Mineral trioxide aggregate and areas of its use in dentistry: A literature review

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Abstract

Mineral trioxide aggregate is a biocompatible dental material consisting of thyroxide powder and hydrophilic particles. In addition to their dentinogenic and osteogenic potentials, it also exhibits good sealing properties. It has a wide range of areas of use in endodontic and restorative dentistry. In clinical practice, it is a material of choice in many applications such as repair of perforations, root-end fillings, apexification and regenerative endodontic treatment procedures, vital pulp treatments.

Keywords: Mineral trioxide, biocompatible dental, thyroxide powder and hydrophilic particles

Introduction

Mineral trioxide aggregate (MTA) is a biomaterial that has been studied for endodontic applications since the early 1990s. MTA was first described in the dental literature in 1993 and was approved for endodontic use by the U.S. Food and Drug Administration in 1998 [1, 2]. MTA is a hydrophilic and biocompatible endodontic material that promotes the growth of cement and bone formation thanks to both its dentinogenic and osteogenic potentials [3]. In studies evaluating the microleakage level of MTA using dye penetration, liquid filtration and bacterial leakage models, it was observed that it provided better sealing compared to conventional endodontic materials [4]. MTA has a wide range of areas of use in both endodontic and restorative dentistry. MTA, which is becoming more widespread in clinical practice, has become the center of attraction for many researchers. Its physicochemical structure, setting reaction, characteristics of biocompatibility are areas requiring further studies. MTA has undergone many changes since its introduction for dental purposes, its physical and chemical properties have been improved, and it seems these properties will be improved further. This review aims to conduct an up-to-date research on the components, properties, clinical applications, and advantages/disadvantages of MTA.

Content and Chemical, Physical, and Mechanical Properties

MTA consists of fine trioxide powder (tricalcium oxide, silicon oxide, bismuth oxide) which sets in the presence of moisture and hydrophilic particles (tricalcium silicate, tricalcium aluminum) [5].

MTA material was first described as hydrophilic fine powder, mainly consisting of calcium and phosphorus ion and bismuth oxide was added into it to provide a higher radio-opacity than dentine [6]. Torabinejad reported that the mean radio-opacity of MTA was sufficient for radiographic visibility and equivalent to aluminum thickness of 7.17 mm [6]. Portland cement (75%), which constitutes a considerable amount of MTA content, is a mixture of tricalcium silicate (CaO)2SiO3, di-calcium silicate (CaO)2SiO2, tricalcium aluminate (CaO)3Al2O3 and tetra-calcium aluminoferrite (CaO)4Al2O4Fe2O5. Bismuth oxide (20%) and gypsum (5%) are other components of MTA [7]. Gypsum stone is an important factor in adjusting the setting time, although it is less effective than tetra-calcium aluminoferrite [8].

Until 2002, there was only one MTA material consisting of gray powder, and in the same year, white mineral trioxide aggregate (WMTA) as ProRoot MTA (Dentsply Endodontics, Tulsa, OK, USA) was introduced to address aesthetic concerns [9]. MTA material was then categorized into two types: conventional gray MTA (GMTA) and white MTA (WMTA).
The results of the studies performed using scanning electron microscopy (SEM) and electron probe microanalysis showed that the biggest difference between GMTA and WMTA was in $\text{Al}_2\text{O}_3$, MgO, and FeO concentrations. WMTA contains less $\text{Al}_2\text{O}_3$, MgO, and FeO than GMTA at a rate of 54.9%, 56.5%, and 92.8%, respectively [10]. It was also reported that WMTA had a smaller particle size than GMTA and that the decrease in the amount of magnesium oxide and iron oxide could contribute to the changing color of WMTA from gray to white [11-13].

During the preparation of MTA, the powder is mixed with sterile water in a powder/liquid ratio of 3:1 and obtaining the consistency of wet sand is the objective. Mixing can be performed using a metal or plastic putty knife on a glass or paper pad. When MTA is first mixed, it forms a colloidal gel structure, which sets in about 3-4 hours [14, 15]. Moisture from surrounding tissues or a wet cotton pellet helps with the setting reaction [16]. Paper cones, pluggers, special carriers of syringe type can be used to move the MTA mixture prepared to the desired area [17, 18]. The mixing time of MTA is very important. The recommended mixing time is less than four minutes. It is also preferable to implement it immediately after mixing to prevent dehydration of the mixture and drying out as a sandy mixture [19].

After mixing MTA, the initial pH level is 10.2, and pH level increases to 12.5 after three hours [6, 20]. The setting process is defined as a hydration reaction of tri-calcium silicate (3CaO·$\text{SiO}_2$) and di-calcic silicate (2CaO·$\text{SiO}_2$) [21]. Hydration has been reported to be adversely affected by exposure to the inflammatory pH (pH 5-7.5) range compared to physiological conditions (pH 7.0) [22].

It was shown that mixing MTA powder with different liquids and additives could have an effect on the setting time and compressive strength [23]. Different solutions may be considered instead of sterile water in the preparation of MTA material, but it should be noted that the physical properties of the material may be lost. Sodium hypochlorite gels shorten the setting time of 3% and 5% calcium chloride solution; however, the compressive strength is significantly lower than those prepared using sterile water. The preparation with saline and 2% lidocaine anesthetic solution prolonged the setting time, but the compressive strength was not affected significantly. A MTA material prepared using chlorhexidine gluconate gel was not set [23]. Moreover, CaCl$_2$, Ca (HCO$_2$)$_2$, and Na$_2$HPO$_4$ may be used as accelerators, but the strength of MTA substantially reduces, while the setting time is shortened [23, 24].

