SURGICAL ENDODONTICS

Policy Number: DCP010.07

Effective Date: March 1, 2020

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Apicoectomy

Apicoectomy may be indicated for the following:
- Failed retreatment of endodontic therapy
- When the apex of tooth cannot be accessed due to calcification or other anomaly
- Where visualization of the Periradicular tissues and tooth root is required when perforation or root fracture is suspected
- Further diagnosis when post endodontic therapy symptoms persist
- A marked over extension of obturating materials interfering with healing

Apicoectomy is not indicated for the following:
- Unusual bony or root configurations resulting in lack of surgical access
- The possible involvement of neurovascular structures
- Teeth with a hopeless prognosis

Periradicular Surgery without Apicoectomy (Includes Surgery and Periradicular Curettage)

Periradicular surgery without Apicoectomy may be indicated for the following:
- Failed retreatment of endodontic therapy
- When the apex of tooth cannot be accessed due to calcification or other anomaly
- When a biopsy of Periradicular tissue is Necessary
- Where visualization of the Periradicular tissues and tooth root is required when perforation or root fracture is suspected
- Further diagnosis when post endodontic therapy symptoms persist
- A marked overextension of obturating materials interfering with healing

Periradicular surgery without Apicoectomy is not indicated for the following:
- Unusual bony or root configurations resulting in lack of surgical access
- The possible involvement of neurovascular structures
- Teeth with a hopeless prognosis

Retrograde Filling

Retrograde Filling is indicated for the following:
- Periradicular pathosis and a blockage of the root canal system that could not be obturated by nonsurgical root canal treatment
- Persistent Periradicular pathosis resulting from an inadequate apical seal that cannot be corrected nonsurgically
- Root perforations
- Resorptive defects

Retrograde Filling is not indicated for teeth with an overall poor prognosis.
**Root Amputation**

*Root Amputation may be indicated for the following:*
- Class III furcation involvement
- Untreatable bony defect (of one root)
- Root fracture
- Root caries
- Root resorption
- Persistent sinus tract or recurrent apical pathology
- When there is greater than 75% bone supporting remaining root(s)
- The tooth has had successful endodontic treatment

**Root Amputation is not indicated for teeth with an overall poor prognosis with or without Root Amputation.**

**Intentional Reimplantation**

*Intentional replantation may be indicated when all of the following clinical conditions exist:*
- Persistent Periradicular pathosis following endodontic treatment
- Nonsurgical retreatment is not possible or has an unfavorable prognosis
- Periradicular surgery is not possible or involves a high degree of risk to adjacent anatomical structures
- The tooth presents a reasonable opportunity for removal without fracture
- The tooth has an acceptable periodontal status prior to the replantation procedure

**Hemisection**

*Hemisection of multirooted teeth may be indicated for the following:*
- Class III or Class IV periodontal furcation defect
- Infrabony defect of one root of a multi-rooted tooth that cannot be successfully treated periodontally
- Coronal fracture extending into the furcation
- Vertical root fracture confined to the root to be separated and removed
- Carious, resorptive root or perforation defects that are inoperable or cannot be corrected without root removal
- The tooth has had successful endodontic treatment

**The following are not indicated for the treatment of lesions that are endodontic in origin, due to insufficient evidence of efficacy:**
- Bone graft in conjunction with Periradicular surgery
- Biologic Materials to aid in soft and osseous tissue regeneration in conjunction with Periradicular surgery
- Guided Tissue Regeneration resorbable barrier in conjunction with Periradicular surgery

**Coverage Limitations**

- These procedures are limited to one per tooth per lifetime

**Exclusions**

- Dental Services that are not Necessary
- Procedures that are considered to be Experimental, Investigational or Unproven

**DEFINITIONS**

**Biologic Materials:** Agents that alter wound healing or host-tumor interaction. Such materials can include cytokines, growth factor, or vaccines, but do not include any actual hard or soft tissue graft material. These agents are added to graft material or used alone to effect acceleration of healing or regeneration in hard and soft tissue surgical procedures. They are also known as biologic response modifiers. (ADA)

