Rush Rodding of Femur in a Child with Osteogenesis Imperfecta Under Caudal Anaesthesia and Sedation: A Case Report

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Osteogenesis imperfecta presents with unique challenges to the anaesthetist. We report a case of osteogenesis imperfecta, posted for rush rodding of femur managed with caudal anaesthesia and sedation using dexmedetomidine and ketamine.

Key words: osteogenesis imperfecta, caudal anaesthesia, dexmedetomidine

Introduction
Osteogenesis imperfecta (OI) is a rare connective tissue disorder caused by mutations of the collagen type I - COLIA 1 and COLIA 2 genes.1 The disease is characterised by fragile bones, hyperextensible joints, blue sclera, improper dentition, and progressive hearing loss.2 OI is classified into four types according to silence classification.3

We present a known case of osteogenesis imperfect posted for rush rodding of femur successfully managed using caudal anaesthesia and sedation using dexmedetomidine infusion.

Case report
An 8-year-old boy, known patient with osteogenesis imperfecta type 1, weighing 10 kg was posted for rush rodding of femur. He had recurrent episodes of fractures for which he was operated two years back under general anaesthesia. There was no other significant history in the past.

On examination his vitals were stable. He was short in stature, had blue sclera, pectus excavatum and kyphoscoliosis. Airway examination revealed mallampati class II with restricted neck movements.

Rest of the examination findings were normal. Blood investigations including liver function tests and coagulation profile were normal. Cardiac evaluation including echocardiogram was normal.

In view of the possible instability of cervical spine, we decided to do the case under caudal anaesthesia and sedation using dexmedetomidine and ketamine. After noting down the baseline vitals, intravenous ketamine 1mg was given. The child was carefully positioned, and caudal anaesthesia was performed using 3ml of 2% lignocaine with adrenaline and 7 ml of 0.25% bupivacaine. The child was positioned supine carefully with adequate padding and was sedated using dexametomidine infusion with a loading dose of 10 microgram over 15 minutes and maintenance dose of 5 microgram per hour. The surgery lasted for two hours. The child remained haemodynamically stable throughout the procedure.

Discussion
OI is the most common skeletal dysplasia occurring in approximately 1 in 20,000 births. The common feature of most of the cases is a gene mutation that leads to defective collagen formation leading to weakening of connective tissue.4

The anaesthetic challenges in OI include difficult airway, susceptibility to fractures, risk of odonto-axial dislocation and breaking or displacement of fragile teeth during laryngoscopy and intubation.

Positioning the patient with OI is another important concern.5 Adequate padding of pressure points and careful handling of these patients play a very important role in the
successful management of these patients. We ensured adequate padding at pressure points and careful handling especially during the injection of local anaesthetic for caudal block.

Use of regional anaesthesia helps to avoid the complications related to difficult airway. Regional anaesthesia, especially central neuraxial anaesthesia in these patients can be challenging owing to the possibility of platelet dysfunction and presence of spine abnormalities.6

We anaesthetised this child with caudal epidural so that we could avoid airway manipulation and complications related to the same. Moreover, positive pressure ventilation may worsen the already present ventilation perfusion mismatch in patients with OI. We chose dexmedetomidine as the sedative agent since it provides excellent hemodynamic conditions and minimal interference to ventilation.

Conclusion
We conclude that caudal anaesthesia with sedation is an alternative technique for anaesthetic management for lower limb surgeries in paediatric patients with osteogenesis imperfecta.

References