The compressive strength of MTA is around 40 MPa in 24 hours and is reported as 67.3 MPa after 21 days. In the literature, it was reported that GMTA exhibited greater compressive strength than WMTA [25]. In a study, it was found that irrigation/oxidizing agents sodium perborate mixed with sodium hypochlorite, saline, and sodium hypochlorite mixed with 30% hydrogen peroxide affected the compressive strength of MTA [26]. There is some evidence that MTA material has a long maturation process which continues following the specified setting period of 3-4 hours. A study showing that MTA used to repair furcation perforations had greater compressive strength in 72 hours compared to 24 hours supports this evidence [27].

Any adhesion between MTA and dentine may be stronger than the cohesive strength of MTA material, as it was reported that MTA-dentine separations were usually occur cohesive separations occurring in MTA material [19].

MTA exhibits excellent sealing as it expands during the setting reaction. Therefore, it is recommended to make MTA contact a moistened cotton pellet before placing permanent restoration to provide better sealing. MTA material of 4 mm thickness was identified to be sufficient for sealing [28]. It was also reported that MTA must have a thickness of at least 3mm to seamlesly seal the apical area [29].

In a study reporting that MTA showed antibacterial activity against enterococcus faecalis and streptococcus sanguis, there was no significant antimicrobial effect against anaerobes [30].

In a different study, it was identified that WMTA and a ZOE preparation had similar antibacterial properties against to Staphylococcus aureus, Enterococcus faecalis, and Pseudomonas aeruginosa in direct contact test [31]. The effect of freshly mixed and set GMA on C. albicans was investigated using the antifungal tube dilution method and it was reported that GMTA inhibited C. albicans [32].

Biocompatibility properties of MTA material were evaluated in many in vitro studies in the literature. In different studies, the mutagenicity of MTA and its genotoxic effect on peripheral human lymphocytes were evaluated. There is no study showing mutagenic and genotoxic properties of MTA [33, 34]. It can also be suggested that MTA with no mutagenic properties is not carcinogenic, considering that all carcinogenic substances are mutagens. Many studies showed that MTA did not adversely affect the activity of periodontal ligament (PDL) fibroblasts. It was identified in SEM analysis that PDL fibroblasts had normal morphology and exhibited growth and bonding to 24-hour MTA surfaces [35, 36]. Moreover, it was reported in a study that PDL fibroblasts showed improved proliferation on MTA [37]. MTA activates osteoblasts by increasing the production of interleukin, promotes the formation of new blood vessels and induces bone resorption, resulting in bone matrix production. Therefore, MTA is a material that has the potential to induce periodontal ligament and bone regeneration [38].

**Areas of Clinical Use**

**Repair of perforations**

Materials widely used to repair perforations include amalgam, Super EBA (Bosworth, Skokie, IL), composites, and recently MTA (Pro-Root MTA, Dentsply, York, PA and MTA Angelus, Londrina, PR, Brazil), Biodentine (Septodont, Saint Maur des Fosses, France) and EndoSequence (Brasseler USA, Savannah, GA) [39].

When MTA was introduced to the market, it became more clear which repair material to use to repair perforations [40, 41]. MTA has many advantages over previous restorative materials when used for the repair of perforations. Its biocompatibility is quite high and has a very low level of probability to create any reaction in periradicular tissues. After the material is placed in the perforation area, a cement-like tissue is shown to expand directly over the material [42, 43]. Moreover, MTA, which is used as a perforation repair material, has been reported to have a positive clinical prognosis in the long term [40, 41].

In a study which included MTA as a perforation repair material, it was shown that MTA did lead to any bacterial leakage based on a 45-day evaluation [44]. On the other hand, when used for furcation repair, there was no significant difference between GMTA and WMTA in resistance to penetration of F. nucleatum [45]. In a different study, microleakage of GMTA and WMTA used in furcation perforation repair was compared in both orthograde and retrograde directions. The results showed that there was no difference in the leakage between the two MTA materials.
However, in the orthograde direction, the microleakage was significantly more. This result suggests that an adequate coronal restoration is required on MTA to prevent coronal microleakage [46].

The material provides good sealing even if the perforation area is contaminated with blood [47]. Although this may seem as an advantage, another study showed that the bond strength to the blood-contaminated root dentine was significantly less than that to the uncontaminated dentine [48].

MTA can also be used for the treatment of perforations occurring as a result of resorption [49, 50]. The main disadvantage of MTA in such cases is the long setting time. The material will remain in contact with oral fluids during the long setting period and move away from the related area. This makes MTA inappropriate for transgingival defects associated with cervical resorption. Therefore, a faster-setting resin ionomer such as Geristore (Den-Mat, Santa Maria, CA) is recommended for lesions going beyond the border of the gums [51-53]. Moreover, both types of MTA cause a discoloration of teeth. Although this is an effect observed in all bioceramics, it is observed much less with new materials such as Biodentine and EndoSequence [54].