**Experimental, Investigational or Unproven Services:** Medical, dental, surgical, diagnostic, or other health care services, technologies, supplies, treatments, procedures, drug therapies or devices that, are determined to be:
- Not approved by the U.S. Food and Drug Administration (FDA) to be lawfully marketed for the propose use and not identified in the American Hospital Formulary Service or the United States Pharmacopoeia Dispensing Information as appropriate for the proposed use; or
- Subject to review and approval by any institutional review board for the proposed use; or
- The subject of an ongoing clinical trial that meets the definition of a Phase 1, 2 or 3 clinical trial set forth in the FDA regulations, regardless of whether the trial is actually subject to FDA oversight; or
• Not demonstrated through prevailing peer-reviewed professional literature to be safe and effective for treating or diagnosing the condition or illness for which its use is proposed; or
• Pharmacological regimens not accepted by the American Dental Association (ADA) Council on Dental Therapeutics

Furcation: The anatomic area of a multirooted tooth where the roots diverge. A furcation involvement refers to loss of periodontal support in a furcation (ADA, 2016).

Glickman Classification of Tooth Furcation Grading (Sims, 2015):
- **Grade I:**
  - Incipient
  - Just barely detectable with examination hand instruments
  - No horizontal component of the furcation is evident on probing
- **Grade II:**
  - Early bone loss
  - Examination hand instrument goes partially into the furcation, but not all the way through
  - Furcation may be grade II on both sides of the tooth, but are not connected
- **Grade III:**
  - Advanced bone loss
  - Examination hand instrument goes all the way through furcation, to other side of tooth
  - Furcation is through-and-through
- **Grade IV:**
  - Through-and-through, plus furcation is clinically visible due to gingival recession

Guided Tissue Regeneration: A surgical procedure with the goal of achieving new bone, cementum, and PDL attachment to a periodontally diseased tooth, using barrier devices or membranes to provide space maintenance, epithelial exclusion, and wound stabilization. (AAP)

Hemisection (bicuspidization): The surgical separation of a multirooted tooth, usually a mandibular molar, through the furcation in such a way that a root and the associated portion of the crown may be removed or retained. (AAE)

Necessary: Dental Services and supplies which are determined through case-by-case assessments of care based on accepted dental practices to be appropriate; and
- Needed to meet your basic dental needs; and
- Rendered in the most cost-efficient manner and type of setting appropriate for the delivery of the dental service; and
- Consistent in type, frequency and duration of treatment with scientifically based guidelines of national clinical, research, or health care coverage organizations or governmental agencies that are accepted; and
- Consistent with the diagnosis of the condition; and
- Required for reasons other than the convenience of you or your dental provider; and
- Demonstrated through prevailing peer-reviewed dental literature to be either:
  - Safe and effective for treating or diagnosing the condition or sickness for which its use is proposed; or
  - Safe with promising efficacy:
    - For treating a life threatening dental disease or condition; and
    - In a clinically controlled research setting; and
    - Using a specific research protocol that meets standards equivalent to those defined by the National Institutes of Health

Periradicular: Surrounding the root. (AAE)

Retrograde Filling: A method of sealing the root canal by preparing and filling it from the root apex. (ADA)

Root Amputation: Surgical removal of all of the root and adherent soft tissues leaving the crown of the tooth intact and supported by remaining root(s). (AAE)

Root End Resection/Apicoectomy: The surgical removal of the apical portion of a root and adherent soft tissues; may be performed in advance of root-end preparation for a root end filling or as a definitive treatment. (AAE)

**APPLICABLE CODES**

The following list(s) of procedure and/or diagnosis codes is provided for reference purposes only and may not be all inclusive. Listing of a code in this policy does not imply that the service described by the code is a covered or non-covered health service. Benefit coverage for health services is determined by the member specific benefit plan.
document and applicable laws that may require coverage for a specific service. The inclusion of a code does not imply any right to reimbursement or guarantee claim payment. Other Clinical Policies and Coverage Guidelines may apply.

