Clinico-Radiological Outcome of Single-Level and Hybrid Total Disc Replacement with Spineart Baguera®-C for Cervical Myeloradiculopathy: Minimum 2-Year Follow-Up Study in Indian Population

Abstract:
Context: Cervical radiculopathy and myelopathy is one of the most frequent ailments encountered by spine surgeon. Motion-preserving surgeries in cervical spine is a standard of care due to its certain advantages such as biomechanical anatomical conformity, reduced chances of adjacent segment degeneration, and revision surgeries. While there is abundant data from some centers, data from developing countries are still limited. Aims: The aim was to study the clinico-radiological outcome of single-level and hybrid total disc replacement (TDR) with Spineart Baguera®-C cervical prosthesis for cervical myeloradiculopathy. Settings and Design: Retrospective study. Materials and Methods: Retrospective analysis of the 29 consecutive patient undergoing single level TDR and hybrid fixation (i.e., TDR with anterior cervical discectomy and fusion) with Spineart Baguera®-C cervical prosthesis for myeloradiculopathy from January 1, 2014 to December 31, 2017, was done. Radiological features and outcome were studied from data collected on Insta-picture archiving and communication system. Statistical Analysis Used: SAS 9.4 was used for all computations. Results on continuous measurements were presented as mean and standard deviation (min-max) and results on categorical measurements were presented as numbers (n) and percentages. Results: Twenty-nine patients were included in the study. The mean age was 43.31 ± 9.04 years with 14 males and 15 females. The most common level of TDR was C5-C6 (72.41%). The mean follow-up duration was 3.14 years ± 1.13 years (2–5 years). The mean hospital stay was 4.93 ± 2.12 days. The mean neck disability index (NDI) at admission was 27.24 ± 7.66 which decreased to 6.41 ± 4.29 at final follow-up. Conclusions: Two-year data on treatment with Spineart Baguera®-C cervical prosthesis shows significantly improved NDI, visual analog scale (arm) with maintenance of movement of the prosthesis.

Keywords: Cervical, myeloradiculopathy, Spineart Baguera®-C, total disc replacement

Introduction
The total disc replacement (TDR) has been in use since the late 1950 due to its proposed biomechanical and anatomical advantages.[1] It offers advantages over fusion such as it maintains the operated segment’s mobility and enables a better strain distribution on the contiguous disc levels as well as on the zygapophysis. Furthermore, it may aid in maintaining cervical lordosis and restoring the disc’s height, which allows a good radicular decompression.[2]

Anterior cervical discectomy and fusion (ACDF) has been considered gold standard since ages for cervical degenerative disorders who have failed conservative treatment.[3,4] Recent understanding of adjacent segment degeneration and pseudoarthrosis in some patients with fusion surgery with similar or even better outcome of TDR in same patients had led to rampant use and research in field of TDR in last decade.[5-9] Various kinds of cervical prosthesis have been developed since then, one among which is Baguera® C cervical prosthesis.[10,11] The Baguera® C cervical prosthesis is considered easier of cervical prosthesis as far as implantation is concerned.[3] The authors have reported outcome of this device previously but no studies has been reported from the Indian subcontinent.[12,13] In this manuscript, we report the clinico-radiological outcome of consecutively operated cases of TDR with Spineart Baguera®-C.
Materials and Methods

Retrospective analysis of clinical and radiological data of patients who underwent TDR or hybrid fixation (TDR with adjacent level ACDF) for cervical myeloradiculopathy from January 1, 2014 to December 31, 2017 was performed.

Inclusion criteria were symptomatic disc disease (radiculopathy/myelopathy) at one to two level in the subaxial cervical spine patients (18–80 years) with symptoms refractory to a thorough trial of conservative management (4–6 weeks).

Exclusion criteria for cervical TDR were pathology at more than two level, prior fusion at an adjacent level, instability on flexion–extension radiographs, inflammatory arthropathy, osteoporosis, infection and severe facet arthrosis at the affected level. History of prior cervical laminectomy and posterior compressive disease not amenable to decompression through an anterior approach were also excluded.

Details on preoperative, operative, and follow-up details of patients were retrieved from medical record section. Radiological data were retrieved and reviewed on Insta-PACS (picture archiving and communication system) version 4.0. Patients were followed up with X-rays in the preoperative, postoperative, 6 weeks, 3 months, 12 months, and final follow-up [Figure 1].

