Late complication of cervical disc arthroplasty: heterotopic ossification causing myelopathy after 10 years. Illustrative case

Che-Han Hsu, MD,1 Yi-Hsuan Kuo, MD,1–3 Chao-Hung Kuo, MD,1,2,4 Chin-Chu Ko, MD, PhD,1,2 Jau-Ching Wu, MD, PhD,1,2 and Wen-Cheng Huang, MD, PhD1,2

1Department of Neurosurgery, Neurological Institute, Taipei Veterans General Hospital, Taipei, Taiwan; and 2School of Medicine, 3Institute of Biomedical Informatics, and 4Department of Biomedical Engineering, School of Biomedical Science and Engineering, National Yang Ming Chiao Tung University, Taipei, Taiwan

BACKGROUND Heterotopic ossification (HO) is a well-documented complication of cervical disc arthroplasty (CDA), although it rarely causes adverse clinical effects. Despite high-grade HO possibly limiting segmental mobility, it is reportedly seldom associated with symptoms.

OBSERVATIONS The authors report a case of a 46-year-old male patient who underwent hybrid CDA and anterior cervical discectomy and fusion for 3-level cervical disc herniation that caused myeloradiculopathy. The surgery was successful; the patient experienced nearly complete recovery postoperatively. The follow-up images, including computed tomography and magnetic resonance imaging scans, showed satisfactory decompression at the indexed levels without residual osteophytes or ossification of the posterior longitudinal ligament. However, 10 years later, the patient presented with symptomatic compressive myelopathy caused by severe HO that prompted a secondary surgery.

LESSONS Although it is generally reported in the literature that HO is clinically innocuous, in this patient, it gradually and progressively developed and caused myelopathy, requiring a secondary surgery. Symptomatic HO can be expected over time, and patients with a high risk of HO deserve long-term follow-up after CDA. Further investigations are warranted to corroborate these risk factors, including multilevel calcified disc herniation, severe spondylosis, and suboptimal placement of the device during primary CDA surgery.

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KEYWORDS myelopathy; heterotopic ossification; cervical disc arthroplasty; Prestige LP; laminectomy

Cervical disc arthroplasty (CDA) has been recognized as a safe and effective alternative to anterior cervical discectomy and fusion (ACDF) for patients with cervical myelopathy and radiculopathy caused by disc herniation or spondylosis.1–5 Many midterm to long-term reports of multicenter prospective randomized controlled trials have demonstrated the effectiveness of CDA in maintaining neurological improvement and preservation of segmental mobility.6–9 The potential to alleviate adjacent segment degeneration (ASD) with CDA has also been suggested but still needs long-term data for corroboration.6,10–13 However, the emerging popularity of CDA has raised the concern of heterotopic ossification (HO), with variable incidence rates ranging from 2.9% to more than half of the patients over time among many series.14–18 Radiographically high-grade HO after CDA could certainly limit segmental mobility at the indexed level, but it rarely alters neurological improvement or clinical outcomes.13,19–24 Although many risk factors and etiologies of HO after CDA have been studied, the reports have unanimously indicated that HO has little clinical consequence and seldom needs revision surgery.25–31

In this report, we present a case of late-onset compressive cervical myelopathy caused by gradually developed HO 10 years after CDA. To date, to our knowledge, this is the first case report on a late-onset development of HO after CDA that caused severe cervical stenosis and required a secondary surgery.

Illustrative Case

Initial Presentation

A 46-year-old male patient presented with clumsiness in both hands, muscle spasms, and unsteady gait after lumbar spine surgery at another
hospital. Upon physical examination, paresthesia over C7–8 dermatomes and hyperreflexia in both legs were observed, accompanied by abnormal Babinski reflexes bilaterally. The radiographs demonstrated mild spondylotic changes with loss of lordosis at C5–7 levels (Fig. 1A). On the computed tomography (CT) scans, calcified discs at C4–5 and osteophytes at C5–6 were seen (Fig. 2A). Magnetic resonance imaging (MRI) showed severe disc herniation and spondylisis at the levels of C4–7 with significant spinal cord compression and increased intramedullary signal intensity on the T2-weighted MRI scan (Fig. 3A). On the basis of the aforementioned clinical symptoms, positive neurological signs, and compatible imaging evidence, the patient underwent a hybrid 2-level CDA at C4–6 (Prestige LP; Medtronic Sofamor Danek) and 1-level ACDF with plate and screws. The original concept was to perform the decompression (anterior discectomy) thoroughly while maintaining the most possible motility (hybrid CDA and ACDF) rather than indirect decompression (posterior laminectomy) and fusion.

The immediate postoperative radiograph revealed satisfactory implant positions and restoration of the cervical lordotic curvature (Fig. 1B). The patient experienced significant improvement after surgery. His clumsiness and unsteady gait greatly recovered after several weeks. He was able to return to work and undertake normal daily activities.

Six months after surgery, a CT scan confirmed the adequacy of decompression for both the canal and the neuroforamen (Fig. 2B). Furthermore, at the 2.5-year postoperative follow-up, MRI demonstrated significant reexpansion of the previously compressed dural sac and almost complete recovery of the increased intramedullary signal intensity of the spinal cord (Fig. 3B). The hybrid CDA and ACDF (C4–6 and C6–7, respectively) surgery was successful for the myeloradiculopathy caused by 3 consecutive levels of disc herniation and severe spondylisis. By performing this procedure, a 3-level fusion (i.e., 3-level ACDF) construct was successfully avoided, and relief of neurological symptoms was attained. Segmental mobility was also successfully preserved up to 10 years after the primary surgery, and the ACDF level appeared solid (Fig. 1C and D).

