Comparison of outcomes of neurosurgical operations performed before and during the COVID-19 pandemic: a matched cohort study

Emma Toman 1,2, Wai Cheong Soon3, Gopiga Thanabalasundaram2, Daniel Burns4,5, Vladimir Petrik2, Colin Watts2,8, Victoria Wykes2,7, Anwen White2

ABSTRACT

Objective To determine how the first wave of the COVID-19 pandemic affected outcomes for all operatively managed neurosurgical patients, not only those positive for SARS-CoV-2.

Design Matched cohort (pairwise method).

Setting A single tertiary neurosurgical referral centre at a large UK Major Trauma Centre.

Participants During the first COVID-19 wave, 231 neurosurgical cases were performed. These cases were matched to cases from 2019. Cases were matched for age (±10 years), primary pathology and surgical procedure. Cases were excluded from analysis if either the age could not be matched to within 10 years, or the primary pathology or procedure was too unique. After exclusions, 191 cases were included in final analysis.

Outcome measures Primary outcomes were 30-day mortality and postoperative pulmonary complications. Secondary outcomes included Glasgow Outcome Score (GOS) on discharge, length of stay (LoS), operative and anaesthetic times and grade of primary surgeon. An exploratory outcome was the SARS-CoV-2 status of patients.

Results There was no significant difference between the pandemic and matched cohorts in 30-day mortality, pulmonary complications, discharge GOS, LoS, operative or anaesthetic times. There was a significant difference in the variation of grade of primary surgeon. Only 2.2% (n=5) of patients had a SARS-CoV-2 positive swab.

Conclusion During the first UK wave of the COVID-19 pandemic, the mortality, morbidity and functional outcomes of operatively managed neurosurgical patients at University Hospitals Birmingham were not significantly affected compared with normal practice. The grade of primary surgeon was significantly more senior and adds to the growing body of evidence that demonstrates how the pandemic has negatively impacted UK surgical training. Mixing COVID-19 positive, unknown and negative cases did not significantly impact on outcomes and indicates that further research is required to support the implementation of evidence-based surgical pathways, such as COVID-light sites, throughout the next stage of the pandemic.

Strengths and limitations of this study

- This is the first study to individually match neurosurgical cases performed during the first pandemic wave to cases performed under 'pre-pandemic' conditions.
- Primary outcomes have been designed to be comparable with prior international surgical cohort studies.
- The numbers of patients analysed are small and from a single unit only.
- Not all neurosurgical cases performed during the first pandemic phase could be matched to pre-pandemic cases.
- Given the evolving understanding of the virus during the first wave, there was no universal swabbing protocol that could be applied to all cases in the pandemic cohort.

INTRODUCTION

Background and rationale

The first cases of COVID-19 were diagnosed in the UK at the end of January 2020.1 Hospitals began to prepare for the anticipated influx of severely unwell patients and surgical departments were advised to adapt their protocols, create additional capacity and protect their vulnerable patient groups. By mid-March, neurosurgical elective procedures were cancelled at the University Hospitals Birmingham (UHB) Foundation Trust. Exceptions were cancer, life-limiting cases or cases that ran a high risk of significant deterioration within 2 weeks if left untreated.2 Neuro-theatre capacity was reduced to one theatre running 24 hours a day and an additional ad hoc emergency daytime list where neuro-anaesthetic support allowed. These changes were necessary as UHB became the Trust ‘worst hit’ by COVID-193 with the Office for National Statistics confirming that Birmingham had the highest peak excess mortality of any major British city at 249.7%.4
While there have been several studies published on the restructuring of neurosurgery services and impact of COVID-19 on workload, there is currently limited evidence as to the impact of COVID-19 on the outcomes that such changes have had for neurosurgical patients. In May 2020, the COVIDSurg collaborative determined that postoperative pulmonary complications occur in 50.0% of neurosurgical patients with perioperative SARS-CoV-2 infection and 30-day mortality was 18.4%. A US group reported the incidence of COVID-19 in neurosurgical patients to be 5.4% with a higher rate of complications in positive patients compared with negative patients. While it is vital to gather evidence on how SARS-CoV-2 affects the outcome of infected neurosurgical patients, the majority of patients (almost 95%) remain uninfected. There is currently no evidence to describe how the pandemic has affected the outcome of the entire neurosurgical cohort when compared with ‘normal’ neurosurgical practice.