Root-end filling material

Cement accumulation is essential for the regeneration of periodontal tissue following apical surgical procedures [55]. Increasing the new cement along the root end and restoration of the root end are necessary for the ideal recovery of the periodontium. This layer also creates a biological barrier that increases the integrity of the apical barrier, making it more resistant to the penetration of microorganisms [56]. The importance of the presence of cement-like tissue adjacent to MTA cannot be underestimated. This situation is mostly observed in sections where MTA is used as a filling material. MTA seems to be able to induce cementoblastic cells to create hard tissue. The results of an analysis (SEM) in a study suggested that cementoblasts could connect to MTA and grow [57]. Furthermore, MTA can increase the production of both proinflammatory and anti-inflammatory cytokines from osteoblasts. The effect of MTA on periradicular tissues is probably due to these reactions. It is necessary to apply MTA with a thickness of at least three mm to seal the apical area [29]. Although there are studies with different results in the literature, it was reported that MTA led to less microleakage compared to amalgam, ZOE, and conventional glass ionomer when used as a root-en filler following an apical resection [1, 33, 47, 58] (65 to be added). There are also studies reporting that the setting of MTA and the ability to prevent microleakage are not affected by the presence of blood [47]. When used as a root-end filling, MTA is less cytotoxic than amalgam, Super EBA, and IRM [59]. In studies on endodontic surgery including dogs and monkeys, it was reported that less periradicular inflammation occurred around the root-end filling material [42, 60-63]. Holland et al. theorized that calcium hydroxide, resulting from the reaction of tricalcium oxide in MTA with tissue fluids, supported hard tissue formation [61-64]. In a clinical study using MTA as a root-end filler following microsurgical applications, it was reported that a clinical success rate of 89% was achieved after a follow-up period ranging from four to 72 months [65].

Canal filling material

Root canal sealers, in which tricalcium silicate and dicalcium silicate content stands out, form the latest bioactive group of the root canal filling materials [66]. These sealers emerged as a result of the popularity of MTA material based on tricalcium silicate and dicalcium silicate, which bursts hydraulic powder [67-69]. Reports on their bioactivity have made these canal sealers even more attractive [70]. It was reported that MTA Fillapex had decreased cytotoxicity following the setting and the sealer offered appropriate bioactivity to stimulate hydroxyapatite nucleation [71, 72].

Radioopacity is a very important property for root canal filling materials. MTA is indistinguishable in radiography, unless a radio-opaque substance is added. It is a significant disadvantage that bismuth oxide providing radioopacity of MTA dyes dentin at the crown in changing colors from brown in the presence of NaOCl, gray in the presence of chlorhexidine, and to black in the presence of glutaraldehyde. This led researchers to search new materials for radioopacity and components such as zirconia dioxide or tantalum oxide were used in different commercial formulations [73].

Sealers including tricalcium and dicalcium silicate sets by reacting with water. The result of this reaction is a mixture of alkaline structure (pH: 12) consisting of calcium silicate hydrates and the matrix Ca (OH)2 [74, 75]. These resulting hydrates are on the surface of the original calcium silicate particles, and hydration gradually penetrates inwards. When tricalcium silicate cement sets, it expands dimensionally at a rate of less than 0.1%. This property helps create a barrier and is very important for uses of endodontic purposes. Although the long setting time of tricalcium silicate cement is a disadvantage for their use in different treatments, they do not have a negative effect on their use as canal sealers. The first MTA introduced to the market was affected by moisture and this prevented its use as a canal sealer. Moreover, the first MTA materials intended for surgical use were not considered suitable for use as canal sealer due to their large particle size and very high film thickness. Negative properties were eliminated in the subsequent studies and many tricalcium and dicalcium silicate-based canal sealers were introduced. Today there are many canal sealers under different trademarks [66, 76-80].

Apexification and Regenerative Endodontic Treatment Procedures

In the treatment of immature teeth with pulp necrosis, conventional approaches using calcium hydroxide for apexification and apical barrier techniques using MTA are preferred.

It was reported that MTA could create both incomplete and complete barrier with mild tissue inflammation when used in apexification. In a study comparing calcium hydroxide and MTA, barrier formation was only observed within the root boundaries of the teeth to which MTA was applied. On the other hand, the formation of an incomplete barrier was observed mostly outside the canal, beyond the previous apical area, in the teeth to which calcium hydroxide was applied [81].

Regenerative endodontic procedures were compared with the apical barrier techniques in which conventional calcium hydroxide apexification and MTA were used and similar results were obtained [82, 83]. In a study comparing conventional and regenerative treatment protocols, it was reported that revascularization led to a significant increase in the length and thickness of the root and had a better prognosis compared to the apical barrier techniques including calcium hydroxide apexification and MTA [84]. Although MTA is the most commonly used material in regenerative endodontics, the discoloration of the front teeth poses a serious aesthetic problem. Therefore, other materials such as Biodentine may be preferred in the front teeth [85].
Vital pulp treatments

MTA is a material showing physicochemical properties that stimulate restorative dentinogenesis through the collection and activation of cells forming hard tissue material and contribute to matrix formation and mineralization [96]. It stimulates restorative hard tissue formation by separating growth factors in the extracellular matrix that mediate wound repair of the dentine-pulp complex and soluble cytokines embedded in the surrounding dentine [87-90].

MTA activates the migration of progenitor cells (fibroblasts) from the central pulp to the site of wound. It promotes the proliferation of progenitor cells without induction of cell apoptosis and the differentiation into odontoblast-like cells. MTA also stimulates the production of in vitro mRNA, increases the protein expression of mineralized matrix genes, which are very important for mineralization, and cellular markers [106].