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<th>CDT Code</th>
<th>Description</th>
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<td>Apicoectomy – anterior</td>
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<td>D3426</td>
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<td>D3427</td>
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<td>D3460</td>
<td>Endodontic endosseous implant</td>
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<td>D3470</td>
<td>Intentional reimplantation (including necessary splinting)</td>
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<td>D3910</td>
<td>Surgical procedure for isolation of tooth with rubber dam</td>
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<td>D3920</td>
<td>Hemisection (including any root removal), not including root canal therapy</td>
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<tr>
<td>D3950</td>
<td>Canal preparation and fitting of preformed dowel or post</td>
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<tr>
<td>D3999</td>
<td>Unspecified endodontic procedure, by report</td>
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DESCRIPTION OF SERVICES

When retreatment of endodontic therapy is unsuccessful or not possible, surgical treatment may be required. Surgical endodontics encompasses the elimination of pathology through periradicular surgery, root amputation and hemisectioning of multirooted teeth. Currently, many surgical endodontic procedures are being performed less frequently, as the high success rate of dental implants makes them an accepted alternative for challenging cases. As a result, new evidence for these procedures is lacking. However, there is emerging evidence showing the potential regeneration capabilities of stem cells and other biological regenerative materials for lesions that are endodontic in origin.

CLINICAL EVIDENCE

Surgical Endodontics

Kim et al. (2018) conducted a comprehensive review discussing current knowledge as well as future directions on regenerative endodontics. The European Society of Endodontology and the American Association for Endodontists have both released position statements and clinical considerations for these procedures. Endogenous stem cells from an induced periapical bleeding and scaffolds using blood clot, platelet rich plasma or platelet-rich fibrin have been utilized in regenerative endodontics. It is hoped that with the concept of tissue engineering, namely stem cells, scaffolds and signalling molecules, that true pulp regeneration is an achievable goal. However, much is still not known about clinical and biological aspects of regenerative endodontics.

Deng et al (2016) conducted a systematic review and meta-analysis to evaluate the effect of regeneration techniques (RTs) on the outcome of periapical surgery with different protocols for different lesion types. Clinical evidence indicates that most of the repair or regeneration of the bone defect takes place during the first year after the procedure and that very few changes occur later, therefore this study only extracted data collected at 1 year after periapical surgery. PubMed, the Cochrane Library, and Embase were searched through December 30, 2014. Studies that met the inclusion criteria were systematically evaluated, and a meta-analysis was performed. Eight randomized controlled trials met the inclusion criteria. A significantly better outcome was found in the combination group (membranes plus bone replacement analogues) and bone replacement analogue-only group whereas no significant beneficial effect was found in the membrane-only group. The results of this meta-analysis showed that use of RTs in periapical surgery yielded better outcomes than traditional periapical surgery, which was inconsistent with a previous meta-analysis (Tsesis et al 2011). The use of RTs favorably affected the outcome of periapical through-and-through lesions and large lesions (≥10 mm). There was no significant benefit of using RTs for 4-wall lesions. Both the isolated use of bone replacement analogues and the combination of membranes and bone replacement analogues can improve
the outcome of periapical surgery, whereas using membranes alone does not have significantly favorable effects. The use of RTs for through-and-through and large lesions should be recommended.

Song et al. (2012) published the results of a study to evaluate the outcomes of cases that were classified as successes in a previous study of the surgical treatment of lesions of endodontic and combined periodontic endodontic origin. Long-term predictability of treated teeth is important in the decision making process between root retention versus extraction and other treatments. The purpose of this study was to evaluate the outcomes over a period of 6 to 10 years, of the 172 cases of the cases that were classified as successes in the previous study. Patients were followed up every 6 months for 2 years and every year up to 10 years. On every follow-up visit, clinical and radiographic evaluations were performed according to the same criteria as in the original study by the same 2 examiners. The results showed a follow-up rate of 104 out of the selected 172 cases. Of these 104 cases, 97 cases were included in the successful group, 91 with complete healing and 6 with incomplete healing, with the overall maintained success rate of 93.3%. The authors concluded that endodontic surgery has a high rate of long term success and is a viable treatment for retaining endodontically involved teeth requiring surgery.

