	al of Neurotrau	uma	Page 66 of 139
1 2 3 4							
5 6 7 8 9 10 11 12	Traumatic Intracranial Hemorrhage Admitted to ICU versus Floor	Age<65 No evidence midline shift CT Present on TBI data base due to suspected TBI/evidence of ICH	to the floor.		Rotterdam CT score	Adjusted analysis, floor admission versus ICU had an odds ratio of 0.77 (95% CI [0.36-1.64]) for a GOS-E score of 8 at six months. Mean/median GCS=15 Mean/median age= 40	Outcome measures: Low risk Prospective follow up by trained staff using validated tool. Not clear what would happen to patients who died or deteriorated and attended a different hospital. Confounding Factors: Patients which are perceived as higher risk
13 14 15 16 17 18							will be put on ICU, likely to be differences in comorbidities Statistical techniques: low risk Well presented- not really relevant to meta- analysis Only GCS15 patients with benign looking CT scans
20 21 22 23 24 25 26 27	Schaller et al 2015 Switzerland	Level 1 Trauma centre Bern Switzerland Jan 2006-Dec 2007 Inclusion criteria: Admission GCS 13- 15	Retrospective cohort study/case series Aim to assess if a specific group of patients with small bleeds can be discharged from hospital without 24 hours of observation	Deterioration in neurological status or need for neurosurgery.	Prognostic factors are the inclusion/exclusion criteria No comparison in risk of deterioration in 2 groups.	110 patients met inclusion and exclusion criteria. None deteriorated within the period of hospital observation, required neurosurgery or re-attended. Mean/median GCS=14.6 Mean/median age= 40 Percent anticoagulated=0	Study Recruitment: Low risk bias Retrospective cohort review- reliant on accuracy of written notes. Attrition: Mod risk Patients may have moved out of catchment area of hospital without the researchers being aware. Loss to F/U if re-presented different hospital.
28 29 30 31 32		Observed for 24H Localised intracranial bleeds up to 5mm- this is from the CCHR paper Exclusion Criteria: Pleads Frage Control Control				6	Prognostic factor measurement: Mod risk Reliability of case notes- may be incomplete Interpretation size of the bleed was taken from written radiology report ?reliability.
33 34 35 36 37 38		 Bleeds > 5mm maximum diameter Multiple bleeds History of bleeding tendency Anti-coagulant or 					Outcome measures: Moderate risk Study dependent on patients re-presenting at the same hospital following discharge if had delayed deterioration. Not clear how patients died in the community would have been identified.
39 40 41 42		anti-platelet medication • Intoxication					Confounding Factors: Low risk No obvious confounding factors Cohort selection criteria including not living

Live greater the 1H from hospital	Peli				alone may select out high risk older patients. Statistical techniques: N/A General comments: Mean age 39.9 years and 25% caused by sporting injuries. ?Age as the confounding low risk prognostic factor. Not generalizable to older populations
Levy et al 2011 Colorado JSA Inclusion criteria: Admission ED GCS 13-15 On trauma registry Blunt head trauma ICD 850-850.99- consistent with concussion (i.e. no detected injury by CT) Admitted to hospital AIS score 2 before 2008 or 1 / 2 in 2008 IC9 code for SAH Exclusion Criteria: Patient admitted directly to hospital Multiple injuries AIS score > 1 head or other regions Age less than 18 Not admitted	Retrospective Cohort Study Aim To assess whether patients admitted with CT -VE mTBI have different outcomes to patients with mTBI and traumatic SAH Univariate and multivariate regression used to examine covariates and relationship to outcomes	ED disposition ICU admission Neurosurgery In-hospital mortality Progression of SAH on CT	Age (18-39)(40-69)(70+) Transfer status Cause of injury GCS Blood alcohol level Presence of skull fracture CT report- divided into small/medium/lar ge based on language included in report	1144 patients admitted with mTBI but negative CT scan 117 with mTBI and traumatic SAH 1/117- progression on repeat CT scan 0/117 required neurosurgical intervention 1/117 died (progression on CT) 4/1144 died All patients died >70 Logistic regression model tSAH versus concussion ICU admit adjusted OR 8.87 (5.62-14.02) P<0.0001 ICU LOS>1D OR0.29 (0.11-0.74) P=0.01 Hosp LOS>1D OR1.07 (0.67-1.69) P=0.79 Mortality OR2.46 (0.27-22.17) P=0.42 Discharge to rehab Age18-39 OR5.48 (0.25-121.70) P=0.28 Age 40-69 7.96 (1.91-33.11) P=0.004 Age >70 1.33 (0.50-3.53) P=0.56	Small numbers Study Recruitment: Low risk bias Patients recruited from trauma registry depends on how good this is Only admitted patients- higher acuity patients then discharged. Likely patients admitted for other reasons if CT negative TBI (although excludes other injuries). Attrition: Low risk All inpatient outcomes Prognostic factor measurement: Mod risk CT findings abstracted from CT reports-severity assigned by language- not actually used in regression model Outcome measures: Moderate risk Only inpatient outcomes- possibility of discharge and deterioration. Confounding Factors: High risk Patients admitted with CT negative TBI likely to be frail or have other reasons for admission- this will affect outcome measures compared to SAH patients admitted due to +ve CT. Statistical techniques: Low risk Well presented.

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1 2 3 4	0,	0					
5 6 7 8 9 10 11 12 13 14 15 16 17 18							Can use for pooling for outcomes SAH-supports low risk sub-population
20 21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40 41	Levy et al 2014 USA	Level III rural non- neurosurgical unit in Rocky mountains April 2007-Dec 2012 April 2007 patients with small bleeds selectively not transferred to neurosurgical unit Inclusion criteria: • Admission GCS 13- 15 • CT positive intra- cranial injury • Not transferred to neurosurg unit in accordance with non-transfer policy. • CT findings of small SAH • Punctate or minimal contusion • Punctate or minimal intra-	Retrospective cohort Study Aim Investigate outcomes after a novel non-transfer policy for mTBI patients with small ICH introduced in a small rural trauma unit without neurosurgical cover	Length of stay Mortality Neurological deterioration Neurosurgery Re-admission in 90 days of discharge Inter-hospital transfer Need for repeat CT	No comparison to patients that were transferred	76/273 patients not transferred >50% injuries due to skiing/snow boarding 71% patients less then 55 No patient deteriorated, died or required neurosurgery or required delayed transfer whilst admitted to hospital. 2 patients re-admitted within 90 days- 1 patient 6 weeks following admission developed an acute on chronic subdural- drained. 1 patient re-admitted with unrelated complaint. Mean/median GCS=14.7 Mean/median age= 36 Percent anticoagulated=0	Study Recruitment: Low risk bias Retrospective cohort review- reliant on accuracy of written notes. CT inclusion criteria are subject and patients may have been transferred despite meeting non-transfer policy if clinicians were concerned. Attrition: low risk Prognostic factor measurement: Mod risk Reliability of case notes- may be incomplete The definitions of bleed size are subjective. Prognostic Factors N/A Outcome measures: Moderate risk Study dependent on patients re-presenting at the same hospital following discharge if had delayed deterioration. Confounding Factors: Low risk Age affect outcome and size of bleed Statistical techniques: N/A
42 43 44 45 46 47			Mary A	nn Liebert, Inc, 140 H	luguenot Stree	t, New Rochelle, NY 10801	

				1		
Joseph et al 2013 USA The acute care surgery model: Managing traumatic brain injury without an inpatient neurosurgical consultation	cranial bleed Small SDH, no mass effect Exclusion Criteria: Any coagulopathy Basilar skull fracture or evidence of CSF leak Extra-dural bleed Any significant contusion or SDH/intra-cerebral haemorrhage Review and discussion of CT and patient with neurosurgeon if unsure if should be transferred Level 1 Trauma centre 2009-2011 (likely subset of patients presented below) Inclusion criteria: GCS13-15 Trauma Positive findings CT- skull fracture and/or ICH Exclusion Criteria: Pre-hospital antiplatelets or anticoagulants	Retrospective cohort study- propensity matching 1:2 ratio patients managed solely by trauma surgeons versus patients that had neurosurgical consultation. Hypothesis Trauma surgeons can manage mTBI patients with CT detected intracranial haemorrhage without neurosurgical invlolvement	Hospital admissions ICU admissions Neurosurgical interventions ED visits after discharge Mortality Progression on CT imaging	Age Sex Initial GCS ISS Head-abbreviated injury score Neurological examination CT scan findings- type of skull fracture/type of ICH/size of bleed- reviewed by study investigator	404-GCS13-15 patients with CT detected injuries in study period. 270/404 used for this study 90/270- had neurosurgical consultations (NC) 180 no neurosurgical consultation. (no-NC) Whether neurosurgical consultation requested as discretion of non-specialist surgeon. Propensity matching in this study between 2 groups. 0/270 neurosurgical interventions, hospital mortality or readmissions either group. 78/90 no-NC and 158/180 NC admitted hospital (P=0.8) 18/90 no-NC and 80/180 NC admitted ICU (P=0.001) Routine repeat CT 18/90 no-NC 155/180 NC (P<0.001) No progression on any repeat CT 8% no-NC and 4% NC group re-attended ED. No readmissions. Mean/median GCS=15	Small numbers. No comparator group- need to compare to transferred patients outcomes. Patient not generalizable- v. young and atypical mechanism of injury (mostly winter sports related). Likely that any patient clinicians felt risky would have been transferred even if did not meet transfer criteria- no way to check this. Study Recruitment: High risk bias Subset of patients that meet inclusion criteria selected in order to facilitate propensity matching. Possible selection out of higher acuity patients as these will have al been referred to a neurosurgeon. Attrition: low risk In patient outcomes and documented ED re-attendances- low risk of patients being lost to follow up Prognostic factor measurement: Low risk All routinely collected clinical data apart from CT imaging which re-reviewed. Outcome measures: Mod risk Study dependent on patients re-presenting at the same hospital following discharge if had delayed deterioration. Confounding Factors: Mod risk Does not exclude patients with additional injuries
					8% no-NC and 4% NC group re-attended ED. No readmissions.	had delayed deterioration. Confounding Factors: Mod risk Does not exclude patients with additional
					Mean/median GCS=15	injuries
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	400				Mean/median age= 30 Percent anticoagulated=0	Statistical techniques: High risk Does not outline how matched groups using propensity scoring
	4	701.				General points Small numbers.
						Likely reporting data reported else where.
AbdelFattah et	Level 1 trauma center	Prospective Cohort Study	Outcome measures during	Comparison	145 patients met inclusion/exclusion criteria.	Study Recruitment: low risk
al	Dallas Texas	rrospective conort study	hospital admission:	between groups:	92/145 for routine repeat CT	Prospective recruitment- states recruited all
2012	Janus Texas	Hypothesis:	The second second	Age	53/145 for CT if deteriorated	eligible patients. Doesn't explain how
	Prospective recruitment	Repeat CT imaging in	Neurologic progression.	Sex	Selective group more likely aspirin use P=0.02	recruitment occurred.
USA	2010-2011	GCS13-15 with ICH,	Medical intervention	Coagulation status	Routine repeat CT worse Head AIS score (P<0.001)	
		without neurological	Neurosurgical intervention	Anti-platelets	Otherwise groups comparable	Attrition: low risk
	Inclusion criteria:	progression, does not	Repeat CT imaging- worse CT	ISS		Follow up only for period in hospital
	Adult with ICH	impact the need for	defined as worse by a	GCS	5/53 deteriorated and had a repeat CT + 1/53 had repeat	
	(note doesn't	neurosurgical	blinded		scan as started on warfarin	Prognostic factor measurement: Low risk
	explicitly state	intervention.	radiologist/neurosurgeon			Blinded appraisal of CT scans by researcher.
	2ndary to trauma-		giving qualitative measure of		1/145 patients died (due to other injuries)	
	but implied)	Patients divided into	bleed.		27/145 radiological deterioration	Outcome measures: Mod risk
	Excluded:	those 2 groups. Patients			9/145 patients intubated- states for other injuries	No F/U following discharge- missed delayed
	• Age<16	with planned repeat CT				outcomes, could have looked for re-
	• GCS<13	imaging and those with CT			Mean/median GCS=14.5	attendance.
	 Undergone 	imaging if deteriorated.			Mean/median age= 41	Doesn't report neurosurgical outcome
	planned or	Allocation by			Percent anticoagulated=6	measures.
	immediate	neurosurgeon-no deviation from normal				Confounding Factors: High risk
	neurosurgery	practice.				Not isolated head injury- other injuries have
	Transferred	practice.				clearly affected outcome measures
	patients					cicarry affected outcome measures
						Statistical techniques: Low risk
						None
						Small study with confoundant regarding
						Small study with confounders regarding outcomes.
Nayak et al	University Hospital	Retrospective Chart	Neurosurgical intervention	Age	321/864 patients GCS13-15 with ICB met inclusion criteria	Study Recruitment: Low risk
2013	Newark New Jersey	Review	after 24 hours- craniotomy,	Sex	20% excluded because incomplete medical notes/transfers	Retrospective case note review- depends
	Level 1 trauma centre		ventriculostomy, ICP	Mechanism of		on information being recorded correctly.
USA	2003-2008	Aim:	bolt/measurement	Injury	0/321 neurosurgical intervention-all within 24 hours of	
		To compare neurologic		GCS on arrival	admission	Attrition: Mod risk
	Inclusion criteria:	outcomes of MHI patients	Death in hospital	ISS		20% excluded because of incomplete notes
		with an intra-cranial bleed		HAIS	No deaths	

8	ormal Discharge disposition	GCS and		Prognostic factor measurement: Mow risk
Blunt trauma neurological examin		neurological	19/142 worse CT on repeat CT after 24 hours of admission	Neuroradiology reports taken at face value-
 Intra-cranial bleed managed with 	and LOS hospital	examination every		no verification
Admitted to without a repeat CT		2 hours- routine	179/321 single CT	
hospital scan	GOS at f/u clinic/ re-	care on a flow	142/321 routine repeat CT	Outcome measures: mod risk
• GCS13-15 on	attendance if applicable	sheet	7C/221 votumed to F/II alinia unavantful	No uniform follow up of potionts post
arrival to ED			76/321 returned to F/U clinic- uneventful	No uniform follow up of patients post
• GCS 15 24 hours			14/321 returned to ED due to symptoms.	discharge. Some patients had F/U clinic others didn't. Patients may presented after
after attendance to			14/321 returned to ED due to symptoms.	discharge to other sites.
ED			Mean/median GCS=14.9	discharge to other sites.
Excluded:			Mean/median age= 41	Confounding Factors: low risk
History brain			Wednymedian age- 41	None obvious
disease, e.g.				None obvious
dementia				Statistical techniques: Low risk
Previous brain inium a process				None completed
injury e.g. CVA				
• Liver cirrhosis,				The inclusion/exclusion criteria have
renal disease, coronary artery				selected out all patients that are not GCS 15
coronary artery disease, bleeding				at 24 hours. Different population than all
or clotting disorder				GCS 13-15 patients with TBI on CT- probably
Unable to assess				unable to pool this data.
GCS due to drugs				
e.g.				Does show patients that are GCS 15 at 24
sedation/intubatio				hours low risk.
n				
Neurological				
deterioration				
leading to repeat				
CT			C 25'	
Aged less than 15				
Incomplete notes				
, , , , , , , , , , , , , , , , , , , ,				
Anandalwar et University Hospital Retrospective c	ohort Repeat CT after 24 hours of	Age	533 patients TBI and ICH	Study Recruitment: High risk
al 2016 Newark New Jersey study	admission due to clinical	Sex	142 met the inclusion/exclusion criteria	Patients at GCS15 at 24 hours- low risk
New Jersey Level 1 trauma centre	concern or deterioration.	Mechanism of	47 underwent a routine repeat CT within 24 hours	group selected out- difficult to extrapolated
USA 2009-20012 Aim		Injury	(violation of policy)- 0/47 neurosurgical, 1/47 had	to all GCS13-15 patients.
Assess the outc	omes Progression on any repeat CT	ISS	incidental finding on CT	
Inclusion criteria: following	the completed.	AIS		Does not compare outcomes in patient that
Aged 18 and over implementation of			95 no repeat routine CT within 24 hours	adhered to and violated non-routine repeat
Blunt trauma policy of observation	,			CT head imaging. Potentially clinicians
Intra-cranial (no repeat CT imaging)			8/95 (non-violation group) had repeat CT >24 hours after	ordered routine repeat CT imaging on
bleed/skull GCS 15 patients	Intubation, ICU admissions,		admission- due to concern.	riskier patients.
fracture	administration of mannitol.		0.40	
Admitted to]	3/8 progression on CT	Attrition: Low Risk

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1 2 3 4							
5 [hospital		ED revisits within 1 year for			Potential for patients to have re-attended
6		• GCS13-15 on		TBI related symptoms.		1 neurosurgical intervention	at other EDs and be missed
7 8		 arrival to ED GCS 15 24 hours after attendance to 				2/8 admitted to ICU due to deterioration- 1 intubated	Prognostic factor measurement: Low risk No risk model developed
9		ED				3/95 patients returned with 1 year to the ED due to TBI	Factors abstracted from case notes
10		Did not receive a				symptoms- all underwent repeat CT. No admissions.	Outcome mecaniman law siek
11		repeat CT head scan	70			Mean/median GCS=14.8	Outcome measures: low risk Re-attendance at other EDs makes re-
12		Excluded:				Mean/median age= 38	attendance a potentially biased outcome
13		History of				Percent anticoagulated=0	measure
14 15		neurological or psychiatric					
16		disorder					Confounding Factors: Mod risk
17		 Immediate neurosurgery 					Cohort includes patients with multiple injuries
18		Previous TBI or					Statistical techniques: Low risk
19		neurosurgery					None presented
20		Spinal injuryCoagulopathy					Is a lower risk population due to selection
21		Pregnancy					for repeat CT imaging and return to GCS15
22		• Transfers					at 24 hours- possibly unable to include in any meta-analysis.
23		Incomplete notes					
24		Patients that did					
25		undergo a repeat CT					
26 27		scan despite meeting the rest of					
28		inclusion/exclusion					
29 30		criteria formed a comparison group					
31	Ditty et al	University Alabama	Retrospective Cohort	Neurological decline- altered	Admission GCS	500 patients met inclusion criteria	Study Recruitment: Mod risk
32	2015 Alabama	Level 1 trauma centre 2003-20013	Study	mental state or focal neurological deficit.	Anti-coagulation Anti-platelets	411/500 isolated SAH 63/500 isolated ICH	High proportion of transferred patients may represent higher or lower acuity patients
33	USA	2003-20013	Aim	nearological deficit.	Transfer Distances	26/500 both	than general population.
34		Inclusion criteria:	Assess the clinical	Inpatient seizure	Sex		
35		 500 consecutive patients present on 	implications of SAH or intraparenchymal	Delayed neurosurgical	Age Haemorrhage type	463 GCS15 30 GCS14	Higher as being transferred to specialist centre, lower as survived /fit to transfer.
36		trauma registry	haemorrhage in mTBI	evacuation as inpatient.	acmorrhage type	8 GCS13	
37		• GCS13-15		Innationt martality		460/500 nationts are bespital medication similable (74/450	No details about inclusion or completeness
38		 ICD9 diagnosis SAH and/or intra- 		Inpatient mortality.		469/500 patients pre-hospital medication available (71/469 taking either anti-coagulants or anti-platelts)	of trauma registry.
39		parenchymal					Attrition: Low Risk
40		contusion-				156/500 transfers	Only inpatient measures
41							

Г	•	confirmed with					
		radiology report and neurosurgical				No patients had seizures.	Prognostic factor measurement: Mod risk Incomplete information regarding
		consult note- if				No patients had neurological decline.	medications.
		disagreement scan re-reviewed if not				No patients underwent delayed neurosurgical intervention.	May be other inaccurate recording of
)		clear patient excluded				No inpatient mortality	factors.
		Excluded:				No inpatient mortality	Outcome measures: Mod risk
<u>'</u>		 Diagnosis extra or subdural 					Only inpatient related outcome measures. Patients may have been discharged and
		hematoma					deteriorated and presented to other
5		 Penetrating injuries 					hospitals. Confounding Factors: Mod risk
3		Fatal extra-cranial					Cohort includes patients with multiple
(injuries • CSF leak					injuries- only excluded if died from other injuries.
é		Aneurysmal SAH Delayed					Statistical techniques: N A
)		 Delayed presentation 					None presented
))							Narrative synthesis- further evidence SAH
- -							low risk.
	Pruitt et al 2016	Level 1 Trauma Centre Chicago	Retrospective cohort study	Clinical deterioration (defined as decrease in	Age Gender	1185 GCS13-15 with CT detected injuries	Study Recruitment: High risk
5	Chicago	2009-2013	stady	mental status, worsening	Method of arrival	814 admitted directly to hospital- poly-trauma, social	Neurosurgeons have admitted higher risk
3	USA	Inclusion criteria:	Aim Assess if mTBI patients	neurologic exam or death)	Whether transfer Comorbidities	reasons or as neurosurgeons felt high risk.	patients we can combine outcomes from both admitted and ED observed patients to
1		Initial GCS13-15	with intra-cranial	Neurosurgery during	Anticoagulant use	371 left under care of ED. Of these, 239/371 transferred ED	give an unbiased estimate.
3		 16 and older Traumatic intra- 	haemorrhage can be managed to an ED	admission.	Mechanism of injury	obs unit. 132/371 discharged directly from the ED after a period of observation.	Attrition: Med Risk
,)		cranial bleed or	observation unit	Progression on CT.	Initial GCS,		Only a proportion of patients are followed
í		skull fracture Identified on			Neurological examination	Admitted patients Clinical deterioration 15/814 Worsening CT 27/814	up- does not describe the mechanism for this or how consistent follow up is e.g. did
2		 Identified on electronic ED 			Alcohol	Neurosurgery® 33/814	they all get repeat CT scans
3		system using ICD 9 classification			intoxication Initial platelet count INR	Composite outcome 75/814	Prognostic factor measurement: Medium
1		system			Initial CT results	ED obs unit	risk
		Admitted to ED			Follow-up CT results,	Clinical deterioration 20/239 Worsening CT 211/239	Dependent on CT scan reports and written
,		observation unit			Neurosurgical	Neurosurgery 3/239	documentation
,		All patients received a			recommendations	Composite outcome 14/239	Outside and a service Mand state
í		neurosurgical consultation			Cranial CT data	Medical admission 4/239 Trauma/neurosurgery admit 8/239	Outcome measures: Mod risk Clinical deterioration not well defined and
)		Consultation			were collected	Follow up 190/239	very broad.
L					from attending	Delayed Neurosurgery 20/239	

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1 2 3 4							
5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23			70/6		radiologist reports- type and size of detected injury	Post traumatic seizure 3/239 Concussive symptoms 16/239 Discharged ED Follow up 111/132 Delayed Neurosurgery 1/132 Post traumatic seizure 2/132 Concussive symptoms 8/132 Figures from table- author has confirmed this is correct: 155 isolate SAH- 0 no clinical or radiological deterioration or cases of neurosurgery. 161 SDH- 6 CT deterioration, 3 planned neurosurgical outcomes. 0 deteriorated clinically 1 neurosurgery greater then 3 weeks later following outpatient assessment. 30 contusion 5 worsening CT scans. Nil clinical deterioration or emergency neurosurgery. 5 extradural- nil deterioration or neurosurgery Of sample 1053 mean/median age=59 11% anticoagulated. Of sample 1185 mean median age=59 10% anticoagulated	Confounding Factors: Low risk Included patients with polytauma and significant comorbidities Statistical techniques: High Risk None presented but data presented in table and text do not match up Paper shows patients admitted to hospital by neurosurgeons have worse outcomes/more likely to require neurosurgery. Does show that in America some of this patient population discharged directly from ED. Consistent with the model used locally in Hull.
24 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40 41	Deepika et al 2013 Bangalore India	Patients admitted tertiary neurosurgical centre 3 months Jan-March 2010. Patients identified on a TBI registry Inclusion criteria: GCS 13-15head injury Underwent CT scan Either negative CT or Isolated traumatic subarachnoid Matched comparison between patients - ve CT and SAH Excluded: Does not state	Retrospective cohort study Aim To assess whether GCS13-15 patients with traumatic subarachnoid haemorrhage have the same outcomes as mTBI patients with -VE CT scans	Prospective 1 year telephone assessment of : GOSE Rivermead post concussion questionnaire Rivermead Head injury follow up questionnaire	Age Sex Mechanism of injury- RTC Fall LOC Seizure Location of SAH Whether multiple bleeds Thickness greater or less than 5mm	34/1628 mTBI patients isolated traumatic subarachnoid haemorrhage 18/34 patients available for follow up at 1 year Good GOSE Rivermead scores comparable to 16 normal CT controls	Study Recruitment: Low risk Cohort identified in TBi registry which is part of normal practice. Is retrospective so limited by accuracy of medical notes. Attrition: High Risk Small sample- with large proportion lost to followup. Prognostic factor measurement: Medium risk Dependent on CT scan reports and written documentation Outcome measures: High risk 1 year too long Confounding Factors: Medium risk No control for other injuries or comorbidities
41 42 43 44 45 46 47			Mary A	nn Liebert, Inc, 140 H	luguenot Stree	t, New Rochelle, NY 10801	