This study compares the outcomes of patients undergoing neurosurgical procedures during the height of the pandemic against a matched cohort from prior to the COVID-19 outbreak.

**Objectives**

1. To compare the rates of 30-day mortality and pulmonary complications.
2. To compare functional outcome and length of hospital stay (LoS).
3. To compare anaesthetic and operative factors.
4. To determine rates of patients screened for COVID-19, incidence of confirmed cases and their clinical course.

**Materials and Methods**

**Study design**

The study was undertaken at the Queen Elizabeth Hospital Birmingham (QEHB) as a single-centre, matched cohort study. The QEHB is the largest hospital within the UHB Foundation Trust, is one of three regional adult neurosurgical units in the West Midlands, and is responsible for a mixed urban and rural population.

**Participants**

Operations performed during the initial stage of the pandemic (19 March 2020–1 June 2020) were matched to pre-pandemic cases (11 March 2019–11 September 2019). The pandemic time period was chosen as this was the interval in which elective neurosurgery cases at UHB were cancelled and will be referred to as the ‘initial pandemic phase’ from here onwards. Cases were matched in a pairwise fashion for age at time of procedure (±10 years), primary pathology and procedure. Primary pathology included severity where appropriate; for example, poor grade subarachnoid haemorrhage (SAH) was matched with another SAH of equivalent poor grade. Procedure was also matched to site where significant; for example a left-sided frontal craniotomy was matched with another left-sided frontal craniotomy rather than a right-sided. Ward-based procedures such as lumbar drain and intracranial pressure bolt insertion were not included. Cases were deemed unmatchable and were excluded from analysis if either the age could not be matched to within 10 years, or the primary pathology or procedure was too unique.

**Procedures**

Cases were identified by performing a search of the Galaxy electronic operating theatre system for all operations carried out under a neurosurgical consultant within the chosen time periods. Galaxy and the Prescribing Information and Communications System were then used to collate relevant variables.

Demographic data included age and gender. Operative data included name of procedure, grade of primary surgeon and operative time points. Grade of surgeon was defined as junior Senior Registrar (SpR) (ST3–5 or equivalent), senior SpR (ST5–8 or equivalent), fellow or consultant. Preoperative anaesthetic time was defined as time ‘into anaesthetic room’ to ‘knife to skin’, operative time was defined as ‘knife to skin’ to ‘skin closure’ and postoperative anaesthetic time was defined as ‘skin closure’ to time ‘into theatre recovery’.

Clinical data included LoS in days, discharge Glasgow Outcome Score (GOS), 30-day mortality, postoperative pulmonary complications and COVID-19 swab status. Swabs were deemed to be positive if SARS-CoV-2 RNA was detected by the standard RT-PCR laboratory test.

COVID-19 swabbing protocols were constantly evolving during this early phase of the pandemic as scientific knowledge and testing increased. For this reason, there is no one single consistent swabbing protocol for the time period interrogated in this study and as such reflects ‘real-life’ practice.

**Outcomes**

**Primary outcomes**

1. 30-day mortality.
2. Postoperative pulmonary complication.

Primary outcomes were designed to be comparable with the COVIDSurg study. For 30-day mortality, day of surgery was treated as day 0. Postoperative pulmonary complication was defined as pneumonia, acute respiratory distress syndrome or unexpected postoperative ventilation (any episode of non-invasive ventilation, invasive ventilation or extracorporeal membrane oxygenation after initial extubation after surgery; or patient could not be extubated as planned after surgery).

**Secondary outcomes**

1. Discharge GOS.
2. LoS in hospital (days).
3. Grade of primary surgeon.
4. Preoperative and postoperative anaesthetic time (minutes).
5. Operative time (minutes).

The GOS is an objective, functional outcome score from 1 to 5. Favourable GOS was defined as a score of
4 (moderate disability) or 5 (good recovery). Unfavourable GOS was defined as a score of 1 (dead), 2 (vegetative state) or 3 (severe disability).

Exploratory outcome
COVID-19 status both preoperatively and postoperatively and the clinical course of positive patients.

