Exercise is a non-pharmacological intervention that can enhance bone regeneration and improve the management of bone conditions like osteoporosis or metastatic bone cancer. Therefore, it is gaining increasing importance in an emerging area of regenerative medicine—regenerative rehabilitation (RR). Osteocytes are mechanosensitive and secretory bone cells that orchestrate bone anabolism and hence postulated to be an attractive target of regenerative exercise interventions. However, the human osteocyte signalling pathways and processes evoked upon exercise remain to be fully identified. Making use of a computer-controlled bioreactor that mimics exercise and the latest omics approaches, RNA sequencing (RNA-seq) and tandem liquid chromatography-mass spectrometry (LC-MS), we mapped the transcriptome and secretome of mechanically stretched human osteocytic cells. We discovered that a single bout of cyclic stretch activated network processes and signalling pathways likely to modulate bone regeneration and cancer. Furthermore, a comparison between the transcriptome and secretome of stretched human and mouse osteocytic cells revealed dissimilar results, despite both species sharing evolutionarily conserved signalling pathways. These findings suggest that osteocytes can be targeted by exercise-driven RR protocols aiming to modulate bone regeneration or metastatic bone cancer.
is well known that different species display different molecular networks linking genes to diseases. This discrepancy led us to postulate that a portion of the signalling pathways and processes in osteocyte mechanotransduction may not be conserved across both species. Thus, another important aim of this work is to ascertain whether human and mouse osteocytic cells show differences from a transcriptome and secretome standpoint. By identifying potential differences, we will be able to make more informed decisions upon using human or mouse cells in basic and preclinical bone studies.

The transcriptome was mapped by RNA-seq which detected ~13,000 gene products in the murine cell line MLO-Y4 and ~14,000 in human stem cell-derived osteocytes (hSCO). The percentage of clean raw reads was homogenous, above 98% for every biological replicate (Supplementary Fig. 1). The hSCO showed 368 gene products exclusive to mechanically active and 619 to static (Fig. 1a). MLO-Y4 exhibited 286 gene products exclusive to mechanically active conditions and 242 exclusive to static (Fig. 1b). Deseq2 analysis revealed 821 differentially expressed gene transcripts in human and 13 in murine (Supplementary Fig. 2). Differentially expressed gene transcripts were analysed using the GO and the KEGG enrichment analysis. In stretched human cells, organelle, nucleus and intracellular were the most overrepresented biological processes while in stretched murine cells, transport, immune system process and extracellular region were the most enriched (Fig. 1c, d). Cell cycle, FoxO, pathways in cancer and DNA repair related pathways were the most significantly enriched signalling pathways in mechanically stretched hSCO whereas TNF, cytokine–cytokine interactions, PI3K-AKT, Jak-STAT, HIF-1 were mostly enriched in murine cells (Fig. 1e, f). Next, we profiled the protein secretome using tandem LC-MSMS. hSCO revealed 36 proteins exclusive to cyclic stretch and 95 proteins exclusive to static conditions (Fig. 2a) whereas murine cells revealed 124 proteins exclusive to mechanical loading and 166 exclusive to static conditions (Fig. 2b). The relations between proteins found exclusively under mechanically active and static conditions are depicted in Supplementary Figs. 3 (human) and 4 (mouse). The cellular location of all identified proteins was found similar under mechanically active and static conditions (Supplementary Fig. 5). Quantitative sequential wind- owed acquisition of all theoretical fragment ion mass spectra (SWATH-MS) analysis of mechanically stimulated cells revealed differential expression of 15 out of 379 secreted factors in hSCO (Fig. 2c) and 12 out of 841 in MLO-Y4 (Fig. 2d). GO enrichment analysis suggests that cholesterol or lipoprotein-related was the most significant biological process found in human cells (Fig. 2e) whereas murine cells revealed 124 proteins exclusive to mechanical loading and 166 exclusive to static conditions (Fig. 2b). The relations between proteins found exclusively under mechanically active and static conditions are depicted in Supplementary Figs. 3 (human) and 4 (mouse). The cellular location of all identified proteins was found similar under mechanically active and static conditions (Supplementary Fig. 5).