Kreitzer et al 2014 2001-2010 Study Cohort Study Study CT head findings Age CT head findings Age hours in ED met the inclusion criteria Too poor quality to include Too poor quality to include Study Recruitment: Mod risk hours in ED met the inclusion criteria Identified through repeat CT imaging in ED			adults only but age range 15-672					Statistical techniques: N/A
2014 202.0200 study 202.0201 study			range 15-6/					Too poor quality to include
Constitution Control		Kreitzer et al	Level trauma center	Retrospective cohort	Death within 30 days	CT head findings	323/1011 patients that under-went 2 CT head within 24	Study Recruitment: Mod risk
USA dentified from colort of patients undergone 2 CT within the ED within 3 hours 1 crit in mTB with CH. If crit in mTB with CH. If CT if mTB with		2014	2001-2010	study	Neurosurgical intervention	Age	hours in ED met the inclusion criteria	Identified through repeat CT imaging in ED-
patients undergone 2 CT at least 6 hours after within the ED within 24 hours Inclusion criteria: • GCS 14.15 and blunt head highry® • Presented within 24 hours injury • Intra-cranial bleed first CT defined extradural, sundural, SAH, intra-cerebral and cerebral contusion • 2" CT within 24 hours Excluded: • Impatient died prior to FD evaluation Abnormal observations • Penetrating injury • CT scans interpreted at different hospital • Coagulopathy either inherited or acquired • NR-1.4 (even if taking wardarin) • Platelets less than 50 • Any non-head injury and adding a part of the properties of the p		Cincinnati			within 2 weeks	Race		relies on all of cohort having repeat scans
within the ED within 24 1" CT if mTBI with ICH. If hours CT and patients stable discharge from ED. Inclusion criteria: • GCS 24-15 and built head injury? • Presented within 24 hours injury • Intra-cranial bleed first CT defined extradural, sundral, SAH, intra-cerebral and cerebral contusion. • 2" CT within 24 hours Excluded: • Incomplete notes • Pregnant • Intubated prior to ED evaluation • Ahnormal observations • Penetrating injury • CT scans interpreted at different hospital • Coagulopathy either inherited or acquired • ININ-1.4 (even if taking warfarin) • Platelets less than 50 • Any non-head injury amendating)	USA	Identified from cohort of		Return to the Ed within 7	Sex	After second CT	·
To and patient stable discharge from ED. Inclusion criteria: • GCS 4.15. and blum head injury? • Persented within 2.4 hours injury. • Intra-creatial bleed first CT defined extradural, saudural, saudural					days of discharge		1 - ,	undergoing second scan being missed
discharge from ED. Apallents died (3 admitted 1 discharged) States death in discharged patients retrured to head injury had further fall. Also 1 other patient with mTBI and little with mTBI and l	,					background	, , ,	
Inclusion criteria: G. G. 14-15 and blunt head rijuryd of Presented within 24 hours injury of Intra-cranial bleed first. CT defined extradural, sundural, SAH, intra-cerebral and cerebral contusion. 2 "CT within 24 hours injury observations observations. Excluded: Incomplete notes Pregnant Inclusion criteria: Alm: Assess outcomes for patients with mTBI and ICH. Assess o			hours				206/323 discharged	
Begin and bluth read injury? Presented within 24 hours injury Intra-cranial bleed first CT defined extradural, sundural, SAH, intra-creebral and cerebral contusion Presented within 24 hours injury Intra-cranial bleed first CT defined extradural, sundural, SAH, intra-creebral and cerebral contusion Presented within 24 hours Excluded: Incomplete notes Pregnant Intubated prior to ED evaluation Abnormal observations Presentating injury CT CS cans interpreted at different hospital Coagulopathy either inherited or acquired NRS-1.4 (even if taking warfaini) Platelets less than 50 Any non-head injury mandating Alm: Assess outcomes for patients related to head injury had further fall. Also 1 other patient dies of septic shock. Alm: Assess outcomes for patients relateding and indicated further fall. Also 1 other patient dies of septic shock. Anexample further fall. Also 1 other patient dies of septic shock. Mean/median age 42 Percent anticoagulated=0 Pregnate Mean/median age 42 Percent anticoagulated=0 Outcome measures: low risk Reasonable outcome measures: Confounding Factors: Low risk Controls for comorbidities and othe injuries Statistical techniques: N/A Statistical techniques: N/A	3			discharge from ED.				. •
blunt head injury® Presented within 24 hours injury Intra-cranial bleed first CT defined extradural, sundural, SAH, intra-cerebral and cerebral contusion 2 d' CT within 24 hours Pregnant Intubated prior to ED evaluation Abnormal observations Pregnant Intubated prior to ED evaluation Abnormal observations Pregnant Coagulopathy either inherited or acquired INRD-1.4 (even if taking warfarin) Predicted sets than 500 Any non-head injury mandating	-							•
Presented within 2 hours injury Intra-cranial bleed first CT defined extradural, sundural, SAH, intra-crebral and cerebral contusion 2 "CT within 24 hours Excluded: Incomplete notes Pergenant Intubated prior to ED evaluation Abnormal observations Penetrating injury CT scans interpreted at different hospital Coagulopathy either inherited or acquired NNR-1.4 (even if taking warfarin) Prognostic factor measurement: Mediur risk States that some CT are reported by radiology trainees overnight and the corrected by attending radiologists the new day- unable to quantify how muc inaccuracy there is. Outcome measures: low risk Resonable outcome measures: Confounding Factors: Low risk Controls for comorbidities and othe injuries Statistical techniques: N/A Statistical techniques: N/A	5							• •
24 hours hijnry Inter-caralial beded first CT defined extradural, sundrul,							further fall. Also 1 other patient dies of septic snock.	attending at other ED
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first CT defined extradural, sundural, SAH, intra-cerebral and cerebral contusion. • 2"d CT within 24 hours Excluded: • Incomplete notes • Pergenant • Intubated prior to ED evaluation • Abnormal observations • Penetrating injury • CT scans interpreted at different hospital • Coagulopathy either inherited or acquired • NR1-1.4 (even if taking warfarin) • Platelets less than 50 • Any non-head injury mandating				ICII			2 nourosurgical interventions (all admitted)	_
extradural, sundural, SAH, intra-crebral and cerebral contusion 2 ° CT within 24 hours Excluded: Intubated prior to ED evaluation Abnormal observations Penetrating injury CT scans interpreted at different hospital Coagulopathy either inherited or acquired None re-admitted and some planned-removal of sutures. Individuals and the corrected by attending radiologists the new day- unable to quantify how muc inaccuracy there is. Does state 32% of repeat scan normal Outcome measures: low risk Reasonable outcome measures Confounding Factors: Low risk Controls for comorbidities and othe injuries Penetrating injury CT scans interpreted at different hospital Coagulopathy either inherited or acquired NNONE re-admitted and some planned-removal of sutures. Mean/median age= 42 Percent anticoagulated=0 Outcome measures: low risk Reasonable outcome measures Confounding Factors: Low risk Controls for comorbidities and othe injuries Statistical techniques: N/A	3							
sundural, SAH, intra-cerebral and cerebral contusion • 2 rd CT within 24 hours Excluded: • Incomplete notes • Pregnant • Intubated prior to ED evaluation observations • Abnormal observations • Penetrating injury • CT scans interpreted at different hospital • Coagulopathy either inherited or acquired • NRS-1.4 (even if taking warfarin) • Platelets less than 50 • Any non-head injury mandating)							
intra-cerebral and cerebral contusion 2 " CT within 24 hours Excluded: Incomplete notes Pregnant Intubated prior to ED evaluation Abnormal observations Penetrating injury CT scans interpreted at different hospital Coagulopathy either inherited or acquired INRS-1.4 (even if taking warfarin) Platelets less than 50 Any non-head injury mandating)		· ·				None re damitted and some planned removal of satures.	
cerebral contusion • 2" C within 24 hours Excluded: • Incomplete notes • Pregnant • Intubated prior to ED evaluation • Abnormal observations • Penetrating injury • CT scans interpreted at different hospital • Coagulopathy either inherited or acquired • INR>1.4 (even if taking warfarin) • Platelets less than 50 • Any non-head injury annadating			, , ,			7 // .	Moon/modian ago- 42	, ,
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observations Penetrating injury CT scans interpreted at different hospital Coagulopathy either inherited or acquired INR>1.4 (even if taking warfarin) Platelets less than 50 Any non-head injury mandating			ED evaluation					
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Ding et al 2012 China Chin							
China control trial comparing to previous part of the properties o	Ding et al	-64	Annears to be a random	GCS at discharge	CT scan results	32/89 nationts in routine CT group GCS13-15	Study Recruitment: High risk
GCS13-15 Excluded: Immediate neurosurgery Died within 3 days Severe multiple injuries Failed to undergo a repeat CT head Huynh et al 2006 USA Huynh et al 2004-2005 USA Level 1 trauma centre 2006 USA Level 1 trauma centre 2006 USA Identified case note review To assess whether neurosurgical review is necessary in GCS 15 Blunt trauma to head GCS15 Abnormal CT head Excluded: Normal initial CT head Legith of admission less than AB hours Retrospective cohort cohor	2012 Neurosurg Center	china 2009-2010 Inclusion criteria: All patients with TBI with evidence of intra-cranial haemorrhage-	control trial comparing outcomes in patients with traumatic intra-cranial haemorrhage assigned either to a routine repeat CT or CT only if	Surgical and medical interventions secondary to	Initial GCS Mechanism of Injury Coagulation INR	2/32 worse CT scans No patients had neurosurgery or altered medical management	Allocation to intervention and rintervention arm not clearly explair states via random number generator Attrition:Low Risk Low risk- inpatient outcomes
Huynh et al 2006 Level 1 trauma centre 2006 2004-2005 Identified Case not review To assess whether neurosurgical review is necessary in GCS 15 establish shad Excluded: Normal initial CT head Excluded: Normal initial CT head Etength of admission less than 48 hours Albanus Al		GCS13-15 Excluded: Immediate neurosurgery Died within 3 days Severe multiple		0,)/.		risk No re-reporting of CTS Outcome measures: Medium risk No outcome measures after discharge Confounding Factors: Low risk
USA Identified case note review Aim To assess whether neurosurgical review is necessary in GCS 15 patients with intra-cranial injuries Blunt trauma to head GCS 15 Abnormal CT head Excluded: Normal initial CT head Length of admission less than 48 hours Aim To assess whether neurosurgical intervention CT Neurosurgical intervention Neurosurgical intervention CT Injury ISS LOC Amnesia Associated injuries Injury ISS LOC Amnesia Associated injuries Injury ISS LOC Amnesia Associated injuries No consistent measure of deterioration O/56 neurosurgical interventions O/56 deaths Prognostic factor measurement: Med risk No re-reporting of CTS Outcome measures: Medium risk No outcome measures after discharge		repeat CT head Level 1 trauma centre	·	_		56 patients met inclusion criteria	Statistical techniques: N/A Study Recruitment: Medium risk
head injuries o GCS 15 o Abnormal CT head Excluded: o Normal initial CT head o Length of admission less than 48 hours head o Jo56 neurosurgical interventions o Jo56 deaths O/56 neurosurgical interventions O/56 deaths O/56 neurosurgical interventions O/56 deaths Prognostic factor measurement: Med risk No re-reporting of CTS Outcome measures: Medium risk No outcome measures after discharge		Identified case note review Inclusion criteria:	Aim To assess whether neurosurgical review is	СТ	Injury ISS LOC Amnesia	Of these 4: 2/56 patients had fall in GCS to 14 from 15 1/56 given mannitol due to worse CT	review Higher risk group as admitted for at leas
head • Length of admission less than 48 hours		headGCS 15Abnormal CT head	•			0/56 neurosurgical interventions 0/56 deaths Mean/median GCS=15	Low risk- inpatient outcomes Prognostic factor measurement: Medrisk
		headLength of admission less than				Mean/median age= 41	Outcome measures: Medium risk No outcome measures after discharge
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						Statistical techniques: N/A
Almenawer et al 2013 Ontario Canada	Neurosurgical centre Ontario, Canada 2006-2011 Identified from trauma database Inclusion criteria: GCS13-15 Blunt traumatic head injury Age>17 Intra-cranial injury CT head Repeat CT scan	Retrospective cohort study + meta-analysis to assess whether repeat CT imaging necessary in mTBI with intra-cranial haemorrhage	Intervention including: Mannitol or hypertonic saline Surgical intervention including ICP bolt or craniotomy Neurological changes: decrease GCS, cranial nerve change, vomiting and headache	Demographics GCS ISS	1121 patients with mTBI and ICH 445 met inclusion criteria 91/445 worse CT 21/445 patients neurosurgical outcomes (all preceded by clinical deterioration prior to repeat ct) 4/445 patients medical intervention 2/4 medical outcomes= treated with mannitol due solely worse CT other 2 treated due to clinical deterioration. Mean/median GCS=14.5	Study Recruitment: High risk Dependent on accuracy of trauma database Large proportion of mTBI patients with ICH did not meet inclusion criteria- selection out of higher risk patients that did not undergo repeat imaging Attrition:Low Risk Low risk- inpatient outcomes Prognostic factor measurement: Medium risk No re-reporting of CTS
	Excluded: No repeat CT scan Previous caniotomy Cranial pathology Coagulopathy Immediate Neurosurgery Patients divided into those underwent intervention due to clinical deterioration or due to repeat CT				Mean/median age= 45 Percent anticoagulated=0	Outcome measures: Medium risk No outcome measures after discharge Confounding Factors: Low risk No control for poly trauma Statistical techniques: N/A
Sifri et al 2004 USA	findings Level Trauma Centre New jersey 1999-2001 Inclusion criteria: GCS 14-15 Blunt traumatic head injury Age>15 Intra-cranial injury CT head Repeat CT Excluded:	Retrospective Cohort Study: To assess the value of routine repeat CT imaging in mTBI patients with intra-cranial haemorrhage	Worse CT Inpatient neurological deterioration- abnormal neurology- confusion, disorientation or drowsiness Inpatient neurosurgical interventions	CT results as abstracted from radiologist and neurosurgeons reports. Best ED GCS Demographics	243 patients with mTBI and ICH 18/243 excluded as no repeat CT- neurosurgeon ruled insignificant lesion 202/243 included as met the rest of inclusion criteria At 24 hours: 151/202 persistently normal or improving neurology 51/202 persistently abnormal or worsening neurological examination	Study Recruitment: Medium risk Selection out of patients not undergoing repeat CT hea dimaging Attrition:Low Risk Low risk- inpatient outcomes Prognostic factor measurement: Medium risk The definition of abnormal neurology is loose and not clear when it developed- not an admission criteria factor

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1 2 3 4							
5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24 25 26 27 28 29 30 31 32 32 32 33 34 34 35 36 36 37 37 37 37 37 37 37 37 37 37 37 37 37	Phelan et al 2014 Dallas USA	History of brain injury Coagulopathy including known bleeding disorder or taking warfarin Immediate neurosurgical intervention including transfer to ICU Level 1 Trauma Centre Dallas Texas 2010-2012 Patients identified on TBI data base Inclusion criteria: Intracranial haemorrhage TBI Patients divided into SAH and non SAH bleed All GCS but data for GCS13-15 patients presented Excluded: Ages less than 18 Pregnant Prisoners	Retrospective Cohort Study Assess whether outcomes for mTBI with isolated traumatic subarachnoid differ for other kinds of intra-cranial bleeds	Worse repeat CT imaging if any Death Craniotomy	CT findings as reread by a study team member Age ISS HAS Emergency department GCS	50/202 worse CT 5/202 required neurosurgery- all had persistent or worsening neurology 1/202 died all in the persistently abnormal/ worsening neurology group No clear measure of deterioration Mean/median GCS=14.7 Mean/median age= 44 Percent anticoagulated=0 77 patients GCS13-15 and traumatic SAH 27/77 scheduled repeat CT 3/27 worse CT 50/77-no routine repeat CT 4/50- unscheduled repeat CT 1/50- clinical deterioration and worse CT 4/77 worse CT 0 neurosurgical intervention	Outcome measures: Medium risk No outcome measures after discharge Confounding Factors: Low risk No control for poly-trauma and comorbidites Statistical techniques: N/A Study Recruitment: Low risk Dependent on accuracy of trauma registry Attrition:Low Risk Low risk- inpatient outcomes Prognostic factor measurement: low risk Does not really assess prognostic value of factors measured Outcome measures: Medium risk No outcome measures after discharge Confounding Factors: Low risk No control for poly-trauma and comorbidites Statistical techniques: N/A
33 34 35 36 37 38 39 40	Homnick et al 2012 New Jersey USA	New Jersey Medical School Level 1 trauma centre 2002-2005 Inclusion criteria:	Retrospective Cohort Study Establish how long intra- cranial bleeds in mTBI continue to expand	Neurosurgical intervention Progression on CT-repeat CTs as discretion of neurosurgeon	Age Sec Pre-injury anti- coagulation Mechanism ISS Initial GCS	341 patients in study (85 mTBI patients with bleeds excluded as no F/U scan) 72/341 intubated in ED 105/341 progression on CT 13/341 death- 9 due to TBI 4 other causes 12/341 neurosurgical intervention Mean/median GCS=14.6 Mean/median age= 47	Study Recruitment: Medium risk Selection out of lower risk patients that did not have repeat CT imaging Attrition:Low Risk Low risk- inpatient outcomes Prognostic factor measurement: low risk Does not really assess prognostic value of factors measured
41 42 43 44 45 46 47			Mary A	nn Liebert, Inc, 140 H	luguenot Stree	t, New Rochelle, NY 10801	

• • • • •	bleed, contusion, subdural, extra-dural or SAH ded: Penetrating trauma Injury >24 hours previously Previous neurosurgery Non-traumatic mass on CT Immediate neurosurgery	Period			Percent anticoagulated=2	Outcome measures: Medium risk No outcome measures after discharge Confounding Factors: Medium risk No control for poly-trauma and comorbidites Statistical techniques: N/A
2011 Karachi Pakistan Inclus Exclus All pa	probability ecutive sampling sion criteria: GCS14-15 All ages-15% sample children mean age 36 2 SD 18 TBI with positive initial CT intra- cranial injury	Retrospective Cross-sectional study Aim: Assess the utility of repeat CT scanning in mTBI patients with intracranial injuries without clinical or neurological deterioration	Worse CT	Age Gender Initial GCS Mechanism of injury CT findings	275 patients met inclusion criteria (note states 255 contusion haematoma) 17/275 worse CT No patients required neurosurgery Mean/median GCS=14.7 Mean/median age= 36 Percent anticoagulated=0	Study Recruitment: Medium risk Does not adequately define deterioratio or over what period Attrition:Low Risk Low risk- inpatient outcomes Prognostic factor measurement: low risk Does not really assess prognostic value of factors measured Outcome measures: Medium risk No outcome measures after discharge Confounding Factors: Medium risk No control for poly-trauma an comorbidites Statistical techniques: N/A Overall Includes kids and quite a different population than North America and Europe
Sates	I I 2 trauma centre 5 2007-2011 sion criteria:	Retrospective Cohort Study Assess whether repeat CT imaging in GCS14-15 mTBI	Increased size of bleed second CT Clinical deterioration-decrease in GCS	Age Sex Initial and follow- up GCS CT findings	68 patients 4 patients transferred to neurosurgery (2 routine) 8/68 patients worse CT	Study Recruitment: Medium risk Identified on trauma data base with patients with incomplete data excluded Does not present number of these patients Also excludes patients transferred

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1 2 3 4							
5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24		GCS14-15 TBI with positive initial CT intracranial injury including subdural, extra-dural, subarachnoid and intra-cerebral bleeds Only data for adults presented Excluded: Patients with incomplete data Transferred to neurosurgery immediately No repeat CT All patients had a repeat CT within 12 hours	with intracranial injury justified	New motor or sensory symptoms Severe headache or vomiting		12/68 mild deterioration 28 patients intra-parenchymal bleed 1/28 worse CT 3/28 neurological deterioration 1/28 transferred to neurosurgery (not patient with worse CT) 7 patients extra-dural 1/7 worse CT 0/7 neurological change 1/7 transferred to neurosurgery 20 patients sub-durals 3/20 worse CT 4/20 neurological deterioration 1/20 neurosurgery 13 patietns SAH 3/13 increase in size bleed 5/13 neurological deterioration 1/13 transferred to neurosurgery Mean/median GCS=14.8 Mean/median age= 56	immediately. Likely to be lower risk smaple than population of interest. Attrition:Low Risk Low risk- inpatient outcomes Prognostic factor measurement: low risk Does not really assess prognostic value of factors measured Outcome measures: Medium risk No outcome measures after discharge Confounding Factors: Medium risk No control for poly-trauma and comorbidites Statistical techniques: N/A
25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40 41	Brown et al 2007 Los Angeles USA	Los Angeles Level 1 trauma center 2003-2004 Inclusion criteria: • All patients with blunt head trauma and intra-cranial bleed initial CT. Presents data for GCS13-15 Excluded: • Immediate neurosurgery • Died within 24 hours • Does not state just adults but seems only for adults	Prospective Cohort Study Aim To identify patients with head injuries that benefit from routine repeat CT imaging	Need for neurological intervention- either medical or surgical (medical= sedatives, mannitol or hyperventilation and surgical= ICP monitor and craniotomy) Mortality	Age Gender Mechanism of Injury ISS Admission GCS Results of CT- interpreted by attending radiologist	354 patients all GCS scores with intra-cranial bleed 37 direct to craniotomy 43 dies within 24 hours 274= study population 142/274= mTBI GCS13-15 15/142 had clinical deterioration 27/142 had worse CT scans (only 72/142 had repeat imaging) 5/142 had medical or neurosurgical intervention 3/142 died Mean/median GCS=14 Mean/median age= 43	Study Recruitment: Mod risk Removal of patients that died within 24 hours may lead to this sample being a lower risk group than population of interest Attrition: Low Risk Low risk- inpatient outcomes Prognostic factor measurement: low risk Does not really assess prognostic value of factors measured Outcome measures: Medium risk No outcome measures after discharge Confounding Factors: Medium risk No control for poly-trauma and comorbidities- Statistical techniques: N/A
42 43 44 45 46 47			Mary A	nn Liebert, Inc, 140 H	uguenot Stree	t, New Rochelle, NY 10801	

Tennesse USA So months from Jan USA To assess whether scheduled repeat CT head imaging is indicated in TBI All patients with blunt head trauma and evidence TBI on initial CT. Presents data for GCS13-15 Age 18- Excluded: Penetrating mechanism Interventions for unclear indications Died before second CT All patients repeat CT at 6-8 hours after admission Klein et al 2010 Klein et al 2010 Klein et al 2010 To assess whether scheduled repeat CT head imaging is indicated in TBI Age 28- Age 48- Age		(mean age 44 +/- 19)					
USA 2001 To assess whether scheduled repeat CT had linclusion criteria: All patients with blunt head trauma and evidence TBI on initial CT. Presents data for GCS13-15 Age 18t Excluded: Penetrating mechanism Immediate neurosurgery Immediate neurosu	2010	Level 1 trauma centre		_	ISS		Dependent on case note review. Patient
Neurological changer-reduced GCS, pupillary change, increased ICP or loss of brain and evidence TBI on initial CT. Presents data for GCS13-15 Age 18+ Excluded: Penetrating mechanism Interventions for unclear indications Died before second CT All patients repeat CT at 6-8 hours after admission Klein et al 2010 Israel Aim: Assess the outcome of low risk patients with ICB Neurosurgical interventions for validation at the interventions based on repeat CT 3/14 medical interventions based on repeat CT 4 Mendical interventions based on repeat CT Antitipated tu se Prognostic factor measurement: M Mean/median age = 42 Mean/median		2001	scheduled repeat CT head		Age Gender	bolt)	
on initial CT. Presents data for GCS13-15 • Age 18+ Excluded: • Penetrating mechanism • Immediate neurosurgery • Interventions for unclear indications • Died before second CT All patients repeat CT at 6-8 hours admission Klein et al 2010 Israel Klein et al 2010 Israel Klein et al 2010 Israel Generating tentric in sirael. None had access to neurosurgery on site. Anticoagulant use Antiplatelet use PT, aPPT, INR CT findings Anticoagulant use Antiplatelet use PT, aPPT, INR CT findings Confounding Factors: Medium risk No control for poly-trauma Statistical techniques: N/A None done Age Als Als Study Recruitment: Low risk Dependent on completeness of registry admission Age Als Als STATISTIC TRANSPORTED Age Als SS 27/323 required neuro-rehab 2/323 died 35/323 neurosurgery Antition: Low Risk Outcome measures: Confounding Factors: Medium risk No control for poly-trauma Statistical techniques: N/A None done Study Recruitment: Low risk Dependent on completeness of registry registry Attrition: Low Risk Only inpatient outcome measures		 All patients with blunt head trauma 	imaging is indicated in TBI	GCS, pupillary change,	injury History of vascular		
Age 18+ Excluded: Penetrating mechanism Interventions for unclear indications Died before second CT All patients repeat CT at 6-8 hours after admission Klein et al 2010 Israel		on initial CT. Presents data for	16		Anticoagulant use Antiplatelet use	Mean/median age= 42	Prognostic factor measurement: Mod risk Does not explain how CT scans reported
mechanism Immediate neurosurgery Interventions for unclear indications Died before second CT All patients repeat CT at 6-8 hours after admission Klein et al 2010 Israel Klein et al 2010 Israel Klein et al 2010 Israel All patients repeat CT at 6-8 hours after admission Retrospective Cohort Study Nortality Neurosurgical intervention Neurological status at discharge Aim: Assess the outcome of Identified ICD9 codes on Identified ICD9 codes on Identified ICD9 codes on Confounding Factors: Medium risk No control for poly-trauma Statistical techniques: N/A None done Study Age AlS Neurosurgical intervention Neurological status at discharge Aim: Assess the outcome of Identified ICD9 codes on Immediate No control for poly-trauma Statistical techniques: N/A None done Age AlS Study Recruitment: Low risk Dependent on completeness of registry Attrition: Low Risk Only inpatient outcome measures		Age 18+ Excluded:			, ,		
Interventions for unclear indications Died before second CT All patients repeat CT at 6-8 hours after admission Klein et al 2010 3 regional trauma centres in Israel. None had access to neurosurgery on site. Aim: Aim: Assess the outcome of Identified ICD9 codes on I		mechanism • Immediate					
All patients repeat CT at 6-8 hours after admission Klein et al 2010 Strael Study Study		 Interventions for unclear indications 					• •
Klein et al 2010 Study S		СТ					
Israel centres in Israel. None had access to neurosurgery on site. Aim: Assess the outcome of Identified ICD9 codes on I		6-8 hours after				(O)*	
neurosurgery on site. Aim: Assess the outcome of Identified ICD9 codes on Identified ICD9 codes		centres in Israel. None	'	Neurosurgical intervention	AIS		Dependent on completeness of trauma
Identified ICD9 codes on low risk patients with ICB Only inpatient outcome measures				<u> </u>	ISS	2/323 died	
			low risk patients with ICB				
Inclusion criteria: hospitals without e GCS13-15 hospitals without neurosurgical services hospitals without nous replain how CT scans repo		Inclusion criteria:	hospitals without			0/77 died 0/77 neurosurgery	Prognostic factor measurement: Mod risk Does not explain how CT scans reported
• ICD9 code for intra-cranial bleed. One hospital transferred Outcome measures: Mod risk Non-transfer on basis of: No F/U after discharge		intra-cranial bleed.					
all patients to single bleed = 5mm or contusion <1cm and noneurosurgical centre. Single bleed </= 5mm or contusion <1cm and nocoagulopathy Confounding Factors: Medium risk</td <td></td> <td>all patients to neurosurgical centre.</td> <td></td> <td></td> <td></td> <td>Single bleed <!--= 5mm or contusion <1cm and no-</td--><td>Confounding Factors: Medium risk</td></td>		all patients to neurosurgical centre.				Single bleed = 5mm or contusion <1cm and no-</td <td>Confounding Factors: Medium risk</td>	Confounding Factors: Medium risk
Other 2 hospitals transferred selected No control for poly-trauma Mean/median age= 39 Comorbidities		· ·				Mean/median age= 39	

