Improved performance of diatomite-based dental nanocomposite ceramics using layer-by-layer assembly

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Abstract: To fabricate high-strength diatomite-based ceramics for dental applications, the layer-by-layer technique was used to coat diatomite particles with cationic [poly(allylamine hydrochloride)] and anionic [poly(sodium 4-styrenesulfonate)] polymers to improve the dispersion and adsorption of positively charged nano-ZrO2 (zirconia) as a reinforcing agent. The modified diatomite particles had reduced particle size, narrower size distribution, and were well dispersed, with good adsorption of nano-ZrO2. To determine the optimum addition levels for nano-ZrO2, ceramics containing 0, 20, 25, 30, and 35 wt% nano-ZrO2 were sintered and characterized by the three-point bending test and microhardness test. In addition to scanning electron microscopy, propagation phase-contrast synchrotron X-ray microtomography was used to examine the internal structure of the ceramics. The addition of 30 wt% nano-ZrO2 resulted in the highest flexural strength and fracture toughness with reduced porosity. Shear bond strength between the core and veneer of our diatomite ceramics and the most widely used dental ceramics were compared; the shear bond strength value for the diatomite-based ceramics was found to be significantly higher than for other groups ($P < 0.05$). Our results show that diatomite-based nanocomposite ceramics are good potential candidates for ceramic-based dental materials.

Keywords: layer-by-layer, diatomite, nanoceramics, zirconia (ZrO2), dental materials

Introduction

Over the past 30 years, growing demand for highly esthetic and natural-looking dental restorations has led to the development of new all-ceramic dental materials with dramatically improved mechanical characteristics to increase their longevity. In particular, high-strength ceramics and related computer-aided design/computer-aided manufacturing (CAD/CAM) techniques have boosted clinical applications of metal-free prostheses with more favorable mechanical characteristics than earlier ceramic versions.

Zirconia (ZrO2) is one of the most promising restorative materials, because of its favorable mechanical properties and reasonable esthetics. Nevertheless, it has many drawbacks, including some unfavorable esthetic characteristics,1,2 and the veneering ceramic has a tendency to chip during machining,3 making it a less-than-perfect material. The field of dentistry, therefore, requires new materials with favorable esthetic and mechanical properties that are also easy to machine.

Diatomite is a type of porous silicate with a large specific surface area, high absorption capacity, low density, and low cost (see Appendix).4,5 It shows excellent biocompatibility and can withstand the layer-by-layer (LBL) assembly process.6,7 These properties allow for versatile applications, including as an adsorbent for heavy metal...
Diatomite has been used to fabricate high-strength porous ceramics. Thus, we investigated whether diatomite could be reinforced with nanoparticles to produce a new all-ceramic dental material with superior properties.

Fabricating nanocomposite ceramics with particles distributed within the matrix grains and along the grain boundaries can be difficult. The key to producing high-performance ceramic materials with high reliability lies in optimized processing. The particle size of the powders plays a critical role in determining the microstructure and mechanical properties of ceramics. Ideal powders have a small particle size and a narrow particle size distribution. However, as the particle size decreases, the powder tends to agglomerate. In this study, “agglomerates” are defined as weakly bonded particles that adhere together as a result of van der Waals forces, whereas “aggregates” are particles bonded together by solid bridges (typically formed during the calcination process). To overcome the problems associated with agglomerates, colloidal techniques have recently been used in ceramic processing.

The LBL technique is used to construct nanometer-scale coatings using the driving force generated by the electrostatic attraction between opposite charges. The LBL technique allows for the deposition of homogeneous films with accurately controlled layer thickness and interlayer separation on arbitrarily shaped objects. The method involves alternately adsorbing polymers on a nano- or micro-sized charged substrate to form a multilayer polyelectrolyte. The LBL technique has a wide range of applications in the modification of structural and surface properties because of its ability to readily tailor particle properties (eg, size, composition, porosity, stability, surface functionality, and colloidal stability). Other advantages include the hybrid organic–inorganic nature of the produced coatings; universality, with regard to both substrate and material to be deposited; and ease of implementation.