Sreedevi et al. (2011) conducted a study to evaluate and compare the clinical and radiographic healing following periapical surgery with and without bone grafting. Twenty patients were selected and randomly divided into two groups: A and B. After periapical surgery, Group A patients had the bony defect filled with hydroxyapatite, while Group B did not. Radiographic angulations were standardized for subsequent followup during the period of the study, and only lesions with 0.5-2cm in dimension were selected. Following surgery all patients were assessed both clinically and radiographically for a period of nine months. Clinical parameters assessed included pain on percussion and palpation, mobility, swelling and vitality of adjacent teeth. Radiographically the graft was assessed by comparing it to surrounding bone (the margin between the bone and the graft, radiopacity of the graft in comparison with the surrounding bone, the presence of trabecular bone formation), and size of the lesion. On clinical evaluation the test group (Group A) did not show any significant immediate or delayed clinical symptoms. Radiographically, in the follow up period of 6 - 9 months the bone graft became indistinguishable from the surrounding bone which indicates complete bone regeneration. Group B showed incomplete bone fill at the end of the nine month evaluation period. The authors concluded that bone regeneration following periapical surgery is effective and can be facilitated using an alloplastic bone graft. Randomized controlled studies with larger patient populations are required to validate these findings.

Dhiman et al. (2015) conducted a prospective randomized controlled trial to evaluate the healing outcomes of platelet-rich fibrin (PRF) in periapical surgeries involving apicomarginal defects and to compare these results with surgeries not using any guided tissue regeneration techniques. Thirty patients with suppurative chronic apical periodontitis and apicomarginal communication were randomly assigned to either the PRF or the control group. Clinical and radiographic parameters including pocket depth (PD), clinical attachment level, gingival marginal position, size of periapical lesion, and percentage reduction of the periapical radiolucency were recorded at baseline and at an interval of 3 months for a period of 12 months. The results showed an overall success rate of 83.33%, with a success rate of 86.66% (13 of 15 teeth) for PRF group and 80% (12 of 15 teeth) for control group. Both the groups exhibited a significant reduction in PD, clinical attachment level, gingival marginal position, and size of periapical lesion at 12-month period. No significant differences were observed between the 2 groups for these parameters except PD, which showed a statistically significant reduction in the PRF group. The authors concluded that the adjunctive use of regenerative techniques may not promote healing of apicomarginal defects of endodontic origin.

von Arx et al. (2011) conducted a literature review of the current clinical and experimental studies to evaluate the outcome of regenerative techniques (RT) in conjunction with apical surgery with regard to type of periapical lesions. A literature search with PubMed and Cochrane databases was conducted in April 2010 and a total of 11 clinical and 10 experimental studies fulfilled the inclusion criteria and included the following: The assessed outcome had to be periapical healing based on radiographic and clinical parameters for clinical studies, or periapical healing based on radiographic, histologic or histomorphometric parameters for experimental studies. The studies had to have a minimum of 10 teeth with a minimum follow-up period of 6 months for clinical studies and of 8 weeks for experimental studies. One clinical and one experimental study of those included differentiated between different types of lesions. The current literature studies do not contain strong evidence demonstrating the need for more research. However, based on the current literature, the authors concluded that the reviewed clinical and experimental studies show no or only minimal benefits of using RT in apical surgery for the treatment of osseous defects limited to the periapical area, (as unimpeded new bone formation can take place) and that the use of RT in apical surgery for treatment of lesions limited to the apical area is not warranted. None of the tested regenerative techniques or materials resulted in a better outcome when comparing test and control sites, except for Expanded Polytetrafluoroethylene Membranes (ePTFE) in two studies. However this material has also been associated with an increased risk of wound dehiscence and site infection in several other studies.
**Intentional Tooth Reimplantation**

Asgary et al. (2014) presented a case series aimed at comprehensively introducing intentional replantation (IR) with a focus on its indications and case selection in endodontics. Twenty teeth were selected and 19 of them had failed endodontic treatment and needed retreatment, surgical treatment, or extraction. The same private practice endodontist provided the IR procedure. Teeth were extracted atraumatically, extraoral time kept to a minimum (<15 minutes), leaving the periodontal ligament and root surface untouched. Root end pathology was treated and teeth reimplanted into extraction socket with position verified radiographically. Teeth were not splinted as they were deemed to be outside of primary occlusion. Patients were given post-operative instructions and returned for oral examination at 1, 7 and 14 days, with follow up beyond 6 months planned. Treatment was deemed successful via clinical and radiographic verification. Subjective symptoms such as pain or discomfort were considered failures, as were teeth that showed symptoms of infection or inflammation. Radiographic examination of teeth that showed no change in size of periapical lesion were also considered failures. Patients were followed up from 8-24 months, with the mean being 15.5 months. Of the original 18 teeth treated with IR, 18 were successful clinically and radiographically. One of the two classified as failures did have some resolution of the periapical lesion, however it was not completely eliminated. The authors concluded that with proper tooth and patient selection and skilled providers, IR can have a high success rate.

