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Julie CHOISNE, Jean-Marc VALIADIS, Christophe TRAVERT, Sami KOLTA, Christian ROUX, Wafa SKALLI - Vertebral strength prediction from Bi-Planar dual energy x-ray absorptiometry under anterior compressive force using a finite element model: An in vitro study - Journal of the mechanical behavior of biomedical materials - Vol. 87, p.190-196 - 2018

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Vertebral strength prediction from Bi-Planar dual energy x-ray absorptiometry under anterior compressive force using a finite element model: An *in vitro* study

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ARTICLE INFO

Keywords:

Osteoporosis
Vertebral strength
Bi-planar dual energy X-ray absorptiometry
Finite element model
Fracture

ABSTRACT

Finite element models (FEM) derived from qCT-scans were developed as a clinical tool to evaluate vertebral strength. However, the high dose, time and cost of qCT-scanner are limitations for routine osteoporotic diagnosis. A new approach considers using bi-planar dual energy (BP2E) X-rays absorptiometry to build vertebral FEM using synchronized sagittal and frontal plane radiographs. The purpose of this study was to compare the performance of the areal bone mineral density (aBMD) measured from DXA, qCT-based FEM and BP2E-based FEM in predicting experimental vertebral strength. Twenty eight vertebrae from eleven lumbar spine segments were imaged with qCT, DXA and BP2E X-rays before destructively tested in anterior compression. FEM were built based on qCT and BP2E images for each vertebra. Subject-specific FEM were built based on 1) the BP2E images using 3D reconstruction and volumetric BMD distribution estimation and 2) the qCT scans using slice by slice segmentation and voxel based calibration. Linear regression analysis was performed to find the best predictor for experimental vertebral strength (F_{expe}); aBMD, modeled vertebral strength and vertebral stiffness. Areal BMD was moderately correlated with F_{expe} ($R^2 = 0.74$). FEM calculations of vertebral strength were highly to strongly correlated with F_{expe} ($R^2 = 0.84$, $p < 0.001$ for BP2E model and $R^2 = 0.95$, $p < 0.001$ for qCT model). The results of this study suggest that aBMD accounted for only 74% of F_{expe} variability while FE models accounted for at least 84%. For anterior compressive loading on isolated vertebral bodies, simplistic loading condition aimed to replicate anterior wedge fractures, both FEM were good predictors of F_{expe} . Therefore FEM based on BP2E X-rays absorptiometry could be a good alternative to replace qCT-based models in the prediction of vertebral strength. However future work should investigate the performance of the BP2E-based model *in vivo* in discriminating patients with and without vertebral fracture in a prospective study.

1. Introduction

Osteoporosis is a bone disease which affects more than 1 in 3 women after their menopause and 1 of 10 men (McDonnell et al., 2007). The risk¹ of bone fracture caused by osteoporosis is predominant

at sites with a lot of trabecular bone such as the proximal femur, distal radius and vertebrae (Seeley et al., 1992). Vertebral fractures are one of the most common clinical manifestations with the major adverse consequences of osteoporosis as they usually occur under non-traumatic loading conditions. Height loss, back pain and functional disability are

