ACDF Using the Solis Cage with Iliac Bone Graft in Single Level: Clinical and Radiological Outcomes in Average 36 months Follow-up

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Objective: To evaluate the utility of anterior cervical disectomy and fusion (ACDF) with polyetheretherketone (PEEK) cage and autograft through long term (average 36 months) follow-up.

Methods: Thirty selected patients (male:20 /female:10) who suffered from cervical radiculopathy, myelopathy or radiculomyelopathy underwent a single level ACDF with PEEK cage and autograft from iliac crest from March 2006 to July 2008 in single institute. We followed patients for an average 36.4±8.1 months (ranged from 23 to 49 months). The Japanese Orthopedic Association (JOA) score for evaluation of myelopathy and visual analogue scale (VAS) for radiating pain was used to estimate postoperative clinical outcome. Plain x-ray on true lateral standing flexion, extension and neutral position view and 3D CT scan were used every 6 months after surgery during follow-up period.

Results: The mean VAS and JOA scoring improved significantly after the surgery and radiological fusion rate was accomplished by 100% 36 months after the surgery. We had no complication related with the surgery except one case of osteomyelitis. There was one case of Grade I fusion, four cases of grade II, and 25 cases of grade III by radiologic evaluation.

Conclusion: This long term follow-up study for ACDF with PEEK cage shows that this surgical method is comparable with other anterior cervical fusion methods in terms of clinical outcomes and radiologic fusion rate.

Key Words: Anterior cervical fusion • PEEK cage

INTRODUCTION

As the cervical spine degenerates with age, there is increased risk of undesirable conditions such as displacement or degeneration of intervertebral disc, slackness of annulus fibrosis, osteophytosis of vertebral body and instability. Degenerative cervical disc may cause posterior neck pain, radiating pain on arm or shoulder, or cervical myelopathy. Anterior cervical disectomy and fusion (ACDF) is a safe and standard operation for the treatment of degenerative cervical disc disease concerned with radiculopathy or myelopathy. The main purposes of intervertebral cage are biomechanical support, restoration of disc height, maintenance of cervical lordosis and ideal osteointegration.

Tricortical iliac crest autograft results in numerous complications including breaking of bone graft, collapsing, pseudoarthrosis, subsidence, angular deformation, protrusion of bone block, pain or bleeding of donor site and infection. The most frequently donor site related complication is pain, and infection of harvest site could be a nettlesome problem. To solve these problems, various types of artificial cages providing immediate firmness without a plate system have been devised and applied to clinical fields.

A standard cage alone for ACDF is an effective method to treat degenerative cervical disease. Short term clinical follow up data has been published less than 18 months after ACDF with the polyetheretherketone (PEEK) cage. In the current study, we present a retrospective long term (mean period: 36 months) study of thirty consecutive patients after ACDF with Solis™ cage with iliac autograft.

MATERIALS AND METHODS

Patient selection

We collected information from charts of patients who visited our neurosurgical department from March 2006 to July 2008, retrospectively. Only patients with single level ACDF
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Fig. 1. Schematic picture of cervical vertebra body (lateral view).

**Surgical procedure**

All surgeries were conducted by the same surgeon using a standard anterior cervical approach. A right side skin incision was done in all cases; fluoroscopy was used to check the target level. We used a Caspar retractor to retract vertebral bodies. Complete removal of the disc, lateral decompression and end plate flattening for maximal contact with cage was performed. After selection of ideal trial size by using intraoperative fluoroscopy, harvested cancellous bone from the iliac crest filled in the hollow space of the cage. The cage was gently placed and tapped into disc space by mallet. Finally, we checked that the radiopaque titanium spike was in an adequate location by fluoroscopy. We recommended a cervical brace for 2 months after the surgery.

**Clinical evaluation**

We used the Japanese Orthopedic Association (JOA) score for evaluation of myelopathy and visual analogue scale (VAS) for radiating pain to estimate postoperative clinical outcome. We assessed degree of pain relief between preoperative and postoperative (last follow-up) status by using two scoring systems.

We cited radiographic assessment which, Lee et al. suggested in 2009 and applied to our study design. Three parameters were used to evaluate radiologic outcome: anterior disc height (ADH), interbody height (IH) and segmental interbody angle (SIA) (Fig. 1). They were checked by plain x-ray on true lateral standing and classified into flexion, extension and neutral position view and 3D CT scan every 6 months after the surgery. Fusion was defined according to trabecular bony formation across interfaces between cage and endplates and bony bridge formation between endplates. Fusion was classified into 3 classifications: grade I, bridging bone partially filling the cage; grade II, bridging bone filling the cage; and grade III, bridging bone within and around the cage.

