- 1 The use of near-infrared systems for investigations of haemodynamics in human *in vivo*
- 2 **bone tissue:** A systematic review
- 3 <u>*Running Title:*</u> NIR for haemodynamic bone measurements.
- 4 *Authors:* Robert Meertens^a, Francesco Casanova^b, Karen M Knapp^a, Clare Thorn^b, William
- 5 David Strain^b,
- 6 <u>Affiliations:</u>
- 7 a. Medical Imaging, University of Exeter Medical School, South Cloisters, St Luke's Campus,
- 8 Heavitree Road, Exeter, United Kingdom, EX2 1LU
- 9 b. Diabetes and Vascular Research Centre, University of Exeter Medical School and
- 10 National Institute of Health Research Exeter Clinical Research Facility, Barrack Rd, Exeter,
- 11 United Kingdom, EX2 5DW
- 12 <u>Correspondence Author:</u>
- 13 Robert Meertens
- 14 University of Exeter Medical School (UEMS)
- 15 Room G.23, South Cloisters, St Luke's Campus
- 16 Heavitree Road, Exeter, EX2 1LU
- 17 Phone: 01392 722511
- 18 Email: <u>r.m.meertens@exeter.ac.uk</u>
- 19 *<u>Competing Interests</u>: None of the authors have any competing interests to declare.*
- 20 <u>Author Contributions:</u> RMM carried out all screening, data extraction and quality assessment
- of studies in the review. FC independently screened full text articles and undertook quality
- 22 assessment of studies. DS and KMK also independently screened full text articles. CET and
- all authors helped in the drafting and revision of this manuscript.

1 Abstract:

2 A range of technologies using near infrared (NIR) light have shown promise at providing real 3 time measurements of haemodynamic markers in bone tissue *in vivo*, an exciting prospect 4 given existing difficulties in measuring haemodynamics in bone tissue. This systematic 5 review aimed to evaluate the evidence for this potential use of NIR systems, establishing their 6 potential as a research tool in this field. Major electronic databases including MEDLINE and 7 EMBASE were searched using pre-planned search strategies with broad scope for any in vivo use of NIR technologies in human bone tissue. Following identification of studies by title and 8 9 abstract screening, full text inclusion was determined by double blind assessment using 10 predefined criteria. Full text studies for inclusion were data extracted using a predesigned 11 proforma and quality assessed. Narrative synthesis was appropriate given the wide 12 heterogeneity of included studies. Eighty eight full text studies fulfilled the inclusion criteria, 57 addressing laser Doppler flowmetry (56 intra-operatively), 21 near infrared spectroscopy 13 and 10 photoplethysmography. The heterogeneity of the methodologies included differing 14 haemodynamic markers, measurement protocols, anatomical locations and research 15 applications, making meaningful direct comparisons impossible. Further, studies were often 16 17 limited by small sample sizes with potential selection biases, detection biases, and wide 18 variability in results between participants. Despite promising potential in the use of NIR light 19 to interrogate bone circulation, the application of NIR systems in bone requires rigorous 20 assessment of the reproducibility of potential haemodynamic markers and further validation of these markers against alternative physiologically relevant reference standards. 21

22

23 Keywords:

24 Bone; haemodynamics; near infrared; emerging technologies; optical systems.

1 Introduction

2 Bone is a dynamic and vascular tissue, dependent on this perfusion to meet its metabolic demands. However, measuring the haemodynamics of the osseous vasculature is difficult 3 4 with existing imaging modalities due to the high density and mineral content of bone. 5 Imaging protocols typically involve nuclear medicine scans, positron emission tomography 6 (PET) or contrast enhanced magnetic resonance imaging (MRI) which are expensive, have 7 limited clinical access, involve injections and do not readily allow longitudinal evaluation. In 8 addition, these techniques do not allow evaluation of oxygen saturation within the tissue, as 9 these modalities can only measure markers of gross perfusion and blood volume based on 10 rates of radiopharmaceutical or gadolinium contrast uptake [1]. 11 Near infrared (NIR) optical systems are a potential solution. They have the advantages of 12 being non-invasive, non-destructive, non-ionising, inexpensive and allowing repeat or 13 continuous measurements. NIR optical systems involve transmitting and receiving 14 designated optical wavelengths using probes at an anatomical site. These systems take 15 advantage of the difference in absorption characteristics of oxygenated and deoxygenated haemoglobin to record markers of bone haemodynamics in real time including tissue 16 oxygenation, perfusion, blood flux and blood volume [2]. Although NIR light only penetrates 17 human tissue superficially, systems have been shown to be able to record data non-invasively 18 19 through *in vivo* tissue to depths of up to 4 cm [3].

20 NIR optical systems are already established research tools in muscle physiology and

transcranial cerebral circulation [4, 5]. The use of transcranial NIR optical systems,

22 demonstrates that NIR light can penetrate bone tissue, and raises the possibility of its use in

assessing bone haemodynamics. Development of NIR optical systems could benefit research

in a range of bone pathologies with suspected vascular components, such as the earlier

detection, prevention and monitoring of haematopoietic malignancies, osteoporosis, nonosteoporotic fragility fractures, slow fracture healing and forms of arthritis. The primary aim
of this systematic review is to gauge the existing knowledge base on the ability of NIR
optical systems to measure haemodynamic markers of blood supply in bone tissue, and to
establish their potential as a research tool in this field.

6 *Methods*

7 An initial scoping review suggests no previous systematic review on this topic has been 8 performed, therefore a diverse but relatively small evidence base was expected. As such 9 broad search criterion were established for studies investigating in vivo bone tissue at any anatomical bony site in human participants. Studies using NIR optical systems in either 10 11 healthy or diseased human populations (or both) were considered eligible. Any optical 12 system utilising NIR wavelengths (600-1050 nm) using a haemodynamic marker that could 13 give insight into the haemodynamic state of the bone tissue sampled was considered eligible. 14 This could include systems measuring haemodynamic markers of blood flux, oxygen 15 saturation, oxygenated or deoxygenated haemoglobin concentration changes or blood volume. Any indication for utilising the NIR optical systems was eligible, including 16 17 validation for a particular application, diagnosis of disease states, prognostic assessment, longitudinal monitoring or screening purposes. 18

Given the broad scope of the review, a wide range of reference standards were eligible, such
as bone biopsy for measuring oxygenation of bone tissue; MRI, nuclear medicine or
angiographic protocols for measuring bone perfusion; or, direct clinical observation of tissue
perfusion intra operatively. Again, allowing for broad search criterion, a comparator test or
external reference standard for results was desirable, but not essential for study eligibility.
There were no restrictions on the geographical location or year of publication for studies, but

only studies published in English were eligible. Case studies of individual participants were
 excluded, as were opinion pieces, reviews and editorials.

