Bioactive dentin replacement

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Abstract: Replacement of dentin with a material popularly marketed as Biodentine (Septodont, Saint Maur des Fossés, France), a new tricalcium silicate-based cement, has recently been commercialized and advertised as a bioactive material. Its clinical application and physical properties have been widely described. Biodentine, a new biologically active cement which has dentin-like mechanical properties and can be used as a dentin replacement in the tooth crown and root region. The cement consists mainly of a tri- and dicalcium silicate powder, which is mixed with an aqueous calcium chloride solution. As regards biocompatibility, long-term impermeability, antibacterial properties, induction of hard tissue regeneration, stability, low solubility, non-absorbability and ease of handling, Biodentine fulfills the requirements found in the literature for a material suitable for these purposes. On the basis of the good material properties of Biodentine, this cement is an interesting alternative to the conventional materials which were hitherto recommended. Biodentine can therefore confer advantages in day-to-day practice and with correct diagnosis contribute to the long-term maintenance of the vitality of the dental pulp and to the retention of teeth. However, little scientific data is available at present.

Keywords: biodentine, dentin, bioactive dentin replacement, tricalcium silicate-based cement, Portland cement

I. Introduction

A new bioactive cement, also called as smart dentin replacement and popularly known as Biodentine (Septodont, St. Maur des Fossés, France), was recently launched as a dentin substitute. It shares both its indications and mode of action with calcium hydroxide, but does not have its drawbacks.

Biodentine is a cement of the same class as MTA: this new calcium silicate-based material exhibits physical and chemical properties similar to those described for certain Portland cement derivatives. On the biological level, it is perfectly biocompatible [1] and capable of inducing the apposition of reactionary dentin by stimulating odontoblast activity [3] and reparative dentin, by induction of cell differentiation [2]. It is in effect a dentin substitute that can be used as a coronal restoration material (for indirect pulp capping), but can also be placed in contact with the pulp. Its faster setting time allows either immediate crown restoration [4], or to make it directly intraorally “functional” without fear of the material deteriorating.

Biodentine can be used both on the tooth crown and also in the region of the tooth root; in the crown region as a base, provisional seal, for deep cavities therapy, as a cervical filling, for direct and indirect pulp capping and in pulpotomies. In the region of the tooth root, Biodentine can be used for the treatment of perforations of the root canal or pulp chamber floor, due to internal and external absorption processes, for apexification and as a retrograde root canal filling material. On the one hand, Biodentine serves as a dentin replacement (in the crown region as a base), and on the other hand for maintaining the vitality of the dental pulp or stimulation of hard tissue regeneration, i.e. both tertiary dentin formation and also bone regeneration, e.g. after root end surgery [5,6].

Bioactive tricalcium silicate has several advantages over calcium hydroxide and mineral trioxide aggregates (MTA). The commercialized tricalcium silicate is different from the usual dental calcium silicate “Portland Cement” materials. The manufacturing process of the active biosilicate technology eliminates the metal impurities seen in the “Portland Cement” calcium silicates. The setting reaction is a hydration of tricalcium silicate which produces a calcium silicate gel and a calcium hydroxide. In contact with phosphate ions, it creates precipitates that resemble hydroxyapatite [7,8]. These precipitates from MTA and tricalcium silicate can be incorporated into root canal dentin. A comparison of the calcium and silica uptake of adjacent root canals treated with MTA versus tricalcium silicate demonstrated a greater uptake of the tricalcium silicate [9]. Using confocal microscopy there was an increase in the carbonate content of interfacial dentin, which suggested intertubular diffusion and mineral tags of Biodentine hydration products creating a hybrid zone [10]. Burgess and co-workers [11] characterized this hybrid zone as being microleakage free.

Histologically, the bioactive tricalcium silicate demonstrated the ability to induce odontoblast differentiation from pulp progenitor cells. The resulting mineralized matrix had the molecular characteristics of dentin. An evaluation comparing the biocompatibility of the tricalcium silicate with MTA and Dycal...
demonstrated that the Biodentine was equivalent to MTA (Dentsply) and more biocompatible than Dycal (Dentsply-Caulk). A clinical evaluation over 6-35 months of Biodentine as a base and for pulp capping demonstrated both biocompatibility and longevity [1,7,8,12]. This review will cover all the aspects of this novel material to be used in dentistry.

