CASE REPORT

A multidisciplinary approach to the management of traumatic intrusion in immature permanent teeth

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SUMMARY

Intrusion in a developing permanent tooth is an extremely complicated traumatic injury and is rarely seen in the permanent dentition. The damage caused by the trauma to the surrounding tooth structures results in a multitude of complications during the healing process. Careful repositioning of the intruded tooth by orthodontic means, revascularisation of the immature tooth to regain vitality, and periodontal treatment to provide the most acceptable aesthetic outcome are described in this case report. It outlines the treatment options to overcome the complications and variable outcomes. The present report emphasises the importance of a multidisciplinary approach to provide complete rehabilitation of a traumatically intruded developing tooth.

BACKGROUND

Intrusion defined as the axial dislodgment of a tooth into its socket, is considered one of the most severe types of dental trauma, and leads to crushing of periodontal ligament (PDL) fibres, the neurovascular bundle and alveolar bone. Intrusive luxation in the permanent dentition is an uncommon event, constituting only 2% of injuries, and is more common in the primary dentition. The nature of the injury is such that it damages all soft and some hard tissue components, causing a multitude of complications. The large amount of energy required to drive the tooth into the socket damages numerous surrounding structures. Shearing stress severs the gingival fibres, compressive forces in the infrabony part of the PDL compress and severe it, and comminution of alveolar bone is evident. Tissues in the apical foramen area are damaged, especially in an open apex, and the crown covered with plaque is displaced into the damaged periodontium increasing the risk of infection.

A multidisciplinary approach is needed to treat such injuries. The pathological outcomes needing management may include radicular inflammatory resorption (external or internal), partial or total pulp canal obliteration, dento-alveolar ankylosis, marginal bone loss and gingival recession. Pulpal necrosis and disturbance of the continued development of the root is commonly seen in immature teeth. The treatment options may include observation for spontaneous eruption (passive repositioning), surgical crown uncovering, orthodontic extrusion (active repositioning) or surgical extrusion (immediate repositioning).

CASE PRESENTATION

A healthy 7-year-old male patient presented with a history of a fall the previous day. The maxillary right central and right lateral incisors were observed to be intruded (figure 1). No previous dental trauma was reported and there was no history of neurological complications. The medical history was non-contributory. Extra-oral findings revealed enlarged bilateral submandibular glands.

INVESTIGATIONS

Intraoral findings: The patient presented with early mixed dentition with a normal overjet and an Angle’s Class I molar relationship. Marginal gingiva around teeth 11 and 12 were red and swollen due to the associated injury. Teeth 11, 12 and 21 were sensitive to percussion. Tooth 11 was severely intruded (the distance from incisal edge of tooth 11 to that of tooth 21 was 7 mm) and firmly lodged into the socket. Tooth 12 was around 4 mm intruded compared to the level of tooth 21 intra-oraly.

Radiographic examination: Extra-oral radiographic examination with the help of a lateral radiograph and an orthopantomogram (figure 2) ruled out alveolar bone fracture and penetration of the root apex though the labial bone plate or the floor of the nose. Periapical radiographs showed incomplete and wide open apices in teeth 11, 12 and 21. The dentinal walls of the affected teeth were thin and incompletely formed (figure 2). There was no evidence of crown or root fracture on the radiograph. A final diagnosis of severe intrusion of tooth 11 and moderate intrusion of tooth 12 was therefore made. Tooth 21 was concussed.

TREATMENT

Immediate care: Oral hygiene instructions were given to the patient. The patient was advised to apply mouth rinses twice daily using a 0.1% chlorhexidine mouthwash and follow a soft diet for a week.

Figure 1 Clinical presentation of the intruded teeth.

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tooth reached its original position in complete alignment with the prepared mixture was then inserted in the canal and temporarily powdered (100 mg) mixed with a base of propylene glycol. The triple antibiotic paste was prepared using ciprofloxacin powder (500 mg), metronidazole powder (500 mg) and minocycline powder (100 mg) mixed with a base of propylene glycol. The paste was used to apply the slow continuous force needed in the required direction (figure 3).

Endodontic management: Ten days after initiation of orthodontic extrusion, endodontic treatment was planned. The high probability of pulpal necrosis due to the severity of the trauma necessitated endodontic treatment to prevent infection or resorption due to the presence of necrotic pulp. Revascularisation was attempted in the involved tooth instead of the conventional calcium hydroxide apexification procedure due to the presence of immature roots and thin dentinal walls. Access to the tooth lingually was now possible since the tooth had extruded from its initial position. The first step was to standardise the radiographs taken by using the same angulation and the same radiograph position at each follow-up visit using film holders (Dentsply XCP RINN) for the paralleling technique.

