Randomized controlled study comparing 2 surgical techniques for decompressive craniectomy: with watertight duraplasty and without watertight duraplasty

Eduardo Vieira, MD, Thiago C. Guimarães, MD, Igor V. Faquini, MD, Jose L. Silva Jr., MD, Tammy Saboia, MD, Rodrigo V. C. L. Andrade, MD, Thais L. Gemir, MD, Valesca C. Neri, MD, Nivaldo S. Almeida, MD, and Hildo R. C. Azevedo-Filho, MSc, PhD, FRCS

Department of Neurological Surgery, Hospital da Restauração, Recife, Brazil

OBJECTIVE  Decompressive craniectomy (DC) is a widely used procedure in neurosurgery; however, few studies focus on the best surgical technique for the procedure. The authors’ objective was to conduct a prospective randomized controlled trial comparing 2 techniques for performing DC: with watertight duraplasty and without watertight duraplasty (rapid-closure DC).

METHODS  The study population comprised patients ranging in age from 18 to 60 years who were admitted to the Neurotrauma Service of the Hospital da Restauração with a clinical indication for unilateral decompressive craniectomy. Patients were randomized by numbered envelopes into 2 groups: with watertight duraplasty (control group) and without watertight duraplasty (test group). After unilateral DC was completed, watertight duraplasty was performed in the control group, while in the test group, no watertight duraplasty was performed and the exposed parenchyma was covered with Surgicel and the remaining dura mater. Patients were then monitored daily from the date of surgery until hospital discharge or death. The primary end point was the incidence of surgical complications (CSF leak, wound infection, brain abscess, or subgaleal fluid collections). The following were analyzed as secondary end points: clinical outcome (analyzed using the Glasgow Outcome Scale [GOS]), surgical time, and hospital costs.

RESULTS  Fifty-eight patients were enrolled, 29 in each group. Three patients were excluded, leaving 27 in the test group and 28 in the control group. There were no significant differences between groups regarding age, Glasgow Coma Scale score at the time of surgery, GOS score, and number of postoperative follow-up days. There were 9 surgical complications (5 in the control group and 4 in the test group), with no significant differences between the groups. The mean surgical time in the control group was 132 minutes, while in the test group the average surgical time was 101 minutes, a difference of 31 minutes (p = 0.001). The mean reduction in total cost was $420.00 USD (a 23.4% reduction) per procedure in the test group.

CONCLUSIONS  Rapid-closure DC without watertight duraplasty is a safe procedure. It is not associated with a higher incidence of surgical complications (CSF leak, wound infection, brain abscess, or subgaleal fluid collections), and it decreased surgical time by 31 minutes on average. There was also a hospital cost reduction of $420.00 USD (23.4% reduction) per procedure.

Clinical trial registration no.: NCT02594137 (clinicaltrials.gov)
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KEY WORDS  decompressive craniectomy; traumatic brain injury; brain edema; duraplasty

Decompressive craniectomy (DC) is very effective in the treatment of refractory intracranial hypertension and is a widely used procedure in neurosurgical practice. In the medical literature, however, there are few well-designed studies focusing on the best surgical technique for the procedure; most studies are retrospective series and expert opinion (Levels 4 and 5). This generates debate and controversy about the best technique to be used, especially regarding how to handle the opened dura mater. In 2011, Güresir et al. published a series of cases introducing the concept of rapid-closure DC, in which, after opening the dura mater, no watertight duraplasty is
performed and the exposed brain parenchyma is covered by the remaining opened dura and Surgicel (Ethicon). Our objective was to conduct a prospective randomized controlled study comparing 2 surgical techniques for DC: with watertight duraplasty and without watertight duraplasty (rapid-closure DC).

**Methods**

**Patient Population**

Institutional review board approval was obtained for a single-center randomized controlled prospective trial. The study population comprised patients ranging in age from 18 to 60 years old who were admitted to the Neurotrauma Service of the Hospital da Restauração from January 2012 to December 2013 to undergo unilateral DC. Patients with intraaxial contusions or hematomas requiring surgical evacuation, in which case injury to the arachnoid mater could lead to an increased risk of CSF leak, were excluded. Patients included in the study had written informed consent obtained from next of kin or guardian as appropriate. This study was registered with clinicaltrials.gov (registration no. NCT02594137).