In this experiment, we modified diatomite-based ceramic powders using poly(allylamine hydrochloride) (PAH) and poly(sodium 4-styrenesulfonate) (PSS), adjusting the surface charge to help solve the problem of agglomeration. Positively charged nano-ZrO₂ particles were adsorbed onto diatomite particles with the aim of generating a microstructure that could help to improve the dispersion of the powder and the mechanical properties of the sintered ceramic bodies. The obtained ceramics were characterized and propagation phase-contrast synchrotron X-ray microtomography (PPCSR-CT) was used for the first time to examine the inner structure of the ceramics. The mechanical properties were also tested and compared with a control specimen, which was prepared by adding nano-ZrO₂ to the unmodified diatomite particles. In addition, we sintered veneer porcelain to diatomite-based ceramics then compared their shear bond strengths with those of dental zirconium dioxide ceramics and IPS e.max (Ivoclar Vivadent AG, Schaan, Liechtenstein) ceramics.

Materials and methods

LBL modification

To form high-quality diatomite-based ceramic powders, a diatomite:bentonite:quartz sand:Na₂CO₃ ratio of 11:4:4:1 was selected. The composite powder was dried after milling for 24 hours with the same amount of absolute ethanol. The diatomite-based powders were dipped in PAH (Mw = 40,000 g/moL; Sigma-Aldrich, St Louis, MO) solution at pH 5, followed by dipping in pH 9 PSS (Mw = 70,000 g/moL; Sigma-Aldrich) solution. The pH values were determined by carefully measuring the zeta potential of the modified powders. The PAH and PSS solutions were 1 wt% and were prepared in deionized water at 2 mg/mL with 0.5 M NaCl. The dipping time was 20 minutes, followed by repeated centrifugation–redispersion cycles (10,000 rpm for 5 minutes), with each successive supernatant being carefully decanted and replaced with deionized water to ensure that all excess polyelectrolyte had been removed. A pH 3 nano-ZrO₂ (3% yttria-stabilized zirconia, 80 nm; Tosoh, Tokyo, Japan) suspension, with positively charged surfaces, was adsorbed onto the negatively charged diatomite powders to produce a uniform structure. Figure 1 illustrates the fabrication process of LBL assembly of diatomite and nano-ZrO₂.

Particle size distribution was measured (Mastersizer 2000, Malvern, UK) before and after modification. The functional groups on the particle surface were analyzed by X-ray photoelectron spectroscopy (XPS; ESCALAB 250, Thermo Scientific, Waltham, MA) at a pass energy of 20 eV, using the Al-Kα line as the excitation source (hv = 1486.6 eV) at 150 W. All binding energies reported were corrected using the signal for the carbonaceous impurities at 284.8 eV as an internal standard. Scanning electron microscopy
Characterization of diatomite-based ceramics

The diatomite-based powders modified by the LBL technique with 0, 20, 25, 30, and 35 wt% nano-ZrO$_2$ were labeled M0Z, M20Z, M25Z, M30Z, and M35Z, respectively. Control specimens composed of the diatomite-based powders with nano-ZrO$_2$ treated in deionized water devoid of PAH and PSS under the same conditions as the experimental groups were labeled U0Z, U20Z, U25Z, U30Z, and U35Z. Both groups were sintered at 1100°C.

Sintered sample surfaces were polished and corroded in acetone for 20 seconds then cross-sections were observed by SEM and energy dispersive X-ray spectroscopy (JSM-5610LV).

The open porosity of the sintered ceramics was measured by the Archimedes’ method using the following equation:\(^2\)

$$P = \frac{M_3 - M_1}{M_3 - M_2} \quad (1)$$

where $P$ is the percentage of open porosity, $M_1$ is the dry weight of the specimen in the air, $M_2$ is the saturated suspended weight of the specimen, and $M_3$ is the saturated weight of the specimen.

Mechanical tests

Sintered rectangular bars (20 x 4 x 1.2 mm) were polished and dried for the three-point bending test by ISO 6872 (span: 15 mm, crosshead speed: 0.5 mm/min; Instron® Universal Testing Machine-3365, Norwood, MA).\(^2\)

The Vickers hardness and fracture toughness were measured with a microhardness tester (HV-10B, Laizhou, YanTai, ShanDong, People’s Republic of China) with an indentation load of 88.2 N and a load time of 15 seconds with the following parameters:\(^3\)

$$H_v = 1.854P/(2a)^2 \quad (2)$$
$$K_{IC} = 0.075Pc^{-3/2} \quad (3)$$

where $P$ is the indentation load (88.2 N), $2a$ is the average length of the two diagonals of the indentation, and $c$ is the average length of the indentation center to the crack tip. The results were analyzed by analysis of variance test and Tukey’s test. A $P$ value $<0.05$ was considered statistically significant.