A local anaesthetic and a rubber dam were used while the access cavity was being prepared. Copious irrigation with 5% sodium hypochlorite (NaOCl) was applied along with normal saline. The canals were not instrumented in order to avoid the formation of a smear layer that can affect the dentinal walls. This is important because it is believed that intact dentinal walls play a role in signalling postnatal mesenchymal stem cells via endogenous growth factor proteins embedded in the walls. A sterile paper point was used to dry the coronal half of the canal. A sterile saline was then condensed gently on the clot and a plug of about 2–3 mm was formed. The MTA was covered with a wet cotton pellet followed by temporary restoration to allow it to set completely. It was later replaced with a glass ionomer base and a composite restoration. The next follow-up visits were scheduled for 1, 3, 6, 9 and 12 months. A retention period of 12 weeks followed orthodontic repositioning. However, a relapse occurred even after this prolonged retention time, stressing the importance of a long retention phase in cases with severe intrusion due to the extensive bone loss that occurs around the involved tooth (figure 4).

Gingival tissue management: A visible gingival lag or recession was observed in the involved tooth after completion of the endodontic treatment even though adequate oral hygiene was maintained by the patient. A high frenum attachment was observed which was suggested to be a possible cause of the gingival recession. A frenectomy and vestibuloplasty were performed to remove the obstacle (figure 4). Following soft tissue healing, the relapsed tooth was moved to its corrected position (figure 5). Improvements in gingival conditions and aesthetics were observed during 6 weeks of prolonged retention with meticulous oral hygiene by the patient (figures 6 and 7).

OUTCOME AND FOLLOW-UP

The radiographs recorded at follow-up visits up to 12 months revealed hard tissue deposition on the canal walls of the immature permanent incisor. The root did not continue to develop particularly in length, but the root apex showed deposition of hard tissue making the root apex blunt and almost closed (figure 8).

Tooth 11 displayed a slight positive response to vitality testing (pulp oximeter) after 12 months, and it was advised that it be kept under observation to check that revascularisation was successful. Heat and cold sensitivity tests and electric pulp testing (EPT) showed negative results, indicating that nerve regeneration had not occurred. However, intervention in the form of root canal treatment would only be required if clinical or radiographic signs of infection were seen in the tooth. The greyish discoloration and translucency of the tooth did not improve, creating an aesthetic problem which would require intervention in the future.
Tooth 12 showed complete apical closure with a positive response to vitality testing, indicating successful recovery from the trauma.

Frenectomy and vestibuloplasty on tooth 11 improved the gingival contour but a slight recession was still observed due to the marginal bone loss around the involved tooth.

DISCUSSION

Intrusion is the most severe form of tooth injury due to the damage it causes to surrounding structures during the traumatic displacement of the tooth into the bone. Healing is complicated by damage to the gingival attachment, contusion of the PDL and bone, and injury to Hertwig’s epithelial root sheath (HERS) (mainly in immature teeth). The treatment of choice is to allow for spontaneous re-eruption, especially when the tooth apex is incomplete or, according to the Royal College of Surgeons of England (RCSE), for cases in which the amount of intrusion is less than 3 mm. Re-eruption occurs particularly when the dental pulp is vital, and seldom occurs when pulp necrosis is established. Orthodontic repositioning represents a biological procedure for teeth with such injury, in addition to allowing access to the chamber for endodontic therapy. There are numerous other studies on the effects of orthodontic forces used to facilitate dental extrusion, and similar results have been observed, along with the potential of this method to prevent the appearance of ankylosis. On the other hand, a few studies have also reported an increased risk of external root resorption and marginal bone loss.

