Minimally invasive laser Doppler flowmetry is suitable for serial bone perfusion measurements in mice

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ABSTRACT

In vivo laser Doppler flowmetry (LDF) has previously been used to quantify blood perfusion accurately at a single timepoint in the murine tibial metaphysis. However, this procedure entailed substantial disruption to soft tissues overlying the bone and caused notable localized inflammation for several weeks after the procedure, impeding serial measurements in the same mouse. In this study, we tested a less invasive technique to measure perfusion in the tibia with LDF and determined that it can be used serially in the same mouse without causing signs of inflammation or gait perturbations. Twenty 14-week-old C57Bl/6J mice were evenly divided into groups that either had daily treadmill exercise or remained sedentary. Within these activity groups, mice were evenly subdivided into groups that received LDF measurements either weekly or only once at the study endpoint. Bone perfusion was measured with LDF in the anteromedial region of the right tibial metaphysis. Serum concentrations of interleukin 6, incision site wound area, and interlimb coordination during gait were measured weekly for four weeks. Tibial perfusion did not differ significantly between exercise and sedentary groupswithin the weekly or endpoint-only LDF groups at any timepoint. Perfusion was significantly increased in the third week in the weekly LDF group relative to measurements in the second and fourth weeks. Ligation of the femoral artery caused consistent, rapid reductions in tibial perfusion, validating that LDF is sensitive to changes in tibial blood supply. Weekly LDF procedures did not adversely affect gait, as interlimb coordination during treadmill locomotion was similar between weekly and endpoint-only LDF groups at every timepoint. Images of the incision site show wound closure within one week, and serum concentrations of interleukin 6 were not significantly different between weekly and endpoint-only groups. Together, these findings demonstrate that our minimally invasive LDF technique is suitable for serial in vivo measurements of intraosseous blood perfusion without inducing localized inflammation or negatively affecting gait patterns in mice.

1. Introduction

Vascularity within bone (osteovasculature) is an essential contributor to bone health, providing nutrients, oxygen, cells, and chemical signals and removing waste products (Parfitt, 2000; Truea, 1963). Adequate vascular perfusion is required for bone development, adaptation in response to loading, and healing after fracture (Truea, 1963; Tomlinson et al., 2013; Wallace et al., 1991). Evidence that vascular pathologies are associated with bone loss is growing. Aortic calcification is associated with decreased lumbar spine bone mineral density (BMD) and increased fracture risk in men and women within four years (Naves et al., 2008), and the incidence of cardiovascular disease increases with reduced BMD in the spine in white men, and hip, trochanter, and femoral neck in black women (Farbat et al., 2007). Osteoporosis is associated with reduced perfusion in the vertebrae for men (Griffith et al., 2005) and in the femoral head for women (Griffith et al., 2008), although the mechanisms responsible for bone loss in these individuals is unknown and needs further examination using animal studies. Although murine models are commonly used to determine the effect of pathologies on bone properties, measuring blood perfusion within mouse bone is complicated due to their small size. Current methods are either experimentally difficult (e.g., hydrogen washout (Whiteside et al., 1977)) or require the animal to be sacrificed (e.g., microspheres, radiolabels, polyoxometalates, barium sulfate, or...
Microfil® (Serrat, 2009; Grundnes and Reikerås, 1992; Reeve et al., 1988; Kerckhofs et al., 2018; Barou et al., 2002; Yao et al., 2004; Boerckel et al., 2011)). Some methods can be performed in vivo but provide poor resolution not suitable for small bones: laser speckle imaging (Briers et al., 2013), laser Doppler perfusion imaging (Roche et al., 2012), contrast-enhanced magnetic resonance imaging (MRI) (Dyke and Aaron, 2009), contrast-enhanced positron emission tomography (PET) (Dyke and Aaron, 2009), and contrast-enhanced micro-computed tomography (Au et al., 2013; Clark et al., 2013). Endpoint and ex vivo measurements only provide a snapshot of vascular network function, missing the timing of vascular changes and any transient changes to vascular supply. A technique that could be used for longitudinal studies of bone perfusion in vivo would enable us to capture temporal changes in bone perfusion for individual subjects, thereby improving understanding of disease progression and intervention effectiveness.