Quantitative sequential windowed acquisition of all theoretical fragment ion mass spectra (SWATH-MS) analysis of mechanically stimulated cells revealed differential expression of 15 out of 379 secreted factors in hSCO (Fig. 2c) and 12 out of 841 in MLO-Y4 (Fig. 2d). GO enrichment analysis suggests that cholesterol or lipoprotein-related was the most significant biological process found in human cells (Fig. 2e) whereas murine cells revealed 124 proteins exclusive to mechanical loading and 166 exclusive to static conditions (Fig. 2b). The relations between proteins found exclusively under mechanically active and static conditions are depicted in Supplementary Figs. 3 (human) and 4 (mouse). The cellular location of all identified proteins was found similar under mechanically active and static conditions (Supplementary Fig. 5).

We successfully mapped the mechanosensitive gene transcripts and proteins secreted by human osteocytic cells after a single bout of cyclic stretch using RNA-seq and LC-MSMS, respectively. This allowed us to identify overrepresented signalling pathways, biological processes and network processes evoked by mechanically stimulated osteocyte-like cells likely to mediate bone regeneration or modulate cancer progression. As mentioned earlier, mechanically stimulated osteocytes can influence bone regeneration through indirect bone remodelling which involves activation of PI3K/AKT and WNT signalling pathways, aiming to target downstream anabolic action provided by osteoblasts. Our results suggest that in addition to this, osteocytes can modulate bone regeneration via direct remodelling of their extracellular environment since the network process ossification and bone remodelling was significantly enriched. This direct remodelling capacity is corroborated by a previous study on lactating mice demonstrating that osteocytes removed the ECM to potentially provide an extra source of calcium to the body.

Exercise or mechanical stimulation of osteocytes appears to offer a protective effect against breast cancer progression, however, the underpinning mechanisms remain unclear. We hypothesised that activation of signalling pathways linked to genome integrity could be one potential mechanism. This study confirmed our hypothesis since the signalling pathways cell cycle, P53, pathways in cancer and DNA-repair like mismatch repair and nucleotide excision repair, were all significantly enriched in stretched osteocytic cells. Another proposed mechanism by which mechanically stretched osteocytes may limit cancer progression is remodelling of the extracellular microenvironment, since ECM remodelling, ossification and bone remodelling and cell–matrix interactions were significantly enriched as mentioned earlier. Remodelling is likely to have an impact on this because cancer cells degrade the normal ECM to replace it with tumour ECM allowing the invasion of the cancer cell. As mechanically stimulated osteocytic human cells remodel their extracellular environment this should compete against breakdown and deposition of tumour ECM and delay or avoid cancer invasion. Although host–tumour interactions are gaining increasing attention in exercise and cancer, functional studies involving cocultures of mechanically stimulated osteocytic cells with cancer cells would be needed to determine their contribution to cancer progression limitation.

A comparative analysis across human and mouse osteocytic cell lines evidenced that the transcriptome and secretome were quite distinctive between the two cell lines. KEGG analysis of the human transcriptome identified as potential mechanosensitive signalling pathways, cell cycle, pathways in cancer, P53, DNA repair related pathways and FoxO, whereas examination of the mouse identified TNF, cytokine–cytokine receptor interactions and PI3K-AKT. As for secretome, two network processes were significantly enriched out of the top ten in both cell lines—ossification and bone remodelling and manganese transport; however, the proteins leading to such enrichment were different. We speculate that variations found between both osteocytic cell lines are because of fundamental differences between the two—one is human and derives from a stem cell model and the other is a mouse immortalised transgenic cell line.

Genetic activation of osteocyte WNT/β-catenin is vital for bone healing. In mouse, WNT/β-catenin activation typically occurs within 1–24 h post-loading (at rest) as demonstrated by β-galactosidase positive staining and increased gene expression of Wnts 1 and 7 and 14,15. In the osteocytic MLO-Y4 cell line, gene expression of WNT1-related gene Wnt3a is upregulated at 1–3 h post-stimulation with pulsatile FFSS (at rest)46. To ascertain whether cold shock was enough to activate WNT/β-catenin signalling in human or mouse osteocytic cells, the relative expression of 84 key genes of the WNT/β-catenin signalling was analysed by RT-qPCR.
RT-qPCR showed that WNT/β-catenin gene expression remained unchanged. A closer analysis of RNA-seq and LC-MSMS indicated potential signs of activation, since the β-catenin target, TCF7 was slightly upregulated in human osteocytic cells and the signalling pathways PI3K-AKT which crosstalk with WNT/β-catenin was enriched in the mouse cells. However, we were expecting to detect a more robust response in preparation of the above events related to WNT/β-catenin activation, in particular, the significant enrichment of the WNT/β-catenin signalling pathway. Given that we analysed WNT/β-catenin immediate after cyclic stretch and did not.
not found a robust response and other studies examining 1–3 up to 24 h post-loading (at rest) did, we postulate that a rest period after mechanical stimulation is fundamental to trigger strong activation of WNT/β-catenin signalling in osteocytes.

