

Murine Axial Compression Tibial Loading Model to Study Bone Mechanobiology: Implementing the Model and Reporting Results

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ABSTRACT: In vivo, tibial loading in mice is increasingly used to study bone adaptation and mechanotransduction. To achieve standardized and defined experimental conditions, loading parameters and animal-related factors must be considered when performing in vivo loading studies. In this review, we discuss these loading and animal-related experimental conditions, present methods to assess bone adaptation, and suggest reporting guidelines. This review originated from presentations by each of the authors at the workshop “Developing Best Practices for Mouse Models of In Vivo Loading” during the Preclinical Models Section at the Orthopaedic Research Society Annual Meeting, San Diego, CA, March 2017. Following the meeting, the authors engaged in detailed discussions with consideration of relevant literature. The guidelines and recommendations in this review are provided to help researchers perform in vivo loading experiments in mice, and thus further our knowledge of bone adaptation and the mechanisms involved in mechanotransduction. © 2019 Orthopaedic Research Society. Published by Wiley Periodicals, Inc. *J Orthop Res* 38:233–252, 2020

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A description of the relationship between mechanical stress (or strain) and internal skeletal structures was first detailed in the mid-late 1800s by von Meyer, Culmann, and Wolff.^{1,2} The culmination of their works can be summarized by the trajectory theory for bone, which describes the alignment of cancellous bone tissues with principal stress trajectories in the bone.² Roux³ added uniquely to this body of work the idea that functional adaptation of the skeleton is a dynamic and self-regulating process, rather than supposing that bones always exist at some static optimum. These theories have since been extended to encompass functional relationships between mechanical loading and cortical bone geometry,^{4–6} bone curvature,^{6,7} collagen fiber orientation,^{8–10} and vascular canal (osteonal) orientation.⁴ Investigation of skeletal functional adaptation through human exercise studies and pre-clinical animal models that used controlled applied loads and functional disuse began in the 1960s^{11–16}; and provided experimental evidence for the adaptive nature of the skeleton in response to mechanical loading. The results of these studies collectively led to mechanistic hypotheses predicated upon components of skeletal loading history in regulating bone modeling and remodeling responses at local skeletal sites (Fig. 1).^{17–22} These hypotheses led to an entire field of computational bone remodeling in response to simulated

mechanical stimuli^{17,23–27} and laid the foundation for much of our current understanding of skeletal mechanobiology (a.k.a., bone adaptation).

Although local bone strain magnitude and loading history (the cumulative product of load-induced bone stress/strain and number of load cycles) were initially identified as key stimuli driving local formation and resorption, studies have reported bone formation at spatial sites experiencing low bone strain magnitude.^{31,32} Thus, to better understand the diversity of factors that influence load-induced bone (re)modeling, beyond simply the magnitude of the applied loads, a number of pre-clinical animal models have been developed. Use of these models has advanced our understanding of the sensitivity of the skeleton’s mechanobiological mechanisms to strain or loading rate, the number of load cycles applied within a given load bout, the duration (days or weeks) of the stimulus, and the role of rest-insertions or refractory periods in re-setting the sensitivity of the skeleton to subsequent load stimuli. Not only have these studies developed the foundation for physical strategies for stimulating bone formation or maintaining bone mass, but they have also led to the elucidation of molecular and cellular mechanisms for bone functional adaptation and tissue anabolism. While the skeletons of larger animal models (rabbits, dogs, pigs, and sheep) may better represent human bone physiology, in the past 20 years, rodents have been predominantly used for these studies because of their relatively low cost, short lifespans, and the ability to manipulate their genomes to study specific cellular pathways. Targeting bone-formative mechanobiological pathways in combination with optimized physical regimens could synergistically provide more effective therapies for

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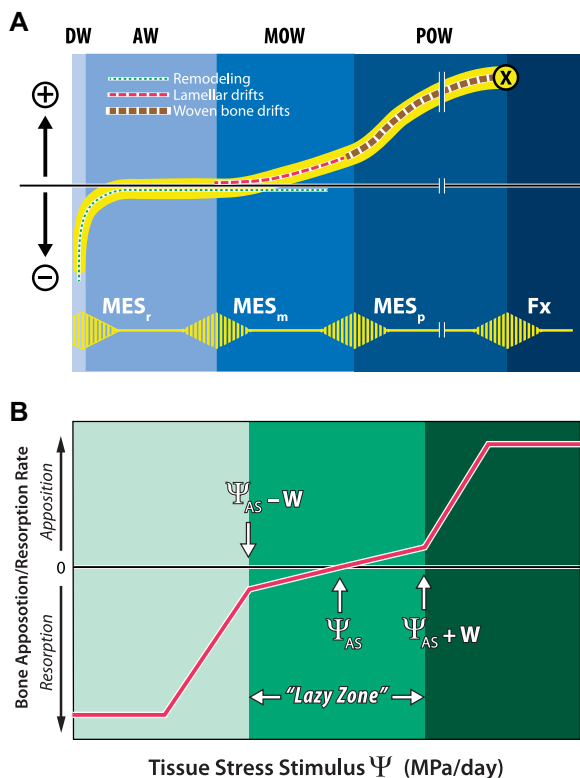


Figure 1. (A) Illustration of the mechanostat hypothesis, modified from Harold Frost's original drawing.²⁸ The horizontal axis depicts peak bone strain, and the vertical axis depicts net loss (−) or gain (+) of bone mass. The lower (yellow) pulsed line shows the threshold values of minimum effective strain (MES) for remodeling (MES_r), modeling (MES_m), microdamage (MES_p), and fracture strain (Fx). The regions labeled at the top represent the disuse window (DW), adapted window (AW), mild overload window (MOW), and pathologic overload window (POW). The dotted line curves illustrate disuse and maintenance of bone and bone formation, originally suggested by Carter.¹⁸ (B) In a model proposed by Carter and colleagues,^{17,29,30} the horizontal axis represents the cumulative daily stress stimulus (a combination of stress magnitude and number of loading cycles experienced per day) relative to the rate of bone formation or resorption on the vertical axis. The attractor stimulus (Ψ_{AS}) is similar to Frost's MES_m and the width of the "lazy zone" is $\Psi_{AS} \pm W$, where W is the half-width of the range of stress stimuli experienced during normal daily activities. [Color figure can be viewed at wileyonlinelibrary.com]

age-related bone loss, fracture healing, and other musculoskeletal diseases in humans and other species.

The goal for this review is to describe one commonly used rodent loading model, the mouse axial compression tibial loading model and key considerations for implementing and reporting the results using this model. Other reviews of the tibial axial compression model convey some of the aspects discussed here.^{33–36} However, where these previous reviews focused on providing detailed methodological guidance for conducting tibial loading experiments, here we emphasize the various pre- and post-experimental factors to consider in the successful implementation of this model. We specifically present (i) the basic software and hardware requirements for successfully conducting this model, (ii) methods for characterizing the load-induced

mechanical environment in the tibia, (iii) the loading protocol parameters to consider in implementing this model, (iv) biological outcome measures that can be assessed using this model, (v) animal welfare considerations, and (vi) a select list of validated protocols for studying tibial adaptation to applied compressive loads.

A REVIEW OF ANIMAL MODELS USED IN SKELETAL LOADING STUDIES

A number of pre-clinical skeletal loading models have contributed to our understanding of skeletal mechanobiology and bone functional adaptation (Table 1). These models maintain a high level of control over the loads applied to the skeleton, which consequently allows for a highly repeatable load stimulus and induced skeletal strains that can be accurately characterized empirically and computationally. Controlled loading models contrast exercise models in which the loads applied to the skeleton are not easily controlled and can be difficult to characterize. Furthermore, controlled loading models are able to isolate the effects of the mechanical stimulus to a target bone element, without inducing systemic physiological loading effects that might influence skeletal remodeling more broadly. Thus investigators are able to use the contralateral non-loaded limb as a within-animal control, which reduces the overall cost of animal life.³⁷ Studies by one group have challenged this notion as they demonstrated that loading of a single limb (ulna) in young rats had systemic skeletal effects that led to changes in the contralateral limbs as well during the loading experiments.^{38,39} However, similar systemic effects were not reported by others using rat ulnar loading⁴⁰ or mouse tibial loading,^{37,41,42} consistent with the notion that loading effects are predominantly local. Thus, the contralateral limb generally can be used as an internal control. We recommend that investigators confirm that the contralateral limb is equivalent to the limbs of non-loaded control animals in their own laboratory. Regardless of whether unilateral loading causes systemic-based effects on the skeleton, comparison of loaded versus contralateral control limbs will account for any systemic effects on the loaded limb's response to load by normalizing the structural and histomorphometric parameters measured in the loaded limb to those measured for the contralateral limb.