In a study conducted using MTA, it was suggested that the tissue formed as a result of pulp capping procedure was not dentine or dentine-like tissue and that it might be dystrophic intrapulpal mineralization in response to pulp capping procedure [91]. The researchers also noted that this tissue had the same properties as the tissue observed following a direct pulp capping with calcium hydroxide.

The biocompatibility of MTA that is set regulates the expression of gene products such as dentin sialoprotein, osteocalcin and alkaline phosphatase, along with transcription and angiogenesis factors [92]. Odontoblast signal proteins are very important in the differentiation of progenitor cells into the cells responsible for the repair and accumulation of hard tissues [92, 93]. Sialoprotein and osteopontin were observed in the mineralized hard tissue matrix in the relevant area following the pulp capping with MTA. [94].

MTA shows superior marginal adaptation to dentine compared to other calcium hydroxide-based materials. The bond strength to dentine is quite high and it shows a bond similar to glass ionomer cement penetrating dentinal tubules thanks to its mineral components [95]. Moreover, another important property of the material is that it creates a sticky interface layer during mineral nucleation on the surface of dentine [86, 96, 97].

In a study, when MTA and calcium hydroxide were used in direct pulp capping, there was very low inflammatory response in the pulp treated with MTA, a thick and continuous dentine bridge appeared and, on the other hand, the dentine bridge was observed only in 1/3 of the samples treated with calcium hydroxide [98]. In another study, MTA used in direct pulp capping induced the matrix of osteodentine which is typically observed in reparative dentine in three weeks [99]. The formation of neodentine bridges was observed between pulpal tissues and MTA crystals in 2 weeks [100].

MTA materials used as pulp capping agents were reported to form calcified bridges in 2 weeks, exhibiting fully calcified bridge formation with mild inflammatory reactions [101].

MTA stimulates the release of dentine matrix components necessary for hard tissue repair and regeneration in healthy and partially inflamed pulps which are mechanically exposed [98, 102].

Data obtained from the studies in the literature generally suggest that MTA is a material promoting biocompatible, non-cytotoxic, antibacterial environment and surface morphology that is favorable for the formation of restorative calcific bridges.

When used in pulpotomy, MTA was reported to lead to less pulpal inflammation compared to the histological pulpal response caused by calcium hydroxide [100]. In a study conducted using MTA in the pulpotomy of 23 permanent teeth with clinical findings of irreversible pulpitis, positive results were obtained regarding prognosis [103].

Conclusion

MTA has a wide range of areas of use in clinical practice. In addition to its biocompatibility, it is a material that has promising properties such as sealing property, antibacterial property, and compressive strength. Although it was reported to show positive biological performance in the studies, there is a further need for a greater number of clinical research including long-term follow-up results.