U30Z and M30Z, which showed superior mechanical properties, were selected for PPCSR-CT and were imaged using a beamline of 13 W at the Shanghai Synchrotron Radiation Facility (Shanghai, People’s Republic of China). We used an isotropic voxel size of 0.7 µm and a beam monochromatized at an energy of 30 keV using a multilayer monochromator. To obtain the phase-contrast effect, we used a sample-detector distance (propagation distance) of around 10 cm, depending on the pixel size and 1500 projections on 180°.

Shear bond strength test

The shear bond strengths between the cores and veneers of the three groups, diatomite-based dental ceramics of M30Z (Dia group), zirconium dioxide dental ceramics (Zir group; Sirona Dental Systems GmbH, Bensheim, Germany), and IPS e.max (IPS group) ceramics were evaluated ($n=8$). The veneer ceramics were sintered in Programat EP3000 (Ivoclar Vivadent AG) in accordance with the manufacturer’s instructions. Shear bond strength was calculated with a universal testing machine.
(Instron Universal Testing Machine-3365) by dividing the load at which failure occurred by the bonding area as follows:\(^3\)

\[
P = \frac{F}{S}
\]

where \(P\) is the shear bond strength (MPa), \(F\) is the failure load (N) and \(S\) is the bonding area (2 \(\times\) 3 mm\(^2\)). Load was applied parallel to the long axis of the specimen through a wedge at the core–veneer interface at a crosshead speed of 0.5 mm/min until delamination of the veneering ceramic occurred. The results were analyzed using a one-way analysis of variance test \((P < 0.05)\).

**Results**

**Modification effects of LBL assembly**

To establish the effect of polyelectrolyte dispersants on the dispersibility of the diatomite powder, we examined the size distribution of the diatomite particles. Before modification (Figure 2A), the size distribution of the diatomite particles was wide, from 0.2 to 90 \(\mu\)m, where \(d_{0.5}\) was 16.42 \(\mu\)m. After PAH/PSS modification, the particle size tended to become smaller, with the main peak shifting to the left, while the size range narrowed (from 0.12 to 35 \(\mu\)m in Figure 2B); \(d_{0.5}\) decreased to 0.42 \(\mu\)m.

XPS is an ideal instrument with which to analyze the chemical state of surfaces in the successive assembly of PAH and PSS because of its high sensitivity and nondestructive nature. The survey scan XPS spectrum of the diatomite particles before modification is shown in Figure 3A, where the Si2p, O1s were detected at 102.66 eV(2p) and 532.05 eV(1s). Figure 3B shows the survey scan XPS spectrum of PAH adsorbed on the negatively charged diatomite particles. A clear N1s signal is present at 399.55 eV(1s) corresponding to the amine group of PAH. Figure 3C of the diatomite/PAH/PSS sample shows characteristic peaks at

![Figure 2](image-url)  
Figure 2 Size distribution of diatomite particles. (A) Before modification, the size distribution of diatomite particles spanned from 0.2 to 90 \(\mu\)m and \(d_{0.5}\) was 16.42 \(\mu\)m. (B) After PAH/PSS modification, the diameter shifted to the left, size distribution narrowed (from 0.12 to 35 \(\mu\)m) and \(d_{0.5}\) decreased to 0.42 \(\mu\)m.  
Abbreviations: PAH, poly(allylamine hydrochloride); PSS, poly(sodium 4-styrenesulfonate).
X-ray photoelectron spectroscopy spectrum of diatomite before and after modification. Before modification (A), Si2p and O1s were detected at 102.66 eV (2p) and 532.05 eV (1s); after modification by PAH (B), a clear N1s signal was present at 399.55 eV (1s), corresponding to the amine of PAH. Note: After modification with PAH/PSS, the spectrum showed characteristic peaks of S2p at 168 eV (S2p), assigned to the sulfur atom of the sulfonate group from the PSS polyelectrolyte.28