Statistical methods
The study was conducted and reported in line with Strengthening the Reporting of Observational Studies in Epidemiology guidelines for cohort studies. Continuous data were tested for normal distribution prior to analysis. Parametric data are presented as mean with 95% CI, and the unpaired t-test used to detect differences between groups. Non-parametric data are presented as median with IQR, and Mann-Whitney used to detect differences between groups. The X² test was used for categorical data. Missing data points were removed from final analyses. Clinical significance was defined as p<0.05.

Analyses were performed using GraphPad Prism software V.8.4.3 for Windows.

Patient and public involvement
While there has been significant patient and public involvement (PPI) with regard to patient pathways during the pandemic, there was no specific PPI consultation for this study. This was due to the acutely evolving nature of the pandemic, time limits and manpower restraints.

RESULTS
Overall, 231 cases were performed during the initial pandemic phase. Of these 17.3% (n=40) were unmatchable and so were removed from the final analysis of primary and secondary outcomes. A final cohort of 191 cases from the initial pandemic phase were successfully matched to 191 cases from 2019.

The mean age for patients in the pandemic group was 55 years and the matched group, 54 years. There was no statistically significant difference between the age of the two cohorts (p=0.795 (95% CI 3.7 to 2.8)). With regard to gender, 42.4% (n=81) and 47.6% (n=91) were women in the pandemic and matched cohorts, respectively. There was no statistically significant difference in gender between the two groups (p=0.304).

In terms of theatre location, during the pandemic 97.4% (n=186) of cases were performed in a dedicated emergency neurosurgical theatre and 2.6% (n=5) performed in a shared-specialties emergency theatre. Cases from the matched cohort were performed across six separate neurosurgery theatres.

Primary outcomes
There was no statistically significant difference in either 30-day mortality or rates of postoperative pulmonary complication between the pandemic and matched cohorts (see table 1).

<table>
<thead>
<tr>
<th>GOS</th>
<th>Pandemic (n=191)</th>
<th>Matched (n=191)</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>30-day mortality</td>
<td>3.7 (7)</td>
<td>3.1 (6)</td>
<td>0.778</td>
</tr>
<tr>
<td>Postoperative complications</td>
<td>10.5 (20)</td>
<td>10.5 (20)</td>
<td>1.000</td>
</tr>
<tr>
<td>Favourable outcome (GOS 4–5)</td>
<td>89.9 (170)*</td>
<td>92.1 (176)</td>
<td>0.366</td>
</tr>
<tr>
<td>Unfavourable outcome (GOS 1–3)</td>
<td>10.4 (20)*</td>
<td>7.9 (15)</td>
<td></td>
</tr>
</tbody>
</table>

Data are presented as % (n).
*One patient still an inpatient at time of analysis and was not included in analysis.
GOS, Glasgow Outcome Score.

Secondary outcomes
The majority of patients in both groups were discharged with a favourable GOS: 89.9% (n=170) and 92.1% (n=176) for pandemic and matched cohorts, respectively. There was no statistically significant difference in discharge GOS between the pandemic and matched cohorts (see table 1 and figure 1).

Median LoS was 4.5 days (IQR 2.0–10.3) for the pandemic cohort and 6.0 days (IQR 2.0–18.0) for the matched cohort. There was no statistically significant difference in LoS between pandemic and matched cohorts (p=0.25).

There was no statistically significant difference in preoperative anaesthetic time, operative time or postoperative anaesthetic time between pandemic and matched cohorts (see table 2). However, there was a significant difference in the variation of grade of primary surgeon between the two cohorts. Although the number of cases that consultants performed remained the same, there was a trend towards more cases being performed by senior SpRs and fellows rather than junior SpRs during the pandemic (see table 2).

COVID-19
For this section, all 231 participants for the pandemic period were analysed, including the unmatched cases. With regard to COVID-19 testing, 60.1% (n=139) were swabbed preoperatively and 37.2% (n=86) were swabbed postoperatively; some were swabbed both preoperatively and postoperatively. Twenty-seven per cent (n=63) of cases had neither preoperative nor postoperative swabs. Five patients in total tested positive for COVID-19 (see table 3 for case details). Four patients with a positive COVID-19 swab were operated on in the same theatre.