References

1. Lee SJ, Monsef M, Torabinejad M. Sealing ability of a mineral trioxide aggregate for repair of lateral root perforations. Journal of endodontics 1993;19(11):541-544.
2. Lee D, Bogen G. Multifaceted use of Pro Root TM MTA root canal repair material. Pediatric Dent 2001;23(4):326-330.
3. Srinivasan V, Waterhouse P, Whitworth J. Mineral trioxide aggregate in paediatric dentistry. International Journal of Paediatric Dentistry 2009;19(1):34-47.
4. Hanna SN, Alfayate RP, Prichard J. Vital pulp therapy: an insight over the available literature and future expectations. European endodontic journal 2020;5(1):46.
5. Lopes MB, Soares VC, Fagundes FH, Gonini-Junior A, Kaneshima RH, Guiraldo RD, et al. Analysis of molecular changes induced by mineral trioxide aggregate on sPLA2. Brazilian dental journal 2019;30(5):453-458.
6. Torabinejad M, Hong C, McDonald F, Ford TP. Physical and chemical properties of a new root-end filling material. Journal of endodontics 1995;21(7):349-353.
7. Kadali NS, Alla KK, Guduri V, Ramaraju A, Sajjan S, Rudraraju VR. Mineral Trioxide Aggregate: An overview of composition, properties and clinical applications. International Journal of Dental Materials 2020;2(1):11-18.
8. Estrela C, Decurcio DdA, Rossi-Fedele G, Silva JA, Guedes OA, Borges ÂH. Root perforations: a review of diagnosis, prognosis and materials. Brazilian oral research 2018, 32.
9. Bakhtiar H, Nekoofar MH, Aminishakib P, Abedi F, Moosavi FN, Esnaashari E, et al. Human pulp responses to partial pulpotomy treatment with TheraCal as compared with Biodentine and ProRoot MTA: a clinical trial. Journal of endodontics 2017;43(11):1786-1791.
10. Asgary S, Parirokh M, Eghbal MJ, Brink F. Chemical differences between white and gray mineral trioxide aggregate. Journal of endodontics 2005;31(2):101-103.
11. Dammashcke T, Gerth HU, Züchner H, Schäfer E. Chemical and physical surface and bulk material characterization of white ProRoot MTA and two Portland cements. Dental Materials 2005;21(8):731-738.
12. Duarte MAH, de Oliveira Demarchi ACC, Yamashita JC, Kuga MC, de Campos Fraga S. pH and calcium ion release of 2 root-end filling materials. Oral Surgery, Oral Medicine, Oral Pathology, Oral Radiology, and Endodontology 2003;95(3):345-347.
13. Parirokh M, Torabinejad M. Mineral trioxide aggregate: a comprehensive literature review-part I: chemical, physical, and antibacterial properties. Journal of endodontics 2010;36(1):16-27.
14. Torabinejad M, Watson T, Ford TP. Sealing ability of a mineral trioxide aggregate when used as a root end filling
material. Journal of endodontics 1993;19(12):591-595.
15. Macwan C, Deshpande A. Mineral trioxide aggregate (MTA) in dentistry: A review of literature. Journal of Oral Research and Review 2014;6(2):71.
16. Kakani AK, Veeramachaneni C. Sealing ability of three different root repair materials for furcation perforation repair: An in vitro study. Journal of Conservative Dentistry: JCD 2020;23(1):62.
17. Arruda RA, Cunha RS, Miguita KB, Silveira CF, De Martin AS, Pinheiro SL, et al. Sealing ability of mineral trioxide aggregate (MTA) combined with distilled water, chlorhexidine, and doxycycline. Journal of oral science 2012;54(3):233-239.
18. Hosoya N, Takigawa T, Horie T, Maeda H, Yamamoto Y, Momoi Y, et al. A review of the literature on the efficacy of mineral trioxide aggregate in conservative dentistry. Dental materials journal 2019;38(5):693-700.
19. Sluyk S, Moon P, Hartwell G. Evaluation of setting properties and retention characteristics of mineral trioxide aggregate when used as a furcation perforation repair material. Journal of endodontics 1998;24(11):768-771.
20. Camilleri J, Montesin FE, Brady K, Sweeney R, Curtis RV, Ford TRP. The constitution of mineral trioxide aggregate. Dental Materials 2005;21(4):297-303.
21. Saghiri MA, Kazerani H, Morgano SM, Gutmann JL. Evaluation of Mechanical Activation and Chemical Synthesis for Particle Size Modification of White Mineral Trioxide Aggregate. European Endodontic Journal 2020;5(2):128.
22. Aksel H, Küçükay Eren S, Askerbeyli Ors S, Karaismailoğlu E. Surface and vertical dimensional changes of mineral trioxide aggregate and biodentine in different environmental conditions. Journal of Applied Oral Science 2019, 27.
23. Kogan P, He J, Glickman GN, Watanabe I. The effects of various additives on setting properties of MTA. Journal of endodontics 2006;32(6):569-572.
24. Prasad A, Pushpa S, Arunagiri D, Sawhny A, Misra A, Sujatha R. A comparative evaluation of the effect of various additives on selected physical properties of white mineral trioxide aggregate. Journal of Conservative Dentistry: JCD 2015;18(3):237.
25. Bortoluzzi EA, Broon NJ, Bramante CM, Felippe WT, Tanomaru Filho M, Esberard RM. The influence of calcium chloride on the setting time, solubility, discintegration, and pH of mineral trioxide aggregate and white Portland cement with a radiopacifier. Journal of endodontics 2009;35(4):550-554.
26. Tiwari N, Borkar AC, Tandale A, Nighot N, Ghare S, Maral S. Comparative evaluation of the effect of various endodontic irrigants on the push-out bond strength of endosequence, Biodentine™, and MTA Plus™ root repair materials: An in vitro study. Journal of the International Clinical Dental Research Organization 2019;11(1):9.
27. Jamshidy L, Amirkhani Z, Sharifi R. Effect of furcation perforation size on fracture resistance of mandibular first molar. Dental Hypotheses 2019;10(1):9.
28. Vajja S, Naik BD, Vummidiidetti SV, Yarlagadda V. Influence of Different Thickness of Mineral Trioxide Aggregate, Resin Modified Glass Ionomer Cement and Intermediate Restorative Material on Sealing Ability of Root End Fillings: An in vitro Study. Journal of Clinical & Diagnostic Research 2018;12(1).
29. Lamb EL, Loushine RJ, Weller RN, Kimbrough WF, Pasheley DH. Effect of root resection on the apical sealing ability of mineral trioxide aggregate. Oral Surgery, Oral Medicine, Oral Pathology, Oral Radiology, and Endodontology 2003;95(6):732-735.
30. Prathita T, Djauharie NK, Meidyawati R. Antimicrobial activity of mineral trioxide aggregate and calcium hydroxide sealer on enterococcus faecalis strain ATCC29212. International Journal of Applied Pharmaceutics 2019, 123-125.
31. Khedmat S, Aminipor M, Pourhajibagher M, Kharazifar MJ, Bahador A. Comparison of antibacterial activities of ProRoot MTA, OrthoMTA, and RetroMTA against three anaerobic endodontic bacteria. Journal of Dentistry (Tehran, Iran) 2018;15(5):294.
32. Al-Hezaimi K, Naghsibandi J, Oglesby S, Simon JH, Rotstein I. Comparison of antifungal activity of white-colored and gray-colored mineral trioxide aggregate (MTA) at similar concentrations against Candida albicans. Journal of endodontics 2006;32(4):365-367.
33. Kettering JD, Torabinejad M. Investigation of mutagenicity of mineral trioxide aggregate and other commonly used root-end filling materials. Journal of endodontics 1995;21(11):537-539.
34. Braz MG, Camargo E, Salvadori DMF, Marques M, Ribeiro D. Evaluation of genetic damage in human peripheral lymphocytes exposed to mineral trioxide aggregate and Portland cements. Journal of oral rehabilitation 2006;33(3):234-239.
35. Cin CP, Chen YJ, Lee YL, Wang JS, Chang MC, Lan WH, et al. Effects of root-end filling materials and eugenol on mitochondrial dehydrogenase activity and cytotoxicity to human periodontal ligament fibroblasts. Journal of Biomedical Materials Research Part B: Applied Biomaterials: An Official Journal of The Society for Biomaterials, The Japanese Society for Biomaterials, and The Australian Society for Biomaterials and the Korean Society for Biomaterials 2004;71(2):429-440.
36. Hajizadeh N, Madani ZS, Zabihi E, Golpour M, Zahedpasha A, Mohammadnia M. Effect of MTA and CEM on mineralization-associated gene expression in stem cells derived from apical papilla. Iranian endodontic journal 2018;13(1):94.
37. Bonson S, Jeansonne B, Lallier T. Root-end filling materials alter fibroblast differentiation. Journal of dental research 2004;83(5):408-413.
38. Rezende TMB, Sobrinho APR, Vieira LQ, da Costa Sousa MG, Kawai T. Mineral trioxide aggregate (MTA) inhibits osteoclastogenesis and osteoclast activation through calcium and aluminum activities. Clinical oral investigations 2020, 1-10.
39. Singla M, Verma KG, Goyal V, Jusuja P, Kakkar A, Ahuja L. Comparison of push-out bond strength of furcation perforation repair materials–Glass ionomer cement Type II, hydroxyapatite, mineral trioxide aggregate, and biodentine: An in vitro study. Contemporary clinical dentistry 2018;9(3):410.
40. Main C, Mirzayan N, Shabahang S, Torabinejad M. Repair of root perforations using mineral trioxide aggregate: a long-term study. Journal of endodontics 2004;30(2):80-83.
41. Roda RS. Root perforation repair: surgical and nonsurgical management. Practical procedures & aesthetic dentistry: PPAD 2001;13(6):467-472; quiz 474.
42. Ford TRP, Torabinejad M, McKendry DJ, Hong C-U,
Kariyawasam SP. Use of mineral trioxide aggregate for repair of furcal perforations. Oral Surgery, Oral Medicine, Oral Pathology, Oral Radiology, and Endodontontology 1995;79(6):756-763.

