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Zoledronic acid prevents pagetic-like lesions and accelerated bone loss in the p62P394L mouse model of Paget’s disease

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ABSTRACT
Paget’s disease of bone (PDB) is an age-related metabolic bone disorder, characterised by focally increased and disorganised bone remodelling initiated by abnormal and hyperactive osteoclasts. The germline P392L mutation of SQSTM1 (encoding p62) is a strong genetic risk factor for PDB in humans, and the equivalent mutation in mice (P394L) causes a PDB-like disorder. However, it is unclear why pagetic lesions become more common with age. Here, we assessed the effect of the p62 P394L mutation on osteoclastogenesis and bone morphometry in relation to ageing, the natural history of lesion progression in p62P394L+ mice and the effect of zoledronic acid (ZA) on lesion development. p62P394L+/+ mice lost 33% more trabecular bone volume in the long bones by 12 months compared with age-matched WT littermates. The p62P394L+/+ mice lost 33% more trabecular bone volume in the long bones by 12 months compared with WT mice (P<0.01), and developed pagetic-like lesions in the long bones which progressed with ageing. ZA prevented the development of pagetic-like lesions, and increased trabecular bone volume tenfold compared with vehicle by 12 months of age (P<0.01). This demonstrates that ageing has a pro-osteoclastogenic effect, which is further enhanced by the p62 P394L mutation, providing an explanation for the increased penetrance of bone lesions with age in this model. Lesions are prevented by ZA, providing a rationale for early intervention in humans.

KEYWORDS: Paget’s disease of bone, Genetic animal models, Ageing, Bone morphometry, Antiresorptives, Zoledronic acid

INTRODUCTION
Paget’s disease of bone (PDB) is the second most common metabolic bone disorder after osteoporosis and its prevalence increases with ageing. PDB is characterised by focally increased and disorganised bone remodelling, which leads to the formation of poor-quality woven bone. Consequently, pain, bone expansion, deformities, secondary osteoarthritis, pathological fractures and, very rarely, sarcoma can develop in the skeletal sites affected by PDB (Ralston et al., 2008).

Pagetic lesions are believed to be initiated by abnormal, enlarged, hypernucleated and hyperactive osteoclasts, which drive the progression of osteolysis, with increased and disorganised osteoblast-mediated new bone formation leading to the production of woven bone. Bisphosphonates (BPs) have been used for decades as treatment for PDB owing to their ability to suppress osteoclast activity, and heal lytic lesions with subsequent restoration of histologically normal new bone deposition and symptomatic improvement (Meunier and Vignot, 1995; Reid and Hosking, 2011; Reid et al., 1996). Currently, the most potent of the BPs, zoledronic acid (ZA), is the treatment of choice for PDB as it effectively suppresses bone remodelling and typically normalises bone turnover markers, in some patients for up to 10 years (Cundy et al., 2017). There is evidence that BPs improve pain (Corral-Gudino et al., 2017); however, intensive treatment of established PDB does not improve disease outcome in terms of quality of life measures or pain (Langston et al., 2010; Tan et al., 2017), which implies that preventive intervention, prior to lesion development, could be a better strategy for susceptible individuals.

It has been shown that genetic predisposition to PDB is mediated through the effects of several common predisposing variants of moderate effect size coupled with influence of rare variants, which have large effect size (Albagha et al., 2010, 2013). The most important predisposing gene for PDB is sequestosome 1 (SQSTM1), which encodes p62, a scaffolding protein involved in a variety of cellular processes including signalling and protein degradation (Rea et al., 2013). PDB-associated mutations of p62 tend to cluster in (but are not limited to) the ubiquitin-associated domain and occur in up to 40% of patients with a family history of PDB and 5-10% of patients with ‘sporadic’ disease (Hocking et al., 2002; Laurin et al., 2002). The most common and most studied PDB-associated p62 mutation is P392L (Hocking et al., 2002; Laurin et al., 2002). Although it is not entirely clear how p62 mutations cause or predispose to PDB, current evidence points to increased receptor activator of nuclear factor kappa-B ligand (RANKL; also known as TNFSF11)-mediated signalling enhancing osteoclastogenesis and dysregulated protein degradation (Rea et al., 2013).