				Journ	nal of Neurotra	uma	Page 82 of 139
1 2 3 4							
5 6 7 8		patients.					Statistical techniques: N/A None done
9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40 40 40 40 40 40 40 40 40 40 40 40 40	Sifri et al 2011 USA	Level 1 Trauma Centre New jersey 2002-2006 Inclusion criteria: Initial GCS 13-15 Blunt traumatic head injury Age 18+ Intra-cranial injury CT head-ICB or skull fracture Repeat CT Abnormal neurological examination at time of repeat CT Excluded: Immediate or planned neurosurgical intervention Normal neurology at time of repeat CT- normal neurology defined as GCS15, orientation to place, person or time, normal neurological exam, no symptoms from head injury- headache, vomiting, dizziness, lethargy Coagulopathy including known bleeding disorder or taking warfarin	Aim: To assess proportion of patients that have worse CT scans and neurosurgical interventions that have abnormal neurology when they have a repeat CT.	Progression of lesion on CT Surgical intervention- includes intubation Medical intervention GOSE at discharge	Demographics Acute deterioration in neurological Exam Persistently Abnormal Neurological exam Unknown whether change as intubated	21/107 deterioration 18/107 unable to assess neurology as intubated. 6 died Mean/median GCS=14.4	Study Recruitment: High risk High risk subgroup that have abnormal neurology at time of repeat CT imaging. Attrition: Low Risk Only inpatient outcome measures Prognostic factor measurement: Mod risk Difficult to assess deterioration in a retrospective study. Outcome measures: Mod risk No F/U after discharge Confounding Factors: Low risk Some control for comorbidities. Statistical techniques: N/A None done
41 42 43 44 45 46 47 48			Mary A	nn Liebert, Inc, 140 I	Huguenot Stree	et, New Rochelle, NY 10801	

Sunnlama	Pregnancy Spinal Cord Injury Ario Matagreea	P: Data Extracted	from Included Stu	dies		
аррістіс	intary wateraar 2				Prevalence of Outcomes N=26	
Reference	Population	Study Design	Outcome Measures	Prognostic factors assessed	Results	Quality Appraisal
ishijima et al 013	Multicenter gical sites Westery Gatrial Level 1	Retrospective Cohort Study	Initial ICU admission from ED	Age Initial GCS	11240 patients coded as bleeds 771 excluded due to missing data	Study Recruitment: Mod risk bias Dependent on accuracy on recording on
ncromento Synon et al 015	disorder Heidelberg Heighereity Sterified Germody codes	Betrespective Cohort \$1 was sess the variability of	Proportion of patients Receipting imaging care Proportion of patients Receipting imaging care	Initial BP Patientsp divided intosthose on no	79 ग्रन्थ मंद्रमाञ्चलाञ्च ineksien झर्चेन्ट्रांने teria. 37 no anticoagulation	trauma registry. Does have some quality Stydy Regruitroans: hyperiston Although high rates of anti-coagulation.
ermany ariability of	ritra-crafilal haemorrhage 2005-2010	ICU use in a cohort of patients with minor	Neurosurge Mintervention Rethanical ventilation	anticonsulants, as Aspirin, in Wasfaria	8명에 나 한국에 Mitted ICU, significant variation between sites 5 warfarin	Note initial GCS 15- lower risk group Attrition: Low Risk
U Use in	Inclusion criteria:	trampate outsomerania	Visop Ressort diagraph use	and POACS.	44944£ (5!1%) Kalaunical care intervention	Anthropatient purcome measures
dult patients	Inclusion en la faction de la	Hatingh hages different types of anti-coagulants	Transfusion blood product	AIS gender,	है,शब्दां २ भं ही ते छंडा हुति हो। intervention	Follow up only during hospital admission Prognostic factor measurement: Low risk
	cranial bleed CT head			trauma mechanism, comorbidities,	25% neurosurgery (18 patients) 43/70 repeat CT imaging-	May be miss-classified in medical notes Outcome measures: Mod risk
				CT findings,	2 deaths both on rivaroxaban	No F/U after discharge
				imaging,	Mean/median GCS=14.5	Confounding Factors: Low risk
				age, GCS scores,	Mean/median age= 67 Percent anticoagulated=16	No control for comorbidities.
				laboratory values	0.4	Statistical techniques: N/A None done
					t, New Rochelle, NY 10801	
		Manus A.		l A Church	t, New Rochelle, NY 10801	

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1 2 3 4							
5 6 7 8 9 10 11 12 13 14 15 16 17 18	with minor traumatic intra- cranial haemorrhages	 Age ≥ 18 years Traumatic ICH Initial ED GCS 15 ISS less than 16 	multiple trauma centres. 2)Estimate the proportion of minor traumatic intracranial haemorrhages patients admitted to ICU that do not receive an ICU intervention	Invasive monitoring		847/888 patients admitted ICU no crit care intervention Mean/median GCS=15 Mean/median age= 48	Prognostic factor measurement: Low risk Doesn't really apply as testing disposition not outcomes Outcome measures: Low risk No measure of outcomes after discharge, but study primarily about disposition. Does not report deaths. Confounding Factors: States IIS increases ICU admission- will be related to other injuries Statistical techniques: low risk N/A Overall Only GCS15 patients with low ISS.
20 21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40 41	Nishijima et al 2015 Sacromento USA Long-term Neurological Outcomes in Adults with Traumatic Intracranial Hemorrhage Admitted to ICU versus Floor	Level1 trauma centre 2008-2013 Inclusion Criteria:	Retrospective Cohort Study Aim compare long-term neurological outcomes in low- risk patients with traumatic intracranial hemorrhage (tICH) admitted to the ICU (intensive care unit) versus patients admitted to the floor.	Prospective long term outcome measure at 6 months Either GOS-E 8 fully recovered or GOS-E 1-7 not fully recovered	age sex, mechanism of injury initial ED GCS score, initial (SBP) heart rate, respiratory rate, blood alcohol level, AIS score ISS score INR Rotterdam CT score	188 met inclusion criteria 151/188 complete data= cohort 106 admitted ICU (70%) 45 admitted ED (30%) 1/151 patients neurosurgical intervention as inpatient 1/151 patient died as inpatient 78 (52%) GOS-E 8 at 6 months Does present analysis for outcome at 6 months GOSE but no inpatient measures of deterioration. Adjusted analysis, floor admission versus ICU had an odds ratio of 0.77 (95% CI [0.36-1.64]) for a GOS-E score of 8 at six months. Mean/median GCS=15 Mean/median age= 40	Study Recruitment: Mod risk bias Dependent on accuracy on recording on trauma registry and accuracy of case notes. Low risk group- GCS 15 and benign CT Attrition: Low risk Loss of 37 patients to follow up Prognostic factor measurement: Low risk As recorded in case notes so dependent on accuracy Outcome measures: Low risk Prospective follow up by trained staff using validated tool. Not clear what would happen to patients who died or deteriorated and attended a different hospital. Confounding Factors: Patients which are perceived as higher risk will be put on ICU, likely to be differences in comorbidities Statistical techniques: low risk Well presented- not really relevant to meta-
42 43 44 45 46 47			Mary A	nn Liebert, Inc, 140 H	luguenot Stree	et, New Rochelle, NY 10801	

							analysis
		CA					Only GCS15 patients with benign looking CT scans
Γ	Schaller et al	Level 1 Trauma centre	Retrospective cohort	Deterioration in neurological	Prognostic factors	110 patients met inclusion and exclusion criteria.	Study Recruitment: Low risk bias
	2015	Bern Switzerland	study/case series	status or need for	are the		Retrospective cohort review- reliant on
)	Switzerland	Jan 2006-Dec 2007		neurosurgery.	inclusion/exclusio	None deteriorated within the period of hospital	accuracy of written notes.
			Aim to assess if a specific		n criteria	observation, required neurosurgery or re-attended.	
<u>'</u>			group of patients with				Attrition: Mod risk
-			small bleeds can be		No comparison in	Mean/median GCS=14.6	Patients may have moved out of catchment
3		Inclusion criteria:	discharged from hospital		risk of	Mean/median age= 40	area of hospital without the researchers
1		 Admission GCS 13- 	without 24 hours of		deterioration in 2	Percent anticoagulated=0	being aware. Loss to F/U if re-presented
5		15	observation		groups.		different hospital.
		 Observed for 24H 					
2		 Localised intra- 					
'		cranial bleeds up					Prognostic factor measurement: Mod risk
3		to 5mm- this is					Reliability of case notes- may be incomplete
9		from the CCHR					Interpretation size of the bleed was taken from written radiology report ?reliability.
)		paper					from written radiology report freliability.
í		Exclusion Criteria: Bleeds > 5mm			7 .		
<u>'</u>		maximum					Outcome measures: Moderate risk
-		diameter					Study dependent on patients re-presenting
3		Multiple bleeds					at the same hospital following discharge if
1		History of bleeding					had delayed deterioration. Not clear how
5 I		tendency					patients died in the community would have
		Anti-coagulant or					been identified.
2		anti-platelet					
′		medication					Confounding Factors: Low risk
3		 Intoxication 					No obvious confounding factors
9		 Other injuries 					Cohort selection criteria including not living
1		Live alone					alone may select out high risk older
.		Live greater the 1H					patients.
.		from hospital					S 1. 1
-		·					Statistical techniques: N/A
3							
1							General comments:
5							Mean age 39.9 years and 25% caused by
							sporting injuries. ?Age as the confounding
2							low risk prognostic factor. Not generalizable
′							to older populations
3							
9							Small numbers
)							

Levy et al	Level 1 Trauma centre	Retrospective Cohort	ED disposition	Age (18-39)(40-	1144 patients admitted with mTBI but negative CT scan	Study Recruitment: Low risk bias
2011	Denver USA	Study	ICU admission	69)(70+)		Patients recruited from trauma registry
Colorado	Jan 1998-Dec 2008		Neurosurgery	Transfer status	117 with mTBI and traumatic SAH	depends on how good this is
USA		Aim	In-hospital mortality	Cause of injury		
	Inclusion criteria:	To assess whether	Progression of SAH on CT	GCS	1/117- progression on repeat CT scan	Only admitted patients- higher acuity
	Admission ED GCS	patients admitted with CT		Blood alcohol level		patients then discharged.
	13-15	–VE mTBI have different		Presence of skull	0/117 required neurosurgical intervention	
	 On trauma registry 	outcomes to patients with		fracture		Likely patients admitted for other reasons if
	 Blunt head trauma 	mTBI and traumatic SAH		CT report- divided	1/117 died (progression on CT)	CT negative TBI (although excludes other
	• ICD 850-850.99-			into		injuries).
	consistent with	Univariate and		small/medium/lar	4/1144 died	
	concussion (i.e. no	multivariate regression		ge based on		Attrition: Low risk
	detected injury by	used to examine		language included	All patients died >70	All inpatient outcomes
	CT)	covariates and		in report		
	Admitted to	relationship to outcomes		•	Logistic regression model tSAH versus concussion	
	hospital				ICU admit adjusted OR 8.87 (5.62-14.02) P<0.0001	Prognostic factor measurement: Mod risk
	AIS score 2 before				ICU LOS>1D OR0.29 (0.11-0.74) P=0.01	CT findings abstracted from CT reports-
	2008 or 1 / 2 in				Hosp LOS>1D OR1.07 (0.67-1.69) P=0.79	severity assigned by language- not actually
	2008 01 1 / 2 111				Mortality OR2.46 (0.27-22.17) P=0.42	used in regression model
					, , ,	
					Discharge to rehab	Outcome measures: Moderate risk
	Exclusion Criteria:				Age18-39 OR5.48 (0.25-121.70) P=0.28	Only inpatient outcomes- possibility of
	Patient admitted				Age 40-69 7.96 (1.91-33.11) P=0.004	discharge and deterioration.
	directly to hospital				Age >70 1.33 (0.50-3.53) P=0.56	discharge and deterioration.
	 Multiple injuries 				Age >70 1.33 (0.30-3.33) F =0.30	Confounding Factors: High risk
	AIS score >1 head					Patients admitted with CT negative TBI
	or other regions					likely to be frail or have other reasons for
	 Age less than 18 					· ·
	 Not admitted 					admission- this will affect outcome
						measures compared to SAH patients
					S S	admitted due to +ve CT.
						Charlestan Landon Invance Landon Internation
						Statistical techniques: Low risk
						Well presented.
						Can use for pooling for outcomes SAH-
						supports low risk sub-population
		Marv A	nn Liebert, Inc. 140 F	luquenot Stree	t, New Rochelle, NY 10801	
		y	,,	3-1-11-00	,,	

Levy et al 2014	Level III rural non-	Retrospective cohort	Length of stay	No comparison to	76/273 patients not transferred	Study Recruitment: Low risk bias
USA	neurosurgical unit in	Study	Mortality	patients that were	>50% injuries due to skiing/snow boarding	Retrospective cohort review- reliant on
	Rocky mountains April		Neurological deterioration	transferred	71% patients less then 55	accuracy of written notes.
	2007-Dec 2012	Aim	Neurosurgery			CT inclusion criteria are subject and
		Investigate outcomes	Re-admission in 90 days of		No patient deteriorated, died or required neurosurgery or	patients may have been transferred despite
	April 2007 patients with	after a novel non-transfer	discharge		required delayed transfer whilst admitted to hospital.	meeting non-transfer policy if clinicians
	small bleeds selectively	policy for mTBI patients	Inter-hospital transfer			were concerned.
	not transferred to	with small ICH introduced	Need for repeat CT		2 patients re-admitted within 90 days- 1 patient 6 weeks	
	neurosurgical unit	in a small rural trauma			following admission developed an acute on chronic	Attrition: low risk
		unit without			subdural- drained. 1 patient re-admitted with unrelated	Prognostic factor measurement: Mod risk
	Inclusion criteria:	neurosurgical cover			complaint.	Reliability of case notes- may be incomplete
	 Admission GCS 13- 					The definitions of bleed size are subjective.
	15				Mean/median GCS=14.7	-
	CT positive intra-				Mean/median age= 36	Prognostic Factors
	cranial injury				Percent anticoagulated=0	N/A
	Not transferred to				Ŭ	
	neurosurg unit in					Outcome measures: Moderate risk
	accordance with					Study dependent on patients re-presenting
	non-transfer					at the same hospital following discharge if
						had delayed deterioration.
	policy.					mad delayed deterioration.
	CT findings of small					Confounding Factors: Low risk
	SAH					_
	 Punctate or 					Age affect outcome and size of bleed
	minimal contusion					S 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1.
	 Punctate or 					Statistical techniques: N/A
	minimal intra-					
	cranial bleed					General points
	 Small SDH, no mass 					
	effect					Small numbers.
	Exclusion Criteria:					No comparator group- need to compare to
	 Any coagulopathy 					transferred patients outcomes.
	Basilar skull					
	fracture or					Patient not generalizable- v. young and
	evidence of CSF					atypical mechanism of injury (mostly winter
	leak					sports related).
	Extra-dural bleed					Likely that any patient clinicians felt risky
	Any significant					would have been transferred even if did
	contusion or					not meet transfer criteria- no way to check
	SDH/intra-cerebral					this.
	haemorrhage					
	Review and discussion of					
	CT and patient with					
	neurosurgeon if unsure					
	if should be transferred					

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1 2 3 4							
5 6 7 8 9 10 11 12 13 14 15 16 17 18		CO.	Perio				
19 20 21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40 41	Joseph et al 2013 USA The acute care surgery model: Managing traumatic brain injury without an inpatient neurosurgical consultation	Level 1 Trauma centre 2009-2011 (likely subset of patients presented below) Inclusion criteria: GCS13-15 Trauma Positive findings CT- skull fracture and/or ICH Exclusion Criteria: Pre-hospital anti- platelets or anti- coagulants	Retrospective cohort study- propensity matching 1:2 ratio patients managed solely by trauma surgeons versus patients that had neurosurgical consultation. Hypothesis Trauma surgeons can manage mTBI patients with CT detected intracranial haemorrhage without neurosurgical invlolvement	Hospital admissions ICU admissions Neurosurgical interventions ED visits after discharge Mortality Progression on CT imaging	Age Sex Initial GCS ISS Head-abbreviated injury score Neurological examination CT scan findings- type of skull fracture/type of ICH/size of bleed- reviewed by study investigator	404-GCS13-15 patients with CT detected injuries in study period. 270/404 used for this study 90/270- had neurosurgical consultations (NC) 180 no neurosurgical consultation. (no-NC) Whether neurosurgical consultation requested as discretion of non-specialist surgeon. Propensity matching in this study between 2 groups. 0/270 neurosurgical interventions, hospital mortality or readmissions either group. 78/90 no-NC and 158/180 NC admitted hospital (P=0.8) 18/90 no-NC and 80/180 NC admitted ICU (P=0.001) Routine repeat CT 18/90 no-NC 155/180 NC (P<0.001) No progression on any repeat CT 8% no-NC and 4% NC group re-attended ED. No readmissions. Mean/median GCS=15 Mean/median age= 30	Study Recruitment: High risk bias Subset of patients that meet inclusion criteria selected in order to facilitate propensity matching. Possible selection out of higher acuity patients as these will have al been referred to a neurosurgeon. Attrition: low risk In patient outcomes and documented ED re-attendances- low risk of patients being lost to follow up Prognostic factor measurement: Low risk All routinely collected clinical data apart from CT imaging which re-reviewed. Outcome measures: Mod risk Study dependent on patients re-presenting at the same hospital following discharge if had delayed deterioration. Confounding Factors: Mod risk Does not exclude patients with additional injuries

					Percent anticoagulated=0	
	6				Percent anticoagulated=0	Statistical techniques: High risk Does not outline how matched groups using propensity scoring
	4					General points
		· C//.				Small numbers.
						Likely reporting data reported else where.
AbdelFattah et	Level 1 trauma center	Prospective Cohort Study	Outcome measures during	Comparison	145 patients met inclusion/exclusion criteria.	Study Recruitment: low risk
al	Dallas Texas		hospital admission:	between groups:	92/145 for routine repeat CT	Prospective recruitment- states recruited all
2012		Hypothesis:		Age	53/145 for CT if deteriorated	eligible patients. Doesn't explain how
USA	Prospective recruitment 2010-2011	Repeat CT imaging in GCS13-15 with ICH,	Neurologic progression. Medical intervention	Sex Coagulation status	Selective group more likely aspirin use P=0.02 Routine repeat CT worse Head AIS score (P<0.001)	recruitment occurred.
		without neurological	Neurosurgical intervention	Anti-platelets	Otherwise groups comparable	Attrition: low risk
	Inclusion criteria:	progression, does not	Repeat CT imaging- worse CT	ISS		Follow up only for period in hospital
	• Adult with ICH	impact the need for	defined as worse by a	GCS	5/53 deteriorated and had a repeat CT + 1/53 had repeat	
	(note doesn't	neurosurgical	blinded		scan as started on warfarin	Prognostic factor measurement: Low risk
	explicitly state	intervention.	radiologist/neurosurgeon			Blinded appraisal of CT scans by researcher.
	2ndary to trauma-		giving qualitative measure of		1/145 patients died (due to other injuries)	
	but implied)	Patients divided into	bleed.		27/145 radiological deterioration	Outcome measures: Mod risk
	Excluded:	those 2 groups. Patients			9/145 patients intubated- states for other injuries	No F/U following discharge- missed delayed
	• Age<16	with planned repeat CT			Many Imadian CCS-14 F	outcomes, could have looked for re- attendance.
	• GCS<13	imaging and those with CT imaging if deteriorated.			Mean/median GCS=14.5 Mean/median age= 41	Doesn't report neurosurgical outcome
	Undergone	Allocation by			Percent anticoagulated=6	measures.
	planned or	neurosurgeon-no			rei cent anticoagulateu-o	measures.
	immediate	deviation from normal				Confounding Factors: High risk
	neurosurgery Transferred	practice.				Not isolated head injury- other injuries have
	patients				10.	clearly affected outcome measures
						Statistical techniques: Low risk
						None
						Small study with confounders regarding
Novel, et al	Hairrania Harris-1	Datus anastius Chart	Nouseausieel interpreting	A = 0	221/964 motionts CCC12 15 with ICD most included a suit and	outcomes.
Nayak et al 2013	University Hospital Newark New Jersey	Retrospective Chart Review	Neurosurgical intervention after 24 hours- craniotomy,	Age Sex	321/864 patients GCS13-15 with ICB met inclusion criteria 20% excluded because incomplete medical notes/transfers	Study Recruitment: Low risk Retrospective case note review- depends
2013	Level 1 trauma centre	VENIEM	ventriculostomy, ICP	Mechanism of	20/0 excluded because incomplete medical notes/transfers	on information being recorded correctly.
USA	2003-2008	Aim:	bolt/measurement	Injury	0/321 neurosurgical intervention-all within 24 hours of	on information being recorded correctly.
03/	2003 2000	To compare neurologic	borg measurement	GCS on arrival	admission	Attrition: Mod risk
	Inclusion criteria:	outcomes of MHI patients	Death in hospital	ISS		20% excluded because of incomplete notes
	Aged 18 and over	with an intra-cranial bleed		HAIS	No deaths	
	G	with a normal	Discharge disposition	GCS and		Prognostic factor measurement: Mow risk
1			0			

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5 6		Blunt trauma Intra-cranial bleed Admitted to	neurological examination managed with and without a repeat CT head	LOS hospital	neurological examination every 2 hours- routine	19/142 worse CT on repeat CT after 24 hours of admission 179/321 single CT	Neuroradiology reports taken at face value- no verification
7 8		Admitted to hospitalGCS13-15 on	scan	GOS at f/u clinic/ re- attendance if applicable	care on a flow sheet	142/321 routine repeat CT	Outcome measures: mod risk
9 10 11		arrival to EDGCS 15 24 hours after attendance to	70.			76/321 returned to F/U clinic- uneventful 14/321 returned to ED due to symptoms.	No uniform follow up of patients post discharge. Some patients had F/U clinic others didn't. Patients may presented after
12 13 14		ED Excluded: History brain disease, e.g.				Mean/median GCS=14.9 Mean/median age= 41	discharge to other sites. Confounding Factors: low risk None obvious
15 16		disease, e.g. dementia Previous brain injury e.g. CVA	, C	<i>h</i> ,			Statistical techniques: Low risk None completed
17 18 19 20		• Liver cirrhosis, renal disease, coronary artery disease, bleeding		0,			The inclusion/exclusion criteria have selected out all patients that are not GCS 15 at 24 hours. Different population than all GCS 13-15 patients with TBI on CT- probably
21 22 23		 or clotting disorder Unable to assess GCS due to drugs e.g. 					unable to pool this data. Does show patients that are GCS 15 at 24
24 25		sedation/intubatio n • Neurological					hours low risk.
26 27 28		deterioration leading to repeat CT • Aged less than 15				0/	
29 30	Anandaliyar at	Incomplete notes	Detroppositive colour	Donact CT often 24 hours of	Ago	F32 actions TDI and IQU	Charles Donniton and High viole
31 32 33 34	Anandalwar et al 2016 New Jersey USA	University Hospital Newark New Jersey Level 1 trauma centre 2009-20012	Retrospective cohort study Aim Assess the outcomes	Repeat CT after 24 hours of admission due to clinical concern or deterioration. Progression on any repeat CT	Age Sex Mechanism of Injury ISS	533 patients TBI and ICH 142 met the inclusion/exclusion criteria 47 underwent a routine repeat CT within 24 hours (violation of policy)- 0/47 neurosurgical, 1/47 had incidental finding on CT	Study Recruitment: High risk Patients at GCS15 at 24 hours- low risk group selected out- difficult to extrapolated to all GCS13-15 patients.
35 36 37		Inclusion criteria: • Aged 18 and over • Blunt trauma	following the implementation of a policy of observation only (no repeat CT imaging) for		AIS	95 no repeat routine CT within 24 hours 8/95 (non-violation group) had repeat CT >24 hours after	Does not compare outcomes in patient that adhered to and violated non-routine repeat CT head imaging. Potentially clinicians ordered routine repeat CT imaging on
38 39		 Intra-cranial bleed/skull fracture Admitted to 	GCS 15 patients	Intubation, ICU admissions, administration of mannitol.		admission- due to concern. 3/8 progression on CT	riskier patients. Attrition: Low Risk
40 <u> </u> 41 42		hospital		ED revisits within 1 year for			Potential for patients to have re-attended