The majority of the non-surgical, controlled skeletal loading models exclusively examine cortical bone adaptation to mechanical loading in rodents. In the rodent four-point bend tibial model (Table 1C), two load points on both the medial and lateral tibial diaphysis are used to induce a pure bending moment in the central diaphysis.⁵⁰ In this model, loads are applied directly to the periosteum which elicits a periosteal contact response that potentially limits the use of this model to analyses of the endosteal response to load.⁵¹ Subsequent models have addressed this limitation by applying load through the joints, far removed from the cortical diaphysis. The rodent ulnar loading model examines cortical bone adaptation by applying compressive loads along the ulna's long axis while taking advantage of the bone's

Table 1. Key In Vivo Animal Models for the Study of Bone Mechanobiology

	Species	Tissue compartment	Key parameters
A	• Rabbit ¹² • Sheep ⁴³ • Dog ¹¹	• Cortical	• Strain rate ⁴⁴ • Load magnitude ⁴⁵
B	• Turkey ⁴⁶ • Chicken ⁴⁷	• Cortical	• Dynamic vs static loads ⁴⁸ • Strain magnitude ⁴⁹
C	• Rat ⁵⁰ • Mouse ⁵¹	• Cortical	• Strain magnitude ⁵² • Rest insertion ⁵³ • # of loading cycles per session ⁵⁰
D	• Mouse ⁵⁴	• Cortical	• Rest insertion ⁵⁵
E	• Rat ⁵⁶ • Mouse ⁵⁷ • Rabbit ⁵⁸	• Cortical	• Strain rate ⁵⁹ • # of cycles per session ⁶⁰ • # of load sessions ⁶⁰ • Recovery time between sessions ⁶¹ • Strain magnitude ⁶² • Dynamic vs. static loading ⁶²
F	• Mouse ⁶³ • Rat ^{64; 65}	• Cortical • Cancellous	• Strain magnitude ^{66; 67} • # of cycles per session ^{66; 68} • Rest insertion ⁶⁸ • # of sessions/wk ⁶⁶ • # of wks ⁶⁶
G	• Rat ⁶⁹ • Mouse ⁷⁰	• Cancellous	• Load magnitude ^{70; 71} • # of wks ⁷¹
H	• Rabbit ⁷²	• Cancellous	• # of loading cycles per session ⁷³ • # of loading sessions ⁷³ • Load magnitude ⁷⁴

References are shown for the initial report for each model as well as parameterization studies using the model. The models shown are (A) tibial axial loading though surgically-placed wires or pins, (B) avian surgically-isolated ulnar loading, (C) rodent four-point tibial bending, (D) mouse tibial cantilever bending, (E) rodent ulnar axial compression, (F) rodent tibial axial compression, (G) rodent vertebral compression, (H) rabbit distal femur cancellous compression.

curvature to induce a bending moment along the diaphysis^{56,57} (Table 1E). Because the carpus and olecranon process are used as the primary points of load application, no direct load is placed on the ulnar diaphysis. This model is the most well-characterized of the current rodent loading models in terms of the interactive effects of

the different load parameters regulating cortical bone response to mechanical load. This model is still commonly used in mice and rats and was recently adapted for rabbits.⁵⁸ While cancellous bone tissue is present in the olecranon and distal ulna, the volume of tissue present is quite small, especially in mice. As a result, no

studies using this model have reported the effects of applied load on cancellous bone. A final purely cortical bone adaptation model is the mouse tibial cantilever model, which secures the knee against medial-lateral movement and applies a load laterally to the distal tibia to induce medial-lateral bending of the tibial diaphysis⁵⁴ (Table 1D).

Fewer animal models are available to examine cancellous bone adaptation, with all but one requiring surgical intervention to isolate cancellous tissue.^{69,75} The rat tail vertebral model is a surgical model that was developed in the early 1990s,⁶⁹ and continues to be used in a scaled-down version in the mouse⁷⁰ (Table 1G). This model applies load to adjacent vertebrae through pins, which can introduce a healing response. Furthermore, this model induces a non-physiological loading mode to generate supraphysiological strain that can produce pin deformation and pull out.⁷⁶ Also, while the lumbar vertebrae (L5) in female C57Bl/6 mice undergo age-related bone loss,⁷⁷ the caudal vertebrae do not.⁷⁸ Thus, comparing adaptation at caudal vertebral sites in aged mice to aged humans may have limited clinical relevance and requires further study.⁷⁹

One of the most recent long bone loading models to be developed and receive widespread use is the mouse axial compression tibial model, which can be used to examine both cortical and cancellous bone adaptation^{63,80} (Table 1F). Like the rodent ulnar model, the tibial model does not apply direct loads to the diaphysis. In this model, the distal femur and foot are fixed with plastic or metal fixtures and compressive loads are applied along the length of the tibia through the knee and ankle joints. The axial compression tibial model uses the tibia's natural curvature proximal to the tibiofibular junction to induce highly repeatable tensile strains on the anterior-medial surface and compressive strains on the posterior-lateral surface of the diaphysis. While the bone strains required to induce bone formation are supraphysiological for the mouse, the strain distribution is fairly physiological, agreeing with the locomotor strain distributions measured *in vivo*⁸⁰ or modeled via finite element (FE) simulations.⁸¹ This controlled loading model has been used to examine the effects of applied strain magnitude,^{66,67,82} load frequency (cycles per load bout),^{66,68} experiment length (number of daily load bouts),⁶⁶ rest-insertion,⁶⁸ sex,⁸³ sex hormones,⁸⁴ and age^{42,85–89} on the cortical bone response to load. Furthermore, this model has also been used in studies seeking to describe the role of specific proteins in bone anabolism and mechanobiology.^{90–95}

The axial compression tibial loading model uniquely enables investigation of these same mechanical and genetic variables in the cancellous bone tissues of the proximal tibia. This specific feature is important because most osteopenia-related bone fractures occur at cancellous-rich bone sites.⁹⁶ The proximal tibial metaphysis has a cancellous bone volume fraction varying between 5% and 20% in C57Bl/6 and BALB/c mice depending upon age and sex.^{42,79,83,86,97–99} While the absolute bone

volume in the proximal tibial metaphysis is fairly similar to the ulnar olecranon, due to anatomical differences, histomorphometric and molecular analyses may be more technically-accessible for the proximal tibia than for the olecranon. Thus, the axial tibial loading model is able to examine the functional adaptation of cortical and cancellous bone tissues to fairly physiological loading as an integrated unit in a single long bone.

IMPLEMENTING THE MOUSE AXIAL COMPRESSION TIBIAL LOADING MODEL

The required elements for successfully implementing the axial compression tibial loading model include animal anesthesia, fixation of the knee and foot during loading, a device for controlled linear actuation, and a load cell. The system must also include software for designating load waveform parameters and that provides force feedback control to apply temporally repeatable peak loads of a specific magnitude to the hindlimb. Both custom and commercial linear actuator devices (hydraulic, electromagnetic, and moving coil) with in-series load cells and software-based force

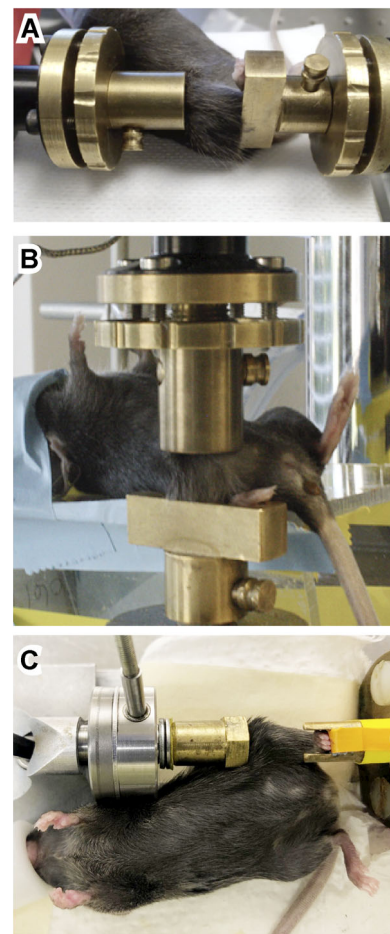


Figure 2. Photos of different loading devices used to perform *in vivo* tibial loading in mice.^{42,63,80,100} (A and C) Mice in supine position with a horizontally-oriented tibia. (B) A supine mouse with vertical tibial orientation. [Color figure can be viewed at wileyonlinelibrary.com]

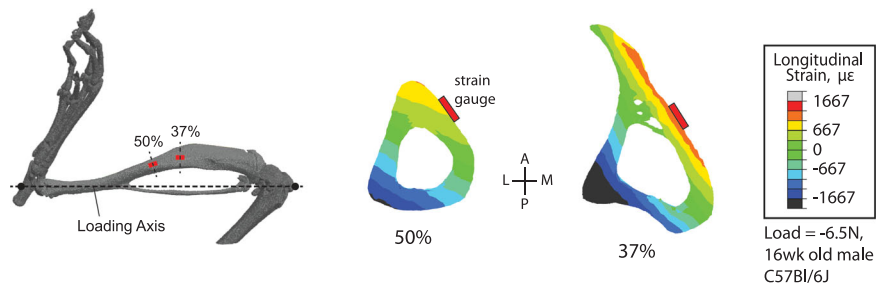


Figure 3. Microcomputed tomography reconstruction of the mouse hindlimb showing common sites for strain gauge attachment for tibial loading studies and calibration of finite element (FE) models (modified from Biewener¹⁰⁶). Transverse FE cross-sections of the tibia showing that the common strain gauge locations do not coincide with peak modeled strains, which typically occur in the compression zone on the posterior-lateral surface of the tibia, a common site for bone formation in this model. [Color figure can be viewed at wileyonlinelibrary.com.]

feedback control have been used to implement this model (Fig. 2). Most commonly, commercial materials testing systems with linear actuators (e.g., Dartec, Instron, TA Instruments/Bose, and Biomomentum) are repurposed for *in vivo* bone loading.