The penetrance of PDB in p62 mutation carriers rises with age to reach between 80% and 90% by the seventh decade of life (Morissette et al., 2006). However, recent observations of a delayed disease onset in the p62 mutation carriers’ offspring, a less severe disease phenotype (Bolland et al., 2007; Cundy et al., 2015) and decreasing incidence of PDB in many countries over the past 25 years (Corral-Gudino et al., 2013; Cundy et al., 1997) suggest
that nongenetic or environmental factors might play a role in triggering the disease and/or affecting its severity. The historical viral hypothesis is controversial owing to conflicting evidence (Friedrichs et al., 2002; Rima et al., 2002; Visconti et al., 2017); nevertheless, transgenic mouse models of PDB-like disorders induced by viral sequences have been reported, suggesting that overexpression of measles virus nucleocapsid protein in osteoclast precursors increased osteoclast activity and bone resorption in vitro and induced areas of high bone turnover in the vertebrae with a 30% penetrance at 12 month of age (Kurihara et al., 2006a,b). One group did not detect evidence of high bone turnover with the characteristics of PDB in the vertebrae of mice bearing a knock-in p62 P394L mutation (equivalent to the human P392L mutation) (Hiruma et al., 2008).

We reported that although the p62 P394L mutation seldom causes vertebral lesions in mice, it frequently causes PDB-like lesions in the long bones, which become increasingly penetrant with ageing (Daroszewska et al., 2011). However, the mechanisms responsible for the age-related increase in penetrance remain unclear and there have been no studies on whether or not BPs could modify this phenotype.

Here, we revisit the p62P394L model of PDB and seek to validate it in the context of age-related osteoclastogenesis. We explore the ‘natural history’ of murine pagetic-like lesion evolution and relate it to human pagetic lesion progression. Finally, we investigate the role of ZA in prevention of the PDB-like phenotype.

RESULTS

Osteoclast formation increases in p62P394L mice with ageing

Studies in vitro showed that macrophage colony-stimulating factor (M-CSF)- and RANKL-induced osteoclast formation from bone marrow-derived macrophages was significantly greater in aged (12-month-old) WT mice when compared with young adult (3-month-old) WT mice (Fig. 1A). The number of osteoclasts generated from young adult p62P394L+/- mice was significantly greater when compared with young adult WT littermates, whereas the number of osteoclasts generated from aged p62P394L+/- mice was greater when compared with young adult and aged WT mice (Fig. 1A,B). This effect was even more striking in the p62P394L+/- mice. The number of osteoclasts generated from aged p62P394L+/- mice increased approximately twofold when compared with young adult p62P394L+/- mice (Fig. 1C) and threefold when compared with aged WT littermates (Fig. 1A,C). Moreover, osteoclast precursors from p62P394L+/- mice showed evidence of increased sensitivity to RANKL as compared with WT cells, at 10 ng/ml, 30 ng/ml and 100 ng/ml RANKL stimulation, which was intensified by ageing (Fig. 1C). A similar effect was seen in osteoclast precursors generated from the p62P394L+/-, although not as pronounced as in the homozygotes (Fig. 1B). Thus, ageing increases RANKL-induced osteoclastogenesis, and the p62 P394L mutation further enhances the age-related increase in osteoclastogenesis with a gene dosage effect.

The p62P394L mutation is associated with accelerated age-related long bone loss

We have previously shown that there was no significant difference in trabecular bone density and structure at the proximal tibial metaphyses of young adult (4-month-old) p62P394L+/+ male mice and WT littermates (Daroszewska et al., 2011). In view of the p62 P394L mutation-induced potentiation of age-related increase in osteoclastogenesis in vitro, we asked whether the p62 P394L mutation had an in vivo effect on age-related bone loss. We examined the distal femoral metaphyses of 12-month-old p62P394L+/- mice and WT littermates using micro computed tomography (μCT). There was a significant decrease in bone volume to total volume (BV/TV) of 33% (P<0.01), a significant decrease in trabecular number (Tb.N) and a significant increase in trabecular separation (Tb.Sp) in aged p62P394L+/- mice compared with WT littermates (Fig. 2), in keeping with accelerated age-related bone loss. There were no significant differences in vertebral (L5) morphometry between p62P394L+/- mice and WT littermates (data not shown).

