SCREENING LOGS FOR TBI TRIALS

<u>ScrTrial-TBI</u> = <u>Screening log for TBI trials</u>

2. CDE Definition This for	Trial_TBI = screening log for TBI trials data element aims to document if enrolment criteria
for ²	
	TBI trials were applied appropriate and monitors for
I tne	potential of selection bias.
	tom made data collection form
instrument for assessment	
4. Description of measure Mul-	tidimensional data collection form
5. Permissible values We form Pati with pote in the (e.g. those crite num crite pote whe app In t conwhite app cap aim exa pati enre Age - ≤ 1 - 18 - 21 - 31 - 41 - 51 - 61 - ≥7 Day Day add clock Reference in the conward of the conwhite state of the conward of	present two examples of possible formats. Both nats are anonymized without patient identification. ents are listed by consecutive number. All patients in TBI admitted/seen at the study facility who could entially meet the study enrolment criteria (e.g. seen ne ER; admitted to ICU) over a given time period g. per week, per month). Both patients enrolled and se not enrolled are listed. The relevant inclusion eria and exclusion criteria for not enrolling patients by inher of the in- and exclusion criteria (e.g. inclusion erion 2,3; exclusion criterion 5). With this format the ential for selection bias can be monitored, but either in- and exclusion criteria were applied ropriately can not be monitored. The second format basic data are documented, cerning the most relevant in- and exclusion criteria ch permit monitoring whether enrolment criteria were lied appropriately or not. Care has been taken to ture these data in a broad categorical format with the of eliminating any potential patient identifiers. The mple presented is aimed at a trial in more severe TBI ents, capturing information on most commonly used oliment criteria: 18 3-20 1-30 1-40 1-50 1-60 1-70

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8. Comments/Special instructions:

Care must be taken to avoid any risk of including patient identifiers in the documentation as for patients not enrolled informed consent will not be requested. The intent to collect screening data and the format used should be reported to the local IRB when submitting the study for IRB approval. From a study perspective it is preferred to transfer the screening log data to the trial coordinating center, but local regulations may require that these data are not transferred outside the study center.

9. Rationale/justification:

Screening logs form an important tool in randomized clinical trials and are essential to reporting trial results according to the consolidated standards of reporting trials (consort guidelines) and to assess the generalizibility of findings. Characteristics of patient populations within a multicenter trial may differ between centers and countries, for example due to aspects of local trauma organisation and consequently may introduce an element of selection bias outside the control of investigators. These observations will limit generalizibility of findings. Further, consistency of accuracy in screening log completion has been shown to be related to center performance in trials. The main reason for collecting screening log data in TBI trials however is to monitor for selection bias and to investigate whether enrolment criteria are being applied appropriately.

10. References:

Slieker FJA, Kompanje EJO, Murray GD, et al. Importance of screening logs in clinical trials for severe traumatic brain injury. Neurosurgery. 2008; 62(6):1321-8. Schulz KF, Altman DG, Moher D for the CONSORT group. CONSORT 2010 statement: updated guidelines for reporting parallel group randomized trials. BMJ 2010; 340:697-702. Kompanje EJ, Maas AI. Is the Glasgow Coma Scale score protected health information? The effect of new United States regulations (HIPAA) on completion of screening logs in emergency research trials. Intensive Care Med. Feb 2006; 32(2):313-4.

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