Beyond selecting proper hardware with software control, another important consideration is the alignment and shape of the fixtures used to hold the knee and foot during loading. For the fixtures, the critical design constraint is that the central points of force application on each end should align along an anatomical axis that transmits load along the tibia and limits flexion and extension of the knee during the load cycle (Fig. 3). A large design space can satisfy these critical specifications. The point of contact at the femur can be fairly localized, while the contact with the foot is necessarily broad. Dorsiflexion of the foot in most established systems ranges from 30° to 56°^{42,83} (Supplementary Fig. S1). The foot holder must be wide enough to not laterally pinch the foot (4–5 mm for mice) to avoid wear sores and calluses over the course of a multi-week loading study. Such calluses can also appear on the heels and should be monitored.

The knee fixture is a critical consideration in conducting tibial loading studies but is rarely described in most protocols. The knee fixture should be designed to hold the distal femur and patella without applying load directly to the tibia. This configuration ensures that the applied load will be transmitted through the distal femoral condyles to the tibial plateau. In many published studies, the knee fixture features a semi-spherical cup to hold the distal femur. The dimensions of the knee cup can have profound effects on the outcome of a study. One of the authors (R.P.M.) has found that a cup with 6 mm inner diameter and 3 mm depth can be used to apply loads as high as 13–15 N with no palpable disruption of the knee anatomy following multiple days of loading, while a knee cup with 10 mm inner diameter and 5 mm depth routinely causes knee dislocation and joint instability, with likely ACL rupture, within 100 applied load cycles using loads as low as 8 N. Similarly, one group reported using a cup with a diameter of 9 mm and a depth of 2–3 mm that could safely be used at 7 N loads, but caused ACL rupture at 9–10 N loads.¹⁰¹ Knee flexion could be altered by variations in fixture cup diameter and

depth. Deeper or wider cups can lead to increased joint mobility during loading that makes injury more likely. A potential link between knee flexion, femoral-tibial translation and knee cup geometry has not been formally investigated. This area deserves more attention since not all the potential users of this model may be trained to identify ACL rupture or other knee injuries. Such injuries should be avoided because of local or systemic pathologic responses that would affect the interpretation of load-induced skeletal changes. We have provided drawings of foot and knee fixtures used currently by some of the authors (Supplementary Fig. S1). Another possible fixturing approach is to make custom-molded pads from hardened modeling clay or PMMA.^{102,103} This approach would allow a snug fit for the foot and knee but would require different sets of molds for mice of different age, sex, or genotype.

Different research groups have positioned mice with different body and limb orientations during loading; supine with the limb held horizontally (supine-horizontal),^{63,68,93} prone with the limb held vertically (prone-vertical),^{88,101} or supine with the limb held vertically (supine-vertical).^{80,100,104} These configurations are often dictated by physical restrictions caused by the hardware used to load the hindlimb. The potential effect of body positioning and limb orientation during loading has not been thoroughly examined. However, adaptive responses reported for prone-vertical and supine-horizontal orientations are generally similar.⁴¹

CALIBRATING AND ASSESSING THE MECHANICAL ENVIRONMENT IN THE TIBIA DURING AXIAL COMPRESSION LOADING

The goal for controlled loading models is to relate a quantifiable, known mechanical stimulus to a biological response. Therefore, the mechanical environment must be known and comparable across animals. The “mechanical environment” is the imposed load, the ensuing deformation, and the resulting stresses and strains in the tissue. For a given load, the stresses and strains induced in the bone are dependent on bone geometry (cross-sectional size and shape) and material properties (elastic modulus). The applied load is almost always the controlled parameter for *in vivo* loading. “Load-matched” protocols apply the same peak load to all

experimental groups, which is appropriate for answering the question, "How does the bone respond to a specific load magnitude?" For load-matched protocols tissue stresses and strains may vary across experimental groups, particularly if inter-group differences in bone size or material properties are present due to sex, age, or genotype. If such differences exist, the "load-matched" approach is typically not the best choice.

"Strain-matched" protocols adjust the peak loads, if necessary so that the peak longitudinal strains are similar across groups. We recommend this approach as the gold standard, based on widespread evidence that the local strain stimulus drives the local (re)modeling response. To accomplish "strain matching," a load-strain calibration is typically determined by attaching a strain gauge on the medial surface of the tibia. Some protocols have used two⁸⁰ or three¹⁰⁵ gauges. Locations for placement of the strain gauge are limited because of the size and shape of the tibia. Along most of the cortical diaphysis proximal to the tibiofibular junction, the medial (or anterior-medial) surface experiences tension (+strain) and the lateral (or posterior-lateral) surface experiences compression (-strain) during axial compression loading.^{86-89,*} Consistency in gauge placement is critical so that load-induced strains at a specific bone site can be directly compared for each group. Detailed protocols for strain gauging have been published previously.^{33,34}

Calibration of the strain can be performed in vivo (in live anesthetized animals) or ex vivo (in cadaveric mice). Although in vivo calibration ensures conditions are exactly as in the experiments, ex vivo calibration is easier to perform. Strain-matched protocols ensure that the mechanical environment in the tissue is comparable between groups, and is often used when comparing mice of different sizes, ages, or genotypes.^{80,93,105} A longitudinal strain of 1,200–2,000 $\mu\epsilon$ on the medial diaphysis is often used as a target value to induce an adaptive response. However, because strain magnitude varies greatly with location on the tibia, this target value is highly dependent on strain gauge placement. Thus, a specific description of calibration gauge placement should be reported for each study in which this calibration is conducted.

Two common sites for strain gauge attachment are the mid-diaphysis (50% bone length) and a site at about 37% of the bone length from the proximal end of the tibia (Fig. 3). Load-induced strain magnitudes differ at these two sites, typically with greater strains at the 37% site. At either of these two sites, the peak strains at that bone level (i.e., the location along the long axis of the tibia) do

not occur at the specific site of gauge placement. If knowing the peak induced strains is an important factor for a given study these can be estimated via FE or planar strain analysis.¹⁰⁵⁻¹⁰⁷ The latter approach requires that three strain gauges be attached to a bone at the same proximal-distal plane which, in the mouse, can only realistically be accomplished *ex vivo*.

Strain gauges measure strain at specific locations on the tibia; however, the strain in the tibia is not homogenous and sampling a single location may not be representative. Digital image correlation (DIC) is an *ex vivo* technique that provides surface strains over a region. A speckle pattern is applied to the surface of the tibia (with soft tissues removed on a cadaveric tibia) and imaged during loading. The deformation of the speckle pattern on the surface allows determination of longitudinal, transverse, and shear strains across the entire surface of the tibia.^{104,108} At least two cameras are used to record the speckle deformation during loading. The resolution of the strain measurement is dependent on the speckle pattern and the processing parameters used in determining strain. Studies using DIC to assess strain have demonstrated that local values of peak strain are much higher than those measured with strain gauges, and the distribution of strain in the tibia becomes more homogeneous with adaptation due to loading.¹⁰⁸ To calibrate applied loads to achieve strain-matched protocols with DIC, one can use peak strain measures or an average strain over a defined area of the bone to achieve greater assurance that a larger area of the strain field is similar across groups than when a strain gauge is used to measure strain over a relatively small area of bone. While average strain values ultimately reduce all the spatial information provided by DIC into a single number, and therefore may not seem much different than strain gauging, the regions of bone that can be assessed by DIC are more extensive than can be achieved with strain gauges. DIC is particularly useful if the spatial characterization of the mechanical environment is needed to answer the research question.

FE modeling of the tibia is often used to determine non-measurable mechanical parameters (Fig. 4). Tibial geometry is readily obtained from microCT scans. Material properties generally are assumed to be linear elastic and may be homogenous or heterogeneous, and isotropic or orthotropic.^{35,105,109-113} The tibial and fibular growth plates may or may not be modeled. The properties of the proximal fibular attachment can affect results.^{110,113} As with any FE model, the assumptions limit the validity of the results, and the effects of assumptions on the results should be examined through sensitivity analyses and model validation with experimental measures.¹¹⁰ Parametric studies have shown that heterogeneous material properties and location and orientation of the applied load significantly influence the strains engendered.^{110,115} If the centroid of the applied load is not aligned with the centroid of the fixed elements at the other end of the bone, excessive bending

*De Souza et al.⁸⁰ reported load-induced compressive strains (-strain) on the medial surface of the tibia and tensile strains (+strain) on the lateral surface via strain gauge measures taken during calibration of their first loading system. This unique strain distribution can be traced to an error in the recording of strain signs during data collection (A. Pitsillides, personal communication).

