Decompressive Craniectomy for Severe Traumatic Brain Injury: A Review of its Current Status

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Abstract

Over the past two decades there has been a resurgence of interest in the use of decompressive craniectomy in the management of severe traumatic brain injury. A number of retrospective studies have demonstrated that in the context of intractable intracranial hypertension surgical decompression can lower the intracranial pressure and a significant number of patients make a good long term recovery. It has been generally assumed that by lowering the intracranial pressure not only are patients more likely to survive but outcome is also improved. However a recent trial has demonstrated that this is not necessarily the case. The DECRA study compared early decompressive craniectomy with standard medical therapy and the results were that there were more unfavourable outcomes in the patients in the surgical arm of the trial. Whilst there were some problems with randomisation and crossover of patients from the standard therapy to the medical arm, overall the trial has provided the first class I evidence in adult neurotrauma and should have been heralded as a landmark study. Unfortunately this has not been the case and commentary on the trial has in most cases been somewhat critical. This review examines the trial in detail and outlines the current role and future direction of decompressive craniectomy in the management of severe traumatic brain injury.

Keywords: Traumatic brain injury; Decompressive craniectomy

Introduction

Over the past two decades there has been a resurgence of interest in the use of decompressive craniectomy in the management of severe traumatic brain injury (TBI) [1-4]. The procedure is technically straightforward and can be performed either unilaterally or bilaterally (or bifrontally). A unilateral decompression is usually performed following evacuation of a mass lesion such as a subdural haematoma or when the cerebral swelling is localized to one hemisphere. A bilateral or bifrontal craniectomy is usually performed when there is diffuse cerebral swelling.

The rationale for surgical decompression is based on the strong association between elevated intracranial pressures and poor outcome and mortality following TBI [5,6]. By providing additional space into which the injured brain can expand the intracranial pressure is lowered and it has been assumed that not only are patients more likely to survive, but also outcome is improved.

Unfortunately this assumption has been seriously questioned by the results of a recently published trial comparing early decompressive surgery with standard medical therapy in which the authors concluded that decompressive craniectomy was associated with more unfavourable outcomes [7]. The aim of this review is to examine these findings in more detail and review the current role of decompressive craniectomy in the management of severe TBI.

The History of Decompressive Craniectomy

The procedure was first described in 1894 by Annandale and its use gained popularity in the early 1970’s only to subsequently fall into disrepute due to poor clinical outcomes [8,9]. At the same time experimental studies suggested that decompression may actually worsen cerebral oedema and this led to use of the procedure being almost abandoned [10].

However, throughout the 1980’s its popularity returned. A number of studies demonstrated that in the context of intractable intracranial hypertension, surgical decompression could reliably lower the intracranial pressure not only following severe TBI but also following ischemic stroke [11-13], subarachnoid haemorrhage [14,15] and cases of severe intracranial infection [16-19]. In addition to these promising clinical results, experimental data suggested that cerebral oedema and secondary brain injury may be reduced following early decompression and whilst this may involve a number of independent, factors a significant contribution comes may be due to maintenance of cerebral perfusion [20,21].

The Physiological Rationale for Decompressive Craniectomy

In 1783 Monroe J [22] deduced that the cranium was a “rigid box” filled with a “nearly incompressible brain” and that its total volume tends to remain constant (Figure 1). The doctrine states that any increase in the volume of the cranial contents (e.g. brain, blood or cerebrospinal fluid), will elevate intracranial pressure. Furthermore, if one of these three elements increases in volume, it must occur at the expense of volume of the other two elements. In 1824 Kellie [23] confirmed many of Monro’s early observations. When the brain is injured and starts to swell or there is a mass lesion such as an intracerebral haematoma, the compensation is made at the expense of blood and CSF volume (Figure 2). As the brain becomes progressively more swollen or a mass lesion increases in size these compensatory mechanisms become exhausted and for incrementally smaller increases in volume there are progressively greater increases in pressure (Figure 3). When viewed...
from this perspective the limitations of traditional medical measures such as hyperventilation, barbiturate coma and more recently hypothermia can be appreciated. There is little doubt that barbiturates and hypothermia have the potential to be neuroprotective due to their influence on many mechanisms known to be important in the cellular response to injury such as calcium mediated toxicity, glutamate excitotoxicity, free radical peroxidation and cellular apoptosis [24-27]. However, the often rapid fall in intracranial pressure that occurs after the application of either of these three therapies occurs as a result of cerebral vasoconstriction [28-31]. Given the well known deleterious effects of ischaemia, it perhaps not unsurprising that numerous studies have failed to demonstrate that any of these therapies provide clinical benefit in terms of improvement in long term outcome [28,32-37].

It is here that decompressive craniectomy may have at least a theoretical advantage. By challenging the Monroe Kellie Doctrine and expanding the “rigid box” the intracranial pressure can be reduced but not at the expense of cerebral blood volume and cerebral perfusion appears to be improved [38-40]. What remains to be scientifically established is whether this improvement in cerebral perfusion and oxygenation is converted into clinical benefit.

