HIGH-RESOLUTION 3D IMAGING OF OSTEOCYTES AND COMPUTATIONAL **MODELLING IN MECHANOBIOLOGY: INSIGHTS ON BONE DEVELOPMENT, AGEING, HEALTH AND DISEASE**

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Abstract

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2 dimensional

Osteocytes are involved in mechanosensation and mechanotransduction in bone and hence, are key to bone adaptation in response to development, ageing and disease. Thus, detailed knowledge of the three-dimensional (3D) structure of the osteocyte network (ON) and the surrounding lacuno-canalicular network (LCN) is essential. Enhanced understanding of the ON&LCN will contribute to a better understanding of bone mechanics on cellular and sub-cellular scales, for instance through improved computational models of bone mechanotransduction. Until now, the location of the ON within the hard bone matrix and the sub-um dimensions of the ON&LCN have posed significant challenges for 3D imaging. This review identifies relevant microstructural phenotypes of the ON&LCN in health and disease and summarises how light microscopy, electron microscopy and X-ray imaging techniques have been used in studies of osteocyte anatomy, pathology and mechanobiology to date. In this review, we assess the requirements for ON&LCN imaging and examine the state of the art in the fields of imaging and computational modelling as well as recent advances in high-resolution 3D imaging. Suggestions for future investigations using volume electron microscopy are indicated and we present new data on the ON&LCN using serial block-face scanning electron microscopy. A correlative approach using these high-resolution 3D imaging techniques in conjunction with in silico modelling in bone mechanobiology will increase understanding of osteocyte function and, ultimately, lead to improved pathways for diagnosis and treatment of bone diseases such as osteoporosis.

Keywords: Osteocyte, 3D imaging, microscopy, biomechanics, lacuno-canalicular network, mechanobiology, mechanosensation, mechanotransduction, osteoporosis.

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CELLS MATERIALS

2D	2-dimensional
3D	3-dimensional
AFM	Atomic force microscopy
ATUM	Automatic tape-collecting
	ultramicrotome
CLSM	Confocal laser scanning microscopy
CT	Computed tomography
ECM	Extracellular matrix
EM	Electron microscopy
FE	Finite element
FSI	Fluid-structure interaction
LCN	Lacuno-canalicular network
LM	Light microscopy
LSM	Light sheet microscopy
μCT	Micro-computed tomography
ON	Osteocyte network
PCM	Pericellular matrix
PCS	Pericellular space
SBF SEM	Serial block-face scanning electron
	microscopy
SEM	Scanning electron microscopy
Serial FIB SEM	Serial focused ion beam scanning
	electron microscopy
SR CT	Synchrotron radiation-based computed
	tomography
TEM	Transmission electron microscopy
TXM	Transmission X-ray microscopy
UHVEM	Ultra-high voltage electron microscopy

Introduction

Through bone adaptation, the skeleton adapts continuously to changed mechanical loading patterns due to development, growth, ageing, disease, disuse or exercise, by removing existing and adding new bone tissue. It has been recognised that osteocytes are the key cells which orchestrate bone adaptation (Tatsumi et al., 2007). Osteocytes are ovoid cells approximately 10 µm long, surrounded by a pericellular matrix (PCM). The osteocytes and their processes form the osteocyte network (ON), which is housed within the lacuno-canalicular network (LCN), a system of voids and channels in the calcified bone matrix (Fig. 1). The presence of the osteocytes throughout the bone tissue and their interconnectedness makes osteocytes ideally placed to



sense changes in mechanical stimuli and to communicate this information through the ON. The osteocyte and the lacuno-canalicular networks (ON&LCN) are thought to be a mechanosensitive organ for external mechanical stimuli, which are translated into biochemical signals that lead to bone resorption and formation by osteoclasts and osteoblasts, respectively. As such, the ON&LCN are central in the bone remodelling process. Thus, changes in the local bone microstructure, including the ON&LCN, can alter and disrupt mechanotransduction and mechanosensation mechanisms, which are implicated in bone diseases such as osteoporosis, osteomalacia/rickets and osteopetrosis (McCreadie *et al.*, 2004; Neve *et al.*, 2012).

Knowledge of the three-dimensional (3D) structure of the ON&LCN would inform conclusions about the function and malfunction of osteocytes for different bone states in development, ageing, disease, disuse or exercise. More specifically, enhanced knowledge would improve the predictive power and accuracy of computational models, which attempt to elucidate the mechanisms of mechanotransduction using 3D geometries for the ON&LCN. Recent finite element and fluid-structure interaction models have used relatively low resolution image data and made a priori assumptions about the actual dimensions of the ON&LCN, such as the pericellular space (PCS) between the ON&LCN (Verbruggen et al., 2012; Vaughan et al., 2014; Verbruggen et al., 2014). One perspective on osteoporosis is that it is a failure of bone's adaptation to functional loading, that is, a failure of its mechanotransduction and/or mechanosensation capacity (Sharma et al., 2012; Ciani et al., 2014). Two mechanisms, direct response to matrix deformation and indirect response to fluid flow alteration have been suggested as possible pathways for mechanotransduction. To determine in detail how these mechanisms enable load transfer and signal transmission, detailed 3D mapping of the ON&LCN is needed. The fine structural details of osteocytes and their processes, the PCS and the connections between the cell and the surrounding bone matrix need to be resolved and quantified. However, structural data of the ON&LCN at cellular and sub-cellular scales of adequate quality for realistic computational modelling approaches for bone mechanotransduction, which is based on experimental data, have not been provided up to now.

This is due to both the inaccessible location of the osteocytes and their processes and to the sub- μ m dimensions. Our review will:

- 1. Identify microstructural phenotypes of the ON&LCN, relevant for bone development, ageing, health and disease.
- 2. Attempt to define the ideal imaging technique for the ON&LCN by reviewing imaging techniques used in studies of osteocyte anatomy, bone pathology and mechanobiology, including light microscopy (LM), electron microscopy (EM) and X-ray computed tomography (CT).
- 3. Examine the history and current state of the art in computational modelling of the ON&LCN and discuss the importance of high-resolution 3D imaging to this field.
- 4. Delineate how future investigations on the ON&LCN can be undertaken, taking into account recent progress in high-resolution 3D imaging.

Osteocyte imaging has been reviewed previously (Schneider *et al.*, 2010; Webster *et al.*, 2013; Kamioka, 2015). These reviews have covered 3D imaging techniques and high-resolution imaging, *in vivo* models, and the use of specialised imaging techniques for computational modelling. Investigations of the role of osteocytes in skeletal development has to date been limited. Osteocyte biology and pathology reviews include the work of Bonewald and Klein-Nulend and colleagues (Bonewald, 2013; Klein-Nulend *et al.*, 2013), while computational modelling approaches have been reviewed in (Fritton and Weinbaum, 2009).

Here we take an integrated and holistic approach, reviewing together osteocyte anatomy, mechanobiology, imaging and computational modelling, their relationships to each other and to ongoing research in these different areas. This more complete and integrated review fits with the recent dominant discussion of the relevance of the Wnt signalling pathway in bone homeostasis and sclerostin as a therapeutic agent for osteoporosis (Niedzwiedzki and Filipowska, 2015). The state of the art in 3D imaging is brought up to date with the inclusion of improved and commercially available technologies for imaging. For instance, serial block-face scanning electron microscopy facilitates concurrent imaging of hard and soft tissues and thus has the potential to further enlighten the discussion of mechanobiology and pathology.





Fig. 2. ON&LCN and typical cell machinery. (**a**) Transmission electron microscopy (TEM) image of murine bone (10-week-old C57BL/6 mouse, tibia). Bone tissue that is perfusion-fixed, decalcified, stained, and embedded in epoxy resin before thin-sectioning and imaging using TEM. Data acquired using a FEI Tecnai 12 TEM at 80 kV. Scale bar = 2 μ m. (**b**) Diagram of the ON&LCN and the typical cell machinery. Abbreviations: actin filaments (AF), endoplasmic reticulum (ER), extracellular matrix (ECM), gap junction (GJ), Golgi apparatus (GA), mitochondria (M), nucleus (N), osteocyte cell process (P), primary cilium (PC), pericellular space (PCS), tethering elements (TE).

Identification of microstructural phenotypes on cellular and sub-cellular scales of developing, ageing, healthy and diseased bone

This section will review the microstructural bone phenotypes and specifically, the structures of the ON&LCN for different bone conditions. For the purposes of this review the following nomenclature will be adopted to describe different hierarchical levels of bone structure: internal bone structure at cellular level, studied by light microscopy, including osteocyte cell bodies and osteocyte lacunae, internal bone microstructure at a level accessible by methods which overcome the diffraction limit of visible light, including osteocyte processes and canaliculi, and bone ultrastructure, including cell organelles, which can be observed by EM methods.

Anatomy and physiology of the ON&LCN

Osteocytes are at once multi-functional communicators, mechanosensors and orchestrators of bone modelling and remodelling as well as regulators of calcium and phosphate homeostasis and endocrine cells. The role of the ON&LCN as an important mechanosensory system of bone has been recognised over the last decade. Variations in the structure of the ON&LCN, such as osteocyte/ lacunar density, number of processes/canaliculi *per* cell and lacunar porosity may have profound effects on bone mechanotransduction and mechanosensation.

Microstructure of the ON&LCN and bone matrix

Lacunar number density is reported to be between 20,000 and 37,000/mm³ in healthy tissue (Carter *et al.*, 2013b; Dong *et al.*, 2014). Osteocyte number density in remodelling bone is elevated compared with inactive tissue in both healthy controls and women with hip fracture. The 2-dimensional (2D) osteocyte density was found to be 647/mm² in forming compared to 470/mm² in quiescent tissue for fracture cases and 538/mm² compared to 401/ mm² in healthy controls, indicating that osteocytes may

contribute to bone homeostasis (Power et al., 2002). Mice with targeted deletion of osteocytes are resistant to bone loss induced by unloading (Tatsumi et al., 2007), and load-related morphological and biochemical responses have been observed in osteocytes, which supports the notion that osteocytes act as mechanosensors (Zhang et al., 2006). Each osteocyte has 50-100 cell processes, that is about one million osteocytes or more per mm³ of bone tissue. Moreover, the shape and size of the osteocyte and the PCS reflect cell viability (Knothe Tate et al., 2004), while studies of rat bone have found a relationship between age and the number of canaliculi, (Okada et al., 2002). At the same time, the number and orientation of osteocyte cell processes are changed over the lifecourse (Holmbeck et al., 2005), with increased numbers of processes extending towards the vascularity than towards the mineralisation front when the cell is fully developed and enclosed in the mineralised matrix (Knothe Tate et al., 2004). Furthermore, time-lapse microscopy has been used to show that cultured osteocytes, observed during the mineralisation process, can expand and contract their cell body within their lacuna, and extend and retract their processes (Veno et al., 2007; Dallas and Bonewald, 2010). Thus, osteocytes could be considered as dynamic structures which may alter in response to external stimuli (Dallas et al., 2009; Dallas and Bonewald, 2010). Finally, changes to the shape, size and organisation of osteocytes have been observed in histological studies of pathological bone conditions (Knothe Tate et al., 2004), which will be discussed in more detail in the section regarding pathology.

It is also interesting to consider the effects of changes to the bone matrix surrounding the ON&LCN. It has been shown that the amplification of strain in osteocyte lacunae is affected by the local bone tissue properties with maximum strain relating inversely to the perilacunar tissue modulus (Nicolella *et al.*, 2006; Bonivtch *et al.*, 2007). Therefore, changes in bone matrix properties will affect osteocyte mechanosensation. A recent study has revealed that the mass density of bone matrix varies with distance





from the LCN (Hesse *et al.*, 2015), with the matrix adjacent to the LCN porosity being most dense. This supports the hypothesis that mineral exchange occurs at the interface of the LCN in bone mineralisation and mineral homeostasis.

ON&LCN at the ultrastructural level

In addition to the typical cell machinery (nucleus, mitochondria, endoplasmic reticulum, Golgi apparatus, *etc.*) the ON&LCN exhibit other ultrastructural features important for mechanobiology, which are discussed in the following and illustrated in Fig. 2.

