Outcomes of Outpatient Hybrid Anterior Cervical Surgery: Safety and Outcome of Outpatient 2-Level Hybrid Anterior Cervical Discectomy and Fusion plus Adjacent Total Disc Replacement

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ABSTRACT

Objective: The emergence of modern technologies and surgical techniques has challenged anterior cervical discectomy and fusion (ACDF) as the gold-standard treatment for cervical spondylosis. In an effort to reduce fusion levels and preserve mobility, combining ACDF and total disc replacement (TDR) has been explored in the literature. No reports were found which investigated the feasibility of this hybrid procedure in an ambulatory surgery centre (ASC). The authors aim to determine the feasibility of performing combined ACDF with TDR in an ASC.

Methods: We evaluated medical records of 15 consecutive patients, who presented with multilevel cervical degenerative disc disease. Single-stage instrumented ACDF with concurrent cervical TDR was performed in an ASC. Outcome measures examined were visual analogue scale (VAS) scores for neck pain, neck disability index (NDI), Nurick grade, quality of life assessment (QoL) through the physical and mental composite scores (PCS and MCS) of the Short-Form 12 (SF-12) health survey and complications. Outpatient spine surgery protocols and guidelines are provided.

Results: Males accounted for 70% of the patient population with overall mean age of 45.13 ± 1.9 years with mean body mass index (BMI) of 28.2 ± 8.5 kg/m². Minimum follow-up was 12 months. Estimated blood loss was 71 ± 23 milliliters and mean operating time was 45 minutes. Clinically significant improvement was achieved in 80% of patients with mean VAS score for neck pain of 8.4 ± 0.8 reducing to 4.5 ± 1.2, which was statistically significant (p = 0.043). Similarly, pre-operative mean NDI of 55 ± 7% reduced to 33 ± 9% postoperatively (p = 0.03). Nurick grades were 0 in each patient by final follow-up and there were no complications reported. Overall improvement in QoL was also accomplished.

Conclusion: Combined ACDF and TDR can be safely done in an ASC with satisfactory clinical and patient-reported outcomes.

Keywords: Ambulatory surgery centre, anterior cervical discectomy and fusion, cervical spondylosis, hybrid surgery, outcomes, outpatient, total disc replacement

Seguridad y resultado de la discectomía y fusión cervical anterior híbrida de dos niveles ambulatoria con reemplazo total del disco adyacente

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RESUMEN

Objetivo: La aparición de modernas tecnologías y técnicas quirúrgicas ha desafiado la discectomía y fusión cervical anterior (DFCA) como el tratamiento de referencia para la espondilosis cervical. En un esfuerzo por reducir los niveles de fusión y preservar movilidad, se explora la literatura en busca de la combinación de DFCA y el reemplazo total de disco (RTD). No se han encontrado reportes de investigación de la viabilidad de este procedimiento híbrido en centros de cirugía ambulatoria (CCA). Los autores tienen por objetivo determinar la viabilidad de realizar la combinación de DFCA con el RTD en un CCA.

Métodos: Se evaluaron las historias clínicas de 15 pacientes consecutivos, que se presentaron con enfermedad degenerativa multinivel del disco cervical. Una DFCA instrumentada en una sola etapa...
INTRODUCTION
The dominance of anterior cervical discectomy and fusion (ACDF) as the “gold standard” treatment for cervical spondylosis for many years is being challenged by emerging technologies and novel techniques available to spine surgeons. Concerns over the risk of adjacent segment disease (ASD), increase incidence of dysphagia and the desire for less invasive surgery with faster recovery has fueled the development of cervical arthroplasty as a viable alternative particularly with multilevel fusions (1–3). Cervical total disc replacement (TDR) attempts to treat radicular pain and simultaneously preserve functional motion in patients suffering from spondylotic degeneration. Results from a prospective, randomized, controlled multicentre FDA trial concluded that when compared with clinical outcomes, the benefits of cervical TDR were either equivalent, or superior to those after ACDF (4).

