Bioactive Glasses in the Management of Dentine Hypersensitivity: A Review

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Abstract
Dentine Hypersensitivity is a common clinical condition, albeit of a low severity, and various in vitro and in vivo studies have been performed to test various approaches in managing the condition. This review investigates the use of bioactive glasses and their efficacy in treating Dentine Hypersensitivity. Significant progress was observed in the introduction of bioactive glasses in previous toothpaste formulations due to its ability to produce a Hydroxyapatite-like layer. Nevertheless, the results of the present review would suggest that a higher quality evidence was required to sufficiently support the use and effectiveness of bioactive glasses in treating Dentine Hypersensitivity. This observation is particularly pertinent in the absence of evidence relating to the effect of abrasivity of the glasses as well as the ability of strontium incorporation into the glasses. And the exact loading of the glass into dentifrice formulations. Overall, in vitro studies do appear to demonstrate that bioactive glass formulations may be an effective material to occlude dentine tubules which may in turn, reduce the fluid flow within the dentinal tubules and subsequently help manage Dentine Hypersensitivity.

Keywords
Dentine Hypersensitivity, Tubular occlusion, Bioactive Glasses, Review

Introduction

Dentine Hypersensitivity (DH) is a common clinical condition which may affect up to 74% of the population [1]. DH can be defined as a "short, sharp pain arising from exposed dentine in response to stimuli typically thermal, evaporative, tactile, osmotic or chemical and which cannot be ascribed to any other form of dental defect or pathology" [2]. The condition is generally prevalent in the age 20-49 year group and may affect the facial or buccal surfaces of incisors, premolars and canines, although molars may also be affected. From a clinical perspective the loss of the overlying enamel and/or cementum of the premolars and canines, although molars may also be affected. From a clinical perspective the loss of the overlying enamel and/or cementum of the premolars and canines, although molars may also be affected.

It was previously reported that not only does Bioglass® bind to bone but it also facilitates the regeneration of bone. Initially the bone substitute was implanted into humans as a replacement for the ossicles of the middle ear and was subsequently used as a material to repair bone defects eventually including those caused by periodontitis as used desensitising toothpaste to alleviate DH [4,5]. Currently treatment approaches are based on the Brännström hydrodynamic theory which required the dentine tubules to act as a capillary bore with the tubeule open to the oral environment [6]. According to this theory a stimuli (e.g., cold air) resulted in the movement of the fluid within the dentine tubules which can be sensed by mechanical receptors near the pulp which in turn transmits the stimulus via Aδ and Aβ fibres triggering a short, sharp pain [7]. A number of in vivo and in vitro studies have reported that in subjects who have either been identified as responding/not responding to a stimulus (through a clinical examination) that there was a significant difference in the appearance of the surfaces of the teeth. For example, die replicas from the clinical impressions of the teeth identified as being sensitive, when viewed under SEM indicated that there were up to 8x more patent tubules per unit area in hypersensitive dentine compared to non-sensitive dentine. These tubules also had a wider aperture (X2) compared to non-sensitive dentine [8]. According to West et al. there was no evidence of any differences between coronal and radicular dentine hypersensitivity, although it was reported that in cervical root dentine there were fewer tubules with narrower apertures compared to mid coronal dentine [6]. Joshi et al. also reported that the hydraulic conductance of coronal dentine was much higher than in radicular dentine [8]. The evidence from conducting in vitro hydraulic conductance measurements would suggest that successfully occluding the dentinal tubules would reduce the dentinal fluid flow. Although there a number of professionally applied and At-Home products that are available for the treatment of DH, there is no current acceptance as to a gold standard product to treat DH [5]. More recently there has been renewed interest in the use of bioactive glass initially developed by Hench to incorporate into toothpaste formulations [9]. The aim of this review was to evaluate the available published literature to determine whether there is sufficient evidence for the effectiveness of bioactive glass formulations in the treatment of DH.