Tethering elements. Fluid flow within the LCN imposes shear stress, which deforms the osteocyte and its processes. A transmission electron microscopy (TEM) image-based model of strain amplification proposed by Han and colleagues (Han et al., 2004) introduced the concept of tethering elements, which structurally connect osteocyte processes and the surrounding mineralised canaliculi with proteins such as CD44, laminins and proteoglycans. The model predicts that fluid flow will deform the tethering elements, creating a drag force that then imposes a hoop strain on the actin bundles in the process. It has later been expanded to include integrin attachments to conical projections on the canalicular wall, which can amplify the mechanical strain (Fig.3) (Wang et al., 2007). Forces applied focally at integrin attachment sites on cultured osteocyte-like cells initiated intercellular signals in vivo (gap junction coupling and purinergic receptor signalling) on the cell processes but not on the cell body (Wu et al., 2013).

Glycocalyx. The strain amplification model described above for the tethering elements (Han *et al.*, 2004; Wang *et al.*, 2007) suggests that only the osteocyte processes are sensitive to mechanical load, and that the cell body is insensitive to strain. Studies on single osteocytes have shown that the processes are more mechanosensitive than the cell body (Adachi *et al.*, 2009). However, studies using cultured cells, with stress applied experimentally, both to the osteocyte body and to the processes, suggest that both are mechanosensitive (Burra *et al.*, 2010). Furthermore, these studies showed that the glycocalyx – a cell membrane coating, made up of glycoprotein and polysaccharide – of the osteocyte processes (but not of the cell body) plays an essential role for bone mechanotransduction (Burra *et al.*, 2010). In particular, the glycocalyx of the osteocyte processes was found to be required for the formation of integrin attachments, which are possibly important mechanotransducers that convey mechanical signals to the osteocyte (Han *et al.*, 2004; Burra *et al.*, 2010). It has also been shown that an intact glycocalyx is necessary for certain biochemical signals to be released in response to fluid flow (Reilly *et al.*, 2003).

Cytoskeleton. The cell cytoskeleton consists of a network of filaments in the cytoplasm, including actin, which gives the cell its shape and stability. Actin filaments are present in the cell body and form the core of osteocyte processes. The cytoskeleton is crucial for the maintenance of cell shape, membrane tension and cell-cell connections (Tanaka-Kamioka *et al.*, 1998). The connection between actin filaments in the cell processes and the PCM is postulated to be important in strain transduction and amplification (You *et al.*, 2001; Han *et al.*, 2004; You *et al.*, 2004).

Gap junctions and Connexin 43. Cells communicate by direct contact and exchange of small molecules through gap junctions formed by connexons that are aligned with those on neighbouring cells. Gap junctions have been found between osteocyte processes and between osteocytes and osteoblasts (Doty, 1981), raising the possibility that they can transmit information from signals sensed by embedded osteocytes to cells at the bone surface, for instance to regulate bone formation and resorption. Additionally, fluid flow has been shown to result in opening of gap junctions in osteocyte-like cells in culture (Cheng *et al.*, 2001). It has been suggested that gap-junctional communication facilitates bone's response to load (Bonewald, 2011). However, other evidence suggests that inhibiting this communication can enhance bone formation during loading



and preserve it during unloading, which is reviewed by Lloyd and colleagues (2014). Recent studies have shown that the osteocyte gap junction channel Connexin 43 plays an integral role in bone mechanobiology and osteocyte viability (Plotkin *et al.*, 2015; Riquelme *et al.*, 2015; Xu *et al.*, 2015).

Primary cilia. Primary cilia are known to act as flow sensors in the embryo during left-right axis determination, and they are linked to mechanotransduction in polycystic kidney disease (Ong and Wheatley, 2003). In line with that, it has been suggested that osteocyte cilia may play a role in bone mechanotransduction and mechanosensation by acting as flow or strain sensors. It is postulated that primary cilia are involved in osteocyte mechanotransduction. Lee and colleagues have shown that fluid flow induces an increase in Ca²⁺ signalling (Lee et al., 2015). This hypothesis is supported by studies in mice, which show that deletion or disruption of ciliary genes leads to a disruption of mechanosensation (Xiao et al., 2011) or reduced bone formation (Qiu et al., 2012). Other studies have provided evidence that the deflection of primary cilia by fluid flow induces osteogenic and bone resorptive responses (Malone et al., 2007). However, different studies propose that cilia are only present on a small fraction of osteocytes and that they act as chemical rather than fluid flow sensors, whose function is related to mineralisation (Coughlin et al., 2015). These results conflict with studies carried out on rat tibiae, where 94 % of osteocytes expressed primary cilia (Uzbekov et al., 2012), leaving the involvement of primary cilia in mechanotransduction open to debate.

Integrins. Integrins are receptors and transducers that connect the cytoskeleton and the extracellular matrix (ECM) as transmembrane cell adhesion proteins. It has been proposed that they play a role in bone mechanotransduction and mechanosensation (Salter *et al.*, 1997; Haugh *et al.*, 2015).

Bone development

The involvement of osteocytes in bone development has been little explored thus far. Endochondral ossification involves the differentiation of mesenchymal stem cells to chondrocytes, which form a cartilaginous matrix, which is subsequently mineralised and vascularised. The blood vessels bring in osteoblasts which bind to the cartilaginous matrix and deposit bone matrix. Some matrix-producing osteoblasts on the surface of bone embed in the osteoid matrix and become osteocytes. The embedding cell develops dendritic processes and changes from a polygonal to a stellate osteocyte, retaining connections with the surface and cells already embedded (Gilbert, 2000). It has been suggested that the embedding osteoid cell and the osteocyte play roles in the mineralisation process and mineral homeostasis (Mikuni-Takagaki et al., 1995; Teti and Zallone, 2009).

Publications on the involvement of osteocytes in developing bone and its relationship to ageing and health and disease are scarce but include a study of the development of craniofacial bone in zebrafish using confocal laser scanning microscopy (CLSM) and light sheet microscopy (LSM) (Jemielita *et al.*, 2013), and further a study of the expression of sclerostin in the developing zebrafish using fluorescence microscopy (McNulty et al., 2012). Moreover, X-ray micro-computed tomography (μ CT) and histology have been used to study skeletal mineralisation in mice (Hafez et al., 2015). Finally, a study of rats at different ages, using acid-etched resin casting and scanning electron microscopy (SEM) to investigate the LCN, showed a developing canalicular system fusing with neighbouring canaliculi and round osteocyte lacunae with short and thick canaliculi (Okada et al., 2002). For acid-etched resin casting, the LCN is filled by a resin and the mineralised bone matrix is etched away by acid so that the relief cast of the LCN can be imaged using SEM after coating with heavy metals (Feng et al., 2006). Investigation and imaging of developing bone phenotypes is an area which requires further exploration in order to clarify the functional role of osteocytes.

Ageing, pathology and other bone states

Various studies have compared ON&LCN morphology, behaviour and changes in terms of age, gender, site, loading and pathology (summarised in Table 1), as outlined in the following.

Age

Lacunar number density decreases with age, by as much as 39 % (Mullender et al., 1996; Qiu et al., 2002; Billings et al., 2012). A study using human bone tissue provided evidence that this decline is associated with the accumulation of microcracks (Vashishth et al., 2000). In the same study, Vashishth and colleagues noted that ageing bone tissue is characterised by increased heterogeneity in the spatial organisation of osteocytes. A 3D study of rat bone at different ages using resin casting of the LCN and subsequent SEM and CLSM demonstrated that the internal structure of compact bone changes with age (Okada et al., 2002). Mandibles of juvenile rats had round lacunae with short, thick canaliculi, while adults had flatter, more ellipsoidal lacunae with more canaliculi, which were branched to a greater extent. On the other hand, animals of advanced age had slender, flat lacunae with fewer, smaller canaliculi. Quantitative analysis of non-human primate bone using SEM provided evidence that lacunar density was decreased, lacunar cross-sectional area was increased and the number of canaliculi per lacuna remained the same (approx. 37) with age (Billings et al., 2012). Studies have found a relationship between age and both the number and orientation of osteocyte processes and canaliculi. A roughly 30 % reduction in the number of canaliculi has been observed in elderly humans (Milovanovic et al., 2013). Conversely, Lai and co-workers have recently reported that the number density of canaliculi in mouse bone was unaffected by age, but that the dimensions of the PCS (in cortical but not in trabecular bone) increased due to an expansion of the canalicular wall (Lai et al., 2015).

Gender

Several studies suggest that there are no differences of the ON&LCN related to gender in the non-osteoporotic population (Vashishth *et al.*, 2000; Qiu *et al.*, 2002; Jordan *et al.*, 2003; Miszkiewicz, 2016). However, it has also been reported that osteocyte number density is 15 % higher in



		Osteocyte/lacuna number density	Lacunar size	Osteocyte connectivity	osteocyte orientation	Osteocyte apoptosis	Microdamage
Age	Ţ	Lacunae 729 \Leftrightarrow 604/mm ² (non-human primate, young \Leftrightarrow old) (Billings et al., 2012) ~900 \Leftrightarrow 500/mm ² (men, 16 \Leftrightarrow 73 years), ~700 \Leftrightarrow ~500/mm ² (women, 28 \Leftrightarrow 63 years) (Vashishth et al., 2000) 8.1 % less (mice, 15 > 32 weeks) (Lai et al., 2015) Up to 24 % less (women, premenopausal > postmenopausal) (Qiu et al., 2002) Osteocytes 210 \Leftrightarrow 150/mm ² (women and men pooled, 30 \Leftrightarrow 91 years) (Mullender et al., 1996) Up to 33 % less (women, premenopausal > postmenopausal) (Qiu et al., 2002)	↑ 119 ⇔ 132 μm ² lacunar area (<i>non-</i> <i>human primate</i> , young ⇔ old) (Billings et al., 2012)	= No significant difference in number of canaliculi (<i>non-human primate, young</i> ⇔ <i>old</i>) (Billings <i>et</i> <i>al.</i> , 2012)			↑ More microdamage with increasing age (human, young < old) (Vashishth et al., 2000)
Gender	=	Equal (<i>human</i>) (Vashishth <i>et al.</i> , 2000; Qiu <i>et al.</i> , 2002; Jordan <i>et al.</i> , 2003) 15 % more osteocytes (<i>healthy males < healthy females</i>) (Mullender <i>et al.</i> , 2005)					= Microdamage independent of gender (<i>human</i>) (Vashishth <i>et</i> <i>al.</i> , 2000)
Osteoporosis (OP)/Oestrogen deficiency	↑	Lacunae 17,100 \Leftrightarrow 12,900/mm ³ (women and men pooled, control \Leftrightarrow <i>OP</i>) (Mullender et al., 1996) 610 \Leftrightarrow 508/mm ² (women and men pooled, control \Leftrightarrow <i>OP</i>) (Jordan et al., 2003) Osteocytes 13,300 \Leftrightarrow 10,500/mm ³ (women and men pooled, control \Leftrightarrow <i>OP</i>) (Mullender et al., 1996) Lacunae No significant difference in lacuna number density (rat, ubia, control \ll <i>OP</i> model) (Sharma et al., 2012) Alterations in the osteocyte lacuna number density (sheep, iliac crest, control \ll <i>OP</i> model) (Zarrinkalam et al., 2012) Lacunae 206 \Leftrightarrow 134/mm ² (<i>Postmenopausal women, healthy</i> \Leftrightarrow <i>OP</i>) (Qiu et al., 2003) 21 % less (sheep, lumbar spine, control $>$ <i>OP</i> model) (Zarrinkalam et al., 2012) Osteocytes 249 \Leftrightarrow 214/mm ² (sheep, iliac crest, control \Leftrightarrow <i>OP</i> model) (Zarrinkalam et al., 2012) 271 \Leftrightarrow 223/mm ² (women, healthy \Leftrightarrow <i>OP</i>), 223 \Leftrightarrow 199/mm ² (men, healthy \Leftrightarrow <i>OP</i>), (Mullender et al., 2005) 188 \Leftrightarrow 125/mm ² (postmenopausal women, healthy $\leqslant > OP$) (Qiu et al., 2003)	 No significant difference in lacunar area (human, OP <=> control) (Mullender <i>et al.</i>, 2005) No significant difference in lacunar area (women, OP <>> control) (McCreadie <i>et al.</i>, 2004) ↓ 44.1 ⇔ 39.1 µm² (women and men pooled, healthy > OP) (Mullender <i>et al.</i>, 1996) 	J Fewer, more tortuous and slack osteocyte processes with decreased connectivity (<i>human</i>) (Knothe Tate <i>et al.</i> , 2004)	Osteocytes less oriented with respect to each other and to the blood supply (<i>human</i>) (Knothe Tate <i>et al.</i> , 2004)	↑ Increased osteocyte apoptosis between 170 % (lumbar spine) and 210 % (liac crest) (sheep, control < OP model) (Zarinkalam et al., 2012) ↑ Increased osteocyte apoptosis (human, healthy < oestrogen withdrawal) (Tomkinson et al., 1997)	
Osteomalacia			↑ Larger lacunae (mice, healthy model < osteomalacia model) (Feng et al., 2006)	= No difference in number of osteocyte processes (<i>human, healthy ⇔</i> <i>asteomalacia</i>) (Knothe Tate <i>et al.</i> , 2004)	↓ Lacunae less oriented (mice, control & asteomalacia model) (Feng et al., 2006) ↓ Osteocytes less organised (human, healthy ⇔ osteomalacia) (Knothe Tate et al., 2004)		
Osteoarthritis (OA)	↑ More lacunae (human, healthy < OA) (Jaiprakash et al., 2012) ↓ Fewer osteocytes (human, healthy > OA) (Knothe Tate et al., 2004)		↓ Smaller lacunar surface area and lacunar volume (human, osteopetrotic and osteopenic > OA) (van Hove et al., 2009)	Lower number of osteocyte processes (human, healthy > OA) (Knothe Tate et al., 2004; Jaiprakash et al., 2012)	= Orientation preserved in OA (human, healthy \Leftrightarrow OA) (Knothe Tate et al., 2004)	↑ Increased osteocyte apoptosis in OA (<i>human</i>) (Wong <i>et al.</i> , 1987)	↑ More microdamage suggested for patients with OA (human, healthy < OA) (Fazzalari et al., 1998)
Diabetes	↓ 10.1 % fewer lacunae (mice, control > diabetes model) (Lai et al., 2015)		= No significant difference in lacunar volume (<i>mice, control</i> ⇔ <i>diabetes model</i>) (Lai <i>et al.</i> , 2015)	= No significant difference in canalicular number density (<i>mice</i> , control ⇔ diabetes model) (Lai et al., 2015)			
Perlecan deficiency			↓ Smaller lacunar volume by 23.6 % for cortical bone (no significant difference for trabecular bone) (mice, control > perlecan deficiency model) (Lai et al., 2015)	↓ Fewer canaliculi by 12-16 % in trabecular and cortical bone, respectively (<i>mice</i> , <i>control</i> ⇔ <i>perlecan</i> <i>deficiency model</i>) (Lai <i>et al.</i> , 2015)			