II. Active Biosilicate Technology And Chemical Composition

This material was developed based on the most biocompatible chemistry available for dental materials: calcium silicates, which can set in the presence of water. Although recognized as highly biocompatible and bioactive, all these materials lack reactivity, with very long setting times (more than 2 hours), low mechanical properties and with very difficult handling (depending on the water ratio, from a sandy consistency to a fluid paste). This calcium silicate chemistry was combined with the requirements of a formulation compatible with classical restorative and endodontic practice. Septodont developed a new technological platform called Active Biosilicate Technology™. This consists in controlling every step of the material formulation beginning with the purity of the raw materials.

Usual dental calcium silicate cements are based on the “Portland Cement” materials, which result from the clinker products manufactured by the building industry from natural stone treatment. This implies that all these products inherently contain unpurifiable mixtures of calcium silicates(C₃S+C₃S), calcium aluminates (C₃A), calcium alumino-ferrites(C₄AF), calcium sulfates (CaSO₄-gypsum), together with low concentrations of metallic impurities coming from the natural minerals used as raw materials.[13,14] The Active Biosilicate Technology is a proprietary technology developed according to state-of-the-art pharmaceutical background applied to the high temperate ceramic mineral chemistry to ensure the purity of the calcium silicate content of the formulation and the absence of any aluminate and calcium sulphate in the final product. In order to reach a formulation with a short setting time (12 minutes) and high mechanical properties in the range of natural dentine, calcium silicates could not be used alone.[13,14]

Usually calcium silicate cements have setting times in the range of several hours, which is too long in most of the protocols in clinical practice. Increasing the setting time was achieved by a combination of different effects. First, particle size greatly influences the setting time, since the higher the specific surface, the shorter the setting. Also, adding calcium chloride to the liquid component accelerates the system. Finally, the decrease of the liquid content in the system decreases the setting time to harden within 9 to 12 minutes. Reaching high mechanical strength is also quite difficult for these systems. The first cause of low mechanical properties of Portland cements are the aluminate components, which make the product fragile. The manufacturer controls the purity of the calcium silicate through the Active Biosilicate Technology™ which consists in eliminating aluminates and other impurities [13].

The second axis of formulation was to adjust the particle size distribution in order to reach an optimal Powder density. The additional charge system selected was calcium carbonate, for both its biocompatibility and calcium content. The paradox of calcium silicate systems is also that water, which is essential for the hardening of the product, canal so affect the strength of the material. On the hand, excess water in the system will create some remaining porosity, significantly degrading the macroscopic mechanical resistance, but on the other hand decreasing the water content leads to reducing the possibility of a homogeneous mix. The addition of hydro soluble polymer systems described as “water reducing agents” or super plasticizers, helps in maintaining the balance between low water content and consistency of the mixture. Radiopacity is obtained by adding zirconium oxide to the final product [13,14].

Biodentine consists of a powder in a capsule and liquid in a pipette. The powder mainly contains tricalcium and dicalcium silicate, the principal component of Portland cement, as well as calcium carbonate. Zirconium dioxide serves as contrast medium. The liquid consists of calcium chloride in aqueous solution with an admixture of poly-carboxylate. The powder is mixed with the liquid in a capsule in the triturator for 30 seconds. Once mixed, Biodentine sets in about 12 minutes. During the setting of the cement calcium hydroxide is formed.[13,14]

**Powder**

<table>
<thead>
<tr>
<th>Component</th>
<th>Function</th>
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<tbody>
<tr>
<td>Tri-calcium Silicate (C₃S)</td>
<td>Main core material</td>
</tr>
<tr>
<td>Di-calcium Silicate (C₂S)</td>
<td>Second core material</td>
</tr>
<tr>
<td>Calcium Carbonate and Oxide</td>
<td>Filler</td>
</tr>
<tr>
<td>Iron Oxide</td>
<td>Shade</td>
</tr>
<tr>
<td>Zirconium Oxide</td>
<td>Radiopacifier</td>
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</table>
III. Setting reaction (2)