The feasibility of performing both ACDF and cervical TDR in a single patient has been demonstrated in a hospital setting (5, 6). The purpose of this paper is to present mid-term follow-up results for patients having a single-stage, combined instrumented ACDF and cervical TDR in an ambulatory surgery centre (ASC) with recommended outpatient guidelines based on our ongoing experience.

SUBJECTS AND METHODS
The medical records of 15 consecutive patients, who presented with multilevel cervical degenerative disc disease (DDD) resulting in myelopathy with or without radiculopathy, were reviewed (Fig. 1).

Patients were only considered for surgery after failed conservative management for at least six weeks, which included anti-inflammatory medication, physical therapy and epidural steroid injections. Pre-operative clinical assessment was made with a comprehensive history and physical examination and the use of the Nurick and neck disability index (NDI) grading systems along with appropriate anteroposterior (AP), lateral radiographs and magnetic resonance imaging (MRI).

Patients with chronic medical conditions were stable and cleared by their physician and/or cardiologist in the case of cardiac disease, prior to surgery. Indications for TDR, included symptomatic spontaneous/degenerative or traumatic herniated cervical nucleus pulposus with or without radiculopathy and cervical degenerative disc degeneration (DDD) without posterior column instability. Indications for ACDF included cervical spondylosis, stenosing herniated discs, degenerative disc disease with instability and facet arthritis, tropism or facetogenic pain. Exclusion criteria for outpatient surgery included, acute severe trauma, fractures, malignancy, infection, unstable chronic medical illnesses, prior anterior cervical fusions or total disc arthroplasty and BMI > 42 (7, 8). The eligibility criteria for surgical intervention for outpatient spine candidates was based on previous study (8).

Informed consent regarding the procedure and its rationale...
was obtained and patients were also made aware that based on intra-operative findings, the proposed surgical procedure could change. Patients were mobilized within hours of surgery and an experienced registered nurse and the attending anaesthesiologist confirmed that patients were fully oriented. All patients were discharged with a responsible adult to drive them home only after confirmation that they were neurologically intact by the attending spine surgeon. Established transfer agreements between the ASC and with neighbouring hospitals within 30 minutes, ensured a mechanism for hospital admission if patients developed any serious problems or cardiac arrest.

**Follow-up**

- Patients were instructed on standardized postoperative protocol (Fig. 2).
- Postoperative, and physical therapy was started. Follow-up continued at six weeks; three, six and twelve-month follow-up thereafter.

All patients had AP and lateral radiographs within the first two weeks postoperatively to ensure implant compliance, and at twelve months to assess for fusion. Functional outcome was assessed by comparison of pre-operative and post-operative patient numeric rating scale/VAS for neck pain as well as the Nurick grade, NDI and quality of life assessment (QoL) through the physical and mental composite scores (PCS and MCS) of the Short-Form 12 (SF-12) health survey.

**Postoperative protocol**

1. Diet is allowed starting with thick liquids and graduate as tolerated over 2–3 days.
2. Soft or hard collar is to be worn as need, while riding a car or walking.
3. It is important to remain mobile, walk as tolerated after first 24 hours.
4. A drain is placed in surgical site; expect dressing to be soiled for the first 24 hours.
5. A member from operative team will remove drain after 24 hours in the office.
6. Most incisions are closed with subcuticular sutures that will dissolve.
7. Cover area to avoid getting incision wet for 2–3 days.
8. While showering avoid allowing water to hit incision directly, apply water resistant bandage.
9. Call if concerned with wound. Pain, reddened, increased drainage after drain removal.
10. Steri-strips fall off in 10–12 days.
12. Pain is expected after surgery. If pain is not relieved by pain medications or getting progressively worse, call office to let us know. Pain medications include: opioid analgesics, antibiotic and muscle relaxant.
13. Avoid anti-inflammatory drugs for 3 months as this delays bone healing.
14. Do not supplement prescription with over the counter analgesics as this may harm your liver.
15. Weakness and tingling in extremities can be part of healing process especially after surgery.