3 <u>Search Strategy</u>

Pre-planned search strategies using medical subject headings and keywords were developed 4 5 around key inclusion criteria for searches using MEDLINE and EMBASE online databases. 6 Grey literature databases including conference abstracts, theses, and unpublished works were 7 searched, including Web of Science, Proquest, OpenSIGLE, OpenGrey, and the British Library EThOS database. At this point the reference list and further citations of any eligible 8 9 studies were hand searched. Authors were contacted where clarification or full text access was required. All searching was carried out by the primary author who then removed 10 11 duplicates and screened the returned results based on title and abstract. The remaining 12 studies were considered for inclusion based on a blinded full text assessment by two independent reviewers (RM and FC). Upon agreement of eligible studies, the primary author 13 14 independently extracted relevant key data using a pre-piloted data extraction proforma (extracted data is presented in Supplementary Materials 1). Searches were performed in 15 September 2015, with a repeated addendum search of MEDLINE and EMBASE performed 16 prior to publication. 17

18 <u>Assessment of Risk of Bias</u>

Identified studies were also scrutinised by the primary author across 6 domains of potential
intrinsic biases using the Cochrane based Risk of Bias Assessment Tool for Non-randomised
Studies (RoBANS) tool to form an overall judgement of reporting quality as either "good",
"fair" or "poor" [6]. External applicability and generalisability of studies was also assessed
using these categorical criterion. Methodological patterns or trends for "unclear" or "high"
areas of risk of bias were identified and their potential effects are discussed in context within

the narrative synthesis of the review (extended assessment results are presented in
 Supplementary Materials 2).

3 Data Synthesis

Due to the clinical and methodological heterogeneity of the identified studies, statistical 4 5 synthesis via meta-analysis was not appropriate. As such, narrative synthesis of identified studies has been performed primarily synthesised around the NIR optical systems of interest 6 7 and grouped around the anatomical locations or target conditions investigated. Discussion is also in context of the haemodynamic markers employed, types of participants, and 8 9 comparators. The review also includes discussion on the strength of the evidence accumulated including consideration of the risk of bias of individual studies, patterns in 10 11 biases, the consistency of results across studies, the applicability of results across the general 12 population, as well as the strengths and limitations of the review process. Further detail on the protocol for the review are registered on the PROSPERO database [7]. 13

14 *Results*

15 **Overview of Findings**

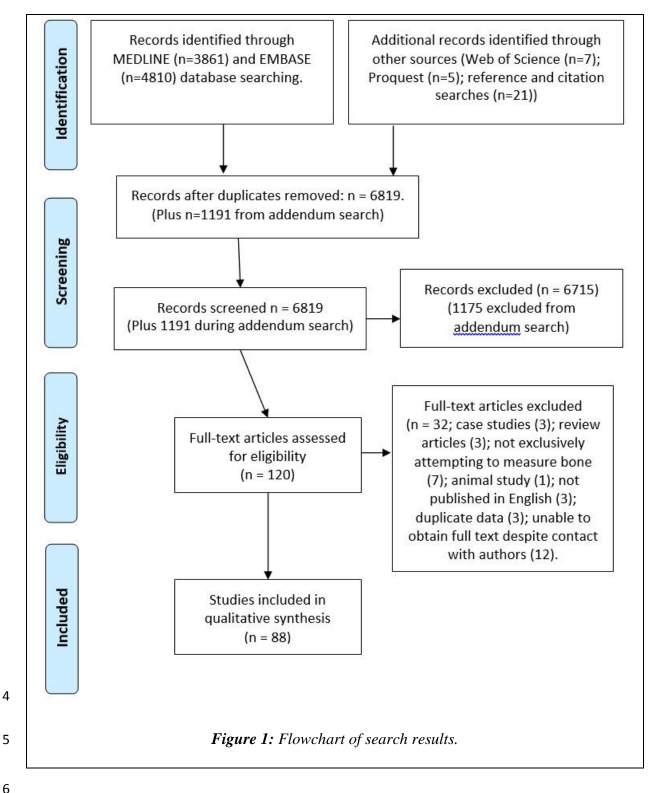
A summary of search results and data extraction is presented in Figure 1 and Supplementary
Materials 1. Eighty eight studies were included for full text analysis. These were
predominately peer reviewed journal articles along with 5 conference proceedings [8-12] and
2 theses [13, 14]. Study publication dates ranged from 1987 to 2017 and there was wide
geographical variation with publications based in 13 different countries.

Twenty one studies were identified using near infrared spectroscopy (NIRS) [3, 8, 11, 13, 15-

30] and 10 studies were identified using photoplethysmography (PPG) non-invasively on *in*

- vivo bone tissue [10, 24, 31-38]. Fifty seven studies were identified using laser Doppler
- flowmetry (LDF) *in vivo* [9, 12, 14, 39-94] of which 56 were intra operative. Only one study

- using LDF attempted to measure bone non-invasively [92]. Most studies were in adult
- populations with only 7 of the 56 LDF studies including paediatric populations [41, 46, 70-
- 72, 78, 91].



1 <u>System Characteristics</u>

2

<u>Near Infrared Spectroscopy (NIRS)</u>

Studies most commonly utilised "continuous wave" NIRS systems using spatially resolved or 3 modified Beer-Lambert law algorithms. These systems utilise at least 2 or 3 discrete 4 5 wavelengths of NIR light in the first NIR window (600 nm-1200 nm) to detect changes in oxygenated and deoxygenated haemoglobin concentrations by taking advantage of their 6 7 different attenuation properties. This also allows these systems to measure oxygen saturation 8 and total haemoglobin concentration changes in the tissue sampled. The oxygen saturation is 9 the mean saturation across the arterioles, capillaries and venules. All were using reflectance spectroscopy (light scattered back from tissue) apart from transmission spectroscopy (light 10 11 scattered through tissue) studies looking at the calcaneus [8, 22, 30] and mandible [21, 23]. 12 For reflectance spectroscopy, probe spacing (affecting the depth of tissue measured) varied from 10 mm [13, 19, 25], 20 mm [13], 25 mm [8, 20], 30 mm [15-18, 26], and 40 mm [3, 18]. 13 14 Temporal resolution typically ranged from 1 second [17] to 12 seconds [20, 30]. Use of time resolved spectroscopy systems are reported in five studies [8, 20, 22, 29, 30]. 15 16 These allow measurements of absorption and scattering coefficients for bone tissue and therefore absolute concentrations of oxygenated and deoxygenated haemoglobin are 17 calculated with participants at rest (in µM) [8, 19, 20, 22, 23]. Farzam et al. 2013 also reports 18 19 on use of a frequency domain based NIRS system utilising 15 NIR laser sources (5 each at 3 different wavelengths) based around 2 photomultiplier detectors [19]. 20 21 Six studies used NIRS systems utilising a broadband spectrum of NIR light enabling the system to calculate absolute concentrations of haemoglobin and in some cases also water, 22

collagen, mineral and lipid content in the bone tissues sampled [8, 20, 22, 23, 28-30].

Additionally, four studies also measured markers of blood flow rates utilising emerging
 diffuse correlation spectroscopy technology [8, 20, 27, 30].

3

<u>Photoplethysmography (PPG)</u>

We identified 10 PPG studies from two research groups investigating the patella and tibia 4 [10, 24, 31-38]. These studies utilised bespoke PPG systems making use of an isobestic 804 5 6 nm NIR wavelength to measure the amplitude of pulsatile flow in bone tissue based on 7 attenuation changes in reflected signal. When reported, inter-probe spacing ranged from 15-25 mm. Some studies also measured overlying skin tissue, utilising a less penetrative 8 9 wavelength in the visible range (either 526 nm or 560 nm). All studies used "peak to peak" amplitude during pulsatile flow as their primary outcome measure, typically measured and 10 11 averaged over a 30-60 second period to gain a mean value representing the strength of 12 pulsatile flow. Changes were then typically compared to baseline measurements during and/or after an intervention in either relative (i.e. percentage change) or absolute terms (in 13 14 Volts).