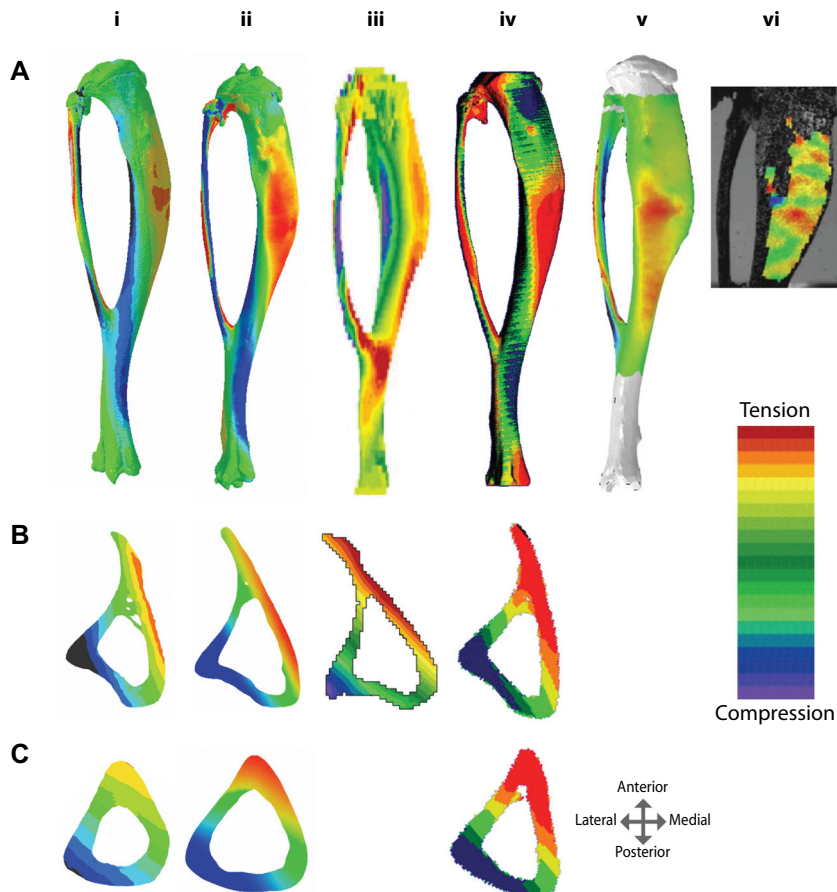


Figure 4. (A) Finite element models of murine tibiae with simulated applied axial compressive loads from (i) based on¹¹⁰ (ii),¹¹⁴ (iii),¹¹¹ (iv),¹⁰⁵ and (v),¹¹² (vi) DIC of surface strains.¹⁰⁸ All tibiae in (A) are viewed from the medial surface, except (iii) which is in lateral view. Tibiae shown are from mice of different ages and model different applied load magnitudes, but one can appreciate that while there are some differences in whole bone strain patterns (particularly ii and iii, which have high strains at the tibiofibular junction), strain distributions at the (B) mid-point between the proximal bone end and the tibiofibular junction (~37% bone length) and (C) mid-shaft (50% of bone length) have quite similar strain distributions, likely owing to the strong effect of the bone curvature on the induced strain environment. [Color figure can be viewed at wileyonlinelibrary.com]

may be imposed and high strains at the tibiofibular junction ensue. Bending occurs even when the centroids are aligned due to the curvature of the tibia, but aligning centroids better ensures that FE-simulated axial loading matches actual loading conditions.

Calibration and characterization of the mechanical environment during tibial loading are not required for every study and depend on study design. For example, groups of mice of the same genotype, age, and gender will have a similar mechanical environment, and it is sufficient to apply a load known to elicit a response for that genetic strain, age, and gender. If comparison of mechanoadaptation is required across groups that differ in genotype, age, gender, or treatment, calibration of the load is critical to ensure that the mechanical environment is the same for all groups. For these studies, *a priori* strain estimation with strain gauges remains the gold standard.^{86,109,116,117}

EXPERIMENTAL PARAMETERS TO CONSIDER FOR IN VIVO LOADING EXPERIMENTS

With the increasing use of electromagnetic materials testing systems to apply mechanical loads to the murine tibia, more detailed manipulation of the loading protocol is possible. Although these systems allow for increased accuracy in delivering the desired loading regimen, they also allow for an increase in the number of customizable parameters, which consequently makes

comparison of different studies from different groups challenging. Comparisons are further confounded because details of the loading protocol are frequently incompletely reported.

A key consideration when designing a mechanical loading protocol is the study rationale. Different research questions will require different loading protocols to generate the desired skeletal response. A single loading protocol will clearly not be suitable for all studies. For example, one investigator may wish to induce microdamage to investigate bone repair or woven bone formation in one study,^{118,119} while in another study lamellar bone formation may be of interest. Generally, a submaximal response may be preferable so that any enhancement of the loading response is detectable for interventions that are expected to cause additive or synergistic effects on bone formation.¹²⁰ Finally, the desired biological response should be considered when deciding upon a loading protocol. For example, enzymatic changes are evident as early as 6 min following a single load bout¹²¹ and changes in gene expression one hour after a single load,^{94,122} while for mineralized bone formation generally 1–2 weeks of loading is required.⁶⁶

A number of different characteristics define any experimental loading protocol and could synergistically affect the skeletal response to mechanical loading, including: the shape of the waveform (triangle, trapezoid,

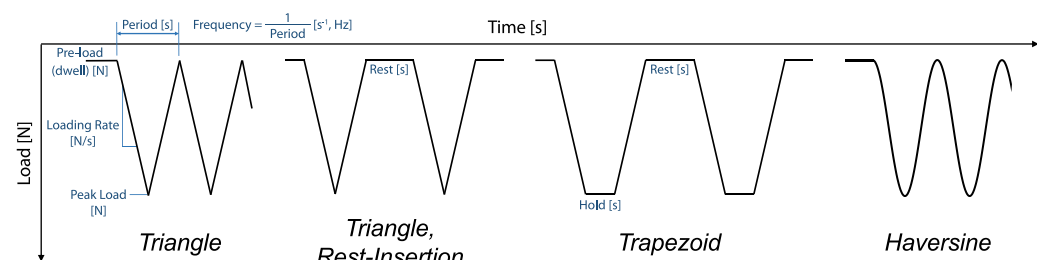


Figure 5. Illustration of the different waveforms used in tibial loading studies, including identification of their key characteristics.

and haversine), the dwell and peak load magnitudes, the rate of load application or duration of a single load cycle, the frequency of the applied loads, the number of cycles applied each day, the number of days over which to load, and the presence of any low-magnitude “rest” phases (Fig. 5). The specific combination of these protocol characteristics should be thoroughly described in any published loading study.

Static Versus Dynamic Load

The first reports of experimentally applying mechanical load to animal bones were by Heirt who used a surgical approach for loading the rabbit tibia¹² (Table 1A). A key finding was that bone formation occurred in response to dynamic but not static loading. These findings were developed further by Lanyon using the isolated turkey ulna loading model (Table 1B) confirming endosteal bone resorption and increased porosity in response to static loading, whereas periosteal bone formation occurred with dynamic loading.⁴⁸ The importance of a dynamic load stimulus has also been shown using the tibial four-point bend and ulnar axial compression models^{62,123} (Table 1C and E). Studies investigating bone formation following loading will therefore almost exclusively apply a dynamic load waveform.

Loading Waveform

Various waveforms have been used to investigate bone formation in response to mechanical stimulation, including triangular,⁶³ sinusoidal,¹²⁴ or

trapezoidal⁵⁷ (Fig. 5). A sinusoidal waveform (e.g., haversine) possesses variation in the loading rate (slope) during loading and subsequent unloading, while triangle and trapezoid waveforms maintain constant loading rates during the entire loading and unloading phases, albeit with abrupt changes at the end of each loading segment. These abrupt changes in slope can introduce higher-order frequencies that have not been characterized in tibial loading studies but may affect the osteogenic potential of the different waveforms. Applying Fourier transform to the different waveforms would reveal the relative importance of different frequency components. The effects of waveform shape and frequency composition on the tissue-level response to load have not been comprehensively examined in the tibial loading model.

A recent study attempted a comparison between triangle and sinusoidal (haversine) waveforms, but found that their actuator system was not able to replicate the triangle waveform command signal in the mouse hindlimb, such that both waveforms appeared fairly sinusoidal as measured by the load cell⁶⁶ (Fig. 6). Success in obtaining reliable triangle and trapezoidal waveforms is typically related to accurate tuning of the actuator-load cell feedback control, which some systems may be better able to achieve. Given the possible variance between command load signal and actual load output, investigators must validate the fidelity of the measured load signal relative to the programmed command signal. Where

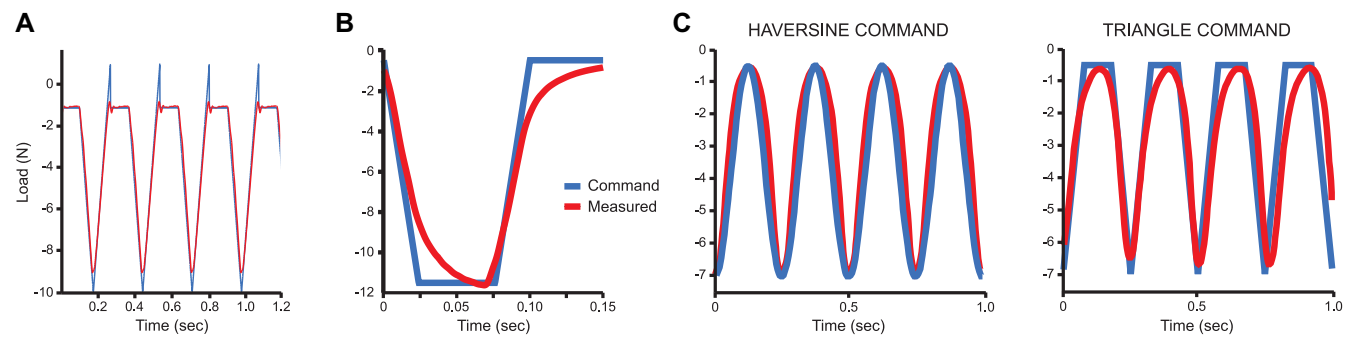


Figure 6. Example load waveforms showing degrees of fidelity between the software-programmed, command waveforms and the load cell-measured waveforms for (A) triangle waveform,⁶⁸ (B) trapezoidal waveform, and (C) haversine and triangle waveforms.⁶⁶ These waveforms were commanded on different mechanical systems: (A) TA Instruments Electroforce TestBench, (B) TA Instruments Electroforce 3100, (C) Instron ElectroPuls E1000. In (A), the small “recovery” spike needed to achieve the correct command pre-load results in a short delay of the signal and a load frequency of 3.75 Hz, as opposed to the intended 4 Hz. Some custom devices do not exhibit this spike when conducting triangle waveforms.⁶³ [Color figure can be viewed at wileyonlinelibrary.com]

possible, the actual and intended load and load rates should be reported.