DISCUSSION
This study has demonstrated that the mortality and morbidity of neurosurgical patients undergoing operative management at UHB was not affected by the initial
COVID-19 outbreak. The patient selection, preoperative, perioperative and postoperative protocols at UHB for neurosurgery during the pandemic have ensured the short-term outcome for patients has not significantly changed compared with normal practice. During the pandemic, all neurosurgical consultants and registrars who were part of the on-call rota underwent fit testing for FFP3 masks. When operating on patients with unknown or pending COVID-19 status, all theatre staff wore FFP3 mask, protective face-shield/visors or eye-goggles perioperatively. When operating on patients with negative COVID-19 result, there was no requirement to wear an FFP3 mask. However, during intubation and extubation in all cases, the anaesthetic team had full personal protective equipment (PPE) due to high risk of exposure to aerosol. In all transsphenoidal surgery cases, all theatre staff wore PPE perioperatively. Such protocols and practices should therefore be reinstated in future pandemic waves.

Discussion surrounding how these results may impact on practice during this recovery phase of the pandemic is more complex. Current guidelines from both the National Institute of Health and Care Excellence (NICE) and the Royal College of Surgeons England (RCS) recommend strict social distancing, swabbing regimes and the separation of COVID-19 positive and negative patients.\textsuperscript{13, 14} The RCS goes further by stating that every hospital should have access to ‘COVID-light sites’ that ‘might be created at independent hospitals, within designated areas in National Health Service (NHS) hospitals or for an entire hospital functioning as an NHS network hub’.\textsuperscript{15} A recent survey however demonstrated that nationally 26% of hospitals have not been able to access COVID-light facilities.\textsuperscript{16}

To comply with NICE and RCS guidance, UHB designed ‘hot’ and ‘cold’ surgical pathways dependant on preoperative COVID-19 status. This new strategy was implemented on 1 June 2020 when elective operating restarted. Designated hot and cold pathways will increase public confidence that the risk of transmission is being reduced as much as is physically possible. The downside to having separate pathways is the huge logistical organisation they require, the associated financial impact of running additional swabbing, admission and surgical pathways, and a greatly reduced theatre capacity at a time where waiting lists are the highest they have been for over a decade.\textsuperscript{17}

This study is the first to demonstrate that mixing COVID-19 positive, unknown and negative cases in a single large centre has not impacted patient outcomes during the initial phase of the pandemic. Our data suggest that it is not necessary to provide ‘COVID-light centres’, and as long as rigorous protocols are adhered to, neurosurgical care can proceed with acceptable healthcare outcomes comparable with pre-pandemic practice. While this small study does not definitively prove that mixing of patients is safe on a national scale, it certainly highlights that more investigation is required to support or refute these findings. The authors therefore strongly recommend that a larger, multicentre study should be undertaken during the next stage of the pandemic to assess whether outcome and infection rate is truly improved by such a costly and time-intensive approach to elective surgery.

![Figure 1](http://bmjopen.bmj.com/)

**Figure 1** Comparison of discharge GOS. GOS, Glasgow Outcome Score.

<table>
<thead>
<tr>
<th>Table 2</th>
<th>Comparison of procedure details</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Pandemic</td>
</tr>
<tr>
<td></td>
<td>n=191</td>
</tr>
<tr>
<td>Grade of primary surgeon</td>
<td></td>
</tr>
<tr>
<td>Junior SpR</td>
<td>5.2 (10)</td>
</tr>
<tr>
<td>Senior SpR</td>
<td>7.3 (14)</td>
</tr>
<tr>
<td>Fellow</td>
<td>9.4 (18)</td>
</tr>
<tr>
<td>Consultant</td>
<td>78.0 (149)</td>
</tr>
<tr>
<td>Preop anaesthetic time</td>
<td>58.5 (44.0–76.0)</td>
</tr>
<tr>
<td>Operative time</td>
<td>101.5 (57.0–150.5)</td>
</tr>
<tr>
<td>Postop anaesthetic time</td>
<td>19 (11.75–28.0)</td>
</tr>
</tbody>
</table>

Data are presented as % (n) for grade of primary surgeon and median (IQR) for anaesthetic and operative times. Units of time presented are minutes.