15

Laser Doppler Flowmetry (LDF)

16 Most studies reported the use of a commercial LDF system utilising a monochromatic wavelength typically at 632 nm, 780 nm, 785 nm, or 830 nm. LDF systems derive a measure 17 of flowmetry from the Doppler frequency shift induced by the moving red blood cells. 18 19 Changes in the frequency power spectrum of the reflected light provide a measure of the concentration of moving red blood cells (blood flux or flowmetry) in arbitrary units 20 21 (described as perfusion units, flux units, or detector signal measured in milliVolts). A mean amplitude of pulsatile flow is calculated over a set period (typically 60s or less). Comparison 22 of these values across participants should be done cautiously, and typically these values are 23

used to assess relative changes following a vascular challenge (either expressed as absolute
 change or relative percentage change).

LDF systems only measure to a depth typically less than 4 mm [95], and therefore 56 of 57 studies involve intra operative use of LDF with probe placement directly on bone or intraosseous measurements. Two studies utilised laser speckle techniques involving laser light interference from a tissue surface and mapping blood flux to produce a 2D flowgraphy "heat" map. These were used intra-operatively to study the flux in the surface of the bone tissue up to a depth of 2mm, namely the femoral head [50] and the cochlea promontory [73].

9 <u>Study Characteristics</u>

10 Studies looking to investigate bone tissue non-invasively used a wide range of superficial 11 anatomical bone sites including the tibia, calcaneus, radius, ulna, greater trochanter of the femur, patella, mandible and manubrium/sternum. In addition, invasive LDF procedures also 12 investigated deeper bone tissue including the proximal femur, acetabulum, cochlea, maxilla, 13 metatarsal, humerus and lumbar vertebrae. Studies could broadly be split into two categories. 14 Firstly, studies investigating the feasibility of NIR technologies for measuring bone tissue in 15 16 healthy adults. Secondly, studies that used previous evidence to defend the validation of NIR technologies, and applied these systems for a physiological research purpose. 17

18 <u>Feasibility Studies</u>

The most substantial work demonstrating NIRS and PPG could truly measure haemodynamic parameters of bone tissue non-invasively was based around studies that demonstrated significantly lower oxygen extraction rates and reperfusion rates in bone tissue compared to adjacent muscle tissue. These two parameters were derived during and after extended arterial occlusions of the leg respectively, reflecting bone's lower metabolic rate compared to muscle [11, 15, 16, 19, 26]. Three studies demonstrate a non-significant confounding contribution of

superficial tissue to PPG and NIRS measurements of the patella and tibia by selectively
altering superficial tissue haemodynamics either through the localised introduction of
vasodilating or vasoconstricting elements such as gentle compression [32], cold packs [13],
nitro-glycerine patches [32] and liniment [34]. However, Klasing et al. 2003 does report a
systematic influence from superficial tissue on NIRS measurements, although it is noted this
is a much earlier study [26].

7 Binzoni et al. 2013 was the only study focusing purely on non-invasive LDF measurements in bone through the skin surface. Feasibility work was undertaken in healthy participants 8 with a source/detector spacing of 1.5 cm giving enough penetrative depth to measure 9 10 superficial bone *in vivo* at four anatomical sites [92]. Diffuse correlation spectroscopy 11 technology with NIRS also provides blood flux information non-invasively but at greater 12 tissue depths and may supersede LDF here given the added advantages of NIRS systems [30]. Other applications of NIRS and PPG systems applied to healthy populations included 13 14 haemodynamic measurements taken during positional changes such as leg extension, or head 15 up/down tilt to demonstrate oxygen saturation and blood volume changes [10, 13, 17, 24, 37, 38]. Four studies examined the effects of positive and negative external pressure changes 16 17 using lower body pressure chambers on the amplitude of PPG pulsatile flow and oxygen saturation [24, 31, 33, 38]. In the case of Larsson et al. 2014, the use of hyperbaric chambers 18 19 and variable respired oxygen levels were also used to observe their effects on bone haemodynamics at the patella [31]. These studies concluded that the use of pressure 20 21 alterations may have applications for therapy by mimicking weight-bearing physiology in a microgravity environment, or for potential therapeutic applications involving vascular 22 mechanisms in bone. Three studies investigated the effects of exercise on tibia and patella 23 haemodynamics using exercise bikes [13], rowing machines [13, 28] or high intensity 24

1	quadriceps workloads [36]. For time resolved NIRS systems, absolute measurements could
2	be taken on healthy adult participants simply at rest [8, 20, 22, 29, 30].

Physiology Research Applications

Studies using NIR optical systems for physiological research in diseased populations were
predominately LDF studies, focusing on dynamic flux changes in response to fixed stimuli or
during various stages of a surgical intervention. Some studies used case-control designs to
correlate these results with other clinically relevant observations assessing the predictive role
of intra-operative LDF on outcomes.

The proximal femur and hip joint represented one of the biggest areas of interest for research 9 10 involving LDF in bone tissue, with 19 studies [9, 39, 41-44, 48-50, 54, 58, 65-67, 76, 77, 82, 11 84, 91] found investigating a range of clinical applications including osteonecrosis (either idiopathic, steroid induced, or following neck of femur fracture) [50, 65, 82, 84], approaches 12 to hip resurfacing arthroplasty [9, 39, 42, 67, 77] or total hip replacement following arthritis 13 [44, 48, 49, 58, 66, 82], femur-acetabular impingement [43], hip joint debridement [76], and 14 congenital developmental problems [41, 54, 91]. Specifically most studies took 15 16 measurements from the femoral head and neck, but others also looked at intra-trochanteric regions [65, 84], the greater trochanter [48, 58, 66], proximal femoral shaft [58], medial 17 calcar [58], or acetabulum [49, 54]. 18

19 There were 6 studies identified that used LDF intraoperatively to investigate patellar blood

20 flux changes before and after positional changes or different surgical manoeuvres in adult

21 patients having total knee arthroplasty [52, 55, 57, 61, 74, 81]. Five studies used LDF

technology to measure blood flux in the manubrium [40] or sternum [51, 59, 60, 75] during

and following open cardiac surgery involving the internal mammary arteries. Seven studies

looked at the use of LDF to measure blood flux in the mandible [14, 62, 88], maxilla [90, 93,