the most encountered consequences of vertebral fractures with repetitive fracture experience more likely occurring within a year after the first fracture (Wadsworth, 1990; Kanis and McCloskey, 1992). Early diagnosis of osteoporosis is therefore important for vertebral fracture prevention as drug treatments are more effective before perforation of the trabeculae (McDonnell et al., 2007; Parfitt, 1987). However difficulties for early diagnosis are associated with the asymptomatic behavior of vertebral osteoporosis as first fracture usually occurs before the patient presents for treatment (Kanis and McCloskey, 1992; Kanis et al., 2013). Diagnosis techniques determine bone mineral density (BMD) from dual energy X-ray absorptiometry (DXA) (Kanis et al., 2013; Kanis and Johnell, 2005) or from quantitative computed tomography (qCT) scans (Buckley et al., 2007). DXA is the most clinically used method to diagnose osteoporosis. However this technique can only predict 40–70% of vertebral fractures as it only measures areal BMD (aBMD) which does not account for three dimensional (3D) geometry and BMD distribution (Sornay-Rendu et al., 2005). Volumetric BMD (vBMD) can be determined from qCT scans but showed modest correlations (R^2 ranging from 0.17 to 0.75) to experimental vertebral strength (Buckley et al., 2007; Crawford et al., 2003; Singer et al., 1995; Edmondston et al., 1997; Ebbesen et al., 1999). Therefore the combination of patient-specific 3D geometry and vBMD distribution is necessary to predict vertebral strength. Finite element (FE) models derived from qCT images demonstrated good correlation ($R^2 = 0.77–0.95$) in the prediction of experimental vertebral strength (Buckley et al., 2007; Crawford et al., 2003; Dall'Ara et al., 2010; Chevalier et al., 2008; Imai et al., 2006). This technique contributed to analyze therapeutic treatments effects (Chevalier et al., 2010; Keaveny et al., 2007), improved vertebral fracture risk assessment in men *in vivo* (Wang et al., 2012) and provided an alternative clinical tool to DXA for patients with osteoporosis (Kopperdahl et al., 2014). However, the main limitations of such approach in routine osteoporotic diagnosis are the high radiation dose associated with high financial cost and time of qCT imaging technique. An alternative to the disadvantages of qCT imaging keeping 3D FE modeling of the spine is bi-planar dual energy (BP2E) X-rays absorptiometry from a low dose imaging system (Dubouset et al., 2010). Combined with 3D reconstruction technique of the spine (Humbert et al., 2009) for volumetric geometry and dual energy absorptiometry for volumetric BMD distribution, this technique allows for patient-specific FE model of the osteoporotic vertebra.

The purpose of this study was to compare the performance of the aBMD from DXA, qCT-based FE model and BP2E-based FE model in predicting experimental vertebral strength. The experimental set up allowed for anterior compression testing on isolated vertebral bodies to ensure repeatable loading condition simulating an anterior wedge-shape fracture.

2. Material and method

2.1. Specimens

Eleven lumbar spine segments (L1–L3) from cadaveric specimens were considered in this study (5 males and 6 females, age: 82 years \pm 7 ranging from 61 years to 87 years). Donors were fresh cadavers and no exclusion criteria was specified. Medical history was unveiled but all donors and vertebrae were screened by a trained clinician to ensure there was no evidence of former surgical procedures or spine conditions (e.g. spine curve deviation, suspicion of bone tumor tissue, compression or traumatic fracture). A total of 28 vertebrae were included (8 L1, 11 L2 and 9 L3), after exclusion of vertebrae presenting anomalies.

2.2. Data acquisition

qCT scans of the lumbar spines were performed on a Scanner ICT 256 (Philips Healthcare, Cleveland, OH, USA) with the following settings; X-ray tube voltage and current: 120 kV, 1489 mA/s,

reconstruction matrix: 512×512 , field of view: 250×250 mm and a voxel size of $0.39 \times 0.39 \times 0.33$ mm. A calibration phantom (QRM-ESP, QRM GmbH, Germany) was used to map gray scale values to bone mineral density.

Dual energy X-ray absorptiometry (DXA) measurements were performed on a standard Hologic® QDR4500A (Hologic Inc, Waltham, MA, USA). The specimens were positioned in a 15 cm water bath and maintained with radio-transparent elastic bands. Bone mineral content (grams of hydroxyapatite equivalent), projected bone area (cm^2) and bone mineral density (g/cm^2) were calculated for L1, L2 and L3 using the standard A-P scanning protocol. Scanner precision was quantified on 5 specimens with measurements repeated 3 times.