The discovery of bioactive glasses by Hench introduced novel synthetic materials that were capable of bonding to living tissues [9]. Prior to this point, synthetic materials were designed to interact as little as possible with the human body and this often resulted in the formation of a non-adherent fibrous layer which encapsulated the implanted material [10]. Hench subsequently defined bioactive materials as those materials which may instigate a biological response at the interface of the material with a subsequent bond forming between the tissue and material. The key differences between bioactive glasses (BG’s) and traditionally soda-lime glasses lie in 3 main features namely 1) Less than 60 mol % of Silica, 2) High levels of Na₂O and CaO, 3) High CaO/P₂O₅ ratio [11]. The original Bioglass® composition known as 45S5 was developed as a bone substitute and not as toothpaste formulation and contained 45% SiO₂, 24.5% Na₂O, 24.5% CaO, and 6% P₂O₅. According to Hench there are two main classifications of bioactive glasses (Table 1) [12].

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well [13]. There are four main methods by which genes involved with osteoblastic proliferation, differentiation, and cell to cell adhesion are upregulated namely: surface chemistry, topography, rate and type of dissolution ions and shear stress at the interface. As a result the rates of osteogenesis, angiogenesis, chondrogenesis, and differentiation of mesenchymal stem cells are increased. However, more advanced studies are required to fully appreciate the effect(s) of the bioactive glass on cell gene expression [14]. Although an increase in the transcription rates of a gene may be recorded for example of vascular endothelial growth factor, its physiological relevance is, however not fully known. Moreover, increased expression does not inherently correspond to increased protein production since post-translational mechanisms may result in a different culmination to that expected.

Further advantageous features of bioactive glasses include antimicrobial properties which may inhibit bacterial growth at the site of implantation, and further oral applications include pulp capping, sinus obliteration, repair of orbital floor fracture, coating for dental implants, topical endodontic disinfectants as well as being used in air-abrasion [15,16].

Therapeutic ions, drugs and growth factors

The original 45S5 Bioglass® contained 45% SiO₂, 24.5% Na₂O, 24.5% CaO, and 6% P₂O₅, however, subsequently more research has been conducted into incorporating various therapeutic ions, drugs and growth factors into the glass structure. Table 2 outlines a variety of products that have been incorporated into the glass structure to provide a beneficial effect [13,15].

Additionally, the BG’s are anti-bacterial in nature due to the pH rise associated with cation release during the surface reactions. For example, pathogens associated with both caries and periodontitis including Streptococcus mutans and Actinobacillus actinomycetemcomitans were demonstrated to be killed during in vitro studies which may demonstrate its potential anti-caries effect in toothpaste [17].

Despite the degree of research, there has been limited data supporting the clinical significance of the incorporation of these products and the use of BG as a method to deliver therapeutic ions, drugs or growth factors is still in its early stage of development.

Bioactive glasses and dentine hypersensitivity

According to the Hydrodynamic theory the resolution of any discomfort from DH would be based on the occlusion of the open dentinal tubules which would then limit or restrict the fluid flow through dentine. Since fluid flow is directly proportional to the fourth power of the radius as indicated by Poiseuille’s law, reduction of the radius of the tubule would subsequently reduce fluid flow and therefore reduce the stimulation of the mechanical receptors and nerve fibres [18]. Thus, halving the tubule radius would result in a 16 times decrease in fluid flow in the tubule.

Table 1: Classification of BG’s (Hench; 1998)

| Class A Bioactive glasses | Results in osteoconduction and osteoproduction. Binds to both soft and hard tissues. |
| Class B Bioactive glasses | Osteoconductive and only able to bind to hard tissues and not soft tissues. |