First proposed as a tool to measure intrasosseous perfusion by Nilsson et al. (1980), laser Doppler flowmetry (LDF) directs a monochromatic light source over a perfused tissue and measures backscattered light from fluid movement with a photodetector to provide a relative measure of blood perfusion. Perfusion is a functional measure of blood flow that is affected not only by the amount and velocity of red blood cells but also capillary density, vascular permeability, and flow direction (Swiontkowski, 1991; Griffith, 2014). LDF was first used to measure blood perfusion in the cancellous bone of pig mandibles by Hellem et al. (1983) and thereafter was rapidly adopted in orthopaedic clinics as an intraoperative tool to aid surgeons in identifying non-viable bone for debridement in patients with osteomyelitis, osteonecrosis of the femoral head, and lower limb traumatic injury (Swiontkowski, 1991). LDF has also been used as an endpoint measure to compare relative intrasosseous perfusion between groups in murine research studies (Okunieff et al., 1998; Melyn et al., 2008). Recently, it was used to quantify perfusion accurately in vivo in the mouse tibia, but the technique used in that study involved a relatively large incision that resulted in inflammation at the incision site up to three months after the procedure (Roche et al., 2013). To monitor longitudinal changes in murine bone perfusion, a less invasive LDF procedure is needed that will not induce significant localized inflammation or limping during gait, which could have both biological and mechanical confounding effects on bone.

We developed a minimally invasive LDF procedure and have used it to measure changes in tibial perfusion in mice in response to diet-induced obesity, ischemic stroke, and treadmill exercise (Hanne et al., 2017a; Hanne et al., 2019b). The objective of this study was to determine if this modified LDF procedure could be performed repeatedly in a longitudinal study without affecting bone perfusion, inducing inflammation, or altering limb coordination during locomotion, which could confound bone metrics of interest and affecting perfusion changes associated with interventions. Developing new in vivo techniques to measure bone blood perfusion is a critical step needed to understand longitudinal changes to osteovasculature, which may contribute to bone loss occurring during progression of various clinical pathologies.

2. Materials and methods

2.1. Study design

The protocol for this study was approved by the Institutional Animal Care and Use Committee at North Carolina State University. Eighteen 14-week-old male C57BL/6J mice (The Jackson Laboratory, Bar Harbor, ME) were acclimated to the animal facility for one week. They were group-housed (4 per cage) on a 12-hour light/12-hour dark diurnal cycle and provided chow and water ad libitum. The mice were randomly assigned to four groups (Fig. 1) based on LDF procedure frequency (weekly, endpoint) and exercise regimen (sedentary, exercise): weekly sedentary (n = 5), weekly exercise (n = 5), endpoint sedentary (n = 4), and endpoint exercise (n = 4). Exercise groups were acclimated to treadmill exercise (Exer-3/6, Columbus Instruments, Columbus, OH) by increasing exercise intensity for 2 days prior to the start of the study (Day 1: 5 m/min for 10 min, 9 m/min for 10 min, and 12 m/min for 10 min; Day 2: 5 m/min for 5 min, 9 m/min for 5 min, and 12 m/min for 20 min). During the study, exercise groups performed daily exercise for 4 weeks (30 min/day, 5 days/week, 12 m/min, 5° incline), while sedentary groups were placed on a stationary treadmill for the same amount of time to equalize handling among the groups.

Laser Doppler flowmetry was used to measure intrasosseous blood perfusion in the right tibial metaphysis, performed either weekly for 4 weeks (weekly groups) or at a single timepoint at the end of the study (endpoint groups). For the weekly LDF groups, starting in Week 2, images of the incision site were taken during the LDF procedure prior to making an incision to assess the wound from the previous week. Blood samples were collected from the submandibular vein of all mice under anesthesia (at the end of the LDF procedure for the weekly groups). Blood samples were centrifuged at 2000 × g for 10 min, and the isolated serum was stored at −80°C until analysis with an enzyme-linked immunosorbent assay (ELISA). At five days after each of the first three LDF procedures, gait patterns were assessed using high-speed video. Immediately following the last LDF procedure and serum collection, mice were euthanized using CO2 asphyxiation followed by cervical dislocation.