Evolution of pagetic-like lesions with ageing in p62P394L mutant mice

In the PBS-treated group, 8/10 (80%) p62P394L+/+ mice developed pagetic-like lesions in the femur and/or tibia with the morphology described before (Daroszewska et al., 2011) by 12 months, as compared with 0/10 (0%) in the ZA-treated p62P394L+/- mice (P<0.001, Fisher’s exact test; see also next section). As femoral pagetic-lesions in patients progress linearly at an estimated rate of 9.4 mm per annum (Renier and Audran, 1997), we monitored PBS-treated p62P394L+/- mice in vivo with μCT to capture and follow up lesion progression. An example of the most severe lesion observed in this cohort and its evolution until the age of 18 months is shown in Fig. 3. The linear progression (Fig. 3D) between the age of 8 and 10 months was from 1.173 to 2.304 mm (change of 1.131 mm); between 10 and 15 months from 2.304 to 4.146 mm (change of 1.842 mm); and between 15 and 18 months from 4.146 to 4.696 mm (change of 0.55 mm). Thus, the average linear progression rate was 0.37 mm per month (4.47 mm per year) to involve ~28.5% of the femur, given the femoral length of 16.5 mm, and the lesion gradually expanded in 3D as well (Fig. 3). Given that mice over 6 months old age 25× faster than humans (www.jax.org), and that a female human femur is, on average, 445 mm long (human femur length to mouse femur length, 445 mm/16.5 mm=26.97), the 1.131 mm change over 2 months in mouse is estimated to correspond to a 7.42 mm change per annum in a human. Likewise, the 1.842 mm (over 5 months) and 0.55 mm (over 3 months) changes in mice correspond to 4.84 mm and 2.41 mm growth per annum, respectively, in a human.
Accordingly, the average mouse lesion progression rate of 4.47 mm per year corresponds to a 4.89 mm annual progression in human.

**Effect of ZA on the development of pagetic-like lesions and bone morphology**

To clarify whether BPs could prevent the development of pagetic-like lesions in the p62P394L+/+ mice, we chose ZA, which is a widely used treatment for PDB and osteoporosis. We monitored for lesion development using in vivo μCT at 1-2 month intervals, but did not detect any pagetic-like lesions in either of the p62P394L+/+ groups treated with ZA (hence the data were pooled). However, a significant change in bone morphology was observed in all ZA-treated p62P394L+/+ compared with vehicle-treated p62P394L+/+ and untreated WT mice (Fig. 4). The cortex was significantly thickened in the ZA-treated p62P394L+/+ mice (Fig. 4B) compared with untreated WT mice, and more so when compared with the vehicle-treated p62P394L+/+ mice affected by pagetic-like lesions (where cortical thickening is part of the picture; Fig. 4C). The metaphyses shape in the ZA-treated p62P394L+/+ mice was cylindrical as opposed to fluted-like in the vehicle-treated p62P394L+/+ and untreated WT mice (Fig. 4D-F). The growth plate looked highly mineralised. Trabeculae were very well preserved and plate-like as opposed to rod-like structures seen with ageing (Fig. 4E). Although no obvious pagetic-like lesions in the ZA-treated p62P394L+/+ mice occurred, we observed lacunae in the significantly thickened cortex (Fig. 4E, compare with F).

On histology examination, tartrate resistant acid phosphatase (TRAcP)-stained osteoclasts were easily seen near the growth plate of ZA-treated p62P394L+/+ mice (Fig. 5A,B); however, no osteoclasts were identified within the lacunacies of the thickened cortex (Fig. 5A,C). Thus, treatment with ZA did not inhibit all osteoclastogenesis, which was ongoing near the growth plate. The cortical lacunacies are unlikely to represent treated pagetic-like lesions, as no disorganised bone or osteoclasts were evident (Fig. 5A,C). Goldner’s trichrome stain showed no osteoid seams (Fig. 5D,E), and very little calcein double label was present (Fig. 5F,H) by 12 months in ZA-treated p62P394L+/+ mice, suggesting suppressed new bone formation. There was very little label in the cortical lacunacies (Fig. 5F,H) [increased labelling would be expected in active pagetic-like lesions (Daroszewska et al., 2011)]. Bone histomorphometry analysis revealed significantly increased bone volume per tissue volume (BV/TV) in ZA-treated versus PBS-treated p62P394L+/+ mice (Table S1). However, bone formation parameters – mineral apposition rate (MAR), mineralising surface per bone surface (MS/BS) and formation rate per bone surface (BFR/BS) – were significantly reduced in the ZA- compared with PBS-treated p62P394L+/+ mice (Table S1).

