A Method for Rapid Demineralization of Teeth and Bones

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Abstract: Tooth and bone specimen require extensive demineralization for careful analysis of cell morphology, as well as gene and protein expression levels. The LacZ gene, which encodes the β-galactosidase enzyme, is often used as a reporter gene to study gene-structure function, tissue-specific expression by a promoter, cell lineage and fate. This reporter gene is particularly useful for analyzing the spatial and temporal gene expression pattern, by expressing the LacZ gene under the control of a promoter of interest. To analyze LacZ activity, and the expression of other genes and their protein products in teeth and bones, it is necessary to carry out a complete demineralization of the specimen before cutting sections. However, strong acids, such as formic acid used for tooth demineralization, destroy the activities of enzymes including those of β-galactosidase. Therefore, most protocols currently use mild acids such as 0.1 M ethylene diamine tetra-acetic acid (EDTA) for demineralization of tooth and bone specimen, which require a longer period of treatment for complete demineralization. A method by which hard tissue specimens such as teeth and bones can be rapidly, but gently, decalcified is necessary to save time and effort. Here, we report a suitable method for rapid demineralization of mouse teeth in 0.1M EDTA at 42°C without any loss of β-galactosidase activity.

Key Words: Demineralization, tooth, LacZ, gene expression

INTRODUCTION

Teeth are the hardest tissues in the body, consisting of enamel, dentin, and cementum, and have a highly mineralized extracellular matrix. Unlike soft tissues, which can be easily sectioned and analyzed, teeth require complete demineralization in order to prepare proper sections for histological analysis using certain stains and antibodies. In the past, many decalcifiers have been tested for different purposes. For example, the Morse’s solution, a strong acid-based decalcifier, has been used for the rapid detection of RNA by in-situ hybridization (ISH) [1]. Another acid-based solution, 5% trichloroacetic acid (TCA), has been used for the analysis of DNA strand breaks by terminal deoxy (d)-UTP nick-end labeling (Tunnel) [2]. EDTA containing solution is a mild demineralization agent which has been mostly used for immunolocalization studies because of its antigen-preserving properties. Treatment of bones with 0.1 M EDTA following initial fixation in glutaraldehyde has been reported as adequate for immunocytochemical localization of certain bone matrix proteins [3]. Furthermore, the treatment of dental tissues with 4.3% EDTA resulted in satisfactory preservation of the fine structures of the cells and matrices [4], however it requires a relatively long incubation time in order to achieve complete demineralization [5,6]. For example, to demineralize human deciduous teeth for analysis, it requires 4 weeks of demineralization using 10% EDTA [7]. To demineralize a 1-month-old mouse skull with similar EDTA solution it takes about 3-4 weeks of treatment to prepare the specimen for analyzing expression levels of the LacZ and growth fac
tor genes [8,9]. To achieve fast demineralization, a microwave-induced demineralization method has been examined [10,11]. While this method has achieved faster demineralization time compared to the traditional method, preserved the fine morphological structures of the bone tissue, and retained RNA, its effect on LacZ gene expression has not been addressed. Moreover it has been speculated that the advantages associated with using the method are due to the higher temperature attained during microwaving the specimen purely due to thermal effect [12].

The LacZ gene is widely used to investigate promoter activity [13-15], however there is not detailed protocol for optimal analysis of β-galactosidase activity in decalcified mouse teeth and bones. Moreover, the effects of acid-based decalcifiers and different fixatives used for preparation of mouse teeth specimen to analyze β-galactosidase activity have not been well established. In this report, we demonstrate the effects of several fixatives and decalcifiers on mouse teeth specimen for LacZ activity, whose expression is driven by a tooth specific DSPP (Dentin Sialophosphoprotein) promoter. Most importantly, we have explored the effects of elevated temperatures on mouse teeth specimen to optimize a rapid but gentle demineralization method suitable for analysis of LacZ activity. This will prove to be a suitable method for rapid demineralization of mouse skulls for analysis using histochemical staining and in-situ hybridization.

MATERIALS & METHODS

Experimental Animals: We reported earlier on the generation and characterization of the Dspp-LacZ transgenic mice, which express LacZ predominantly in the odontoblasts, were used for all the experiments in this study [16]. The skulls from 2-week-old, 1-month-old, 3-month-old, and

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1-year-old mice were removed after cervical dislocation and were cut into two symmetrical halves. The mouse skulls were subsequently treated using the indicated procedures. Studies were performed in compliance with the National Institutes of Health (NIH) guidelines on the care and use of laboratory and experimental animals. All experimental procedures were approved by the Animal Care and Use Committee of the National Institute of Dental and Craniofacial Research.