To summarise, this study suggests that mechanically stimulated osteocytes support bone regeneration via ossification and ECM remodelling and modulate cancer through transcriptional activity linked to genome integrity and ECM remodelling. Transcriptomic

**Fig. 2 Secretome analysis of murine and human osteocytic cells.** Venn diagram of hSCO (a) and MLO-Y4 (b) showing the secreted molecules exclusive to mechanically active and static cell culture conditions; Protein expression fold change given for the stretch condition vs control group (static) in hSCO (c) and MLO-Y4 (d); Chart depicting top ten overrepresented biological processes in hSCO (e) and MLO-Y4 (f); Chart depicting top ten overrepresented process networks in hSCO (g) and MLO-Y4 (h); Number of overexpressed proteins in each enrichment score within brackets; n = 4.
and secretomic data show dissimilar results between human and mouse cell lines in response to the same cyclic stretch regimen suggesting that mouse cells could be suboptimal predictors of human response to non-pharmacological interventions like exercise and should only be used when the signalling pathways or processes to be investigated are highly evolutionary conserved.

Despite the importance of these findings, we do recognise the importance of repeated bouts of mechanical loading and the performance of functional studies involving stretched osteocytes and cancer cells. Therefore, future studies addressing both aspects are warranted. As our understanding of the transcriptome and secretome in stretched cells and human bone cell models progresses, RR protocols based on exercise may be efficiently translated to the clinic.

METHODS

Cell culture

Human osteocyte-like cells derived from a stem cell culture model: Adipose stem cells (ASCs) were purchased from ATCC (ATCC® PCS-500-011® and expanded in complete growth medium (ATCC® PCS-500-030® and ATCC® PCS-500-040®) supplemented with 1% penicillin-streptomycin until reaching 70–80% confluence and passed to a 6-well plate at a cell density of 18,000 viable cells/cm². After 48 h, the complete growth medium was removed, cells were washed twice with PBS and differentiation medium (ATCC® PCS-500-052®) was added. The differentiation medium was changed every 3–4 days. ASCs differentiated between 13 and 15 days. Osteocyte-like cells derived from ASCs stained positive for Alizarin red, FGF23 and exhibited dendritic-like extensions like primary osteocytes.

MLO-Y4: This cell line was acquired from Kerafast (EK0002) and expanded in MEM-a (Fisher) supplemented with 2.5% foetal calf serum (FBS) and 2.5% newborn calf serum (NCS) both inactivated (Fisher) and 1% penicillin-streptomycin until reaching 70% confluence. The medium was changed every 2–3 days. Both cell lines were cultured in a humified incubator at 37 °C and 5% CO₂.

Mechanical loading

Multiple studies reported that a single bout of exercise is enough to observe positive regenerative and health benefits, e.g. skeletal muscle satellite cells numbers increase and vascular function is normalised and therefore we opted for a single bout of mechanical loading rather than a multi bout one.

Upon exercise, bone experiences a myriad of forces, tension, compression, torsion and FFSS. A study comparing FFSS with cyclic tension demonstrates that human-derived bone cells react differently to both stimuli and that tension is a more potent stimulus for bone formation than FFSS as evidenced by the increased production of the type I collagen. The anabolic action of tension and its importance in RRPs, we decided to observe positive regenerative and health beneﬁts, e.g. skeletal muscle satellite cells numbers increase and vascular function is normalised and therefore we opted for a single bout of mechanical loading rather than a multi bout one.

