Outcome Prognostication of Acute Brain Injury using the Neurological Pupil Index (ORANGE) study: protocol for a prospective, observational, multicentre, international cohort study

ABSTRACT

Introduction. The pupillary examination is an important part of the neurological assessment, especially in the setting of acutely brain-injured patients, and pupillary abnormalities are associated with poor outcomes. Currently, the pupillary examination is based on a visual, subjective and frequently inaccurate estimation. The use of automated infrared pupillometry to measure the pupillary light reflex can precisely quantify subtle changes in pupillary functions. The study aimed to evaluate the association between abnormal pupillary function, assessed by the Neurological Pupil Index (NPI), and long-term outcomes in patients with acute brain injury (ABI).

Methods and analysis. The Outcome Prognostication of Acute Brain Injury using the Neurological Pupil Index study is a prospective, observational study including adult patients with ABI requiring admission at the intensive care unit. We aimed to recruit at least 420 patients including those suffering from traumatic brain injury or haemorrhagic strokes, over 12 months. The primary aim was to assess the relationship between NPI and 6-month mortality or poor neurological outcome, measured by the Extended Glasgow Outcome Score (GS-E; poor outcome=GOS-E 1–4). Supervised and unsupervised methods and latent class mixed models will be used to identify patterns of NPI trajectories and Cox and logistic model to evaluate their association with outcome.

Ethics and dissemination. The study has been approved by the institutional review board (Comitato Etico Brianza) on 16 July 2020. Approved protocol V.4.0 dated 10 March 2020. The results of this study will be published in peer-reviewed journals and presented at conferences.

Trial registration number. NCT04490005.

INTRODUCTION

Pupillary examination, including pupillary light reactivity (PLR), is a fundamental part of the clinical examination in patients suffering from acute brain injury (ABI), with both diagnostic and prognostic values.1 As an example, the oculomotor nerves might be compressed due to displacement of the brainstem, and clinicians have accepted fixed and dilated pupils as part of the ‘herniation’ syndrome.

Elevated intracranial pressure (ICP) may alter midbrain function and cause abnormalities in pupil size, symmetry and PLR.2–5 Sustained or newfound pupillary abnormalities are associated with a worse outcome,6 and indeed PLR is a robust validated predictor in several prognostic models, such as the Corticosteroid Randomization after Significant Head Injury and the International Mission for Prognosis and Analysis of Clinical Trials (IMPACT) scores.7 However, in current clinical practice, the pupillary examination is performed using a manual, hand-held light source (eg, pen torch), implying that the evaluation of pupillary size and reactivity remains essentially based on a visual qualitative assessment. This traditional approach has several limitations, such as limited precision (eg, due to small pupil size or specific administered treatments), significant intraobserver and interobserver variability, differences in ambient light exposure between measurements or the technique used to direct the stimulus (ie, intensity, proximity, duration and orientation of the light source).8–10
Quantitative, automated, infrared technology for pupillary examination has been used for years in ophthalmology and anaesthesiology research. Its interest in neurocritical care has progressively grown, in parallel with the advancements in device technology. In this regard, the use of the non-invasive NPi−200 pupillometer (Neuroptics, Laguna Hills, California, USA) allows the measurement of a series of dynamic pupillary variables which can be integrated into an algorithm to compute the Neurological Pupil Index (NPI). The NPI is calculated by the handheld device using a set of variables including size, latency, constriction velocity and dilation velocity. Each variable from an individual pupil measurement is compared against the mean of a reference distribution of healthy subjects, taking the difference and then standardising it by the corresponding SD. Finally, the sets of all healthy subjects, excluding variables (eg, ICP max, ICP 20 index: number of end hourly measures of ICP of >20 mm Hg divided by the total number of measurements, multiplied by 100) and interventions (eg, osmotherapy, therapy intensity level and neuroimaging). The GOS-E will be collected at ICU/hospital discharge and 6 months from ICU admission. The latter will be collected via telephone-structured interviews with patients and/or family members using a validated questionnaire. In case of death, details on the cause and date will be collected as well.