1	94] or both [63, 89] for patients undergoing maxilla-facial or dental surgery including
2	wisdom tooth removal [14], dental implants [62, 88, 94], osteomyelitis [89] or corrective
3	maxillary surgery [63, 90, 93]. Eleven studies reported on the use of LDF to assess blood
4	flow in the bony wall of the cochlea [45, 46, 68-73, 79, 80, 87] to investigate a range of inner
5	ear pathologies including otosclerosis [73, 80], Meniere's disease [79], curative treatment for
6	uncontrolled drooling [68, 69] or idiopathic or congenital hearing loss and surgical
7	implantation of hearing aid devices [46, 70-72]. Four studies from the same group
8	investigated the use of LDF technology for assessing bone perfusion during the surgical
9	debridement of bone tissue in adult patients compromised by acute trauma (such as open
10	fracture) and/or osseous infections such as osteomyelitis [47, 83, 85, 86].
11	LDF has also been used to investigate successful post-operative monitoring of fibular bone
12	grafts [78]; intra-osseous haemodynamic measurements of thoracic and lumbar vertebral
13	bodies during mimicked unilateral and bilateral ligation [53]; for investigating the predictive
14	ability of humeral head fracture patterns for determining the risk of humeral ischemia [56];
15	and for investigating the blood flux in the first metatarsal head during corrective surgery [12,
16	64].
17	The clinical utility of NIRS has been explored in the mandible [18, 21, 23] and sternum [25].
18	In a case-control manner, mandibular conditions such as osteoradionecrosis post radiotherapy
19	[21, 23], and fibular grafting post tumour removal [18], as well as chest wall measurements
20	taken post cardiac surgery [25] were explored. Results showed potential for the use of NIRS
21	for post-operative monitoring using oxygen saturation measurements, however the small
22	sample sizes and shortage of adverse outcomes precluded the ability to identify a diagnostic

23 predictive threshold.

Naslund et al., 2007 used PPG to investigate haemodynamic differences between cases of
Patello-Femoral Pain Syndrome with age, gender and body mass index (BMI) matched
controls. This study demonstrated cases of Patello-Femoral Pain Syndrome had significantly
reduced PPG pulsatile amplitude when flexing their affected knee to 90 degrees for 5
minutes, supporting the hypothesised ischaemic element to pathogenesis [35].

6 <u>Reliability of NIR systems</u>

7 Broadly speaking, study results were predictive of the expected haemodynamic changes during interventions. However, across most studies wide variability in results between 8 9 participants was evident, especially across applications of LDF. Crucially, no studies were identified specifically addressing the reliability or reproducibility of NIR optical systems. 10 11 Studies that did attempt to assess reliability typically did this in a superficial or *ad hoc way* in 12 small samples without statistical analysis of agreement or reliability. Approaches included comparing contralateral results, analysing repeat measurements for their variability (either in 13 14 the same or different sessions), or simple comparison of results with existing literature in animal studies or involving different tissue types. 15

This wide variation in results may prohibit the development of useful diagnostic thresholds
for NIR optical systems [60, 76]. Wide biological variability in vascular measurements is a
known barrier to research in this field [96] and this is also reflected in suboptimal
reproducibility in alternative modalities for measuring bone tissue haemodynamic markers
such as MRI [97, 98].

Variability is likely to be affected by the heterogeneity of bone tissue and the small sampling
volume of NIR optical systems (particularly LDF) with sampling of small arterioles or less
vascular trabecular striate potentially altering readings. It is important to tease out what
variation in results is attributable to measurement error and what reflects normal

physiological ranges. Likewise, most studies typically used only one, or a small group, of
 operators. Few studies took repeat measurements, or multiple readings at adjacent sites,
 which is suggested due to this small penetration depth and sampling volume of NIR optical
 systems.

Other potential factors affecting reliability specific to LDF studies include the comparison of 5 6 results in studies where the probe is placed non-invasively, directly on the bone surface, or 7 intra-osseously. The sensitivity of probes to movement also means stable probe placements 8 during surgical operations is important and some studies reported holding probes by hand or using bespoke probe holders [40, 51]. Some study designs involved having to move probes 9 10 in between measurements to facilitate surgery, which can introduce measurement error given 11 the small sampling volume and potential for measuring a different vascular bed when probes 12 are repositioned [77, 81]. In addition, as these LDF studies were intra operative the effects of anaesthesia, blood loss during surgery and direct impact of surgery cannot be discounted as 13 sources of error [41]. When placing probes intra-osseously, the effect of drilling on intra-14 osseous blood flow is unknown and flushing of the probes is required to remove clotting 15 around the measurement site, which could otherwise prohibit flow and affect readings. 16

17 *Quality Assessment*

An overview of quality assessment ratings is presented in Supplementary Materials 2. A general methodological issue with many studies was the poor description of inclusion and exclusion criteria, recruitment strategies, and summary information of important participant demographics. In the absence of specific details, this opens many studies to potential criticism of selection biases. Similarly, most feasibility studies involved small cohorts of healthy, young and predominately normal BMI participants, raising concerns around the generalisability of non-invasive applications of NIR systems in wider demographics and

disease states. Many studies also included small sample sizes of participants without sample
 size justification.

3 Attrition or incomplete results were rarely directly addressed. How incomplete results are 4 handled can lead to biases in the accuracy of results and so it is important these are reported in context with study findings, especially at the early stages of the technological development 5 6 of NIR optical systems. Generally, most studies had a low risk of performance and reporting 7 bias as testing protocols were clearly pre-stated and all participants received the same testing. 8 In the case of studies investigating the potential feasibility of NIR optical systems on healthy 9 participants, detection bias was considered generally low, despite studies not always reporting if testing and data analysis was strictly protocoled, or if acquisition and data 10 analysis was blinded to participant information/status. However, in studies investigating 11 12 different sub populations for physiological differences, this was deemed a more significant potential bias risk, especially when LDF or oxygen saturation with NIRS was used as an 13 14 outcome measure, where probe placement can be easily adjusted for minor subjective corrections in results, or the most suitable data could be selectively sampled for data analysis. 15 It is acknowledged that with intra operative LDF studies that in most cases LDF operators 16 were not blinded to their participant status as they were likely performing the surgical 17 procedures. As such, detection biases are hard to avoid. 18

Studies were all also deemed at risk of "other biases" as there are inherent unknowns around the use of NIR optical systems for these applications in bone tissue, as relatively weak validation exists. Along similar lines many studies involving NIRS and PPG systems reported use of bespoke made systems, reducing the applicability of results as there is evidence for systematic differences in haemodynamic measurements across commercial systems from different manufacturers [99].

1 There was also an applicability issue evident when considering intra-operative LDF findings 2 to guide clinical practice or normal physiology. Often studies found reduced blood flux at 3 various stages during surgical interventions, but without the ability to take pre and post-4 operative readings the clinical importance of these findings is hard to distinguish. Likewise, 5 obtaining healthy control data is also ethically difficult given the invasive approach taken. 6 Similarly, some studies only presented relative percentage changes in haemodynamic 7 markers with time. Whilst this demonstrates the responsiveness of NIR optical systems in 8 bone, the applicability of these results is limited without development of absolute 9 haemodynamic quantitative markers and threshold results for normal physiology that can guide research into diseased states. 10

11 Discussion

A large number of studies have been identified utilising NIRS, PPG and LDF systems to investigate bone tissue either non-invasively or intra-operatively. The wide heterogeneity in anatomical sites and investigated applications demonstrates the demand for the types of information NIR technology promises. The studies identified are predominately early stage proof of concept type work which often illustrate the promise for future clinical and research applications. However, there are a wide range of challenges that require addressing to advance this field of research in bone health.

