Processing extracted teeth for immediate grafting of autogenous dentin

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Tooth extraction is one of the most widely performed procedures in dentistry, and it has been historically well documented that it can induce significant dimensional changes of the alveolar ridge.

In their review, Horowitz, et al. (2012), stated that less ridge resorption occurs when alveolar ridge preservation procedures are used, compared to leaving fresh alveolar sockets without placing graft material. If performed inadequately, the resulting deformity can be a considerable obstacle to the esthetic, phonetic, and functional results.

In dentistry, allogeneic bone and synthetic mineral materials are the main source for grafting in bone. However, fresh autogenous bone graft is still considered the gold standard since it exhibits bioactive cell instructive matrix properties and is non-immunogenic and non-pathogenic, in spite of the need for harvesting bone and possible morbidity resulting from it.

It is well-known that jawbones, alveolar bone, and teeth develop from cells of the neural crest and that many proteins are common to bone, dentin, and cementum (Donovan, et al., 1993; Qin, et al., 2002). It is, therefore, not surprising that dentin, which comprises more than 85% of tooth structure, can serve as native bone grafting material.

Interestingly, Schmidt-Schultz and Schultz (2005) found that intact growth factors are conserved even in the collagenous extra-cellular matrix of ancient human bone and teeth.

Methods for processing bovine dentin into particulate and sterile grafting material for preserving of alveolar bone have been described and used in several animal studies (Fugazzotto, et al., 1986; Nampo, et al., 2010; Qin, et al., 2014). It is, therefore, evident that teeth can become grafts that are slowly and gradually replaced by bone (Hasegawa, et al., 2007).

Currently, all extracted teeth are generally considered clinical waste and, therefore, are simply discarded. Recently, however, several studies have reported that extracted teeth from patients, which undergo a process of cleaning, grinding, demineralization, and sterilization, can be a very effective graft to fill alveolar bone defects in the same patient (Kim, et al., 2010; Kim, et al., 2011; Murata, et al., 2011). However, this procedure is extremely time-consuming since the graft is only ready several hours or days after extraction.

This article, therefore, aims to present a modified procedure that employs freshly extracted teeth in a clinical setting by recycling them into bacteria-free particulate autogenous mineralized dentin for immediate grafting.

A Smart Dentin Grinder® (SDG) (KometaBio) was devised, which grinds and sorts extracted teeth into dentin particulate of a specific size. A chemical cleanser is then applied to process the dentin particulate into a bacteria-free graft over the course of about 15-20 minutes.

This novel procedure is indicated mainly in cases when teeth are extracted because of periodontal reasons and for partially or totally impacted teeth. Teeth that have undergone root canal fillings should not be employed in this procedure because of the risk of contamination by foreign materials. On the other hand, crowns and fillings can be reduced, and the clean dentin of the tooth crown can be processed for immediate grafting.

Method: from extraction to grafting particulate dentin

Teeth without root canal fillings, which have been extracted due to advanced periodontal bone loss or other reasons, such as wisdom teeth extraction or orthodontic indications, can be prepared for immediate grafting.

Immediately after extraction, restoration or removal of carious lesions and discolored dentin, or remnants of periodontal ligament (PDL) and calculus should be reduced by using a tungsten bur (Figures 1A and 1B). The authors have found that high-speed tungsten carbide burs are most efficient for this process. The roots could be split in case of multi-rooted teeth.

Clean teeth, including crown and root dentin, are dried by air syringe and put into the grinding sterile chamber of the newly designed Smart Dentin Grinder (Figure 2A). The SDG can grind the roots in 3 seconds...
and then uses the vibrating movement of the grinding chamber to sieve any particles smaller than 1,200µm into a lower chamber that collects particles between 300µm and 1,200µm (Figure 2B). Particles smaller than 300µm fall into a waste drawer, as this fine particulate is not considered to be an efficient size for bone grafting. This grinding and sorting protocol is repeated to grind the remaining teeth particles left in the grinding chamber, still collecting particles between 300µm and 1,200µm.

The particulate dentin from the drawer is immersed in basic alcohol for 10 minutes, in a small sterile glass container. The basic alcohol cleanser consists of 0.5M of NaOH and 30% alcohol (v/v) for defatting, dissolving all organic debris, bacteria, and toxins of the dentin particulate.