43. Holland R, Otoboni Filho JA, de Souza V, Nery MJ, Bernabé PFE, Junior ED. Mineral trioxide aggregate repair of lateral root perforations. Journal of endodontics 2001;27(4):281-284.

44. Tsesis I, Elbahary S, Venezuela NB, Rosen E. Bacterial colonization in the apical part of extracted human teeth following root-end resection and filling: a confocal laser scanning microscopy study. Clinical oral investigations 2018;22(1):267-274.

45. Ferris DM, Baumgartner JC. Perforation repair comparing two types of mineral trioxide aggregate. Journal of endodontics 2004;30(6):422-424.

46. Hamad HA, Tordik PA, McClanahan SB. Furbation perforation repair comparing gray and white MTA: a dye extraction study. Journal of endodontics 2006;32(4):337-340.

47. Torabinejad M, Higa RK, McKendry DJ, Ford TRP. Dye leakage of four root end filling materials: effects of blood contamination. Journal of endodontics 1994;20(4):159-163.

48. VanderWeele RA, Schwartz SA, Beeson TJ. Effect of blood contamination on retention characteristics of MTA when mixed with different liquids. Journal of endodontics 2006;32(5):421-424.

49. Simon S, Rilliard F, Berdal A, Machtou P. The use of mineral trioxide aggregate in one-visit apexification treatment: a prospective study. International Endodontic Journal 2007;40(3):186-197.

50. Sarris S, Tahmassebi JF, Duggal MS, Cross IA. A clinical evaluation of mineral trioxide aggregate for root-end closure of non-vital immature permanent incisors in children—a pilot study. Dental Traumatology 2008;24(1):79-85.

51. Scherer W, Dragoo M. New subgingival restorative procedures with Geristore resin ionomer. Practical periodontics and aesthetic dentistry: PPAD 1995;7(1 Suppl):1-4.

52. Dragoo MR. Resin-Ionomer and Hybrid-Ionomer Cements: Part II. Human Clinical and Histologic Wound Healing Responses in Specific Periodontal Lesions. International Journal of Periodontics & Restorative Dentistry 1997;17(1).

53. Behnia A, Strassler HE, Campbell R. Repairing iatrogenic root perforations. The Journal of the American Dental Association 2000;131(2):196-201.

54. Mozyńska J, Metlerski M, Lipski M, Nowicka A. Tooth discoloration induced by different calcium silicate–based cements: A systematic review of in vitro studies. Journal of endodontics 2017;43(10):1593-1601.

55. Lindskog S, Blomlöf L, Hammarström L. Repair of periodontal tissues in vivo and in vitro. Journal of Clinical Periodontology 1983;10(2):188-205.

56. Andreassen J, Munksgaard E, Redbo L, Rud J. Periodontal tissue regeneration including cementogenesis adjacent to dentin-bonded retrograde composite fillings in humans. Journal of endodontics 1993;19(3):151-153.

57. Thomson TS, Berry JE, Somerman MJ, Kirkwood KL. Cementoblasts maintain expression of osteocalcin in the presence of mineral trioxide aggregate. Journal of endodontics 2003;29(6):407-412.

58. Torabinejad M, Hong C-U, Ford TRP, Kariyawasam SP. Tissue reaction to implanted super-EBA and mineral trioxide aggregate in the mandible of guinea pigs: a preliminary report. Journal of endodontics 1995;21(11):569-571.