		• GCS13-15 on		TBI related symptoms.		1 neurosurgical intervention	at other EDs and be missed
		arrival to ED					
		• GCS 15 24 hours				2/8 admitted to ICU due to deterioration- 1 intubated	Prognostic factor measurement: Low risk
		after attendance to					No risk model developed
		ED				3/95 patients returned with 1 year to the ED due to TBI	Factors abstracted from case notes
_		Did not receive a				symptoms- all underwent repeat CT. No admissions.	
0		repeat CT head				Many location CCC 44.0	Outcome measures: low risk
1		scan				Mean/median GCS=14.8 Mean/median age= 38	Re-attendance at other EDs makes re- attendance a potentially biased outcome
2		Excluded:				Percent anticoagulated=0	measure
3		History of				reicent anticoagulateu-o	lileasure
		neurological or psychiatric					
4		disorder					Confounding Factors: Mod risk
5		Immediate					Cohort includes patients with multiple
6		neurosurgery					injuries
7		Previous TBI or					Statistical techniques: Low risk
8		neurosurgery					None presented
9		Spinal injury					
-		 Coagulopathy 					Is a lower risk population due to selection
0		 Pregnancy 					for repeat CT imaging and return to GCS15
1		 Transfers 					at 24 hours- possibly unable to include in
2		 Incomplete notes 					any meta-analysis.
3							
4		Patients that did					
5		undergo a repeat CT					
		scan despite meeting					
6		the rest of					
7		inclusion/exclusion criteria formed a					
8		comparison group					
9		companison group					
0	Ditty et al	University Alabama	Retrospective Cohort	Neurological decline- altered	Admission GCS	500 patients met inclusion criteria	Study Recruitment: Mod risk
1	2015	Level 1 trauma centre	Study	mental state or focal	Anti-coagulation	411/500 isolated SAH	High proportion of transferred patients may
	Alabama	2003-20013	,	neurological deficit.	Anti-platelets	63/500 isolated ICH	represent higher or lower acuity patients
2	USA		Aim		Transfer Distances	26/500 both	than general population.
3		Inclusion criteria:	Assess the clinical	Inpatient seizure	Sex		
4		• 500 consecutive	implications of SAH or		Age	463 GCS15	Higher as being transferred to specialist
5		patients present on	intraparenchymal	Delayed neurosurgical	Haemorrhage type	30 GCS14	centre, lower as survived /fit to transfer.
6		trauma registry	haemorrhage in mTBI	evacuation as inpatient.		8 GCS13	
7		• GCS13-15		la antique de la contraction de		ACO/FOO activate and benefit and inting continue (74/ACO	No details about inclusion or completeness
•		ICD9 diagnosis SAH		Inpatient mortality.		469/500 patients pre-hospital medication available (71/469 taking either anti-coagulants or anti-platelts)	of trauma registry.
8		and/or intra-				taking either anti-coagulants of anti-platerts)	Attrition: Low Risk
9		parenchymal contusion-				156/500 transfers	Only inpatient measures
0		confirmed with					
1	<u> </u>	1 commined with	l	I	I		- (/± .
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b			Mary A	nn Liebert, Inc, 140 H	luguenot Stree	t, New Rochelle, NY 10801	
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5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21		radiology report and neurosurgical consult note- if disagreement scan re-reviewed if not clear patient excluded: • Diagnosis extra or subdural hematoma • Penetrating injuries • Fatal extra-cranial injuries • CSF leak • Aneurysmal SAH • Delayed presentation	Police			No patients had seizures. No patients had neurological decline. No patients underwent delayed neurosurgical intervention. No inpatient mortality	Prognostic factor measurement: Mod risk Incomplete information regarding medications. May be other inaccurate recording of factors. Outcome measures: Mod risk Only inpatient related outcome measures. Patients may have been discharged and deteriorated and presented to other hospitals. Confounding Factors: Mod risk Cohort includes patients with multiple injuries- only excluded if died from other injuries. Statistical techniques: N A None presented Narrative synthesis- further evidence SAH low risk.
22 23 24 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40 41	Pruitt et al 2016 Chicago USA	Level 1 Trauma Centre Chicago 2009-2013 Inclusion criteria: Initial GCS13-15 16 and older Traumatic intracranial bleed or skull fracture Identified on electronic ED system using ICD 9 classification system Admitted to ED observation unit All patients received a neurosurgical consultation	Retrospective cohort study Aim Assess if mTBI patients with intra-cranial haemorrhage can be managed to an ED observation unit	Clinical deterioration (defined as decrease in mental status, worsening neurologic exam or death) Neurosurgery during admission. Progression on CT.	Age Gender Method of arrival Whether transfer Comorbidities Anticoagulant use Mechanism of injury Initial GCS, Neurological examination Alcohol intoxication Initial platelet count INR Initial CT results Follow-up CT results, Neurosurgical recommendations Cranial CT data were collected from attending radiologist	1185 GCS13-15 with CT detected injuries 814 admitted directly to hospital- poly-trauma, social reasons or as neurosurgeons felt high risk. 371 left under care of ED. Of these, 239/371 transferred ED obs unit. 132/371 discharged directly from the ED after a period of observation. Admitted patients Clinical deterioration 15/814 Worsening CT 27/814 Neurosurgery 33/814 Composite outcome 75/814 ED obs unit Clinical deterioration 10/239 Worsening CT 11/239 Neurosurgery 3/239 Composite outcome 14/239 Medical admission 4/239 Trauma/neurosurgery admit 8/239 Follow up 190/239 Delayed Neurosurgery 20/239 Post traumatic seizure 3/239	Study Recruitment: High risk Neurosurgeons have admitted higher risk patients we can combine outcomes from both admitted and ED observed patients to give an unbiased estimate. Attrition: Med Risk Only a proportion of patients are followed up- does not describe the mechanism for this or how consistent follow up is e.g. did they all get repeat CT scans Prognostic factor measurement: Medium risk Dependent on CT scan reports and written documentation Outcome measures: Mod risk Clinical deterioration not well defined and very broad. Confounding Factors: Low risk
42 43 44 45 46 47			Mary A	nn Liebert, Inc, 140 H	luguenot Stree	et, New Rochelle, NY 10801	

		1			T
			reports- type and	Concussive symptoms 16/239	Included patients with polytauma an
			size of detected	D: 1 15D	significant comorbidities
			injury	Discharged ED	Chatistical to shairman High Diels
				· ·	Statistical techniques: High Risk None presented but data presented in tab
					and text do not match up
				•	and text do not match up
				Concussive symptoms of 132	Paper shows patients admitted to hospit
				Figures from table- author has confirmed this is correct:	by neurosurgeons have wor
					outcomes/more likely to requi
				or cases of neurosurgery.	neurosurgery.
				161 SDH- 6 CT deterioration,	
				3 planned neurosurgical outcomes.	Does show that in America some of the
					patient population discharged directly fro
					ED. Consistent with the model used loca
				•	in Hull.
				•	
				5 extrauarar- nii aeterioration or neurosurgery	
				Of sample 1053 mean/median age=59 11% anticoagulated	
				, , , , , , , , , , , , , , , , , , , ,	
				, , , , , , , , , , , , , , , , , , ,	
Patients admitted	Retrospective cohort	Prospective 1 year telephone	Age	34/1628 mTBI patients isolated traumatic subarachnoid	Study Recruitment: Low risk
tertiary neurosurgical	study	assessment of :	Sex	haemorrhage	Cohort identified in TBi registry which
centre 3 months Jan-		GOSE	Mechanism of		part of normal practice.
March 2010.	Aim	Rivermead post concussion	injury-		Is retrospective so limited by accuracy
					medical notes.
	· ·		-	Rivermead scores comparable to 16 normal CT controls	Associations (11) do Disclar
		follow up questionnaire			Attrition: High Risk Small sample- with large proportion lost
	•				followup.
					Tollowap.
	, parameter 11 and 12 a		bleeds		Prognostic factor measurement: Mediu
			Thickness greater		risk
or Isolated			or less than 5mm		Dependent on CT scan reports and writte
traumatic					documentation
subarachnoid					
 Matched 					Outcome measures: High risk
comparison					1 year too long
•					Confermation Footows Madisum viels
					Confounding Factors: Medium risk No control for other injuries
					comorbidities
					comorbidities
adults only but age					Statistical techniques: N/A
	Patients admitted tertiary neurosurgical centre 3 months Jan-March 2010. Patients identified on a TBI registry Inclusion criteria: GCS 13-15head injury Underwent CT scan Either negative CT or Isolated traumatic subarachnoid Matched comparison between patients - ve CT and SAH Excluded:	Patients admitted tertiary neurosurgical centre 3 months Jan-March 2010. Patients identified on a TBI registry Inclusion criteria: GCS 13-15head injury Underwent CT scan Either negative CT or Isolated traumatic subarachnoid Matched comparison between patients - ve CT and SAH Excluded: Retrospective cohort study Aim To assess whether GCS13-15 patients with traumatic subarachnoid haemorrhage have the same outcomes as mTBI patients with -VE CT scans	tertiary neurosurgical centre 3 months Jan-March 2010. Aim To assess whether GCS13- 15 patients with traumatic subarachnoid lnclusion criteria: GCS 13-15head injury Underwent CT scan Either negative CT or Isolated traumatic subarachnoid Matched comparison between patients - ve CT and SAH Excluded: study Aim To assess whether GCS13- Rivermead post concussion questionnaire Rivermead Head injury follow up questionnaire Rivermead Head injury follow up questionnaire Rivermead Head injury follow up questionnaire	Patients admitted tertiary neurosurgical centre 3 months Jan-March 2010. Aim To assess whether GCS13-15 patients with raumatic subarachnoid haemorrhage have the same outcomes as mTBI patients CT or Isolated traumatic subarachnoid • Matched comparison between patients - ve CT and SAH Excluded: Rivernead post concussion questionnaire Rivernead Head injury follow up questionnaire Rivernead Head injury follow up questionnaire Age Sex Mechanism of Rivernead Head injury follow up questionnaire IoC Seizure Location of SAH Whether multiple bleeds Thickness greater or less than 5mm	Patients admitted tertiary neurosurgical centre 3 months Jan-March 2010. Patients and mitted tertiary neurosurgical centre 3 months Jan-March 2010. Patients identified on a TBI registry Inclusion criteria: 1

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Neurosurgical intervention within 2 weeks repeat Return to the Ed within 7 days of discharge for	323/1011 patients that under-went 2 CT head within 24 hours in ED met the inclusion criteria After second CT 92/323 admitted 25/323 observed in ED and subsequently discharged 206/323 discharged 4 patients died (3 admitted 1 discharged) States death in discharged patient unlikely to be related to head injury had further fall. Also 1 other patient dies of septic shock. 3 neurosurgical interventions (all admitted) 28/206 discharged patients returned to ED within 1 week. None re-admitted and some planned- removal of sutures. Mean/median age= 42 Percent anticoagulated=0	Too poor quality to include Study Recruitment: Mod risk Identified through repeat CT imaging in EDrelies on all of cohort having repeat scans and patients deteriorate and not undergoing second scan being missed Attrition:Low Risk Followed up through social security system for deaths and the rest are inpatient outcome. Possibility of patients reattending at other ED Prognostic factor measurement: Medium risk States that some CT are reported by radiology trainees overnight and then corrected by attending radiologists the next day- unable to quantify how much inaccuracy there is. Does state 32% of repeat scan normal Outcome measures: low risk Reasonable outcome measures Confounding Factors: Low risk Controls for comorbidities and other injuries Statistical techniques: N/A
e r urs th I t	Neurosurgical intervention within 2 weeks e repeat urs after th ICH. If t stable Neurosurgical intervention Age Race Race Sex Medical background	Neurosurgical intervention within 2 weeks Return to the Ed within 7 days of discharge Return to the Ed within 7 days of discharge Return to the Ed within 7 days of discharge Sex Medical background Sex Medical background Medical background Sex Medical background Age Race Sex Medical background Sey After second CT 92/323 admitted 25/323 observed in ED and subsequently discharged 206/323 discharged 4 patients died (3 admitted 1 discharged) States death in discharged patient unlikely to be related to head injury had further fall. Also 1 other patient dies of septic shock. TBI and 3 neurosurgical interventions (all admitted) 28/206 discharged patients returned to ED within 1 week. None re-admitted and some planned- removal of sutures. Mean/median age= 42

Ding et al 2012	Neurosurgical Centre China	Appears to be a random control trial comparing	GCS at discharge Surgical and medical	CT scan results Initial GCS	32/89 patients in routine CT group GCS13-15	Study Recruitment: High risk Allocation to intervention and non-			
Neurosurgical Center China	2009-2010 Inclusion criteria: All patients with TBI with evidence of intra-cranial haemorrhagesome data for GCS13-15 Excluded: Immediate neurosurgery Died within 3 days Severe multiple injuries Failed to undergo a	outcomes in patients with traumatic intra-cranial haemorrhage assigned either to a routine repeat CT or CT only if deteriorates	interventions secondary to CT	Mechanism of Injury Coagulation INR and platelets	2/32 worse CT scans No patients had neurosurgery or altered medical management Mean/median age= 48	intervention arm not clearly explained- states via random number generator Attrition:Low Risk Low risk- inpatient outcomes Prognostic factor measurement: Medium risk No re-reporting of CTS Outcome measures: Medium risk No outcome measures after discharge Confounding Factors: Low risk Controls for other injuries			
Huynh et al 2006 USA	repeat CT head Level 1 trauma centre 2004-2005 Identified case note review Inclusion criteria:	Retrospective cohort study Aim To assess whether neurosurgical review is necessary in GCS 15 patients with intra-cranial injuries	Changes on follow up CT- all patients had routine repeat CT Neurosurgical intervention	Demographics Mechanism of Injury ISS LOC Amnesia Associated injuries	56 patients met inclusion criteria 4/56 patients worse repeat CT Of these 4: 2/56 patients had fall in GCS to 14 from 15 1/56 given mannitol due to worse CT 1/56 loaded with phenytoin for seizures No consistent measure of deterioration 0/56 neurosurgical interventions 0/56 deaths Mean/median GCS=15 Mean/median age= 41	Statistical techniques: N/A Study Recruitment: Medium risk Weaknesses of a retrospective case note review Higher risk group as admitted for at least 48 hours Attrition: Low Risk Low risk- inpatient outcomes Prognostic factor measurement: Medium risk No re-reporting of CTS Outcome measures: Medium risk No outcome measures after discharge Confounding Factors: Low risk No controls for other injuries Statistical techniques: N/A			
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Almenawer et	Neurosurgical centre	Retrospective cohort	Intervention including:	Demographics	1121 patients with mTBI and ICH	Study Recruitment: High risk
al 2013	Ontario, Canada	study + meta-analysis to	Mannitol or hypertonic	GCS		Dependent on accuracy of trauma database
Ontario	2006-2011	assess whether repeat CT	saline	ISS	445 met inclusion criteria	
Canada	Identified from trauma	imaging necessary in mTBI	Surgical intervention			Large proportion of mTBI patients with ICH
	database	with intra-cranial	including ICP bolt or		91/445 worse CT	did not meet inclusion criteria- selection
		haemorrhage	craniotomy			out of higher risk patients that did no
					21/445 patients neurosurgical outcomes (all preceded by	undergo repeat imaging
	Inclusion criteria:		Neurological changes:		clinical deterioration prior to repeat ct)	
	• GCS13-15		decrease GCS, cranial nerve			Attrition:Low Risk
	Blunt traumatic head injury		change, vomiting and headache		4/445 patients medical intervention	Low risk- inpatient outcomes
	Age>17				2/4 medical outcomes= treated with mannitol due solely	Prognostic factor measurement: Mediun
	Intra-cranial injury				worse CT other 2 treated due to clinical deterioration.	risk
	CT head					No re-reporting of CTS
	Repeat CT scan				Mean/median GCS=14.5	
	Excluded:				Mean/median age= 45	Outcome measures: Medium risk
	No repeat CT scan				Percent anticoagulated=0	No outcome measures after discharge
	 Previous 					
	caniotomy					Confounding Factors: Low risk
	 Cranial pathology 					No control for poly trauma
	 Coagulopathy 					S 1. 1
	 Immediate 					Statistical techniques: N/A
	Neurosurgery					
	Patients divided into					
	those underwent					
	intervention due to					
	clinical deterioration or					
	due to repeat CT					
	findings					
Sifri et al 2004	Level Trauma Centre	Retrospective Cohort	Worse CT	CT results as	243 patients with mTBI and ICH	Study Recruitment: Medium risk
USA	New jersey	Study:		abstracted from	18/243 excluded as no repeat CT- neurosurgeon ruled	Selection out of patients not undergoin
	1999-2001	To assess the value of	Inpatient neurological	radiologist and	insignificant lesion	repeat CT hea dimaging
		routine repeat CT imaging	deterioration- abnormal	neurosurgeons		
	Inclusion criteria:	in mTBI patients with	neurology- confusion,	reports.	202/243 included as met the rest of inclusion criteria	Attrition:Low Risk
	• GCS 14-15	intra-cranial haemorrhage	disorientation or drowsiness	D . FD 666		Low risk- inpatient outcomes
	Blunt traumatic		l	Best ED GCS	At 24 hours:	
	head injury		Inpatient neurosurgical	Demographics	151/202	Prognostic factor measurement: Mediu
	• Age>15		interventions		151/202 persistently normal or improving neurology	risk The definition of abnormal neurology
	Intra-cranial injury				51/202 parsistantly abnormal or warraning revertable	The definition of abnormal neurology
	CT head				51/202 persistently abnormal or worsening neurological	loose and not clear when it developed- no
	Repeat CT				examination	an admission criteria factor
	Excluded:				50/202 worse CT	Outcome measures: Medium risk
	History of brain				JU/ ZUZ WUISE CI	Outcome measures: Medium risk
	injury			1		No outcome measures after discharge

	Coagulopathy including known bleeding disorder or taking warfarin Immediate neurosurgical intervention including transfer to ICU	Ŷe₁			5/202 required neurosurgery- all had persistent or worsening neurology 1/202 died all in the persistently abnormal/ worsening neurology group No clear measure of deterioration Mean/median GCS=14.7 Mean/median age= 44 Percent anticoagulated=0	Confounding Factors: Low risk No control for poly-trauma and comorbidites Statistical techniques: N/A
Phelan et al 2014 Dallas USA USA	Level 1 Trauma Centre Dallas Texas 2010-2012 Patients identified on TBI data base Inclusion criteria: Intracranial haemorrhage TBI Patients divided into SAH and non SAH bleed All GCS but data for GCS13-15 patients presented Excluded: Ages less than 18 Pregnant Prisoners	Retrospective Cohort Study Assess whether outcomes for mTBI with isolated traumatic subarachnoid differ for other kinds of intra-cranial bleeds	Worse repeat CT imaging if any Death Craniotomy	CT findings as reread by a study team member Age ISS HAS Emergency department GCS	77 patients GCS13-15 and traumatic SAH 27/77 scheduled repeat CT 3/27 worse CT 50/77-no routine repeat CT 4/50- unscheduled repeat CT 1/50- clinical deterioration and worse CT 4/77 worse CT 0 neurosurgical intervention	Study Recruitment: Low risk Dependent on accuracy of trauma registry Attrition:Low Risk Low risk- inpatient outcomes Prognostic factor measurement: low risk Does not really assess prognostic value of factors measured Outcome measures: Medium risk No outcome measures after discharge Confounding Factors: Low risk No control for poly-trauma and comorbidites Statistical techniques: N/A
Homnick et al 2012 New Jersey USA	New Jersey Medical School Level 1 trauma centre 2002-2005 Inclusion criteria: • Age>17 • GCS>12 • TBI with positive initial CT-intracerebral bleed, contusion, subdural, extra-	Retrospective Cohort Study Establish how long intra- cranial bleeds in mTBI continue to expand	Neurosurgical intervention Progression on CT-repeat CTs as discretion of neurosurgeon	Age Sec Pre-injury anti- coagulation Mechanism ISS Initial GCS	341 patients in study (85 mTBI patients with bleeds excluded as no F/U scan) 72/341 intubated in ED 105/341 progression on CT 13/341 death- 9 due to TBI 4 other causes 12/341 neurosurgical intervention Mean/median GCS=14.6 Mean/median age= 47 Percent anticoagulated=2	Study Recruitment: Medium risk Selection out of lower risk patients that did not have repeat CT imaging Attrition:Low Risk Low risk- inpatient outcomes Prognostic factor measurement: low risk Does not really assess prognostic value of factors measured Outcome measures: Medium risk No outcome measures after discharge
2 3 4 5 5 5 7		Mary A	nn Liebert, Inc, 140 H	luguenot Stree	t, New Rochelle, NY 10801	On

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1 2 3 4							
5 6 7 8 9 10 11 12 13 14 15	Nasir et al	dural or SAH Excluded: Penetrating trauma Injury >24 hours previously Previous neurosurgery Non-traumatic mass on CT Immediate neurosurgery Specialist Centre	Retrospective Cross-	Worse CT	Age	275 patients met inclusion criteria (note states 255	Confounding Factors: Medium risk No control for poly-trauma and comorbidites Statistical techniques: N/A Study Recruitment: Medium risk
16 17 18 19 20 21 22 23 24 25 26 27 28 29 30 31 32 33 34	2011 Karachi Pakistan	Karachi Non-probability consecutive sampling Inclusion criteria: GCS14-15 All ages-15% sample children mean age 36 2 SD 18 TBI with positive initial CT intra- cranial injury Excluded: Clinical deterioration Immediate neurosurgery Isolated pneumocephalus All patients had a repeat CT within 72 hours	sectional study Aim: Assess the utility of repeat CT scanning in mTBI patients with intracranial injuries without clinical or neurological deterioration		Gender Initial GCS Mechanism of injury CT findings	contusion haematoma) 17/275 worse CT No patients required neurosurgery Mean/median GCS=14.7 Mean/median age= 36 Percent anticoagulated=0	Does not adequately define deterioration or over what period Attrition:Low Risk Low risk- inpatient outcomes Prognostic factor measurement: low risk Does not really assess prognostic value of factors measured Outcome measures: Medium risk No outcome measures after discharge Confounding Factors: Medium risk No control for poly-trauma and comorbidites Statistical techniques: N/A Overall Includes kids and quite a different population than North America and Europe.
35 36 37 38 39 40 41	Boris et 2013 Israel	Israel Level 2 trauma centre Sates 2007-2011 Inclusion criteria: GCS14-15 TBI with positive	Retrospective Cohort Study Assess whether repeat CT imaging in GCS14-15 mTBI with intracranial injury justified	Increased size of bleed second CT Clinical deterioration-decrease in GCS New motor or sensory symptoms	Age Sex Initial and follow- up GCS CT findings	68 patients 4 patients transferred to neurosurgery (2 routine) 8/68 patients worse CT 12/68 mild deterioration	Study Recruitment: Medium risk Identified on trauma data base with patients with incomplete data excluded. Does not present number of these patients. Also excludes patients transferred immediately. Likely to be lower risk smaple than population of interest.
42 43 44 45 46 47			Mary A	nn Liebert, Inc, 140 H	luguenot Stree	t, New Rochelle, NY 10801	

Brown et al 2007 Los Angeles USA	initial CT intracranial injury including subdural, extra-dural, subarachnoid and intra-cerebral bleeds Only data for adults presented Excluded: Patients with incomplete data Transferred to neurosurgery immediately No repeat CT All patients had a repeat CT within 12 hours Los Angeles Level 1 trauma center 2003-2004 Inclusion criteria: All patients with blunt head trauma and intra-cranial bleed initial CT. Presents data for GCS13-15 Excluded: Immediate neurosurgery Died within 24 hours Does not state just adults but seems only for adults (mean age 44 +/-19)	Prospective Cohort Study Aim To identify patients with head injuries that benefit from routine repeat CT imaging	Need for neurological intervention- either medical or surgical (medicalsedatives, mannitol or hyperventilation and surgical= ICP monitor and craniotomy) Mortality	Age Gender Mechanism of Injury ISS Admission GCS Results of CT- interpreted by attending radiologist	28 patients intra-parenchymal bleed 1/28 worse CT 3/28 neurological deterioration 1/28 transferred to neurosurgery (not patient with worse CT) 7 patients extra-dural 1/7 worse CT 0/7 neurological change 1/7 transferred to neurosurgery 20 patients sub-durals 3/20 worse CT 4/20 neurological deterioration 1/20 neurosurgery 13 patietns SAH 3/13 increase in size bleed 5/13 neurological deterioration 1/13 transferred to neurosurgery Mean/median GCS=14.8 Mean/median age= 56 354 patients all GCS scores with intra-cranial bleed 37 direct to craniotomy 43 dies within 24 hours 274= study population 142/274= mTBI GCS13-15 15/142 had clinical deterioration 27/142 had worse CT scans (only 72/142 had repeat imaging) 5/142 had medical or neurosurgical intervention 3/142 died Mean/median GCS=14 Mean/median age= 43	Attrition:Low Risk Low risk- inpatient outcomes Prognostic factor measurement: low risk Does not really assess prognostic value of factors measured Outcome measures: Medium risk No outcome measures after discharge Confounding Factors: Medium risk No control for poly-trauma and comorbidites Statistical techniques: N/A Study Recruitment: Mod risk Removal of patients that died within 24 hours may lead to this sample being a lower risk group than population of interest Attrition: Low Risk Low risk- inpatient outcomes Prognostic factor measurement: low risk Does not really assess prognostic value of factors measured Outcome measures: Medium risk No outcome measures after discharge Confounding Factors: Medium risk No control for poly-trauma and comorbidities- Statistical techniques: N/A
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	Thomas et al 2010	Tennesse Level 1 trauma centre	Retrospective Cohort Study	Neurosurgical interventions- craniotomy or ICP monitor	Initial GCS ISS	457/836 in included sample population GC\$13-15	Study Recruitment: Mod risk Dependent on case note review. Patient
-	Tennesse	50 months from Jan		,	Race	14/457= neurosurgical intervention (craniotomy or ICP	with "unclear" indications for interventions
- 1	JSA	2001	To assess whether	Medical interventions-	Age	bolt)	removed.
			scheduled repeat CT head	mannitol/hypertonic saline	Gender	3/457 medical management	
		Inclusion criteria:	imaging is indicated in TBI		Mechanism of		
		All patients with		Neurological change-reduced	injury	5/14 neurosurgical interventions- based on repeat CT	Attrition: Low Risk
		blunt head trauma and evidence TBI		GCS, pupillary change, increased ICP or loss of brain	History of vascular disease	3/14 medical interventions based on repeat CT	Only inpatient outcome measures
		on initial CT.		stem reflexes	Anticoagulant use	Mean/median age= 42	Prognostic factor measurement: Mod risk
		Presents data for		Stem renexes	Antiplatelet use	Wedny median age 42	Does not explain how CT scans reported
		GCS13-15			PT, aPPT, INR		
		 Age 18+ 			CT findings		Outcome measures: Mod risk
		Excluded:					No F/U after discharge
		 Penetrating 					
		mechanism					Confounding Factors: Medium risk
		 Immediate 					No control for poly-trauma
		neurosurgery					Statistical techniques: N/A
		 Interventions for unclear indications 			7 //.		None done
		Died before second					None done
		CT					
		Ci					
		All patients repeat CT at					
		6-8 hours after					
		admission					
- 1	Klein et al 2010	3 regional trauma	Retrospective Cohort	Mortality	Age	323 patients all 3 hospital intra-cranial bleed and GCS13-15	Study Recruitment: Low risk
	srael	centres in Israel. None	Study	Neurosurgical intervention	AIS		Dependent on completeness of trauma
		had access to		Neurological status at	ISS	27/323 required neuro-rehab	registry
		neurosurgery on site.	Aim:	discharge		2/323 died	Attrition: Low Risk
		Identified ICD9 codes on	Assess the outcome of low risk patients with ICB			35/323 neurosurgery	Only inpatient outcome measures
		national trauma registry.	managed in district			77/323 not transferred-	Only inpatient outcome measures
		Inclusion criteria:	hospitals without			0/77 died	Prognostic factor measurement: Mod risk
		• GCS13-15	neurosurgical services			0/77 neurosurgery	Does not explain how CT scans reported
		• ICD9 code for	-			2/77 delayed transfer	
		intra-cranial bleed.					Outcome measures: Mod risk
		One hospital transferred				Non-transfer on basis of:	No F/U after discharge
		all patients to				Single bleed = 5mm or contusion <1cm and no-</td <td></td>	
		neurosurgical centre.				coagulopathy	Confounding Factors: Medium risk
		Other 2 hospitals				Mean/median age= 39	No control for poly-trauma or
		transferred selected patients.				wiean/median age= 39	comorbidities
		le s sections					Statistical techniques: N/A