**RESULTS**

**Clinical outcome**

We compared VAS for neck and arm between preoperative and last follow-up status, additionally, JOA score was also used. Table 2 shows a summary of clinical parameters. The mean preoperative VAS for neck was 7.4±0.7 and for arm was 7.9±0.8. The mean VAS for neck at the last exam was 2.5±0.8 and for arm was 3.2±1.1 (p-value <0.05). In JOA scoring assessment for 9 patients who were suffered from myelopathy, score improved from 11.1±2.9 at preoperative status to 14.5±2.6 at last exam (p-value <0.05).

**Radiologic findings**

Among 30 cases, 22 patients were performed 3 dimensional CT scan (3D CT) at last follow-up and remainders were checked by cervical dynamic plain x-ray. Radiological fusion rate was

| Characteristic          | n=30                     |
|-------------------------|--------------------------|
| Sex & Age               |                          |
| Male/female             | 20/10                    |
| Age                     | 47.6±7.1 yrs (28-63)     |
| Clinical manifestation  |                          |
| Radiculopathy           | 21                       |
| Myelopatheit            | 2                        |
| Myeloradiculopathy      | 7                        |
| Level                   |                          |
| C3-C4                   | 5                        |
| C4-C5                   | 4                        |
| C5-C6                   | 11                       |
| C6-C7                   | 10                       |
| f/u period              | 36.4±8.1 months (23-49)  |

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Table 2. The mean values of clinical parameters measured before surgery, and at the last follow-up

| Parameters   | Preoperative | Last f/u | p (t-test) |
|-------------|--------------|----------|------------|
| VAS (neck)  | 7.4±0.7      | 2.5±0.8  | <0.05      |
| VAS (arm)   | 7.9±0.8      | 3.2±1.1  | <0.05      |
| JOA score   | 11.1±2.9     | 14.5±2.6 | <0.05      |

Table 3. Mean values of radiological parameters

| Parameters | Preoperative | Postoperative | Last f/u period | p (t-test) |
|------------|--------------|---------------|-----------------|------------|
| ADH (mm)   | 6.27±1.47    | 10.63±1.32    | 8.32±1.08       | <0.05\*    |
| IH (mm)    | 36.7±3.35    | 41.2±4.23     | 38.7±3.75       | <0.05\*    |
| SIA (°)    | -3.35±3.44   | -6.85±3.01    | -3.21±3.61      | <0.05\*    |

Values are means±SD.

Complications

We had no complications concerned with the surgery itself except one case of osteomyelitis. During the follow-up period, there was no cage related complication or soft tissue injury. The superficial osteomyelitis of iliac bone donor site developed in 1 patient and a culture study of the infection site confirmed Methicillin resistant staphylococcus aureus (MRSA) infection. We performed local debridement and irrigation and treated with vancomycin intravenous treatment for 10 days. Infection was controlled and no problem occurred.

DISCUSSION

ACDF has been performed for treatment of cervical degenerative disease and applied to infectious disease, cervical trauma and tumorous condition. Clinically, ACDF with cage system began to be used regularly after clinical success of prospective research year 2000. Single level ACDF with autograft shows over 95% of fusion rate and approximately 80% of neurologic improvement. Some reports postulated there was no need for fusion after cervical discectomy and fusion should be considered only when instability occurs. The authors believe that discectomy alone aggravates instability and certain mechanical substitution is needed. The loss of disc height and increased motion affects pathophysiologic spondylosis. The limitation of a cage-alone procedure is weak initial me-

Fig. 2. A-C: Serial plain X-ray of one patient shows bony bridge formation between C4 and C5 body. (A: preoperative X-ray film, B: postoperative X-ray film, C: last follow-up X-ray film) D, E: 3D CT scan shows grade III fusion comparing preoperative and last exam in one patient. (D: preoperative 3D CT, E: last follow-up 3D CT)
Mechanical stability and subsidence to endplate. A cervical anterior plate and screw system provides reduction of kyphotic angulation, prevention of implant migration, and increases fusion rate and durable fixation. There may be, however, dysphasia due to plate and loosening, breakage of screw, plate migration and stress shield. Innovative development of surgical instrument have reduced complications of the anterior plate system. However, complications are still reported even though dynamic plate or self locking screw is suggested.