Table 1. Anatomy of the osteocyte and the lacuno-canalicular network with ageing, pathology and other bone states.



healthy females compared to healthy males (Mullender *et al.*, 2005).

Site/Loading

Here, variations in osteocytes between bones from different sites are assumed to be due to the influence in loading direction and intensity and they are discussed together. Increased loading triggers the addition of new bone while unloading or disuse results in bone resorption. Various biochemical responses including sclerostin have been shown to be affected by loading of different frequency, timing and intensity, which are reviewed in (Bonewald, 2013).

Fibular osteocytes are reported to be elongated in shape and aligned with each other, parallel to the loading direction. On the other hand, calvarial osteocytes are more spherical or oblate and not aligned in any particular direction (Vatsa *et al.*, 2008). A study of women aged from 20-86 years, where a reduction in loading with increasing age has been assumed, found that lacunae became smaller and more spherical (Carter *et al.*, 2013a). In a synchrotron radiation-based CT (SR CT) study (Carter *et al.*, 2014) it has been observed that lacunar number density of healthy human femora varied significantly within individuals, with the anterior regions. These variations are postulated to be related to loading and bending of the femoral shaft, but the reasons are unclear (Carter *et al.*, 2014).

In summary, studies up to date suggest that lacunar number density decreases and lacunar area increases with age, but conflicting evidence exists regarding change in the numbers of canaliculi. With age, cell shape changes from round to slender and the size of the PCS increases, while the level of osteocyte organisation decreases. Studies on the influence of gender provide contradictory results, where most studies did not identify significant differences between males and females. Finally, osteocytes have been reported to align in the direction of loading and they are smaller when loading is reduced.

Pathology

Oestrogen deficiency and osteoporosis. The hormone oestrogen is essential for maintenance of the skeleton as it suppresses bone turnover and balances the formation and resorption of bone tissue, preventing bone loss (Cummings and Melton, 2002). Ovariectomy has been shown to increase bone turnover and induce bone loss (Wronski et al., 1985; Frost and Jee, 1992). Therefore, ovariectomised mice act as a model for osteoporosis, which mimics the reduction in oestrogen after the menopause, leading to an osteoporotic-like bone phenotype and related changes in bone structure and material properties, which are detrimental for bone mechanics (Frost and Jee, 1992). Studies have shown that ovariectomised rats have differences in bone matrix collagen arrangement, possibly changing the permeability of the matrix and affecting interstitial fluid flow (Sharma et al., 2012). Oestrogen deficiency has also been shown to induce apoptosis, which could alter osteocyte communication and interconnectedness (Tomkinson et al., 1997; Brennan et al., 2014a; Brennan et al., 2014b). Any or all of these changes could affect bone mechanics by

modifying bone mechanotransduction pathways and with that, signals perceived by osteocytes from strain or fluid flow, eventually triggering osteocyte biochemical outputs.

A study demonstrated higher lacunar and osteocyte number densities in osteoporotic individuals and significantly reduced lacunar area in osteoporotic individuals (Mullender *et al.*, 1996). Another study using human biopsies confirmed that osteocyte number density declines with age, where this decline occurs only in older bone and significantly in both pre- and post-menopausal women (Qiu *et al.*, 2002).

Bone cells derived from patients with osteoporosis show an impaired osteocyte response to mechanical loading (Sterck *et al.*, 1998). Osteoporotic patients have been reported to have 34 % fewer osteocytes than healthy controls (Qiu *et al.*, 2003). Overall, patients whose bone houses fewer osteocytes are more likely to sustain fracture, demonstrating their crucial role in bone maintenance (Qiu *et al.*, 2003). To date, and to the best knowledge of the authors, it is not known yet if the lower osteocyte number is a cause or rather a consequence of osteoporosis.

The proportion of occupied lacunae was found to be the same in healthy and diseased patients, indicating that the change is not the result of premature osteocyte death, but a consequence of reduced osteocyte formation (Qiu et al., 2003). Using a sheep model for osteoporosis, reduced osteocyte number density, reduced bone volume, osteoid surface area and bone formation rate were observed, while the number density of empty lacunae increased, when compared to controls (Zarrinkalam et al., 2012). Another study found that osteocyte density was lower in osteoporotic females compared to healthy females (Mullender et al., 2005). A sample of osteoporotic bone exhibited a decrease in connectivity of the ON&LCN compared to healthy bone and the orientation of osteocyte processes to each other and to the blood supply was decreased (Knothe Tate et al., 2002).

Osteoarthritis. Results based on acid-etched resin casting have shown that the osteocyte lacunar morphology is rougher and rounder in osteoarthritis compared to control samples, (Jaiprakash *et al.*, 2012). A histological investigation showed an increase in the number of nonviable osteocytes in osteoarthritic patients compared to controls (Wong *et al.*, 1987). A sample of osteoarthritic bone exhibited a decrease in connectivity of the ON&LCN compared to healthy bone, while the orientation of osteocyte processes to each other and to the blood supply was preserved (Knothe Tate *et al.*, 2002). Osteocytes in osteoarthritic human bone were found to be elongated and their lacunae smaller than those from osteopenic and osteopetrotic bone (van Hove *et al.*, 2009).

Osteomalacia. Osteomalacia (known as rickets in children) is softening of bone due to hypomineralisation, through lack of vitamin D, or overactive resorption of calcium from the bone. The LCN in osteomalacia, characterised by an acid-etched resin casting study of mouse bone, has rough surfaces on the interior of the osteocyte lacunae and fewer canaliculi (Feng *et al.*, 2006), where the surface roughness is most likely due to reduced and/ or uneven mineral content in the perilacunar matrix, which allows more resin to penetrate into the bone matrix. In ad-



dition, the osteocyte lacunae in this study were larger and organised to a lesser extent than in control bone (Feng *et al.*, 2006). In a sample of osteomalacic human bone the osteocytes appeared highly connected, but had tortuous processes (Knothe Tate *et al.*, 2002, 2004). In addition, the ON observed in the same sample was found to be less organised ('chaotic') than in other pathological states.

Sclerostin-related bone diseases. Studies of sclerosteosis and van Buchem disease, both rare high-bone mass diseases caused by mutations in the SOST gene, have led to the identification of sclerostin as a key negative regulator of bone mass (Balemans et al., 2001; Staehling-Hampton et al., 2002). Sclerostin is a glycoprotein produced by osteocytes (and chondrocytes), which is encoded by the SOST gene. Sclerostin binds to LRP5/6 co-receptors and inhibits the intracellular Wnt signalling pathway, leading to decreased bone formation by osteoblasts, reviewed by Bonewald and Johnson as well as by Klein-Nulend and co-workers (Bonewald and Johnson, 2008; Klein-Nulend et al., 2013). It is also suggested that sclerostin plays a role in the regulation of perilacunar mineralisation by osteocytes, enabling extraction of calcium from the perilacunar area when needed, followed by restoration of the mineralisation when the demand has been met (Atkins and Findlay, 2012; Kogawa et al., 2013).

Studies in rats found that treatment of osteoporotic animals with sclerostin antibody produced marked increases in bone formation, reversing the osteoporotic changes induced by ovariectomy (Li *et al.*, 2009; Ominsky *et al.*, 2014). In contrast to most other therapeutic treatments, which involve anti-resorptive agents and prevent or decelerate further bone loss, sclerostin has the potential to restore lost bone mass and strength. For example, two humanised sclerostin monoclonal antibodies have had positive effects on bone density, suggesting that neutralising sclerostin antibodies have potential as an osteoporosis treatment (Shah *et al.*, 2015a). Phase III studies are ongoing in osteoporotic patients and results are awaited (Becker, 2014).

Osteonecrosis. Osteonecrosis is characterised by missing or dead osteocytes, where necrotic bone does not have the capacity to remodel. Osteonecrosis can be caused by alcohol abuse, radiation, sickle cell anaemia and also by treatment with glucocorticoids or bisphosphonates, reviewed in (Bonewald, 2013) and (Hesse *et al.*, 2014). The mechanisms of osteonecrosis are unclear but may be linked to osteocyte apoptosis (Calder *et al.*, 2004).

Diabetes. Impaired osteoblast differentiation and activity are associated with diabetes and Type 1 diabetic osteoporosis (Motyl *et al.*, 2012). In another study CLSM and TEM has been used to examine quantitatively the topology and structure of the ON&LCN in diabetic mice (Lai *et al.*, 2015). The results for lacunar volume, lacunar spacing, canalicular number density, PCS dimensions and canalicular space dimensions were similar to healthy animals; however, the lacunar number density was reduced approximately 10 %, probably linked to osteoblast differentiation impairment. The canalicular wall area and cell process area were significantly reduced in diabetic cortical bone.

Perlecan deficiency. (Human Schwartz-Jampel syndrome) Perlecan is an essential component of the PCM that maintains an adequate fluid/solute transport pathway by possibly inhibiting mineralisation of the PCS (Farach-Carson and Carson, 2007). An investigation of the ON&LCN morphology in perlecan-deficient mice showed that the dimensions of the pericellular area between the osteocyte process and the canalicular wall decreased (Thompson *et al.*, 2011).

Imaging of the ON&LCN

Current techniques to image the ON&LCN include LM, TEM and SEM, CLSM, CT, transmission X-ray microscopy (TXM), ptychographic X-ray CT, X-ray phase tomography (holotomography)/X-ray phase nanotomography, serial focused ion beam SEM (serial FIB SEM) and atomic force microscopy (AFM). For a summary of the various imaging techniques discussed, relevant references and their advantages and disadvantages please see Table 2.