Strain Magnitude

Load-related bone formation correlates with the peak strain magnitude attained during loading. A clear dose:response relationship can be observed that is linear within the lamellar bone formation window.^{52,67,125} Above the threshold for woven bone formation the linear relationship can change depending on whether damage occurs to the cortical bone.²² Microdamage to human cortical bone occurs at 2,500 μe in tension and 4,000 μe in compression,¹²⁶ although values for rodent bone may be different. The bone formation will only be observed if the peak strain magnitude attained during loading exceeds that of normal habitual cage activity (a threshold limit sometimes referred to as the minimum effective strain [MES] Fig. 7), or if the direction of loading is sufficiently novel to stimulate bone formation.^{89,98} Below this level, bone mass will not change leading to a phenomenon known as the lazy zone, dead zone, or adapted window.^{22,30}

Similar to the practical limitations of achieving an idealized waveform, target load magnitudes can sometimes be difficult to reach depending upon the tuning of the loading system, due to a combination of the high rates of loading used and the viscoelastic nature of the mouse hindlimb. In these situations, some level of load "over-command" or "under-command" may be necessary to achieve the exact desired load (± 0.2 N). Iterative adjustment to the command load signal to match the intended and actual peak load magnitudes is more easily done in multi-day load protocols in which the command load magnitude can be adjusted from day-to-day. For single bout protocols, we suggest tuning the

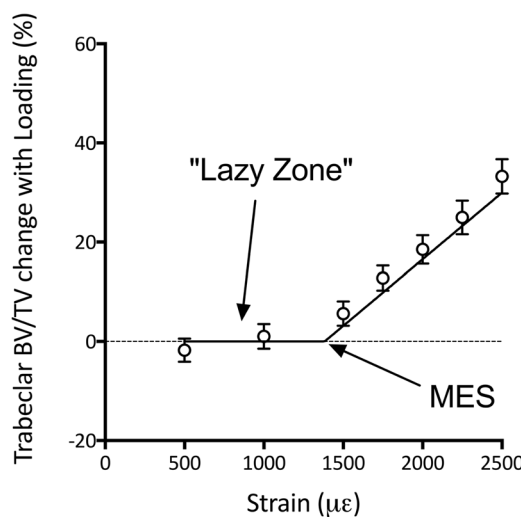


Figure 7. In animals which ambulate normally between loading bouts a minimum effective strain (MES) is apparent where bone formation is only observed above this threshold, referred to as the lazy or dead³⁰ zone; adapted from Meakin et al.⁸⁹ This phenomenon was first reported experimentally for cortical bone using the tibial four-point bend model⁵² and the ulnar axial compression model.¹²⁵

loading device and establishing a load protocol in fresh cadaveric material prior to loading in live mice.

Strain Rate

Consistent with extracellular fluid flow around osteocytes being proposed as a possible mechano-stimulus in bone,¹²⁷ the rate at which strain (or load) is applied has also been positively correlated to the bone formation response.^{44,59,128} Using sinusoid waveforms, strain rate is impossible to alter without also altering load frequency, strain magnitude, or the dynamic strain range experienced by the bone. Changes in strain rate can be isolated from changes in strain magnitude and load frequency using trapezoidal or triangle waveforms, but only by altering the length of time spent at peak load or in the "rest" phase.⁵⁹ Because some studies have shown length of the rest phase to enhance load-induced bone formation, changing the duration of these static periods in the waveform could impact the response of the bone to load as well.^{55,129} Thus, determining the relative contribution of each of these variables independently (strain rate, strain magnitude, and load frequency) is challenging.

Frequency (Number of Loading Cycles per Second)

Studies using the turkey ulna loading model,⁴⁶ rat tibia four-point bending model,^{123,128} and rat ulna axial loading model^{130,131} have all shown that increasing loading frequency causes enhanced bone formation (Tables 1B, C, and E). Cortical bone adaptation to mechanical loading increased with loading frequency up to 5–10 Hz and then plateaued with frequencies from 10 to 30 Hz.¹³⁰ Increasing loading frequency may lead to an enhanced rate of fluid flow in the bone's lacunar-canalicular system, with frequencies greater than 10 Hz leading to less efficient fluid flow and transduction processes. Load-response curves for *in vivo* osteocyte Ca^{2+} signaling were also shown to be dependent upon loading frequency (0.5, 1, and 2 Hz), using a novel mouse metatarsal three-point bend model.¹³²

Number of Loading Cycles per Session

The number of cycles required to maintain bone mass or generate a bone formation response is surprisingly few, ranging from 5 to 60 depending on the study.^{47,66,68,133} Increasing the cycle number beyond this does not dramatically increase the osteogenic response, suggesting that saturation of the stimulus occurs after relatively few cycles. Including excessive numbers of loading cycles in the loading regimen may unnecessarily increase anesthesia time and could increase the likelihood of inducing fatigue damage in the bone. Articular cartilage in the knee is also prone to damage, which could cause lameness and affect ambulation between loading bouts that would alter habitual loading and potentially bone mass.^{134–136} Furthermore, one study demonstrated that the bone formation response in trabecular bone actually

decreased slightly when the cycle number was increased from 216 to 1,200.⁶⁸ Therefore, minimizing loading cycle number is recommended whenever possible.

Number of Loading Sessions per Week

A recent study demonstrated that the total number of load sessions (or bouts) over which loading is applied to the limb also affects the adaptive response.⁶⁶ Loading for 5 days per week induced a 38% increase in cortical bone volume compared with 15% when loading was performed 3 days per week. Furthermore, reducing the length of loading from 2 weeks to 1 week did not reduce the bone formation response in the cortical bone at the 37% site. Therefore, it would seem sensible to limit loading studies to a single week with loading on 5 consecutive days. This study measured only cortical bone response to load by dynamic histomorphometry and did not assess cancellous bone response to load, which may be differentially sensitive in response to various loading variables relative to cortical bone.⁶⁸

Rest Insertion

Partitioning the number of loading cycles into short bouts with rest periods in between or inserting a period of rest between every cycle can increase the bone-formative response to loading in different loading models.^{53,137} More recently, the effectiveness of short rest insertions for enhancing bone formation was challenged in a study using tibial axial compression, which did not find any benefit of inserting rest between every four cycles.⁶⁸ A similar result was noted in an earlier rat ulnar loading study, albeit with more than one loading parameter varied between the two protocols compared.⁴⁰

Pre-Load or Inter-Cycle “Resting” Load Magnitude

Typically, a pre-load is applied through the loading system software prior to the start of any loading protocol, to hold the hindlimb firmly in place. This same load magnitude is often used as the “rest” (or “dwell”) load in any rest-inserted protocol. When reported, this load has varied from -0.2 to -12.4 N across different axial compression tibial loading studies.^{42,63,66,86,100} The effect of altering this initial (or resting) load level itself, or in combination with other loading parameters has not been systematically examined in the axial tibial loading model. A recent study using a novel off-axis compression

tibial loading model in 4-month-old female BALB/c mice showed that even when keeping the change from pre-load to peak load similar (-3.8 N), increasing the static pre-load from -0.03 to -1.5 N had detrimental effects on bone formation, even though the greatest pre-load (-1.5 N) also corresponded to the greatest applied peak load (-5.3 N).¹³⁸ These results may depend upon the genetic strain, age of mouse, and loading protocol employed, but suggest that choice of pre-load magnitude can have important effects upon the load-induced anabolic response of both cortical and cancellous bone tissue. To this end, we recommend that the pre-load magnitude be minimized to below -1.0 N where possible.

Minimum Loading Protocol Parameter Reporting

As part of this review, we have suggested guidelines regarding the minimum parameters that should be reported in a bone mechanical loading study (Table 2).

In addition to reporting these parameters, ensuring that the desired parameters (e.g., peak load, loading rate, and frequency) are achieved and the actual values reported is important.

OUTCOME MEASURES TO CONSIDER FOR TIBIAL LOADING STUDIES AND WHEN THEY ARE MOST EFFECTIVE

Microcomputed Tomography (microCT)

MicroCT is the standard method to assess rodent volumetric bone mineral density and microstructure; reporting guidelines and nomenclature have been previously outlined.¹³⁹ Although microCT is often performed *ex vivo*, after euthanasia, an increasing number of studies are implementing *in vivo* microCT imaging.