The calcium silicate has the ability to interact with water leading to the setting and hardening of the cement. This is a hydration of the tricalcium silicate (3CaO·SiO2 = C3S) which produces a hydrated calcium silicate gel (CSH gel) and calcium hydroxide (Ca(OH)2).

\[
2(3\text{CaO} \cdot \text{SiO}_2) + 6\text{H}_2\text{O} = 3\text{CaO} \cdot 2\text{SiO}_2 \cdot 3\text{H}_2\text{O} + 3\text{Ca(OH)}_2
\]

\[\text{C}_3\text{SCSH}\]

This dissolution process occurs at the surface of each grain of calcium silicate. The hydrated calcium silicate gel and the excess of calcium hydroxide tend to precipitate at the surface of the particles and in the pores of the powder, due to saturation of the medium. This precipitation process is reinforced in systems with low water content.

The unreacted tricalcium silicate grains are surrounded by layers of calcium silicate hydrated gel, which are relatively impermeable to water, thereby slowing down the effects of further reactions. The C-S-H gel formation is due to the permanent hydration of the tricalcium silicate, which gradually fills in the spaces between the tricalcium silicate grains. The hardening process results from the formation of crystals that are deposited in a supersaturated solution. The working time of Biodentines is up to 6 minutes with a final set at around 10-12 minutes.

Biodentine has a consistency after mixing which enables manipulation with a spatula, with an amalgam carrier or with carriers which are used for endodontic cements in retrograde fillings (Messing gun, MTA gun).

IV. Physical And Mechanical Properties

1. Density: 2.260(0.002) g/cm³

2. Porosity: 6.8 (0.2)%. 

3. Compressive strength:

There is a sharp increase in the compressive strength reaching more than 100 MPa in the first hour. The mechanical strength continues to improve to reach more than 200 MPa at 24h which is more than most glass ionomers values. A specific feature of Biodentine is its capacity to continue improving with time over several days until reaching 300 MPa after one month. This value becomes quite stable and is in the range of the compressive strength of natural dentin (297 MPa). This maturation process can be related to the decrease of porosity with time, which was illustrated previously. Biodentine is an evolutive biomaterial which improves its mechanical properties with time. This demonstrates the superiority of Biodentine for building in short time (9-12min) sufficient mechanical resistance to be used as a dentine substitute, compatible with dental restorations.[3,6,13,14]

4. Flexural strength: 22 MPa

5. Vickers micro hardness: The hardness increases in time when cements are immersed in distilled water. After 2 hours, the hardness of Biodentine was 51 HVN and reached 69 HVN after 1 month. The reported micro hardness values for natural dentin are in the range of 60-90 HVN (O’Brien 2008). Biodentine has surface hardness in the same range as natural dentin [3,6,13-15].

6. Radiopacity: displays a radiopacity equivalent to 3.5 mm of aluminium [13,14].

7. Stability in oral Enviroment: The material properties of Biodentine are similar to those of dentin. Both the elasticity modulus of the cement and also the pressure resistance, bending strength and Vickers hardness are comparable with dentin and, except for the Vickers hardness, lie above the values that can be measured on glass ionomer cements. However, Biodentine is not as stable as a composite material, so that Biodentine is not suitable for a permanent enamel replacement. However, in comparison to other Portland cement-based products, Bio dentin™ is stable enough to find use as a temporary filling even in the chewing load bearing region [3,6,13-15].
8. **Induces the formation of hard tissue**: Tricalcium silicate is a main component of Biodentine, MTA and Portland cement. In addition to their biocompatibility, materials of this type are known to be biologically active (Laurent et al. 2009). Biological activity refers to a positive effect of a medicine or material on living tissue. A material is described as biologically active if it interacts with cells of the human body or has an advantageous action on cells [3,6,13-15]. In studies of bio-mineralisation, bioactivity mostly refers to the promotion of hard tissue formation which is induced by a material. As regards the biological activity of Biodentine, it could be shown in vitro that dental pulp fibroblasts form so-called mineralisation nuclei after the cement has been added to the cell medium. These mineralisation nuclei have the molecular characteristics of dentin. This indicates a promotion of the transformation of the dental pulp fibroblasts by Biodentine to odontoblast-like cells which can then form hard tissue.