<table>
<thead>
<tr>
<th>Ion, Drug or Growth Factor</th>
<th>Effect</th>
</tr>
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<tbody>
<tr>
<td>Fluoride</td>
<td>Formation of Fluorapatite which is more chemically stable than HCA at lower pH values and so more acid resistant. Fluoride containing BG result in increased alkaline phosphatase activity, an indication of increased bone mineralisation. Nevertheless, low fluoride containing bioactive glasses demonstrated the greatest cell proliferation Inhibits bacterial enzymes. Stimulates osteoblasts at a concentration of 25–500 ng ml⁻¹, whilst over 500 ng ml⁻¹ inhibits osteoblast activity.</td>
</tr>
<tr>
<td>Strontium</td>
<td>Improves demineralized dentine mineral density. Increases osteoblast activity and inhibits osteoclast activity.</td>
</tr>
<tr>
<td>P₂O₅</td>
<td>Favours FAP formation instead of CaF₂ Assists in the maintenance of network connectivity.</td>
</tr>
<tr>
<td>Zinc</td>
<td>Anti-inflammatory and anti-microbial. Promotes osteoblast activity and inhibits osteoclast activity</td>
</tr>
<tr>
<td>Silver</td>
<td>Anti-bacterial</td>
</tr>
<tr>
<td>Lithium</td>
<td>Promotes osteogenesis and cementogenesis</td>
</tr>
<tr>
<td>Copper</td>
<td>Anti-bacterial and promotes osteogenesis and angiogenesis</td>
</tr>
<tr>
<td>Iron</td>
<td>Promote osteogenesis</td>
</tr>
<tr>
<td>Dexamethasone</td>
<td>Promote osteogenesis</td>
</tr>
<tr>
<td>Vascular Endothelial Growth Factor</td>
<td>Promote angiogenesis</td>
</tr>
<tr>
<td>Ampicillin</td>
<td>Anti-bacterial</td>
</tr>
<tr>
<td>Doxorubicin</td>
<td>Anti-cancer drug. Inhibits activity of osteosarcoma cells.</td>
</tr>
</tbody>
</table>

Table 2: Ion, Drug or Growth Factor and their Effects.
Consequently, there are two main mechanisms through which DH may be managed. The first mechanism has been proposed to operate via Potassium ions, e.g., Potassium Nitrate (KNO₃), Potassium Chloride or Potassium Citrate and as such does not involve tubule occlusion or affect fluid movements but rather targets the excitation of the nerve fibres. The increase in concentration of extracellular Potassium around the nerve fibres prevents their repolarization thereby blocking their excitation by inactivating their action potential [19]. The second method involves occluding the exposed dentinal tubules to produce a similar effect by preventing any stimuli initiating fluid flow to trigger the mechanoreceptors. Abi et al. and Yoshiyama et al. identified that most dentine tubules were occluded in the non-sensitive dentine in the replica dies of patients examined clinically [20,21]. BG toothpaste formulation would therefore be through this method of occluding tubules to relieve DH. The negative surface charge of the glass formed during its contact with water or saliva enables it to bind to the Type 1 collagen fibres found in dentine tubules. The deposition of the HCA layer that forms following the surface reactions of the glass as well as the presence of residual BG particles result in the occlusion of the tubules. Gillam et al. stated that to treat DH via tubule occlusion was the logical conclusion from Brännström’s hydrodynamic theory [22].

**In vitro studies**

There is *in vitro* data supporting the ability of bioactive glasses, currently marketed under Sensodyne® toothpaste as Novamin®, to occlude dentinal tubules. For example, Gillam et al. demonstrated via SEM micrographs that the original Bioglass® was capable of partially occluding the tubules. However, more interestingly, it showed that when bioactive glasses were used to form new toothpaste by replacing the silica component to different degrees, it was consistently more effective in occluding tubules [23]. Tests were conducted on dentine discs from caries-free third molars comparing new toothpaste compositions containing 0%, 2.5% and 7.5% bioactive glass particles. Silica is the standard filler in most toothpastes, thus the replacement of silica with BG particles ensured the validity of the study as any differences demonstrated in the SEM images of the 2.5% and 7.5% toothpastes compared to the 0% BG control must be due to the BG particles. All compositions showed a greater tubule occlusion than the original Bioglass® formulation with increased bioactive glass levels corresponding to significantly greater tubule occlusion due to increased particle deposition on the surface and within the tubules [23].