Low dose bi-planar dual energy (BP2E) X-rays absorptiometry were acquired for all spine segments using a dual energy prototype of the EOS® system (EOS imaging, Paris, France) which can simultaneously take a pair of X-rays in the sagittal and frontal planes in upright position (Dubouset et al., 2010), allowing for 3D reconstruction of the spine (Humbert et al., 2009). Two levels of energy can be achieved with the EOS® prototype by quickly changing the X-ray tube settings between two fast passes scans (approximately 20 s depending on the size of the lumbar spine). The computed projected areal Bone Mineral Density (aBMD) images of the vertebrae are similar to DXA images (Sapin et al., 2008; Sapin-De Broses et al., 2012). aBMD measurement was previously validated by comparing EOS® accuracy and reproducibility with the dual x-ray absorptiometry densitometers' characteristics (Sapin et al., 2008). X-ray tube voltage and current were 140 kV and 149 mA for the high energy images and 70 kV and 298 mA for the low energy images.

2.3. Mechanical testing

The 28 vertebrae were extracted from the spinal segment by cutting the intervertebral disc with a scalpel and disarticulating the posterior facets. The vertebrae were then cleaned from all soft tissue and the posterior elements were transected at the pedicles junction to the anterior body. The vertebral end plates were embedded in polymethyl methacrylate (PMMA) for parallelism to ensure uniform loading conditions. Anterior compressive tests were conducted using a quasi-static traction-compression Instron 5500 R device (Instron Ltd., High Wycombe, UK) in combination with a 10 kN load cell and a spherical seating loading platen (Instron Ltd., High Wycombe, UK) allowing for uniform load application onto the upper surface of the specimen (Fig. 1). The center of rotation of the spherical seating loading platen was aligned with the anterior third of the vertebral body to apply anterior compressive load to the vertebra. The anterior third of the vertebral body was measured using EOS® radiographs of the vertebral body embedded in the PMMA layer before the destructive compression test. The specimen was bolted to the inferior test platen with the mold

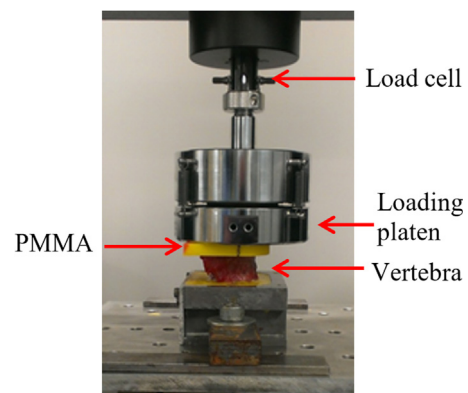


Fig. 1. Experimental setup for anterior compressive test of an isolated vertebral body.

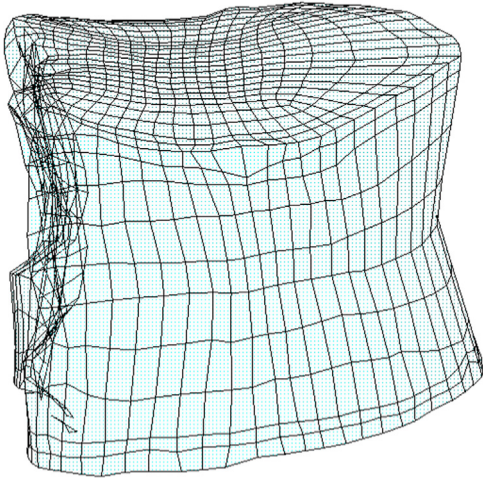


Fig. 2. Finite element model of a vertebral body. This model consisted of 8300 elements.

used for embedding the PMMA.

To avoid shear forces, the inferior test platen was mounted on an antero-posterior sliding connection and an initial compressive preload of 50 N was applied. Specimens were preconditioned by applying 10 cycles of compressive force between 100 N and 250 N at 0.1 Hz followed by destructive compression test at 1 mm/min until ultimate force was achieved (Buckley et al., 2007). Experimental vertebral strength was defined as the ultimate load achieved and axial stiffness was calculated as the slope of the linear region of the force-displacement curve.