Sifri et al 2011 Level 1 Trauma Centre Retrospective Cohort Progression of lesion on CT De USA New jersey Study Surgical intervention- Ac		1
· · · · · · · · · · · · · · · · · · ·		
USA New Jersey Study 1 Surgical Intervention- Ac	emographics 107 patients met inclusion criteria	Study Recruitment: High risk
, ,	· ·	High risk subgroup that have abnorm neurology at time of repeat CT imaging.
	eterioration in 7/107 neurosurgical group eurological Exam 21/107 deterioration	hedrology at time of repeat C1 imaging.
	ersistently 18/107 unable to assess neurology as intubated.	Attrition: Low Risk
	onormal 6 died	Only inpatient outcome measures
	eurological exam	Only inpatient outcome measures
	nknown whether Mean/median GCS=14.4	Prognostic factor measurement: Mod risl
, , , , , , , , , , , , , , , , , , ,	nange as Mean/median age= 48	Difficult to assess deterioration in
1.04 = 4	tubated Percent anticoagulated=0	retrospective study.
CT head-ICB or they have a repeat CT.	Tercent uniteouguided o	retrospective study.
skull fracture		Outcome measures: Mod risk
Repeat CT		No F/U after discharge
Abnormal		The type direct discharge
		Confounding Factors: Low risk
neurological examination at		Some control for comorbidities.
examination at		
time of repeat CT Excluded:		Statistical techniques: N/A
		None done
• Immediate or planned		
· · · · · · · · · · · · · · · · · · ·		
neurosurgical intervention		
at time of repeat CT- normal		
neurology defined as GCS15,		
as GCS15, orientation to		
place, person or time, normal		
neurological exam,		
no symptoms from head injury-		
headache,		
vomiting, dizziness,		
lethargy • Coagulopathy		
, , , , ,		
including known		
bleeding disorder		
or taking warfarin		
Pregnancy Sping Conduction		
Spinal Cord Injury		
Mary Ann Liebert Inc. 140 Hus	juenot Street, New Rochelle, NY 10801	
waiy Aini Liebeit, iic, 140 Hug	jacinot otroct, new reconcile, NT 10001	

			1	T	_	_
	 Prior brain surgery 	'				
	 Acquired or 	'				
	congenital cerebral		[
	pathology or		[
	existing					
	neurological or					
	psychiatric					
	disorder					
Beynon et al	Heidelberg University	Retrospective Cohort	Repeat CT imaging	Patients divided	70 patients met inclusion criteria	Study Recruitment: Low risk
2015	Hospital Germany	Study	Progression on CT	into those on no	37 no anticoagulation	Although high rates of anti-coagulation.
Germany	2013-2014		Neurosurgery	anticoagulants,	27 anti-platelets	
		Aim:	Death	Aspirin, Warfarin	5 warfarin	Attrition: Low Risk
	Inclusion criteria:	Compare outcomes in	Mean GCS at discharge	and DOACS.	6 DOACS (rivaroxaban)	Only inpatient outcome measures
	 Initial GCS 13-15 	patients on different			1 patient dabigatran	
	Traumatic Intra-	types of anti-coagulants		gender,		Prognostic factor measurement: Low risk
	cranial bleed CT	7,6		trauma	25% neurosurgery (18 patients)	May be miss-classified in medical notes
	head			mechanism,	43/70 repeat CT imaging-	
	nedd			comorbidities,	15/701021111111111111111111111111111111111	Outcome measures: Mod risk
				CT findings,	2 deaths both on rivaroxaban	No F/U after discharge
				repeated CT	E deditio both on management	No 1/6 dite. dissilarge
				imaging,	Mean/median GCS=14.5	Confounding Factors: Low risk
				age,	Mean/median age= 67	No control for comorbidities.
		'		GCS scores,	Percent anticoagulated=16	110 001.0101 001.0101.0101.01
						Statistical techniques: N/A
		'		lustrator, interest		None done
						None as
					et, New Rochelle, NY 10801	
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Supplementary Material 2: Data Extracted from Included Studies

Studies with univariate or multivariate risk factors N=21 (also included in pooled estimates outcome prevalence)

Reference	e Population Study Design Outcome Prognostic factor				Results	Quality Appraisal
Reference	Population	Study Design	Measures	assessed	nesuits	Quality Appliaisai
Nieleiiinee	Single-site: Level 1	Prospective			600 patients	Study Recruitment: Mod risk bias
Nishijima et al 2014	trauma centre	cohort study	critical care invention within	Age <u>></u> 65years	71% male	Missed 20% eligible patients- not
		conort study		Sex		5 1
	2009 – 2013	A !	48 hours of arrival	D	0.5% died + 6.5% neurosurgery + 8.3% intubated	completely clear individuals in cohort identified. Otherwise clear
o USA		Aim:	ED:	Dangerous	68% GCS 15	
	Inclusion Criteria:	Derive a clinical	Intubation	mechanism (any	030/	inclusion and exclusion criteria.
	 Age ≥ 18 years 	decision	 Neurosurger 	non-fall from	93% admitted ICU	
	 Consecutive 	instrument for	y including	standing	19.3% had crit care intervention	Attrition: Low risk
	patients	patients with mild	ICP	mechanism)	9.2% transfusion	Follow up only 48 hours so low risk
	 Initial ED GCS 	ICH low risk	monitoring/		8.3% intubation	of attrition bias.
	13-15	requiring critical	giving	Pre-injury	6.5% Neurosurgical	D
	• CT +ve ICH-	care intervention.	mannitol/hy	antiplatelet use	16 11 11 11 11 11 11 11	Prognostic factor measurement:
	SAH, SDH,		pertonic	(aspirin or	4 predictors need for crit care intervention: (Recursive partitioning)	Low risk
	EDH, intra-	Statistical	saline	clopidogrel)	GCS<15 (RR 2.95; 95% CI 2.21-4.12)	Standardised and objective
	ventricular,	Method:	 Transfusion 		≥ 65years (RR 1.46; 95% Cl 1.05-2.03)	prognostic factor measurement.
	intra-	Derived clinical	RBC/FFP	High risk co-	CT midline shift/absence cisterns (RR 4.11; 95% CI 3.08-5.48)	Collected all patients.
	parachymal	decision	 Vasopressor 	morbidity	Non-isolated head injury (RR 2.74; 95% CI 1.99-3.78)	
	bleed/contusi	instrument with	/ionotrope	50 VC: 1 :		Outcome measures: Low risk
	on, diffuse	binary recursive	use	ED Vital signs	Sensitivity of decision rule to predict intubation/neurosurgery within 48 hours of	Recorded in uniform way for all
	axonal injury	partitioning	 Cardiac 	GCS <15 at	admission ED.	patients. Only 48 hours.
		(misclassification	arrest/arrhy	admission	98.6% specificity 36.6%	
	Exclusions:	cost 20:1).	thmia	BP<90 at any point	To any crit care inteverntion	Confounding Factors: Mod Risk
	 Patients with 		(HR<40,	ED	Sensitivity 98.3% 95% C.I. (93.9-99.5%)	Additional severe injury may be
	DNACPR	Performance of	HR>120)	Sats <95% at any	Specificity 39.7% 95% C.I. (35.4-44.1%)	related to prognostic factors and
	 Patients pre- 	instrument	 Intervention 	point ED	Positive predictive value 28.1% 95% C.I. (23.9-32.6%)	outcome measures. Not accounted
	injury anti-	compared to	al		Negative predictive value 99% 95% C.I. (96.3-99.7%)	for in in analysis.
	coagulant use	clinical	angiography	Lab results:		
		impression.		Platelet count	Clinician impression:	Statistical techniques: low risk
				INR	Do you think patient needs ICU?	Good presentation of methods
				Haematocrit	Sensitivity 90.1% 95% C.I. (83.1-94.4%)	
					Specificity 49.2% 95% C.I. (44.7-53.8%)	Overall summary
				Initial CT:		Risk factors identified by case note
				Midline	Clinical impression deterioration in 48 hours?	review/d/w treating physicans
				shift/absence	Sensitivity 91% 95% C.I. (84.2-95.0%)	where not clear. Radiology
				cisterns	Specificity 39.5% 95% C.I. (35.1-44.1%)	attending written report used for
				Depressed skull		CT findings. No independent
				fracture	Presence of swelling or shift on initial cranial CT RR (95% CI) 4.11 (3.08-5.48)	quality verification- could
					Admission GCS score less than 15 RR (95% CI) 2.95 (2.12-4.12)	introduce bias. CT end point also
				Non-isolated head	Non-isolated head injury RR (95% CI) 2.74 (1.99-3.78)	missed spectrum of possible

				injury AIS score 3 or	Hypotension prior to admission RR (95% CI) 2.70 (1.61-4.54)	findings.
				more additional	Presence of depressed skull fracture RR (95% CI) 2.44 (1.46-4.08)	
				injury	Presence of any high-risk co-morbidity	Outcomes out 48 hours too short,
					1.58 (1.07-2.33) RR (95% CI) Pre-injury antiplatelet use	also crit care intervention
					1.54 (1.04-2.30) RR (95% CI) Hypoxia prior to admission	definition very broad- e.g.
					1.52 (1.03-2.24)	transfusion. No blinding to
					Age 65 years or older RR (95% CI) 1.46 (1.05-2.03)	exposure/outcomes.
					Non-fall from standing mechanism of injury RR (95% CI) 1.12 (0.80-1.57)	
					Mean/median GCS=14.6	Overall good internal validity of
					Mean/median age= 52	study.
					Percent anticoagulated=0	But issues with generalising
						results:
						Exclusion of anti-coagulated
						patients.
						Short outcome measurement 48 hours.
						Outcome measures of critical care
						intervention quite soft- including
						transfusion of blood products.
						No external validation of results.
Sweeney et	Identified on	Retrospective	Neurosurgical	ISS (measure of	50496 patients met criteria	Study Recruitment: High risk bias
al 2015	national trauma	Cohort study	Intervention:	head injury severity	4474/50496 neurosurg	Eligible patients recruited through
USA	data base 2007-	, ,	Defined as	due to exclusion	58% admitted to ICU	a relatively new national trauma
	2012	Hypothesis that	operative	criteria).		data base by ICD9 coding. Potential
	Inclusion criteria:	injury type	procedure, or		EDH-N=901 18% Neurosurg	selection bias as to which hospitals
	 Age ≥ 18 years 	associated with	placement of an	Coagulopathy	SDH-N=18784 16% Neurosurg	upload data. Also uncertain how
	ED inital GCS	deterioration in	ICP monitor.	(pooled measure of	Mixed N=11984 8% Neurosurg	accurate coding is.
	14-15	isolated TBI.	Identified by ICD9	Vit K deficiency,	SAH N=13191 1.5% Neurosurg	
	• ICD 9 code		coding.	haemophilia,	Contusion N=5636	Excluded patients with incomplete
	intra-cranial	Multiple logistic		thrombocytopaenia,		data, they may be systemically
	injury=	regression used		chronic anti-		different.
	cerebral	to assess risk of		coagulant therapy)	Data set split into 2/3 training set and 1/3 test set.	
	contusion,	outcomes.		Chronic aspirin use	Adjusted adds vetics for neurosurgical precedures Multiple logistic assessing and a 2/2	Assuisia and a constitute
	SAH, SDH,	Mixed effects		not included.	Adjusted odds ratios for neurosurgical procedures. Multiple logistic regression run on 2/3	Attrition: Low risk
	EDH, multiple	Mixed effects model to explore		Type of intra-cranial	training set (n = 33,327)	As a trauma registry represents routine information that should be
	TBI	potential		injury as per ICD 9	Age (years) OR=1.002 (95% CI0.999 – 1.01) P=0.18	consistently on all eligible patients.
	Admitted to	differences		code.	Age (years) OK-1.002 (95% Cl0.999 – 1.01) P-0.18 Anticoagulation Disorder OR=0.853	consistently on all eligible patients.
	hospital Exclusions:	between		couc.	(95% CI 0.66 – 1.09) P=0.21	Prognostic factor measurement:
	ICD9	hospitals.		ED vital signs	ED GCS OR=0.894 (95% CI 0.781 – 1.03) P=0.11	Mod risk
	diagnoses skull				ED Systolic Blood Pressure OR=1.004 (95% CI 1.002 – 1.01) P<0.001	Grouping of coagulopathy
	fractures			Age	ED Pulse OR=0.99 (95% CI0.986 – 0.993) P<0.0001	problematic, different likely risk of
l	Penetrating				ED Respiratory Rate OR=0.962	warfarin versus ITP for example. CT
	mechanism of				(95% CIO.944 – 0.98)	findings watered down to code for
1	incentation of				P<0.0001	injury, misses important

	injur				ISS 7 11 OP-2 25 (OF9) CL1 44 4 00) P<0 01	information
	injury				ISS 7-11 OR=2.35 (95% CI 1.44 – 4.09) P<0.01 ISS 12-18 OR=3.37 (95% CI 2.06 – 5.86) P<0.0001	information.
	AIS score>1				ISS 19-27 OR=18.9 (95% CI 11.6 – 33) P<0.0001	Outcome measures: Moderate
	any other				,	risk
	body region				ISS > 27 OR=7.01 (95% CI 3.79 – 13.4) P<0.0001	
	Data missing				Injury Category (vs. Contusion)	Need for neurosurgery only as recorded on trauma data bank.
	ED vital signs				Isolated SAH OR=0.95 (95% CI 0.64 – 1.41) p=0.79	1
					Isolated SDH OR=4.9 (95% CI 3.61 – 6.84) P<0.0001	possibly unreliable. Misses other
					Isolated EDH OR=6.42	important adverse outcome e.g.
					(95% CI 4.15 – 9.97) P<0.0001	death and intubation. Does not
					Multiple Injury Types OR=2.34	include time scale from
					(95% CI 1.7 – 3.29) P<0.0001	presentation or what happens to patients who are discharged and
					After adjustment injury severity, age, coagulopathy and ED vital signs: injury pattern	re-attend with adverse outcome.
					significantly associated need for neurosurgery:	Follow up not clear
					OR EDH versus contusion 6.4(95% CI 4.1-9.9).	•
						Confounding Factors: Low risk
					Age no association.	Excluded other injuries and made
						adjustments in logistic regression
					ED vital signs also predictive.	model. No attempt to control for
						co-morbidities.
					In test AUC ROC curve= 0.81 in test set	
					Hosmer-Lemeshow P = 0.8 in test set	Statistical techniques: low risk
						Good presentation of methods
					38% expected and observed rate of neurosurgery highest risk decile. 0.5 % in lowest risk	
					decile.	Finds that injury type significantly
						associated with need for
					Mean/median age= 61	neurosurgery -provides candidate
					Percent anticoagulated=5	factors. There are methodologica
						problems with paper.
Joseph et al	Level 1 trauma	Retrospective	Progression on	Age	876 patients met inclusion criteria	Study Recruitment: Mod risk
2015	center	Chart Review	repeat CT	Gender		Retrospective identification of case
	Arizona			Race	115 (13.1%)=progression on CT	notes- depends on accuracy o
		Aim	Neurosurgical	Ethnicity		case notes
USA	Retrospective case	Identify factors	intervention=	Mechanism of injury	Univariate predictors:	
	note review 2009-	that predict	craniotomy or	GCS		Excludes patients on anti-
Is MTBI	2012	progression on CT	craniectomy as	BP	Age 65+ p=0.07 OR1.5(0.9-2.5)	coagulatants and anti-platelts
defined by		imaging and	inpatient	HR	Male p=0.8 OR1.1 (0.6-1.7)	
GCS: is it	Inclusion criteria:	neurosurgical		FBC	Intoxication p=0.9 OR1.3 (0.3-4.7)	Attrition: low risk
really mild?	Initial GCS13-	intervention in		Serum lactate	Mechanism of injury p=0.5 OR 1.1 (0.3-2.8)	Outcomes only as inpatients
	15	GCS13-15 patients		Base deficit	HR>100 P=0.7 OR1.1 (0.6-1.8)	
	 Aged 18+ 			AIS	BP<90 p=0.35 OR 1.3 (0.45-1.9)	Prognostic factor measurement
	 Initial scan +VE 			ISS	LOC p=0.2 OR1.2 (0.6-2)	Low risk
	ICH/skull	Method			Displaced skull fractue P=0.02 OR 1.9 (1.1-3.3)	Relies on accuracy of medical
	fracture and	All patients		CT findings-	SDH >10mm p=0.004 OR3.4 (1.5-8)	notes.
	routine repeat	underwent		reviewed by an	EDH >10mm p=0.01 OR3.8 (1.2-7.6)	
	1	routine repeat CT		investigator that	Hgb<10 P=0.4 OR 1.5 (0.76-3.1)	Re-examines CT images

K					Jou	rnal of Neurotrauma		Page 106 of
1 2 3 4 5	0,-	scan still	imaging within 6		was part of the	Platelets less than 100000 p=0.04 OR 1.5 (1.1-3.9)		
6		showed injury	hours of initial CT		team- classified size	Lactate =/<2.5 p=0.18 OR2.6 (1.2-5.5) (?!)	О	utcome measures: Mod risk
7 8 9		Isolated TBI as defined head AIS greater/equal 3 and AIS <3	imaging. Univariate analysis to identify risk		of lesion and whether progression on CT	Base deficit>4 p=0.02 OR 3.1 (1.2-7.6) Multi-variate Analysis: Age 65+ P=1.4 OR 1.4(0.7-2.7)	Po	nly measures as inpatient. otential for discharge and eterioration.
10 11 12 13 14		other body regions Excluded: On Antiplatelets	factors for progression on CT or neurosurgery. P=/<0.2 included			LOC P=0.8 OR1.1 (0.5-2) Displaced skull fracture P=0.08 OR 2.3 (0.9-3.5) SDH>10mm P=0.007 OR 4.8 (1.9-9.6) EDH>10mm P=0.001 P=7.9 (2.4-12.6) Platelets less than 100000 p=0.1 OR 1.3 (0.9-3.6)	Po ot	onfounding Factors: low risk ossibility of confounding due to ther comorbidities- does not djust for this,
15 16 17 18		 On Anti- coagulants Transfers Needed immediate 	multivariate analysis	10	2	Lactate =/<2.5 p=0.2 OR 2.1 (0.89-2.5) Base deficit>4 p=0.01 OR 2.8 (1.6-4.1) 47 (5.4%)= neurosurgery	Sc	catistical techniques: Mod risk ome of the results appear to be aported wrong. E.g. Lactate
19 20 21		neurosurgery.				Univariate predictors: Age 65+ p=0.3 OR 1.08 (0.8-1.3) Male P=0.19 OR 1.2 (0.8-1.3)	Pr	verall resents useable data for analysis
22 23 24						Intoxication P=0.3 OR1.8 (0.9-3.4) BP<90 p=0.35 OR 1.3 (0.45-1.9) Mechanism P=0.34 OR1.2 (0.4-1.8) LOC p=0.19 OR1.4 (0.7-3.2) HR>100 P=0.26 OR 1.5 (0.9-2.8)	hi	ote base deficit found to be ghly prognostic- only study to ssess this.
25 26 27 28						Displaced skull fractue P=0.01 OR 16 (7.6-19.6) SDH >10mm p=0.001 OR3.9 (2.4-5.1) EDH >10mm p=0.03 OR4.8 (2.9-5.6) Hgb<10 p=0.51 OR 1.2 (0.6-2.5)		
29 30 31						Platelets less than 100000 p=0.31 OR 2.5 (1.15-5.1) Lactate =/<2.5 p=0.12 OR3.6 (0.7-6.5) Base deficited p=0.01 OR 23 (1.6-31)		
32						Multi-variate Analysis:		
33 34 35						Male p=0.1 OR 1.6 (0.8-2.1) LOC P=0.3 OR1.2 (0.5-1.9) Displaced skull fracture P<0.001 OR 10 (6.7-12)		
36 37						Multi-variate Analysis: Male p=0.1 OR 1.6 (0.8-2.1) LOC P=0.3 OR1.2 (0.5-1.9) Displaced skull fracture P<0.001 OR 10 (6.7-12) SDH>10mm P<0.001 OR 3.4 2.1-4.46) EDH>10mm P=0.006 P=3.5 (1.4-5.5) Platelets less than 100000 p=0.09 OR 1.3 (0.98-4.8)	2/4	*
38 39 40						Lactate =/<2.5 p=0.21 OR1.9 (0.62-3.1) Base deficit>4 p=0.001 OR 21 (1.6-27)		6.
41						Mean/median GCS=14.3		

						Mean/median age= 54	
						Percent anticoagulated=0	
	Borczuk et	Level 1 trauma	Described as a	Deterioration	Data extracted from	404/863 TBI patients met inclusion criteria (46.8% patients with traumatic bleeds).	Study Recruitment: low risk
	al 2013	centre Boston	cross sectional	whilst in hospital	case notes by 2 ED		Dependent on how good electronic
	USA		study	including:	researchers. Not	11.8%(48) deteriorated	coding is and case note review
		Case note review		Decrease in GCS	blinded to the	5.9% neurosurgical	was.
		2009-2010 patients	Seems more like a	Worsening	hypothesis	Deterioration stratified by injury:	
		identified through	retrospective	neurological		24/136 isolated SDH	Attrition: Low risk
		ED electronic coding	cohort study	examination	Age	0/1 isolated EDH	Follow up only for period in
		ICD9 coding for	Aims	Worsening CT	Method of arrival	1/75 isolated SAH	hospital
		intra-cranial	Develop a set of	result on repeat	History of HTN	2/31 contusions	Prognostic factor measurement:
		haemorrhage.	criteria to identify	CT	Anti-coagulation	22/161 mixed lesions	Low risk
			patients who are	Neurosurgery	Mechanism		Written CT reports from attending
		Inclusion criteria	at low risk for	Death	Initial GCS	Univariate predictors of deterioration:	radiologist used for data
		• GCS 13-15	deterioration and		Neurological		extraction. No verification of
		• Age 15 or	thus may not	Composite	examination	Age 65+ OR 0.93 95%CI 0.5-1.69	accuracy or consistency.
		older	require	outcome	Alcohol Intoxication	Sex OR 0.77 95%CI 0.41-1.41	
		 CT positive 	neurosurgical	All outcomes	Initial platelet count	Fall OR 0.57 95%CI 0.29-1.09	Outcome measures: Mod risk
		traumatic	evaluation	whilst in hospital-	INR	Assault OR 1.07 95% CI 0.45-2.51	No F/U following discharge- missed
		intra-cranial		no discharge	Initial CT result	RTC OR 0.51 95%CI 0.12-2.21	delayed outcomes, could have
		haemorrhage	Method	outcomes	F/U CT result	Pedestrian Struck OR1.12 95% CI0.32-3.92	looked for re-attendance.
		Excluded:	Univariate			Bicycle Struck OR 1.51 95%CI 0.42-5.44	GCS and neurological examination
		 Isolated Skull 	analysis to predict		CT categorised by	HTN OR0.94 95%C.I. 0.51-1.73	also potentially subjective.
		fractures	composite		attending	Aspirin OR 0.79 95% CI0.41-1.51	
			outcome of		radiologist type,	Warfarin OR0.87 95% CI 0.33-2.32	Confounding Factors: Mod risk
			deterioration		location and size of	Clopidogrel OR1.25 95% CI 0.27-5.75	No attempt to control or exclude
					bleed/contusion.	005 45 00 2 42 05% 014 04 42	polytrauma patients or patients
			3 factor		Presence of midline	GCS<15 OR 2.12 95% CI 1.01-4.43	with multiple comorbidities
			multivariate		shift		
			model derived			CT findings	Statistical techniques: Mod risk
			from univariate			Any lesions	Good univariate analysis
			analysis			SDH OR 2.64 95% CI 1.20-5.83	Small number prevented large
						EDH OR 2.4 95% CI 0.91-6.31	enough multi-variate model
						SAH OR 0.42 95% CI 0.22-0.81 Contusion OR 0.79 95% 0.39-1.62	
						Contusion Ok 0.79 95% 0.39-1.62	
						Isolated lesions	
						SDH OR 1.62 95% CI 0.88-2.96	
						EDH OR only 1 patient	
						SAH OR 0.078 95% CI 0.01-0.59	
						Contusion OR 0.46 95% 0.11-1.96	
						Containon on 0.40 55/0 0.11-1.50	
						Multiple logistic regression with 3 variables GCS=15, presence SDH and presence isolated	
						SAH:	
						All remained significant predictors of deterioration. Sensitivity 97.9% and specificity 20.8%	
L			ı			0	C/X

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1 2 3 4	0,,						
5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36 37 37 38 38 38 38 38 38 38 38 38 38 38 38 38	Washingto n et al 2012 USA	Level I trauma center Washington Retrospective case note 2-year period (January 2007-December 2008) Inclusion criteria:	Retrospective Cohort Study Aim To determine if there exists a sub- population of mild TBI patients with an abnormal head CT scan that requires neither repeat brain imaging nor admission to an ICU Standard of care is to admit these patients to ICU and routinely re- CT Methods: Univariate and multivariate analysis for outcomes of interest	Neurological or medical decline. The need for neurosurgical intervention. The GOS score. Neurological decline was defined remaining in the ICU or transfer back to an ICU or intervention as a result of a decline in mental status or the development of a neurological deficit. Medical decline was defined as an increase in monitoring or intervention due to cardiac, pulmonary, or renal decline. Outcome measures during admission and at discharge.	Age Sex, Injury mechanism Initial GCS score Duration of hospital stay. Aspirin/Clopidogrel/ Warfarin use Ttransfusion of blood products Intubation CT scans classified into Marshall and Rotterdam Criteria- blinded assessment by author	Negative predictive value 99.6% Positive predictive value 38.8% Mean/median GCS=14.8 Mean/median age=60 Percent anticoagulated=10 321 patients met the inclusion criteria Neurological decline 1% 4 Surgical intervention 1% Medical decline 6% 18 Cardiac event 7% Respiratory event 4% Seizure event 2% CT progression™6% GOS score at discharge: 1 1% 2 0% 3 4% 4 10% 5 85% Age + transfusion predictors of a medical decline (p < 0.01). Odds ratio of having a medical decline after undergoing a blood product transfusion was 12.55 (95% CI 4.3–36.7). Cardiac and respiratory events the odds ratios were 5.6 (95% CI 2.4–13.1) and 8.8 (95% CI 2.6–30.4). Significantly higher mortality transfused group as compared with the non-transfused group (6% vs 0%, respectively, p < 0.0001, Fisher exact test). Higher rate of brain injury progression in the transfused patients (13% vs 5%, p = 0.04). Predictors of bleed progression univariate analysis: ICH vol >10 ml OR 20.13 95% CI (5.67–71.44) subfrontal/temporal contusion™ OR 5.73 95% C.I.(2.20–14.89) age 265 yrs™ORA.00 C.I>(1.40–11.42) antiplatelet &/or Coumadin therapy OR 2.94 C.I. (1.12–7.71) Unclear which other factors assessed.	Study Recruitment: low risk Through case note review- potential for patients without notes to be missed Attrition: low risk Follow up only for period in hospital Prognostic factor measurement: Low risk Case note extraction- potentially incomplete CT scans re-reported. Uses Marshall classification Outcome measures: Mod risk Outcome measures: only during hospital admission. No measure of re-attendance or community outcome F/U The outcome measures of neurological and medical decline are subjective. Confounding Factors: Medium risk No control for other injuries and comorbidities Statistical techniques: High risk Selective reporting of significant risk factors and does not present full analysis. No analysis to predict neurosurgical outcomes.
39 40 41						States: "multivariate analysis was performed, only an ICH volume > 10 ml was	Potentially can re-analyse the data from what is presented
42 43 44 45 46 47				Mary Ann	Liebert, Inc, 140	Huguenot Street, New Rochelle, NY 10801	