The ideal imaging method

2D images provide only a small part of the complete context of any cell. Accurate morphology including cell shape, size or location cannot be derived reliably from 2D images. Furthermore, the relationship between structures – both within and without a cell – are often invisible in 2D. Viewing and analysing structures in a 3D context avoids misinterpretation. It is however a challenge to obtain 3D data, especially at high spatial resolutions. Novel (highresolution) 3D imaging techniques are currently being developed as discussed at the end of this section on imaging of the ON&LCN and at the end of this review on the future role of high-resolution 3D imaging in osteocyte anatomy and pathology.

In addition to providing 3D data, the ideal technique for imaging bone cells and their ECM at cellular and subcellular scales would be a high-resolution, non-destructive method, capable of imaging large volumes *in situ* and *in vivo* and offering concurrent hard and soft tissue image contrast. Furthermore, the imaging technique(s) of choice would be widely available at a reasonable cost and produce quantitative hallmarks of healthy and diseased tissue, which are easy to interpret and analyse.

To meet all of these criteria compromises have to be made in practice. The ON&LCN extend throughout the bone matrix. To reveal the true degree of connectedness between contiguous osteocytes the ideal imaging method should be able to visualise a volume containing many cells. However, in the real world, different imaging techniques depend on various sample preparation protocols and there exists a broad range of experimental limitations during imaging. Most imaging techniques require tissue to be excised from the body, many require mechanical sectioning to some degree, and some of the techniques need harsh chemical treatment to render the tissue suitable for imaging, which can change the state of the native biological sample and thus, complicate interpretation of the gathered image data (Weston *et al.*, 2010).



Table	2.1	maging	techniques f	or the	assessment	of the	osteocyte	and	lacuno-o	canalicula	r network.
		00	1				~				

Technique	2D/3D	Resolution	Typical volume *	Destructive?	Comments	Reference		
LIGHT MICROSCOPY								
light microscopy (LM)	2D	200 nm	-	No	 Functional imaging through fluorescent labelling Soft tissue contrast 	Marotti et al., 1995		
High-resolution episcopic microscopy	3D	1 μm	$> 1 \text{ cm}^3$	No	• Specific labelling possible	Mohun and Weninger, 2012		
Light sheet microscopy	3D	10 µm	$> 0.5 \text{ cm}^3$	No	Live cell imaging possibleRequires optical clearing	Weber <i>et al.</i> , 2014		
Confocal laser scanning microscopy (CLSM)	3D	200 nm	0.1 mm ³	No	 Functional imaging through fluorescent labelling Soft tissue contrast Limited range of depth 	Kamioka et al., 2001		
		•	ELECTRON	MICROSCO	PY			
Scanning electron microscopy (SEM)	2D	3 nm	-	No	• Inherently 2D	Okada et al., 2002		
Serial section SEM/ automated tape-collecting ultramicrotome (ATUM) SEM	3D	2 nm (x/y) and 60- 90 nm (z)	$10^9 \ \mu m^3$	No	Complex sample preparation	Tapia <i>et al.</i> , 2012		
Transmission electron microscopy (TEM)	2D	< 1 nm	-	No	 Limited field of view Complex sample preparation 	You <i>et al.</i> , 2004		
TEM tomography and serial TEM tomography	3D	2 nm	$> 10 \ \mu m^{3}$	No	Long acquisition timesMissing wedge problem	Bonetta, 2005; Kamioka <i>et al.</i> , 2009		
Serial section TEM	3D		$> 10^3 \ \mu m^3$	No	Technically difficultTime-consuming	Harris <i>et al.</i> , 2006; Palumbo <i>et al.</i> , 1990		
Serial focused ion beam SEM (serial FIB SEM)	3D	< 3 nm	10 ⁴ µm ³	Yes	Slowfield of view	Schneider et al., 2011		
Serial block-face SEM (SBF SEM)	3D	< 10 nm	10 ⁵ µm ³	Yes	• Extensive sample preparation	Peddie and Collinson, 2014		
			X-RAY T	ECHNIQUES				
X-ray absorption CT (lab-based µCT and synchrotron radiation- based CT)	3D	< 1 µm	$> 10^9 \mu m^3$	No	 High radiation dose Simple sample preparation Relative sub-μm mineral density quantification 	(Schneider et al., 2007)		
Ptychographic X-ray CT	3D	< 100 nm	$1.7\times 10^3\mu m^3$	No	 High radiation dose Absolute sub-µm mineral density quantification 	(Dierolf <i>et al.</i> , 2010)		
Transmission X-ray microscopy (TXM) CT	3D	< 50 nm	$4.5\times 10^3\mu m^3$	No	 Chemical information High radiation dose Relative sub-µm mineral density quantification 	(Andrews et al., 2010)		
X-ray phase tomography (holotomography)/X-ray phase nanotomography	3D	120- 150 nm	$> 10^9 \mu m^3$	No	 Requires access to synchrotron radiation facility Simple sample preparation 	(Langer et al., 2012)		
SCANNING PROBE MICROSCOPY								
Atomic force microscopy (AFM)	2D	20-50 nm	-	No	 Information on mechanical properties Imaging in aqueous medium possible 	(Reilly <i>et al.</i> , 2001)		

* without tiling.

Many imaging modalities have been used in highresolution studies of the ON&LCN. Each has advantages and disadvantages. For comprehensive comparisons of these techniques see (Schneider *et al.*, 2011), Fig. 4, Table 2 and the following sections. In brief, LM, including CLSM, allows functional imaging of cells but is limited in spatial resolution by the diffraction limit of visible light (~ 200 nm). EM provides high-resolution images of cell ultrastructure, including the organelles, cytoskeletal components and membranes, but requires complex sample preparation. In addition, EM is an inherently two-dimensional (2D) technique. In contrast, X-ray CT has enabled timely imaging of large volumes of bone tissue without the need for complex sample preparation. Standard X-ray imaging techniques however lack sufficient resolution for substantial sub-micrometre imaging and do not provide enough imaging contrast for weakly X-ray absorbing soft tissues such as the ON. This limitation for soft tissues means that using standard (*i.e.*, absorption-based) X-ray imaging, only the hard mineralised bone matrix – the negative imprint of the ON – can be assessed at sufficiently high contrast levels, but not the weakly X-ray-



Resolution





Fig. 4. Summary of imaging techniques for the assessment of the ON&LCN. The various techniques are grouped regarding their dimensionality (2D or 3D) and currently achievable spatial resolution. Please see Table 2 for further details regarding the individual techniques. Abbreviations: Atomic force microscopy (AFM), confocal laser scanning microscopy (CLSM), light microscopy (LM), scanning electron microscopy (SEM), serial block-face SEM (SBF SEM), serial focused ion beam SEM (serial FIB SEM), transmission electron microscopy (TEM), transmission X-ray microscopy CT (TXM CT). Figure adapted from (Schneider *et al.*, 2011) with kind permission of Elsevier.

absorbing soft cells within the LCN. This weak vision for soft tissues reduces the informative value of such results when used for instance in computational modelling studies aimed at understanding mechanotransduction processes using cell geometries based on experimental data.

When evaluating imaging modalities used for the assessment of biological tissue, and particularly for the ON&LCN, there are many factors to consider, which are discussed in the following.

Sample preparation and method validation

Imaging tissue in its native state is an often elusive ideal. Chemical interventions to preserve tissue and/or enhance image quality, such as heavy metal staining or decalcification, can have unwanted effects such as tissue shrinkage (Buytaert et al., 2014). Where artefacts are introduced the microscopist must be aware of their impact and be able to compensate for them experimentally or post hoc during data processing. Nanophosphor particles delivered by bioballistic particle delivery systems can be used to monitor morphological changes in tissue during processing (Bushong et al., 2015). As bone is a composite material made up of a hard mineralised and a soft collagenous phase, the task of measuring artefactual change and of identifying and applying validating techniques, in order to ensure that the native tissue state is preserved and assessed, is more challenging than in homogenous tissues. Novel fixation techniques have been applied to osteocytes to optimise preservation. Acrolein, a small molecule which penetrates the tissue rapidly, has been used as fixative to improve the cell membrane preservation and reduce shrinkage (McNamara et al., 2009). Rutheniumbased fixatives and stains have also been investigated and found to enhance the preservation of cell and matrix proteoglycans (You *et al.*, 2004; Shah *et al.*, 2015b).

Validation of sample preparation and imaging techniques, to make sure that results accurately reflect the tissues and cells being examined, is important. Using more than one technique on the same sample and comparing the results can increase confidence in a novel method (Smith et al., 2006). Comparisons between the same tissues prepared for 'standard' EM and using cryo methods provide insight into changes which occur during sample preparation. Statistical methods have been used to overcome limitations of some EM data (Russo and Passmore, 2014). Beyond that, the issue of validating new 3D EM techniques has been reviewed by a task force set up to provide recommendations in 2012 (Henderson et al., 2012). It resulted in the creation of a unified data resource called EMDataBank (http://www. emdatabank.org), with the aim to develop validation reports and establish validation standards for 3D EM.

Adequate spatial resolution of microstructural and ultrastructural detail

Imaging techniques are needed that can resolve the detail of the cell processes (diameters in the order of ~ 100 nm) and ultrastructural cell details (You *et al.*, 2004).

Adequate volume to visualise cell networks and tissue matrix

For most imaging techniques there exists an inverse relationship between spatial resolution and maximum possible sample volume to be assessed, due to experimental limitations. These limitations can be found on the side of image acquisition, such as the dimension of the detector or the time available to record images at a sufficient





Fig. 5. Light microscopy (LM). A reflected polarised light micrograph of a secondary osteon in human tibia showing only the canaliculi lying in the light-reflecting plane. From (Marotti *et al.*, 1995) with kind permission from Elsevier. (**Editor's note**: Normally, eCM Journal insists on such figures showing a meaningful scale bar. As this figure is a reproduction of a previously published figure, this was not possible)

signal-to-noise ratio. Consequently, sampling smaller volumes allows working at higher spatial resolutions, while mitigating to some extent the problem of the limited penetration depth of the light and electrons used for imaging. However, small volume analysis introduces sampling errors and reduces the validity of results as the smaller number of cells examined cannot be considered to be representative (anymore) of the tissue. Due to the high tissue penetration depth of hard X-rays (energies typically > 10 keV), X-ray CT offers the largest volumes of interest of all the imaging techniques under consideration here, with whole-bone scanning being common. The volume of tissue which may be analysed using other techniques can also be limited by physical conditions such as the technical challenges of producing serial TEM sections, the restriction of the field of view in serial FIB SEM to sometimes less than 1000 µm² (Knott and Genoud, 2013; Peddie and Collinson, 2014) and the loss of the 'missing wedge' in electron tomography (Ercius et al., 2015).

3D visualistation

Elucidation of the mechanisms of mechanotransduction will require images of the ON&LCN in 3D at different levels (cellular, sub-cellular, ultrastructural). LM and CT techniques produce 3D data at low (~ 300 nm) and high (< 100 nm) spatial resolutions. While traditional EM techniques are 2D, the relatively new techniques of serial FIB SEM and serial block-face SEM (SBF SEM), produce 3D data sets by repeatedly imaging a resin block-face, as nanoscopic tissue sections are removed from the surface, either by an ion beam (serial FIB SEM) or by a remotely controlled ultramicrotome inside an SEM (SBF SEM) (Peddie and Collinson, 2014).

Light microscopy

'Standard' light microscopy (LM)

Since Hooke first observed and coined the term 'cell' in 1665, light microscopy (LM) has succeeded in elucidating details of the structure and function of bone tissue (Fig. 5). Much work has been carried out using LM on stained histological bone sections, fundamentally a 2D observation method (Marotti et al., 1995; Remaggi et al., 1998; Ferretti et al., 1999; Hirose et al., 2007). Improved optics and the advent of digital imaging have enhanced the capabilities of LM, but as with all light-based microscopy methods, the spatial resolution is restricted by the diffraction limit of visible light to around 200 nm. Additionally, as light penetration is limited in calcified tissue, decalcification may be necessary to optimise LM results. The spatial resolution of LM is insufficient to image the ON&LCN completely, as the cell process diameters are around 100 nm (You et al., 2004). Nevertheless, specialised techniques such as reflected polarised LM have been used to quantify canaliculi (Marotti et al., 1995). Highresolution episcopic microscopy is a technique designed to provide 3D histological information by consecutive imaging of a wax block as it is sectioned (Mohun and Weninger, 2012), which has not been applied to image the ON&LCN to date.