IF THERE ARE ANY ISSUES DO NOT HESITATE TO CALL

**RESULTS**

Mean age was 45.13 ± 1.9 years, with males accounting for 70% of the patient population. Minimum follow-up was seven months with a mean of 19.34 ± 6.34 months. Mean BMI was 28.2 ± 8.5 kg/m². History of smoking and narcotics use was statistically non-contributory. Estimated blood loss was 71 ± 23 millilitres and mean operating time was 45 minutes. Clinically significant improvement was achieved in 80% of patients with mean VAS score for neck pain of 8.4 ± 0.8 reducing to 4.5 ± 1.2, which was statistically significant $[p = 0.043]$ (Fig. 5).

Similarly, pre-operative mean NDI of 55 ± 7% reduced to 33 ± 9% postoperatively $[p = 0.03]$ (Fig. 6).

Nurick grades were 0 in each patient by final follow-up and there were no complications reported. Quality of life assessment through SF-12 interpretation revealed that a minimum clinically important improvement was achieved: mean

**Surgical technique**

Signed consent was obtained for the procedure and under general anaesthesia; patients were prepped and draped under sterile conditions. A modified approach to the standard Smith-Robinson operative technique was used (9). Surgical exposure of the desired vertebral level was achieved through a transverse midline anterior cervical incision. Following discectomy with pituitary ronguers, curette and burr drill to remove affected disc, the posterior longitudinal ligament was retained in situ (10). The centre of the disc was identified to begin trial- ing. A keel was made using burr after which trial was removed and disc arthroplasty performed with Prodisc-C®, Synthes Inc, and West Chester, PA, USA). Discectomy was repeated at the other affected level. Appropriately sized PEEK interbody cage (Arena-C®, SpineFrontier Inc, Malden, MA, USA) packed with DBM pure placed and fusion aided with anterior cervical plate (ACP) (Inset®, SpineFrontier Inc, Mal-den, MA, USA).

Once haemostasis was achieved and the wound was completely dry (Fig. 3), a Penrose drain was placed above the implants and brought though the incision and secured with a sterile safety pin in outpatients for wound drainage to prevent postoperative haematoma development at home. Statistical analysis was performed using Microsoft Excel version 14.1.3. Comparisons were expressed as counts or means with standard error. Tests were considered significant if $p < 0.05$. 

![Fig. 3: Intra-operative photograph showing an artificial disc above an adjacent anterior cervical plate.](image-url)
pre-operative PCS 35.9 ± 3.9, MCS 45.5 ± 3.2 improved to mean postoperative PCS 40.5 ± 3.3, MCS 45.9 ± 3.5. Although improvements in QoL were observed, these changes were not statistically significant ($p = 0.41$ for PCS and $p = 0.93$ for MCS). Fusion was demonstrated at 12 months follow-up (Fig. 4) at the level where ACDF was performed.

Fig. 4: Postoperative anteroposterior and lateral radiographs showing total disc replacement in satisfactory position at C3-4 and an interbody fusion device at the level of C4-5 where anterior cervical discectomy and fusion was performed.

DISCUSSION
Cervical spondylosis is a common disease of the cervical spine, particularly in older adults, as a consequence of ageing (11). Treatment is initiated with conservative measures for at least six months prior to surgical intervention, which can be broadly classified into fusion and non-fusion techniques (12). Although both techniques have many described advantages and disadvantages (13), ACDF has traditionally been the most common surgical treatment method described, especially since the emergence of ACPs, which have been shown to enhance fusion rates (14).

A commonly expressed concern within the literature is that of the influence of ACPs on the development of ASD and moderate to severe dysphagia following ACDF (15, 16). Several studies have shown that this risk increases with the more levels fused though class one evidence is yet to be established (17). Concerns over these complications and the uncertain sequelae of ACDF lead to the development of total (or artificial) disc replacement as a feasible alternative (12).