Surgical details included hospital stay, operative time, blood loss, and any intraoperative or postoperative complication. Clinical details included-Japanese Orthopaedic Association (JOA) score, visual analogue scale (VAS) arm, neck disability index (NDI). Radiological assessment included-range of motion (ROM) of prosthesis (The difference in the angular measurement between the lower endplate of upper vertebra and upper endplate of lower vertebra in flexion and extension X-rays), ROM of adjacent level (Difference in the angular measurement at the supra-adjacent level), C2-C7 alignment (Cobb angle between C2 inferior end plate and C7 superior end plate on neutral X-ray), Implant subsidence was measured in neutral X-ray. Middle disc height was measured at TDR level and upper adjacent level to quantify subsidence. All radiological measurements were done using Insta PACS version 4.0 (Meddiff Technologies Pvt. Ltd, Bengaluru, Karnataka, India), computed tomography scan was used to assess for heterotopic ossification (HO) at final follow up. HO were assessed according to the Mehren/Suchomel modification of McAfee scale.[^14] It consisted of zero to four grades. Grade 0: No HO present, Grade I: HO is detectable in front of the vertebral body but not in the anatomic interdiscal space, Grade II: HO is growing into the disc space with possible affection of the function of the prosthesis, Grade III: Bridging ossifications which still allow movement of the prosthesis, Grade IV: Complete fusion of the treated segment without movement in flexion. Radiological data set was blinded and assessed by a fellowship trained spine surgeon.

Surgical details

The patient was placed on the operating table in the supine position with the neck in the neutral position. The patient’s head was secured in place with tape across the forehead to maintain a neutral position, and his or her shoulders are secured down with tape to allow for proper visualization with fluoroscopy. The endplates was examined on lateral fluoroscopy to visualize that they are positioned in a parallel position. The Smith–Robinson approach was used. Excessive exposure was avoided. Distraction pins were placed into the vertebral bodies directly superior and inferior to the disc space. The pins are placed in the midline of the vertebral body on the anteroposterior view. The discectomy and decompression was then performed. Once the discectomy was completed, release of the posterior longitudinal ligament to allow for a thorough assessment of the posterior and postero-lateral decompression. Care was taken to remove all osteophytes as required. Effort was given to minimize the use of high speed burr to prevent any bone debris. Abundant irrigation was done and all bleeding edges were meticulously bone waxed.[^5] Trials are then used to determine proper implant size in all dimensions (height, width, and depth) and ensure proper alignment with the guidance of fluoroscopic imaging. Largest possible size implant in the anteroposterior and mediolateral planes was chosen. Once the proper implant size has been selected, it was placed in the disc space on lateral fluoroscopy and confirmed on the AP view. The size of the implant was
based on a trial that occupies maximum width and depth with the posterior border of the trial flush to the posterior vertebral border [Figures 2-4].

Hybrid surgery, consisting of TDR at the mobile level, along with ACDF at the spondylotic level, was done for patients with multilevel cervical degenerative disc disease [Figure 5]. Postoperatively patients were mobilized on first postoperative day, advised soft cervical collar for 2 weeks. All cases were started on Indomethacin 150 mg for 14 days.

**Implant feature**

The Baguera® C cervical prosthesis (Spineart SA, Geneva, Switzerland) is a biomechanical device designed to be used for TDR. It consists of a high-density polyethylene (PE) nucleus that articulates between two titanium endplate components, with a porous coated exterior and a diamond-like carbon-coated interior. The implant allows a physiological rotation as well as translation in both the anterior-posterior (±0.3 mm) and rotational (±2°) directions. The controlled mobility of the PE nucleus is designed to prevent excessive constraints on the facet joints, and its curve is designed to respect axial rotation movements. The inferior plate and PE design allow 0.15 mm elastic deformation to absorb shocks and vibrations.

**Statistical analysis**

SAS 9.4 was used for all computations. Results on continuous measurements are presented as mean, Standard deviation (min-max) and results on categorical measurements are presented as numbers (n) and percentages. The significance is assessed at 5% level of significance. Student’s t-test (2 × 1 tailed)/Mann–Whitney test has been utilized to find the significance of parameters on a continuous scale between two groups as per the distribution of data.

**Results**

Twenty-nine patients were included in the study. Eighteen patients underwent TDR while 11 patients underwent hybrid fixation. Twenty-seven patients were operated for symptoms of radiculopathy, while two patients were operated for radiculomyelopathy. The mean age was 43.31 ± 9.04 years with 14 males and 15 females [Table 1]. The most common level of TDR was C5–C6 (72.41%). The mean follow-up duration was 3.14 years ± 1.13 years (2–5 years). The mean hospital stay was 4.93 ± 2.12 days. The average surgical time was 193.64 ± 28.73 min in hybrid group and 170.83 ± 46.15 min in TDR only group, and blood loss was 155.55 mL ± 60.96 mL.