Country Every Line Line Cohert study Using trauma data base Cohert study Using trauma data base Cohert patients in Identified trauma data base Cohert patients Cohert patients	rt study g trauma data Delayed neurological deterioration defined as: Ororal course outcomes of ents who riorate hours Ologically measures: Delayed neurological deterioration defined as: GCS drop 2 or more points for more than 1 hours New focal	Age, Sex, Ethnicity, Mechanism of injury, GCS, AIS,	151 not included due to incomplete notes or meeting exclusion criteria 757= final cohort 31/757= delayed deterioration at inpatient. 4.1% (21 due to progression ICH, 10 due to medical causes)	Retrospective identification of patients on trauma database. Relies on patients being correctly recorded on this. Patients with incomplete notes excluded- may
results Statistical techniques: right risk	MHI and ICH deficit nods ents briate and ivariate risk wath Worse CT if performed- worsening in Marshall criteria or significant expansion in volume- neuroradiologist GOS outcome at 6		21/757 patients worse CT scans Univariate analysis outcome death Age>/=60 P=0.001 Coagulopathy P=0.02 Increase Marshall classification repeat CT P=0.001 Decline in consecutive GCS scores more than 6 P=0.02 Deterioration within 9 hours P=0.04 H-AIS>3 P=0.32 ISS>20 P=0.38 Initial GCS<15 P=0.40 Initial Marshall classification >II P=0.41 Age>60 predicted deterioration due to expansion of bleed and death in stepwise logistic regression (p<0.01)	Reports no loss to F/U at 6 months routine clinic- may form part of group of patients excluded due to incomplete notes Prognostic factor measurement: Low risk Relies on accuracy of medical notes Outcome measures: Mod risk Outcome measure of delayed deterioration- relies on adequate checks on patients and neurological examinations in a consistent way. Assumes this is baseline level of care- likely to vary dependent on where the patients were admitted (e.g. ICU versus normal hospital bed) Confounding Factors: low risk Doesn't explicitly say for patients with only a head injury, if does include other injuries high risk for confounding.
results		months	months	Mary Ann Liebert, Inc, 140 Huguenot Street, New Rochelle, NY 10801

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1 2 3 4							
5 6 7 8 9 10 11 12 13 14 15 16			* Ac	V/e			presented only as P values. Performed multivariate stepwise regression- for mortality reports only one result without confidence intervals. Overall Compares patients with medical and neurosurgical deterioration and that died and didn't die with worsening CT scans. Much more pertinent to compare patients that deteriorated and didn't
17 18 19 20 21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40 41	Kim et al 2014 South Korea	University hospital Seoul South Korea Case note review from Jan 2002-Dec 2012 Inclusion criteria: All patients with acute traumatic subdural bleeds Excluded: Neurosurgery within 24 hours of admission GCS<13 on admission Patients with vascular abnormalities Subdural localised to the falx/tentorium cerebelli Bilateral subdurals Aged less than	chart review	Delayed surgical evacuation of subdural haematoma	Age Gender Cause of trauma Presence of other CT findings GCS Neurological deficit Comorbidities History of antiplatelets Anticoagulation therapy INR Platelet count	98 patients included 51/98 progression on CT either at 1 week , 2 weeks or 3-10 weeks. 34/98 delayed surgical evacuation up to 10 weeks following trauma Univariate comparison between conservative and delayed neurosurgical group: Mean age P=0.375 Male, P=0.950 Glasgow Coma Scale P= 0.647 Hypertension P= 0.883 Diabetes P= 0.785 Smoking P=0.107 Alcohol abuse P=0.840 Use of anticoagulant P= 1.000 Use of anticoagulant P= 1.000 Use of antiplatelet agent P= 0.546 Thrombocytopenia (<50,000) P= 1.000 Prolonged prothrombin time (INR> 1.4) P=0.656 Cause of head trauma P0.651: Fall from standing Motor vehicle accident Fall from a height Assault Bicycle accident Mean SDH maximal thickness (mm, range) P<0.001* Mean midline shift (mm) P<0.001* Presence of cerebral contusion P= 0.003*	Study Recruitment: Low risk Retrospective case note review- depends on information being recorded correctly. Attrition: low risk All patients appeared to have been followed up appropriately Prognostic factor measurement: Low risk Appears CTs have been reviewed and volume measurements conducted by member of study team Outcome measures: Low risk All patients followed up until clinic. No reports of deaths. Confounding Factors: Low risk None obvious-exclude patients with other injuries Statistical techniques: Low risk Well presented Overall Only patients with subdural- have been shown to high risk in other
42 43 44 45 46 47				Mary Ann	Liebert, Inc, 140	Huguenot Street, New Rochelle, NY 10801	

15 • Other significant injuries • Patients refusing surgery	* 76			Presence of SAH, P=0.003* Diffuse cortical atrophy Mean bifrontal ratio (range)P= 0.345 Mean Sylvian fissure ratio (range) P=0.602 Multivariate analysis of prediction of delayed haematoma evacuation. Maximal thickness P=0.527 OR 2.5 (0.5-41.1) Volume haematoma P=0.01 OR= 1.1 (1.02 -1.17) Midline shift P=0.01 OR=1.43 (1.09-1.89) Cerebral contusion P=0.92 OR 0.85 (0.18-3.97) SAH P=0.43 OR 0.53 (0.11-2.56)	studies. The neurosurgical rate for these injuries appears v. high ?length of follow up. These patients have been discharged and then undergone reimaging as outpatients. Doesn't preclude early discharge of some of these patients but they will need to be followed up.
Overton et al 2014	Retrospective Cohort Study Aim Reports initial experience with the management of MTBI by trauma surgeons alone. Hypothesize that patients with MTBI managed by trauma surgeons will be the same as outcomes for patients managed by neurosurgeons.	Outcome measured GOS score at discharge 1= death 2=severe disability 3=mod disability 4= full recovery Method Mulitvariate regression analysis to assess whether admission under trauma surgeons affected likelihood of GOS >3 (good recovery)	trauma versus neurosurgical management age, sex, race/ethnicity, injury severity, insurance status GCS	171 patients 8 deaths 4 severe disability 24 moderate disability Neurosurgeons managed 120 Trauma surgeon 51 Multivariate regression analysis to predict GOS >3 (full recovery) Admission Trauma surgeon P=0.3OR 1.74(0.61–4.92) Age P<0.001®OR0.94 (0.91–0.96) ISS P<0.001 OR0.87 (0.81–0.94) GCS P=0.005 OR13.96(2.23–87.3) Other factors in model but no results reported: sex, ethnicity, ISS, insurance status Mean/median GCS=14.7 Mean/median age= 49	Study Recruitment: Mod risk Retrospective case note review- depends on information being recorded correctly. Only patients with bleed less than 1cm Attrition: Mod risk Not clear when outcomes measured- if at discharge low risk Prognostic factor measurement: Low risk Doesn't explain how CT reports interpreted and how 1cm cut off decided. Outcome measures: mod risk States GOS- but not when or who determined score ?self reported Confounding Factors: Mod risk None obvious Statistical techniques: Mow risk States backward step binary logistic regression analysis performed to assess trauma

admissions- controlled for age, sex,

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1 2 3 4							
5 6 7		60	<u></u>				race, ISS, insurance status and GCS motor scores- presents the analysis for only some of these.
0 1 2			196	L;			Overall Limited by inclusion criteria of less than 1cm and even though no difference in outcomes with who patients were admitted under, potentially the patient groups received different care.
4	Schwed et	UCLA California	Retrospective	Favorable	Vital signs	380 TBI patients in study period	Study Recruitment: Mod risk
5 6	al 2016 California USA	Level 1 trauma centre 2012-2015	Aim	outcome- composite outcome of	AIS ISS CT findings-Marshall	19 missing records 201 remaining cohort met inclusion/exclusion criteria	Only admitted to ICU- higher risk group than total population.
7 8 9		Inclusion criteria: • Patients	Identify admission variables associated with	following: Alive at discharge ICU admission for	and Rotterdam scores	4/201 deaths (2 attributable to bleed progression) 129/201 GCS15	Attrition: Low Risk Only inpatient measures
С		identified on trauma registry and	favourable outcomes with mTBI and intra-	less than 24 hours No in hospital complications		6/201 neurosurgical outcomes	Prognostic factor measurement: Mod risk
1 2 3		case note review	cranial haemorrhage	Did not require neurosurgery		21% (42) in hospital complication	Does not assess pupillary response or anticoagulation/antiplatelets
4 5		Initial GCS13-15Intra-cranial	Method	Failed to achieve this if required		78/201=met conditions favourable outcome 0/1 EDH favourable outcome 1/4 ICH favourable outcome	Outcome measures: Mod risk Only inpatient related outcome
6 7		bleed any variety	Univariate and multi-variate	ventilation or ionotropic		18/36 SDH favourable outcome 30/57 SAH favourable outcomes	measures.
3		identified by CT imaging Excluded:	regression analysis prediction of	support at any point.		22/83 mixed lesions favourable outcome 123/201=unfavourable outcome	Confounding Factors: Mod risk Cohort includes patients with multiple injuries- 2 deaths appear
) I		TransfersNot admitted	"favourable outcome composite			Univariate comparison between patients with favourable and unfavourable outcomes: Age P=0.01	due to factors unrelated to head injury
2 3		to ICU • Required emergent	measure"			ISS P=0.001 Head AIS P=0.026	Statistical techniques: Mod Risk Selective reporting of significant
4 5		neurosurgeryPatients less				Time to first head CT (hours) non-significant ED systolic blood pressure P= 0.01 ED heart rate P=0.48	results. Does present statistical
6 7		than 18 In police custody				Marshall score P=0.11 GCS at time of admission ICU P <0.0001	comparison between the groups with favourable and unfavourable
8 9		 Pregnant 				GCS 15 at admission P=0.0001 Type of hemorrhage Epidural P=0.42	outcomes
10 11						IVH P=0.55 SDH P=0.1	0/,
12 13							
44 45							
16 17				Mary Ann	Liebert, Inc, 140	Huguenot Street, New Rochelle, NY 10801	

				T	ı		1
						SAH P=0.02 Combination P=0.002	
						55.118.118.116.11	
						All factors statistically significant in univariate analysis were assessed in multivariate	
						analysis	
						Multivariate model predicting favourable outcome: including ED BP, Marshall score,	
						Isolated SAH, Head AIS, ISS<25, GCS15 at ICU admission and age<55	
,						GCS 15 at ICU admission OR 5.5 95% CI (1.6-18.8) P=0.006	
						Isolated SAH 5.1 95% C.I. (1.5-17.6) P=0.01	
						Age<55 OR 3.5 95% C.I. (1.1-11.2) P=0.03	
						Mean/median age= 60	
	Thorson et	Miami	Retrospective	Progression of	CT findings-	1510 patients with GCS13-15 and head injury	Study Recruitment: High risk
	al 2012	Level 1 trauma	cohort study	initial lesion or	including type of		Neurosurgeon have selected out
	Miami	centre	•	new lesion	injury, presence of	537/1510 +ve initial CT scans	patients with "trivial" injuries-
	USA	1996-2010	Aim	identified.	oedema, mass	62 proceeded immediately to surgery and 115 no repeat CT in 24 hours- (mostly as the	makes this a higher risk group than
			To test whether		effect or herniation.	neurosurgeon deemed injury insignificant).	population of interest
		Inclusion criteria: Initial GCS13-	routine CT imaging in mTBI	Neurosurgical intervention.	Age Sex	360/537 had repeat CT imaging.	Attrition: Low Risk
		15	with detected	intervention.	ISS	300/337 Had repeat of imaging.	Only inpatient measures
		 Present on 	intra-cranial	Death.	GCS	11% of repeat CT scans-recalled (i.e.no actual injury)	,,
		trauma	injuries provides		Abnormal	108/360- progression on CT imaging	Prognostic factor measurement:
		registry	useful		neurological		Low risk
		Head	information in the absence of		examination- change in GCS	Mean/median GCS=14.5 Mean/median age= 47	Loose definition for abnormal neurology
		abbreviated AIS 1 or	neurological		greater than 1, GCS	Percent anticoagulated=3	neurology
		greater	deterioration		less than	r el celle di lescagalates d	Outcome measures: Mod risk
		No other			13,Neurological	Age No change 46 SD 20 Progression 50 D 23 P=0.13	Only inpatient related outcome
i		injuries (AIS=0	Methods		deficit, or significant	Sex No Change Male 178 Progression 79 PO.11	measures.
		other body	Step wise multi-		symptoms including	Intubated No Change 22 Progression 17 P=0.05	Conformation Footomer Lawrence
		regions)	variate regression for factors P<0.2		headache, lethargy, visual disturbance.	ISS No change 12 SD 5 Progression 15 SD 6 P<0.01 GCS 15 arrival No Change 158 Progression 37	Confounding Factors: Low risk None obvious
		 Repeat CT head scan if 	associated with		visual distarbance.	GCS 14 No Change 65 Progression 43	None obvious
		intracranial	progression on CT			GCS 13 No Change 31 Progression 28	Statistical techniques: Mod Risk
		injury	and craniotomy			Anticoagulant Use No Change 17 progression 11 0.29	Selective reporting of outcomes in
		detected. (4-6				Aspirin No Change 7 Progression 3	regression model
		hours after				Plavix No Change 1 Progression 2 Coumadin No change 2 Progression 4	Paper concludes all patients should
		initial CT).				LMWH No Change 2 Progression 0	have a repeat CT as 7/360 patients
		Note neurosurgeons				Multiple No Change 5 Progression 2	had neurosurgery based solely on
		decided				PT No Change 12.2 Progression 12.6 P= 0.443	repeat CT head findings.
		whether a				PTT No Change 25.2 Progression 24.8 P=0.85	
							Possibly include but is a higher risk

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1 2 3 4	0/-						
5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24 25 26 27 28 29 30 31		lesion was to insignificant to warrant a repeat CT Excluded: Penetrating trauma Pregnant Age<18 Incarcerated Transfers	***		7 0	Age No Neuro Surg 47 SD 21 Neuro Surg 51 D 23 P=0.97 Sex No Neuro Surg Male 241 Neuro Surg 22 P0.11 ISS No Neuro Surg 13 SD 5 Neuro Surg 17 SD 6 P<0.01 GCS 15 arrival Neuro Surg 180 Neuro Surg 13 GCS 14 No Neuro Surg 100 Neuro Surg 8 GCS 13 No Neuro Surg 50 Neuro Surg 9 Anticoagulant Use No Neuro Surg 22 Neuro Surg 6 0.024 Aspirin No Neuro Surg 9 Neuro Surg 13 Plavix No Neuro Surg 2 Neuro Surg 2 Coumadin No Neuro Surg 2 Neuro Surg 2 Coumadin No Neuro Surg 2 Neuro Surg 4 LMWH No Neuro Surg 4 Neuro Surg 2 PT No Change 12.1 Progression 12.0 P= 0.35 PTT No Change 25 Progression 27.5 P=0.45 7/30 operated patients solely on basis of worse CT (no prior neurological decline) 22/360 deaths Logistic regression analysis: unclear which factors were tested in the model Predictors of worse 2 nd CT AU ROC curve 0.703 GCS=13 OR4 95% CI 2.02-7.93 P<0.001 GCS=14 OR 3.11 95% CI 1.07-5.48 P<0.001 ISS OR 1.07 95% CI 1.02-1.11 P<0.001 Mass effect OR 2.02 2.02-3.78 P<0.001 Predictors of craniotomy: AUC ROC 0.849 Initial mass effect OR 5.24 95%C.I. (1.96-14.1) P=0.001 New/worse EDH 2 nd CT OR 23.3 3.67-148.3 P=0.001 New/worse EDH 2 nd CT OR 23.3 3.67-148.3 P=0.001 New/worse mass effect 2 nd CT 5.73 95% 1.64-20)	population given selection out of patients with "non-significant" findings. Note also 11% of 360 repeat CTs recalled-i.e. initial finding not present (4/6 hours after injury).
32 33 34 35 36	Quigley et al 2012 Pennsylvani a USA	Level 1 trauma centre	Retrospective Cohort Study Aim To assess if traumatic subarachnoid haemorrhage more benign form of mTBI Multivariable	Discharge home Clinical deterioration CT progression Neurosurgery	Demographics Mechanism of injury Number and results of follow up CT Length of hospital and ICU admission ISS CTs re-reviewed by study radiologist	New/worse herniation 32.1 95% C.I. 7.83-131.6 P=0.001 547 patients identified as subarachnoid 478/547 isolated subarachnoid 470/478 repeat CT imaging 15/470 worse CT (1 is new stroke) 342/478 discharged home 51/478 discharged rehab or nursing home 4/478 self discharge 4/479 long term care facility 1/479 other facility	Study Recruitment: Low risk Identified from prospective trauma registry- dependent on how accurate this is Attrition: Mod Risk Not clear whether and when all patients followed up but presents outcomes from outpatient clinic Prognostic factor measurement:

	Present on trauma registry Initial GCS13-15 Isolated subarachnoid haemorrhage	analysis computed with step-down logistic regression- discharge home primary outcome			1/479 to hospice 6 week follow up 1/478 bilsteral subdural- drained States surgical intervention 0.2% Step down Multivariate regression with outcome discharge home Age P<0.0001 Admission GCS P=0.0018	Low risk Ct scans reviewed Outcome measures: Mod risk Not clear if uniform outpatient followup Confounding Factors: High risk Clearly an old patient population-
	Does not state adult only but mean age 65.7		Vie,	1	ISS P=0.0088 Not progression of bleed on CT	discharge to rehab/nursing home like related comorbidities or other injuries Statistical techniques: High Risk Selective reporting of outcomes in regression model No confidence intervals or odds ratios. No explanation of high the model
Volmahas	Macrachusotte	Detroppestive	Surgical	Demographics	602 nations had CT for hard injury	was derived General comments: Discharge outcomes contradict low level of intervention. Unable to pool risk factors as are. Can pool to confirm Subarachnoids are low risk.
Velmahos et al 2006 Massachus etts USA	Massachusetts Level 1 trauma centre 2003-2004 All patients with intra-cranial injuries	Retrospective cohort study Comparison univariate characteristic patients with	Surgical or medical intervention following repeat CT (caniotomy, ICP monitoring, intubation or magnital	Demographics ISS Admission observations Time interval between admission and 1st CT and	692 patients had CT for head injury 179/692- for scheduled repeat CT 154/692 repeat CT due to intracranial injury 25 no lesion- repeat CT due to anti-coagulation 37/154 worse CT	Study Recruitment: Low risk Identified from trauma registry- dependent on how accurate this is Standard model of care for all patients
	identified reviewed by a neurosurgeon and repeat CT scheduled within 24 hours. Inclusion criteria: Present on	worse CT scans compared with the same or improved. Where P value 0.2 or less included in stepwise logistic regression model	mannitol, increased ventilation, CSF drain, sedation, transfer to ICU) Worse repeat CT	subsequent CT scans	7/154- medical or surgical intervention due to deterioration 4/154 neursourgical 8/179 deaths 1/44 subdurals neurosurg 0/33 SAH neurosurg 1/13 intra-parenchymal neurosurg 0/7 extra-durals	Attrition: Low Risk Appears only inpatient outcomes Prognostic factor measurement: Mod risk Assessment of time to CT- not clear biological mechanism how this affects outcome or how measured
	trauma registry Initial GCS13-				2/57 multiple neurosurgical Male P=0.44 Age (years) P0.01	Outcome measures: Mod risk Takes reports from attending at face value.
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1 2 3 4							
5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24 25 26		Blunt head injury Repeat CT for intra-cranial injury Presumably adults age presented as mean 48 and SD 25	***************************************		*O	≤65 P<0.01 Mechanism of blunt trauma P= 0.31 Fall Road traffic accident Other 0.31 Injury Severity Score P=0.01 ISS>16 0.09 Glasgow Coma Scale score on arrival P=0.02 Systolic Blood Pressure on arrival (mm Hg) P= 0.63 Anticoagulation therapy P=0.25 Time from arrival to CT P<0.01 First head CT findings solitary or multiple findings P<0.01 Time between first and second CT P=0.10 Stepwise logistic regression model to predict worse CT Time from injury to CT <90 mins OR6.37 95% CI 2.29-17.76 P<0.1 Age>65 OR3.33 95% CI 1.29-8.60 P=0.01 GCS<15 OR 3.13 95% 1.23-8.01 P=0.02 Multiple lesions OR 11.03 95% CI 1.32-92.06 P=0.03 AUC ROC curve 0.83 If all 4 factors present 83% chance worse CT If none present 2% chance worse CT Mean/median GCS=14.7 Mean/median age= 51 Percent anticoagulated=10	Does not report deaths as a primary outcome but included in table- not clear what the cause of deaths is. Confounding Factors: High risk Not isolated head trauma and no selection out of comorbid patients-does not appear deaths related to head injury but clear Statistical techniques: Mod Risk Selective reporting of outcomes in regression model General comments: Time to initial CT highly significant-slightly odd for this study population- not examined any other study. No explanation for deaths given in paper.
27 28 29 30 31 32 33 34 35 36 37 38 39 40 41	Fabbri et al 2013 Italy- multicenter	Multi-centre 32 Italian hospital- both specialist and general 2009 Inclusion criteria:	Retrospective multicentre cohort study Aim To assess whether pre-injury antiplatelet use lead to worse outcome in patients with intra-cranial injuries detected by CT imaging	Worse repeat CT defined as increase point on Marshal criteria within 24 hours Neurosurgery within 7 days GOS at 6 months	Age Sex Mechanism Coagulation GCS Anti-platelet medications Type of injury on CT Marshal Classification	Study of all GCS patients but present data for GCS14-15: 1123/1558 patients GCS14-15 Antiplatlet therapy increased the risk of a worse CT: When 2 or less lesions RR 1.86 95% CI 1.06-3.30 P=0.032 When 3+lesions RR 3.34 95% CI 1.74-6.40 P=0.003 87/1123 Worse Characteristic on CT Mean/median age= 65	Study Recruitment: Mod risk The paper is not clear about how patients were identified and data extracted Also patients requiring emergency surgery within 7 days based on initial CT excluded- may select out higher risk groups- in practice excluded Marshall 5/6 patients which is reasonable Attrition: Low Risk No loss to follow up and standard care for all patients to be reviewed at 6 months Prognostic factor measurement:

Shih et al	Marshal category 2-4 Within 24 hours of injury Excluded: Need immediate neurosurgery GCS 3 fixed dilated pupils Unclear history of mechanism Hypotension< 90 systolic Penetrating Injuries Discharge against medical advice	Potromoctive Neurologic	Say	240 patients mot inclusion criteria	Low risk Scans all re-reported Outcome measures: Low risk Good outcome end points Confounding Factors: Mod risk Not isolated head trauma and state no need to control for comorbidities as shown not to affect head injury outcome Statistical techniques: Low Risk Appropriate and well presented General comments: Good study Fabbri previously shared data- ?request GCS13-15 subset
Shih et al Taiwan 2016	Tertiary referral Teaching hospital Taiwan No time frame given Inclusion criteria:	Retrospective cohort study Aim Determine the potential risk factors of delayed neurosurgical intervention in mTBI with intracranial haemorrhage Stepwise logistic regression to identify variables that predicted failure of conservative treatment Neurologic deterioration-GCS drop 2+ points seizures, sign raised ICP Repeat CT identify deterioration-whether worse Neurosurgical intervention-including craniotomy, craniectomy	Mechanism of injury GCS ISS Laboratory results	340 patients met inclusion criteria 13/340 neurosurgical outcomes 25/340 neurological decline 7/118 mixed lesions neurosurgery 34/340 worse CT 3/340 died Univariate analysis: delayed neurosurgery versus non-neurosurgery Median age P=0.082 Male/female P=0.573 OR 0.648 95% CI 0.196–2.149 GCS P= 0.189 Anti-platelet and/or warfarin therapy P=0.403 OR 2.188 95% CI 0.263–18.222 Statin therapy P= 1.000 Hypotension 0 4 P= 1.000 WBC count (1000/mL)P=0.023 RBC count (1000/mL) p=0.401 Hemoglobin, P=0.606 Coagulopathy P=1.000 Hypertension P=0.526 OR 0.484 95% CI 0.105–2.228 Diabetes mellitus P=1.000 OR 1.028 95% CI 0.221–4.780 (!?)0 Old cerebral vascular accident=1.000 Coronary artery diseases P=1.000 Liver cirrhosis P=1.000	Study Recruitment: Lod risk No uniform criteria for which patients undergo immediate neurosurgery- just selected by neurosurgeon Attrition: Low Risk Only inpatient measure Prognostic factor measurement: Low risk Scans all re-reported Outcome measures: Mod risk Only inpatient measures- potential for discharge and deterioration Confounding Factors: Mod risk Not isolated head trauma Statistical techniques: Mod Risk Mod risk selective reporting of significant prognostic factors. Does not report whole model.
	·	Mary Ann	Liebert, Inc, 140	Huguenot Street, New Rochelle, NY 10801	Y/jon