We then analysed the dynamic bone morphometry of the ZA- and PBS-treated p62P394L+/+ mice using the obtained in vivo μCT scans. Between 4 and 12 months of age, the PBS-treated p62P394L+/+ mice lost endosteal, while gaining periosteal, bone at the femoral mid-shaft, which led to an increase in bone diameter and marrow space (Fig. 6A,C,D). In contrast, the ZA-treated mice gained both endosteal and periosteal bone, although the periosteal bone gain was reduced compared with that of PBS-treated control animals (Fig. 6B-D). Treatment with ZA led to a rapid increase in trabecular bone volume during the first 2 months (0.38±0.09 mm³/month; Fig. 6F,G), owing to almost complete suppression of bone resorption and a substantial increase in bone formation. Over the next 6 months, the bone volume further increased, but at a much slower rate (0.05±0.04 mm³/month, P<0.001; Fig. 6E-H). The PBS-treated control p62P394L+/+ animals showed a small increase in trabecular bone volume; however, this was partially offset by trabecular bone loss at 6 months (net bone gain between 4 and 6 months, 0.08±0.04 mm³/month; Fig. 6E,G), and completely offset by 12 months of age (net bone gain between 6 and 12 months, −0.02±0.03 mm³/month; Fig. 6H).

**ZA increases bone mass in p62P394L+/+ mice**

In view of the striking changes in the ZA-treated p62P394L+/+ mice bone morphology, we carried out a μCT morphometry analysis. At 12 months, BV/TV was approximately tenfold higher in the ZA-compared with PBS-treated p62P394L+/+ mice (P<0.001; Fig. 7), in keeping with histomorphometry findings (Table S1). Tb.Th increased by 20% (P<0.01), Tb.Sp decreased by 60% (P<0.001) and Tb.N increased by over sixfold (P<0.01) in the ZA-treated group (Fig. 7). There was no significant difference between the four mice that received six doses, and the six mice that received five doses, of ZA; therefore, the data were pooled. Thus, ZA not only prevented the development of pagetic-like lesions and protected against the accelerated bone loss in the p62P394L+/+ mice, but substantially increased bone volume and enhanced bone structure.

**Long-term treatment with ZA leads to highly mineralised bone in the p62P394L+/+ mice**

Long-term administration of ZA to young adult p62P394L+/+ mice led to suppression of bone turnover, which can result in hypermineralisation of bone matrix (Allen and Burr, 2011). Furthermore, during sectioning for histology, we observed that the bone samples were brittle and caused damage to the microtome knives, suggesting high mineralisation. To investigate whether the bones of ZA-treated mice were indeed highly mineralised, we performed additional scans of the distal femurs, whether the bones of ZA-treated mice were indeed highly mineralised, we performed additional scans of the distal femurs, with increased averaging and camera binning to reduce image noise, and analysed tissue mineralisation. The ZA-treated p62P394L+/+ mice had significantly higher bone tissue mineralisation compared with the PBS-treated p62P394L+/+ mice (Fig. 8A). The mean bone mineralisation density in the ZA-treated cohort was 1.47±0.031 g/cm³, compared with 1.416±0.032 g/cm³ in the PBS-treated cohort (P<0.01). There was also an increased width of the density distribution in the treated group (standard deviation of the distribution 0.105±0.004 g/cm³) compared with that in the control group (standard deviation of the distribution 0.093±0.004 g/cm³, P<0.001) (Fig. 8B).
DISCUSSION

We have previously shown that the p62 P394L mutation is sufficient to induce a pagetic-like phenotype in mice, characterised by focal, asymmetric, mixed osteolytic/osteosclerotic lesions predominantly affecting the long bones, femur and tibia, with increased penetrance with ageing (Daroszewska et al., 2011). We also demonstrated the presence of microtubular structures in osteoclasts within PDB-like lesions similar to those previously reported in human PDB (Daroszewska et al., 2011). In the present study, we examined the effect of the p62 P394L mutation on age-related osteoclastogenesis and bone loss, the development and progression of pagetic-like lesions, and the prophylactic use of ZA.