SEM images of diatomite-based ceramic powders and sintered ceramics

Diatomite-based composite dental ceramic powders without modification showed obvious agglomeration (Figure 4A); after modification, the agglomeration decreased. In addition, nano-ZrO₂ particles were adsorbed around the diatomite powder, as marked by a circle in Figure 4B. The microstructures of sections of sintered diatomite-based nanocomposite ceramic bodies were also examined (Figure 5). During sintering, polymer surfactants were burned off, leaving only diatomite and nano-ZrO₂ particles. The sintered body of unmodified diatomite-based powder containing 30 wt% nano-ZrO₂ (U30Z; Figure 5A) showed a large volume of irregular pores. Furthermore, the unmodified ceramic particles agglomerated significantly (Figure 5B).

Figure 3 X-ray photoelectron spectroscopy spectrum of diatomite before and after modification. Before modification (A), Si2p and O1s were detected at 102.66 eV (2p) and 532.05 eV (1s); after modification by PAH (B), a clear N1s signal was present at 399.55 eV (1s), corresponding to the amine of PAH.

Note: After modification with PAH/PSS, the spectrum showed characteristic peaks of Si2p at 168 eV (Si2p), assigned to the sulfur of the sulfonate group of PSS.

Abbreviations: PAH, poly(allylamine hydrochloride); PSS, poly(sodium 4-styrenesulfonate).

Figure 4 Scanning electron microscopy images of diatomite-based particles containing 30 wt% nano-ZrO₂. (A) Image shows agglomeration of unmodified diatomite particles containing agglomerated nano-ZrO₂. After modification (B), the diatomite particles are more uniformly distributed and nano-ZrO₂ particles are adsorbed.

Note: A typical structure is circled in (B).
Figure 5 Scanning electron microscopy analysis of sections of the sintered ceramics. (A and B) show sintered unmodified ceramics containing 30 wt% nano-ZrO$_2$ (U30Z), with a large volume of irregular pores. (C and D) show sintered modified ceramics containing 30 wt% nano-ZrO$_2$ (M30Z). Compared with those shown in (A and B), these ceramics exhibit high density with low porosity.

Note: The element of a typical structure was detected by EDS, confirming that it was composed of diatomite with adsorbed nano-ZrO$_2$ particles.

However, in the cross-section of sintered-modified diatomite-based ceramics containing 30 wt% nano-ZrO$_2$ (M30Z), the porosity had obviously decreased, while well-distributed microcrystalline phases were found embedded in the amorphous matrix (Figure 5C). A uniformly dispersed structure of nano-ZrO$_2$ adsorbed around diatomite particles was also observed (Figure 5D). The element of this structure was detected by energy dispersive X-ray spectroscopy. The results showed that point 1 was mainly composed of silicon, while point 2 was mainly composed of zirconium, confirming the
above results. Particle size decreased and agglomeration lessened after modification, leading to the formation of the compact green body shown in Figure 4B.

Open porosity
The open porosity values of the sintered ceramics in the modified group were statistically lower compared with those of the unmodified ones \( (P < 0.05) \). The porosity of both groups decreased with increasing levels of nano-ZrO\(_2\) (Figure 6). Before modification, the open porosity of U0Z showed a maximum of 3.15% ± 0.63%. After modification, M35Z showed a minimum of 0.10% ± 0.06%, indicating a near-perfect compact body. Open porosity declined because the particle size decreased and size distribution narrowed after modification. Smaller powders could more easily fill in the green body, so a more compact structure was formed. However, the volume of the closed pores of sintered dental porcelain could not be measured directly, so we combined the data with observations of the internal structure through SEM and PPCSR-CT to obtain a more detailed picture.

Mechanical properties
The mechanical properties of the modified and unmodified groups containing different amounts of nano-ZrO\(_2\) are shown in Figure 7. After modification, agglomeration decreased and nano-ZrO\(_2\) particles were adsorbed around the diatomite powder, which led to the formation of a compact green body. The mechanical properties of the modified sintered ceramic groups were statistically higher than those of the unmodified ones \( (P < 0.05) \). Flexural strength in the modified groups increased with increasing levels of nano-ZrO\(_2\), reaching a peak value of 276.80 ± 16.27 MPa when 30 wt% nano-ZrO\(_2\) was adsorbed (M30Z), and decreasing again when 35 wt% nano-ZrO\(_2\) was adsorbed (M35Z). The Vickers hardness of the modified groups was also statistically higher \( (P < 0.05) \), reaching a peak of 12.82 ± 0.37 GPa (M35Z). The fracture toughness of modified groups improved \( (P < 0.05) \) and reached its highest level of 5.10 ± 0.24 MPA m\(^{1/2}\) in M30Z.