**Study Intervention and Surgical Technique**

After clinical indication for unilateral DC, patients were randomized by numbered envelopes into 2 groups: with watertight duraplasty (control group) and without watertight duraplasty (rapid-closure DC) (test group). In all cases, a large trauma flap (i.e., large reverse question mark starting from the tragus and extending to the midline) was performed. The skin, galea, and muscle layers were elevated according to the surgeon’s preference. Thus, a wide (at least 12 × 15 cm) craniotomy was performed, and the temporal bone was removed until it was flush with the middle fossa floor (Fig. 1). After dural opening, watertight duraplasty with pericranium or an artificial graft (at the surgeon’s discretion) was performed in the control group (Fig. 2), while no watertight duraplasty was performed in the test group, in which the exposed brain parenchyma was covered with Surgicel (Fig. 3). Usual closure was then performed. Patients were then monitored daily by evaluators blinded to the randomization (control or test group) from the date of surgery until hospital discharge or death.

The primary end point of our study was the incidence of complications, namely CSF leak (CSF drainage through surgical wound), subgaleal fluid collections (CSF drainage to the subcutaneous/epidural space, but not through surgical wound), wound infection (limited to the subcutaneous/epidural space; not involving the brain parenchyma), or brain abscess (infection involving the brain parenchyma). Cranial CT scans were obtained routinely between the 7th and 10th postoperative days to evaluate the occurrence of subgaleal fluid collections. The following were analyzed as secondary end points: clinical outcome (analyzed using the Glasgow Outcome Scale [GOS]), surgical time, and...
procedure costs. To estimate procedure costs, the mean costs of a DC procedure per minute were calculated based on anesthetic drug usage and operating room utilization time (data supplied by hospital administration). Costs analysis further included cost of artificial dural grafts (when used).

**Statistical Analysis**

For categorical variables we used the Pearson chi-square test or Fisher’s exact test if the chi-square test was not applicable; the Student t-test or Mann-Whitney test was used for numeric variables. Based on previous studies, we found that the surgical mean time for decompressive DC with watertight dural closure is 129 minutes. For our study, we assumed that the use of rapid-closure DC without watertight duraplasty would reduce surgical time by 40%. Using a 2-sided alpha of 0.01, 25 patients per treatment group would give a 99% power to detect at least a 40% decrease in surgical time.

**Results**

Fifty-eight patients with a mean age of 33.4 years (18–59 years) were included. Thirty-five patients were male. The main indication for DC was traumatic brain injury (TBI) (Table 1). The mean Glasgow Coma Scale (GCS) score at the time of surgery was 8.6 points. Overall, 19 patients presented with signs of herniation that manifested as anisocoria. The mean GOS score for the study population was 2.85.

Twenty-nine patients were randomized to each group. Three patients were excluded (1 due to consent error and 2 due to lack of appropriate data). Of the remaining 55 patients, 28 were assigned to the control group (with duraplasty/watertight closure) and 27 to the test group (without duraplasty/rapid-closure DC) (Fig. 4).

There was no significant difference between groups in terms of age, GCS score at the time of surgery, and GOS score (Table 2). The length of follow-up also did not differ between groups (27 vs 26.4 days, $p = 0.95$) Overall, there were complications in 9 patients, 5 in the control group and 4 in the test group, with no significant difference (Table 3). Two patients in each group developed a CSF leak. Three of them were treated conservatively with acetazolamide and/or lumbar CSF drainage, while 1 patient (in the control group) required surgery for correction. One patient in each group developed wound infection and both were treated surgically. Two patients in the control group and 1 patient in the test group developed subgaleal fluid collection seen on postoperative CT scans. These collections were probably caused by CSF leakage through duraplasty or opened dura but contained by the skin closure. Patients were asymptomatic and required no treatment.

**TABLE 1. Etiology leading to the indication of decompressive craniectomy**

<table>
<thead>
<tr>
<th>Etiology</th>
<th>No. of Patients</th>
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<tbody>
<tr>
<td>TBI</td>
<td>36</td>
</tr>
<tr>
<td>MCA infarction</td>
<td>15</td>
</tr>
<tr>
<td>Vasospasm due to subarachnoid hemorrhage</td>
<td>5</td>
</tr>
<tr>
<td>Dural venous sinus thrombosis</td>
<td>2</td>
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</table>

MCA = middle cerebral artery.

![Flowchart of enrollment, randomization, and follow-up of the patients.](image)
In the test group, the mean surgical time was 101 minutes and in the control group it was 132 minutes; the difference was 31 minutes, which was statistically significant (p = 0.001). Considering the decrease in surgical time, operating room utilization time, anesthetic drugs, and artificial graft, there was a mean total cost reduction of $420.00 USD (cost reduction of 23.4%) per procedure in the test group (Table 4).