2.2. Tibial perfusion

All mice were fasted for 6–8 h before each LDF procedure. Anesthesia was induced and maintained with isoflurane (2%) in pure oxygen throughout the procedure (about 15 min). After anesthesia induction, the fur over the right knee was shaved, mice were placed supine on a heated pad, and the right leg was taped to the surgical platform. A 2-5-mm long incision was made over the anterior medial surface of the right proximal tibial metaphysis, the bone was exposed, and a small region of the periosteum was scraped away. LDF measurements were recorded using an LDF monitor with a 785-nm light source (MoorVMS-LDF, Moor Instruments Ltd., Axminster, UK) and a 3-kHz lowpass filter. A VP4 Needle Probe (0.8 mm outer diameter, 0.25 mm fiber separation) was placed directly on the exposed bone surface (Fig. 2) and held in place using a micromanipulator (MM3-ALL, World Precision Instruments, Sarasota, FL) to reduce signal noise from probe movement. Each weekly measurement was composed of the weighted mean of three 30-s readings, with repositioning of the probe between readings. Readings that contained noisy data with spikes or deviated substantially from a steady value (i.e., had a nonzero slope) were re-taken. The incisions were closed using VetBond™ tissue glue (3M, St. Paul, MN) and covered with triple antibiotic cream.

2.3. Femoral ligation validation

At the end of the study, just prior to euthanasia, additional LDF measurements were performed in a subset of mice (n = 12) during arterial ligation to confirm the association of LDF perfusion measurements with changes in blood supply to the bone. While still anesthetized during the final LDF procedure, the skin incision over the proximal tibia was extended to expose the entire inner thigh and the femoral artery. The LDF probe was again positioned over the tibial metaphysis, and a suture was tied around the femoral artery but not tightened. A 30-s baseline measurement was taken, the suture was tightened to ligate the artery, and another 30-s measurement was recorded. The reduction in tibial perfusion was calculated as the ratio of the ligated measurement to the baseline measurement, expressed as a percent.

2.4. Wound area

As mentioned above, for the weekly LDF groups, pictures were
taken of the incision wounds immediately prior to each LDF procedure to assess localized inflammation and healing at the wound site from the procedure performed in the preceding week. Wound area was calculated by tracing the edge of the wound in ImageJ (version 1.51k, National Institutes of Health, Bethesda, MD). The wound was considered closed if no moist granulation tissue was visible and the wound was covered with new epithelium (Goova et al., 2001).

2.5. Serum concentration of interleukin 6

Systemic inflammation was examined by quantifying serum concentrations of the proinflammatory marker interleukin 6 (IL-6) with an ELISA (IL-6 Mouse ELISA kit, KMC0061, Invitrogen, Carlsbad, CA). A subset of the serum samples (Table 1) was prepared according to the manufacturer’s instructions and measured using a plate reader (Synergy H1, BioTek Instruments, Inc., Winooski, VT). Due to limited serum, some samples were diluted 2–4 times to allow samples to be run in duplicate. Since no increase in IL-6 was detected in this subset, which included serum from all groups at each timepoint, the remaining serum samples were not analyzed.

2.6. Gait pattern analysis

The effect of the LDF procedures on interlimb coordination was examined weekly in all mice, five days after each procedure day, because limping or other gait asymmetries could alter the strain experienced by hindlimb bones and confound bone outcome measures (Frost, 2003). During a short treadmill session (12 m/min for 60 s), high-speed video was collected in the sagittal plane at 240 frames per second (HERO4, GoPro, Inc., San Mateo, CA). Gait was analyzed using Kinovea (version 0.8, Kinovea Open Source Project) to quantify duty cycle for both hindlimbs and phase dispersions for ipsilateral, diagonal, and contralateral limbs with relation to the LDF limb (right hindlimb) (Fig. 3) (Leblond et al., 2003; Kloos et al., 2005; Redondo-Castro et al., 2013). Duty cycle for a given limb is the ratio of the time that limb is on the ground (measured from paw strike to lift off) to the total time of an entire gait cycle (measured from paw strike to paw strike). Phase dispersion between two limbs is a measure of the time between paw strikes for those two limbs within a gait cycle. Five consecutive gait cycles were analyzed for each treadmill session. Duty cycle and phase dispersion were averaged over the five gait cycles at each timepoint.