9. **Bacteriostatic**: During the setting phase of Biodentine, calcium hydroxide ions are released from the cement. This results in a pH of about 12.5 and a basification of the surroundings. This high pH inhibits the growth of microorganisms and can disinfect the dentin [16].

**V. Indications**

Biodentine, the first biocompatible and bioactive material can be used wherever dentine is damaged. Biodentine uniqueness not only lies in its innovative bioactive and “pulp-protective” chemistry, but also in its universal application, both in the crown and in the root.

**Dentine substitute under a composite** [3,6,13-15]
- temporary enamel restoration,
- permanent dentin restoration,
- deep or large carious lesions,
- deep cervical or radicular lesions,
- pulpotomy

**Endodontic repair material** [3,6,13-15]
The endodontic indications of Biodentine are similar to the usual calcium silicate based materials, like the Portland cements (i.e.ProRoot MTA).This type of product is already well documented. Several physical, chemical and biological properties are comparable. However, Biodentine has some features which are superior to MTA.

Biodentine consistency is better suited to the clinical use than MTA’s.

Biodentine presentation ensures a better handling and safety than MTA.

Biodentine does not require a two step obturation as in the case of MTA. As the setting is faster, there is a lower risk of bacterial contamination than with MTA.[3,6,13-15]

- Direct pulp capping following carious pulp exposure
- Direct pulp capping following dental trauma/injury to healthy pulp(partial pulpotomy)
- Repair of perforated root canals and/or pulp chamber floor
- Retrograde endodontic surgery
- Pulpotomy in primary molars
- Apexification

**VI. Benefits Of Biodentine** [1,12-15]
- Versatile: Usable for pulp caps to bulk fill, does not stain, has excellent radiopacity, and there is no surface preparation or tedious bonding required due to the micro-mechanical anchorage.
- Saves teeth: Has higher compressive strength than dentin, preserves pulp, and promotes pulp healing - eliminating RCT (root canal therapy) in most cases.
- Excellent sealing properties: Outstanding microleakage resistance, enhanced by the absence of shrinkage due to the resin-free formula.
- Sets in ten minutes: Allows full restorations to be completed in one office visit! Proven clinical results.
- No surface preparation: micro-mechanical anchorage, no tedious bonding is required.
- Excellent sealing properties: mineral tags in the dentin tubules.
Outstanding microleakage resistance, enhanced by the absence of shrinkage thanks to the resin-free formula.
High dimensional stability: long lasting sealing properties.
Biodentine prevents caries recurrence in deep cavities thanks to its alkaline pH (pH=12) giving it bacteriostatic properties.
Excellent radiopacity