The PEEK cages are biologically inactive which means they do not induce a corrosive reaction and provide high versatility and good mechanical properties while causing minimal artifacts on MRI and CT. Furthermore, excellent radioluency gives information that surgeon can use to adjust the cage at an optimal location during operation, and bone fusion can be readily evaluated by plain x-ray. Titanium spikes on upper and bottom can make easy to anchor the vertebral body and provide immediate fixation, therefore cage migration is prevented. The hollow inner space of cage provides an interface that promotes bone fusion. Elastic modulus is defined as the capacity of returning to its initial shape when load is removed. The higher the elastic modulus the higher the forces needed to deform certain material. Lower elastic modulus shows a more natural characteristic with respect of bone material and lead to reduction of stress shield and increase bony fusion rate. Among several cervical implants, the elastic modulus of the PEEK approximates that of bony structure. High elastic mismatch between cage and bone can affect bone growth, promoting cortical thinning and subsidence in the case of metallic cages. Generally, it is acceptable that bone fusion is achieved around 3 to 6 months after the surgery when using the PEEK cage. Cho et al. reported 100% of bone fusion in forty cases was achieved by using PEEK cage comparing 93.1% of bone fusion when using iliac tricortical bonegraft. In this study, we achieved 100% of fusion rate. Only cancellous bone from the iliac crest might be great role of increase fusion rate and small size of patient group might be another cause. The frequency of harvest autograft has decreased gradually, because of the relatively high rate of complications. Complications of tricortical iliac bone graft are various such as displacement bone graft, pseudarthrosis, hematoma, pain, nerve injury and infection of donor site. The incidence of related complications has been reported to be as high as 20% to 50%. Moreover, the low stabilization support of iliac crest grafts requires instrumentation with anterior plate. To reduce complications related with donor site, many types of material have been studied to substitute autologous bone, but none of these have showed advantage over autologous bone. Many complications related with iliac bone graft have been reduced with the use of cage system. In our patient group, the wound incision on the donor site was performed minimally and harvested only small amount of cancellous bone to minimize related complications.

The question of the best implant for ACDF is still controversial. Numerous materials have been tried to substitute for autologous bone graft to reduce donor site related complications. The ultimate material could provide immediate structural support and osteogenic intergration.

Since introduction of the cervical cage, numerous types of cage have been designed; however subsidence of cage was reported frequently. Ha et al. studied subsidence in standalone Solis cage and found 8.1% of cages had subsidence greater than 3 mm. We observed several cases where subsidence occurred, but it did not affect prognosis. Matge et al. noted that a decrease of disc height after operation did not affect clinical outcome. Lee et al. found that subsidence occurred on radiologic evaluation at last follow-up when they used DMB rather than autologous bone graft. However, our data shows that at 36 months follow-up IH is still maintained compared to preoperative IH which implies autologous bone graft helps earlier bone fusion than DBM.

Cervical interbody fusion causes acceleration of degeneration at adjacent levels by increased stress. Symptomatic adjacent segment problems occur at rate of 2.9% per year during ten years after ACDF with autologous bone graft, the risk was lower in multilevel ACDF than in single level, and the fusion rate declines as the number of involved levels increase. In our study, we could not find degeneration of adjacent levels, but we concluded that this is due to small size of patient group.

In current study, the complications of ACDF with the PEEK cage did not include breaking, pseudarthrosis, kyphotic angulation, protrusion of cage itself, and severe subsidence. However, one patient suffered from infection of donor site. Local heating and pus discharge occurred at right iliac crest where cancellous bone was harvested and culture study performed. Methicillin-resistant staphylococcus aureus (MRSA) was detected and vancomycin was prescribed for a week.

To establish a role of the PEEK cage in ACDF, we need prospective larger series and longer follow up studies. The authors searched many articles to find a long term follow-up study for clinical outcome after ACDF with PEEK cage longer than 36 months but we believe that this study is the first report for long term follow-up.

**CONCLUSION**

ACDF with the Solis cage provides favorable clinical outcomes and radiologic fusion rates comparing with other clas-
cal or alternative cervical fusion methods. It does not cause plate relating complication and radiologic opacity of cage spike enable to locate a cage at optimal location by using fluoroscope. There have been several reports that describe the efficacy of ACDF with PEEK cage for short term follow-up, but long term (more than 24 months) follow-up studies are rare. We present here a long term follow-up study for ACDF with PEEK cage and its clinical efficacy is favorable.

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