				Jou	rnal of Neurotrauma	Page 118 of
1 2 3 4						
5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24 25 26 27	reviewed neurosurgeon determined whether immediate neurosurgery conservative management	***************************************			Chronic renal disease P=1.000 Renal failure P=1.000 ISS score, Median P=0.005 Single intracranial heamorrhageP=0.149 Multiple intracranial heamorrhage P=0.149 EDH P ≤0.001 OR 9.923 95% CI 3.105–31.708 SDH P=1.000 OR 0.906 95% 0.298–2.753 IPH P=0.366 OR1.812 95% CI 0.594–5.526 SAH P=0.044 OR0.251 95% CI 0.068–929 IVH P= 0.111 OR13.542 95% CI 1.147–159.876 Midline shift P≤0.001 OR19.813 95% CI5.495–71.435 Skull fracture P≤0.001 OR21.750 95% CI4.707–100.510 Pneumocranium P=0.621 Volume of EDH P≤0.001 Volume of SDH P=0.092 Volume of IPH P=0.657 Stepwise logistic regression: model included WBC count, midline shift, skull fracture large volume EDH and higher ISS- significant predictors of delayed neurosurgery. Volume of extra-dural haemorrhage associated with delayed neurosurgery Increase volume EDH 1 cubic cm increase risk of neurosurgery by 16% (p=0.022 OR 1.190 95% CI 1.041-1.362) AUC volume EDH=0.917 (95% CI 0.797-1.00) Mean/median GCS=14.7 Mean/median age= 50	Also some apparent mistakes in univariate analysis General comments: Does not report outcomes by single lesion type
Bardes et al 2016 USA USA 2016	Level 1 trauma centre West Virginia 2009-2011 All mTBI patients with bleeds admitted to general surgical ICU with a neurosurgical consultation Inclusion criteria: Blunt TBI Age>18 GCS13-15	Retrospective Cohort study Aim: Identify low risk mTBI patients with intra-cranial bleeds that do not require admission to ICU	Documented neurological decline Medical intervention Neurosurgical intervention	Admissions GCS GCS 6, 12, and 24 hours Type of bleed Bleed progression on CT Aspirin Clopidogrel Warfarin Admission Coag ISS	389 patients met inclusion criteria 5.1% (20) in hospital mortality 53/389 patients neurological decline 376/389 scheduled repeat CT 69/376 worse CT 35/389 craniotomy 46/389 patients required medical or neurosurgical intervention Univariate comparison patients with decline versus no neurological decline GCS<15 P=0.002 SDH P=0.0025 Age≥55 P=0.001 Use Warfarin P=0.039 ISS P=0.22 AIS=P=0.12 SAH P=0.15 EDH P=0.18	Study Recruitment: Lod risk Representative sample of population of interest. Limitations of retrospective data collection Attrition: Low Risk Only inpatient measure Prognostic factor measurement: Low risk Scans not re-reported Outcome measures: Mod risk Only inpatient measures- potential for discharge and deterioration
12 13 14 15 16 17			Mary Ann	Liebert, Inc, 140	Huguenot Street, New Rochelle, NY 10801	

	ISS<25 Excluded: Penetrating injury GCS<13 States in results all patients had evidence of intracranial haemorrhage on bleed- doesn't define what this includes			ICB P=0.051 Aspirin P=0.54 Clopidogrel P=0.17 PT P=0.042 aPPT P=0.0028 Admision INR P=0.42 Decision tree subgroup analysis: No GCS15 patient ≤ 55 underwent neurological decline= low risk group Mean/median GCS=14.8 Mean/median age= 63 Percent anticoagulated=12	Confounding Factors: Mod risk Not isolated head trauma or control for comorbidities Does use ISS to exclude severe polytrauma Statistical techniques: Mod Risk Mod risk selective reporting of significant prognostic factors. Does not present decision tree analysis transparently
Sharifuddi et al 2012 Malaysia	Patients admitted under neurosurgeons 2008-2009 specialist centre Inclusion criteria: GCS 13-15½ 12 years and older½ positive initial head CT½ isolated blunt head injury½ presented within 24 hour of initial injury Excluded: previous history of head injury½ on anticoagulatio n therapy (aspirin, heparin or warfarin)½ polytrauma Major comorbidity	Prospective observational study Aim To evaluate whether the repeat head CT were useful in providing information that leads to any neurosurgical intervention To evaluate whether the repeat head CT were useful in providing information that leads to any neurosurgical intervention Worsened (increase in size or evidence of new intracranial lesion). Surgical interventions: craniotomy, intracranial pressure monitor placement or intubation.	Admission GCS Associated symptoms®Post- traumatic amnesia Headache Vomiting Dizziness Type of injury identified	Neurological decline 66 patients (23.7%) Worse CT in 58 patients (20.8%). 31 (11.1%) patients neurosurgical outcome. 3 deaths. Univariate comparison patients with progression on CT and without: Male P=0.189 Age ≥ 65 P < 0.001 Ethnic groups P=0.624 Mechanism of injury MVA versus others P=0.333 GCS<15 P=0.003 Post-traumatic amnesia P=0.069 Headache P=0.019 Vomiting P=0.441 Dizziness P=0.262 Multiple lesion P=0.001 Base of skull fracture P=0.842 Hb (g/litre) on admissionP0.009 INR on admission P=3 0.388 Stepwise multiple logistic regression model	Study Recruitment: Low risk Retrospective case note review-depends on accuracy of notes. Not clear if all patients with ICH admitted under neurosurgeon-potential for selection of high risk population. Note age 12+ does not strict meet inclusion criteria. Attrition: Low RIsk Outcomes only during hospital admission- no loss to F/U Prognostic factor measurement: Mod risk The mechanism of injury- doesn't discriminate between high and low risk mechanisms. CT interpreted once by attending radiologist or neurosurgeon. No quality control. Outcome measures: low risk As reported outcomes of worse CT, neurosurgery or death as an inpatient low risk for bias. However, no follow up outcome measures for delayed deterioration. Confounding Factors: Mod risk

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1 2 3 4 5 6 7 8 9 10	0,-	 suspected drug or alcohol intoxication, Neurological impairment trauma Immediate 	^_^_			Risk factors for progression on CT: Age ≥ 65 P<0.001 95%C.I. (0.098- 0.364) Multiple lesions on initial CT P=0.018 95% C.I.(0.239- 0.877) GCS score < 15⊞P= 0.016 95% C.I. (1.164 - 4.333) 44/144 multiple lesion worse CT	Possibility of anti-coagulants. Not recorded. Statistical techniques: Mow risk Stats do not present what the risk measure is- presumably an OR. Also selective reporting of significant results.
11 12 13 14		 neurosurgery Admitted ICU for close observation 	, C	Vio		Mean/median GCS=14.6 Mean/median age= 39 Percent anticoagulated=0	Only for progression on CT-dubious value
15 16 17 18 19 20 21 22	Sumritpradi t et al 2016 Bangkok Thailand	Patients admitted to an Acute Care Unit surgery 2009-2013 Inclusion criteria: Admission<72 hours hours 16 years and older positive initial	Retrospective cohort study Aim: To determine the value of repeat CT imaging in TBI for risk stratification of patients	Neurologic deterioration: reduced consciousness, limb weakness, lateralizing signs, severe headache, vomiting, and dizziness.	Age Sex Co-morbidities Medications Initial GCS AIS Medications CT findings	145 patients matched inclusion criteria 98/145 GCS13-15 74/98 routine repeated CT scans (36/98 worse) (1/74 neurosurgical) 24/98 clinically deteriorated and underwent CT imaging (7/28 neurosurgery) Overall	Study Recruitment: High risk Only recruited patients that neurosurgeons had planned a repeat CT scan (293/442 patients with injuries no repeat CT versus 149/442 for repeat CT) Selection bias of higher risk group then all GCS13-15 patients with CT detected injuries
23 24 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39		Non-surgical initial management Includes all GCS score but presents data for GCS13-15 patients Patients under went repeat CT imaging-determined after neurosurgical review		Neurosurgery		8/98 GCS13-15 patients neurosurgery 24/98 some clinical deterioration-prompting repeat CT GCS13-15 Univariate comparison patients underwent neurosurgery and did not. Age>50 P=0.478 Mean age P=0.295 Male P=0.706 Traffic injury=0.256 Diabetes mellitus P=0.354 Hypertension P=0.135 Ischemic heart disease P=0.070 Cerebrovascular disease P=0.592 Aspirin =1.000 Warfarin P=1.000 Clopidogrel P=0.017 ISS, mean p= 0.405 ISS > 19 P= 0.282 Brain AIS, mean P=0.080 AIS > 4 P=0.073	Attrition: Low Risk Outcomes only during hospital admission- no loss to F/U Prognostic factor measurement: Mod risk No outline of how CT scans reported and risk stratified b Outcome measures: low risk As reported outcomes of worse CT, neurosurgery or death as an inpatient low risk for bias. However, no follow up outcome measures for delayed deterioration. Confounding Factors: Mod risk Does not state how patient with other injuries delt with
40 41 42 43 44 45 46		1	1	Mary Ann	Liebert, Inc, 140	O Huguenot Street, New Rochelle, NY 10801	Statistical techniques: Low risk

1					Heart rate on admission, mean n= 0.00E	Droconte cimplo universista anclusta
ļ					Heart rate on admission, mean p= 0.095 Epidural hematoma P= 1.000	Presents simple univariate analysis
ŀ					Subdural hematoma P= 1.000 Subdural hematoma P=0.136	between neurosurgical and non- neurosurgical patients
ŀ					Subarachnoid haemorrhage P=0.464	neurosurgicai patients
ŀ					Hemorrhagic contusion P=0.715	Is a higher risk population due to
ŀ					•	
ŀ					Intraventricular hemorrhage P=1.000	selection for repeat CT imaging-
ŀ					Diffuse axonal injury P=) 1.000 Skull fracture P=1.000	possibly unable to include in any meta-analysis.
ŀ					Base of skull fracture=0.409	illeta-allalysis.
ŀ					Midline shift > 2 mm P=0.003	!
ŀ					Duration from injury to 1st CT P=0.603	
ŀ					Duration from finjury to 1st CF F-0.003	
			1/6		Odds ratios associated with these factors reported separately:	
l					Subdural hematoma OR 5.3 95%CI (0.63–45.33) P=0.136	
ļ					Hypertension OR 4.1 95% CI (0.78–21.46) P=0.135	ļ
ŀ					AIS > 4 OR 4.0 95%CI (0.91–17.55) P=0.073	
					Ischemic heart disease OR 4.8 95% C.I. (0.99–23.19) P=0.070	
					Clopidogrel OR 10.2 95C.I. (1.87–55.38 P=0.017	
					Midline shift > 2 mm OR11.9 95% C.I. (2.50–57.20) P=0.003	
ļ					Neurological deterioration resulting in CT OR 30.0 95% C.I. (3.46–280.83) P<0.001	
l					rearbioglear acterioration resulting in er on 50.0 55% c.i. (5.40 250.05) 1 40.001	
l					Mean/median age= 57	
l					Percent anticoagulated=4	
ifri et al	New Jersey	Prospective	Neurosurgery	Abnormal	161 patients GCS13-15 with intra-cranial bleed	Study Recruitment: Mod risk
006	Level 1 trauma	Cohort Study	following second	neurological		
lew Jersey	centre	•	scan	examination prior	10 excluded due to co-morbidities.	Only patients with repeat CT- likely
JSA	2002-2003 12	Aim		to repeat CT	5 required immediate neurosurgery	to be a higher risk group
ŀ	months	Prospectively	Admission to ICU	(GCS<15 or severe	16 did not undergo repeat imaging	
ŀ		assess the value	or administration	headache/vomiting/		Attrition: Low Risk
ŀ	Inclusion criteria:	of a repeat CT in	of mannitol	gross motor or	130 in study population	Only inpatient measures
ŀ	Initial GCS13-	patients with	following second	sensory deficits)		
ŀ	15	mTBI and intra-	scan		99 normal neurology at time of repeat CT; 31 abnormal neurology at time of repeat CT.	Prognostic factor measurement:
ŀ	 Intra-cranial 	cranial		Sex		Mod risk
ŀ	bleed- intra-	haemorrhage and	In hospital	Age	0/99 neurosurgery	Does not try and grade severity of
ŀ	cerebral,	normal	mortality.	GCS	1/99 death (unrelated to intra-cranial injury)	CT findings as predictor.
ŀ	extra-dural,	neurological		Mechanism	13% 99 CT scans worse	
	subdural	examination	GOS at discharge.	Type of injury	2/31 neurosurgery	Loose definition for abnormal
	subarachnoid			identified by CT	5/31 deaths	neurology- sometimes prompted
	or contusion	Repeat CT within	Discharge		14/31 repeat CTs worse	repeat CT and no uniformed time
	Excluded:	24 hours	destination			when all CT scans performed.
ļ	 Previous brain 				Abnormal neurological exam predicts changes repeat CT OR 5.28 CI2.08-13.4 P=0.002	
	surgery or					Outcome measures: Mod risk
ļ	cerebral				Mean/median GCS=14.6	Only inpatient related outcome
	pathology or				Mean/median age= 45	measures.

Medication Suppose Medication Suppose Medication Suppose Suppose Medication Suppose Su		admission Data only presented for adults (15-94)	<u>^</u>				
Risk of Neurosurgery Unstable GCS unadjusted 4.16 (0.51-33.63) adjusted 2.98 (0.35-25.18) P=0.32 ISS Unadjusted 1.04 (1.01-1.07) adjusted 1.05 (0.99-1.12) P=0.10 Age Unadjusted 1.01 (1.00-1.02) ajusted 1.11 (0.96-1.28) Fabbri et al 2008 Bistrict general hospital rural Italy Cohort study Follow up GOS at 6 months (includes Charlson Co- Charlson Co- Charlson Co- Anonymised individual patient made available by authors and used for analysis.	Thesis 2015	centre California 2007-2011 Patients identified on a hospital trauma registry Inclusion criteria: Initial GCS13- 15 Blunt head trauma Positive CT scan. 2 or more CT scans 18+ Excluded: Pregnant Age<18 Penetrating	Cohort Study: To assess whether GCS 15 patients with intra-cranial haemorrhage that maintain a GCS of 15 benefit from routine CT	imaging Neurosurgical	Anti-coagulant Medication ISS LOC Skull fracture displaced/undisplac ed Neurological symptoms Time interval between scans GCS/deterioration	88 incomplete notes 201 only 1 CT scan Study population 369 patients with at least 2 CT scans. 111/369 GCS 15 at presentation and throughout. 0/111 neurosurgery 20.7% of 111 worse CT 0.9% mortality 258 GCS<15 at some point during hospital admission 37.6% 258 worse CT 11/258 neurosurgery 2.7% 258 deaths Overall 11/369 neurosurgical interventions Mean/median age= 53 Progression of Injury: Unstable GCS < 15 Unadjusted OR 2.21 (95% C.I. 1.33-3.68) adjusted 1.71 (95 % C.I.1.00-2.91) P=0.05 ISS Unadjusted 1.04 (95% C.I. 1.01-1.07) Adjustede 1.1 (0.99-1.05) P=0.27 Age Unadjusted1.01 (95% C.I. 1-10.2) Adjustede 1.01 (0.99-1.02) P=0.08	Approximately 1/3 of patients with injuries detected by CT imaging not included either because incomplete or only 1 CT scan. Patients on which multiple scan conducted likely to be higher risk. Attrition:Low Risk Low risk- inpatient outcomes Prognostic factor measurement: Medium risk No re-reporting of CTS Does not include CT findings as a prognostic factor. Outcome measures: Medium risk No outcome measures after discharge Confounding Factors: Medium risk No control for comorbidities
(includes Charlson Co- Anonymised individual patient made available by authors and used for analysis.			'	•		Unstable GCS unadjusted 4.16 (0.51-33.63) adjusted 2.98 (0.35-25.18) P=0.32 ISS Unadjusted 1.04 (1.01-1.07) adjusted 1.05 (0.99-1.12) P=0.10 Age Unadjusted 1.01 (1.00-1.02) ajusted 1.11 (0.96-1.28)	Performs different analysis for neurosurgical outcomes compared
recruitment from Evaluate the Injury Severity Score	Italian	Prospective	Aim:		Charlson Co- morbidity Index,	Anonymised individual patient made available by authors and used for analysis.	0//*

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5	1999-2006	effects on	Neurosurgical	GCS	
3		outcome of a	intervention	CT scan results-	
7		model based on	within 7 days.	Marshall category	
<u>'</u>	Inclusion criteria:	observation in a	•	Type of Injury	
3	Admission GCS	neurosurgical unit			
9	score ≥ 9	versus			
10	Age over 10	observation in a			
11	Initial head CT	peripheral			
	scan positive	hospital with			
12	for any type of	neurosurgical			
13	trauma	expertise via a			
14	• Initial non-	teleradiology			
15	operative	system and a NSU			
16	management.	transfer time of		1 -	
	Excluded:	30–60 min			
17	Persistent				
18	hypotension				
19	caused by				
20	additional				
24	injuries				
21	Patients				
22	requiring immediate				
23	surgery				
24					
22 23 24 25	 Penetrating injuries 				
20	Patients that				
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Suppleme	upplementary Material 2: Data Extracted from Included Studies Papers deriving and validating the BIG criteria N=3 (not included in meta-analysis)							
Reference	Population	Study Design	Outcome Measures	Prognostic factors assessed	Results	Quality Appraisal		
Joseph et al 2014 USA Study 1: defining the BIG criteria	Level 1 Trauma centre 2009-2011 Inclusion criteria:	Retrospective Cohort Study- Aim: Define guidelines for based patients' history, examination and initial CT head findings regarding which patients require observation in ED, RHCT or neurosurgical consultation. Local consensus for categories	Neurosurgical intervention Progression of CT findings on a repeated scan Neurological deterioration if BIG 1 or 2- GCS<12, abnormal focal neurology or abnormal pupils	_	Results 1232 patients TBI with positive CT scan 121=BIG 1 313=BIG 2 798=BIG 3 888/1232 underwent repeat CT 13% (159) patients neurosurgical outcome- all in BIG 3 category. No BIG 1 patients had neurological deterioration No Big 1 patient worsening CT 2.6% (9) BIG 2 patients worsening CT 2/313 BIG 2 patients deteriorated neurologically- transferred to neurosurgical care. No BIG2 patient needed neurosurgery BIG3 patients 21.6% worsening CT 3% neurosurgical intervention	Study Recruitment: Low risk bias Retrospective cohort review- reliant on accuracy of written notes. Cohort identified by case note review but no details of how this was done- possible selection bias. What constitutes emergent surgical intervention- how many from BIG 1/BIG2 criteria excluded by this. Attrition: low risk Inpatient outcomes only Prognostic factor measurement: Mod risk Radiology report double checked by one person, only. Definition of neurological deterioration is defined differently as altered mental state and focal deficit and GCS less then 13 in different places. Outcome measures: Mod risk No routine follow up of all patients- must re-attend at same hospital to register Confounding Factors: Low risk Age affect outcome and size of bleed		
	coagulated or antiplatelets single ICH <5mm and no skull fracture single IPH BIG 2 (admit to hosp. not neurosurgeon) GCS 13-15, normal pupils and no focal neurological deficit Can be intoxicated Non-displaced Skull fracture Bleed 5-7mm le 2 intra cerebral					Statistical techniques: N/A		

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1 2 3 4						
5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21		bleeds 3-7mm Not anticoagulated or antiplatelets BIG 3 (repeat CT and admit under neurosurgeon HDU) GCS <13 or abnormal pupils or focal neurological deficit Taking anticoagulant or antiplatelets Multiple types of injury on CT Bleeds >7mm Displaced skull fractures Intubated patients				
22 23 24 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40 41	Joseph et al 2014 USA Study 2 validating the BIG criteria Identified Search Strategy	March 2012-Dec 2013 Level 1 Trauma centre Inclusion criteria BIG 1 patients: GCS 13-15, normal pupils and no focal neurological deficit Not intoxicated not anti- coagulated or anti- platelets single ICH <5mm and no skull fracture single IPH Excluded: Patients transferred from other hospital Intubated Patients undergoing emergent Prospective Cohort Study Aim To evaluate the established BIG 1 category for managing patients with traumatic brain injury	Patients remained in ED for observation for 6 hours. If no neurological deterioration-discharged. Repeated neurological assessment every 2 hours- if GCS<13, unequal pupils or focal neurological deficit-neurological deterioration Need for neurosurgical intervention. Need for Repeat CT due to neurological deterioration. Hospital or ICU admission. In-hospital mortality. 30 day readmission	Prospectively recorded: Age Sex Admission observations Neurological assessment of GCS, examination and pupils. Intoxication Anti-platelet or anti-coagulation Intubation LOC Initial CT findings by attending radiologist-confirmed by study radiologist	States 148 patients met criteria prospectively. 127/148 patients included and matched 127 patients with matched characteristics of demographics, medications and CT findings before implementation of BIG criteria. No patients underwent neurosurgery, had neurological deterioration or died, both of the 127 prospectively recruited and those matched retrospectively. Statistically significant reduction in hospital admissions, ICU admissions and repeat CT imaging in prospective cohort post implementation of BIG criteria. 0 30 day readmissions although 5 ED visits	Study Recruitment: mod risk States GCS13-15 and range presented as GCS13-15 but also excludes unexaminable patients and patients with altered mental state- appears cohort does not contain all GCS 14 and 13 patients. Not clear about how the cohort was prospectively recruited. Attrition: mod risk Disregards 21 of recruited cohort in analysis to match with retrospectively available patients. Prognostic factor measurement: Mod risk Reliability of case notes- may be incomplete The definitions of bleed size are subjective. Abnormal focal neurology is subjective and clinician dependent. CT scan re-reviewed by a single researcher-possible bias. Outcome measures: Mod risk Measures: no structured follow up of every patient. Patients could have been discharged and died in the community- study would have missed this. States over 50% admitted but that all discharged from the ED in the abstract.

Joseph et al 2015 USA Study 2:further validation of BIG criteria	Pre BIG TBI March 2011- Feb 2012 Post BIG July 2012-June 2013 Level 1 Trauma centre Inclusion criteria: All patients with blunt trauma mechanism and ICH/Skull fracture Excluded: Transfers Dead on arrival Needed immediate neurosurgery. Presents subgroup analysis of BIG 1 patients: GCS 13-15, normal pupils and no focal neurological deficit Not intoxicated	Prospective cohort study Compare outcomes in TBI before and after implementation of BIG criteria	Number of routine repeat CT head scans Neurosurgical consultations Progression of bleed on CT Neurosurgical intervention during hospital admission (craniotomy, craniectomy ICP monitoring) ICU admission 30 day readmission	Prospectively recorded: Age Sex Admission observations Neurological assessment of GCS, examination and pupils. Intoxication Anti-platelet or anti- coagulation Intubation LOC Initial CT findings by attending radiologist- confirmed by study radiologist	Pre BIG 87 BIG 1/415 0 neurosurgery 0 deaths 3 progression on CT 68 (78%)admitted 24 (27.5%) admitted ICU 76 (87.4%) neurosurg consultations 59 (67.8%) repeat CT Post Big 83 BIG 1/381 0 neurosurgery 0 deaths 1 progression on CT 42 admitted (50.6%) 6 ICU admission (7.2%) 7 (8.4%) neurosurg consultation 6 (7.2%) repeat CT Statistically significant (P<0.001 admission hospital, ICU, repeat CT imaging and neurosurgical consultation post introduction of BIG criteria)	Confounding Factors: Mod risk Age not part of BIG1 but could affect outcome and size of bleed Statistical techniques: N/A General Points: Small numbers of patients in this specific setup. Would support small CT findings low risk, but risk stratification very dependent on accuracy and consistency of radiology report. Study Recruitment: Low risk States all patients with TBI prospectively recorded on datanot cleat how patients identified and recruited. Emergent neurosurgical patients excluded- no definition given Attrition: low risk Outcomes only as inpatients or if re-present Prognostic factor measurement: Mod risk Ct are reviewed by a member of study group- the cut offs are slightly subjective on CT measurement Outcome measures: Mod risk Only measures as inpatient/re-presentation. Potential for discharge and deterioration. Confounding Factors: low risk Age Statistical techniques: Mod risk Presents data for all patients or BIG 1 patients- not all GCS13-15 patients
	GCS 13-15, normal pupils and no focal neurological deficit				admission hospital, ICU, repeat CT imaging and neurosurgical consultation post introduction of BIG	

single IPH				
	Mary		New Rochelle, NY 10801	

Supplementary Material 3: Table of Full Studies Retrieved and Excluded

No.	Study	Reason Excluded
1.	Anonymous et al ³¹	Unable to differentiate initial GCS13-15 patients
	(Full study revealed duplicate of	
	Corrigendum et al ¹⁴⁶)	
2.	Bajsarowicz et al ³⁴	Abstract only
3.	Bajsarowicz et al ³³	Unable to differentiate initial GCS13-15 patients
4.	Baldawa et al ³⁵	Letter about included study
5.	Basahm et al ³⁶	Unable to differentiate initial GCS13-15 patients
6.	Carlson et al ³⁸	Included paediatric patients and patients with no
		injuries identified by CT imaging
7.	Chen et al ³⁹	Uses lumbar puncture to diagnose brain injury
8.	Choudhry et al ⁴¹	Duplicate study ⁴⁰
9.	Flaherty et al ⁴³	Abstract only
10.	Gore et al ⁴⁴	Abstract only
11.	laccarino et al ⁴⁵	Unable to differentiate initial GCS13-15 patients
12.	Inamasu et al ⁴⁶	Unable to differentiate initial GCS13-15 patients
13.	Jacobs et al ⁴⁷	Includes patients no injuries on CT imaging
14.	Jiang et al ⁴⁸	Included patients of initial GCS<13
		Not clear if all GCS13-15 patients have injuries
		present on CT imaging.
15.	Jiang et al ⁴⁹	Included patients of initial GCS<13
		Not clear if all GCS13-15 patients have injuries
		present on CT imaging.
16.	Joseph et al ⁵⁰	Unable to differentiate initial GCS13-15 patients
17.	Joseph et al ⁵¹	Unable to differentiate initial GCS13-15 patients
18.	Joseph et al ⁵³	Unable to differentiate initial GCS13-15 patients
19.	Kim et al ⁵⁶	Unable to differentiate initial GCS13-15 patients
20.	Kreitzer et al ⁵⁸	Abstract only (full study included ⁸⁶)
21.	McCutcheon et al ⁶¹	Unable to differentiate initial GCS13-15 patients
22.	Nishijima et al ⁶⁴	Abstract only and associated paper included
	The injuries of all	patients of initial GCS<13
23.	Nishijima et al ⁶⁷	Unable to differentiate initial GCS13-15 patients
24.	Nishijima et al ⁶⁸	Unable to differentiate initial GCS13-15 patients
25.	Penn et al ⁷⁰	Abstract only (full study included ³⁷)
26.	Rubino et al ⁷²	Outpatient Setting
27.	Orringer et al ⁷⁹	Unable to differentiate initial GCS13-15 patients
28.	Yuan et al ⁸⁰	Unable to differentiate initial GCS13-15 patients
29.	Zare et al ⁸¹	Includes paediatric population
30.	Zhao et al ⁸²	Not clear about inclusion criteria and definition of
30.	21140 Ct 41	non-operative-no response from authors when
		contacted.
31.	Park et al ⁸³	Unable to differentiate initial GCS13-15 patients
32.	Schuster et al ⁸⁴	Unable to differentiate initial GCS13-15 patients Unable to differentiate initial GCS13-15 patients
33.	Smith et al ⁸⁵	Unable to differentiate initial GCS13-15 patients Unable to differentiate initial GCS13-15 patients
34.	Choudhry et al ⁸⁸	Abstract only (full paper included ⁴⁰)
35.	Tong et al ¹⁴⁷	Unable to differentiate initial GCS13-15 patients
	Yadav et al ⁹¹	·
36.	rauav et al	Unable to differentiate initial GCS13-15 patients and
		included children