59. Torabinejad M, Hong C, Ford TP, Kettering J. Cytotoxicity of four root end filling materials. Journal of endodontics 1995;21(10):489-492.

60. Apaydin ES, Shahabang S, Torabinejad M. Hard-tissue healing after application of fresh or set MTA as root-end-filling material. Journal of endodontics 2004;30(1):21-24.

61. Holland R, de Souza V, Nery MJ, Otoboni Filho JA, Bernabé PFE, Dezan Jr E. Reaction of dogs’ teeth to root canal filling with mineral trioxide aggregate or a glass ionomer sealer. Journal of endodontics 1999;25(11):728-730.

62. Torabinejad M, Hong C-U, Lee S-J, Monsef M, Ford TRP. Investigation of mineral trioxide aggregate for root-end filling in dogs. Journal of endodontics 1995;21(12):603-608.

63. Torabinejad M, Ford TRP, McKenzie DJ, Abedi HR, Miller DA, Kariyawasam SP. Histologic assessment of mineral trioxide aggregate as a root-end filling in monkeys. Journal of endodontics 1997;23(4):225-228.

64. Holland R, de Souza V, Nery MJ, Otoboni Filho JA, Bernabé PF, Dezan Jr E. Reaction of rat connective tissue to implanted dentin tubes filled with mineral trioxide aggregate or calcium hydroxide. Journal of endodontics 1999;25(3):161-166.

65. Saunders WP. A prospective clinical study of periapical surgery using mineral trioxide aggregate as a root-end filling. Journal of endodontics 2008;34(6):660-665.

66. Jafari F, Jafari S, Etesamnia P. Genotoxicity, bioactivity and clinical properties of calcium silicate based sealers: a literature review. Iranian endodontic journal 2017;12(4):407.

67. Awawdeh L, Al-Qudah A, Hamouri H, Chakra RJ. Outcomes of vital pulp therapy using mineral trioxide aggregate or biodentine: a prospective randomized clinical trial. Journal of endodontics 2018;44(11):1603-1609.

68. Parirokh M, Torabinejad M, Dummer P. Mineral trioxide aggregate and other bioactive endodontic cements: an updated overview—part I: vital pulp therapy. International Endodontic Journal 2018;51(2):177-205.

69. Torabinejad M, Parirokh M, Dummer P. Mineral trioxide aggregate and other bioactive endodontic cements: an updated overview—part II: other clinical applications and complications. International Endodontic Journal 2018;51(3):284-317.

70. Camilleri J, Pitt Ford T. Mineral trioxide aggregate: a review of the constituents and biological properties of the material. International Endodontic Journal 2006;39(10):747-754.

71. Chang S-W, Lee S-Y, Kang S-K, Kum K-Y, Kim E-C. In vitro biocompatibility, inflammatory response, and osteogenic potential of 4 root canal sealers: Sealapex, Sankin apatite root sealer, MTA Fillapex, and iRoot SP root canal sealer. Journal of endodontics 2014;40(10):1642-1648.

72. Salles LP, Gomes-Cornelio AL, Guimarães FC, Herrera BS, Bao SN, Rossa-Junior C, et al. Mineral trioxide aggregate–based endodontic sealer stimulates hydroxyapatite nucleation in human osteoblast-like cell culture. Journal of endodontics 2012;38(7):971-976.

73. Aminoshariae A, Johnson WT, Kulidc JC, Tay F. Cohen’s Pathways of the Pulp. In: Berman LH, Hargreaves KM, editors. Obturation of then Cleaned and Shaped Root Canal System 2020, 317-320.

74. Cuesta A, Zea-Garcia JD, Londono-Zuluaga D, Angeles G, Santacruz I, Vallcorba O et al. Multiscale
understanding of tricalcium silicate hydration reactions. Scientific reports 2018;8(1):1-11.

75. Darvell B, Wu R. “MTA”-an hydraulic silicate cement: review update and setting reaction. Dental Materials 2011;27(5):407-422.

76. Lim E-S, Park Y-B, Kwon Y-S, Shon W-J, Lee K-W, Min K-S. Physical properties and biocompatibility of an injectable calcium-silicate-based root canal sealer: in vitro and in vivo study. BMC oral health 2015;15(1):1-7.

77. Mendes AT, Silva PBd, So Bb, Hashizume LN, Vivan RR, Rosa Rad, et al. Evaluation of physicochemical properties of new calcium silicate-based sealer. Brazilian dental journal 2018;29(6):536-540.

78. Siboni F, Taddei P, Zamparini F, Prati C, Gandolfi M. Properties of BioRoot RCS, a tricalcium silicate endodontic sealer modified with povidone and polycarboxylate. International Endodontic Journal 2017;50:e120-e136.

79. Urban K, Neuhaus J, Donnermeyer D, Schäfer E, Dammaschke T. Solubility and pH value of 3 different root canal sealers: a long-term investigation. Journal of endodontics 2018;44(11):1736-1740.

80. Walsh RM, He J, Schweitzer J, Opperman LA, Woodmansey KF. Bioactive endodontic materials for everyday use: a review. Gen Dent 2018;66(3):48-51.

81. Felippes W, Felippe M, Rocha M. The effect of mineral trioxide aggregate on the apicification and periapical healing of teeth with incomplete root formation. International Endodontic Journal 2006;39(1):2-9.