2.4. Finite elements models

A qCT-based finite element model was built from vertebral geometry obtained by a semi-automatic segmentation method (Le Pennec et al., 2014). A hexahedral mesh of the vertebral body (Fig. 2) was generated from this geometry using a multiblock meshing program wrote in C++ (Grosland et al., 2009). Briefly, the multiblock meshing technique consists in multiple building blocks composed of meshing seeding arranged in rows, columns and layers. The mesh seeds are then projected on the vertebral surface and morphed to each vertebral surface as nodes to lay the foundation for the FE mesh (Grosland et al., 2009). In this 8304-element mesh the average element size was controlled to range between 1 mm and 1.5 mm. All meshes were generated with the same topology for each vertebral level allowing the same element to be located approximately at the same position in the vertebra. Convergence analysis was performed to determine the ideal number of elements needed (Choisne et al., 2017).

Once the mesh generated, the average BMD of a single finite element was defined on the basis of the qCT scan voxels that fall inside the element. BMD value for each element was converted to linear elastic mechanical properties from an experimental relationship between BMD and elastic modulus (Choisne et al., 2017) as shown in Eq. (1). The Poisson ratio, ν , was set to 0.4 (Imai et al., 2006).

$$E(\text{MPa}) = 3230 \text{ BMD} (\text{g}/\text{cm}^3) - 34.7 \quad (1)$$

A bi-planar dual energy X-rays absorptiometry based (BP2E-based) FE model was built from vertebral geometry obtained by 3D reconstruction of the spine from sagittal and frontal plane X-rays (Choisne et al., 2017). FE meshes similar to the qCT-based model were generated using the same element numbering and topology.

Volumetric BMD for each element was estimated from the lateral dual energy X-ray absorptiometry areal BMD image using a method previously described (Choisne et al., 2017). Shortly, a vBMD distribution based on the qCT density database was used to build a generic FEM model. Then, a digitally reconstructed radiography (virtual aBMD

image) was made based on the generic model. Finally, an iterative vBMD adjustment was performed to maximize pixel similarity between the virtual and BP2E aBMD image. Once the image similarity was optimized, the personalized vBMD distribution from BP2E images was set. BMD values were then converted to material properties using the same Eq. (1) for each element.

2.5. Boundary conditions

Previously described boundary conditions and failure criterion (Sapin-De Brosses et al., 2012) were considered to build the qCT-based and BP2E-based FE models in order to reproduce the experimental testing procedure. Briefly, all models were virtually added layers of PMMA (about 0.5–1 cm thick, $E = 2500 \text{ MPa}$, $\nu = 0.3$) to both vertebral endplates. Lower nodes of the lower PMMA layer were constrained in all degrees of freedom. Anterior compressive load was applied to the upper PMMA layer joined by rigid elements to a node located at the anterior third of the vertebra. Simulations were run on ANSYS software (ANSYS Inc., Canonsburg, PA, USA). Vertebral strength was defined when a contiguous region of 1 mm^3 of elements reached 1.5% deformation (Sapin-De Brosses et al., 2012).

2.6. Analysis

Shapiro-Wilk tests were used to assess the normality of the distributions. Data were presented as the mean, standard deviation, and range. Linear regression analysis was performed to find the best predictor for experimental vertebral strength estimation; aBMD determined from DXA, modeled vertebral strength and vertebral stiffness. The Pearson determination coefficients of these regressions (R^2) were calculated both in their raw and sample size-adjusted forms (adj. R^2). R^2 values were statistically compared using Steiger's Z test for dependent samples (Meng et al., 1992). $P < 0.05$ was considered the cut-off for significance in all statistical analyses.

3. Results

Descriptive data derived from the DXA analysis, finite element models analysis, and mechanical testing are summarized in Table 1.

According to patients classification criterion for DXA, eight specimens had osteoporosis with a t-score below -2.5 , one specimen had osteopenia and the two remaining specimen were considered normal (Assessment, 1994). The estimate of the 95% confidence interval of the DXA's precision error was $0.010 \text{ g}/\text{cm}^2$. Bland and Altman plots, linear regression analysis coefficient of determination (R^2), p-values, slope, intercept and standard error of the estimate (SEE) values are summarized in Table 2 and displayed in Figs. 3 and 4.