Methods and analysis

Study design and setting

This is a prospective, observational cohort study involving 13 centres worldwide that routinely use pupillometry. Recruitment will last 12 months, and patients will be followed up for 6 months.

Study population

Consecutive participants will be recruited at the participating centres according to the inclusion and exclusion criteria reported as follows.

Inclusion criteria

- Intensive care unit (ICU) admission after ABI, including TBI, SAH and ICH requiring intubation and ventilation for neurological reasons/deterioration.
- Age ≥ 18 years old.
- Pupillometry is available as a standard evaluation tool.

Exclusion criteria

- Facial trauma not allowing pupils’ evaluation.

Screening and data collection

All patients admitted to the participating ICUs after ABI will be screened daily and entered into a screening log (online supplemental appendix, Screening Log–Registry). Each ICU will recruit consecutive eligible patients and collect data for each included patient daily in an expanded electronic CRF developed in REDCap (online supplemental appendix, eCRF).

Both common data elements and aetiology-specific data will be recorded. Demographic characteristics and medical history information will be extracted from patients’ medical records including gender, age, comorbidities, diagnosis, timeline and clinical presentation of ABI. All NPI and ICP data, as well as additional neuromonitoring and neuroimaging data, will be extracted from patients’ medical records too and documented in the eCRF. The patients admitted to the units will be screened by research staff, and the pupillometry evaluation, part of the clinical practice in all the centres, will be performed by trained staff. The two eyes specific NPI and the matched ICP will be collected every 4 hours from admission up to day 7. Data collected will also include additional ICP-derived variables (eg, ICP max, ICP 20 index: number of end hourly measures of ICP of >20 mm Hg divided by the total number of measurements, multiplied by 100) and interventions (eg, osmotherapy, therapy intensity level and neuroimaging). The GOS-E will be collected at ICU/hospital discharge and 6 months from ICU admission.

Sample size and statistical analysis

As no data on NPI trajectory in time and its association with outcome is available for sample size calculation, we referred to pupil reactivity as a proxy of NPI behaviour and to results of a study recruiting patients with similar characteristics (Intracranial Pressure monitoring in the ICU: An International Prospective Observational Study on Intracranial Pressure in Intensive Care, trial registration number: NCT03257904). Assuming a 6-month mortality of 53% in patients with one or both unreactive pupils and 29% in patients with both reactive pupils, a sample of 420
patients achieves 94% power to detect a HR of nearly 2 at a 0.05 significance level. As far as recruitment, each of the 13 centres involved in the study is expected to include nearly 35 patients in a recruitment period of 1 year, and this is in line with their potentiality of recruitment for the three pathologies that is of at least 80 patients per year.

Qualitative variables will be summarised by counts and percentages, while quantitative characteristics will be summarised by quartiles or mean and SD, as appropriate. Supervised and unsupervised methods (eg, pattern recognition and cluster trajectory analyses) will explore the possibility to identify patterns of NPi trajectories associated with different prognosis on the individual NPi longitudinal measurements. NPi trends will be also modelled by longitudinal mixed models using splines and latent class mixed models.

A Cox and a logistic model will be applied to evaluate the association between the NPi process and the 6-month mortality and neurological recovery (GOS-E≤4 vs GOS-E>4) at 6 months, respectively. This will be done considering NPI in categories that identify different potential patterns in NPi longitudinal profiles or using summary measures that have been already introduced in this context, such as the percentage of NPi≤3 observed in the time course or the area under the trajectory in time. This analysis will be done both overall, on a multivariable model that will explore the interaction with the different pathologies, and on the three specific pathologies. A longitudinal linear model that will explore the interaction with the time. This analysis will be done overall, on a multivariable model that will explore the interaction with the different pathologies, and on the three specific pathologies that is of at least 80 participants per year.


data confidentiality

The main limitation of this study is its observational nature, which makes it impossible to draw causal inferences reliably. We try to overcome this limitation with a preplanned statistical plan and a rigorous analysis of the findings.