82. Alobaid AS, Cortes LM, Lo J, Nguyen TT, Albert J, Abu-Melha AS, et al. Radiographic and clinical outcomes of the treatment of immature permanent teeth by revascularization or apicification: a pilot retrospective cohort study. Journal of endodontics 2014;40(8):1063-1070.

83. Kahler B, Rossi-Fedele G, Chung N, Lin LM. An evidence-based review of the efficacy of treatment approaches for immature permanent teeth with pulp necrosis. Journal of endodontics 2017;43(7):1052-1057.

84. Jeeruphan T, Jantarat J, Yanpiset K, Suwannapun L, Khewsawai P, Hargreaves KM. Mahidol study 1: comparison of radiographic and survival outcomes of immature teeth treated with either regenerative endodontic or apexification methods: a retrospective study. Journal of endodontics 2012;38(10):1330-1336.

85. Bakhtiar H, Esmaeili S, Tabatabayi SF, Ellini MR, Nekoofar MH, Dummer PM. Second-generation platelet concentrate (platelet-rich fibrin) as a scaffold in regenerative endodontics: a case series. Journal of endodontics 2017;43(3):401-408.

86. Okiji T, Yoshiba K. Reparative dentinogenesis induced by mineral trioxide aggregate: a review from the biological and physicochemical points of view. International journal of dentistry 2009.

87. Koh E, Ford TP, Torabinejad M, McDonald F. Mineral trioxide aggregate stimulates cytokine production in human osteoblasts. In: Journal of Bone and Mineral Research; 1995:

88. Koh E, Torabinejad M, Pitt Ford T, Brady K, McDonald F. Mineral trioxide aggregate stimulates a biological response in human osteoblasts. Journal of Biomedical Materials Research: An Official Journal of The Society for Biomaterials and The Japanese Society for Biomaterials 1997;37(3):432-439.

89. Tomson PL, Lumley PJ, Alexander MY, Smith AJ, Cooper PR. Hepatocyte growth factor is sequestered in dentine matrix and promotes regeneration-associated events in dental pulp cells. Cytokine 2013;61(2):622-629.

90. Tziafas D, Pantelidou O, Alvanou A, Belibasakis G, Papadimitriou S. The dentinogenic effect of mineral trioxide aggregate (MTA) in short-term capping experiments. International Endodontic Journal 2002;35(3):245-254.

91. Dammaschke T, Nowicka A, Lipski M, Ricucci D. Histological evaluation of hard tissue formation after direct pulp capping with a fast-setting mineral trioxide aggregate (RetroMTA) in humans. Clinical oral investigations 2019;23(12):4289-4299.

92. Paranje A, Smoot T, Zhang H, Johnson JD. Direct contact with mineral trioxide aggregate activates and differentiates human dental pulp cells. Journal of endodontics 2011;37(12):1691-1695.

93. Paranje A, Zhang H, Johnson JD. Effects of mineral trioxide aggregate on human dental pulp cells after pulp-capping procedures. Journal of endodontics 2010;36(6):1042-1047.

94. Kuratate M, Yoshiha K, Shigetani Y, Yoshina N, Ohshima H, Okiji T. Immunohistochemical analysis of nestin, osteopontin, and proliferating cells in the reparative process of exposed dental pulp capped with mineral trioxide aggregate. Journal of endodontics 2008;34(8):970-974.

95. Kaup M, Dammann CH, Schäfer E, Dammaschke T. Shear bond strength of Biodentine, ProRoot MTA, glass ionomer cement and composite resin on human dentine ex vivo. Head & face medicine 2015;11(1):1-8.

96. Sarkar N, Caicedo R, Ritwik P, Moiseyeva R, Kawashima I. Physicochemical basis of the biologic properties of mineral trioxide aggregate. Journal of endodontics 2005;31(2):97-100.

97. Torabinejad M, Smith PW, Kettering JD, Ford TRP. Comparative investigation of marginal adaptation of mineral trioxide aggregate and other commonly used root-end filling materials. Journal of endodontics 1995;21(6):295-299.

98. Ford TRP, Torabinejad M, Abedi HR, Bakland LK, Kariyawasam SP. Using mineral trioxide aggregate as a pulp-capping material. The Journal of the American Dental Association 1996;127(10):1491-1494.

99. Choi S-H, Kang J-H, Koh J-T, Chang H-S, Hwang Y-C, Hwang I-N, et al. Effect of leptin on odontoblastic differentiation and angiogenesis: an in vivo study. Journal of endodontics 2019;45(11):1332-1341.

100. Chacko DV, Kurikose DS. Human pulpal response to mineral trioxide aggregate (MTA): a histologic study. Journal of Clinical Pediatric Dentistry 2006;30(3):203-209.

101. Kang C-M, Hwang J, Song JS, Lee J-H, Choi H-J, Shin Y. Effects of three calcium silicate cements on inflammatory response and mineralization-inducing potentials in a dog pulpotomy model. Materials 2018;11(6):899.

102. Ainehchi M, Eslami B, Ghanbariha M, Saffar A. Mineral trioxide aggregate (MTA) and calcium hydroxide as pulp-capping agents in human teeth: a preliminary report. International Endodontic Journal 2003;36(3):225-231.

103. Witherspoon DE, Small JC, Harris GZ. Mineral trioxide aggregate pulpotomies: a case series outcomes assessment. The Journal of the American Dental Association 2006;137(5):610-618.