VII. Critical Analysis
The studies mentioning the role of this new material are discussed and critically analysed.

<table>
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<tr>
<th>Authors (year)</th>
<th>Analysis of the Inference</th>
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<tr>
<td>Shamkalov GSet al, 2013 [17] In vitro</td>
<td>Examined materials (&quot;Biodentine&quot; (Septodont), &quot;Rootdent&quot; (TehnoDent) and adhesive &quot;Futurabond HP&quot; (Voco)) for antibacterial activity against E. coli, S. aureus, C. albicans, Str. faecalis. The results confirmed the analyzed materials to be a useful tool for deep caries lesions treatment.</td>
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<tr>
<td>Camilleri J., 2013 [18]</td>
<td>Biodentine demonstrated both structural and chemical changes when etched with 37% phosphoric acid. Biodentine exhibited a lower calcium to silicon ratio and a reduction in the chloride peak height when etched. When used as a dentine replacement material in the sandwich technique overlayed with composite, significant leakage occurred at the dentine to material interface. On the other hand materials based on glass ionomer cement were etched successfully and no chemical and physical changes or micro-leakage were detected when the materials were used as bases under composite restorations. The micro-hardness of all the materials was unaffected by etching.</td>
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<tr>
<td>Nowicka A et al, 2013 [19]</td>
<td>The majority of specimens showed complete dentinal bridge formation and an absence of inflammatory pulp response. Layers of well-arranged odontoblast and odontoblast-like cells were found to form tubular dentin under the osteodentin. Within the limitations of this study, Biodentine had a similar efficacy in the clinical setting and was considered an interesting alternative to MTA in pulp-capping treatment during vital pulp therapy.</td>
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<td>Vallès Met al, 2013 [20] In vitro</td>
<td>The combination of light and anaerobic conditions (similar to those in clinical situations) results in differences in color of the tested calcium silicate-based materials (CSMs) during a period of 5 days, of which Biodentine and White Portland Cement (PC) demonstrated color stability.</td>
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<tr>
<td>Zhou HMet al, 2013 [21] In vitro</td>
<td>Cells exposed to extracts from Biodentine and MTA showed the highest viabilities of human gingival fibroblasts. at all extract concentrations, whereas cells exposed to glass ionomer cement extracts displayed the lowest viabilities. Biodentine caused gingival fibroblast reaction similar to that by MTA. Both materials were less cytotoxic than glass ionomer cement.</td>
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<td>Han Let al, 2013 [22] In vitro</td>
<td>Compared white ProRoot MTA (WMTA), EndoSequence BC sealer (BC sealer) and Biodentine with regard to their ability to produce apatites and cause Ca and Si incorporation in adjacent human root canal dentine after immersion in phosphate-buffered saline (PBS). Compared with Biodentine and WMTA, BC sealer showed less Ca ion release and did not show Ca and Si incorporation as deeply in human root canal dentine when immersed in PBS for up to 90 days.</td>
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<tr>
<td>Grecch Let al, 2013 [23] In vitro</td>
<td>The hydrated materials were composed of a cementitious phase that was rich in calcium and silicon and a radiopacifying material. Biodentine included calcium carbonate, and Bioaggregate included silica and calcium phosphate in the powders. IRM was composed of zinc oxide interspersed in a matrix of organic material.</td>
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<td>Shayegan A et al, 2012 [24] In vivo</td>
<td>Assessed and compared, in primary pig teeth, the pulp response after a pulpotomy using either Biodentine (a new tricalcium-silicate cement), white mineral trioxide aggregate (WMTA), or formocresol (FC) and repeat the same after direct pulp capping using either Biodentine, WMTA, or calcium hydroxide. They concluded that Biodentine and white mineral trioxide aggregate are both suitable, biocompatible materials for pulp capping in primary teeth of pigs.</td>
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<tr>
<td>Grecch Let al, 2013 [25] In vitro</td>
<td>The physical properties of prototype radiopacified tricalcium silicate cement, Bioaggregate and Biodentine were investigated. Intermediate restorative material was used as a control. All the materials tested had a radiopacity value higher than 3mm thickness of aluminum. IIRM exhibited the highest radiopacity. Biodentine demonstrated a high washout, low fluid uptake and sorption values, low setting time and superior mechanical properties. The fluid uptake and setting time was the highest for Bioaggregate.</td>
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<tr>
<td>Tran XV et al, 2012 [26] In vitro</td>
<td>Evaluated the capacity of a new calcium-silicate-based restorative cement (biodentine) as compared to calcium hydroxide and MTA to induce pulp healing in a rat pulp injury model. At day 7, results showed that both the evaluated cement and MTA induced cell proliferation and formation of mineralization foci, which were strongly positive for osteopontin. At longer time-points, we observed the formation of a homogeneous dentin bridge at the injury site, secreted by cells displaying an odontoblastic phenotype. In contrast, the reparative tissue induced by Ca(OH)₂ showed porous organization, suggesting a reparative process different from those induced by calcium silicate</td>
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VIII. Conclusion

Overall, Biodentine is an interesting, very promising product, which with correct diagnosis can certainly contribute to a high degree to maintenance of the vitality of the dental pulp or to the retention of a tooth. Unfortunately at present little scientific published data on Biodentine is available. More scientific studies on Biodentine are therefore absolutely necessary.

References


[7]. Atmeh AR. A Dynamic bioactive interface with dental tissues. 45th Meeting of the Continental European Division of the IADR (CED-IADR) with the Scandinavian Division (NOF). 2011; Abstract no. 1.


Bioactive dentin replacement

[16]. Flrla M.; Dentin-Ersatzmaterial auf der Basis der Active Biosilicate Technology. DZW Kompakt 2011;14: (1) 11-14.