2.7. Statistical analysis

All data analyses were performed using SAS (SAS University Edition v. 9.4, SAS Institute Inc., Cary, NC) with a significance level of 0.05. Models were chosen to answer five questions: 1) Does performing weekly LDF procedures affect bone perfusion? To answer this question,
LDF data from the final timepoint (Week 4) were compared across LDF frequency (weekly, endpoint) and exercise regimen (sedentary, exercise) using a two-way ANOVA with interaction. Tukey’s post-hoc tests were used to compare group means. 2) Does exercise affect bone perfusion? For this question, LDF data for the weekly group were compared across exercise regimen and timepoint (Weeks 1–4) to examine differences between exercise groups within each timepoint (e.g., sedentary vs. exercise at Week 1). A mixed effects general linear model (procedure MIXED) with interaction was used, with exercise group as a fixed factor and timepoint as a repeated factor. The covariance matrix was modeled using compound symmetry. Exercise effect differences were calculated based on least squares means (LSM) with Tukey-Kramer adjustments for multiple comparisons. For the endpoint-only group (Week 4 data), the effect of exercise was examined with one-way ANOVA. 3) Do weekly LDF procedures alter exercise effects on bone perfusion? Effect differences between timepoints (e.g., Week 1 vs. Week 2) were evaluated in the same mixed effects model using LSM with Tukey-Kramer adjustments for multiple comparisons. 4) Do LDF readings directly correspond to changes in blood supply, assessed by femoral artery ligation? LDF data before and after the femoral artery was ligated were compared using a paired t-test. 5) Do weekly LDF procedures alter gait and interlimb coordination? This question was addressed by comparing gait parameters between LDF groups within exercise groups and timepoints (e.g., weekly sedentary vs. endpoint sedentary at Week 1). Four gait parameters were compared across LDF groups, exercise groups, and timepoints (Weeks 2–4) with mixed effects linear models (procedure MIXED) with interaction, where LDF frequency and exercise regimen were fixed factors and timepoint was a repeated measure. The covariance matrices were modeled using either compound symmetry (duty cycle, contralateral phase dispersion) or a first-order autoregressive (diagonal and ipsilateral phase dispersion), based on which covariance matrix yielded a lower corrected Akaike’s Information Criterion. Effect differences were calculated based on LSM with Tukey-Kramer adjustments for multiple comparisons. All data are presented as mean ± standard deviation, except LDF and gait data, which are presented as LSM ± 95% confidence interval.

### Table 1
Serum concentrations of interleukin-6 in a subset of mice.

| Group           | Week 1          | Week 2          | Week 3          | Week 4          |
|-----------------|-----------------|-----------------|-----------------|-----------------|
| Weekly-exercise | ND, n = 1       | ND, n = 3       | ND, n = 3       | ND, n = 3       |
| Weekly-sedentary| ND, n = 154.0 pg/mL, n = 1 | ND, n = 1       | ND, n = 3       | ND, n = 3       |
| Endpoint-exercise| ND, n = 2       | ND, n = 2       | ND, n = 2       | ND, n = 2       |
| Endpoint-sedentary| ND, n = 3       | ND, n = 2       | ND, n = 3       | ND, n = 252.1 pg/mL, n = 1 |

ND: not detected.
placement between the recordings.

3.2. Femoral ligation validation

After the femoral artery was ligated, perfusion in the tibia dropped rapidly from 15.3 ± 9.2 perfusion units (PU) to 3.8 ± 1.4 PU within 30 s (31 ± 19% of the baseline perfusion, p = 0.004) (Fig. 5), validating that the LDF perfusion measurements are directly associated with blood supply within the bone. Two measurements were not included in the analysis, because the LDF probe slipped off the tibia when the ligature was tightened.

3.3. Wound area

Wound images taken one week after each weekly LDF procedure showed minimal signs of inflammation for all but one incision for both exercise and sedentary groups, with either full closure (Fig. 6A) or a small, dry scab (Fig. 6B) resulting in zero wound area recorded. Wet granulation tissue was observed in only one incision site, for a sedentary mouse in Week 2 (Fig. 6C, wound area = 0.39 mm²), and the wound was healed within the subsequent week.

3.4. Serum concentration of IL-6

Circulating levels of proinflammatory marker IL-6 were below the detectable limit for all tested animals at each timepoint, except one mouse in Week 1 and one mouse in Week 4 (Table 1). Since the lower threshold of the test is 7.8 pg/mL, and serum samples were diluted up to four times to allow for duplicate measurements, serum levels of IL-6 were below 31.2 pg/mL. These levels agree with normal physiologic concentrations of IL-6, which are below 100 pg/mL in C57Bl/6J mice (Amar et al., 2007; Krause et al., 2012), suggesting the mice in our study experienced little to no systemic inflammation in response to the weekly LDF procedures. Pathologic inflammation can increase IL-6 levels up to 200–1000 pg/mL (Amar et al., 2007; Krause et al., 2012).