It is well established that osteoclasts and osteoclast precursors from patients with PDB, with or without the p62 P392L mutation, are hypersensitive to RANKL (Chamoux et al., 2009; Menaa et al., 2000; Neale et al., 2000). We have previously shown that osteoclast precursors generated from the bone marrow of 3-month-old p62P394L+/− mice were hypersensitive to RANKL, and that both heterozygous and homozygous osteoclasts showed higher bone resorption than the WT (Daroszewska et al., 2011). Here, we have extended our analysis to assess the effect of ageing on p62P394L osteoclast RANKL hypersensitivity, and found that it increased incrementally in all groups, i.e. WT, heterozygotes and homozygotes, with a p62 P394L allele dose effect. The in vitro
findings of an age-related increase in RANKL-induced osteoclast formation in WT concur with previous findings in mice (Cao et al., 2005; Perkins et al., 1994) and humans (Chung et al., 2014; Koshihara et al., 2002). The role of p62 in osteoclastogenesis has been established by Duran and colleagues, who reported that targeted disruption of p62 in mice impaired osteoclastogenesis mediated by PTHrP (also known as PTHLH) (Duran et al., 2004), whereas overexpression of the human p62 P392L mutation in murine osteoclasts has been shown to increase osteoclastogenesis and induce a low bone mass phenotype in young adult mice, which progressed with ageing (Kurihara et al., 2007). Human studies have shown that the p62 P392L variant increased osteoclast differentiation, nuclearity and longevity (Chamoux et al., 2009). However, our finding of age-related enhancement of already high RANKL-induced osteoclast formation owing to the p62 P394L mutation (with an allele dose effect) is novel, and suggests that the p62 P394L mutation significantly potentiates osteoclastogenesis in an otherwise already pre-osteoclast-rich ageing skeleton (Farr et al., 2017), which intuitively might be permissive for relatively minor stimuli to induce further ‘uncontrolled’ osteoclastogenesis and pagetic lesions. In line with these observations, the known age-related increase of constitutive RANKL expression in stromal cells, osteoblasts and osteocytes (Cao et al., 2005; Chung et al., 2014; Piemontese et al., 2017) could be a contributory factor to the development of an osteoclast formation-permissive environment (Hiruma et al., 2008).

As age-related bone loss coincides with increased osteoclast activity, which is potentiated by the p62 P394L mutation, we hypothesised that the p62 P394L mutation would cause accelerated bone loss with ageing. We previously showed that long bone morphometry of young adult (4-month-old) p62<sup>P394L<sup>+/+</sup> mice was no different to WT (Daroszewska et al., 2011). In a previous study of the equivalent to our p62<sup>P394L<sup>+/+</sup> transgenic mouse, Hiruma and colleagues did not see any morphometric differences in the spine for up to 12 months of age (Hiruma et al., 2008), and we similarly did not find evidence of increased bone loss at the spine in the present study (data not shown). However, we found a significant increase in bone loss of the hind limb bones of p62<sup>P394L<sup>+/+</sup> mice by 12 months of age, which suggests that the p62 P394L mutation enhances age-related bone loss, which likely becomes first apparent in the long bones, as age-related bone loss takes place in the long bones ahead of the vertebrae in mice (Glatt et al., 2007). Interestingly, pagetic-like lesions also develop preferentially in the long bones (Daroszewska et al., 2011) rather than vertebrae (Daroszewska et al., 2011; Hiruma et al., 2008), which raises the possibility that biomechanical factors might interact with the p62 P394L mutation to influence where and when bone lesions develop. Locomotion differences between human (bipedal) and mouse (quadrupedal) carry different mechanical loading, which coincides with differences in pagetic lesion distribution: axial skeleton and the long bones are preferentially affected in human, whereas long bones (but not the axial skeleton) are preferentially affected in mice. Interestingly, development of pagetic lesions in humans has been described in bones subject to decades of supraphysiological repetitive mechanical loading (Gasper, 1979; Solomon, 1979). Animal work has shown that mechanical loading-induced bone fatigue or microfractures (Cardoso et al., 2009; Noble et al., 2003; Verborgt et al., 2000, 2002) led to apoptosis of osteocytes (key mechanosensing cells in bone), which occurs focally. As osteocyte apoptosis promotes focal osteoclast activation (Kogianni et al., 2008), it is possible that in the ageing skeleton affected by the p62 P392L mutation, highly primed for RANKL-mediated osteoclastogenesis, a trigger for focal pagetic lesion development could be localised osteocyte apoptosis; for example, due to microcracks developing with ageing, or repetitive mechanical loading-induced bone fatigue. This hypothesis warrants further investigation because, if confirmed, it could have significant translational implications for carriers of the p62 P392L mutation and affected individuals.