ETHICS AND DISSEMINATION

This study will be conducted in compliance with the protocol V.4.0 dated 10 March 2020 approved by the ethics committee ‘Brianza’ at the ASST-Monza (approval date: 16 July 2020). Each National Coordinator will notify the relevant ethics committee, in compliance with the local legislation and rules. The approval of the protocol (if required by local authorities) must be obtained before any participant is enrolled. Any amendment to the protocol will require review and approval by the three chief investigators before the changes are implemented into the study. In case of patients not able to provide an informed consent at the time of study recruitment, each country will refer to the local/national law on the matter of lack of capacity. Generally, if patients will regain capacity at the follow-up, they will be asked to provide the informed consent for the acute data and follow-up or deny further research participation without any objection against use for research of data collected during the acute phase or deny further research participation and require the destruction of acute data collected. The study will be performed according to the Helsinki Declaration and the International Conference on Harmonisation for Good Clinical Practice.

The University of Milano–Bicocca has the property of all the data collected. The data reside at the University Milano–Bicocca as study sponsor; all procedures will comply with the EU regulation on data protection 2016/679 on the protection of natural persons regarding personal data processing and movement. Local site data will be co-owned by each participating centre, and they will be given access to local data for any scientific purpose on request. By entering data into the Outcome Prognostication of Acute Brain Injury using the Neurological Pupil Index (ORANGE) study database, each centre agrees that the chief can use these data for scientific purposes. Any requests for the use of the data set for subsequent studies will be made to the ORANGE study chief investigators. Any requests for the use of the data set for subsequent studies will be made to the ORANGE study chief investigators. A formal data monitoring committee is not needed since it is not an interventional controlled study. A dedicated staff from the University of Milano–Bicocca will monitor the data included in the eCRFs checking for inconsistencies.

Patient and public involvement

It was not appropriate or possible to involve patients or the public in the design, conduct, reporting or dissemination plans of our research.

Limitations

The main limitation of this study is its observational nature, which makes it impossible to draw causal inferences reliably. We try to overcome this limitation with a preplanned statistical plan and a rigorous analysis of the findings.

Data confidentiality

Participant confidentiality is strictly held in trust by the participating investigators and their staff. All medical or administrative staff with access to the data is subject to a duty of confidentiality and data protection. Therefore, the study protocol, documentation, data and all other information generated will be held in strict confidentiality agreement protocols. The study sponsor and representatives of local authorities may inspect all documents and records required to be maintained by the local institutions.
investigator for the participants in this study. The clinical study site will permit access to such records.

Study participant research data, which is for purposes of statistical analysis and scientific reporting, will be transmitted to the Data Manager and the Statisticians of the study. For this purpose, data will be deidentified at input into the eCRF by the local centres/investigators. Individual participants and their research data will be identified by a unique study identification number. The eCRF system used by clinical sites and by research staff will be secured.

Lack of capacity and delayed consent
Patients recruited in this study will not be able to provide informed consent at the time of recruitment. Consent procedures will follow local policies.

At follow-up, patients who have regained capacity will be asked to provide informed consent and will be given the possibility to
- Provide informed consent for the acute data and follow-up.
- Deny research participation and request destruction of acute data collected.

Medical care related to the study
The medical care of the participant in the study is performed as per the local standard of care, without any deviation from clinical protocols. All the procedures follow the latest recommendations for ABI.

Premature termination or suspension of study
This study may be suspended or prematurely terminated for reasonable cause agreed by the investigators. Written notification, documenting the reason for study suspension or termination will be provided by the suspending or terminating party. If the study is prematurely terminated or suspended, the local principal investigator will promptly inform the ethics committees or other local authorities according to local legislation and will provide the reasons for the termination or suspension. Circumstances that may warrant termination could be recruitment that will be prolonged for >5 years or insufficient compliance with any deviation from clinical protocols. All the procedures follow the latest recommendations for ABI.

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Contributors MO, FT and GC conceived and designed the study. SG and PR conceived the statistical plan of the study. SG and GC drafted the manuscript. All authors revised the manuscript and gave the final approval of the version to be submitted.

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