Confocal laser scanning microscopy (CLSM)

In confocal laser scanning microscopy (CLSM) a point source laser light excites tissue which is either autofluorescent (*e.g.* bone matrix through high level of collagen content) or has been stained with fluorescent dyes specific for cell components (*e.g.* actin filaments stained with phalloidin). Tissue is imaged at sequential focal planes using a pinhole detector to exclude light outside of the focal plane. A stack of 2D optical sections is acquired, which enables production of 3D representations of internal





Fig. 6. Confocal laser scanning microscopy (CLSM). A fluorescence image of a chick calvarial fragment reconstructed from 40 confocal images. Cells labelled in green are osteocytes, and cells in red are osteoblasts on the surface of the bone. The staining clearly delineates the osteocyte processes. Scale bar = 20 μ m. Adapted from (Kamioka *et al.*, 2001) with kind permission of Elsevier.

bone microstructures (Fig. 6). CLSM images have a spatial resolution of at best 200 nm in-plane and slightly worse (up to 450 nm) out-of-plane, dependent on the objective lens employed. As in standard LM, this is not sufficient to image the ON&LCN structure fully as the diameter of the osteocyte processes is in the order of 100 nm (You et al., 2004). The limitations of CLSM, when imaging in dense mineralised tissue, include the limited working distance (90-300 µm) of high-quality, high-numerical aperture objectives and the decrease in signal with depth from light absorption and scattering. The practically achievable penetration depth of CLSM in mineralised tissue is 100-150 µm (Jones et al., 2005). The difference in resolution in the x/y-direction and the z-direction can also make 3D reconstructions, quantification and interpretation of data more challenging. Finally, CLSM requires tissue to be cut, processed and cleared, thus moving it away from its native state. CLSM has been successfully used to characterise the spatial organisation, orientation and morphology of the ON&LCN (Kamioka et al., 2001; Vatsa et al., 2008; Sugawara et al., 2011; Cardoso et al., 2013; Kamel-ElSayed et al., 2015).

Live cell imaging

Time-lapse LM is used to observe and capture cellular dynamics by imaging live cells at predetermined regular time intervals in bright-field, phase-contrast and CLSM systems. A camera captures sequences of images which are later viewed at faster speed. Live cell imaging has shown that the osteocyte embedding process in living bone tissue is highly dynamic and that the cells can extend and retract their dendrites, making temporary connections with already embedded osteocytes (Dallas *et al.*, 2009; Dallas and Bonewald, 2010). In studies of unloaded rat bone with the tracers procion red and microperoxidase diffusion transport mechanisms have been examined (Knothe Tate *et al.*, 1998). Tracer injections have also been used to demonstrate the connection between the blood and canalicular fluid (Wang *et al.*, 2004).

Light sheet microscopy (LSM)

In light sheet microscopy (LSM) a sheet of laser light is used in place of the point source in CLSM (Weber *et al.*, 2014). Tissue is illuminated and imaged in slices as the laser scans through the sample. LSM provides images of intermediate spatial resolution (~ 10 μ m) from optically cleared samples up to 1 cm³ in size. LSM has so far not been used extensively for bone imaging. Studies to date include the development of bone in zebrafish embryos (Jemielita *et al.*, 2013) and simultaneous imaging of the soft and hard tissue of mouse cochlea (Buytaert *et al.*, 2013). The reduced amount of photodamage induced and the ability to image larger volumes than for CLSM are both advantages of LSM, which should lead to LSM being more widely used in future osteocyte studies.

Electron microscopy

Electron microscopy (EM) takes advantage of the shorter wavelength of electrons compared to visible light to extend resolving power. EM uses a beam of accelerated electrons to improve the theoretical spatial resolution to the atomic range with aberration corrected instruments (Pennycook *et al.*, 2015). In practice, the best spatial resolution for biological tissue is approximately 2 nm.





Fig. 7. Transmission electron microscopy (TEM) (**a**) TEM image showing an osteocyte process in longitudinal section. Transverse elements (arrows) are shown connecting the cell process to the matrix. Scale bar = 300 nm. (**b**) TEM image showing an osteocyte process in a transverse section. Dark spots (arrow) are cytoskeletal filaments, likely actin, sectioned transversally. Scale bar = 100 nm. From (You *et al.*, 2004) with kind permission of John Wiley & Sons.

Transmission electron microscopy (TEM)

In transmission electron microscopy (TEM) electromagnetic lenses focus a beam of electrons which is transmitted through an ultrathin (≤ 100 nm) section of fixed, stained, dehydrated and resin-embedded tissue, where some electrons pass through the tissue and others are scattered to form an image. This has been the favoured technique for high-resolution imaging of cell ultrastructure since its development in the 1930s. For TEM, tissue must be prepared as ultrathin sections capable of withstanding low pressure (i.e. high vacuum) and bombardment by highenergy electrons. This requires involved sample processing using chemicals and heat that can introduce artefacts such as shrinkage. Bone presents the further challenge of a hard, mineralised matrix requiring an additional step of decalcification to facilitate sectioning (An and Martin, 2003). Cryofixation is known to preserve tissue in a more life-like state but has the disadvantage that only small ($\sim 200 \ \mu m \ deep$) volumes of tissue can be preserved. The first published record of TEM bone studies dates back to 1950 (Barbour, 1950). More recently it has been used to investigate tethering fibres between osteocytes and the PCM on a nanoscopic scale (Fig. 7) (You et al., 2004; McNamara et al., 2009) and the organisation of collagen and crystals in the bone matrix (Rubin and Jasiuk, 2005).

The 2D nature of TEM imaging has been somewhat overcome by the introduction of serial section TEM, TEM tomography and ultra-high voltage EM (UHVEM). Serial section TEM is technically difficult and time-consuming but provides high-resolution results (White *et al.*, 1986; Harris *et al.*, 2006) such as those used to elucidate the 3D shape of osteocytes (Palumbo *et al.*, 1990). TEM tomography involves imaging a section repeatedly while it is incrementally tilted around its axis (Bonetta, 2005; McIntosh *et al.*, 2005). The series of projections is used to reconstruct a 3D image of the sample. One disadvantage is the 'missing wedge' caused by the limited tilt range of about 60° in most TEMs. UHVEM is not widely available and can only be used to image small samples (typically 3-5 μ m sections (Takaoka *et al.*, 2008)) due to the substantially limited penetration depth of electrons in hard tissues.

Nevertheless, UHVEM studies (Fig. 8) have produced 3D images of osteocyte processes at 16 nm resolution (Kamioka *et al.*, 2009) and portions of the canaliculi down to around 2 nm resolution, providing significant information which contradicts previous studies (Kamioka *et al.*, 2012). This is discussed further in the section of this review entitled 'The importance of high-resolution 3D imaging to computational modelling in bone mechanobiology'.

Scanning electron microscopy (SEM)

Scanning electron microscopy (SEM) employs a beam of electrons which is raster-scanned over a sample, usually fixed, dehydrated, dried and made conductive with a metal coating. The incident beam interacts with the sample producing secondary electrons, backscattered electrons and X-rays (among other signals) that are collected to form an image. SEM resolution, which can range from about 20 µm to 0.4 nm, depends on the accelerating voltage, spot size, scanning speed and vacuum level. SEM has been used to evaluate the number density of canaliculi in human bone (Marotti et al., 1995). Resin-casted and acid-etched samples have been examined using SEM, showing the distribution, size and surface details of osteocyte lacunae and the vasculature (Fig. 9) (Feng et al., 2006; Kawakami et al., 2009; Kubek et al., 2010). Although SEM images have a characteristic depth of field and convey a 3D impression, owing to the fact that SEM can produce images in focus over a range of distances, the technique is essentially 2D.



Fig. 8. Transmission electron microscopy tomography (TEM tomography). A 3D reconstruction of both the ON and the LCN based on TEM tomography data. Images (ac) show an osteocyte process in orange within a translucent canalicular wall in blue. Images (d-f) show the same views with collagen fibrils in green and red and a solid canalicular wall. Images (g-i) show the same views with a translucent canalicular wall. From (Kamioka et al, 2012) with kind permission from The Royal Society of Chemistry.



Fig. 9. Scanning electron microscopy (SEM). SEM image of an acid-etched plastic embedded section of mouse femur showing osteocyte lacunae and canalicular processes. From (Kubek *et al.*, 2010) with kind permission from Wiley (online publishing).

Volume SEM

Serial section SEM or 'array tomography' is a complex technique where multiple ribbons of serial ultrathin sections of tissue are collected on a substrate and subsequently viewed by SEM (Wacker and Schroeder, 2013). This technique is suitable for examining large areas at high *x*-*y*-resolution (*z*-resolution is limited by the section thickness). The sections can be re-examined

many times (Tapia *et al.*, 2012). Currently in development as a commercial product, the automated tape-collecting ultramicrotome (ATUM) facilitates the collection of serial sections on conductive tape for subsequent SEM imaging (Wacker *et al.*, 2015).

Both serial FIB SEM and SBF SEM produce high-resolution 3D data using automated sectioning techniques. x/y-resolution is comparable to TEM (< 5 nm), whereas the





Fig. 10. Serial focused ion beam scanning electron microscopy (serial FIB SEM). (**a**) A 3D representation of the LCN of the mid-diaphysis of a murine femur produced by serial FIB SEM. Osteocyte lacunae are shown as yellow ellipsoids and canaliculi in green. (**b**, **c**). The junction of an osteocyte lacuna and a canaliculus with the canalicular bone interface is shown in red. From (Schneider *et al.*, 2011) with kind permission from Elsevier.

z-resolution is limited by the sectioning technique, typically 10 nm for serial FIB SEM and 15 nm for SBF SEM. In serial FIB SEM a focused ion source (usually gallium) is used to mill the resin-embedded tissue by scanning over the surface. The block-face is imaged using SEM and the process repeated until the volume of interest has (partly) been milled away. A proof-of-concept study by Schneider et al. for serial FIB SEM on bone tissue has successfully imaged parts of two adjacent osteocyte lacunae (Fig. 10) (Schneider et al., 2011). SBF SEM studies of bone tissue have not been reported, but it offers a promising 3D imaging approach since initial work in the neuroscience field a decade ago (Denk and Horstmann, 2004). Using a remotely controlled ultramicrotome within an SEM, nanoscopic layers are removed from the tissue block, which is consecutively assessed by a back-scattered electron detector to extract morphologies from 3D SEM stacks with spatial resolutions comparable to TEM. SBF SEM can cover large fields of view (~ 1 mm²) and in principle, provides imaging contrast for soft and hard tissues at the same time, such as the ON&LCN, respectively.

X-ray imaging

Conventional X-ray computed tomography (CT)

Unlike light, X-rays penetrate bone easily and without causing image artefacts. Attenuation of X-rays by bone tissue provides image contrast for (2D) X-ray projections at different angular positions, which are used to reconstruct the 3D map of the specimen. Sample preparation requirements are minimal, the technique is non-destructive and once

set up, operator involvement is low. Bone can be imaged and analysed at different levels (organ, tissue, cellular) using micro-computed CT (μ CT) at isotropic resolutions down to 1 μ m and below (Müller, 2009). It is notable that CT studies do generally not provide soft tissue contrast; reconstructions depict the LCN only, which is the negative imprint of the mineralised bone tissue surrounding the ON.

Lab-based X-ray µCT. Lab-based X-ray µCT systems are commonly used to investigate bone microstructure (Müller, 2009; Draenert et al., 2012; Peyrin et al., 2014). A range of spatial resolutions from about 100 µm to below 1 µm can be reached, comparable to the resolution obtained using LM, but not reaching the resolution of EM. Thus, µCT allows visualisation of lacunae, but not the fine detail of canaliculi. Commercially available µCT systems with improved spatial resolution have recently become available, some of which claim voxel sizes in the order of 100 nm at volumes of 1 mm. Studies using this equipment have investigated differences in morphology of lacunae from different sites in the mouse (Vatsa et al., 2008) and compared morphology of lacunae in humans with different bone diseases, where significant variations have been observed between osteoarthritis, osteopenia and osteopetrosis (Vatsa et al., 2008; van Hove et al., 2009) (see also Table 1). A disadvantage of CT at these particular low voxel sizes below the micrometre range is the long acquisition time of many hours that are required for a signal-to-noise ratio that is sufficient to resolve individual osteocyte lacunae.