The need for further work on the reliability and reproducibility of NIR optical systems for
repeat measurements across different operators and participants has been identified.
Likewise, continued investigation around whether variability is biological or equipment
based (or both) is important, as this is still not clear from the literature. If physiological
variability is wide at an individual level, NIR optical systems may be unhelpful for
development of individual diagnostic thresholds, but NIR research could perhaps still

elucidate important haemodynamics differences between sub populations of interest given the
 current lack of alternative research tools.

Few studies attempted to validate NIR optical system results against an external comparator or reference standard, such as microangiography, MRI protocols, nuclear medicine and PET protocols. However, those that did presented promising results [50, 56, 82]. External validation remains crucial to give credence to future NIR optical systems. Alternatively, confidence could be gained through correlation of other relevant indicators of bone health, such as bone density, blood markers of bone metabolism, or longitudinal patient follow up when used to guide operative cases.

Establishing the generalisability of results is also crucial to the development of these 10 11 technologies. This includes gauging the expected normal physiological variability expected 12 between different ethnicities, ages, genders and body habitus as well as during the operative state. It appears the influence of overlying tissue on non-invasive measurements requires 13 14 careful consideration but may be overcome with continued technological advances. As always, patient tolerability of protocols should also be considered, as well as ruling out any 15 potential negative impact of intra-operative use of LDF, which was not addressed by the 16 17 studies identified.

Technological advances in NIR optical systems holds the key to the future of this application.
The development of time-resolved NIRS systems promises the ability to measure absolute
concentrations of oxygenated and deoxygenated haemoglobin non-invasively with the patient
at rest. This facilitates easier measurement protocols and more appropriate data comparison
between participants. It may also help to address some of the variability in measurements
between different tissue types resulting from the significant variability in light scattering.
Likewise, the possible development of commercially available broadband NIRS systems can

allow the potential measurement of other relevant components of bone tissue such as mineral,
 lipid, collagen and water concentration [8, 22, 29, 30]. With improved probe design the
 ability to measure deeper tissues may also be feasible in the future.

Further work is required before NIR optical systems can be considered a valid and reliable
research tool of vascular bone health. However, the wide and varied literature base identified
in this review ultimately highlights the strong promise of this application of NIR optical
systems, which potentially offers real time, safe and inexpensive measurements of bone tissue
haemodynamics.

9

10 Acknowledgements:

This systematic review was supported by the College of Radiographers Industry Partnership
Scheme (CORIPS) Doctoral Fellowship Grant (Applicant 003). The CORIPS are providing
financial support but have no input into the design, performance or analysis of this systematic
review.

Thanks goes to Chris Hyde, Zhivko Zhelev and Jamie Peters from the University of Exeter
Medical School Institute of Health Research for their support with methodological aspects of
this systematic review. Thanks also to Sarah Merson for providing her support to this
project.

WDS, FC and CT would like to acknowledge the NIHR Exeter Clinical Research Facility and
the NIHR Collaboration for Leadership in Applied Health Research and Care (CLAHRC) for
the South West Peninsula. The views expressed in this publication are those of the author(s)
and not necessarily those of the NIHR Exeter Clinical Research Facility, the NHS, the NIHR
or the Department of Health in England.

<u>References:</u>

2	1.	Dyke, J.P. and R.K. Aaron, Noninvasive methods of measuring bone blood perfusion.
3		Annals of the New York Academy of Sciences, 2010. 1192: p. 95-102.
4	2.	Binzoni, T. and L. Spinelli, Near-infrared photons: a non-invasive probe for studying
5		bone blood flow regulation in humans. J. Physiol. Anthropol., 2015. 34(1): p. 1-6.
6	3.	Aziz, S.M., F. Khambatta, T. Vaithianathan, et al., A near infrared instrument to
7		monitor relative hemoglobin concentrations of human bone tissue in vitro and in vivo.
8		Rev. Sci. Instrum., 2010. 81(4): p. 043111.
9	4.	Vardi, M. and A. Nini, Near-infrared spectroscopy for evaluation of peripheral
10		vascular disease. A systematic review of literature. European journal of vascular and
11		endovascular surgery : the official journal of the European Society for Vascular
12		Surgery, 2008. 35 (1): p. 68-74.
13	5.	Scheeren, T.W., P. Schober and L.A. Schwarte, Monitoring tissue oxygenation by
14		near infrared spectroscopy (NIRS): background and current applications. Journal of
15		clinical monitoring and computing, 2012. 26(4): p. 279-87.
16	6.	Kim, S.Y., J.E. Park, Y.J. Lee, et al., Testing a tool for assessing the risk of bias for
17		nonrandomized studies showed moderate reliability and promising validity. Journal of
18		clinical epidemiology, 2013. 66(4): p. 408-414.
19	7.	Meertens, R.M., C. Hyde, K.M. Knapp, et al. The use of near infrared spectroscopy
20		for measuring haemodynamics of the microvascular blood supply in bone tissue:
21		protocol for a systematic review. 2015; Available from:
22		http://www.crd.york.ac.uk/PROSPERO/display_record.asp?ID=CRD42015024463.
23	8.	Konugolu Venkata Sekar, S., A. Farina, E. Martinenghi, et al. Time-resolved diffused
24		optical characterization of key tissue constituents of human bony prominence

locations. in *European Conferences on Biomedical Optics*. 2015.

1	9.	Beaule, P.E., P. Campbell and P. Shim, Femoral head blood flow during hip
2		resurfacing. Clin. Orthop. Relat. Res., 2007(456): p. 148-152.
3	10.	Becker, R.L., J. Siamwala, B.R. Macias and A.R. Hargens, Tibial bone microvascular
4		flow changes as compared to anterior tibial macrovascular flows during body tilt. J.
5		Orthop. Res., 2017. 35 (no pagination).
6	11.	Meertens, R., K. Knapp, D. Strain and F. Casanova, Near infrared spectroscopy: A
7		potential tool for assessing haemodynamic markers of the microvascular blood supply
8		within bone tissue. Osteoporos. Int., 2016. 27 (2 Supplement): p. S636-S637.
9	12.	Minokawa, S., M. Naito, I. Yoshimura, et al., Effect of blood flow of the metatarsal
10		head with hallux valgas after minimally invasive distal linear metatarsal osteotomy. J.
11		Orthop. Res., 2016. 34 (no pagination).
12	13.	Alneami, A.I., Measuring blood perfusion in bone using NIRS (bone optical
13		spectroscopy). 2015, Northeastern University: Ann Arbor. p. 69.
14	14.	Al-Kassab, B.A.M., Laser Doppler flowmetry in measuring microvascular responses
15		in the oral mucosae and mandibular bone. 1995, The University of Manchester
16		(United Kingdom): Ann Arbor.
17	15.	Binzoni, T., S. Blanchi, J.H. Fasel, et al., Human tibia bone marrow blood perfusion
18		by non-invasive near infrared spectroscopy: a new tool for studies on microgravity. J.
19		Gravit. Physiol., 2002. 9(1): p. P183-4.
20	16.	Binzoni, T., T. Leung, V. Hollis, et al., Human tibia bone marrow: defining a model
21		for the study of haemodynamics as a function of age by near infrared spectroscopy. J.
22		Physiol. Anthropol. Appl. Human Sci., 2003. 22(5): p. 211-218.
23	17.	Binzoni, T., T.S. Leung, C. Courvoisier, et al., Blood volume and haemoglobin
24		oxygen content changes in human bone marrow during orthostatic stress. J. Physiol.
25		Anthropol., 2006. 25 (1): p. 1-6.