Longitudinal microCT imaging allows one to monitor changes in bone density and microstructure over time but comes with special considerations, such as radiation exposure. Since frequent or excessive exposure to X-rays can lead to bone loss or skeletal abnormalities,¹⁴⁰ a trade-off must be made between radiation dose and image quality if microCT will be used for *in vivo* assessment. Decreased trabecular bone volume fraction was reported in irradiated limbs (4 scans over 5 weeks, 0.5 Gy per scan) compared with contralateral limbs (one scan) in 8–10 week old C3H/He (-8%), C57BL/6 (-13%), and BALB/cBy (-20%) female mice.¹⁴¹ Willie et al.⁴² also showed that radiation (approximate dose of 0.48 Gy per scan) caused decreased trabecular bone volume fraction and increased trabecular

Table 2. Important Loading Protocol Variables to Report for Tibial Loading Studies

Number of loading cycles/bout	Number of loading bouts/week	Number of weeks loading
Peak load magnitude	Peak strain engendered and location in the bone	Frequency (e.g., # of load cycles/second)
Peak loading rate	Peak strain rate	Pre-load magnitude
Waveform used	Rest insertion (duration and load magnitude)	Animal/bone orientation: supine-horizontal, supine-vertical, prone-vertical

separation in young (10-week-old), but not adult (26-week-old) C57BL/6 mice after comparing morphometric parameters of animals being radiated four times with an interval of 5 days to mice being radiated only once. Interestingly, a greater increase in trabecular bone volume fraction due to loading was measured in mice scanned four times over 2 weeks compared with mice scanned only once (+140% vs. +88% load-induced increase, respectively).⁴² These data and subsequent studies^{42,142,143} suggest that exposure to ionizing radiation from microCT imaging at these doses does not hamper the response of cancellous and cortical bone to mechanical loading.

Studies performing repeated *in vivo* microCT analyses should either measure the effect of radiation on bone formation in their individual study or use established protocols,¹⁴⁴ with special attention when using skeletally immature mice. When choosing a volume of interest and voxel size, one must consider if the resulting scanning time and radiation exposure is acceptable. It is recommended that when using new protocols, an age-matched non-scanned or minimally scanned control group be included to determine the influence of radiation on bone (re)modeling.

Furthermore, because the contralateral hindlimb is commonly used as a non-loaded control, mice should be examined at baseline “time zero” prior to loading and imaging, by performing an additional *in vivo* microCT scan at baseline or including an additional group to examine with *ex vivo* microCT at baseline. These baseline data allow one to control for handedness, that is, the assumption that the bone density, morphology, and microstructure of both left and right limbs are not significantly different at the onset of the study. This same age-matched non-loaded group (non-loaded left and right limb) that is scanned at baseline to determine handedness could also be scanned at end-point to compare with the non-loaded control limb of the loaded group, thus lending confidence to the use of your contralateral limb as a control that is unaffected by loading of the opposite limb. In some studies, the contralateral limb by itself is not an adequate control, such as when a tumor or systemic drug is injected into the mice, and thus a separate non-loaded group is essential.¹⁴⁵

Histomorphometry

While microCT is considered the gold standard to assess static bone microarchitecture, histomorphometry is still considered the gold standard to evaluate bone (re)modeling. Fluorochrome labeling has been used since the 1950s,¹⁴⁶ when it was observed that tetracycline and other fluorescent dyes, such as alizarin and calcein green bind to newly forming hydroxyapatite, and thereby provide labels, which can be used to quantify dynamic active bone formation over a known time period. Histomorphometry guidelines and nomenclature have been reported,¹⁴⁷ and continue to be updated.¹⁴⁸

Fluorochrome-derived histomorphometry must be performed on undecalcified sections, which are usually

embedded in polymethyl methacrylate. The embedded mice tibiae should be sectioned in the frontal plane of the proximal metaphysis to assess trabecular bone morphology and in the transverse plane of the diaphysis to assess cortical bone morphology. The proximal metaphysis should not be sectioned in the transverse plane, since it is difficult to assess how far the region of interest is from the growth plate compared with having the entire proximal bone section within the analyzed field. In contrast to humans, mice do not normally undergo Haversian (intracortical) remodeling, but rather experience surface remodeling. Thus, the cortical diaphyseal region should be analyzed at both the endo- and periosteal surface. Intracortical remodeling has been observed in very old mice¹⁴⁹ or after fatigue loading which produces microdamage.¹⁵⁰ At least the following parameters should be reported: single- and double-labeled surface per bone surface (sLS/BS and dLS/BS), mineralizing surface (MS/BS), mineral apposition rate (MAR), and bone-formation rate (BFR/BS), calculated as $0.5 \times \text{sLS}/\text{BS} + \text{dLS}/\text{BS}$. An alternative metric that has been reported is inter-label area, which provides a simple measure of bone accrual in cases where a mix of lamellar and woven bone is induced.^{59,80} In cases where woven bone is noted, the incidence and location of woven bone should be described, and if applicable, values of woven bone surface (Wo.S/BS) and area (Wo.Ar/Ct.Ar) should be reported.^{66,67,85} Recommendations on how one should express MS/BS, BFR/BS, and MAR in the absence of double and/or single labels in a sample are addressed elsewhere.^{151,152}

Alternative Methods for Analyzing Bone Remodeling

Although histomorphometry has greatly advanced our understanding of bone formation, limitations include many necessary assumptions and the two-dimensional nature of the analyses. Label escape can occur if bone formation is initiated after the first label is given or if formation stopped before the second label was administered, resulting in only one label. Also, differentiating between surface modeling and surface remodeling in rodents using histomorphometry is extremely challenging.^{148,153–155} Resorption cannot be assessed with dynamic histomorphometry, but rather is commonly assessed using tartrate-resistant acid phosphatase (TRAP) staining of osteoclasts. However, TRAP staining is a static measure that does not allow investigation of the temporal or spatial distribution of resorption. Serial block-face imaging has been reported as an alternative to assess both formation and resorption since it allows three-dimensional (3D) fluorochrome mapping¹⁵⁶ and quantification of the number and size of resorption cavities.¹⁵⁷ Although this approach for 3D mapping is often limited to measuring small sample volumes and only one-time point within a particular specimen, this method remains a valuable tool that combines the benefits of classical bone histomorphometry with the ability to examine volumes rather than areas.

Advances in microCT allow one to follow structural changes in the bone of living animals in 3D space over time.^{158,159} A series of 3D high-resolution images of the bone architecture of the living animal are taken with a time-lapse, the duration of which depends on how rapidly the bone is (re)modeling (e.g., up to 15 days or less in young growing mice tibiae^{87,160}). A later 3D microCT data set is superimposed (registered) onto an earlier data set of the same mouse in a common coordinate system by rotating and translating one data set with respect to the other using an optimization criterion. This method identifies formation (F), resorption (R), and quiescent (Q) sites over given time intervals. Bone volumes only present in the earlier data set are considered resorbed, while volumes only present in the latter data correspond to formed bone volumes. Formed and resorbed bone volumes (MV/BV and EV/BV) surface areas (MS/BS and ES/BS), and thicknesses (MTh and ED) are reported. Time-lapse *in vivo* microCT-based morphometry was used in mice to assess cortical⁸⁷ and trabecular¹⁶¹ bone formation and resorption after mechanical loading. By combining time-lapse *in vivo* morphometry with micro-FE analysis, the local mechanoregulation of bone formation and resorption was reported for the mouse caudal vertebra^{162,163} and tibia.^{164,165} This approach was extended to spatially and temporally track formation, resorption, and quiescent sites over time to distinguish between surface modeling (spatially not correlated formation and resorption) and surface remodeling (resorption followed by new formation at the same site) after tibial loading in adult female mice.¹⁶⁶ These microCT-based time-lapse methods hold a great deal of promise but are limited by voxel size, which is typically on the order of the thickness of newly formed (or resorbed) bone. Additionally, image registration may be difficult in cases of rapid longitudinal bone growth and/or bone formation and resorption, such that the scanned VOI at the two-time intervals does not include sufficient mutual information (similar architecture) to align and rigidly register the data.¹⁶⁷

Whole Bone and Material Characterization

Mechanical loading can alter the bone mass, geometry, and microstructure and thus measuring these changes at the whole bone and tissue level are often performed. Jepsen et al.¹⁶⁸ introduced guidelines to evaluate phenotypic changes in mouse long bones using biomechanical testing and recommended minimum reportable information for experimental testing conditions and outcome variables. Other whole-bone morphometric properties to consider reporting are the tibial length and anterior-posterior and medial-lateral radii of curvature (C_{AP} and C_{ML}, mm),¹¹⁶ which can be measured from microCT scans.