RNA extraction

Immediately after the mechanical loading, cells were washed twice with ice-cold PBS and lysed with RLT buffer (Qiagen) for 1 min. Next, RNA was extracted using the RNeasy kit (Qiagen) according to the manufacturer instructions. RNA quality and quantity were estimated using the NanoDrop™ 2000 spectrophotometer (Thermo Scientific™).

cDNA synthesis and quantitative gene expression

Gene expression was analysed by RT-qPCR. After RNA extraction, cDNA was synthesised using the RT® First Strand Kit (Qiagen) according to the manufacturer instructions. As for the RT-qPCR, the RT2 Profiler® PCR Array Human WNT Signalling Pathway PAHS-043Y (Qiagen) and the RT® Profiler PCR Array Mouse WNT Signalling Pathway PAMM-043Z (Qiagen) were used to quantify relative expression genes of the WNT signalling pathway. RT-qPCR was performed in the Rotorgene 6000 thermocycler (Corbett Research) using the SYBR Green ROX master mix (Qiagen) following the manufacturer instructions. CT values were exported to an Excel file to create a table of CT values and this table uploaded to the data analysis web portal at http://www.qiagen.com/geneglobe. CT values were normalised based on an automatic selection from a full panel of reference genes. Fold-Change (2^(-ΔΔCT)) was calculated based on the normalised gene expression (2^(-ΔΔCT)) in the mechanically stimulated samples divided by the normalised gene expression (2^(-ΔΔCT)) in the control sample. The Student’s t-test (two-tailed) was used to perform comparisons between stretched cells and static control. Three independent experiments were performed.

RNA-seq

RNA degradation and contamination were monitored on 1% agarose gels. RNA purity was checked using the NanoPhotometer® spectrophotometer (IMPLEN, CA, USA). RNA integrity and quantitation were assessed using the RNA Nano 6000 Assay Kit of the Bioanalyzer 2100 system (Agilent Technologies, CA, USA). A total amount of 1 μg RNA per sample was used as input material for the RNA sample preparations. Sequencing libraries were generated using NEBNext® UltraTM RNA Library Prep Kit for Illumina® (NEB, USA) following the manufacturer’s recommendations and index codes were added to attribute sequences to each sample. The RNA was purified from total RNA using poly-T oligo-attached magnetic beads. Fragmentation was carried out using divalent cations under elevated temperature in NEBNext First Strand Synthesis Reaction Buffer (5X). First-strand cDNA was synthesised using random hexamer primer and M-MuLV Reverse Transcriptase (RNase H). Second strand cDNA synthesis was subsequently performed using DNA Polymerase I and RNase H. Remaining overhangs were converted into blunt ends via exonuclease/polymerase activities. After adenylation of 3’ ends of DNA fragments, NEBNext Adaptor with hairpin loop structure was ligated to prepare for hybridisation. To select cDNA fragments of preferentially 150–200 bp in length, the library fragments were purified with AMPure XP system (Beckman Coulter, Beverly, USA). Then 3 μl USER Enzyme (NEB, USA) was used with size-selected, adaptor-ligated cDNA at 37 °C for 15 min followed by 5 min at 95 °C before PCR. Then PCR was performed with Phusion High-Fidelity DNA polymerase, Universal PCR primers and Index (X) Primer. At last, PCR products were purified (AMPure XP system) and library quality was assessed on the Agilent Bioanalyzer 2100 system. The clustering of the index-coded samples was performed on a cBot Cluster Generation System using PE Cluster Kit cBotHS (Illumina) according to the manufacturer’s instructions. After cluster generation, the library preparations were sequenced on an Illumina NovaSeq 6000 S4 platform and paired-end reads were generated. Reference genome and gene model annotation files were downloaded from the genome website browser (NCBI/UCSC/Ensembl) directly. Paired-end clean reads were mapped to the reference genome using HISAT2 software. Differential expression analysis between stretch and static control was performed using the DESeq2 R package. The resulting P values were adjusted using the Benjamini and Hochberg’s approach for controlling the false discovery rate (FDR). Genes with an adjusted P value < 0.05 found by DESeq2 were assigned as differentially expressed. Three independent experiments were performed.

Secretome and proteomic quantitative mass spectrometry

Osteocyte-like cells were cultured in a collagen-coated flexible bottom 6-well plate (Biorfix, Flexcell) with a cell density of 18,000 viable cells/cm² for 24 h. The plates were transferred to the bioreactor (Flexcell) to initiate the mechanical loading protocol, 34,000 µs, 2 Hz for 5 h. Cells seeded on the same plates but cultured under static conditions were used as control. Mechanically loaded and static osteocyte-like cells were cultured in a humified incubator at 37 °C and 5% CO₂.

