Bone Changes in Femoral Bone of Mice Using Calcein Labeling

Moon-Jung Shim

Department of Clinical Laboratory Science, Ansan University, Ansan 15328, Korea

Mice에서 Calcein 표지를 이용한 골 변화 관찰

심문정

안산대학교 임상병리과

Introduction

Bone in a live skeletal system constantly changes. Approximately 7.5% of bone turnover occurs annually. Bone modeling and remodeling are inherently involved in the alteration of bone. Traditional methods of in vivo bone growth evaluation in small animal models utilize noninvasive radiological, nuclear, and bioluminescence imaging methods. However, these do not allow detailed examination of the tissue. Fluorochrome labeling is a widely used standard technique in skeletal research, which is simple and efficient for the investigation of the dynamics of bone formation in combination with plain histology [1,2]. The use of fluorochrome labeling techniques for in vivo bone studies is not a new development. In 1956, the first application of fluorochromes in vivo experiments was reported [3].

Bone formation and remodeling in vivo can be assessed by polychrome labeling using calcium–binding fluorescent dyes. Newly forming bone tissue can be measured after vital staining with fluorochromes, such as calcein or tetracycline, which forms fluorescent complexes with calcium and can be measured with a high accuracy by use of a fluorescence...
microscope. Calcein green is a calcium chelator that adheres to regions of mineralizing bone thereby permitting localization of new bone.

In human studies, for examples, fluorochromes are used for the assessment of cancellous or endocortical bone turnover in iliac bone biopsies [4]. In animal research this technique is used for a much broader range of applications such as the measurement of the cancellous or endocortical, intracortical, and periosteal bone formation rate for the evaluation of bone elongation and for the assessment of bone resorption [5,6]. However, most previous studies were performed in rats [7–9], rabbits [10], and dogs [11] because bone of mice is brittle than other animals. Even though mice were used in the histological study, tibia bone [12] or longitudinal sections [13] were common. Therefore, in this study, 2 month old young mice were utilized and femurs were cross-sectioned at the mid-diaphysis.

The aim of the present investigation is to evaluate the deposition of calcium at both 4th and 8th weeks in the femoral bone of normal mice model using calcein labeling.

Materials and Methods

Mice were labeled by with calcein (25 mg/kg; Sigma-Aldrich Co., St. Louis, MO, USA) in PBS every 5 days for 4-week-old mice, 7 days for 8-week-old mice by intraperitoneal injection, and 2 days prior to sacrifice. All procedures were approved by the Institutional Animal Care and Use Committee of Ohio State University.

Femurs were dissected, fixed in 70% ethanol and femoral bone sections were obtained using a low speed saw with two parallel diamond blades cutting under water irrigation. The sections were dehydrated at 4°C in alcohol solution up to 100% ethanol and infiltrated in a media containing methyl methacrylate for 24 hr under vacuum. And then polymerization was started by warm water heating of the samples. After polymerization, the polymethyl methacrylate blocks were exposed to the UV light source for 1 week in refrigerator and sectioned about 80 μm thick using a Leica SP 1600 microtome (Leica Microsystems, Nussloch, Germany). To mount the glass slide, a UV light curing glue was used because of its short curing time and high bond strength. Calcein levels on the surface of bone were analyzed using a Olympus DP 70 fluorescence microscope (Olympus America Inc., Center Valley, PA, USA).

Results

With the vital labeling technique, the new forming bone area was marked with the calcein–calcium complex, which can be seen in a fluorescence microscope. Therefore with this technique it is possible to determine the new bone area qualitatively and quantitatively.

In this study, calcein green labeling has confirmed the presence of newly formed bone tissue. And calcein has a high calcium affinity which translates into a relatively broad fluorescent band. After 4 and 8 weeks, microscopic analyses of the surfaces verified as good new bone formation which showed that new bone was formed as a surface-related phenomenon. The double layer after 8 weeks were observed broader than 4 weeks, and showed intense deposition of calcium in the bone after the application of 8 weeks (Fig. 1, 2). It means that was deposited at the site of active mineralization.

These days this technique of fluorochrome labeling is used in order to investigate bone formation and remodeling processes in vivo at different time interval.

Fig. 1. Fluorescence image of the femoral cross-section at 4–week-old-mouse. Bone formation was visualized by double calcein labeling in the femoral bone. Representative picture: ×10, scale bar: 200 μm.
Fig. 2. Fluorescence image of the femoral cross-section at 8-week-old-mouse. Bone formation was visualized by double calcein labeling in the femoral bone. Representative picture: ×10, scale bar: 200 μm.