1	18.	Cai, Z.g., J. Zhang, J.g. Zhang, et al., Evaluation of near infrared spectroscopy in
2		monitoring postoperative regional tissue oxygen saturation for fibular flaps. Journal
3		of Plastic, Reconstructive and Anaesthetic Surgery, 2008. 61(3): p. 289-296.
4	19.	Farzam, P., P. Zirak, T. Binzoni and T. Durduran, Pulsatile and steady-state
5		hemodynamics of the human patella bone by diffuse optical spectroscopy. Physiol.
6		Meas., 2013. 34 (8): p. 839-857.
7	20.	Farzam, P., C. Lindner, U.M. Weigel, et al., Noninvasive characterization of the
8		healthy human manubrium using diffuse optical spectroscopies. Physiol. Meas., 2014.
9		35 (7): p. 1469-1491.
10	21.	Hutchison, I.L., M. Cope, D.T. Delpy, et al., The investigation of osteoradionecrosis
11		of the mandible by near infared spectroscopy. Br. J. Oral Maxillofac. Surg., 1990.
12		28 (3): p. 150-154.
13	22.	Pifferi, A., A. Torricelli, P. Taroni, et al., Optical biopsy of bone tissue: a step toward
14		the diagnosis of bone pathologies. Journal of biomedical optics, 2004. 9(3): p. 474-
15		480.
16	23.	Reher, P., B.R. Chrcanovic, R. Springett and M. Harris, Near infrared spectroscopy:
17		A diagnostic tool to evaluate effects of radiotherapy in the mandible? Spectroscopy,
18		2011. 26 (1): p. 11-32.
19	24.	Siamwala, J.H., P.C. Lee, B.R. Macias and A.R. Hargens, Lower-body negative
20		pressure restores leg bone microvascular flow to supine levels during head-down tilt.
21		J. Appl. Physiol., 2015. 119(2): p. 101-109.
22	25.	Takami, Y., K. Tajima and H. Masumoto, Near-infrared spectroscopy for noninvasive
23		evaluation of chest wall ischemia immediately after left internal thoracic artery
24		harvesting. Gen. Thorac. Cardiovasc. Surg., 2008. 56(6): p. 281-287.

1	26.	Klasing, M. and J. Zange. In vivo quantitative near-infrared spectroscopy in skeletal
2		muscle and bone during rest and isometric exercise. in European Conference on
3		Biomedical Optics 2003. 2003. International Society for Optics and Photonics.
4	27.	Binzoni, T., B. Sanguinetti, D. Van de Ville, et al., Probability density function of the
5		electric field in diffuse correlation spectroscopy of human bone in vivo. Applied
6		optics, 2016. 55 (4): p. 757-762.
7	28.	Draghici, A.E., D. Potart, J.L. Hollmann, et al., Near infrared spectroscopy for
8		measuring changes in bone hemoglobin content after exercise in individuals with
9		spinal cord injury. J. Orthop. Res., 2017.
10	29.	Sekar, S.K.V., A. Dalla Mora, I. Bargigia, et al., Broadband (600–1350 nm) Time-
11		Resolved Diffuse Optical Spectrometer for Clinical Use. IEEE Journal of Selected
12		Topics in Quantum Electronics, 2016. 22(3): p. 1-9.
13	30.	Sekar, S.K.V., M. Pagliazzi, E. Negredo, et al., In Vivo, Non-Invasive
14		Characterization of Human Bone by Hybrid Broadband (600-1200 nm) Diffuse
15		Optical and Correlation Spectroscopies. PLoS One, 2016. 11(12): p. e0168426.
16	31.	Larsson, A., J. Uusijärvi, J. Näslund, et al., Bone and Soft Tissue Blood Flow during
17		Normobaric and Hyperbaric Oxygen Breathing in Healthy Divers. Journal of
18		Biomedical Science and Engineering, 2014. 7: p. 973-981.
19	32.	Mateus, J. and A.R. Hargens, Photoplethysmography for non-invasive in vivo
20		measurement of bone hemodynamics. Physiol. Meas., 2012. 33(6): p. 1027-1042.
21	33.	Mateus, J. and A.R. Hargens, Bone hemodynamic responses to changes in external
22		pressure. Bone, 2013. 52(2): p. 604-610.
23	34.	Naslund, J., J. Pettersson, T. Lundeberg, et al., Non-invasive continuous estimation of
24		blood flow changes in human patellar bone. Med. Biol. Eng. Comput., 2006. 44(6): p.
25		501-509.

1	35.	Naslund, J., M. Walden and L.G. Lindberg, Decreased pulsatile blood flow in the
2		patella in patellofemoral pain syndrome. Am. J. Sports Med., 2007. 35(10): p. 1668-
3		1673.
4	36.	Näslund, J., S. Näslund, E. Lundeberg, et al., Bone blood flow is influenced by muscle
5		contractions. Journal of Biomedical Science and Engineering, 2011. 4: p. 490-496.
6	37.	Howden, M., J.H. Siamwala and A.R. Hargens, Bone microvascular flow differs from
7		skin microvascular flow in response to head-down tilt. J. Appl. Physiol., 2017.
8		123 (4): p. 860.
9	38.	Siamwala, J.H., B.R. Macias, P.C. Lee and A.R. Hargens, Gender differences in tibial
10		microvascular flow responses to head down tilt and lower body negative pressure.
11		Physiological Reports, 2017. 5 (4) (no pagination)(e13143).
12	39.	Amarasekera, H.W., M.L. Costa, P. Foguet, et al., The blood flow to the femoral
13		head/neck junction during resurfacing arthroplasty: A comparison of two approaches
14		using Laser Doppler flowmetry. Journal of Bone and Joint Surgery - Series B, 2008.
15		90 (4): p. 442-445.
16	40.	Bahn, C.H. and G.A. Holloway, Jr., Effect of internal mammary artery mobilization
17		on sternal blood flow. Chest, 1990. 98(4): p. 878-80.
18	41.	Bassett, G.S., K.L. Barton and D.L. Skaggs, Laser Doppler flowmetry during open
19		reduction for developmental dysplasia of the hip. Clin. Orthop. Relat. Res.,
20		1997(340): p. 158-164.
21	42.	Beaule, P.E., P.A. Campbell, R. Hoke and F. Dorey, Notching of the femoral neck
22		during resurfacing arthroplasty of the hip. Journal of Bone and Joint Surgery - Series
23		B, 2006. 88 (1): p. 35-39.