BMD measured from DXA was moderately related to experimental vertebral strength (F_{expe}). Experimental vertebral strength was better predicted by variables estimated from the FE models compared to aBMD measured from DXA as indicated by higher R^2 and smaller standard errors of the estimate (Table 2).

Table 1
Descriptive statistics of 28 vertebrae.

		Mean \pm SD	Range
DXA	T-score	-2.7 ± 1.4	-5.1 – 0.7
	BMC (g)	10.19 ± 3.88	5.50 – 21.09
	BMD (g/cm^2)	0.724 ± 0.182	0.440 – 1.186
qCT-based FEM	Failure load (N)	3067 ± 1362	1275 – 6331
	Stiffness (kN/mm)	$10,775 \pm 4332$	4998 – $19,358$
BP2E-based FEM	Failure load (N)	3033 ± 1484	995 – 6223
	Stiffness (kN/mm)	$11,103 \pm 4526$	4340 – $18,995$
Mechanical testing	Failure load (N)	3145 ± 1573	1270 – 6561
	Stiffness (kN/mm)	5473 ± 1892	1643 – 8523

Table 2

Linear regression analysis between the experimental vertebral strength and the following variables: areal bone mineral density (aBMD) measured from DXA, qCT-based FEM strength (F_{qCT}) and stiffness (K_{qCT}), BP2E-based FEM strength (F_{BP2E}) and stiffness (K_{BP2E}) and experimental vertebral stiffness (K_{expe}).

	R^2	Slope	Intercept	SEE % (N)
aBMD (mg/cm ²)	0.74 [*]	7.44	- 2244	25 (783)
F_{qCT} (N)	0.95 [*]	1.13	- 313	11 (333)
F_{BP2E} (N)	0.84 [*]	0.97	199	20 (617)
K_{expe} (kN/mm)	0.85 [*]	0.77	- 1062	19 (588)
K_{qCT} (kN/mm)	0.92 [*]	0.35	- 597	14 (450)
K_{BP2E} (kN/mm)	0.78 [*]	0.31	- 268	23 (720)

* $p < 0.001$.

The coefficient of determination for F_{expe} VS aBMD was significantly lower than for F_{expe} VS K_{expe} , ($p < 0.01$), F_{qCT} ($p < 0.01$) and K_{qCT} ($p < 0.05$). qCT-based finite element model calculations of vertebral strength (F_{qCT}) and stiffness (K_{qCT}) were strongly correlated with F_{expe}

($R^2 = 0.95$, $p < 0.001$ and $R^2 = 0.92$, $p < 0.001$ respectively). Statistical error in qCT-based model vertebral strength estimation was $78 N \pm 381$ compared to F_{expe} with a RMSE of 12% (382 N) and a SEE of 11% (333 N). BP2E-based FE model vertebral strength (F_{BP2E}) and stiffness (K_{BP2E}) determination were highly correlated with F_{expe} ($R^2 = 0.84$, $p < 0.001$ for F_{BP2E} and $R^2 = 0.78$, $p < 0.001$ for K_{BP2E}) and F_{qCT} ($R^2 = 0.90$, $p < 0.001$ for F_{BP2E} and $R^2 = 0.83$, $p < 0.001$ for K_{BP2E}). Statistical error in for F_{BP2E} estimation was $112 N \pm 629$ compared to F_{expe} with a RMSE of 20% (628 N) and a SEE of 20% (617 N).

4. Discussion

The purpose of this study was to compare the performance of the aBMD from DXA, qCT-based FEM and the bi-planar dual energy X-rays absorptiometry based (BP2E-based) FEM in predicting experimental vertebral strength.