**Discussion**

In 1908, Cushing stated, “an accurate approximation of the dura in its two layers is desirable and should be painstakingly done!” This statement became the goal to be reached at the end of any case by generations of neurosurgeons aiming to prevent CSF leakage and infection. Recently, however, studies have demonstrated that a nonwatertight dural closure does not always result in a higher incidence of complications. Barth et al. intraoperatively randomized patients who were undergoing supratentorial craniotomies in which primary dural closure could not be accomplished into 2 groups. In the first group, watertight closure was achieved using grafts, while in the second group, the dura mater was partially sutured with 3–5 interrupted single sutures with an interspace of 2–3 cm. Between the groups, there was no difference in the incidence of CSF leak and/or infection, and, in the nonwatertight dural closure group, surgeries were faster and less expensive ($213 vs $678 USD). Sade et al. also showed that in surgery for supratentorial meningiomas, watertight dural closure was not superior to dural reconstruction using an artificial dural graft applied in an onlay, nonwatertight fashion.

In DC, watertight dural closure, by definition, cannot be achieved in primary fashion, i.e., solely by bringing the dural edges together. Once the decision is made to perform a watertight dural closure in a DC, some type of graft will always be needed. Such a graft can be autologous (pericranium, fascia lata) or artificial. The rationale for using such grafts is, at least theoretically, to prevent CSF leaks and infection.

Some authors, however, have proposed nonwatertight duraplasty for DC. Ragel et al. describing the treatment of wartime TBI during conflicts in Iraq and Afghanistan, as well as Holland and Nakaji, did not perform a watertight dural closure and instead utilized a dural substitute onlay graft, with no increase in the complication rates. We do not use artificial dural grafts; instead, we cover the exposed brain parenchyma with Surgicel. Güresir et al. described the rapid-closure DC, a procedure that is similar to the one we used in our study. With a total of 341 procedures, the authors concluded that, compared with previously reported studies in which DC was performed with watertight dural closure, the rapid-closure technique did not increase the incidence of CSF leak, infection and/or healing problems; in addition, surgical time was significantly decreased (69 vs 120 minutes). More recently, Kolias et al., in a review article, advocated the same technique for performing DC.

**Review of the Surgical Technique for DC**

Decompressive craniectomy is a familiar procedure for most neurosurgeons, and most of the studies focusing on its surgical technique are expert opinions or retrospective series, which creates controversy about the optimal technique. There are 2 possible cutaneous incisions through which to perform DC: the expanded reverse question mark incision and the “T” incision, as described by Kempe for hemispherectomy. The first incision begins no more than 1 cm anterior to the tragus and curves posteriorly around the pinna to the posterior mastoid line, where it turns superiority, reaching the midline and then running parallel to the sagittal suture to reach the hairline. This incision provides good bone exposure but often leads to coagulation of the posterior auricular artery and occipital artery, which makes preservation of the superficial temporal ar-
tery and its branches (frontal and parietal) crucial for an adequate skin flap vascularization and wound healing. Furthermore, healing disorders may occur on the posterior (occipital) portion of the incision due to compression by the weight of the patient’s head in the supine position. The T incision consists of 2 straight incisions: a frontooccipital incision parallel to or at the midline and an incision perpendicular to the first one, starting 1 cm anterior to the tragus and running toward the midline until it reaches the frontooccipital incision. This incision technique may lead to injury of the parietal branch but preserves the frontal branch of the superficial temporal artery as well as the posterior auricular artery and occipital artery, leading to a better flap vascularization. It also eliminates potential healing difficulties related to compression of the wound. In our department, we often opt for the reverse question mark, with possible exceptions in elderly patients (> 60 years old), for which we favor the T incision. The temporalis muscle can be elevated in a single layer along with the skin flap or separately. Our preference is to raise the skin and temporalis muscle separately because a better exposure of the squamous portion of the temporal bone can be achieved in this manner, and a better temporal lobe decompression can then be performed.

After adequate bone exposure, raising the bone flap is the next step. Jiang et al.8 carried out a randomized controlled prospective trial comparing 2 sizes of bone flaps for DC: a frontotemporoparietal flap (12 × 15 cm) and a temporoparietal flap (6 × 8 cm). At the 6-month follow-up, 39.8% of patients in the frontotemporoparietal group had a favorable outcome according to the GOS scores (4 and 5), compared with only 28.6% in the temporoparietal group. In addition, patients in the temporoparietal group had a higher incidence of intracranial hematomas, incisional hernias, and CSF fistulas. Since this study was published, this craniectomy size has been the standard for all DCs performed in our service. Moreover, we believe that it is important to remove temporal bone until it is flush with the middle fossa roof. Herniation of the brain parenchyma through a small craniectomy leads to compression of vessels, especially veins, against the bone edges, which makes the venous drainage difficult, exacerbating brain edema and leading to postoperative bleeding. Recently, Hartings et al.15 compared the effectiveness of the treatment of traumatic brain injury at 2 centers. In cases requiring DC, the center in which larger bone flaps were performed (82.4 vs 52.4 cm²) produced better clinical outcomes (67% good results [GOS Score 4 or 5] vs 46%). Tamrikulu et al.18 compared different sizes of DC (12–15, 15–20, and 20–24 cm) and found no additional benefit in wider craniectomies provided that the lower limit of 12 cm is observed.