In terms of mechanisms underlying the increased age-related RANKL-induced osteoclast formation potentiated by the p62 P392L mutation, it is possible that alterations in the autophagy pathway play a role. Whilst DeSelm and colleagues provided evidence for the noncanonical role of autophagy in the resorptive function of osteoclasts (DeSelm et al., 2011), we have previously shown increased expression of key regulatory autophagy genes – SQSTM1, autophagy-related gene-5 (ATG5) and microtubule-associated light chain 3 (LC3; also known as MAP1LC3A) – as well as increased accumulation of LC3-II after treatment with bafilomycin in pre-osteoclasts and osteoclasts, respectively, generated from young adult p62<sup>P394L<sup>+/+</sup> mice compared with WT mice (Daroszewska et al., 2011), in keeping with induction of the autophagy pathway. However, the effect of ageing on canonical and noncanonical autophagy pathways in osteoclasts is currently unknown.

The finding of the p62 P394 mutation’s ability to accelerate age-related RANKL hypersensitivity of osteoclasts, paralleled by increased bone loss in p62<sup>P394L<sup>+/+</sup> mice, is also interesting from the translational perspective. Although in vitro, the p62<sup>P394L<sup>+/+</sup> osteoclasts showed the highest RANKL hypersensitivity, heterozygous osteoclast formation was also significantly increased
compared with WT, implying that p62P394L+/− mice could also show accelerated bone loss with ageing. Indeed, we have previously shown an allele dose effect of the p62 P394L mutation on the PDB-like phenotype severity (Daroszewska et al., 2011) and, as such, focused our investigations on homozygotes. Thus, we have not aged heterozygotes in the current study in an effort to use the minimum number of mice necessary in keeping with the principles of the 3Rs (Replacement, Reduction and Refinement of Animals in Research; https://nc3rs.org.uk). As the vast majority of patients with the p62 P392L-associated PDB are heterozygous, although rare homozygous or compound heterozygous cases with severe disease have been reported (Collet et al., 2007; Eekhoff et al., 2004; Laurin et al., 2001, 2002; Morissette et al., 2006), it is unclear whether our findings could be translated to a potentially increased risk of osteoporosis in the carriers of the p62 P392L mutation. Intriguingly, whilst PDB is classically considered a focal disease, there is evidence of increased bone remodelling in sites unaffected (Meunier et al., 1980), and possibly an increased risk of vertebral and rib fractures, again at unaffected sites (Melton et al., 2000). However, whether patients with PDB, p62 P392L mutation-linked PDB or unaffected mutation carriers are at an increased risk of osteoporosis is currently unknown.