The results of this study suggest that FE models are better

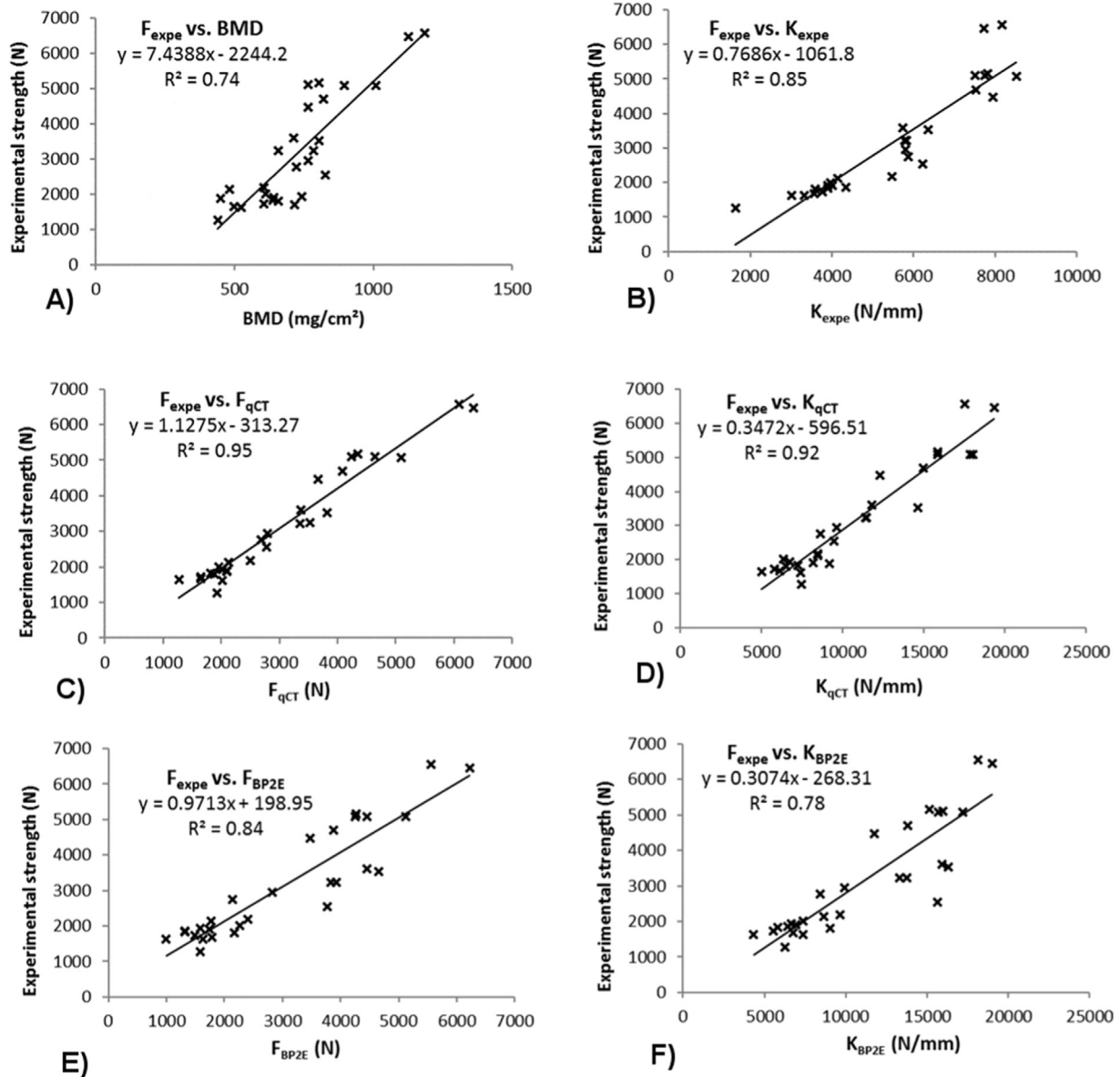


Fig. 3. Linear regressions of experimental vertebral strength as a function of A) bone mineral density (BMD) from DXA, B) experimental stiffness (K_{expe}), C) qCT-based FE model strength (F_{qCT}), D) qCT-based FE model stiffness (K_{qCT}), E) bi-planar dual-energy FE model strength (F_{BP2E}), and F) bi-planar dual-energy FE model stiffness (K_{BP2E}).