Particulate Bioglass® has been used for periodontal osseous repair using a size range of 70–910 μm and 300–355 μm and this original 4555 Bioglass® was initially used in the Gillam et al. study to determine its ability to occlude the dentinal tubules [24,25]. However it was easily displaced by rinsing and handling and subsequently the investigators incorporated the glass into standard toothpaste formulations to determine its effectiveness in occluding the tubules. Test dentine disc halves were treated with two Over-the-Counter toothpastes; Macleans Freshmint – a calcium carbonate-based paste (Smithkline Beecham, now GSK) and Elmex Amine Fluoride – an abrasive free paste (GABA, now Colgate), mixed in a ratio 5:1, toothpaste with the silica component to different degrees, it was consistently more resistant to acid/abrasive challenges although currently this concept has not been proven.

Further studies have demonstrated the benefits of different ionic compositions of bioactive glasses in its use as a dentifrice. Brauer et al. reported that fluoride containing glasses resulted in Fluorapatite (FAP) formation which is more acid resistant than HCA due to its lower Ka value which would suggest that incorporating bioactive glasses would provide a more effective dentifrice for treating DH [30]. However, these investigators reported that high fluoride containing glasses mainly formed CaF₂ in simulated body fluid (SBF). It was hypothesised that increasing the phosphate concentration would increase the Fluorapatite formation instead of CaF₂ formation. This was confirmed by Mneimne et al. who compared a 6 mol% phosphate BG to a 1 mol % phosphate BG and reported that increased phosphate content did favor Fluorapatite instead of CaF₂ formation in a Tris buffer solution [31]. Furthermore, it was reported that an increased phosphate levels resulted in a much more rapid apatite formation, within 6 hours for the 6 mol % phosphate compared to within 3 days for the 1 mol % phosphate concentration. A quicker rate of deposition would theoretically provide a faster relief from DH, and a greater acid resistance would lead to reduced loss of the apatite layer. It was suggested therefore that the use of a high phosphate content and fluoride containing BG would effectively provide a Fluorapatite...
formation which in turn would be more acid resistant than those dentifrices forming CaF$_2$ only. O’Donnell et al. further supported these results in a study and explained these results by suggesting that by increasing the phosphate levels would in turn reduce the pH to a more near optimum level of 7.2 pH which is ideal for apatite deposition [32]. Mohammed et al. refuted the claims of previous studies that CaF$_2$ formation was beneficial due to its ability to exhibit an anti-caries effect; form a physical barrier on enamel surface; reduce rate of demineralization and act as a fluoride reservoir [33]. Instead these investigators reported that CaF$_2$ formation reduced the Ca$^{2+}$ available for remineralization which further resulted in loss of PO$_4^{3-}$ and concluded that CaF$_2$ formation was “potentially detrimental” as it prevents apatite formation. It was further noted that Strontium as well as Fluoride, both at concentrations 45 ppm were capable of inhibiting the CaF$_2$ formation. However, the study only used static models not dynamic cyclic models which would have better mimicked the oral environment. Moreover, an increase in Ca$^{2+}$ and PO$_4^{3-}$ deposition may also lead to an increase in calculus formation. Nonetheless, the addition of Fluoride to BG’s was concluded to be beneficial to oral health.

**Abrasivity**

Ali et al. identified that BG used in a dentifrice e.g. Novamin based on the 45S5 glass was significantly harder than enamel and dentine (Table 3) [15]. This data, together with other data would suggest that some BGs could have a Mohs hardness value of 7 GPA or more. However, this figure should ideally be below 3.5 GPA, which is the hardness of enamel, in order to prevent any tooth wear. These investigators also noted that the Fluoride and Strontium addition would result in a reduction of both hardness and abrasivity values which would be highly beneficial to be incorporated in a BG dentifrice. There are however limited and contradictory data analysing the abrasivity of BG in the published literature which also suggests the need for further work in this area. Additionally, Ali et al. indicated that there was a lack of evidence in the literature on the effect of salivary proteins on apatite deposition and emphasized the need for further research into this area [15].