**Hemisection and Amputation**

Park et al. (2009) conducted a 10 year retrospective study on the long term outcome of root resection of molars. From December 1992 to March of 2006, 579 patients received root resection on 691 molar teeth at the Institute of Oral Health Science, Samsung Medical Center in Seoul, Korea. Cases were chosen based on root resection therapy for periodontal problems, endodontic problems, caries and root fracture. Ultimately 60 cases were excluded due to missing clinical information, and a retrospective review was done of all clinical and radiographic documentation. Data collected included type of prosthetic abutment, opposing dentition, furcation classification, and amount of bone support on remaining root. They also included clinical information in regards to the presence of periapical lesions, endodontic status and total number of teeth remaining in the dentition. The amount of bone was measured using radiographs taken with the same film holding device to minimize operator differences in film and tubehead placement providing standardization. The study showed a 10 year survival rate of molar resected teeth of 29.8% which is similar to previous studies. The researchers concluded that root resection is still a valid treatment option for retention of teeth with loss of bone due to periodontal disease or endodontic lesions, with periodontal defects showing a slightly higher long term prognosis. Success is highly dependent on patient case selection, careful prosthetic planning and practitioner skill level. The authors also concluded that further studies are needed in this area, but not likely due to more dentists and patients choosing extraction and implants as a treatment modality with higher long term success.

Zafiropoulos et al. (2009) conducted a retrospective non randomized study on the long term success of mandibular molar resectioning and implant procedures in a private practice setting. A retrospective chart review was performed. In one group of patients 56 mandibular first or first and second molars were treated by hemisection (Group H). A second group received 36 implants in the mandible to replace periodontally involved first or first and second molars (Group I). All patients had been in maintenance for at least 4 years after treatment and the occurrence and timing of posttreatment complications were evaluated. The majority of hemisected teeth (68% of Group H) and implants (89% of Group I) remained free of complications for the entire observation period. Group H had a greater incidence of overall complications. The results indicated that both root resected mandibular molars and mandibular molar implants could be expected to have, on average, a complication-free survival of 6 years. Although root resected molars were at a significantly greater risk for complications, approximately 80% of root resected mandibular molars were retained overall, and almost 70% of root resected mandibular molars remained complication free for an average of 5 years.

The authors concluded that within the limitations of this retrospective, practice-based study, implants replacing periodontally involved mandibular molars had fewer complications than hemisected mandibular teeth, but hemiseected teeth have an acceptable long term survival rate.

**U.S. FOOD AND DRUG ADMINISTRATION (FDA)**

Products used for bone grafting and guided tissue regeneration for endodontic applications are extensive. See the following website for more information and search by product name in device name section: http://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfPMN/pmn.cfm. (Accessed December 2018)

**REFERENCES**

American Dental Association (ADA). Glossary of Clinical and Administrative Terms.
American Dental Association CDT Codebook 2020.


Royal College of Surgeons Guidelines for Surgical Endodontics. 2012.


UnitedHealthcare Insurance Company Dental Certificate of Coverage. 2018


### POLICY HISTORY/REVISION INFORMATION

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<th>Date</th>
<th>Definitions</th>
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<td>03/01/2020</td>
<td>Updated definition of:</td>
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<td>o Hemisection (biscuspidization)</td>
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### Supporting Information

- Updated References section to reflect the most current information
- Archived previous policy version DCP010.06

### INSTRUCTIONS FOR USE

This Dental Clinical Policy provides assistance in interpreting UnitedHealthcare standard dental benefit plans. When deciding coverage, the member specific benefit plan document must be referenced as the terms of the member specific benefit plan may differ from the standard dental plan. In the event of a conflict, the member specific benefit plan document governs. Before using this policy, please check the member specific benefit plan document and any applicable federal or state mandates. UnitedHealthcare reserves the right to modify its Policies and Guidelines as necessary. This Dental Clinical Policy is provided for informational purposes. It does not constitute medical advice.