3.5. Gait pattern analysis

Weekly LDF procedures did not affect gait parameters during treadmill locomotion. Limb coordination did not differ between weekly and endpoint-only groups at any timepoint for either sedentary or exercise groups (Fig. 7). Sedentary groups did have small alterations in gait parameters in Week 2 compared to exercise groups (hindlimb duty cycle and diagonal and contralateral phase dispersion), possibly indicating slight discomfort or unfamiliarity with treadmill locomotion. Diagonal phase dispersion was lower in the exercise groups compared to sedentary groups during Week 4 (8.6% lower, p = 0.012).

4. Discussion

Our minimally invasive laser Doppler flowmetry technique measured in vivo intraosseous perfusion in the tibia weekly without inducing localized or systemic inflammation. LDF measures of tibial perfusion were similar between groups that received weekly procedures and groups that received only an endpoint procedure, indicating that the procedure itself did not impact measurements of intraosseous perfusion. A previous study demonstrated that LDF could be used to quantify blood perfusion in murine tibiae but noted that signs of inflammation were observed at the incision site up to three months following the procedure and suggested that it be used as an endpoint-only measure to avoid influencing bone outcomes (Roche et al., 2013). In this study, we limited the invasiveness of the procedure by reducing the incision size and preserving muscle tissue, thereby enabling repeated measurements to be performed without causing chronic increases in proinflammatory marker IL-6. Furthermore, all but one incision from the procedures were fully closed within the one-week period before the next procedure, and no visual signs of inflammation were observed throughout the study. The fast healing times observed in the small incisions from this procedure (2–5 mm) are consistent with wound closure studies, where even large (9–28 mm²), unclosed dermal biopsies heal in 7–14 days (Keylock et al., 2008).

The femoral ligation test confirmed that LDF measures are sensitive to changes in blood supply and thus are directly related to perfusion within the underlying cortical bone and marrow space. We demonstrated that femoral artery ligation reliably decreased the measured
perfusion to about 3.8 PU (31% of baseline) within 30 s in all mice. The variation in perfusion observed in each 30-s long measurement was due primarily to probe movement (when the mouse breathes), equipment noise, and changes in blood supply within the tibia. Measurement duration was not optimized in this study, but the variation around the mean value appears constant for most readings. The 30-s measurement is fairly short yet provides enough data to determine if the reading deviates from a steady value (i.e., has a substantial slope) or is excessively noisy (i.e., with large spikes), both of which are criteria for excluding the reading and repositioning the probe. Inter-measurement variation can be minimized with better probe placement as the researcher becomes more familiar with the LDF procedure. In a subsequent study, we used only two measurements per bone and achieved a standard deviation of 2.1 PU, only 15% of the mean value (Hanne et al., 2019b).

Cortical thickness in the tibia affects the depth into the marrow space that LDF can measure. Another study quantified this relationship in the murine tibia and reported that small variations in cortical thickness had minimal effect on LDF perfusion measurements (Roche et al., 2013), suggesting that LDF can be used to track longitudinal changes in bone perfusion. Because cortical thickness in the tibial diaphysis changes by about 10% from skeletal maturity at 16 weeks of age to 52 weeks of age (Ferguson et al., 2003), and can differ by sex (Somerville et al., 2004), longitudinal LDF measurements should only be performed in age- and sex-matched subjects. Finally, we found no interaction effect of daily treadmill exercise on LDF readings. Taken together, these results suggest that our minimally invasive LDF procedure is suitable for monitoring and comparing blood perfusion longitudinally in murine studies involving exercise therapy. This procedure can be used to track changes to osteovascular function, which is known to play an important role in bone development, remodeling, and repair (Marks and Odgren, 2002; Gerber et al., 1999), yet remains understudied.