Biomechanical and clinical studies comparing the two procedures have suggested that increased intradiscal pressures and shear forces at the adjacent levels contribute to accelerated degeneration following fusions with ACP whereas physiologic cervical motion can be preserved with TDR, unlike ACDF (15, 18, 19). Currently, there exists no clear evidence for the superiority of one procedure over the other, however, one major advantage of using TDR over a fusion for less severely degenerated segments is a young and active patient with a high life expectancy (13).

The rationale for the use of this combined fusion-non-fusion technique to treat these patients was based on numerous factors. Varying stages of degeneration exist in patients with spondylotic disease at multiple levels hence, we felt it prudent to limit the extent of surgery by fusing only the segment with the most degenerative changes or patients with facet-mediated pain. Degenerative changes were evidenced by pain and limited segmental motion on physical examination and pre-operative radiologic evidence of partial fusion or osteophytes intimately involved with the facet joints, facet reactive changes or hypertrophy. A TDR in the case of facet pathology will not address the patient’s pain and should be rendered futile.

Secondly, we felt that younger and middle-aged patients were more likely to benefit due to the likelihood of development of adjacent segment disease over-time, with a higher life expectancy, in addition to the motion sparing benefit of a TDR in terms of quality of life with less levels fused. Although desirable, two level cervical TDR has been studied and reported but is not currently FDA-approved in the cervical spine (20). Older patients would also benefit due to a less morbid procedure such as a three-level ACDF. Furthermore, upper cervical levels are more challenging to access surgically, especially with an ACP due to more complex anatomy and less mobile upper oesophagus. Thus, in order to limit prolonged surgery and increased complication risks, we chose TDR for higher diseased cervical levels, which were less severe and did not warrant full fusion. The risk of postoperative dysphagia has been shown to be increased after ACDF as well as with the more cervical levels fused (16, 21), hence, our preference to
use a TDR in the superior less degenerated segments. Finally, with more recent evidence revealing mid and long-term follow-up data showing that the incidence of ASD is not increased after TDR (12), we were encouraged to offer this to our patients at levels where we believed surgical intervention was warranted but ACDF was too aggressive.

Midterm follow-up revealed that patient-reported outcomes were excellent with this combined fusion-non-fusion technique in an ASC. Statistically significant improvement was achieved in reducing pain and disability as evidenced by the mean postoperative Nurick grades and NDI scores.

Recently, authors have begun looking at the feasibility of combining ACDF with TDR for a variety of indications (5, 6). We found one study in the literature that reported the suc-cess of a single-stage “hybrid fusion-non-fusion procedure” as described, which was almost identical to our study (5). However, this was a hospital-based study with slightly differ-ent eligibility criteria and did not utilize ACP as part of their fusion technique.

To our knowledge, this is the first paper, which reports on the outcomes of these cases being done in a single operation, in an outpatient setting. Further, long-term follow-up studies are warranted to conclude whether this procedure does, in fact, reduce the incidence of adjacent segment disease. Additionally, whether this combined technique may lead to biomechanical compromise of the artificial disc adjacent to a fused segment is yet to be determined, however, based on a recent biomechanical study (22) and our clinical experience thus far, this has not been shown to occur. We have had no cases of hardware failure as a direct or indirect consequence of this combined technique to date. Limitations of this study acknowledged are its retrospective nature and small sample size with no control group. This paper adds to the body of knowledge on outpatient spine surgery and the use of a hybrid technique in patients with multilevel disease.

CONCLUSION
Through a consistent operating team and strict follow-up protocols, we were able to achieve satisfactory midterm clinical and radiologic results after combining instrumented ACDF with cervical TDR for the treatment of multilevel cervical spondylosis in an ambulatory surgery centre.

AUTHORS’ NOTES
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REFERENCES