Genotyping: Genotyping was performed by PCR analysis of genomic DNA extracted from tail snips. Tail biopsies were lysed overnight at 55°C in a lysis buffer containing 100mM Tris pH 7.4, 5mM EDTA, 200mM NaCl, 0.2% SDS, and 100 μg/ml Proteinase K/ml (Invitrogen) in distilled water. DNA was purified using phenol and chloroform, and PCR was performed for genotyping Dspp-LacZ transgenic mice using the following primers: the forward primer (5'-CGTAATAGCGAAGAGGCCCGACC-3’) and the reverse primer (5’-CTGCCTTTTACCCC TGCCATAAG-3’). The PCR conditions were set as follows: 30 cycles at 94°C for 1 min., 70°C for 30 sec., and 72°C for 1 min.

β-galactosidase Assay: The skulls were cut from 3-month-old Dspp-LacZ transgenic mice, and skins were removed and then fixed with 0.25% glutaraldehyde in PBS, 4% paraformaldehyde (PFA) in PBS, aqueous buffered zinc formalin (Anatech Ltd, Battle Creek, MI), or 10% formalin, for 1 hour at room temperature, and decalcified in 0.1M EDTA (0.5M EDTA, pH 8.0 buffered by 1X DPBS, pH 7.5) for 3 weeks. Decalcified skulls were then washed in PBS three times and embedded overnight in O.C.T. compound (Sakura, Torrance, CA) to prepare frozen sections for cutting. Next day, the sections that were 12 μm in thickness were cut using a cryostat (Leica Microscopy, model CM3050 S), and transferred onto Vectobond treated slides (Vector Laboratories, Burlingame, CA). These sections were stained with LacZ solution containing 0.02% NP-40 (IPEGAL), 0.01% sodium deoxycholate, 2mM MgCl2, 1mM spermidine, 5mM potassium ferricyanide, 5mM potassium ferrocyanide, and 0.5mg/ml blue-Gal (Invitrogen, Carlsbad, CA) in phosphate buffered saline (PBS) overnight at 37°C. The stained sections were counterstained with eosin and then photographed (Axioplan 2, Zeiss, Thornwood, NY).

Post Fixation Assay: The skulls were cut from 3-month-old mice, fixed, decalcified, and sections were arranged on the slides as described above. The slides containing tooth incisor sections were post-fixed with 4% PFA for 0 min., 2 min., 5 min., 10 min., 30 min., and 2 hours at RT, before being stained with LacZ solution as described above.

Decalcifier Comparison Assay: The skulls were cut from 3-month-old mice, fixed with 4% PFA for 1 hour at room temperature and decalcified with Formalin-4 for 2 days, with Immunocal (Decal Chemical Corporation, Congers, NY) for 2 days, or with 0.1 M EDTA solution for 3 weeks. The decalcified skulls were embedded in O.C.T., sectioned, and stained with LacZ solution as described above.

Temperature Assay: The skulls were removed from 3-month-old mice, cut in half, fixed with 4% PFA for 1 hour at room temperature and decalcified with the 0.1M EDTA solution at RT, 37°C, 42°C or 50°C, as follows: first, the skull specimens of different ages were placed in one-liter bottles filled with 0.1M EDTA. The bottles containing skulls were then placed in the multi-stirrer water bath (Lab-line, Model 1286, Melrose Park, IL) and filled with water warmed at a desired temperature with a constant mixing of the demineralization agent. The demineralization of the mouse skulls of different ages was performed one temperature at a time. As an alternate method, the skull specimens were placed in a 50 ml tube filled with 0.1M EDTA. The 50 ml tube containing skull specimens was placed and rotated in a temperature-controlled hybridization oven (Little Shot II, Boekel Scientific, Feasterville, PA). The skull specimens were x-rayed (model MX20, Faxitron x-ray Corporation, Wheeling, IL) daily at an optimized setting of 20 KV X 45 seconds to monitor the progression of the demineralization process.

Von Kossa Staining: The frozen sections from the decalcified skulls were prepared as described above, and the completion of the demineralization process was confirmed by incubating the slides in 5% silver nitrate solution for 1 hour. They were then rinsed with distilled water three times and counterstained with Nuclear Fast Red solution. The slides were then dehydrated first in 95%, and then in 100% alcohol, mounted with a xylene-based mounting medium, and the images were photographed.