**Clinical outcome**

The mean NDI improved from 27.24 ± 7.66 to 6.41 ± 4.29 at final follow up (P < 0.0001). The mean JOA (two patient) improved from 13 to 15.5. VAS (arm) improved from 7.17 ± 1.26 to 0.34 ± 0.55 (P < 0.0001) at final follow-up [Table 2]. Significant improvement was also seen in the VAS and NDI from preoperative to 3 months.
Radiological outcome

The mean ROM of prosthesis was 9.85 ± 0.86 (8.4–12.2) at 3-month follow-up which reduced to 8.47 ± 0.80 (7–10.4) at final follow-up ($P < 0.0001$). The mean ROM of adjacent level was 10.33 ± 0.81 which increased to 10.45 ± 0.81 at final follow-up ($P = 0.0001$). Mean C2-C7 alignment preoperatively was 52.34 ± 2.31 which was 51.46 ± 3.04 at final follow-up ($P = 0.0001$) [Table 3]. Disc height at prosthesis level was 4.09 ± 0.39 mm which increased to 6.77 ± 0.33 mm at 3 months of follow-up ($P < 0.0001$) and reduced to 6.58 ± 0.28 at final follow-up ($P = 0.0001$) [Figure 6]. Disc height at upper adjacent level was 4.17 ± 0.41 preoperatively, and 4.10 ± 0.39 at final follow up ($P = 0.0005$). At final follow-up HO was seen in 8 (27.6%) patients. There was Grade 0 HO in 21 (72.4%) patients, Grade I HO in 3 (10.34%), Grade II HO in 2 (6.90%), Grade III HO in 1 (3.45%) patient. HO restricting mobility was seen in 3 patients, Grade III HO in 2 (6.9%) patients and Grade IV HO in 1 (3.45%) patient. Hence at the final follow-up, mobility on prosthesis was seen in 89.65%.

Table 1: Demographics of patients enrolled in the study

| Parameters                  | Observations |
|-----------------------------|--------------|
| Sex distribution            |              |
| Male                        | 14           |
| Female                      | 15           |
| Mean age (years)            | 43.31±9.04   |
| Symptom distribution        |              |
| Radiculopathy               | 27           |
| Myeloradiculopathy          | 2            |
| Surgical procedure          |              |
| TDR alone                   | 18           |
| Hybrid fixation             | 11           |
| Level of TDR                |              |
| C3-C4                       | 1            |
| C4-C5                       | 4            |
| C5-C6                       | 21           |
| C6-C7                       | 3            |

TDR – Total disc replacement

Discussion

In our study using Baguera device with minimum 2-year follow-up, 89.65% cases had maintained mobility at the disc with an average movement of over 8° at final follow up. Significant improvement in VAS and NDI was seen. Heterotrophic ossification was seen in 27.6% of cases with severe (Grade 3 and Grade 4) grade in 10.34% cases.

ACDF, a safe and reliable technique, is regarded as the gold standard procedure for single or multilevel cervical spondylosis leading to radiculopathy and/or myelopathy.$^{[15,16]}$ Over the years, cervical disc replacement has become an alternative in select cases over anterior cervical fusion. Fusion has been found to be associated with a risk of around 2.4% cumulative rate of symptomatic adjacent segment with 22.2% cases needing revision surgery at 10 years.$^{[17]}$ Studies have shown significantly lesser re-operation rates and improved NDI after 10 years of disc replacement compared to fusion.$^{[18,19]}$

Over the past 40 years, there have been several designs for cervical disc prosthesis. Metal-on-metal prostheses...
include the Bristol Disc, Cummins design, Prestige Disc (Medtronic Sofamor Danek, Memphis, Tennessee), and Cervicore (SpineCore, Stryker Spine, Allendale, New Jersey) system. Another commonly used design uses metal end plates with a “plastic” center. Prostheses using a PE center include the ProDisc-C (Synthes-Stratec) and the Porous-Coated Motion (PCM) cervical artificial disc (Cervitech, Inc., Rockaway, New Jersey). The Bryan Cervical Disc Prosthesis (Medtronic Sofamor Danek, Memphis, Tennessee) is another metal-polymer implant that uses a polyurethane center.

The Baguera®C cervical prosthesis (Spineart SA, Geneva, Switzerland) is a biomechanical device designed to be used for TDR. It consists of a high-density PE nucleus that articulates between two titanium endplate components, with a porous-titanium-coated exterior and a bio ceramic coated interior, in contact with the PE nucleus.