In addition to altering bone mass and microstructure, some studies have indicated that mechanical loading can also alter bone mineral and matrix properties.^{169–172} The material bone mineral density or

so-called tissue mineral density (TMD) represents a volume of bone matrix that does not include marrow spaces, osteonal canals, lacunae or canaliculi, thus reflecting the degree of mineralization of organic bone matrix. Although microCT can be used to assess tissue mineralization, lower resolutions are often used that include contributions from porosity. Also, the polychromatic beam used in lab-based microCT can suffer from beam hardening, which limits the accuracy of tissue mineralization measurements. Thus, other methods are often used to assess degree and heterogeneity of tissue mineralization including: quantitative backscattered electron imaging (qBEI), Fourier transform infrared spectroscopy (FTIR), and Raman spectroscopy. Mineral particle thickness, collagen crosslinks and microdamage, for example, can be measured using some of the aforementioned techniques.

Gene and Protein Expression Analysis

The cellular and molecular mechanisms driving bone formation caused by mouse tibial loading have been examined using immunohistochemistry, quantitative real-time polymerase chain reaction, microarray, and RNA-sequencing.^{91,94,122,124,173–176} Load-induced alterations in gene expression can be measured within one hour following loading.⁹⁴ Therefore, studies that focus on gene-level mechano-transduction responses will typically perform a single bout of loading and then sacrifice the mice at various time points, up to 24 or 72 h post-loading.^{122,173} A single load bout is usually sufficient to induce a detectable immediate or near-term response to load, albeit the number of differentially expressed genes at 1–4 h after loading may be few.¹²² To assay genes related to downstream bone formation and resorption effects following loading, gene expression should be measured within the first week following the initiation of loading, after either a single or multiple loading bouts.^{173,175} Studies examining protein expression generally mimic either the single loading protocol used for gene expression analyses or the week(s) long loading protocol to assess bone mass gains.¹¹¹ Processing of whole bones, diaphyses, or separation of the cancellous bone volume from the cortical bone volume have been reported.¹⁷⁶ If the focus of the study is bone cell response to loading, we recommend removal of bone marrow (by flushing or centrifugation) so that the transcriptional response reflects bone cell activity and is not diluted by the large number of marrow cells.¹⁷⁶ If separately analyzing the cortical and cancellous bone is desirable, mice characterized by low bone mass phenotypes may require pooling of cancellous bone samples from multiple mice to obtain an adequate amount of RNA. When initially establishing a loading protocol, one should first determine if the protocol is anabolic in cortical and/or cancellous bone of the particular mouse strain being studied using standard methods including microCT and conventional histomorphometry before assessing outcome parameters such as gene or protein expression.

ANIMAL WELFARE AND HUSBANDRY CONSIDERATIONS IN TIBIAL LOADING STUDIES

Local and national animal welfare committee guidelines should be followed for animal housing, anesthesia, analgesia, and all other animal procedures. If choosing to individually house mice during the loading study, separation of mice no later than 5 days prior to loading is advised to reduce their time alone. Extended periods of pre-study isolation should be avoided as group housing is considered less stressful by most institutional animal welfare committees. The potential effects of cage enrichment (e.g., play tunnels and cardboard houses) on load-induced bone (re)modeling have not been explicitly examined. However, treadmill exercise prior to initiation of axial tibial loading altered some aspects of the bone adaptive response,¹⁷⁷ so enrichment options that offer opportunities for highly repetitive limb loading (e.g., running wheels) should be avoided. Isoflurane anesthesia is most commonly used during loading and additional procedures such as *in vivo* microCT imaging due to ease in controlling the depth of consciousness and lower mortality risk, in our experience, compared with injected ketamine-based cocktails.

Analgesics are not used in most loading protocols, but may be required by institutional animal care committees. While typically not visible by eye, evidence suggests that axial tibial loading can cause minor gait asymmetries following loading. Interestingly, changes in gait kinematics (stance time and stride length) do not occur in the loaded limbs, but in the non-loaded contralateral limb, perhaps as compensation for trauma done to the loaded limb.¹³⁶ Development of protocols for gait outcome measures in loading studies would be a valuable advancement for monitoring behavior and cage activity following loading. While the analgesic Carprofen was effective in returning contralateral limb gait parameters to normal following a single loading bout, analgesics did not correct the altered gait in mice loaded three times a week for 2 weeks. We recommend avoiding the use of Carprofen and other COX-2 inhibitors that have been implicated in delayed bone healing in animals,¹⁷⁸ blocking the activation of β -catenin signaling, and reducing sclerostin expression post-loading.¹⁷⁹ If an analgesic is required by your institutional animal care committee or a more invasive procedure is coupled with tibial loading (e.g., bone healing), we recommend using buprenorphine or tramadol.¹⁸⁰ Analgesic dosage information should always be provided by the authors.

Care should be taken to avoid using load levels or fixture designs that cause pain or tissue damage such that the animal alters its normal gait and/or weight-bearing status because the anabolic response to loading is a combination of the applied loading protocol and normal “background” loading. Application of high load magnitudes are a particular concern when using mice with high bone mass phenotypes that require higher load levels to engender comparable strains to those in wild-type mice. Pflanz et al.⁹³ reported excessive

limping and ankle swelling in 10-week-old *Sost* KO mice after 3 days of loading mice at -17 N to engender $1,200\text{ }\mu\text{e}$ on the medial midshaft. In this case, the investigators terminated the experiment at 3 days, euthanized the mice and proceeded with a new experiment in which the *Sost* mice were loaded at -12.9 N engendering $900\text{ }\mu\text{e}$ on the tibial mid-shaft. Similarly, the loads for the control mice were reduced from -11 to -7 N to match the $900\text{ }\mu\text{e}$ induced in the *Sost* KO mice. At these loads, the mice did not exhibit any ankle swelling or limping during the experiment.

Each study should describe the methods used for induction and maintenance of anesthesia during loading, the use of any post-loading analgesics, and provide assurance of approval for the study by the relevant institutional animal care and use review board. All excluded animals should be reported as well as the reason for their removal from the study, which can include, for example, anesthesia overdose, euthanized to relieve distress, euthanized after sustaining a tibial fracture during loading. A poor response by the animals to the mechanical loading protocol will often be reflected in their body mass. Body mass should be monitored daily for loading studies of 3 days or longer and animals given nutritional supplement if their mass drops by 10–15%. Animals should be excluded from the study (or euthanized) if body mass decreases by 25% of the initial body mass.

WHAT WE HAVE LEARNED ABOUT MECHANORESPONSE FROM TIBIAL AXIAL LOADING STUDIES?

Several studies have used this model to examine the effects of age, sex, mouse strain, and background activity levels on the cortical and cancellous bone response to loading. The bone formation response to loading is reduced with age in female C57Bl/6 mice tibiae.^{42,85,86,89,97,160,181} Less is known about age-related changes in adaptation to loading in males or other mouse strains.⁷⁹ The Silva group⁸⁸ reported that old (22 months) male BALB/c mice had a significantly greater response to loading at the endocortical surface than mature (7 months) mice, while responses at the periosteal surface did not differ between age groups. They also examined female BALB/c mice¹⁷⁵ and observed at cortical sites that the young-adult skeleton (4 months) is more responsive to loading in terms of bone formation than the mature to the middle aged skeleton (7–12 months). Interestingly, their loading protocol was not anabolic in these studies and actually diminished trabecular BV/TV in adult mice (4–12 months old).^{88,175} In a follow-up study, they observed a mild anabolic response in trabecular bone in 4-month-old mice, although when they compared their loading protocol to one used by the Cornell/HSS group, regardless of mouse strain (C57BL/6 or BALB/c), the Cornell/HSS protocol induced a greater accrual of trabecular bone and an earlier peak in cortical bone volume accrual.⁴¹ In terms of bone volume, mouse strain did not affect the

overall tibial response to loading, but rather the manner of relative changes in trabecular thickness and cortical accrual, which were greater in C57BL/6 than BALB/c mice.⁴¹ However, the estimated peak bone strains differed by approximately 20% between C57BL/6 and BALB/c mice, so results for the bone tissue response to load between these two strains may not be directly comparable in this study. Additional studies are needed to further investigate mouse strain-related differences in mechano-adaptation using the axial compression tibial loading model.

Studies have used this tibial loading model to investigate the role of specific proteins on bone mechano-transduction by subjecting transgenic and conditional knock-in/knock-out mice to tibial loading. An experimental design consideration is whether to use littermate or wild-type controls when examining transgenic mice. To avoid the effects of genetic drift within a mouse strain, littermate control mice are the gold-standard. Wild-type mice may display more variability in terms of basal expression of certain proteins, due to differing genetic backgrounds compared with littermates who share the same parents as the transgenic strain of interest.

Studies of tibial loading have been performed using both male and female mice, although far fewer studies have examined how male mice respond to loading. The effect of mechanical load on the cancellous response to load did not show any differences by sex, when strain-matched loading was performed in 10-week-old male and female C57BL/6 mice.⁸³ Other studies observed an elevated mechanoresponse in the cortical bone of female compared with male wild-type mice.^{89,91,98,117} In keeping with NIH guidelines, the inclusion of sex as a biological variable is encouraged. However, certain differences between males and females must be considered. First, studies working with adult male mice must consider the possible impact of group housing on study results (see below). Second, important sex-related differences exist in cortical and cancellous bone mass between male and female mice.

Strain gauge calibration allows one to equalize cortical bone strain stimuli at a specific cortical location. Relative cancellous bone volume can vary by sex and genotype^{77,83,89} and may induce different cancellous mechanical stimuli between experimental groups. If the primary goal is to compare the cancellous tissue response to load between different sexes or genotypes, the strains induced in cancellous tissues during loading should be modeled for all study groups. If two load values alone are not sufficient to equalize strain stimuli in the cancellous and cortical tissues simultaneously between two groups, one can prioritize the experimental design for one tissue type or conduct two independent studies in which strain stimuli are equalized in cortical and cancellous tissues separately.