Clinical Studies

There have now been numerous publications that have demonstrated the role of decompressive craniectomy in the management of intractable intracranial hypertension following severe traumatic brain injury. However, none of these can be adjudged to have provided class 1 evidence on which to base clinical practice.

The DECRA study – clinical findings

The DECRA study is the first randomised controlled trial for adult patients with severe TBI and as such represents the only high-level evidence pertaining to the surgical management of this group of patients. The trial compared early decompressive craniectomy for diffuse traumatic brain injury with standard medical therapy [7]. The results of the trial were that the patients undergoing craniectomy had lower intracranial pressures (ICP) and spent less time in intensive care, however at six month follow up 51 (70%) of patients in the craniectomy group had an unfavourable outcome compared with 42 (51%) of patients in the standard care group ($OR = 2.21 [95% CI: 1.14-4.26]$ $p=0.02$). Based on these findings the authors concluded that not only was the use of decompressive craniectomy associated with more unfavourable outcomes, but also that by using standard medical therapy rather than surgical decompression, healthcare systems would save millions of dollars per year [41]. Unfortunately these conclusions have not been shared by the global community and overall the observations regarding the trial have been fairly critical [42-45].

The DECRA study – critical appraisal

In general the debate has centred around three issues [46,47]. In the first instance there were some problems with randomisation such that the patients in the surgical arm of the trial had sustained a slightly more severe primary brain injury. More patients in the surgical group had bilateral non-reactive pupils (27% versus 12%; $p=0.04$), radiological findings as adjudged by the Marshall grading were more severe (grade III & non-evacuated haematoma: total 77% versus 67%) and the GCS was lower (median 5 versus 6). All these factors are significant prognostically and when the pupil reactivity was adjusted for in the multivariate analysis, there was no statistically significant difference between the two groups. Whilst in isolation the differences in radiological findings and GCS fail to reach univariate statistical significance the cumulative affect of these adverse presentation variables mean that the surgical group had a significantly more severe primary brain injuries. Secondly, there was a significant crossover of patients such that 19 (23%) patients in the medical arm had a surgical decompression. Finally there was criticism regarding the relatively low ICP for a relatively short time period that was used as the clinical indicator for surgical decompression and this meant that the patients enrolled into the study were not representative of current
Based on these observations the overall global response has been fairly critical, indeed some commentators have gone so far as to state that “no conclusions regarding management of the use of decompressive craniectomy in patients with traumatic brain injury should be drawn from this trial and clinical practice should not be changed on the basis of these results” [44]. However, it may be a little premature and certainly a little unfair to adopt a diametrically opposite position and disregard the trial entirely. An alternative approach would be to objectively analyse the results in order to obtain information that can be clinically useful. This raises the following issues that require careful consideration:

**The trial hypothesis:** In the first instance it must be recognised that this was a well organised, multicentre study, based on genuine clinical equipoise. The trial hypothesis was that early decompressive surgery would lower the intracranial pressure and prevent secondary brain injury. At the time of the trial inception a number of a number of observational clinical studies had shown that in the context of intracranial hypertension surgical decompression could successfully lower ICP [3,4]. In addition there had been considerable advances in the understanding of the complex cellular response to trauma and it was becoming progressively more apparent that a substantial amount of cell death is due to a series of deleterious neurochemical cascades that are initiated either at the time of injury or fairly soon thereafter and these can be amplified by secondary insults such as cerebral ischaemia [50,51]. Based on these observations it would appear perfectly reasonable to perform an early decompression in order to limit secondary brain injury.

**Timing of surgical decompression:** What trial has clearly demonstrated is that an ICP > 20mmHg for 15 minutes provides insufficient evidence that there are significant ongoing secondary insults and therefore any benefit conferred by decompression is offset by surgical morbidity. This finding would appear to be unequivocal. Whilst there may be some confounding of the results introduced by some problems with randomisation of patients and crossover between the surgical and standard care arms, it has to be accepted that the current scientific evidence is that early surgical decompression in these particular circumstances does not improve outcome. The observation that to perform a decompressive craniectomy in these circumstances is not representative of the current clinical practice is not unreasonable [42-44,49]. However, it should perhaps be acknowledged that the aim of the trial was not to confirm the efficacy of current practice but rather to change that practice. If the trial had confirmed that early decompression provided benefit then although the indication for decompression in the trial was not representative of current clinical practice, the patients in the trial would in fact come to represent the clinical practice of the future. This would have had significant impact on neurosurgical practice and resources.

**Why did early decompression provide no benefit?:** The question remains as to why the trial failed to show benefit and this must be attributed to surgical morbidity [52-54]. Whilst technically straightforward it is becoming increasingly apparent that there are significant complications associated not only with the decompressive craniectomy but also with the subsequent cranioplasty [52,55,56]. These include, herniation of the cerebral cortex through the cranial defect, subdural effusion, hydrocephalus, syndrome of the trephined and infection. Given the high incidence of complications that have been reported it is perhaps not entirely unsurprising that patients in the surgical arm of the trial had a worse outcome than those patients who had relatively mild and transient intracranial hypertension that in most cases was managed adequately with standard medical care. What remains to be established is if what point does any benefit provided by surgical decompression outweigh the morbidity of the approach.