There are also limited data on the particle size and shape and relative abrasivity of bioactive glass particles and it has been previously reported that particles that were more rounded and less angular could be expected to have a reduced abrasivity and a direct correlation was observed between both particle size and abrasivity. LaTorre et al. also demonstrated that finer particle sizes of Calcium Sodium Phosphosilicates (CSPS) resulted in a greater Ca$^{2+}$ and PO$_4^{3-}$ deposition [34]. However, these studies were conducted in vitro and ideally in vivo studies are required since these variables would be considerably different in a final formulation [15]. This would be particularly relevant for research into the effect on abrasion (by tooth brushing) following an acid challenge. Furthermore, as in vitro studies generally use healthy or unerupted teeth for SEM investigation, clinically teeth exposed to the norms of daily life may exhibit tooth wear and the thickness of the enamel may be reduced. This is evident particularly in the cervical area of the tooth where the enamel is at its thinnest and the dentine may be exposed. Ideally in vitro studies need to mimic the normal oral environment but sometimes this is very difficult to achieve for a number of reasons and care should be taken not to extrapolate these results in vitro studies into the clinical environment. It was also apparent that further investigation was required to analyse both particle size and shape in order to reduce abrasivity. For example, Mahmood et al. reported that increased particle size resulted in a greater enamel loss when comparing three different particle sizes of BG dentifrices: <38, 38-63 and 63-110 microns [35]. Particles formed from percussion milling with sharp edges were reported to be significantly more abrasive than round ball milled particles. Preferential wear was observed at the Enamel Dentine Junction; considering that this area relevant for both gingival recession and DH, it would be worthwhile to reduce the abrasivity of the dentifrice as much as possible otherwise greater wear will occur in the cervical area of the tooth. These investigators concluded that the abrasivity of a BG dentifrice was not a major problem, nevertheless, it is still advantageous to lower the abrasivity component especially when considering that the Fluorapatite layer formed is thin and can potentially be removed via abrasion [35]. Abrasives in a dentifrice can however contribute to the removal of tooth substance and initiate DH and as such it should be recommended to have low abrasive toothpaste which will minimise tooth surface loss while being able to remove stain and plaque from the tooth surface.

The original Novamin® dentifrice did not contain strontium, and Ali et al. suggested that it would be advantageous to include Strontium in BG dentifrice due to its anti-caries effect, ability in remineralization and apatite formation; and help to reduce the abrasivity of a dentifrice [15]. Studies specifically comparing Strontium containing Bioactive glasses to non-strontium containing dentifrices should be conducted in vitro and in vivo to determine whether the incorporation of Strontium into a dentifrice is relevant and clinical relevant. Further research analysing different ionic compositions of bioactive glass dentifrices including different concentrations of Strontium, Fluoride, Zinc and other ions Table 2 should therefore by conducted to determine the optimum concentration to be used in a bioactive glass dentifrice to treat DH.

**In vivo studies**

Litkowski et al. conducted a proof of principle study to test the bioactive glass’s safety in vivo and to assess whether the in vitro occlusion of dentinal tubules translated to a clinically significant reduction in DH [36]. This randomized, double-blinded study compared the following interventions namely 2.5% Novamin®, 7.5% Novamin® and a placebo (0% Bioglass) dentifrice formulations. Using thermal air and tactile tests on 66 subjects, the Novamin® dentifrices resulted in a statistically significant greater reduction in sensitivity scores compared to the placebo. The 7.5% Novamin® formulation resulted in the greatest reduction with no adverse events reported which appeared to support the use of a Novamin® dentifrice to relieve DH. The clinical evidence was supported by Gillam et al. in an in vitro study where the same interventions were evaluated under SEM investigation in that the 7.5% Novamin® occluded most of the tubules compared to 2.5% and 0% Novamin® which provided evidence of the potential mechanism (tubule occlusion) of the Novamin® dentifrice formulation [4]. Gendreau et al. also provided a comprehensive summary of previous in vivo studies [37]. In all the studies reviewed by Gendreau et al. Novamin® consistently resulted in a significantly greater reduction in all sensitivity tests compared to either a placebo