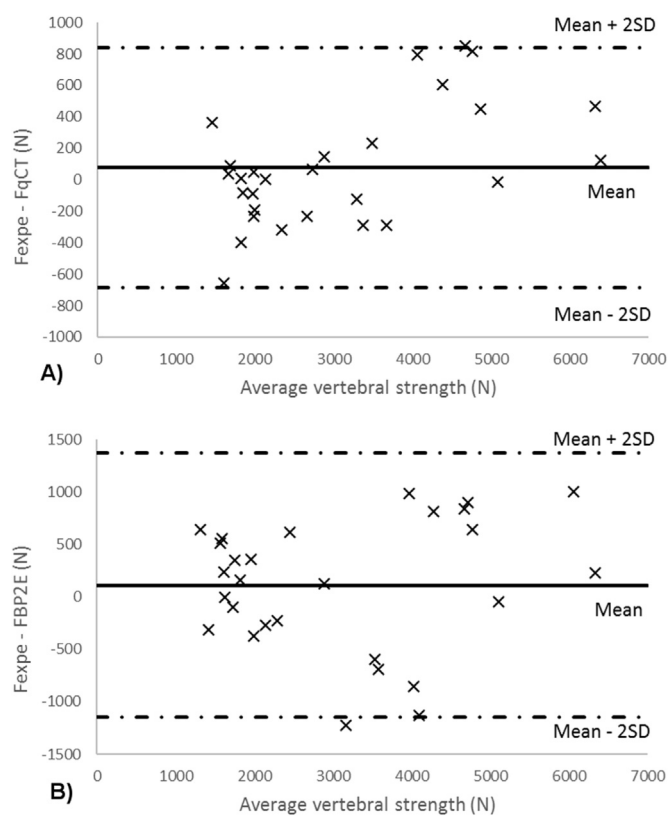


Fig. 4. Bland and Altman plots between experimental vertebral strength and vertebral strength determined from A) FE-based qCT (F_{qCT}) and B) FE-based BP2E (F_{BP2E}).

experimental vertebral strength predictors than aBMD measured with DXA. For anterior compressive loading on isolated vertebral bodies, simplified loading condition applied at the anterior third of the vertebra, aBMD accounted for 74% of the variability in experimental vertebral strength while FE models accounted for at least 84% of the strength variability. This 10% difference is clinically relevant as it can diagnose osteopenia bone earlier compared to DXA that can mostly distinguish between healthy and very osteoporotic bone.

4.1. BMD from DXA

BMD from DXA was the worst predictor for experimental compressive vertebral strength with a coefficient of determination of 0.74 and SEE of 25%. Previous studies showed similar correlation ranging from 0.46 to 0.69 between aBMD and experimental ultimate compressive load (Edmondston et al., 1997; Chevalier et al., 2008; Faulkner et al., 1991; Eriksson et al., 1989; Moro et al., 1995; Roux et al., 2010). PA-projection DXA is the method of choice to diagnose osteoporosis, however BMD measurement was shown to account for less than 75% of vertebral strength variability *in vitro*. Moreover *in vitro* DXA measurements are not influenced by extra-vertebral tissues and might therefore appear to have higher accuracy than DXA when performed *in vivo* (Ebbesen et al., 1999). Furthermore, an *in vivo* study aiming to prospectively assess the risk of new clinical vertebral fractures demonstrated that DXA measurement hazard ratio change between fracture and non-fracture for aBMD was significantly lower than for strength determined from finite element models based on qCT images (Wang et al., 2012).