Published in partnership with the Australian Regenerative Medicine Institute
for normalisation of the data was the most likely ratio normalisation. 51% confidence limit and a minimum of two peptides per protein were used as criteria for assessing fold change. Four independent experiments were performed.

Bioinformatics
Transcriptome. GO enrichment analysis of differentially expressed genes was implemented by the GOseq R package, in which gene length bias was corrected. GO terms with corrected P value less than 0.05 were considered significantly enriched by differential expressed genes. We used KOBAS software to test the statistical enrichment of differential expression genes in KEGG pathways.

Secretome. Biological processes of significantly changed proteins were analysed by gene ontology using the curated database Metacore (Clarivate Analytics). The percentage of overlap between proteins secreted from mechanically stimulated and static cells was estimated using the Venny 2.1 software. Protein interaction maps were generated using the Cytoscape STRING App and String DB. Protein location was estimated using Panther.

Reporting summary
Further information on research design is available in the Nature Research Reporting Summary linked to this article.

DATA AVAILABILITY
The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium (http://proteomexchange.vproteome.com) via the PRIDE partner repository with the dataset identifier PXD020774. Transcriptomic data are available in the ArrayExpress database (http://www.ebi.ac.uk/arrayexpress) under accession number E-MTAB-9487.

Received: 28 May 2020; Accepted: 10 May 2021; Published online: 07 June 2021

REFERENCES
1. Rose, L. F. et al. The convergence of regenerative medicine and rehabilitation: federal perspectives. npj Regen. Med. 3, 19 (2018).
2. Kubiak, E. N. et al. Early weight bearing after lower extremity fractures in adults. J. AAOS 21, 727–738 (2013).
3. Pagnotti, G. M. et al. Combating osteoporosis and obesity with exercise: lever-aging cell mechanosensitivity. Nat. Rev. Endocrinol. 15, 339–355 (2019).
4. Cannioto, R. A. et al. Physical activity before, during, and after chemotherapy for high-risk breast cancer: relationships with survival. J. Natl Cancer Inst. 113, 54–63 (2021).
5. Tu, X. et al. Osteocytes mediate the anabolic actions of canonical Wnt/β-catenin signalling in bone. Proc. Natl Acad. Sci. USA 112, E478–E486 (2015).
6. Fonseca, H. et al. Voluntary exercise has long-term in vivo protective effects on osteocyte viability and bone strength following ovarioectomy. Calcif. Tissue Int. 88, 443 (2011).
7. You, L. et al. Osteocytes as mechanosensors in the inhibition of bone resorption due to mechanical loading. Bone 42, 172–179 (2008).
8. Moustafa, A. et al. Mechanical loading-related changes in osteocyte sclerostin expression in mice are more closely associated with the subsequent osteogenic response than the peak strains engendered. Osteoporos. Int. 23, 1225–1234 (2012).
9. Vazquez, M. et al. A new method to investigate how mechanical loading of osteocytes controls osteoblasts. Front Endocrinol. 5, 208 (2014).
10. Mann, V. et al. The influence of mechanical stimulation on osteocyte apoptosis and bone viability in human trabecular bone. J. Musculoskelet. Neuronal Interact. 6, 408–417 (2006).
11. Lu, J. Y. & Wang, Y. F. Osteocyte apoptosis: the roles and key molecular mechanisms in resorption-related bone diseases. Cell Death Dis. 11, 840 (2020).
12. Lynch, M. E. et al. In vivo tibial compression decreases osteolysis and tumor formation in a human metastatic breast cancer model. J. Bone Min. Res. 28, 2357–2367 (2013).
13. Mei, X. et al. Microfluidic platform for studying osteocyte mechanoregulation of breast cancer bone metastasis. Integr. Biol. 11, 119–129 (2019).
14. Lara-Castillo, N. et al. In vivo mechanical loading rapidly activates β-catenin signalling in osteocytes through a prostaglandin mediated mechanism. Bone 76, S8–66 (2015).