**The Future of Decompressive Craniectomy in the Management of Severe TBI**

**Advances in multimodal monitoring**

As demonstrated by the DECRA study, using the ICP as an independent indicator of secondary brain injury has some limitations and over recent years a number of multimodal monitoring techniques have been developed such that it is now possible to obtain continuous data regarding a number of physiological and biochemical parameters. The aim of these monitors is to gather as much information as possible in order to assess the severity of secondary insults and there a number of modalities available, the most common being cerebral oxygenation monitoring [57-59] microdialysis [60,61] and continuous EEG monitoring [62,63]. Whilst there continues to be debate regarding the precise role of each type of monitor it is becoming increasingly apparent that information obtained from a single modality that is interpreted independently from other physiological and metabolic parameters is unlikely to provide significant clinical benefit [64]. Over the next few years there are likely to be further advances in real time data analysis which integrates information gained from a number of parameters and presents it in a clinically user friendly fashion such that a more accurate assessment of secondary brain injury is provided [64]. This can then be used to guide appropriately targeted therapeutic intervention such as surgical decompression.

**Ongoing clinical trials**

The role of decompressive surgery as a life saving procedure for those patients who’s ICP continues to rise beyond 25mmHg is currently being addressed by the RESCUEicp (Randomised Evaluation of Surgery with Craniectomy for Uncontrollable Elevation of Intra-Cranial Pressure) trial [65]. Notwithstanding the outcome of this trial the interpretation of any study attempting to demonstrate an improvement in outcome over and above standard medical therapy must be tempered with the realisation that in most centres a decompressive craniectomy is carried out once all medical therapy has either failed or is in the process of failing and the patient is though, unlikely to survive without surgical intervention. In these circumstances attempting to scientifically establish that a decompressive craniectomy provides clinical benefit may be extremely difficult. Once a patient is adjudged to have failed medical therapy can they realistically be randomised to continue that therapy?

**Ethical considerations**

A final consideration is long term outcome following decompressive surgery. Whilst a significant number of patients survive following surgery and go on to make a good functional recovery, a significant number remain severely disabled. To what degree that outcome is acceptable to those individuals is difficult to determine however a recent analysis of the patients who had survived having had a decompressive hemicraniectomy for ischaemic stroke found that those patients that had a severely reduced functional status would not have provided consent for the procedure if they had known their eventual outcome [66]. Currently this issue has not been addressed for those patients that survive following TBI however, there has to come a point where...
Outcome prediction

Until recently the difficulty has always been how to accurately assess the severity of the primary brain injury and thereby deciding at what point serious consideration must be given to this issues. However, the CRASH collaborators (corticosteroid randomization after significant head injury) web based outcome prediction model has gone some way to addressing this problem [68]. The model is based on the data obtained from the CRASH study that investigated whether steroids would improve outcome following TBI [69]. Whilst the results of the trial were negative, the significant amount of clinical data enabled the investigators to develop a prediction model incorporating those factors such as: age, initial GCS, pupil reactivity, extracranial injuries and radiological appearances, that are known to have prognostic significance [70-72].

The model provides a percentage predicted risk of unfavourable outcome at six months (defined by the Glasgow Outcome Scale as; dead, persistent vegetative state or severely disabled [73]. Previous studies have demonstrated how the predicted risk can be used as a surrogate index of injury severity [74,75]. These studies compared the percentage predicted risk of an unfavourable outcome with the observed long term outcome in a cohort of patients who had had a decompressive craniectomy following severe traumatic brain injury. It can be seen from Figure 4 how the predicted risk can be used to stratify patients according to injury severity and this can be used as an objective assessment of the most likely long term outcome following decompressive surgery. For example, it can be seen from these results that once the percentage risk of unfavourable outcome is greater than 80%, the observed long term outcome for those patients that survive is one of severe disability.

Clinical Applications of Outcome Prediction Models

The applications of this type of objective assessment have yet to be explored however a number of small studies have demonstrated how this type of information could influence clinicians opinion when considering what may be considered life saving but none restorative surgical intervention [76,77]. In addition, whilst we fully agree with the CRASH collaborators that the this model should only be used to support and not replace clinical judgement, it could provide supportive information which could be used to facilitate the discussion of realistic outcome expectations.

Conclusions

The current role of decompressive craniectomy in the management of severe traumatic brain injury has yet to be established. The results of the recent DECRA have provided important clinical information and demonstrated some of the difficulties encountered when planning and executing trials of this nature. Whilst the results may not significantly alter the current neurosurgical practice the trial has in some ways suggested that more surgical judgement may be required prior to considering surgical decompression because in certain situations it may provide no benefit and may in fact do more harm. Prior to the findings of the study it was almost assumed that lowering the intracranial pressure by surgical decompression would be beneficial and whilst many authorities remain convinced of clinical efficacy it has to be accepted that this remains scientifically unproven.

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