Another variable to consider is the habitual activity of the mice between loading bouts. Reducing hindlimb loading through unilateral sciatic neurectomy can increase the unloaded tibia's response to loading, even in

aged mice.¹⁸² Conversely, increasing activity through treadmill exercise prior to loading reduced the trabecular response to mechanical loading.¹⁷⁷ Finally, the response to loading in adult male mice was reduced when mice were group-housed. The reduced adaptive response was believed to be caused by increased strains engendered in the tibiae during group-housed fighting activities that increased the MES and masked the response to loading.⁹⁸ Adult male mice should be individually housed during loading studies. If group housing is used, care should be taken to observe if fighting is occurring. The influence of group housing in interpreting the results of skeletal loading studies is less for younger mice or females, who are less likely to fight. If male mice are over 7 weeks of age and group housed, placing them into separate cages 5–7 days prior to the loading experiment is sufficient to permit a significant load-induced bone formation response (R. Main, unpublished data).

RECOMMENDATIONS FOR TIBIAL LOADING PROTOCOLS (CONSENSUS FROM OUR GROUP)

About 100 published studies have used the axial compression tibial loading model to study various aspects of bone formation and bone mechanobiology. Many of these studies employ different protocol parameters to accomplish their goals. Thus, no single loading protocol has been used consistently in axial compression tibial loading. Assuming that one protocol would be "right" for all studies is unrealistic. Protocol parameters should be chosen according to the study goals. Furthermore, a lot of unexplored parameter space remains and our understanding is limited regarding interactions of the various protocol parameters with one another and animal intrinsic factors (e.g., animal age, sex, genetic strain, and hormone status) to form either lamellar or woven bone. As mentioned earlier, one problem with all of the unique loading protocols used is that comparison of results across studies is difficult. Thus, despite the near limitless options for combinations of protocol parameters and the number of different protocols that have successfully produced load-induced bone formation, we provide here four protocols that have been used successfully on numerous occasions in an effort to facilitate inter-study comparison, or provide a starting point for new investigators. These protocols reliably result in bone formation and/or elicit changes in gene expression related to bone formation. Please note that applied load (or strain) levels may need to be altered from those presented here depending on animal intrinsic factors.

I.^{83,97}

Target longitudinal bone strain and strain rate: +1,200 $\mu\epsilon$ on medial mid-shaft (50%) site (+16,000 $\mu\epsilon/s$)
Waveform: triangle

Ramp/unload timeframe: 0.15 s (0.075 s ramp, 0.075 s unload)

Load application frequency: 4 Hz (0.1 s "rest" phase between consecutive ramp/unload cycles)

Inter-cycle "rest" phase load: -1.5 N

Long-term rest insert: None

Number of load cycles per day: 1,200

Number of loading days: 10 days over 2 weeks (e.g., Monday–Friday loading for 2 weeks)

Euthanasia: Mice euthanized on the third day following the last day of load application

Notes: Protocol induces cancellous and cortical anabolic responses detected by endpoint microCT in female and male C57Bl/6 mice. Short-term load-induced changes in gene expression by RNA-Seq and qPCR have been found in mice euthanized within 3–24 h following a single loading bout.¹²²

II.^{66,105}

Target longitudinal bone strain and strain rate: +1,000 $\mu\epsilon$ on medial 37% site (+13,333 $\mu\epsilon/\text{s}$); -1,800 $\mu\epsilon$ on posterior–lateral cortex 37% site (-24,000 $\mu\epsilon/\text{s}$).

Waveform: Haversine

Load application frequency: 4 Hz

Pre-load: -0.5 N

Rest Insert: None

Number of load cycles per day: 60

Number of loading days: 5 days consecutive over 1 week

Euthanasia: Mice euthanized 1 week following last load application

Notes: Protocol induces increased bone formation indices by dynamic histomorphometry, and changes in gene expression by qPCR and RNA-Seq in female C57Bl/6 mice; the number of loading days may be increased to 10 over a 2-week period to better induce detectable bone accrual by microCT.¹⁴

III.^{42,114}

Target longitudinal bone strain and strain rate: +1,200 $\mu\epsilon$ on medial mid-shaft (50%) site (+16,000 $\mu\epsilon/\text{s}$); +1,750 $\mu\epsilon$ on anterior–medial cortex at the 37% site (26-week-old female C57Bl/6 mice).

Waveform: triangle

Ramp/unload timeframe: 0.15 s (0.075 s ramp, 0.075 s unload)

Load application frequency: 4 Hz (0.1 s “rest” phase between consecutive ramp/unload cycles)

Inter-cycle rest phase load: -1 N

Rest insert: 5 s at -1 N after every fourth cycle

Number of load cycles per day: 216

Number of loading days: 10 days over 2 weeks (e.g., Monday–Friday loading for 2 weeks)

Euthanasia: Mice euthanized 3 days after the last day of load application

Notes: Protocol induces cancellous and cortical lamellar bone formation detected by endpoint microCT, time-lapse morphometry and histomorphometry in female 10, 26, and 78-week-old C57Bl/6 mice.^{42,87,160} Short-term load-induced changes in gene expression by microarray and qPCR have been found in female 78-week-old C57BL/6 mice euthanized within 1–24 h following a single loading bout (B. Willie, unpublished data).

IV.^{67,89,98,108}

Target longitudinal bone strain and strain rate: +1,800 $\mu\epsilon$ on medial 37% site (+31,000 $\mu\epsilon/\text{s}$);

-4,000 $\mu\epsilon$ on posterior–lateral cortex at the 37% site (-160,000 $\mu\epsilon/\text{s}$).

Waveform: trapezoid

Ramp/unload timeframe: 0.10 s (0.025 s ramp, 0.05 s hold at peak load, 0.025 s unload)

Load application frequency: Not applicable, a single load cycle then long rest phase

Inter-cycle “rest” phase load: -0.5 N

Rest insert: 10 s after every cycle

Number of load cycles per day: 40

Number of loading days: 3 days per week (e.g., Monday, Wednesday, and Friday) or alternate days for 2 weeks

Euthanasia: Mice euthanized 2 days following the last load application

Notes: Woven bone can be observed on the periosteal surface of the posterior–lateral cortex of 16–19-week-old female C57Bl/6 mice.^{67,89,111} Woven or lamellar bone is formed on this surface in male mice of similar age.^{89,98} Lamellar bone is formed on the medial periosteal surface. Older mice (19 months) of both sexes show primarily lamellar periosteal bone formation.⁸⁹ Cancellous BV/Tv increases with load. Load-related changes in sclerostin and other bone anabolic markers have been described using this protocol or subtle variants of it after one or two loading bouts^{89,111} and gene expression changes as early as one hour after a single loading bout.⁹⁴

CONCLUSIONS

The guidelines and recommendations provided in this review are intended to help researchers successfully perform *in vivo* loading experiments in mice, and thus contribute to the extension of our knowledge of the mechanisms of bone mechanotransduction. Furthermore, the reporting of mechanical loading studies needs to become more complete and transparent, using the defined parameters that we have outlined in this review. Adopting a single loading protocol for all studies using the tibial axial compression model is unrealistic. However, limiting protocol variation to a small set of defined protocols upon which minor changes can be made as dictated by study requirements would enhance the uniformity of basic results and increase our ability to compare study results between different research groups.

Prior studies with the tibial axial compression model and other skeletal loading models have laid a strong foundation for our understanding of bone mechanotransduction. However, much remains to be learned about the physical and chemical processes of mechanotransduction at the level of the single cell (osteocyte, osteoblast, etc.). Adopting a broader view of bone as a heterogeneous organ, instead of focusing on discrete tissue envelopes (cancellous bone, cortical bone, periosteum, marrow tissues), will determine how cross-talk between these tissues regulate bone (re)modeling and the unique contributions of different tissue microenvironments to this process. Studies examining interactions between skeletal loading and factors such as age, sex, obesity, reproductive

status, stage in estrus cycle, circulating protein and hormone levels, and genetic modification are expected to continue to reveal important insights regarding disease- or age-related changes in skeletal mechanotransduction and contribute to our understanding of age-related skeletal wasting diseases. Novel applications of this model, such as studies examining the role of mechanical loading on cancer tumor growth and metastasis in bone,¹⁴⁴ the role of repetitive loading in the development of knee osteoarthritis,^{134,135} or solute transport in the lacunar-canalicular network¹⁸³ are expected to continue to develop.

While *in vitro* and computational studies are critical to continued advancements in bone mechanobiology, presently such approaches are not able to capture the complexity of the three-dimensional environment in bone, the interaction of different cell types, or the complexities of relevant genetic pathways to answer all questions. Thus, for the foreseeable future, much of our understanding of bone mechanobiology at the cell and tissue levels will continue to be based on *in vivo* loading models, including the mouse axial compression tibial loading model.

AUTHORS' CONTRIBUTION

All authors played a significant role in drafting and editing the figures and text of this manuscript in its entirety. All authors have read and approved of the final submitted manuscript.

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SUPPORTING INFORMATION

Additional supporting information may be found in the online version of this article.