We found significant and sustained improvement in clinical results. Irrespective, of the device type, most studies report similar improvement in clinical parameters. In a 2-year follow-up study using Baguera, Fransen et al. found decrease by around 2° in the ROM. Our study showed an average reduction on 1.64° at final follow up. Mobility at the treated level after 2 years of TDR using Baguera was evaluated by the ROM between flexion and extension; mobility is present when ROM value is at least 2°, or better 4° as suggested by Vital et al. The preserved ROM is from the semi-constraint design of the implant which consist of semi mobile nucleus that allows for near physiological cervical motion compared to a constraint device. The adjacent level ROM in the Fransen study was in contrast to ours, where they showed an increased ROM of the adjacent segment from 10.5° to 13.6° at 2 years. This could be from differences in the study cohort. Two years studies of other disc devices show an average of 6.5° of ROM with Prodisc C, 7.55° for Prestige and 8.1° for Bryan. These slight differences could be because of the cohort difference, difference in the pre-operative ROM, difference rate of occurrence of HO and the device design itself. Few devices need to have a keel prepared or some bony work done on the end plates.
in order to place the device. This is in contrast to Baguera, which does not need any end plate preparation.

There was also a nonsignificant subsidence seen with the use of the implant in our study. Similar subsidence was also seen in study by Fransen et al.[12,13] Though they did not discuss the reason for this late nonsignificant subsidence, it is perhaps from the sinking of the implant spikes into the end plates over time.[Figure 4] Other devices, that use a keel or screws for fixation, do not show subsidence.

We found advanced HO occurring in 10.35% of our cases and no or Grade I HO in 72.40% cases. Fransen et al. had 19.3% cases of Grade 3 and 4 HO in their 2-year follow-up study using Baguera device.[13] Wide variation is seen in the occurrence of HO (7.3%–69.2%).[14,28‑30] Yi et al. found certain factors as significant such as male sex, and implant type. They reported Prodisc-C having the highest rate followed by Mobi-C then Bryan.[29] Though, advanced stages of HO are associated with reduced movement of the prosthesis, its clinical significance is yet to be conclusively determined. A study looking at the impact of HO on patient outcomes showed no statistical difference in terms of VAS neck pain, VAS arm pain, and NDI score when compared to a control group, though ROM was less in patients who developed HO.[18] With longer follow ups coming, presence of HO may be a rule rather than a exception in TDR. Its occurrence has been influenced by patient selection, type of implant and surgical technique. Noriega et al. in a comparative study found Grade 1 or no HO in 67% cases of Baguera compared to 10.5% with Prodisc C. Advanced HO (Grade 3 or 4) was seen in 18.5% cases using Baguera compared to 73.8% using Prodisc and 65% using PCM.[32]

Certain factors have been implicated to reduce the occurrence of HO. Placement of the device as posterior as possible to allow maximum motion of the device, avoiding unnecessary bony exposure, liberal use of bone wax, avoiding use of drill helps to prevent release of bone growth factors.[12,29‑32] Additionally implant that are placed without the need of drilling or keel also help in reducing the incidence of HO due to reduction of release of bone growth factors from marrow. Baguera design avoids need of drilling and keel may assist in lesser incidence of higher grades of HO.[12,13] Use of anti-inflammatory medication for prevention of HO has been looked in cervical disc and hip replacement surgery literature. Although, there is no conclusive evidence in support for this, we used Indomethacin in our series.[33‑35]

Hybrid constructs has become vogue currently for multilevel cervical spondylosis with adequate literature available on its efficacy and good clinical results. It has been considered safe, reliable and effective modality of treatment for multilevel spondylosis.[36‑37] Our results were also similar to other studies with significantly better clinical and radiological outcomes with none having any complications at 2 years follow up.

The limitation of this study was a small sample size. It was a retrospective and noncomparative study with no control arm. Our follow up period was also relatively small and we were not able to assess revision surgeries, however we did see relative preservation of adjacent segment ROM and disc height in our study. In spite of the limitations, we feel that our study might set up a benchmark of applicability of TDR and H-TDR in developing nation with good clinico-radiological outcome.

Conclusions

Spineart Baguera®-C cervical prosthesis used alone or as hybrid construct has good clinico-radiological outcome. Average ROM was maintained at over 8° and advanced heterotrophic ossification was seen in 10.35% cases. Significant improvement was seen in VAS (arm) and NDI.

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Conflicts of interest

There are no conflicts of interest.

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