15. Holguin, N., Brodt, M. D. & Silva, M. J. Activation of Wnt signaling by mechanical loading is impaired in the bone of old mice. J. Bone Min. Res. 31, 2215–2226 (2016).
16. Tan, S. D. et al. Fluid shear stress inhibits TGF-β-induced osteocyte apoptosis. J. Dent. Res. 85, 905–909 (2006).
17. Joeng, K. S. et al. Osteocyte-specific WNT1 regulates osteoblast function during bone homeostasis. J. Clin. Invest. 127, 2678–2688 (2017).
18. Cho, S. et al. Mechanosensing by the lamina protects against nuclear rupture, DNA damage, and cell cycle arrest. Developmental Cell 49, 920–935 (2019).
19. Kato, Y. et al. Establishment of an osteocyte-like cell line, MLO-Y4. J. Bone Min. Res. 12, 2014–2023 (1997).
20. Maupin, K. A., Droscha, C. J. & Williams, B. O. A comprehensive overview of skeletal phenotypes associated with alterations in WNT/β-catenin signalling in humans and mice. Bone 1, 27–71 (2013).
21. Perlman, R. L. Mouse models of human disease: an evolutionary perspective. Evol. Med. Public Health 2016, 170–176 (2016).
22. Qing, H. et al. Demonstration of osteocytic perilacunar/canalicular remodelling in mice during lactation. J. Bone Min. Res. 27, 1018–1029 (2012).
23. Cox, T. R. & Erler, J. T. Remodeling and homeostasis of the extracellular matrix: implications for fibrotic diseases and cancer. Dis. Models Mechanisms 4, 165–178 (2011).
24. Koehwyn, G. et al. Exercise-dependent regulation of the tumour microenvironment. Nat. Rev. Cancer 17, 620–632 (2017).
25. Schupbach, D. et al. Wnt modulation in bone healing. Bone 38, 115491 (2020).
26. Santos, A. et al. Pulsating fluid flow modulates gene expression of proteins involved in Wnt signaling pathways in osteocytes. J. Orthop. Res. 27, 1280–1287 (2009).
27. Snijders, T. et al. A single bout of exercise activates skeletal muscle satellite cells during subsequent overnight recovery. Exp. Physiol. 97, 762–773 (2012).
28. Bouchter, Y., Hopp, J. & Bouchter, S. Acute effect of a single bout of aerobic exercise on vascular and baroreflex function of young males with a family history of hypertension. J. Hum. Hypertens. 25, 311–319 (2011).
29. Mullender, M. et al. Mechanotransduction of bone cells in vitro: mechanobiology of bone tissue. Med. Biol. Eng. Comput. 42, 14–21 (2004).
30. Han, Y., Cowin, S. C., Schaf, L. S. and Weinbaum, S. Mechanotransduction and strain amplification in osteocyte cell processes. Proc. Natl Acad. Sci. USA 101, 16689–16694 (2004).
31. Robinson, J. A. et al. Wnt/β-catenin signalling is a normal physiological response to mechanical loading in bone. JBC 281, 31720–31728 (2006).
32. Ugur-Kluscek, A. et al. Monoamine oxidase-A promotes protective autophagy in human SH-SYSY neuroblastoma cells through Bcl-2 phosphorylation. Redox Biol. 20, 167–181 (2019).
33. Lambert, J. et al. Mapping differential interacomes by affinity purification coupled with data-independent mass spectrometry acquisition. Nat. Methods 10, 1239 (2013).
34. Perez-Riverol, Y. et al. The PRIDE database and related tools and resources in 2019: improving support for quantification data. Nucleic Acids Res. 47, D442–D450 (2019).
35. Ather, A. et al. ArrayExpress update - from bulk to single-cell expression data. Nucleic Acids Res. 47, D711–D715 (2019).

ACKNOWLEDGEMENTS
The authors would like to thank Nottingham Trent University for the financial support and facilities made available for this study.

AUTHOR CONTRIBUTIONS
L.S. designed this study, cultured osteocytic cells under mechanically active and static conditions, undertook bioinformatic analysis and performed RNA extraction, cDNA and real-time PCR. A.U.-K quantified the protein content of the supernatant and prepared it for LC-MS/MS. C.C. undertook LC-MS/MS and D.J.B. analysed LC-MS/MS data and performed bioinformatic analysis. RNA was sequenced by Novogene, Cambridge, UK. All authors contributed with ideas, reviewed, edited and agreed with the final version of this manuscript.

COMPETING INTERESTS
The authors declare no competing interests.