Fig. 11. Synchrotron radiation-based computed tomography (SR CT). SR CT images of human femur acquired at three different spatial resolutions. (a) Shows a CT slice showing osteocyte lacunae as small black dots, (b) illustrates the osteocyte lacunae (yellow) after 3D rendering and (c) shows a coloured map of the lacunae indicating their aspect ratio. (d) Displays a region of interest in the original CT slice at smaller voxel size, showing osteocyte lacunae and canaliculi, (e) a 3D rendering of the LCN (lacunae = yellow, canaliculi = cyan) and (f) a segmented 3D data set of five lacunae and their canaliculi. (g) Shows a high-resolution phase CT section with lacunae and canaliculi in black, (h) a 3D rendering an apparent texture of the collagen fibres (lacunae in orange) and (i) detailed rendering of one lacuna and its canaliculi. From (Peyrin *et al.*, 2014) with kind permission of Springer Science and Business Media.

Synchrotron radiation-based CT (SR CT). Synchrotron sources, operated as large-scale electron accelerators at a small number of sites worldwide (http://www.lightsources.org/regions), offer quasimonochromatic X-rays and much higher photon fluxes with small X-ray source spot sizes, thereby providing spatial resolutions sufficient to resolve trabecular and internal bone microstructures on a sub-µm scale (Salome *et al.*, 1999). SR CT has been used to investigate trabecular architecture and osteocyte lacunae (Fig. 11) (Peyrin *et al.*, 1998; Schneider *et al.*, 2007). In addition to X-ray absorptionbased CT, coherent synchrotron light also provides phase contrast imaging capability, where the induced phase shift by the sample is exploited in comparison to the use of X-ray absorption only, generally involving a phase retrieval step before actual CT reconstruction. Phase contrast-based CT



Fig. 12. Transmission X-ray microscopy CT (TXM CT). (a) 3D tomographic reconstruction of TXMCT data of a lacuna and surrounding canaliculi from mouse tibia trabecula. Scale bar = 5 μ m. (b,c) Reconstructed slices showing inner structure and larger processes extending from the lacuna. Scale bars = 2 μ m. From (Andrews *et al.*, 2010) with kind permission of Cambridge Journals.

Fig. 13. X-ray ptychographic CT. Osteocyte lacuna retrieved from ptychographic CT imaging of a mouse femur (mid-diaphysis). (a) Volume rendering with the bone matrix in translucent colours to show the osteocyte lacunae (L) and connecting canaliculi (C). (b) Isosurface rendering of the LCN. Long edges of 3D scale bars = 5 μ m. From (Dierolf *et al.*, 2010) with kind permission of Nature Publishing Group.



imaging yields images with improved contrast, particularly useful for weakly X-ray absorbing portions of the studied object such as collagen. Experimental SR-based imaging techniques include transmission X-ray microscopy (TXM), ptychographic X-ray CT and X-ray phase tomography (holotomography)/X-ray phase nanotomography.

Transmission X-ray microscopy (TXM). In TXM a zone plate acts as an X-ray lens to magnify the image onto the detector, which overcomes the limited spatial resolution of conventional μ CT, given by the diffraction limit of visible light around 200 nm. TXM has produced 3D reconstructions of single osteocyte lacunae and associated canaliculi at reported spatial resolutions below 50 nm, (Fig. 12), along with information regarding the composition of the surrounding bone matrix, for small fields of view (15-30 μ m²) (Andrews *et al.*, 2010). However, the canaliculi were interrupted, indicating that the provided spatial resolution of TXM needs improvement for a reliable assessment of the LCN.

Ptychographic X-ray CT. Ptychographic X-ray CT is a lensless imaging technique, which involves recording a sequence of X-ray diffraction patterns from a sample, exploiting the high phase sensitivity to produce quantitative and absolute 3D electron density maps (Dierolf *et al.*, 2010). Excellent spatial resolution, presently 16 nm (Holler *et al.*, 2014), at high contrast levels is possible, while the volume of interest is restricted to about 100 μ m³ and scanning times can be very long (hours up to days in extreme cases). In a study using X-ray ptychographic CT the LCN has been imaged at a voxel size of 65 nm, showing continuous canaliculi (Fig. 13) (Dierolf *et al.*, 2010).



Fig. 14. X-ray phase tomography (holotomography)/X-ray phase nanotomography. (**a**) 3D rendering of the LCN of human femur (middiaphysis) assessed by X-ray phase tomography (holotomography)/ X-ray phase nanotomography overlaid on a single greyscale slice. (**b**) Higher magnification of a single lacuna from (**a**) showing the branching of canaliculi. From (Langer *et al.*, 2012) with kind permission of PLoS One.



X-ray phase tomography (holotomography)/X-ray phase nanotomography. In X-ray phase tomography (holotomography)/X-ray phase nanotomography, developed as a SR imaging technique by a group operating at the European Synchrotron Radiation Facility (Langer et al., 2012), a focused monochromatic X-ray beam is used. The sample is placed behind the focal spot and imaged onto a scintillator and a camera situated at a fixed distance from the focal spot. Due to the divergence of the X-ray beam, moving the sample and recording the projections at varying distances from the focal spot varies the magnification factor. The 2D phase maps are retrieved following a holographic reconstruction approach (Cloetens et al., 1999), separating the object information from the defocused X-ray projections, which is employed for subsequent CT reconstruction to provide the 3D representation of the sample. X-ray phase nanotomography has been used by Langer and colleagues to image the LCN in 3D at a voxel size of 60 nm and a field of view of 90 µm² (Fig. 14) (Langer et al., 2012). Details of canalicular structure and branching, collagen fibre orientation and degrees of local bone mineralisation beyond microscopic scales were mapped in this study. X-ray phase nanotomography has also been applied by the same group to study the 3D orientation of mineralised collagen fibrils in human lamellar bone (Varga et al., 2013).

In summary, SR CT has been used to image and quantify the LCN, including image-based modelling approaches of bone mechanotransduction on sub-micrometre scales. Nonetheless, these newer X-ray CT imaging methods all operate for the moment at resolutions close to the dimensions of the relevant internal bone microstructures (*e.g.* osteocyte cell processes and canaliculi), and thus, care must be taken when interpreting X-ray CT image data with respect to bone mechanobiology.

Scanning probe microscopy

Atomic force microscopy (AFM) is a high-resolution scanning probe technique, which generates both images of surface topography and information on mechanical properties of tissues and materials. 2D images are recorded by scanning the surface of a sample (or a resin cast replica thereof) with a cantilever probe, where the cantilever deflection is measured with a laser. AFM can be used to image across a range of levels (tissue and cellular) and, where spatial resolutions of 20-50 nm can be reached in both air and liquid (Goldsbury et al., 2009; Usukura et al., 2012). AFM can resolve step heights of > 1 nm in crystal structures (Vavouraki et al., 2008). The ability to image in aqueous media is an important advantage of AFM, enabling closer to native-state imaging than other techniques. Additionally, it allows measurement of forces between molecules and investigation of mechanical properties (hardness, roughness) of biological materials. As the AFM does not have a lens system it does not suffer from the limitations in spatial resolution, given by the diffraction limit of visible light such as for LM or μ CT for instance. Staining or coating of the sample is not required for AFM imaging. Compared to SEM, AFM offers a small field of view (in the order of 100 μ m²) and the scan times are longer. The occurrence of thermal drift makes measurement of distances less accurate. AFM studies have imaged the LCN and the collagen structure of the matrix (Fig. 15) (Reilly et al., 2001; Lin and Xu, 2011; Katsamenis et al., 2013). In a recent study on rat femora AFM has been used to analyse the mineral phase of rat bone over the lifecourse for comparison with SEM, µCT and chemical analysis (Zhang et al., 2015). In addition, AFM has been used to investigate the crystal structure of the bone matrix (Tong et al., 2003; Kwon et al., 2009).

Novel imaging methods

Dual-purpose fluorescent and electron-dense labels for EM (SOG, miniSOG, APEX2) are genetically encoded tags for correlative light and electron microscopy (Shu *et al.*, 2011; Lam *et al.*, 2015), which could be applied to the ON&LCN. Through-focus scanning optical microscopy (Attota *et al.*, 2013), confocal soft X-ray scanning transmission microscopy (Spath *et al.*, 2015) and intravital microscopy (Masedunskas *et al.*, 2012; Alexander *et al.*, 2013) – subcellular imaging in living animals – are techniques which may add to the repertoire for osteocyte investigation.

Our group has recently adopted SBF SEM to retrieve 3D data from the ON and the LCN simultaneously. We believe that this technique will facilitate the collection of quantitative 3D information by visualising the cell morphology and internal ultrastructure, the association with the PCM and ECM and the distribution of osteocytes within the bone matrix. Fig. 16 and Supplementary video 1 contain new data obtained from mouse bone, showing individual slices from the image stack and part of an osteocyte in a 3D reconstruction.





Fig. 15. Atomic force microscopy (AFM). An AFM image of demineralised osteon in the midshaft of a bovine femur which reveals microstructural and ultrastructural features such as osteocyte lacunae, the canalicular network and lamellar interfaces. Adapted from (Katsamenis *et al.*, 2013) with kind permission of Elsevier.

Computational models

A fundamental paradox exists in bone mechanobiology, where mechanical strains applied to the whole bone are much lower than the strain levels required to cause intracellular signalling. Therefore, whole-tissue strains must be amplified to elicit a biochemical response (You *et al.*, 2000). In order to scrutinise this paradox, relevant mechanical pathways in bone mechanobiology must be understood.

Proposed mechanisms for strain amplification

Candidates for the mechanosensory system in bone were initially reviewed in the early 1990s, which included changes in whole tissue strain, hydrostatic pressure, and streaming potentials generated by fluid flow through pores in a charged matrix (Cowin *et al.*, 1991). More recently, theoretical and experimental studies have provided more evidence in favour of interstitial fluid flow between the OC and the LCN, driven by extravascular pressure and loading and also of direct cell strain, where mechanical signals are transmitted from the whole-bone to the ON as opposed to hydrostatic pressure or streaming potentials. A further possibility is the sensation of fatigue microdamage which stimulates removal and/or repair of damaged bone.

Interstitial fluid flow

In terms of fluid flow, it was proposed in the 1970s that mechanical loading-induced fluid flow can facilitate cell nutrition and waste removal (Piekarski and Munro, 1977). Early mathematical models of stress-generated fluid flow in the LCN neglected the cell process and the PCM, while empty canaliculi were assumed (Kufahl and Saha, 1990). More refined models followed, which took into account both the PCM and the osteocyte processes. These models have shown that fluid flow is driven inwards to the centre of osteons and that a molecular sieve is present in the PCS (Zeng *et al.*, 1994). This theoretical prediction was confirmed when tracer studies delineated the interstitial fluid space (Wang *et al.*, 2004). Evidence for cellular excitation and mechanotransduction by the interstitial fluid flow model reviewed in (Fritton and Weinbaum, 2009) includes theoretical models, tracer experiments and computational fluid dynamics studies.

In the mid-1990s attention was focused on interstitial fluid flow as the key mechanism for bone mechanotransduction (Weinbaum et al., 1994; Cowin et al., 1995; Klein-Nulend et al., 1995). Investigators first suggested (Cowin and Weinbaum, 1998) and then developed (You et al., 2001) a model for strain amplification due to fluid flow for the mechanical stimulation of osteocyte processes. It was hypothesised that the cell process was like a cable attached to the bone matrix by proteoglycan tethering fibres and adhesion proteins. It was assumed that the glycosaminoglycans in the PCM are attached to the fibres. Further, it was proposed that during bone deformation, the fluid passing through the PCS creates a hydrodynamic drag on the bone matrix, which in turn creates tension on the tethering fibres and leads to strain that is experienced by the osteocyte process membrane. This model predicted cellular -level strain amplifications from tenfold to hundredfold. This finding corroborated earlier predictions (Weinbaum et al., 1994) with the difference that the activating signal was the fluid flow-induced drag on the tethering fibres, and not the fluid shear stress.

TEM studies subsequently revealed an organised actin filament bundle within the process and less organised actin within the cell body (Tanaka-Kamioka *et al.*, 1998).