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**Table 3: Mohs harness values in selected materials and structures.**

<table>
<thead>
<tr>
<th>Material</th>
<th>Mohs hardness</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bioglass®</td>
<td>4.5</td>
</tr>
<tr>
<td>Enamel</td>
<td>3.5</td>
</tr>
<tr>
<td>Dentine</td>
<td>2.0-2.5</td>
</tr>
<tr>
<td>Calcium Carbonate</td>
<td>3.0</td>
</tr>
<tr>
<td>Alumina</td>
<td>9.25</td>
</tr>
</tbody>
</table>

or other dentifrices [37]. However, the Talioti et al. systematic review identified the various biases present in many of these and other studies [1]. For example, Litkowski et al. failed to provide any information concerning their randomization process or allocation concealment nevertheless, their study was still included in the Talioti et al. review unlike the study which was excluded due to its negative control group containing Fluoride whereas the intervention group, Novamin®, did not contain Fluoride [3,36]. Studies that were only reported as an abstract were also excluded. The Pradeep et al. and Pradeep and Sharma, studies were also considered to be of a high quality and fitted the inclusion criteria to be incorporated in the review [38,39]. However, Pradeep et al. did not disclose any information concerning calibration over the measure of DH and failed to sufficiently describe the placebo group. Additionally, Pradeep and Sharma did not provide any further details regarding their randomization method which was performed via a lottery process. Most of the studies obtained in the search strategy, however, were excluded by Talioti et al. [1]. For example, 31/34 Novamin® studies were excluded and the reasons for exclusion were extensive, the investigators also commented on the heterogeneity between the various studies which made further analysis impossible. Overall, the Talioti et al. review included only three randomized controlled trials involving CSPS [1]. Although randomized controlled trials are a high level of evidence, the small number of included studies rendered it impossible to make any definitive conclusions on the efficacy of Novamin® for treating DH. According to these investigators there was a need for further high quality studies in order to assess the effects of Novamin® formulations. It was evident that the current Novamin® products contain 5% CSPS although Litkowski et al. clearly showed that the 7.5% formulation was more efficacious than the 2.5% formulation [36]. There were, however limited data on studies comparing the various concentrations of bioactive glasses based on the Novamin® patent but one possible reason for using the lower percentage of glass in the formulation was the cost of the glass particles.

Several studies have recently compared a Novamin® formulation with Potassium Nitrate (5% KNO3) dentifrice formulations; for example, Acharya et al. compared a 5% CSPS dentifrice to a 5% KNO3 dentifrice [40]. It was reported that although the 5% CSPS group did result in a lower VAS score after 2 weeks of use, there were no statistically significant differences between the two groups at 4 and 8 weeks. Satyapal et al. also compared 5% CSPS to 5% KNO3 and reported that the CSPS group resulted in a statistically significant lower VAS scores after 3 weeks [41]. The 60 participants were subsequently asked to discontinue using the dentifrice and then recalled after a further 3 weeks. The VAS scores increased following this discontinuation due to the immense placebo effect present in DH studies [42]. A further problem in this field is the lack of a so-called gold standard product to compare in these types of studies and although some investigators have used KNO3 dentifrice formulations in this way there is no evidence to support the use of KNO3 dentifrices in treating DH [5,43].

Conclusions

Although the exact number of individuals suffering DH is controversial, it is undoubtedly a significant proportion of the population that complain about the problem. Overall, there appears to be promising evidence supporting the use of bioactive glasses to treat DH. In vitro evidence supports the ability of CSPS to occlude dentinal tubules whereas currently there appears to be limited in vivo evidence to support its efficacy in alleviating DH. There are also limited data in the published literature on the exact loading of CSPS glass into dentifrice formulations, abrasivity of CSPS, particle size and shape and the incorporations of various ions such as strontium and active ingredients into the bioactive glass’ network structure.

Reference


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