The durotomy also must be large to avoid compression of the parenchyma against dural edges. There are several ways to perform the durotomy, each with advantages and disadvantages. In our department, the most performed durotomies are the C-shaped and the stellate durotomies. Regardless of the durotomy shape, it is important that the major vessels are not compressed against the dural edges. When this occurs, relief cuts perpendicular to the initial durotomy should be performed. Csókay et al.2 described the creation of vascular tunnels with hemostatic sponge and absorbable sutures that, at least in theory, could improve circulation to the herniated brain.

Alternative procedures for brain decompression have been proposed to replace standard DC to avoid a second surgery to replace the removed bone flap (cranioplasty). The most popular and performed procedure is the “hinge” craniotomy in which the bone flap is secured to the skull near the midline through a “Y” plate and 2 other plates are attached only to the bone flap to avoid its subsiding after brain edema resolves.4,12,13 Modifications of this technique have been proposed, such as dividing the bone flap into 2 or even 4 pieces or hinging the bone flap with sutures instead of plates, but all of them maintain the same principle of hinge.15,19 Although some studies have shown that this technique is as effective as DC, the brain expansion volume is lower (77.5 ml vs 105.1 ml), and in 1 study, the hinge craniotomy was associated with greater hospital mortality.10,11 In our department, we abandoned the use of this technique due to patients requiring reoperation because of inadequate cerebral decompression, with persistent midline shift and intracranial hypertension. We believe that a possible utility for this procedure is after drainage of acute subdural hematomas in which the midline shift is smaller or equal to the thickness of the hematoma, i.e., with no important associated brain edema (Zumkeller index ≤ 0).20

### The Nonwatertight Duraplasty in DC (rapid-closure DC)

We demonstrated that DC without watertight duraplasty is not associated with a higher incidence of complications when compared with the control procedure with watertight dural closure. Overall, there were 5 complications in the control group and 4 in the test group. It is important to emphasize that the evaluators in our study were blinded to randomization. The most feared complication after DC without dural closure, a CSF leak, occurred in 2 patients in the control group and only in 1 patient in the test group. Furthermore, the incidence of subgaleal fluid collections, possibly representing the occurrence of CSF leaks contained by the skin closure, was higher in the control group (2 vs 1). We concluded that once the arachnoid is intact, there is no increased risk of CSF leaks. Moreover, attempts to achieve watertight closure may lead to small defects on suture lines, causing a “one-way valve” effect that could potentially facilitate the development of CSF leakage.

Patients undergoing DC are often critically ill, especially victims of severe TBI, who are not uncommonly polytrauma patients. In these patients, faster surgical procedures focusing on the restoration of physiological parameters, known as damage control surgery (an already common concept in abdominal trauma), can be of ben-

### Table 4. Mean surgical time and costs per procedure

<table>
<thead>
<tr>
<th>Variable</th>
<th>Mean Op Time (mins)</th>
<th>Mean Cost*</th>
</tr>
</thead>
<tbody>
<tr>
<td>w/ watertight duraplasty</td>
<td>132</td>
<td>$1790</td>
</tr>
<tr>
<td>w/o watertight duraplasty</td>
<td>101</td>
<td>$1370</td>
</tr>
</tbody>
</table>

* In USD.
ever, we believe that our results can help other centers and neurosurgeons to conduct new studies in different populations.

Conclusions

In our study, DC without watertight duraplasty was not associated with a higher incidence of postoperative complications (i.e., CSF leaks, wound infections, abscesses, and subgaleal collections) and decreased surgical time by 31 minutes on average, which can be beneficial in critically ill patients, especially in victims of severe TBI. We also noted a reduction in hospital costs ($420 USD [a 23.4% reduction] per patient on average). Our study is a randomized controlled trial addressing the surgical technique for DC, a widely used procedure in neurosurgical practice; it provides data for the rational choice of the best technique to be employed in each case.

References


**Disclosures**

The authors report no conflict of interest concerning the materials or methods used in this study or the findings specified in this paper.

**Author Contributions**


**Correspondence**

Eduardo Vieira, Department of Neurological Surgery, Hospital da Restauração, Avenida Agamenon Magalhães SN, Recife PE 52010040, Brazil. email: evcj2005@gmail.com.