Fig. 16. First serial blockface scanning electron microscopy (SBF SEM) images for bone tissue. Osteocyte from mouse tibia diaphysis imaged by SBF SEM. (a-c) Individual images of section numbers 0, 40 and 80. (d,e) 3D reconstruction of 150 SEM sections of an osteocyte from mouse bone visualised using Amira software (FEI, Hillsboro, Oregon, USA). Tissue prepared according to the 'Ellisman' protocol (Deerinck et al., 2010). Data assessed using the 3View® system of Gatan Inc. (Pleasanton, USA) within an FEI Quanta 250 FEG-SEM at 2.5 kV, vacuum = 40 Pa, spot size = 3, slice thickness = 50 nm, pixel size = 22 nm. Total volume imaged = 211 μ m³. Scale bars = $2 \mu m$

Modified sample fixation techniques using ruthenium resulted in improved images of both the central actin filament bundle and the PCM (You *et al.*, 2004). Tethering elements were observed in these experiments, which had been hypothesised in the previous theoretical model (You *et al.*, 2001). Evidence was also found that the PCM fills the whole space between the osteocyte and the bone matrix.

The original strain amplification model (You *et al.*, 2001) assumed a radial arrangement of actin filaments, where the structural details of the cross-linking of the filaments were simplified. The improved ultrastructural mapping (You *et al.*, 2004) led to a structurally more refined model (Han *et al.*, 2004). Drawing on this refined model for strain amplification, the fundamental paradox in bone mechanobiology mentioned earlier (You *et al.*, 2000) could directly be addressed, by demonstrating that mechanical strains applied to the whole bone can be amplified enough to elicit cell signalling and namely, that the stiffness of the actin filament bundle is three times higher than previously predicted (You *et al.*, 2001).

None of the molecules mentioned in the studies of the original and the refined strain amplification model (You

et al., 2001; Han et al., 2004) (CD44, proteoglycans, hyaluronic acid) are known to initiate intracellular signalling. This led to an investigation into integrinbased focal adhesion complexes on the cell processes, as they have been identified as mechanical transducer sites in other cells. A TEM study of mouse bone identified conical structures protruding from the ECM, which were connected with the membrane of the osteocyte process (Wang et al., 2007) (see Fig. 3). Immunohistochemical analysis revealed punctate staining of $\alpha V\beta 3$ integrins on the cell process and β 1 integrins on the cell body but absent from the processes. Additionally, tethering fibres were co-located with the ECM projections (McNamara et al., 2009). The creation of a new strain magnification model was thus triggered, introducing the concept of integrin-based initiation of intracellular signalling (Wang et al., 2007) (see Fig. 3). The model exhibits rigid integrin attachments located at the tip of the conical structures, the tethering complex at the osteocyte process membrane surface linked to the central actin bundle by cross-linking molecules, and actin filaments that slide relative to each other. The motion of the actin and the flexibility of the



tethering elements allow local stress concentrations and large axial strains to build up in the vicinity of the more rigid integrin attachment sites when there is interstitial fluid flow through the PCM. This model leads to greatly amplified axial strains in the membrane near the integrin attachments, which are sufficient to open stretch-activated ion channels and to initiate electrical signalling. The role of $\alpha V\beta 3$ integrin attachments has been further investigated using cultured cells showing that blocking integrin function changes osteocyte morphology and disrupts the release of biochemical responses to shear stress (Haugh et al., 2015). A finite element (FE) model using focal adhesion density and stiffness measurements from AFM showed that the density of focal adhesions is related to the cell morphology and that both focal adhesion density and morphology can modulate the effect of substrate stiffness on internal cell tension (Mullen et al., 2014). It has been suggested by analytical models that the viscosity of the interstitial fluid and the permeability of the PCM significantly affect flow in the LCN (Sansalone et al., 2013). Models based on UHVEM predict that the geometry of the PCS affects the velocity of fluid flow (Kamioka et al., 2012). Fluidstructure interaction (FSI) studies take into account the elasticity and potential deformation of the cell membrane in response to fluid flow. CLSM data has been used with FSI modelling to predict the mechanical environment of osteocytes during loading and showed that stimulation is greater in the cell processes than the cell body (Verbruggen *et al.*, 2014).

Direct matrix-cell deformation

While fluid flow theory is an important concept for bone mechanotransduction, an alternative mechanism is direct matrix-cell deformation (Nicolella et al., 2005). The effect of direct bone matrix deformation has been studied by in vivo strain measurements during physical activity (Hoshaw et al., 1997; Milgrom et al., 2004). The disadvantage of these measurements is that they do not account for microstructural inhomogeneity and variations in local tissue/cell strains (Nicolella et al., 2006). Therefore, FE models have been used to quantify local tissue deformation and response to changes in the local environment (Bonivtch et al., 2007; Deligianni and Apostolopoulos, 2008). It has been proposed that the osteocyte lacuna acts as a strain amplifier and that the amplification factor is related to the properties of the surrounding bone matrix (Nicolella et al., 2006; Bonivtch et al., 2007). Using FE modelling it was found that changes in the osteocyte (body and process) stiffness had little effect on strain levels but that changes in the material properties of the surrounding bone matrix influenced the level of strain transmitted to the osteocyte, with the maximum strains resulting from decreased tissue modulus (Bonivtch et al., 2007). In a recent publication (Wang et al., 2015b) a simplified 3D FE model of the ON&LCN was used to explore the relationship between strain amplification and loading. The results from this study suggest that the strain amplification factor increases with loading frequency and increased strain.

A multilevel FE approach, using idealised LCN geometry, related loads applied to whole bone with strains at a single lacuna and canaliculi (Deligianni and

Apostolopoulos, 2008). This study revealed that osteocytes may undergo much larger strains than measured *in vivo*, which may be sufficient to elicit cell responses, suggesting that loads directly mediated through the mineralised bone matrix may produce a reaction similar to fluid flow.

Microcrack theory

Disruption or deformation of osteocytes by microcracks in the bone matrix caused by loading and fatigue has been shown to trigger firstly localised apoptosis and subsequently, targeted bone remodelling (Martin, 2000; Martin, 2002; Noble et al., 2003; Heino et al., 2009). A relationship between microdamage, osteocyte apoptosis and the initiation of bone resorption was demonstrated by showing that prevention of osteocyte apoptosis leads to complete inhibition of osteoclastic remodelling (Cardoso et al., 2009). Concomitantly, microcracks are likely to affect the fluid flow within bone and consequently, bone mechanotransduction and mechanosensation. In a study measuring microstructural strain in cortical bone based on light microscopy and digital image correlation, Nicolella and co-workers found that the presence of a microcrack caused strain amplification that is greatly increased (Nicolella et al., 2005). In a FE modelling study for a simplified Haversian system, where strain-induced interstitial fluid velocities in osteons have been evaluated, Nguyen and colleagues showed that the presence of a microcrack would reduce the fluid velocity, modifying the cellular environment and thus mechanobiological mechanisms (Nguyen et al., 2011). It is also suggested that microcracks do not cause a reduction in local signal sensation but a reduction in transduction (Ridha et al., 2015). Computational studies incorporating both mechanical strain and microdamage have been carried out. One such study has suggested that there is a damage threshold below which osteocyte processes can sense changes in strain and fluid flow, above which osteocyte signalling is disrupted and apoptosis occurs (McNamara and Prendergast, 2007). In line with that, a FE model based on ruptured osteocyte connections suggests that microcracks inhibit transmission of signals through the osteocyte network (Ridha et al., 2015).

Proposed mechanotransduction mechanisms

Mechanotransduction in the broader sense is the conversion of a mechanical signal into a biochemical response, which is transmitted to an effector cell, resulting in a coordinated response causing appropriate architectural changes. Osteocytes (processes and/or cell body) detect load via a mechanical signal, which is perceived by one or more of the mechanisms discussed above. Osteocytes respond to the mechanical signal with a range of biomolecules such as intracellular calcium signals, nitric oxide which modulates the activity of osteoblasts and osteoclasts, prostaglandins and Wnt proteins which stimulate osteoblast recruitment and activity, bone morphogenetic proteins which induce bone formation and sclerostin which inhibits bone formation reviewed in (Klein-Nulend et al., 2013). These signalling molecules modulate the recruitment and activity of osteoblasts and osteoclasts. The way how mechanical stimuli become processed and integrated by the osteocytes



and their network, and transmitted as signals to osteoblasts and osteoclasts, is suspected to depend largely on the network topology and the precise network architecture and connectivity of the ON&LCN. Moreover, in a novel perspective on the complexity of the ON&LCN, Buenzli and Sims emphasise the extent of the mechanosensory system within bone and the large number of osteocytes in the human skeleton (several billions), comparable to the numbers of neurons in the brain (Buenzli and Sims, 2015). Most likely, this allows for a large capacity of information to be processed and transmitted by the ON, which should be explicitly addressed in the future.

The importance of high-resolution 3D imaging to computational modelling in bone mechanobiology

Many computational modelling studies on bone mechanotransduction are limited by the use of idealised ON&LCN geometries, which have been shown to be inappropriate when investigating local strains (e.g. simplifying the lacuna as an ellipsoid) (McCreadie et al., 2004). Comparing the use of actual and idealised pericellular and canalicular geometries in computational fluid dynamics shows the influence of physiological geometry on force magnitudes imparted to cells through load-induced fluid flow. Idealisation resulted in underestimation of flow and shear stresses imparted by fluid drag on osteocytes (Anderson and Knothe Tate, 2008). A numerical study of murine osteocytes highlighted large differences between the results derived from idealised and from more realistic models for the ON&LCN, the PCM and ECM (Verbruggen et al., 2012). However, LCN geometry in this study was approximated based on CLSM images, which offers a spatial resolution insufficient to capture the fine details of the canaliculi. In a FSI study the same authors applied CLSM on fluorescently stained rat bone sections to create a realistic ON&LCN model, which was further informed by earlier TEM data in terms of the conical ECM projections (Verbruggen et al., 2014). This study predicted that the cell process is the most mechanically active and mechanosensitive region of the osteocyte (Verbruggen et al., 2014), confirming the results from an earlier study using an idealised model (Anderson et al., 2005). Some limitations of this study are the spatial resolution (~ 100 nm) not being adequate to detect the narrower processes, and the use of 2D TEM data to create a 3D model. Furthermore, assumptions were made about material properties of the cell and the bone matrix, the PCM was excluded and the anisotropy of the cell and the influence of the actin cytoskeleton were not reflected in the model. This does not undermine the value of the study of Verbruggen and colleagues, but rather underlines the complexity of modelling mechanotransduction processes at cellular and sub-cellular scales.

As the reliability of FE analyses is largely determined by the accuracy of the model input, accurate spatial description and quantification of the ON&LCN geometry is an essential prerequisite for more realistic models. In line with this, computational modelling of the LCN using high-resolution imaging techniques has been carried out. An LCN model for FE analysis has been derived from ptychographic X-ray CT data of mouse bone (Dierolf *et al.*, 2010; Schneider *et al.*, 2012), which provided evidence that high strain magnifications are limited to pericanalicular regions, and that FE modelling based on experimentally derived images of the LCN leads to higher strain magnification in comparison with idealised models.

A recent fluid dynamics modelling approach employed ultra-high voltage EM tomography to reconstruct a small section of a canaliculus in 3D (ignoring tethering elements and proteoglycans in the PCM) (Kamioka *et al.*, 2012). Contrary to previous work (Wang *et al.*, 2007; McNamara *et al.*, 2009), no conical projections from the matrix wall were found (based on a single sample). In addition, the study of Kamioka and co-workers showed that variations in the roughness of the canalicular wall influenced the loading-induced flow of the interstitial fluid. One conclusion of this study was that fluid flow simulations based on 2D data may lead to inaccurate findings about mechanotransduction mechanisms.

In a recent FE study human bone was imaged using synchrotron X-ray phase nanotomography at a voxel size of 50 nm (Varga et al., 2014). The LCN morphology was quantified and the osteocyte and processes artificially reconstructed using a priori assumptions from TEM and CLSM studies to approximate cell shape, canalicular occupation as well as PCM and osteocyte process dimensions. FE models were generated, which were subjected to uniaxial compression, confirming the existence of strain magnification through direct matrix-cell mechanotransduction. Furthermore, it was reported in the same study that cell deformations were localised at the cell-process junctions. These results confirm findings of previous investigations (Bonivtch et al., 2007; Deligianni and Apostolopoulos, 2008). A limitation of this study was the lack of original imaging data describing the cell shape.