We have previously shown that the PDB-like phenotype penetrance in the p62P394L+/− mice increased with ageing and reached 70% and 95% by 8 and 12 months, respectively (Daroszewska et al., 2011). Here, we were interested to capture the moment of the PDB-like lesion occurrence and progression over time. Using an in vivo μCT approach, we confirmed that the lesions...
had mixed osteolytic/osteosclerotic morphology and enlarged over time, but generally did not occur before the age of 6 months. We did not observe purely sclerotic lesions (to indicate ‘burned out’ PDB), which can occur in long-standing PDB in some patients (Siris et al., 1980); however, this is not unexpected given the higher rate of bone turnover in mice, compared with humans and a relatively short observation period, as our mice were culled at the age of 12-18 months. We have previously shown that the lesions most commonly developed at the distal femur and proximal tibia and less commonly in the shaft of the long bones (Daroszewska et al., 2011), which is comparable with a similar distribution in humans in the long bones, albeit with more predilection for the proximal ends of the long bones in the latter (Renier et al., 1996). Here, we presented a lesion in the diaphysis of the femur, which progressed over time. The estimated rate of its linear progression was equivalent to 7.42 mm per annum growth in human at early stages and later to between 4.84 mm and 2.41 mm per annum, which compares to an ∼9.4 mm per annum linear progression of a femoral pagetic lesion in human, according to estimations made two decades ago (Renier and Audran, 1997), when PDB was more severe than currently. Thus, although the murine pagetic-like lesions appear to progress at a slower rate than the human ones, we argue that their progression is comparable. Indeed, the slowing down of progression with ageing, arguably could suggest eventual ‘burning out’ of the phenotype.

There is evidence that treatment with BPs inhibits PDB lesion progression and promotes formation of histologically normal bone (Reid et al., 1996). BPs also suppress the raised bone turnover that is characteristic of active PDB, and ZA is the most potent BP (Corral-
Thus, we were interested to assess whether ZA could prevent the development of pagetic-like lesions in our mouse model. No lesions were detected in the ZA-treated p62<sup>P394L</sup>+/+ mice, providing evidence of the efficacy of this agent in suppressing the raised bone turnover that occurs in this mouse model of PDB. Furthermore, the effectiveness of ZA given early, prior to lesion development, provides pre-clinical evidence, to support the approach used in the ZiPP study (ISRCTN11616770; https://doi.org/10.1186/ISRCTN11616770), in which ZA is being investigated as a means of preventing development of PDB lesions in SQSTM1 mutation carriers.

Although studying the general skeletal effect of ZA was not the primary objective of our experiments, our study led to a number of important observations with potential translational implications. The positive effect of ZA on bone volume was striking, and ZA had a profound inhibitory effect on bone resorption. Although the brittle nature of the sections from ZA-treated mice precluded formal measurement of bone resorption parameters by histomorphometry, observation of the (mostly badly damaged) sections indicated a significant reduction in osteoclast numbers in 12-month-old ZA-treated p62<sup>P394L</sup>+/+ mice. Moreover, ZA attenuated new bone formation, but with a net effect of increased bone volume. Furthermore, bones of the ZA-treated p62<sup>P394L</sup>+/+ mice showed significantly higher mineralisation compared with controls, which suggests that ZA significantly slowed down the bone remodelling process, which allowed for increased mineralisation over time. On the other hand, the change of shape at the mid-femoral shaft consequent to treatment with ZA suggests that modelling was still taking place, although it was impaired. The dose-dependent protective effect of ZA on bone has been described previously in animal models of ovariectomy-induced osteoporosis (Gasser et al., 2011). Thus, given incomplete penetrance and declining prevalence and severity of PDB, careful consideration would have to be given when contemplating prophylactic use of ZA in susceptible individuals, in the light of potential negative ramifications of prolonged suppression of bone remodelling. It is expected that the outcome of the ZiPP trial will be instrumental in future clinical decision making.

The current study has confirmed our previous findings that the p62 P394L mutation is sufficient to cause a PDB-like disorder in mice and has provided new insight into the pathophysiology of the increased age-related penetrance of this disorder. The increase in osteoclastogenic potential of bone marrow cells from p62<sup>P394L</sup> mice with age provides an explanation for the increased penetration with age that we previously observed in this model (Daroszewska et al., 2011), and for the increased penetrance of PDB with age in humans (Morissette et al., 2006). Furthermore, the preferential targeting of the lower limb bones with sparing of the vertebrae suggests that biomechanical factors could play a key role in determining where and when lesions develop (Daroszewska et al., 2011).

In summary, we have shown that osteoclastogenesis is enhanced with ageing and the p62 P394L mutation further increases age-related osteoclastogenesis and age-related bone loss. In the ageing skeleton, the p62 P394L mutation causes PDB-like lesions, which progress over time. It is unclear what triggers these lesions; however, the increased age-related osteoclastogenesis potentiated...
by the p62 P394L mutation seems to allow for lesion development, thus creating a permissive environment. ZA prevents the development of the PDB-like lesions, significantly increases bone volume and bone mineralisation, and interferes with age-related long bone shape changes.