PPCSR-CT
In this study, we attempted to determine the internal structure of the ceramics. Figure 8 shows 3D tomographic representations of U30Z and M30Z and their cross-section images from the x, y, and z axes. U30Z contained many pores, while no pores were visible to the naked eye on M30Z,
Discussion

Narrow size distribution powders give consistently more uniform and finer pore structures and are sintered to high density at relatively low temperatures with a uniform and fine-grained microstructure. Polyelectrolyte dispersants can contribute both static and steric stability at a certain pH to prevent the agglomeration of ceramic particles. Polyelectrolytes in solution can adsorb onto the powder surface, which changes the charge properties of the powder surface and the electrostatic repulsion among particles, as reflected by the increasing zeta potential. Thus, electrostatic repulsion is a well-known method of dispersion stabilization for both particles and polyelectrolytes. However, increasing the thickness of the polymer adsorption layer on the particle surface enhances the steric effects. Good stability is demonstrated through a synergistic effect between polymer dispersions.

We hypothesized that we could solve the problem of agglomeration of diatomite particles to form a uniformly dispersed nanocomposite ceramic powder by using polyelectrolyte dispersants, the LBL technique, and nano-ZrO₂ particles. After LBL modification, the particle size of the diatomite powders decreased and the size distribution range narrowed; dₜ₅ decreased from 16.42 to 0.42 µm (Figure 2).

In addition, nano-ZrO₂ particles, which showed a positive charge on the surface, were adsorbed around the negatively charged P AH/PSS-modified diatomite powders (Figure 4B). In this study, thicker coatings were not needed, so only two-layer polymer coatings were prepared. The smaller particle size is conducive to the formation of denser and more uniform diatomite/nano-ZrO₂ composite ceramic powders, which can significantly improve the mechanical properties of ceramics.

To detect whether P AH/PSS polyelectrolyte had adsorbed onto the surface of the diatomite particles, we examined the particles by XPS. The XPS spectrum of the diatomite/P AH/PSS sample revealed the presence of nitrogen from the amine groups of P AH linkers. Considering that the modification of the diatomite by P AH occurred because P AH has an amine group, a peak indicating the presence of nitrogen could be expected. The N₁s peak at 399.55 eV can be deconvoluted into two nitrogen species: a species that looks like an unprotonated amine at 399.2 eV and the protonated amine (NH₃⁺) at 401.2 eV in Figure 4B. In addition, nano-ZrO₂ particles, which showed a positive charge on the surface, were adsorbed around the negatively charged PAH/PSS-modified diatomite powders (Figure 4B). In this study, thicker coatings were not needed, so only two-layer polymer coatings were prepared. The smaller particle size is conducive to the formation of denser and more uniform diatomite/nano-ZrO₂ composite ceramic powders, which can significantly improve the mechanical properties of ceramics.

Shear bond strength

The results of shear bond strength were significantly different between the three groups (P < 0.05). The Dia group had the highest shear bond strength of 21.26 ± 4.23 MPa. The average value in the IPS group was 14.07 ± 4.28 MPa, while the Zir group had the lowest value of 6.85 ± 1.11 MPa.
expected to be protonated at pH conditions of PSS adsorption (at pH 5, the amine groups of PAH are all accessible), while a fraction of the protonated amine groups of PAH that strongly interact with the sulfonate groups (possibly via H-bonding) would have a corresponding N1s peak at a lower binding energy than the amine groups not matched to sulfonates.