In conclusion, reliable quantitative characterisation of both the ON and the LCN based on advanced 3D imaging across different length scales will inform computational models of bone mechanotransduction and will allow realistic predictions of this complex dynamic system and its emergent behaviours (Buenzli and Sims, 2015). For example, data on canalicular occupancy and the number of cells, which make connections with vascular and other structures, would improve accuracy when calculating the number of connections between osteocytes and related cell networks.

The future role of high-resolution 3D imaging in osteocyte anatomy and pathology

Much research has been carried out into microcracks, loadinduced fluid flow and direct strain from the matrix to the osteocyte and their posited roles in mechanotransduction and mechanosensation ; however, many issues still remain to be discovered and thoroughly investigated. It is necessary to clarify the role of (ultra)structural differences between the cell processes and the cell body of osteocytes in sensation of mechanical stimuli and in the initiation of cell signalling. The involvement of primary cilia is unclear and it is not known whether it is crucial for bone mechanosensation. The co-localisation of integrins with



Table 3. Choice of imaging technique for the assessment of the osteocyte and lacuno-canalicular network. Abbreviations: Transmission electron microscopy (TEM), micro-computed tomography (μ CT), confocal laser scanning microscopy (CLSM), serial focused ion beam scanning electron microscopy (serial FIB SEM), serial block face scanning electron microscopy (SBF SEM).

Choice of imaging technique 3D		High resolution	Automated	Hard and soft tissue contrast	Volume of interest
Serial section TEM	yes	yes	no	yes	$> 10^{3} \mu m^{3}$
μCT	yes	intermediate	yes	no	$> 10^{9} \mu m^{3}$
CLSM	yes	no	no	yes	0.1 mm ³
X-ray ptychographic CT	yes	intermediate	yes	no	$> 10^{3} \mu m^{3}$
Serial FIB SEM	yes	yes	yes	yes	$10^4\mu m^3$
SBF SEM	yes	yes	yes	yes	$10^{5} \mu m^{3}$

the protrusions, connecting the cell membrane and the mineralised bone matrix, also requires further investigation.

Data from recent research, where new imaging techniques have been employed, were used by Buenzli and Sims (Buenzli and Sims, 2015) for theoretical mathematical calculations of the dimensions of the entire ON and LCN in an average human adult skeleton. This has led to insights into the extent of the influence of osteocytic osteolysis on bone and its potential to alter the structural properties and permeability of the ON&LCN. The use of cutting-edge techniques, such as UHVEM, has raised questions about the impression produced by 2D imaging that osteocyte processes are directly attached to the canalicular walls (Kamioka *et al.*, 2012) although the small field of view for UHVEM data has to be taken into account for such considerations.

Improvements will help to validate theoretical and computational models for strain magnification as well as quantifying bone interstitial fluid movement in response to mechanical loading. Questions to be addressed include: What is the 3D structure of the ON&LCN at different sites of an entire bone sample? How are the ON and the LCN and thus, the interstitial fluid flow altered in disease? To answer these questions, existing 3D imaging techniques need refinement and further development, and novel highresolution 3D imaging techniques will be applied in the field of bone research.

Refinements and developments of high-resolution 3D imaging techniques

The features of a number of currently available 3D imaging techniques are set out in Table 3, showing the relative strengths of each as an imaging approach for the ON&LCN. The number of potentially useful techniques is rising however, as imaging techniques traditionally applied in physics and materials science are increasingly applied to address biological and medical questions. Some recent developments include the improved resolution of CT techniques using synchrotron sources or the development of cryo-TEM (Weston et al., 2010; Ercius et al., 2015), which facilitates high-resolution imaging of samples in a near-to-native state. Cryo-soft X-ray tomography is a promising technique for close-to-native state imaging (Carzaniga et al., 2014; Wang et al., 2015a). Improvements in the sensitivity of EM detectors used in SBF SEM may reduce the amount of specimen preparation, needed and beam deceleration could allow imaging at lower energies, and therefore causing less specimen damage. Furthermore, conductive resins would improve the quality of SBF SEM data quality and it may even be possible to carry out SBF SEM on samples up to 10 mm in width (Marx, 2013; Ellisman, 2015). New refinements of super-resolution LM are breaking the diffraction limit of visible light (Betzig, 2015; Wang *et al.*, 2015a). Other recent LM improvements include better depth penetration and reduction of light scattering (Wang *et al.*, 2015a). The development of multibeam SEM with more than 60 parallel beams will greatly accelerate the rate of data collection (Marx, 2013). A multienergy deconvolution SEM is currently in development which can perform 'sub-surface' imaging using different energy beams, which improves *z*-resolution and reduces the risk of missing fine structures (Marx, 2013).

Techniques previously confined to the realm of engineering are increasingly being applied to biological systems and pathology research (Friedl *et al.*, 2012). For example, electrochemical strain microscopy (a variation of AFM) captures dynamics in nanometre scale volumes (Jesse *et al.*, 2012) and ferroelectric switching has been observed in ECM components (Liu *et al.*, 2014).

Conclusions and outlook

At present, high-resolution 3D native-state imaging of the ON&LCN is impossible. Achieving the goal of concurrent imaging of both hard and soft tissue networks would undoubtedly have a significant impact on the understanding of bone adaptation and pathology processes. Many new high-resolution 3D imaging techniques are advancing the ability to achieve this aim incrementally.

Given that mechanotransduction mechanisms act across scales (from the whole-bone down to the cell ultrastructure) bone tissue must be assessed at different hierarchical levels (Lucas *et al.*, 2012). No technique exists that can cover so many different scales at the same time, due to physical limitations. Even to image the different features of the ON&LCN in a restricted field of view, several length scales must be covered, from the osteocyte and the lacuna (~ 10 µm) down to the cell processes and canaliculi (< 1 µm), the PCS (< 100 nm), *etc.* A correlative imaging approach combining light, X-ray and electron microscopy modalities might prove to be useful. Large volumes could be analysed and high-resolution quantitative information obtained and used for realistic *in silico* simulations.



Comparison of 3D quantitative hallmarks of osteocytes in healthy and pathological tissue would enhance the current knowledge of mechanotransduction processes. The wealth of current data on bone physiology and the ON and LCN indicate the importance of a multi- and inter-disciplinary approach, combining the expertise and differing experimental methodologies of life scientists, microscopists, engineers and mathematicians.

We have identified relevant phenotypes of developing, ageing, healthy and diseased bone on cellular, sub-cellular and ultrastructural levels. Techniques used for ON&LCN imaging have been discussed and assessed, including their relevance for computational modelling approaches. In conclusion, we have considered how developments in high-resolution 3D imaging technology and novel imaging approaches may contribute to the study of osteocyte structure. First attempts to visualise the ON&LCN using SBF SEM have been reported. As high-resolution 3D imaging techniques mature, become more widely available and are used to image bone from different species, ages, and (pathological) states, the nature of the elusive role of the osteocyte for bone development, ageing, health and disease will become clearer in the future.

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Discussion with Reviewers

Peter Varga: Authors mention the importance of validation. I completely agree that this is an important, but at the same time very challenging step. In your view, how could these models be validated? What would be a meaningful and doable approach?

Author: Validation of sample preparation and imaging techniques is important, to make sure that results accurately reflect the tissues and cells being examined. Comparing tissue prepared for 'standard' electron microscopy (EM) and cryo-EM (Dubochet et al., 1988) would provide insight into changes that occur during sample preparation. Conventional EM preparation involves aldehyde fixation, a combination of heavy metal stains, dehydration, and resin embedding. As water is the main constituent of biological material and dehydration leads to material aggregation and consequent loss of structural information (Dubochet and Sartori Blanc, 2001), maintaining cells in their hydrated state is an advantage. Cryofixation by plunge or slam freezing removes heat quickly, forming vitreous ice, which minimises the amount of crystal damage and immobilises water molecules within milliseconds. The limited depth of vitrification (several µm) makes the technique eminently suitable for small particles and thin parts of cells, but less good for tissue samples. A better alternative for bone tissue would be cryo-electron microscopy of vitreous sections (CEMOVIS) (Al-Amoudi et al., 2004) which - when combined with high-pressure freezing - eliminates the effects of chemical fixation, staining, and dehydration. CEMOVIS gives close-to-native-state results at excellent spatial resolutions (Bouchet-Marquis and Fakan, 2009). It must be remembered however, that cryo-EM methods can introduce cutting artefacts such as compression, and that the act of imaging with an electron beam can also affect the samples by thinning or 'bubbling' reviewed in (Mielanczyk et al. (2014).

Peter Varga: The general problem with osteocyte imaging is still the lack of data. At the same time, we need proper statistical description on this cell network. You mention the need for comparing healthy and pathological conditions. Would you agree with the statement that we first would need to identify the normal variation of these features in healthy condition (sex, age, tissue age, anatomical site, loading condition, *etc.*)?

Author: We agree that it is important to understand bone anatomy on different hierarchical levels during development and ageing, as well as in health and disease. An initial approach, using novel three-dimensional (3D) highresolution imaging techniques for the assessment of the osteocyte network and the surrounding lacuno-canalicular network, should encompass comparisons between healthy and pathological conditions for different anatomical sites, thus elucidating inter-individual and intra-individual variability. A significant amount of work has already been published to identify and quantify normal variation of anatomical features of bone at different hierarchical levels, thus a meta-analysis of these investigations would help to consolidate the current knowledge and strengthen the evidence collected, specifically in terms of the osteocyte and the lacuno-canalicular network. Standardised measures would facilitate these comparisons, and the collection of data would feed into computational models in the context of mechanobiology for instance. To date, most studies have been based on small numbers of samples. We suggest that the pre-clinical and clinical studies on sclerostin (Clarke, 2014) are a potential source of significant amounts of data and tissue samples for validation studies. Large numbers of human and animal subjects have been treated with antisclerostin antibodies (romosozumab and blosozumab), which have been found to increase bone formation and bone mineral density. We believe that analysis of this cohort would provide much valuable data.

Pascal Buenzli: The authors emphasise the role of the osteocyte network anatomy for the network's mechanosensing and mechanotransducing capabilities. Comparative tables such as Table 1 are invaluable tools to help understand this role. However, being still in an exploratory phase, the field has no accepted standard for measuring ON&LCN properties. Could the authors briefly discuss what network properties they believe future studies should aim to report in order to start building a standard and ease comparative studies?

Author: Studies of network characteristics in the brain have related modifications in Alzheimer's disease and schizophrenia (Bullmore and Sporns, 2009). As in principle, it may be possible to relate changes in the properties of the osteocyte and the lacuno-canalicular network (ON&LCN) to different pathological bone conditions, it is crucial therefore how (3D) spatial characteristics are subsequently quantified. To create a standard for measuring ON&LCN properties, which will facilitate comparative studies, a consistent and well-defined set of quantitative measures for



describing the ON&LCN is needed, which is independent of the employed imaging methods and the experimental conditions that are required. The quantification should ideally be user-independent and automated to ensure consistency and efficiency, although some human interpretation and intervention will presumably remain necessary. There will always be a compromise between field of view and spatial resolution as well as acquisition time, data volume, and related financial implications.

A framework has been proposed for the quantitative characterisation of osteocyte lacunae and their 3D spatial relationships (Mader et al., 2013), which included shape, orientation, number density, spatial distribution, and alignment. This framework is a valuable first step and should be expanded to encompass finer cellular detail such as cell processes and ultrastructure. In a study, where the osteocyte network has been related to the surrounding bone mineral quality, measures related to connectivity and spatial distribution were compared for the ON, including canalicular details (Kerschnitzki et al., 2013). Dong and colleagues used automated quantification to extract 3D shape descriptors from synchrotron radiation-based microcomputed tomography data (Dong et al., 2014). Combining the measures in these studies and including ultrastructural detail would lead to a comprehensive standard data set. In summary, we believe that a quantitative framework should include, although not be limited to, detail such as; osteocyte number density, spatial distribution and alignment, microstructural information including cell, peri-cellular space and lacunar shape and volume as well as ultrastructural detail such as process and canaliculus dimensions and tortuosity.

BoneJ, an ImageJ (Schneider *et al.*, 2012) plug-in designed specifically to provide standard measures for bone image analysis (Doube *et al.*, 2010) in an easily accessible and open-source approach, has provided a significant contribution to the field.

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