1	43.	Beck, M., K.A. Siebenrock, B. Affolter, et al., Increased intraarticular pressure
2		reduces blood flow to the femoral head. Clin. Orthop. Relat. Res., 2004(424): p. 149-
3		152.
4	44.	Bogehoj, M., C. Emmeluth and S. Overgaard, Blood flow and microdialysis in the
5		human femoral head. Acta Orthop., 2007. 78(1): p. 56-62.
6	45.	Degoute, CS., MP. Preckel, C. Dubreuil, et al., Sympathetic nerve regulation of
7		cochlear blood flow during increases in blood pressure in humans. Eur. J. Appl.
8		Physiol. Occup. Physiol., 1997. 75(4): p. 326-332.
9	46.	Drinias, V., G. Granstrom and A. Tjellstrom, High age at the time of implant
10		installation is correlated with increased loss of osseointegrated implants in the
11		temporal bone. Clin. Implant Dent. Relat. Res., 2007. 9(2): p. 94-99.
12	47.	Duwelius, P.J. and A.H. Schmidt, Assessment of bone viability in patients with
13		osteomyelitis: preliminary clinical experience with laser Doppler flowmetry. J.
14		Orthop. Trauma, 1992. 6(3): p. 327-332.
15	48.	ElMaraghy, A.W., E.H. Schemitsch and J.P. Waddell, Greater trochanteric blood
16		flow during total hip arthroplasty using a posterior approach. Clin. Orthop. Relat.
17		Res., 1999(363): p. 151-157.
18	49.	ElMaraghy, A.W., E.H. Schemitsch and J.P. Waddell, Acetabular blood flow during
19		total hip arthroplasty. Can. J. Surg., 2000. 43(3): p. 197-201.
20	50.	Fukuoka, S., T. Hotokebuchi, S. Jingushi, et al., Evaluation of blood flow within the
21		subchondral bone of the femoral head: Use of the laser speckle method at surgery for
22		osteonecrosis. J. Orthop. Res., 1999. 17(1): p. 80-87.
23	51.	Green, G.E., D.G. Swistel, J. Castro, et al., Sternal blood flow during mobilization of
24		the internal thoracic arteries. Ann. Thorac. Surg., 1993. 55(4): p. 967-970.

1	52.	Hasegawa, M., G. Kawamura, H. Wakabayashi, et al., Changes to patellar blood flow
2		after minimally invasive total knee arthroplasty. Knee Surg. Sports Traumatol.
3		Arthrosc., 2009. 17(10): p. 1195-1198.
4	53.	Hempfing, A., M. Dreimann, S. Krebs, et al., Reduction of vertebral blood flow by
5		segmental vessel occlusion: an intraoperative study using laser Doppler flowmetry.
6		Spine, 2005. 30 (23): p. 2701-5.
7	54.	Hempfing, A., M. Leunig, H.P. Notzli, et al., Acetabular blood flow during Bernese
8		periacetabular osteotomy: An intraoperative study using laser Doppler flowmetry. J.
9		Orthop. Res., 2003. 21 (6): p. 1145-1150.
10	55.	Hempfing, A., R. Schoeniger, P.P. Koch, et al., Patellar blood flow during knee
11		arthroplasty surgical exposure: Intraoperative monitoring by laser Doppler
12		flowmetry. J. Orthop. Res., 2007. 25(10): p. 1389-1394.
13	56.	Hertel, R., A. Hempfing, M. Stiehler and M. Leunig, Predictors of humeral head
14		ischemia after intracapsular fracture of the proximal humerus. J. Shoulder Elbow
15		Surg., 2004. 13 (4): p. 427-33.
16	57.	Hughes, S.S., A. Cammarata, S.P. Steinmann and V.D. Pellegrini Jr, Effect of
17		standard total knee arthroplasty surgical dissection on human patellar blood flow in
18		vivo: an investigation using laser Doppler flowmetry. J. South. Orthop. Assoc., 1998.
19		7 (3): p. 198-204.
20	58.	Hupel, T.M., E.H. Schemitsch, S.A. Aksenov and J.P. Waddell, Blood flow changes
21		to the proximal femur during total hip arthroplasty. Can. J. Surg., 2000. 43(5): p. 359-
22		364.
23	59.	Kamiya, H., P. Akhyari, A. Martens, et al., Sternal microcirculation after skeletonized
24		versus pedicled harvesting of the internal thoracic artery: A randomized study. J.
25		Thorac. Cardiovasc. Surg., 2008. 135(1): p. 32-37.

1	60.	Knobloch, K., A. Lichtenberg, M. Pichlmaier, et al., Microcirculation of the Sternum
2		Following Harvesting of the Left Internal Mammary Artery. Thorac. Cardiovasc.
3		Surg., 2003. 51 (5): p. 255-259.
4	61.	Kohl, S., D.S. Evangelopoulos, M. Hartel, et al., Anterior knee pain after total knee
5		arthroplasty: Does it correlate with patellar blood flow? Knee Surg. Sports
6		Traumatol. Arthrosc., 2011. 19(9): p. 1453-1459.
7	62.	Kokovic, V., E. Krsljak, M. Andric, et al., Correlation of Bone Vascularity in the
8		Posterior Mandible and Subsequent Implant Stability: A Preliminary Study. Implant
9		Dent., 2014. 23 (2): p. 200-205.
10	63.	Kretschmer, W.B., G. Baciut, M. Baciut, et al., Changes in bone blood flow in
11		segmental LeFort I osteotomies. Oral Surgery, Oral Medicine, Oral Pathology, Oral
12		Radiology, and Endodontology, 2009. 108(2): p. 178-183.
13	64.	Kuhn, M.A., F.G. Lippert, 3rd, M.J. Phipps and C. Williams, Blood flow to the
14		metatarsal head after chevron bunionectomy. Foot Ankle Int., 2005. 26(7): p. 526-9.
15	65.	Lausten, G.S. and C.C. Arnoldi, Blood perfusion uneven in femoral head
16		osteonecrosis. Doppler flowmetry and intraosseous pressure in 12 cases. Acta
17		Orthop. Scand., 1993. 64(5): p. 533-536.
18	66.	Lausten, G.S., T. Kiaer and B. Dahl, Laser Doppler flowmetry for estimation of bone
19		blood flow: studies of reproducibility and correlation with microsphere technique. J.
20		Orthop. Res., 1993. 11(4): p. 573-80.
21	67.	Lorenzen, N.D., M. Stilling, M. Ulrich-Vinther, et al., Increased post-operative
22		ischemia in the femoral head found by microdialysis by the posterior surgical
23		approach: A randomized clinical trial comparing surgical approaches in hip
24		resurfacing arthroplasty. Arch. Orthop. Trauma Surg., 2013. 133(12): p. 1735-1745.