Discussion

Histomorphometry is the gold standard method for the assessment of bone remodeling. Although other methods, such as serum or urine biomarkers, can provide whole body estimates of remodeling, only histology can provide site-specific remodeling data. In vivo labeling of bone with fluorochromes is a widely used method to assess mineral apposition rate and bone formation rate in bone histomorphometry [14-16]. In sections of undecalcified bone, fluorochromes can be visualized by their specific fluorescence under UV excitation. Prolonged exposure to UV light results in irreversible fading of fluorochromes bound to bone mineralization sites. Therefore, possible fading of fluorochromes precludes the use of light for polymerization of the methacrylate in a routine plastic embedding of undecalcified bones suitable for both bone histomorphometry and histochemical and immunohistological techniques.

Fluorochrome labels, when bound to calcium ions, can be incorporated at sites of mineralization in the form of hydroxyapatite crystals [17]. This means that the label indicates all sites of bone formation or dentin deposition, and hypertrophic cartilage in the growth plate. In the first 24~36 hr after administration, the label is stabilized. As a result, the fluorescent label demarcates the mineralization front at the time of administration and can be detected in histological sections without any further staining or decalcification.

During growth, bone continually undergoes changes of shape and structure. Accretion in length and thickness, modeling, and drift activities lead to different morphologies and determine the relative position among various skeletal parts. After completion of growth, pathological processes, aging, and degeneration are responsible for structural alterations, and healing processes and adaptation to new functional demands require new structural characteristics. Certain stains are able to bind to sites, which are calcifying. These stains stay there for a long time and serve as markers, allowing the identification of tissues, which are mineralized at the time of dye administration.

Animal models are commonly used in the study of skeletal biology and serve as useful tools to delineate mechanisms underlying bone loss and skeletal fragility. Bone-seeking fluorochromes provide a useful tool for analyzing changes of bone morphology. The presence of the fluorochromes indicates site, time and amount of bone deposition, and enhances the information continued in bone specimens. The administration of several fluorochromes, distinguishable by color, offers distinct advantages over single labels. With the method of vital labeling, bone forming areas can be measured. The apposition rate was calculated from the distance between the periosteal surface and the first calcein injection line for the first two time points of sacrifice of each group. Mice are used widespread in biological studies of musculoskeletal disease. It is possible to determine parameters, such as mineral apposition rate, mineralizing surface, and bone formation rate. However, unfortunately in this study, mice were too young and bone were easy breaking to determine parameters.

As I mentioned, most previous studies were performed in rats, rabbits, and dogs because bone of mice is brittle. However mice model can obtain significant experimental results within a short period, incurring less side effects, and be suitable to apply in vivo experiment using genetically modified mouse such as knock-out mouse.

In this study, calcein has been confirmed as a marker for the identification of bone growth, which is indicated by the green color on the photomicrographs in the femoral bone of
4-week, 8-week old normal mice model. Furthermore, examining human bone biopsies has shown different fluorochromes have different binding affinities, so other fluorochromes should be studied, including tetracycline which is used clinically. This methodological progress may provide basic information for interpreting the bone formation and regeneration to pharmacologic or genetic manipulation in mice.

요 약

골은 일생에 걸쳐 지속적인 재형성과정을 거쳐면서 유지되고 이러한 기전에 대한 연구는 골다공증을 비롯한 골대사 질환의 병태생리와 치료에 있어 큰 진전을 이루고 있다. 특히 생체 내 골형성 및 재생과정을 연구하는데 있어 형광표지자를 이용하는 방법이 널리 알려져 있는데, 그 중 calcein은 갈슘 킬레이터로 골이 새롭게 형성하는 부위에 녹색을 띠므로 유용한 마커로 사용된다. 그러나 대부분의 골형성 연구에서 실험동물의 경우 표본제작을 할 때 크기가 작고 뼈가 부숴지기 쉬워 rat이나 rabbit을 이용하였으며, mice의 femur를 cross-section해서 관찰한 연구는 거의 없는 실정이다. 그래서 본 연구에서는 어린 mice를 실험동물로 이용하였으며, 생체 내 calcein을 4주간, 8주간 투여한 후 골 형성 변화를 형광현미경으로 관찰한 결과 8주차 쥐에서 4주차보다 진하고 골 형성 간격도 넓게 관찰된 것을 확인 할 수 있었다. Mice는 빠른 시일 내에 결과를 얻을 수 있고 부작용이 적은 장점이 있어서 knock-out mice를 이용한 생체 내 실험에 활용하기 적합하다고 생각되며, 골 형성 속도 평가 등 다양한 분야에서의 골 형성과 재생연구에 있어 기초 정보를 제공할 것으로 기대한다.

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