37.	Cohen et al ⁹²	Includes patients with no injury on initial CT
38.	Stein et al ¹⁰⁵	Theoretical study-no data
39.	Borovich et al ¹¹⁰	Case reports
40.	Knuckey et al ¹¹¹	Pre-1996
41.	Chen et al ¹¹²	Pre-1996
42.	Mertol et al ¹¹³	Case reports pre-1996
43.	Brown et al ¹¹⁵	Unable to differentiate initial GCS13-15 patients
44.	Fainardi et al ¹¹⁷	Unable to differentiate initial GCS13-15 patients
45.	Karasu et al ¹¹⁸	Unable to differentiate initial GCS13-15 patients and
		includes children
46.	Türedi et al ¹²⁰	Includes patients with no injury on initial CT
47.	Connon et al ¹²¹	Unable to differentiate initial GCS13-15 patients
48.	Chang et al ¹⁴⁸	Unable to differentiate initial GCS13-15 patients
49.	Chao et al ¹²³	Unable to differentiate initial GCS13-15 patients
50.	Sullivan et al ¹²⁴	Unable to differentiate initial GCS13-15 patients
51.	Innocenti et al ¹²⁶	Includes patients with no injury on initial CT
52.	Muszynski et al ¹²⁷	Includes Children
53.	Patel et al ¹²⁸	Unable to differentiate initial GCS13-15 patients
54.	Lingsma et al ¹²⁹	Includes patients with no injury on initial CT
55.	Wong et al ¹³¹	Case studies and pre-1996
56.	Offner et al ¹³²	Unable to differentiate initial GCS13-15 patients
57.	Wong et al ¹³³	Duplicate of 55
58.	Bhau et al ¹³⁴	Unable to differentiate initial GCS13-15 patients
59.	Chen et al ³⁹	Includes Children and patients without CT identified
		injuries
60.	Gaetani et al ¹³⁵	Unable to differentiate initial GCS13-15 patients
61.	Greene et al ¹³⁶	Unable to differentiate initial GCS13-15 patients
62.	Son et al ¹³⁷	Unable to differentiate initial GCS13-15 patients
63.	Pradeep et al ¹³⁸	Unable to differentiate initial GCS13-15 patients
64.	Alahmadi et al ¹⁴⁹	Unable to differentiate initial GCS13-15 patients
65.	Chieregato et al ¹¹⁶	Includes Children
66.	Kehoe et al ⁹⁵	Unable to differentiate initial GCS13-15 patients
67.	Lesko et al ⁹⁶	Unable to differentiate initial GCS13-15 patients
68.	Lawrence et al ⁹⁴	Includes Children
69.	Roka et al 2008 ¹¹⁹	Includes Children

Supplementary Material 4: Characteristics of included studies

No.	Study	Туре	Size	Outcomes	Estimate of Outcome of interest	Univariate of analysis of any Prognostic factor	Multivariable Model of several prognostic factors
1	Sifri et al 2006 ⁷⁵	Prospective Cohort	130	Death Neurosurgery Progression CT	√	√	
2	Brown et al 2007 ¹¹⁴	Prospective Cohort	142	Death Deterioration Neurosurgery Progression CT	√		
3	Fabbri et al 2008 ¹³⁹	Prospective Cohort	723	Death Neurosurgery	✓	✓	
4	AbdelFattah et al 2012 ²⁸	Prospective Cohort	145	Death Deterioration Progression CT	√		
5	Sharifuddin et al 2012 ⁷⁴	Prospective Cohort	279	Death Deterioration Neurosurgery Progression CT	√	√	✓
6	Ding et al 2012 ⁹⁰	Prospective Trial	32	Neurosurgery Progression CT	√		
7	Nishijima et al 2014 ⁶⁶	Prospective Cohort	600	Deterioration Neurosurgery	✓	✓	✓
8	Sifri et al 2004 ¹⁰²	Retrospective Cohort	202	Death Deterioration Neurosurgery Progression CT	0		
9	Velmahos et al 2006 ⁷⁷	Retrospective Cohort	154	Deterioration Neurosurgery Progression CT	√	1	√
10	Huynh et al 2006 ⁹⁷	Retrospective Cohort	56	Deterioration Neurosurgery Progression CT	✓	7	
11	Bee et al 2009 ⁹⁹	Retrospective Cohort	207	Death Neurosurgery	✓	√	
12	Klein et al 2010 ⁵⁷	Retrospective Cohort	323	Death Neurosurgery	✓		
13	Schaller et al 2010 ⁸	Retrospective Cohort	110	Death Deterioration Neurosurgery	✓		
14	Nasir et al 2011 ¹⁰⁶	Retrospective Cross sectional	275	Neurosurgery Progression CT	✓		
15	Sifri et al 2011 ¹²⁵	Retrospective Cohort	107	Deterioration Neurosurgery Progression CT	√		

			,		,		1
16	Levy et al 2011 ⁵⁹	Retrospective Cohort SAH only	117	Death Neurosurgery Progression CT	√		
17	Washington et al 2012 ⁷⁸	Retrospective Cohort	321	Deterioration Neurosurgery Progression CT	✓	✓	✓
18	Homnick et al 2012 ¹⁰⁴	Retrospective Cohort	341	Death Deterioration Neurosurgery Progression CT	√		
19	Nayak et al 2013 ⁶²	Retrospective Cohort	321	Death Neurosurgery Progression CT	√		
20	Borczuk et al 2013 ³⁷	Retrospective Cohort	404	Deterioration Neurosurgery	√	✓	✓
21	Almenawer et al 2013 ¹⁸	Retrospective Cohort study and meta- analysis	445	Neurosurgery Progression CT	✓		
22	Joseph et al 2013 ⁵²	Retrospective Cohort	270	Death Neurosurgery	✓		
23	Thorston et al 2012 ⁶	Retrospective Cohort	360	Neurosurgery Progression CT	✓	✓	√
24	Choudhry et al 2013 ⁴¹	Retrospective Cohort	757	Death Deterioration Progression CT	√	√	√
25	Deepika et al 2013 ⁴²	Retrospective Cohort SAH only	34	Unable to extract			
26	Fabbri et al 2013 ⁸⁷	Retrospective Cohort	1123	Progression CT	1	✓	
27	Boris et al 2013 ¹⁰⁷	Retrospective Cohort	68	Deterioration Neurosurgery Progression CT	1		
28	Thomas et al 2010 ⁷	Retrospective Cohort	457	Deterioration Neurosurgery	✓		
29	Nishijima et al 2013 ⁶³	Retrospective Cohort	1412	Deterioration Neurosurgery	✓		
30	Quigley et al 2013 ⁷¹	Retrospective Cohort SAH only	478	Neurosurgery Progression CT	✓		4
31	Levy et al 2014 ⁶⁰	Retrospective Cohort	76	Deterioration Neurosurgery	√		
32	Overton et al 2014 ⁶⁹	Retrospective Cohort	171	Deterioration	√		1
33	Phelan et al 2014 ¹⁰³	Retrospective Cohort SAH only	77	Death Deterioration Neurosurgery Progression CT	1		
34	Kreitzer et al 2014 ⁸⁶	Retrospective Cohort	323	Death Neurosurgery	✓		
35	Kim et al 2014 ⁵⁵	Retrospective Cohort Subdurals	98	Neurosurgery Progression CT	✓	✓	√

		only		1	1			7
36	Sweeney et	Retrospective	50493	Neurosurgery	√	√	✓	
	al 2015 ⁹⁸	Cohort		0 ,	V	•	*	
37	Nishijima et al 2015 ⁶⁵	Retrospective Cohort	151	Deterioration	✓			
38	Darby et al 2015 ¹³⁰	Retrospective	369	Death	✓		✓	
	2015	Cohort		Neurosurgery Progression CT				
39	Beynon et al 2015 ⁹³	Retrospective Cohort	70	Death Neurosurgery	✓			-
40	Joseph et al 2015 ⁵⁴	Retrospective Cohort	876	Neurosurgery Progression CT	✓	√	√	
41	Ditty et al 2015 ³²	Retrospective Cohort SAH/ICB only	500	Death Neurosurgery Progression CT	✓			
42	Anandalwar et al 2016 ³⁰	Retrospective Cohort	142	Deterioration Neurosurgery	✓			
43	Bardes et al 2016 ¹⁰¹	Retrospective Cohort	389	Death Deterioration Neurosurgery Progression CT	√	√	√	
44	Shih et al 2016 ¹⁰⁰	Retrospective Cohort	340	Deterioration Neurosurgery Progression CT	✓	√	1	
45	Schwed et al 2016 ⁷³	Retrospective Cohort	201	Deterioration Neurosurgery	✓	√	√	
46	Sumritpradit et al 2016 ⁷⁶	Retrospective Cohort	98	Deterioration Neurosurgery Progression CT	√	✓		
47	Pruitt et al 2016 ¹⁰⁸	Retrospective Cohort	1053	Deterioration Neurosurgery	1			
48	Jospeph et al ^{9, 27, 109}					fication tool and a	combination of	-
49	al ^{9, 27, 109}	retrospective a	nd prospect	tive data followin	g its implement	ation.		
50								
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Supplementary Material 5: Table of Risk Factors Assessed

Risk Factor		Assessed	Univariate	Multivariate	Recursive
		Number of studies			partitioning
1 Age	Continuous	10 ^{6, 55, 69, 71,}	7 ^{6, 55, 73, 76, 77,}	4 ^{69, 71, 98, 130}	
		73, 76, 77, 98-100,	99, 100, 130		
		130			
	≥65	6 ^{37, 54, 66, 74, 77,}	6 ^{37, 54, 66, 74,}	3 ^{54, 74, 77}	1 ⁶⁶
		78	77, 78		
	≥60	1 ⁴¹	1 ⁴¹	1 ⁴¹	
· ·	≥55	2 ^{73, 101}	1 ¹⁰¹	1 ⁷³	1 ¹⁰¹
	≥50	1 ⁷⁶	1 ⁷⁶		
2 Gender		10 ^{6, 37, 54, 55,}	9 ^{6, 37, 54, 55, 74,}	2 ^{54, 69}	
		69, 74, 76, 77, 98,	76, 77, 98, 100		
		100			
3 Initial GCS	<15	7 ^{37, 41, 66, 73, 74,}	6 ^{37, 41, 66, 73,}	4 ^{37, 73, 74, 77}	2 ^{66, 101}
		77, 101	74, 101		
	GCS	7 ^{6, 55, 69, 73, 77,}	4 ^{6, 55, 73, 77,}	2 ^{69, 98}	
		98, 100	100		
	GCS=14	1 ⁶		1 ⁶	
	GCS=13	1 ⁶		1 ⁶	
4 CT Findings	Midline shift	5 ^{6, 55, 66, 76, 100}	4 ^{6, 66, 76, 100}	4 ^{6, 55, 76, 100}	1 ⁶⁶
	CT/Mass effect				
	Marshall	2 ^{41, 73}	2 ^{41, 73}		
	Classification				
	SDH>10mm	1 ⁵⁴	1 ⁵⁴	1 ⁵⁴	
	EDH>10mm	1 ⁵⁴	1 ⁵⁴	1 ⁵⁴	
	ICH vol>10ml	1 ⁷⁸	1 ⁷⁸	1 ⁷⁸	
	Mean Vol	1 ⁵⁵	1 ⁵⁵	1 ⁵⁵	
	Maximal thickness	1 ⁵⁵		1 ⁵⁵	
	Volume ED	1 ¹⁰⁰	1 ¹⁰⁰	1 ¹⁰⁰	
	Volume SDH	1 ¹⁰⁰	1 ¹⁰⁰		
	Volume ICB	1 ¹⁰⁰	1 ¹⁰⁰		
5 Type of	Contusion	1 ^{37, 78}	1 ^{37, 78}		
isolated		_	-		
injury					
	SDH	3 ^{37, 73, 98}	2 ^{37, 73}	1 ⁹⁸	
	EDH	3 ^{37, 73, 98}	2 ^{37, 73}	1 ⁹⁸	
	SAH	3 ^{37, 73, 98}	2 ^{37, 73}	2 ^{73, 98}	
	Mixed	1 ^{73, 98}	1 ⁷³	1 ⁹⁸	
	ICB	1 ⁷³	1 ⁷³		
6 Presence of	Contusion	3 ^{37, 76}	3 ^{37, 76}		
(includes					, in the second
mixed					
injuries)		6 27 76 400	6 27 76 422	27	
	SDH	5 ^{6, 37, 76, 100,}	5 ^{6, 37, 76, 100,}	1 ³⁷	
		101	101		
	EDH	5 ^{6, 37, 76, 100,}	5 ^{6, 37, 76, 100,}		

		101	101		
		4 ^{6, 37, 76, 100,}	4 ^{6, 37, 76, 100,}		
	SAH	101	101		
				- 100	
	fracture	4 ^{6, 74, 76, 100}	4 ^{6, 74, 76, 100}	1 ¹⁰⁰	
	Displaced/depressed	2 ^{54, 66}	2 ^{54, 66}	1 ⁵⁴	
	fracture	-74 76	-74.76		
	Base of skull	2 ^{74, 76}	2 ^{74, 76}		
	fracture	- 100	- 100		
	pneumocranium	1 ¹⁰⁰	1 ¹⁰⁰		
	ICB	3 ^{6, 100, 101}	3 ^{6, 100, 101}		
	IVH	3 ^{6, 76, 100}	3 ^{6, 76, 100}		
	Diffuse Axonal Injury	1 ⁷⁶	1 ⁷⁶	74 77	
	2+ lesions	4 ^{6, 74, 77, 100}	4 ^{6, 74, 77, 100}	2 ^{74, 77}	
	3+ lesions	1 ⁶	1 ⁶		
7 Subdural	contusion	1 ⁵⁵	1 ⁵⁵	1 ⁵⁵	
with					
	SAH	1 ⁵⁵	1 ⁵⁵	1 ⁵⁵	
8 Non-isolated	head Injury	1 ⁶⁶	1 ⁶⁶		1 ⁶⁶
9 BP		7 ^{54, 73, 76, 77, 98-}	6 ^{54, 73, 76, 77,}	2 ^{73, 98}	
		100	99, 100		
10 Pre-admission	on Hypotension	1 ⁶⁶	1 ⁶⁶		
11 HR		4 ^{54, 73, 98, 99}	3 ^{54, 73, 99}	1 ⁹⁸	
12 RR		1 ⁹⁸	1 ⁹⁸		
13 Pre-injury Hy	/poxia	1 ⁶⁶	1 ⁶⁶		
14 Intoxication		2 ^{54, 55}	2 ^{54, 55}		
15 Coagulopath	y: including any anti-	6 ^{6, 41, 55, 77, 98,}	5 ^{6, 41, 55, 77,}	1 ⁹⁸	
coagulant use		100	100		
16 Warfarin Use	2	3 ^{37, 76, 101}	3 ^{37, 76, 101}		
20 Warfarin or	anti-platelet	2 ^{78, 100}	2 ^{78, 100}		
17 PT/INR		3 ^{6, 74, 101}	3 ^{6, 74, 101}		
18 aPPT		1 ^{6, 101}	2 ^{6, 101}	•	
19 Platelet cour	nt<100000	1 ⁵⁴	1 ⁵⁴	1 ⁵⁴	
20 Platelet cour	nt<50000	1 ⁵⁵	1 ⁵⁵		
21 Hb<10		1 ⁵⁴	1 ⁵⁴		
22 Hb		2 ^{74, 100}	2 ^{74, 100}		
23 WCC		1 ¹⁰⁰	1 ¹⁰⁰	1 ¹⁰⁰	
24 Aspirin		3 ^{37, 76, 101}	3 ^{37, 76, 101}		
25 Clopidogrel		3 ^{37, 76, 101}	3 ^{37, 76, 101}		
25 Any Anti-pla	telet	2 ^{55, 66, 87}	1 ^{55, 66}	1 ⁸⁷	
26 ISS		11 ^{6, 69, 71, 73,}	9 ^{6, 41, 73, 76, 77,}	7 ^{6, 69, 71, 73, 98,}	
		76, 77, 98-101, 130	99-101, 130	100, 130	
27 (H)AIS		5 ^{41, 73, 76, 99,}	5 ^{41, 73, 76, 99,}	1 ⁷³	U X
		101	101		
28 LOC		1 ⁵⁴	1 ⁵⁴	1 ⁵⁴	
29 Mechanism	of Injury	2 ^{54, 55}	2 ^{54, 55}		
(unqualified)	· ·				
30 Non-fall from	n standing	1 ⁶⁶	1 ⁶⁶		
31 Fall	<u> </u>	2 ^{37, 77}	2 ^{37, 77}		
32 Assault		1 ³⁷	1 ³⁷		
33 RTC		4 ^{37, 74, 76, 77}	4 ^{37, 74, 76, 77}		
33 KIC		-7	7		

24 Dodostnian Ct	ole	1 ³⁷	1 ³⁷			
34 Pedestrian Struck	LK	1 1 1 3 7	1 1 1 3 7			
35 Bicycle struck		1 1 54	1 1 1 1 1 1 1 1 1 1	1 ⁵⁴		
36 Lactate 37 Base deficit		1 1 54	1 1 1 1 1 1 1 1 1 1	1 1 1 1 1 1 1 1	+	
38 Comorbidities	HTN	3 ^{37, 76, 100}	3 ^{37, 76, 100}	1		
38 Comorbialties	Diabetes	2 ^{76, 100}	2 ^{76, 100}	+	+	
	Old CVA	2 ^{76, 100}	2 ^{76, 100}			
	IHD	2 ^{76, 100}	2 ^{76, 100}			
	Arrhythmia	1 ¹⁰⁰	1 ¹⁰⁰			
	Liver disease	1 ¹⁰⁰	1 1 1 1 1 1 1 1 1 1			
	CKD	1 ¹⁰⁰	1 ¹⁰⁰		+	
	AKI	1 ¹⁰⁰	1 ¹⁰⁰		+	
	Any high risk	1 ⁶⁶	1 ⁶⁶			
39 Smoking	/my mgm ion	1 ⁵⁵	1 ⁵⁵			
40 Time to first CT		2 ^{73, 76}	2 ^{73, 76}			
41 Statin Therapy		1 ¹⁰⁰	1 ¹⁰⁰			
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Supplementary Material 6: Forest plots of within study risk factors' effect on the risk of neurosurgery or clinical deterioration

Meta-analysis of effect of initial GCS=15 on Risk of Clinical Deterioration/Neurosurgery

	Initial GC	S=15	Initial GC	S<15		Odds Ratio	Odds Ratio			
Study or Subgroup	Events	Total	Events	Total	Weight	IV, Random, 95% CI		IV, Randor	m, 95% CI	
Bardes et al 2016	31	310	22	79	18.6%	0.29 [0.16, 0.53]				
Borczuck et al 2013	37	344	11	60	15.6%	0.54 [0.26, 1.12]		-	-	
Fabbri et al 2008 (Neurosurgery)	62	493	47	226	24.3%	0.55 [0.36, 0.83]		-		
Nishijima et al 2014	46	406	70	194	24.1%	0.23 [0.15, 0.35]		-		
Schwed et al 2016	66	129	57	72	17.3%	0.28 [0.14, 0.54]		-		
Total (95% CI)		1682		631	100.0%	0.35 [0.23, 0.52]		•		
Total events	242		207							
Heterogeneity: Tau2 = 0.13; Chi2 =	10.65, df = 4	4 (P = 0)	$.03$); $I^2 = 63$	2%			0.02	0.1 1	10	50
Test for overall effect: Z = 5.16 (P <	0.00001)						0.02	Initial GCS=15	10 Initial GCS<15	50

Meta-analysis effect of isolated Subarachnoid haemorrhage versus any other injury on Clinical Deterioration/Neurosurgery

	Inclotes									
	isolated	HAZ b	Any Other Injur	y Type		Odds Ratio		Odds Ratio)	
Study or Subgroup	Events	Total	Events	Total	Weight	IV, Random, 95% CI		IV, Random, 95	5% CI	
Borczuck et al 2013	1	76	47	328	14.2%	0.08 [0.01, 0.59]				
Pruitt et al 2016 (neurosurg.)	0	155	4	216	8.2%	0.15 [0.01, 2.84]	\leftarrow	•	-	
Schwed et al 2016	27	57	96	144	32.3%	0.45 [0.24, 0.84]		-		
Sweeney et al 2015 (neurosurg.)	197	13191	4315	37305	37.2%	0.12 [0.10, 0.13]		•		
Velmahos et al 2006 (neurosurg.)	0	33	4	121	8.2%	0.39 [0.02, 7.42]	_	•		
Total (95% CI)		13512		38114	100.0%	0.19 [0.07, 0.50]		•		
Total events	225		4466							
Heterogeneity: Tau2 = 0.63; Chi2 = 1	7.98, df=	4 (P = 0.	001); I² = 78%				0.01	01 1	10	100
Test for overall effect: $Z = 3.39$ (P = 0	0.0007)						0.01		other Injury	100

Meta-analysis effect of Isolated Extradural versus any other injury on Clinical Deterioration/Neurosurgery

	Isolated	EDH	Any other	Injury		Odds Ratio		Odds	Ratio		
Study or Subgroup	Events	Total	Events	Total	Weight	IV, Random, 95% CI		IV, Rando	m, 95% C	CI	
Borczuck et al 2013	0	1	42	378	0.3%	2.64 [0.11, 65.82]			÷		_
Pruitt et al 2016 (neurosurg.)	0	5	4	366	0.3%	7.32 [0.35, 153.20]				*	\longrightarrow
Schwed et al 2016	1	1	122	200	0.3%	1.92 [0.08, 47.79]					_
Sweeney et al 2015 (neurosurg.)	159	901	4315	49595	98.8%	2.25 [1.89, 2.68]					
Velmahos et al 2006 (neurosurg.)	0	7	4	144	0.3%	2.08 [0.10, 42.34]		-			_
Total (95% CI)		915		50683	100.0%	2.26 [1.90, 2.68]			•		
Total events	160		4487								
Heterogeneity: Tau2 = 0.00; Chi2 = 0	.60, df = 4	(P = 0.9)	$(6); I^2 = 0\%$				0.04			-10	400
Test for overall effect: $Z = 9.22$ (P < 0	0.00001)	020	2.2				0.01	0.1 Any Other Injury	Isolated	10 Extra-du	100 ral

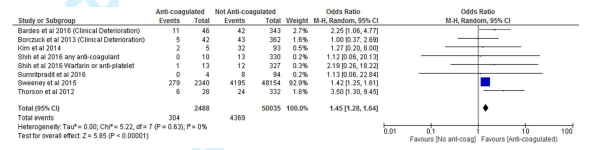
Meta-analysis Isolated subdural versus any other Injury on Clinical Deterioration/Neurosurgery



Meta-analysis Isolated contusion versus any other Injury on Clinical Deterioration/Neurosurgery

	Isolated Co	ontus	Any Other Injur	у Туре		Odds Ratio		Odds	Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	IV, Random, 95% CI		IV, Rando	m, 95% CI	
Borczuck et al 2013	2	33	46	271	1.3%	0.32 [0.07, 1.37]			-	
Pruitt et al 2016 (neurosurg.)	0	30	4	341	0.3%	1.23 [0.06, 23.38]			-	
Sweeney et al 2015 (neurosurg.)	139	5636	4335	44860	98.3%	0.24 [0.20, 0.28]				
Total (95% CI)		5699		45472	100.0%	0.24 [0.20, 0.28]		•		
Total events	141		4385							
Heterogeneity: Tau2 = 0.00; Chi2 = 1	1.34, df = 2 (F	P = 0.51	; I² = 0%				0.01	0.1	10	100
Test for overall effect: Z = 16.54 (P	< 0.00001)						0.01		Isolated Contusion	

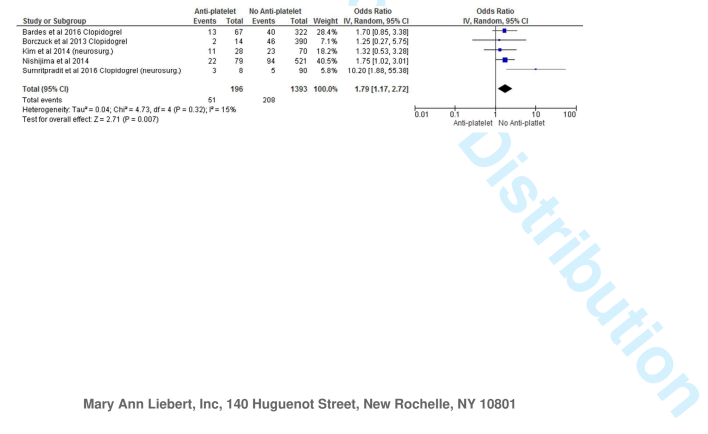
Meta-analysis of effect of coagulopathy use on Clinical Deterioration/Neurosurgery:



Meta-analysis effect of aspirin/anti-platelet use on Clinical Deterioration/Neurosurgery

	No Anti ni	No Anti-platelet Anti-platelet			Odds Ratio		Odds Ratio		
	NO Allu-pi	atelet	Allu-pia	telet		Ouus Rauo			
Study or Subgroup	Events	Total	Events	Total	Weight	IV, Random, 95% CI		IV, Random, 95% CI	
Bardes et al 2016 Aspirin	23	154	30	235	28.3%	1.20 [0.67, 2.16]		-	
Borczuck et al 2013 Aspirin	15	130	33	274	23.1%	0.95 [0.50, 1.82]		-	
Kim et al 2014 (neurosurg.)	11	28	23	70	11.8%	1.32 [0.53, 3.28]			
Nishijima et al 2014	22	79	94	521	33.3%	1.75 [1.02, 3.01]		-	
Sumritpradit et al 2016 Aspirin (neurosurg.)	2	23	6	75	3.5%	1.10 [0.21, 5.84]			
Total (95% CI)		414		1175	100.0%	1.30 [0.95, 1.78]		*	
Total events	73		186						
Heterogeneity: Tau2 = 0.00; Chi2 = 2.17, df = 4	(P = 0.70); I	$^{2} = 0\%$					0.01	01 1 10	100
Test for overall effect: Z = 1.66 (P = 0.10)							0.01	No Anti-platelet Anti-platelet	100

Meta-analysis effect of clopidogrel/anti-platelet use on Clinical Deterioration/ Neurosurgery



Supplementary Material 7: Pooled risk of clinical deterioration stratified by the injury type identified by initial CT imaging