1	68.	Miller, J.M., E.A. Laurikainen, R.A. Grénman, et al., Epinephrine-induced changes in
2		human cochlear blood flow. Otol. Neurotol., 1994. 15(3): p. 299-306.
3	69.	Miller, J.M., TY. Ren and A.L. Nuttall, Studies of inner ear blood flow in animals
4		and human beings. Otolaryngology-Head and Neck Surgery, 1995. 112(1): p. 101-
5		113.
6	70.	Nakashima, T., T. Hattori, E. Sato, et al., Blood flow measurements in the ears of
7		patients receiving cochlear implants. Ann. Otol. Rhinol. Laryngol., 2002. 111(11): p.
8		998-1001.
9	71.	Nakashima, T., T. Hattori, M. Sone, et al., Cochlear blood flow and speech perception
10		ability in cochlear implant users. Otol. Neurotol., 2012. 33(2): p. 165-168.
11	72.	Nakashima, T., E. Sato, T. Hattori, et al., Blood flow in the ears of patients receiving
12		cochlear implants. Ann. Otol. Rhinol. Laryngol., 2004. 113(6): p. 426-430.
13	73.	Nakashima, T., M. Sone, H. Fujii, et al., Blood flow to the promontory in cochlear
14		otosclerosis. Clin. Otolaryngol., 2006. 31(2): p. 110-115.
15	74.	Nicholls, R.L., D. Green and M.S. Kuster, Patella intraosseous blood flow
16		disturbance during a medial or lateral arthrotomy in total knee arthroplasty: A laser
17		Doppler flowmetry study. Knee Surg. Sports Traumatol. Arthrosc., 2006. 14(5): p.
18		411-416.
19	75.	Nishi, H., M. Mitsuno, H. Tanaka, et al., Decreasing sternum microcirculation after
20		harvesting the internal thoracic artery. Eur. J. Cardiothorac. Surg., 2011. 40(1): p.
21		240-244.
22	76.	Notzli, H.P., K.A. Siebenrock, A. Hempfing, et al., Perfusion of the femoral head
23		during surgical dislocation of the hip. Journal of Bone and Joint Surgery - Series B,
24		2002. 84 (2): p. 300-304.

1	77.	Schoeniger, R., N. Espinosa, R.J. Sierra, et al., Role of the extraosseus blood supply in
2		osteoarthritic femoral heads? Clin. Orthop. Relat. Res., 2009. 467(9): p. 2235-2240.
3	78.	Schuurman, A.H., K.E. Bos and Y.H. Van Nus, Laser Doppler bone probe in
4		vascularized fibula transfers: a preliminary report. Microsurgery, 1987. 8(4): p. 186-
5		9.
6	79.	Selmani, I.P., H. Ishizaki, TI Marttila, Z, Cochlear Blood Flow Measurement in
7		Patients with Me´niere's Disease and Other Inner Ear Disorders. Acta Otolaryngol.,
8		2001. 121 (545): p. 10-13.
9	80.	Sone, M., T. Yoshida, H. Otake, et al., Evaluation of vascular activity in otosclerosis
10		by laser Doppler flowmetry: comparison with computed tomographic densitometry.
11		Otol. Neurotol., 2013. 34 (9): p. 1559-1563.
12	81.	Stoffel, K.K., G. Flivik, P.J. Yates and R.L. Nicholls, Intraosseous blood flow of the
13		everted or laterally-retracted patella during total knee arthroplasty. The Knee
14		Journal, 2007. 14(6): p. 434-438.
15	82.	Sugamoto, K., T. Ochi, Y. Takahashi, et al., Hemodynamic measurement in the
16		femoral head using laser Doppler. Clin. Orthop. Relat. Res., 1998(353): p. 138-147.
17	83.	Swiontkowski, M.F., Criteria for bone debridement in massive lower limb trauma.
18		Clin. Orthop. Relat. Res., 1989(243): p. 41-47.
19	84.	Swiontkowski, M.F., R. Ganz, U. Schlegel and S.M. Perren, Laser Doppler flowmetry
20		for clinical evaluation of femoral head osteonecrosis. Preliminary experience. Clin.
21		Orthop. Relat. Res., 1987(218): p. 181-5.
22	85.	Swiontkowski, M.F., K. Hagan and R.B. Shack, Adjunctive use of laser Doppler
23		flowmetry for debridement of osteomyelitis. J. Orthop. Trauma, 1989. 3(1): p. 1-5.
24	86.	Swiontkowski, M.F., D.P. Hanel, N.B. Vedder and J.R. Schwappach, A comparison of
25		short- and long-term intravenous antibiotic therapy in the postoperative management

1		of adult osteomyelitis. Journal of Bone & Joint Surgery, British Volume, 1999. 81-
2		B (6): p. 1046-1050.
3	87.	Tono, Y.U., Naoto Nagata, Atsushi Haruta, Shizuo Komune, Tetsuya, Effects of
4		trimetaphan-induced deliberate hypotension on human cochlear blood flow. Acta
5		Otolaryngol., 1998. 118(539): p. 40-43.
6	88.	Verdonck, H.W., G.J. Meijer, P. Kessler, et al., Assessment of bone vascularity in the
7		anterior mandible using laser Doppler flowmetry. Clin. Oral Implants Res., 2009.
8		20 (2): p. 140-4.
9	89.	Wannfors, K. and B. Gazelius, Blood flow in jaw bones affected by chronic
10		osteomyelitis. Br. J. Oral Maxillofac. Surg., 1991. 29(3): p. 147-153.
11	90.	Wong, K., Laser Doppler flowmetry for clinical detection of blood flow as a measure
12		of vitality in sinus bone grafts. Implant Dent., 2000. 9(2): p. 133-142.
13	91.	Ziebarth, K., M. Leunig, T. Slongo, et al., Slipped capital femoral epiphysis: Relevant
14		pathophysiological findings with open surgery hip. Clin. Orthop. Relat. Res., 2013.
15		471 (7): p. 2156-2162.
16	92.	Binzoni, T., D. Tchernin, J.N. Hyacinthe, et al., Pulsatile blood flow in human bone
17		assessed by laser-Doppler flowmetry and the interpretation of photoplethysmographic
18		signals. Physiol. Meas., 2013. 34(3): p. N25-N40.
19	93.	Sakharia, A. and M.R. Muthusekar, A comparative assessment of maxillary perfusion
20		between two different Le Fort I osteotomy techniques. Int J Oral Maxillofac Surg,
21		2015. 44 (3): p. 343-8.
22	94.	Vasovic, M., V.S. Todorovic, E. Krsljak, et al., Assessment of bone vascularity in the
23		posterior maxilla during dental implant insertion by laser doppler flowmetry.
24		Biomedical Research, 2017. 28(9): p. 4228-4232.

1	95.	Notzli, H.P., M.F. Swiontkowski, S.T. Thaxter, et al., Laser Doppler flowmetry for
2		bone blood flow measurements: helium-neon laser light attenuation and depth of
3		perfusion assessment. Journal of orthopaedic research : official publication of the
4		Orthopaedic Research Society, 1989. 7(3): p. 413-24.
5	96.	Roustit, M. and J.L. Cracowski, Non-invasive assessment of skin microvascular
6		function in humans: an insight into methods. Microcirculation, 2012. 19(1): p. 47-64.
7	97.	Griffith, J.F., D.K.W. Yeung, S.K.K. Chow, et al., Reproducibility of MR perfusion
8		and 1H spectroscopy of bone marrow. Journal of Magnetic Resonance Imaging, 2009.
9		29 (6): p. 1438-1442.
10	98.	Padhani, A.R., C. Hayes, S. Landau and M.O. Leach, Reproducibility of quantitative
11		dynamic MRI of normal human tissues. NMR in Biomedicine, 2002. 15(2): p. 143-
12		153.
13	99.	Ghosh, A., C. Elwell and M. Smith, Cerebral near-infrared spectroscopy in adults: a
14		work in progress. Anesthesia & Analgesia, 2012. 115(6): p. 1373-1383.