**Compressive Myelopathy From HO Formation**

Ten years after the primary surgery (i.e., hybrid CDA and ACDF), and having missed regular follow-up visits for reasons, the patient presented with progressive paresthesia over 4 limbs and recurrent gait disturbance for 3 months. There was such rapid deterioration of neurological function that he had to come to the clinic on crutches. Physical examination revealed gait ataxia and hyperreflexia in both legs. The dynamic lateral radiographs of the cervical spine demonstrated McAfee grade 2 HO formation at C4–6 levels and a herniated disc at C3–4, which could be attributed to ASD (Fig. 1C and D).

The growth of HO significantly narrowed the spinal canal at C4–6 where the CDA levels were, whereas the C6–7 ACDF level had solid fusion and little stenosis (Fig. 2C). On the MRI scan taken 10 years after the primary surgery, there was severe spinal cord compression at levels C4–6 with worsened myelopathy, which could be indicated by the lengthy increased intramedullary signal intensity (Fig. 3C). Moreover, the electrophysiological examinations, including motor evoked potential evaluations, revealed bilateral corticospinal tract lesions above the C8 level, a mild degree for bilateral hands, and a moderate degree for bilateral feet. These results suggested recurrent multiple-level compressive cervical myelopathy caused by gradual formation of HO at the sites where previous CDA was successfully performed.

Because the compressive myelopathy was significant and lengthy, the patient was taken for posterior decompression and fusion 10 years after the primary surgery that had attempted decompression and preservation of motion. The patient then underwent a...
C3–7 laminectomy and instrumentation at C3–7 (Fig. 1E). The posterior surgery also went smoothly, and there were no surgical complications. Fortunately, the patient had neurological improvement after rehabilitation and was independently ambulatory after 3 months.

**Discussion**

We present a case of late-onset HO formation after CDA surgery that caused compressive myelopathy after 10 years. The patient had a primary anterior cervical discectomy for 3 levels that consisted of a hybrid reconstruction, including 2-level CDA and 1-level ACDF at C4–6 and C6–7, respectively. By coincidence, the patient had an internal control for comparison of the 2-level CDA to 1-level ACDF for the subsequent results after anterior cervical discectomy. The final images, including those from dynamic lateral radiographs, CT scans, and MRI, showed functioning (preserved segmental motility) of the CDA levels, despite massive HO formation that caused spinal stenosis and myelopathy. Moreover, the ACDF level demonstrated solid fusion and a well-decompressed canal size. The bottom-line observation from this patient indicated that, although the CDA preserved segmental mobility, its complication of HO formation was eventually problematic and required secondary decompression and fusion. The above-described history of the disease intuitively makes one wonder if the progression of cervical myelopathy could have been halted if the patient had received a 3-level ACDF up front. Although this phenomenon was inadequate to generalize opposition to CDA, it warrants further evaluation and longer-term follow-up of patients who undergo CDA surgery. It also implies that HO after CDA might not be as innocuous as the literature has suggested. Moreover, it should be noted that the currently available reports on U.S. Food and Drug Administration trials have only included 1- and 2-level CDA surgery; therefore, any CDA for more than 2 contiguous levels is considered off-label use.

The etiologies of HO remain elusive, and it has been rarely reported in industry-supported large-scale randomized controlled trials comparing CDA with ACDF. However, in several retrospective cohorts outside of the United States, HO has been evaluated in smaller sample sizes. Potential risk factors for HO could be theoretically categorized into aspects of the patient-dependent, surgeon-dependent, and device-dependent risk factors. On the one hand, the reported risks included multilevel cervical spondylosis, calcified disc herniation, nonsmoker, advanced age, suboptimal installation of the device, lack of postoperative consumption of nonsteroidal anti-inflammatory drugs, preoperatively limited segmental mobility, and semiconstrained design of the device. On the other hand, the clinically significant consequences caused by HO have rarely been reported. To the best of our knowledge, only one case was presented by Wenger et al. in 2016, in which HO-associated myelopathy nearly 9 years after CDA surgery with a Bryan disc (Medtronic Sofamor Danek) led to conversion to ACDF.
However, the report provided little imaging support of the follow-up after the primary surgery. Thus, it inevitably raised the concern of residual osteophytes or preexisting segmental ossification of the posterior longitudinal ligament (OPLL).

The present report is the first, to our knowledge, to document progression of HO that eventually became clinically significant and required a secondary surgery. Furthermore, in terms of the type of prosthesis, the present report is also, to our knowledge, the first description of HO-associated myelopathy after 2-level Prestige LP (Medtronic Sofamor Danek) CDA surgery.

The true etiologies, risk factors, and consequences of HO warrant further investigation. Although many reports have demonstrated that HO might not be problematic for years after CDA, long-term evaluations are necessary to corroborate the eventual outcomes of CDA and to clarify the incidences of clinically significant HO with reference to its risk factors.

**Observations**

The authors report a case of a 46-year-old male patient who underwent hybrid CDA and ACDF for 3-level cervical disc herniation and spondylosis that caused myeloradiculopathy. The surgery was successful; the patient experienced almost complete recovery postoperatively. The follow-up images, including CT and MRI scans, showed satisfactory decompression at the indexed levels without residual osteophytes or OPLL. However, 10 years later, the patient presented with symptomatic compressive myelopathy caused by severe HO that prompted a secondary surgery.

**Lessons**

Although it is generally reported in the literature that HO is clinically innocuous, in this patient, HO gradually and progressively developed into a massive growth and caused myelopathy requiring a secondary surgery. Symptomatic HO may be expected over time, and patients with a high risk of HO deserve long-term follow-up after CDA. Further investigations are warranted to corroborate these risk factors, including multilevel calcified disc herniation, severe spondylosis, and suboptimal placement of the device during primary CDA surgery.

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