There may be several reasons why the mechanical properties of our ceramics were enhanced. First, the ceramic powders obtained after modification were composed of particles with a narrower size distribution and less agglomeration; the diatomite particles coated by nano-ZrO2 were homogeneous and could form a denser green body resulting in extremely greater mechanical properties nearly equal to those of conventional glass ceramics used in dental applications.33–36 As is the case for many high-performance ceramics, the strength of sintered ceramics can be increased by decreasing the porosity and number of flaws caused by processing. Secondly, the nano-ZrO2 particles adsorbed around the diatomite and embedded into the gaps could form a protective barrier to prevent the extension of cracks.32 Mechanical performance was further improved by crack deflection by mica plates and phase transformation of ZrO2 particles. Due to crack deflection and the inhibition of grain boundary sliding at elevated temperatures, Nano-ZrO2 is a toughening material with excellent mechanical properties at ambient temperatures.33–36

Flexural strength was highest in M30Z, decreasing in M35Z when levels of nano-ZrO2 rose further; this might be because the excess nano-ZrO2 could not be adsorbed on the surface of the diatomite or packed into the gaps but instead agglomerated, resulting in an uneven density and reduced strength. In unmodified groups, there was nearly no adsorption of nano-ZrO2, only mechanical mixture. The fracture toughness of modified groups was somewhat improved, reaching a peak of 5.10 ± 0.24 MPa m^1/2 in M30Z. From a practical standpoint, microstructural effects were found to be important, but only within a limited range; the chemistry appeared to define the band of achievable property values.37

The traditional method for microstructure characterization is SEM, which offers the advantage of precision for nanometer-scale microstructures. However, it can only show a single 2D cross-sectional view of the interior of the specimen. Observations of internal structures are very important for ceramic materials because the dispersion of internal sintered compounds, porosity, and pore volume are essential to the study of ceramics. X-ray synchrotron microtomography is a nondestructive technique for visualizing features from any vantage point, visualizing internal characteristics in virtual cross-sections in any plane, and generating 3D virtual extractions of internal structures useful for the analysis of a wide range of materials including rock, bone, ceramic, metal, and soft tissue.38–41 The contrast in absorption tomography is related to the density of the material, and thus different material phases can be distinguished. Pores in the ceramic sintered body should have a low density, which can be clearly observed from the 3D image (Figure 8).

Since adhesion of the veneering ceramic to the ceramic substructure is key to the successful performance of veneer/core bilayered restorations, the initial bond strength can provide useful information on the behavior and predictability of all-ceramic systems in clinical applications. The diatomite-based substrate ceramic of M30Z after LBL modification (Dia group) statistically showed the highest mean shear bond strength of 21.26 ± 4.23 MPas compared with the other clinical all-ceramic materials (P < 0.05). Unmodified diatomite-based dental ceramics were not included in this study, as they did not meet the basic strength requirements of clinical substrate ceramics. Many variables can affect the core–veneer bond strength, such as residual stresses generated by a mismatch in the thermal expansion coefficient, flaws and structure defects at the core–veneer interface, and wetting properties and volumetric shrinkage of the veneer.32,43 Usually, the core–veneer bond or the veneering material itself has to be strong enough to withstand the stresses of mastication to prevent delamination and fracture of the veneering material. Experimental veneer with a higher thermal expansion coefficient resulted in massive fractures in both the core and veneering material. Failures occur, in part, because of transient and residual stress concentrations at the core–veneer interface that are formed during cooling. Residual stresses can assist the propagation of micro-cracks along the interface, significantly decreasing the bond strength or causing veneer delamination or chipping.44 As a silicate ceramic, diatomite contains the same ingredients as veneering porcelain, which may help to form a chemical bond between them. If a good chemical match between the core and veneer porcelain can be obtained, the structure should be able to withstand not only transient and residual stresses but also the greater forces of functional processes. In our next study, we will further investigate the nature of the interface between the diatomite-based dental ceramic and the veneering porcelain.

Conclusion
The agglomeration of diatomite-based powders improved with the use of the LBL technique, which led to a significantly smaller particle size and a narrower size distribution.
After modification, the surface of the diatomite-based powder was negatively charged, thus, could adsorb positively charged nano-ZrO₂ particles to improve the sintered ceramics. A highly dense microstructure was obtained and the mechanical properties of the modified diatomite-based nanocomposite ceramics were significantly improved. The shear bond strength of the diatomite-veneer ceramic was statistically higher than that of other clinical all-ceramic dental materials.

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Disclosure

The authors declare no conflicts of interest in this work.

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Appendix
Diatomaceous earth has recently attracted a great deal of attention from industry because of its abundance, low cost, and the unique structure of diatom shells. Our research group developed a new type of diatomite-based machinable dental ceramics in 2008 (China patent 200910115734.3).