Figure 3 shows the efficiency of the cleanser to dissolve all the organic debris from dentin particulate, including dentin tubules. The scanning electron microscope (SEM) picture shows open and clean tubules after 10 minutes of cleanser treatment (Figure 3C). After decanting the basic alcohol cleanser, the particulate is washed twice in sterile phosphate-buffered saline (PBS). The PBS is decanted, leaving wet particulate dentin ready to graft into freshly extracted sockets, alveolar bone defects, or in procedures involving augmenting the maxillary sinus.

The process from tooth extraction until grafting takes approximately 15-20 minutes.

It should be noted that the efficiency of selecting the dentin particulate of specific size for grafting is more than 95%. It is also obvious that the volume of the particulate dentin is more than twice of the original root volume. Alternatively, the wet particulate can be put on a hot plate (140°C) for 5 minutes to produce dry, bacteria-free particulate autologous dentin that can serve for immediate or future grafting procedures.

**Results: clinical evaluation**

Over a period of 2 years, more than 100 dentists have employed the present procedure for preparing autogenous dentin particulate from extracted teeth for immediate grafting in the same patient. It should be noted that teeth that underwent root canal treatment were discarded. When intact teeth were processed, the enamel and cementum were included. Figures 4 to 7 show a number of typical case presentations where teeth were extracted and processed into bacteria-free particulate dentin.

**Figures 1A-1C:** From extraction to clean particulate: 1A. Tooth after extraction, debris, and calculus. 1B. Same tooth after reducing debris with tungsten carbide bur. 1C. Particulate dentin after grinding and sorting. The particulate dentin size is 300µm-1200µm

**Figures 2A-2B:** Smart Dentin Grinder and drawer with particulate dentin of 300µm-1200µm size ready for cleanser treatment: 2A. Smart Dentin Grinder and sorter. 2B. Drawer that collects particulate dentin after grinding and sorting. The size of particles in this drawer is 300µm-1200µm

**Figures 3A-3C:** 3A. Scanning electron microscope (SEM) x750 of particulate dentin when cleanser is added. 3B. SEM x750 of particulate dentin at 3 minutes after treatment with cleanser. 3C. SEM x750 of particulate dentin at 10 minutes after treatment with cleanser. Note the wide-open tubuli openings. Bacteriological tests revealed no bacteria growth at this point

**Figures 4A-4D:** Extraction sites at LR8 filled with particulate dentin prepared from extracted tooth by the Smart Dentin Grinder procedure: 4A. Clinical view of the extraction site. 4B. X-ray of impacted tooth LR8. 4C. After extracting the LR8, particulate of extracted tooth was prepared and placed in extraction site. 4D. By 4 months, the particulate and newly formed bone completely restored the void next to the distal root of tooth LR7
autogenous tooth dentin for immediate grafting in same patient.

**Wisdom tooth extraction**

A total of 16 wisdom teeth, including partially impacted, horizontally impacted, and caries-affected teeth, were processed using the SDG procedure during this study. Figure 4 shows a horizontally impacted LR8 tooth that was in close proximity to the distal root surface of the LR7, creating a deep void. The surgically extracted LR8 exposed the distal root surface of the LR7, almost denuded from bone tissue. The LR8 was processed immediately into the particulate graft, which totally filled the extraction site (Figure 4C). Healing and recovery after the surgical procedure and grafting took place without complications.

A follow-up after 4 months revealed a normal pattern of marginal gingiva around the LR7. Probing was normal: 1 mm-2 mm in depth. On the X-ray distal to the LR7, new bone and particulate dentin was integrated into bone, completely restoring the extraction site and distal bone support of the LR7 (Figure 4D).

**Periodontal extractions**

A further 37 teeth were extracted because of poor periodontal attachment, bone loss, and mobility. Figure 5 illustrates the case of a 56-year-old male patient with a localized, advanced periodontal condition in posterior parts of the mandible.

The LR7 and LR8 were extracted, and the granulation tissue was removed exposing bone tissue walls. The LR7 had a root canal filling and was therefore discarded. The LR8 was processed into particulate dentin by the SDG device and prepared for immediate grafting in the extraction sites.

The grafting of one tooth produced sufficient volume of particulate dentin to overfill the extraction site of both sockets. A Choukroun PRF (platelet rich fibrin) membrane was prepared from the patient’s blood (Cieslik-Bielecka, et al., 2012) to cover the graft. The mucoperiosteum was sutured overfill the extraction site of both sockets. 6D. After preparation of particulate from tooth UL6, the socket was grafted and the oroantral opening filled with particulate dentin. 6F. After 3 months, three implants were placed, and immediate solid anchorage was achieved.