Reverse Transcription- PCR: The skulls were cut from 3-month-old mice, fixed, decalcified at 37°C, 42°C or 50°C, and cut by a cryostat into slides of 12 μm thickness as described above. Odontoblasts were taken directly from the slides of the frozen tooth sections using toothpicks, and the total RNA was prepared using TRIZOL (Invitrogen). The RNA specimens were subjected to first strand cDNA synthesis using the Superscript first strand synthesis system, and RT-PCR (Invitrogen) was performed using GAPDH standard primers (Forward-CCATCACCATCTTCAGGAG and Reverse-GCATGACTGTGTCATGAG). All PCR reactions were carried out in a PerkinElmer gene PCR system 600 (PerkinElmer, Waltham, MA).

RESULTS

Evaluation of fixatives and fixation time on β-galactosidase activity: The effects of different fixatives on β-galactosidase activity in soft tissues were reported earlier [17], however there are no such reports about the effects of different fixatives on hard tissues such as teeth. To examine which fixative agent is suitable to analyze β-galactosidase activity in tooth sections, we first compared the effects of 1 hour fixation by 0.25% glutaraldehyde, 4% PFA, zinc formalin, and formalin. As shown in Fig. (1), we could detect similar levels of β-galactosidase activity in the odontoblasts of the incisors from the skulls fixed with different fixatives, indicating that treatment with these agents retained the enzyme activity at a similar level. To further investigate the optimum fixation time for analyzing β-galactosidase activity in teeth, we post-fixed the tooth section with 4% PFA for different periods of times. All the time periods tested, except for the 2 hour time period, yielded a strong signal for β-galactosidase activity (Fig. 2).

EDTA based demineralization maintains β-galactosidase activity: The acidity of demineralization agents has been
reported to affect the stability of proteins, enzyme activities and tissue integrity [8]. Therefore, we analyzed the effects of various demineralization reagents on the β-galactosidase activity. We fixed the skulls from 3-month-old Dspp-LacZ mice with 4% PFA and then decalcified them in Formical-4 and Immunocal solution, both of which are formic acid-based fixative decalcifiers. We then prepared the frozen sections, stained them with LacZ solution and compared the β-galactosidase activity with the sections from the skulls that were fixed by 4% PFA and decalcified with 0.1M EDTA. Only the tooth sections from the skulls treated with EDTA-based decalcifiers preserved the activity, as shown by the strong blue staining in the odontoblasts in Fig. (3c).

Since moderate increase in incubation temperature could shorten the demineralization process time period for the analysis of β-galactosidase activity, we tested the effects of elevated temperatures during the demineralization process using 0.1M EDTA as a decalcifier. The degree of opacity
was well-correlated with the degree of skull demineralization. The undecalcified areas of the skulls appeared as radioopacity in the x-ray image. At RT, 10 days of incubation were required for complete demineralization, however just 6, 5, and 3 days were required at 37°C, 42°C, and 50°C, respectively (Fig. 4). Next, in order to test the effects of each temperature on the stability of β-galactosidase activity at the time of complete demineralization, we decalcified the skulls with 0.1M EDTA for a sufficient period of time, which was 10 days at RT, 6 days at 37°C, 5 days at 42°C, and 3 days at 50°C, and then we analyzed the β-galactosidase enzyme activity in the sections. We found that the skulls decalcified at 37°C and 42°C, but not at 50°C, showed equivalent β-galactosidase enzyme activity, although at 50°C the incubation time was the shortest (Fig. 5). The successful completion of the demineralization process was indicated by the light brown precipitate on tooth sections in Fig. (5 b, d, f and h), as opposed to the dark brown calcium deposit shown in the undecalcified control tooth section (Fig. 5i).

Longer time required for demineralization of mouse skulls from older mice: Having demonstrated that we can achieve a rapid demineralization at 42°C with 0.1M EDTA, we then investigated the effects of aging on the time required for complete demineralization of the skulls. The time required for a complete demineralization of the skulls of 2-week-old and 1-month-old mice was 3 to 4 days, whereas for 3-month-old and 1-year-old mouse skulls it was 5 to 6 days (Fig. 6).

Elevated incubation temperature used for demineralization process did not affect stability of nucleic acids: To evaluate the effects of elevated temperatures on the stability
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of nucleic acids, we performed RT-PCR using mRNA extracted from the tooth sections. The amplification of Gapdh mRNA transcripts at 37°C, 42°C or 50°C all showed similar levels (Fig. 7).