**MATERIALS AND METHODS**

**p62P394L+/+ mice**
The p62P394L+/+ mice were generated by gene targeting as previously described (Daroszewska et al., 2011) and housed in a standard animal facility with free access to water and food. The study was conducted in accordance with institutional, national and European regulations of laboratory animal care and use, and approved by the Home Office (UK). The mice were on a mixed 129sv and C57BL/6 background, and the colonies were maintained by breeding heterozygotes; WT, heterozygous and homozygous animals used in this study were littermates. As in this mouse model of PDB, there is no phenotypic difference between males and females (Daroszewska et al., 2011), for the assessment of ZA (Novartis) in prevention of pagetic-like lesion development, ten female p62P394L+/+ mice were randomly allocated to receive treatment with ZA in phosphate-buffered saline (PBS) at a dose of 85 µg/kg subcutaneously (equivalent to a human 5 mg/60 kg dose) as of 4 months of age, at which age mice are fully mature, but have no detectable pagetic-like lesions (Daroszewska et al., 2011). Of the ZA-treated p62P394L+/+ mice, four mice received three doses of ZA at monthly intervals and three subsequent doses at 2 month intervals (six doses in total). The remaining six mice received a total of five doses at 2 month intervals. The initial frequency of ZA administration at monthly intervals is approximately equivalent to 5 mg given at 2 year intervals to a human three times [given that a mouse ages 25× faster than a human (www.jax.org)]. Likewise, the frequency of administration of ZA 2 months apart to mice corresponds to 5 mg given 4.2 years apart to a human. This frequency corresponds broadly to the approach used in clinical practice for treatment of osteoporosis (three to six times annually) and PDB.

**Bone histology**
Bone samples were processed and stained for histology as described by van ‘t Hof et al. (2017). Briefly, animals received calcein intraperitoneal injections 4 days and 1 day before culling. The skin was removed, hind limbs were fixed in 4% formalin-buffered PBS overnight in water, and the distal femur scanned inside drinking straws using a Skyscan1272 scanner (resolution 4.5 µm, X-ray source at 50 kV and 200 µA, 0.5 mm Al filter, camera binning 2×2, rotation step size 0.3°, averaging at 3). Hydroxyapatite standards (Skyscan), were scanned using identical settings to calibrate mineral density. Next, datasets consisting of 300 slices at the mid shaft of the femur were thresholded to identify bone, and the binary was eroded (3D space) with a sphere with a radius of 2 to remove voxels affected by partial voxel effects. This binary was then used as a mask to measure the mineral density in mineralised tissue only [the tissue mineral density (TMD)].

**Statistical analysis**
Statistical analyses were performed using SPSS version 21. Differences between genotype or treatment groups were determined by ANOVA, Student’s t-test or Fisher’s exact test. All data are presented as means±s.d. unless stated otherwise.

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**Competing interests**
S.H.R. has received honoraria to his institution from Novartis, is in receipt of research grants from Eli Lilly and UCB, and was an investigator in clinical trials sponsored by Abbvie, Ultradynex, Amgen, Gilead and Eli Lilly. The other authors have no competing interests to declare.

**Author contributions**
Conceptualization: A.D., S.H.R., R.J.v.H.; Methodology: A.D., S.H.R., R.J.v.H.; Software: R.J.v.H.; Validation: A.D., R.J.v.H.; Formal analysis: A.D., R.J.v.H.; Investigation: A.D., L.R., N.S., G.C., A.P., K.R., R.J.v.H.; Resources: A.D., S.H.R., R.J.v.H.; Data curation: A.D., R.J.v.H.; Writing - original draft: A.D., R.J.v.H.; Writing - review & editing: A.D., S.H.R., R.J.v.H.; Visualization: A.D., R.J.v.H.; Supervision: A.D., R.J.v.H.; Project administration: A.D., S.H.R., R.J.v.H.; Funding acquisition: A.D., S.H.R., R.J.v.H.

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