**Sinus lifts**

Autogenous dentin particulate can serve as a superior grafting matrix for augmenting bone in maxillary sinuses, as presented in the next case.

The patient presented with alveolar bone loss, with infrabony pockets that extended into the maxillary sinus of tooth UL6 (Figure 6). Two months after grafting with the particulate dentin from tooth LL8, three implants were inserted (Figure 5H), and 1 year later, the bone density and bone level with no signs of bone resorption at the crest after restoration could be observed (Figure 5J).

**Closure of the wound and sutures of mucoperiosteum flap was performed.**

Healing was normal, and 3 months later, an alveolar ridge of minimum 8.3 mm height was achieved, allowing placement of three implants. It should be noted that one molar — the UL6 — produced at least 2 cc of particulate dentin, which allowed augmentation of the extraction socket and part of the sinus.

Moreover, we found that autogenous dentin grafting allowed the placement of implants after 3 months in the upper jaw because the new bone that was integrated with particulate dentin produced a solid support for implants.

Loading of implants followed. During preparation of the site for implant placement, a core of bone was recovered from the grafted socket site. The histology revealed new bone integrated with grafted dentin, producing a bone-dentin interface and connectivity (Figure 7).
Discussion
More than 40 years ago, autogenous teeth were routinely transplanted into extraction sockets when possible. It is evident that transplanted teeth that are ankylosed in the jawbone undergo replacement resorption over 5 to 8 years (Sperling, et al., 1986).

In addition, it is well documented that avulsed teeth that are implanted back into their sockets undergo reattachment to bone, which is formed directly on root dentin or cementum, leading to ankylosis (Andersson et al., 1989). An ankylosed root is continuously resorbed and replaced by bone, eventually resorbing the entire root, while the alveolar process is preserved during this period and later.

In a recent review, Malmgren (2013) stressed that in ankylosed teeth that are treated by decoronation, the alveolar ridge is maintained in the buccal/palatinal direction, while vertical height is even increased (Park, et al., 2007). Our results reveal similar interaction between the mineralized dentin and osteogenic cells that attach and produce mineralized bone matrix directly on the dentin graft.

A tooth bank in Korea provides a service that prepares autogenic demineralized dentin matrix graft in block or granular types (Kim, et al., 2011; Murata, et al., 2011; Kim, 2012), delaying the grafting procedure from several hours to several days and, therefore, requiring an additional surgical session. Although demineralized dentin exposes matrix-derived growth and differentiation factors for effective osteogenesis, the newly formed bone and residual demineralized dentin are too weak to support implant anchorage. In contrast, the SDG procedure allows preparation of bacteria-free particulate dentin from freshly extracted autologous teeth, ready to be employed as autogenous graft material immediately.

Mineralized dentin particles have the advantage of maintaining mechanical stability, allowing early loading after grafting in fresh sockets and bone defects. Moreover, in spite of its delayed inductive properties (Yeamans and Urist, 1967; Huggins, et al., 1970), the mineralized dentin is firmly integrated with newly formed bone, creating a solid site for anchorage of dental implants. In fact, our clinical data indicates that implant insertion and loading can be performed in both lower and upper jaws 2 to 3 months after dentin grafting.

Since the mineralized dentin is very slowly remodeled (Yeamans and Urist, 1967; Kim, et al., 2014; Andersson, 2010) in comparison to cortical bone or most biomaterials, the esthetic and structure pattern of the alveolar crest and mucoperiosteum is maintained for years. Teeth and jawbone have a high level of affinity, having a similar chemical structure and composition. Therefore, the authors and others (Kim, et al., 2011; Murata, et al., 2011; Kim, 2012) propose that extracted non-functional teeth or periodontally involved teeth should not be discarded any more.

Extracted teeth can become autogenous dentin, ready to be grafted within 15 minutes after extraction. We consider autogenous dentin as the gold standard graft for socket preservation, bone augmentation in sinuses, or filling bone defects.

Disclosure
The Smart Dentin Grinder is distributed by Kometa Bio. Drs. Itzhak Binderman and Lari Sapoznikov helped to develop the Smart Dentin Grinder and have shares in Kometa Bio Ltd., the company responsible for distributing the device.

Drs. Gideon Hallel, Casap Nardy, and Avinoam Yaffie have no conflict of interest. They participated actively in providing clinical cases and their follow-ups. 

REFERENCES