More instances of pulp necrosis are encountered with intrusions compared other luxation injuries. Andreasen observed 100% pulp necrosis in intruded teeth with open apices and 62.5% in teeth with incomplete root formation. Tronstad et al recommended endodontic treatment of intruded teeth with open apices at 1–2 weeks after the occurrence of trauma. In the present case, endodontic treatment was initiated at 4 weeks after trauma, following evaluation of pulpal sensitivity using a cold spray (−0°C) which showed negative results. Although no sign of active infection was present, the severity of trauma, a progressive decreased translucency revealed through transillumination, and grey colour changes in the crown indicated necrotic changes in the pulp. The traditional apexification procedure with calcium hydroxide or MTA does not strengthen the walls; indeed, calcium hydroxide can actually further weaken thin walls and leave the tooth more prone to fracture. Root wall strengthening with composite limits the possibility of root canal retreatment if the need arises in future. Numerous case reports and series have indicated that it is possible to treat necrotic and immature permanent teeth with revascularisation procedures, leading to clinically symptom-free teeth, along with radiographic evidence of resolution of apical periodontitis. The result is a continuous increase in the thickness of dentinal walls with apical closure or further development of root length. In a similar case report, revascularisation was achieved in an immature necrotic permanent incisor after a severe intrusive luxation injury, and complete healing of the periapical area with apical closure was seen, but vitality tests after 27 weeks showed negative results indicating that revascularisation had not occurred, and root canal treatment had to be performed. Although there is little information about the histological features of the tissue produced in the root canal space of teeth after treatment for revascularisation, animal studies have shown that in cases of thickened walls, a cementum-like tissue was formed along the inner dentin wall (termed intracanal cementum-like tissue), and the lumen was often filled with bone-like tissue (termed intracanal bone-like tissue). However, two recent reports have described the presence of pulp-like tissue in human teeth extracted following revascularisation procedures.

Figure 4 Gingival management and orthodontic correction of the relapsed tooth.

Figure 5 Orthodontic correction of the relapsed tooth after gingival healing.

Figure 6 Final presentation of the tooth after the 6-week retention period.
Therefore, due to the potential maturation of the immature root along with formation of a biological barrier, a revascularisation procedure was performed in the present case. Out of the five responses to revascularisation procedures reported by Chen et al. in immature permanent teeth with infected necrotic pulp tissue, the current case reports a type 2 reaction to the procedure with hard tissue deposition on the walls of the root and apex but no continuation of root formation.

A root consists of dentin and cementum which are formed from odontoblasts and cementoblasts differentiated from ectomesenchymal cells in the apical papilla and dental follicle, respectively, after they receive a signal from HERS. A functional root cannot be formed without cementum because acellular cementum and cellular cementum provide the primary and minor anchorage, respectively, of PDL fibres to the alveolar socket. The viability of HERS, apical papilla and dental follicle depends on the severity of the trauma and duration of infection in immature necrotic permanent teeth. Therefore, failure of continued root development may be due to damaged HERS during intrusion.

Gingival recession is also a very common complication in cases of traumatic intrusion. Due to the associated traumatic injury, crestal bone surrounding the involved tooth is damaged. Orthodontic extrusion is usually performed slowly with the help of elastics at a rate which allows bone growth. However, any soft tissue attachments can inhibit movement of the marginal gingiva and prevent it following the normal gingival contour. Therefore, in the present case frenectomy was performed to allow the marginal gingiva to heal without stress. It has been widely believed that appropriately applied orthodontic forces do not damage the periodontium. However, it is suggested that insufficient width of attached gingiva is a predisposing factor for recession. Lang and Löe concluded from their study that 2 mm of keratinised gingiva is adequate to maintain gingival health. Mucogingival surgery may thus be needed during orthodontic treatment to maintain sufficient width of attached gingival. Therefore, vestibuloplasty was also performed in this case to improve the width of attached gingiva and thus prevent recession.

Oral hygiene was also maintained satisfactorily to prevent inflammation and further disruption of the attachment. However, at 12 months after orthodontic and endodontic treatment, gingival recession of the tooth with clinical attachment loss was observed. A loss in the crestal bone level compared to the adjacent teeth was also seen and it was concluded that this had made a large contribution to the formation of the gingival defect. As the child was still growing a further correction of the gingival level was anticipated, and because the patient had a low smile line, the defect did not cause major cosmetic concern. Treatment for tooth discoloration includes conservative chemical bleaching and complete coverage restorations. The ‘walking bleach technique’ could be used in this case, but has the disadvantage of cervical resorption, especially in trauma cases. External bleaching could be performed either at home or in the clinic which would require several visits to the dental office. Treatment options such as veneers and porcelain crowns can also be used for aesthetic improvement.

Figure 7  Improved gingival conditions with healing after soft tissue surgery.

Figure 8  Post-operative radiographs at 3, 6, 9 and 12 months showing apical closure and thickening of the dentinal walls.
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