In addition to providing aerobic exercise, treadmill activity also mechanically loads the bones and increases the functional strain experienced by the bones (Wallace et al., 2007; Prasad et al., 2010; Berman et al., 2019). Even slight changes to functional strain can affect osteogenesis (Frost, 2003; Ellman et al., 2013) and angiogenesis (Yao et al., 2004; Steward et al., 2016). Since other studies have shown that changes to gait kinematics and limb patterning affect bone strain (Prasad et al., 2010; Hurwitz et al., 1998), we were concerned the LDF procedure could affect locomotion patterns and confound exercise effects by altering functional strain. We found no differences in duty cycle or interlimb coordination between weekly and endpoint-only groups at any timepoint, indicating that weekly LDF measurements do not alter gait patterns (and thus functional strain) during treadmill exercise. Although not the main focus of this study, aerobic treadmill exercise was anticipated to cause increased tibial perfusion over time due to vascular growth and adaptation, as we have previously found in studies using the same exercise regimen and LDF technique (Hanne et al., 2019a; Hanne et al., 2019b). Perfusion is a functional measure of not only the amount and direction of blood flow but also vascular permeability and capillary density (Griffith, 2014). Treadmill exercise may be affecting the size, number, or cellular makeup of the vasculature without causing changes in perfusion. Rats that performed a similar treadmill routine for two weeks had a 19% increase in the number but not total area of blood vessels in the proximal tibial metaphysis compared to the sedentary group (Yao et al., 2004). A more rigorous aerobic

![Fig. 7. Treadmill locomotion patterns involving the LDF affected limb (right hindlimb). Gait patterns were not significantly different between weekly and endpoint-only groups at any timepoint for A) hindlimb duty cycle ratio, B) diagonal phase dispersion, C) contralateral phase dispersion, or D) ipsilateral phase dispersion. Diagonal and contralateral phase dispersion were lower in exercise than sedentary groups at some timepoints. Data are presented as least squares mean ± 95% confidence interval. *p < 0.05 exercise vs. sedentary. ^p = 0.066 exercise vs. sedentary.](image-url)
exercise intervention, such as free access to running wheels where mice will run 4–10 km daily (Gertz et al., 2006; Styner et al., 2017), may have a larger and more detectable effect on perfusion. Nevertheless, our study confirmed that weekly LDF procedures will not confound perfusion measurements in future exercise studies.

Stress, like inflammation and aerobic exercise, can also affect vascular function. Neuropeptide Y, which is expressed during stress response, has been shown to be both angiogenic and vasoconstrictive, which could increase blood pressure and the amount of vasculature in bone (Kuo and Zukowska, 2007). Although we found almost no detectable increases in IL-6 or visible signs of inflammation at the incision site, both sedentary and exercise groups had a significant increase in tibial perfusion in the third week that was resolved by the fourth week. Stress may have played a role in increasing perfusion in the third week; all mice were handled daily for either treadmill exercise or the sedentary treadmill activity and had blood drawn weekly, which could induce a stress response. The increased perfusion lasted only one week and was present in both exercise and sedentary groups. The low serum yield in these small animals limited our assessments to one inflammation marker in a subset of mice – other markers of inflammation may have been elevated, or IL-6 may have been elevated in the subset of mice that were not measured. IL-6 is known to be elevated following musculoskeletal trauma (Reikeras et al., 2014) and has been shown to stimulate bone resorption (Ishimi et al., 1990), which would preclude the use of LDF as a weekly tool in most bone experiments.

This study had several limitations that warrant attention in future studies. A primary concern of this technique is the removal of a small area of the periosteum, a highly vascularized tissue that contains osteoblast precursor cells (Colnot et al., 2012). LDF measures of intracortical perfusion in the tibiae of juvenile ewes dropped by 25% immediately following the removal of the periosteum from the medial aspect (Kowalski et al., 1996). Although the amount of periosteum removed in our procedure is small (about the size of our probe, 0.5 mm² area), the effects of periosteal removal were not examined and may affect bone tissue function. The effects of weekly LDF procedures on bone remodeling and homeostasis were not examined in this study. This study did not compare LDF results to other promising emerging techniques for examining osteovasculature in vivo. Several new higher resolution PET scans can be used in rodent bones (Tomlinson et al., 2012), and emerging MRI techniques (e.g., blood oxygen level-dependent MRI and intravoxel incoherent motion MRI (Griffith, 2014)) greatly improve resolution and do not require contrast agents, but these techniques remain prohibitively expensive.

5. Conclusions

Weekly LDF procedures performed over four weeks did not induce measurable signs of inflammation or significantly alter gait patterns during treadmill exercise. Unlike other existing methods used for measuring the vascular network in bone, this procedure can be performed in vivo, is repeatable without confounding study controls, is relatively simple to perform, and is inexpensive. Monitoring intracortical perfusion serially with LDF provides a functional measure of blood flow, enabling researchers to track changes to the osteovasculature noninvasively during disease progression and interventions.

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Declaration of competing interest

None.

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CRediT authorship contribution statement

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