DISCUSSION

We set out to optimize the demineralization method in order to make it shorter and safer for histological analysis of mouse teeth. In the process of this optimization, we addressed some of the key issues related to the effects of the decalcifiers and fixatives used for preparing the mouse skull specimen. Our study indicates that the demineralization of mouse skulls at 42°C using 0.1M EDTA can significantly shorten the time required for complete demineralization, while also retaining sufficient β-galactosidase activity.

DSPP, a key dentin extracellular matrix protein secreted by odontoblasts, plays a crucial role in the mineralization of predentin to form mature dentin [18-21]. We previously reported the isolation and characterization of the murine Dspp gene [22] and the validation of its promoter sequence [16]. The Dspp-LacZ transgenic mice used in the present study display spatial and temporal expression patterns similar to endogenous DSPP expression profiles [16]. In these mice, the LacZ gene, which is placed under the DSPP promoter sequence, is expressed as early as embryonic day 17.5 (E17.5) in preameloblasts and odontoblasts, and at E18.5 in molars. Its expression in preameloblasts is transient, whereas it stays robust in odontoblasts throughout tooth development. Because of this expression profile, the tooth specimens from these mice are ideal for testing the effects of decalcifiers, fixatives and incubation conditions on the stability of β-galactosidase activity. However, because the LacZ gene, which encodes β-galactosidase, has been widely used as a reporter gene to analyze the expression of a gene of interest, the demineralization protocol that we have optimized in the current study can be applied to other transgenic mouse models expressing β-galactosidase in teeth and bones.

We first verified the effects of various fixatives and the fixation time on the stability of the β-galactosidase enzyme in the tooth sections. We tested 0.25% glutaraldehyde, 4% PFA, zinc formalin and formalin for their effects, and found that all of them work well as fixatives to analyze β-galactosidase activity in the tooth sections (Fig. 1). As for the fixation time, we found that post-incubation of the skulls with 4% PFA for up to an hour did not have any adverse effects on β-galactosidase activity. However, extending the
incubation time to 2 hours adversely affected the enzyme activity (Fig. 2). As previously reported [17], LacZ enzymatic activity in kidney tissue was also retained during a 1-hour incubation time with 4% PFA, but not during a longer period of time.

Next, we tested the acid-based demineralization reagents, Formical-4 and Immunocal, in order to analyze their effects on β-galactosidase activity. As you can see in Fig. (3), it is clear that 0.1% EDTA, but not the acid-based demineralization reagents, maintained β-galactosidase enzyme activity after complete demineralization (Fig. 3). Chelating agents such as EDTA can bind to calcium and other ions and remove them from mineralized tissue. This reaction occurs gently, and enzymatic activity can therefore be retained during demineralization. In contrast, long-term exposure to acidic conditions could potentially destroy enzymatic activity such as β-galactosidase. Another possibility is that, because Formical-4 and Immunocal contain fixatives, β-galactosidase may be over-fixed and thus result in a lack of enzymatic activity. As shown in Fig. (4), we then explored the potential for shortening the demineralization process by increasing the temperature during demineralization, since it has already been reported that bone demineralization was accelerated by microwave treatment [10,11]. Our data indicated that incubation of mouse skulls with 0.1M EDTA at 42°C results in demineralization that is 2-fold faster than incubation at room temperature. Our data also revealed that, although an incubation temperature of 50°C can achieve demineralization the shortest incubation time, it adversely affects the enzyme activity (Fig. 5). These data suggest that incubation at 42°C is most useful for analysis of enzymatic activity in decalcified teeth. Moreover, as shown in Fig. (6), 6 days of incubation with 0.1M EDTA at 42°C was sufficient for complete demineralization of 1-year-old mouse teeth. These observations suggest that any specimens can be demineralized by incubation with 0.1M EDTA at 42°C for 6 days.

To assess the effects of elevated temperatures on the stability of nucleic acids, we extracted total RNA from the tooth sections of the skulls subjected to different temperatures for the demineralization process, and performed RT-PCR analysis. The amplification of Gapdh mRNA transcripts at 37°C, 42°C or 50°C showed similar levels, indicating that a higher temperature had no affect on the stability of nucleic acids (Fig. 7).

In summary, we have optimized the conditions for rapid demineralization of mouse skulls, which significantly shortens the time required for complete demineralization but does not affect the enzyme activity or nucleic acids in the tooth sections. We recommend 0.1M EDTA as a safe decalifier, to be used at 42°C for successful demineralization of mouse skulls. These conditions could be widely applied for the enzymatic and immunohistochemical analysis of other proteins in hard tissues. Moreover, rapid demineralization using this method will promote quicker analysis of genes and proteins implicated in tooth and bone diseases.

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