SpR, Senior Registrar.
It may seem intuitive that by increasing the amount of PPE anaesthetic and surgical teams must wear, the anaesthetic and operative time should also increase. Unpublished data from UHB analysing neuro-oncology operations during the first month of initial phase of the pandemic suggested that the average length of operation was significantly increased. However, our study has demonstrated that there was no significant difference in the length of operation or anaesthetic when compared with normal practice. It must be noted however that although actual times for anaesthetic and procedure have not been impacted, turnaround time between cases has anecdotally increased at UHB. This is due to an additional 15-minute theatre ventilation air change prior to each new case, supplementary cleaning protocols and the removal of the communal ‘holding bay’ prior to surgery. These data were not available retrospectively and so should be included in any future prospective study when analysing length of theatre time and planning realistic theatre lists.

The only statistically significant finding in this study was a difference in the grade of primary surgeon. Although there was no difference in the proportion of patients operated on by a consultant, there was a shift towards seniority among operations performed by non-consultants. During the pandemic, more operations were primarily performed by senior SpRs and fellows than junior SpRs compared with normal practice. One reason for this may be that initially, more junior SpRs and senior house officers were redeployed to Intensive Care Unit and proning teams compared with senior SpRs. These data add to the growing body of evidence that COVID-19 has negatively impacted on surgical training in the UK. The national implementation of COVID-19 surgical pathways for the remainder of this national and global health disaster. This study is based on data from a single centre and focuses on neurosurgery patients only. Given the variation in national practices, we would encourage individual hospitals and Trusts to analyse outcomes in the same way. This would provide larger datasets and could identify which protocols have been most effective. In particular, it is important to ascertain whether mixing positive, unknown and negative cases in a shared surgical pathway is detrimental. If not, the national implementation of COVID-light sites should be called into question as these currently limit patient flow in a time of waiting list crisis.

**Limitations**

All matched cohort studies recognise the limitations of matching cases as not all variables can be taken into account. For a COVID-19 study in particular, we recognise that ethnicity and body mass index are contributing factors to morbidity and mortality and these variables were not matched.

The sample size included in this analysis is relatively small and so may be underpowered. Larger studies and meta-analysis would provide further data to support or refute these findings.

During the pandemic period, the UHB swabbing protocol evolved and as a result was not consistent across the whole study period. While a true reflection of our
CONCLUSION

During the initial phase of the COVID-19 pandemic, the mortality, morbidity and functional outcomes of operatively managed neurosurgery patients were not significantly affected compared with normal practice. Although anaesthetic and operative time was not altered, the grade of primary surgeon was significantly more senior which is likely to have negatively impacted on training opportunities for junior surgeons. This adds to the growing body of evidence that surgical training in the UK has been significantly affected by the pandemic. Mixing COVID-19 positive, unknown and negative cases did not significantly impact the outcomes of neurosurgical patients and calls into question the need for COVID-light sites. Larger multicentre studies are needed to confirm or refute these findings and collect prospective data to support the implementation of evidence-based surgical pathways now that we are entering the next wave of the pandemic.

Author affiliations

1Institute of Inflammation and Ageing, University of Birmingham College of Medical and Dental Sciences, Birmingham, UK
2Department of Neurosurgery, University Hospitals Birmingham NHS Foundation Trust, Birmingham, UK
3Department of Neurosurgery, Queen Elizabeth Hospital, Birmingham, UK
4Department of Infectious Diseases and Tropical Medicine, University Hospitals Birmingham NHS Foundation Trust, Birmingham, UK
5Department of Medicine, Royal Centre for Defence Medicine, Birmingham, UK
6Institute of Cancer and Genomic Studies, University of Birmingham, Birmingham, UK
7Institute of Cancer and Genomics, University of Birmingham College of Medical and Dental Sciences, Birmingham, UK

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Patient consent for publication Not required.

Provenance and peer review Not commissioned; externally peer reviewed.

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ORCID iD Emma Toman http://orcid.org/0000-0003-2142-1923

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13 NICE. Nice guideline NG-179. COVID-19 rapid guideline: arranging for the COVID-19 positive patients to enable inclusion for the COVID-19 positive patients, means that it is unsuitable for performing inferential statistics on these data. However, the authors felt it important to include descriptive data for the COVID-19 positive patients to enable inclusion in any future meta-analysis or power calculations.