4.2. qCT-based finite element model

Vertebral strength determined from qCT-based FE model was the

best predictor for experimental vertebral strength with a coefficient of determination of 0.95 and SEE of 11%. These results are in agreement with the literature regarding the efficacy of FE models to predict experimental vertebral strength with R^2 value ranging from 0.77 to 0.95. Moreover this model was a good predictor of bone elastic behavior with model stiffness highly correlated to experimental stiffness ($R^2 = 0.86$). Compared to existing FE models (Buckley et al., 2007; Crawford et al., 2003; Chevalier et al., 2008; Imai et al., 2006; Liebschner et al., 2003), this model was tested in a configuration replicating anterior wedge fractures, commonly seen on osteoporotic patients, by applying anterior compression on the vertebral body instead of centered compression. One study (Dall'Ara et al., 2010) used a similar mechanical setup by including a loading plate with a ball joint to allow for rotation and therefore creating anterior wedge fractures. Using a non-linear finite element model, they estimated experimental vertebral strength and stiffness with coefficients of determination of 0.79 and 0.49. The present study showed higher correlation with experimental output using a simplified linear FE model in a more time efficient manner, although our model is less adapted to determine fracture detailed mechanism oppositely to the nonlinear FE model (Dall'Ara et al., 2010).

4.3. BP2E based finite element model

BP2E-based model output was a good predictor for experimental vertebral strength and stiffness with coefficient of determination of 0.84 and 0.78 respectively. This model using bi-planar dual energy X-rays absorptiometry (BP2E) performed as good as existing FE models based on qCT images (Buckley et al., 2007; Crawford et al., 2003; Dall'Ara et al., 2010; Chevalier et al., 2008; Imai et al., 2006; Liebschner et al., 2003). Compared to the present qCT-based FE model, the coefficient of determination between BP2E-based model and experimental vertebral strength was statistically lower ($p < 0.05$) but with a slope closer to one (0.97 for BP2E vs 1.13 for qCT) and intercept approaching 0 (199 for BP2E vs -313 for qCT). Both models were highly correlated with $R^2 = 0.90$ for vertebral strength and 0.83 for stiffness. FE models based on BP2E X-rays absorptiometry could become a good alternative to replace qCT-based model in the prediction of vertebral strength avoiding qCT scan disadvantages such as high cost and radiation exposure. Indeed the EOS[®] device is a low dose X-ray system with a fast acquisition time and an effective dose received of ~ 0.3 mSv (Damet et al., 2014) compared to 5 mSv with qCT scan (Wang et al., 2012). Lateral and AP projection DXA images could be used with the same approach keeping in mind that DXA images resolution is low (Sapin et al., 2008; Ahmad et al., 2010; Whitmarsh et al., 2011, 2013).

4.4. Potential clinical applications

This study shows improved prediction in the vertebral strength assessed experimentally by using FE models compared to DXA, the most clinically used method to detect osteoporosis. Moreover this study highlights potential clinical applications of BP2E-based FE models and the advantages of using FE models rather than DXA in the diagnosis of osteoporosis. Further, these findings support those of previous studies showing that vertebral FE models are better predictors for experimental vertebral strength than BMD measurements alone (Buckley et al., 2007; Crawford et al., 2003).

If bi-planar dual energy (BP2E) X-rays absorptiometry performs as well *in vivo* as it did *ex vivo*, BP2E-based FE model could potentially become an alternative to qCT-based models involving high cost and radiation exposures. Due to qCT scan disadvantages most clinicians prefer using less irradiant methods such as DXA instead of qCT images in the diagnosis of osteoporosis (Kanis et al., 2013). The present study demonstrated that BP2E-based FE model is a better predictor for experimental vertebral strength than DXA *ex vivo*. A previous study demonstrated that the EOS[®] system has a good diagnostic value for the diagnosis of vertebral fracture with a better legibility of upper thoracic

spine than DXA *in vivo* (Briot et al., 2015). With its low radiation dose and cost, the EOS® system is a potential useful technique in the diagnosis of osteoporosis. However, it will be important to undertake prospective studies to determine the predictive strength of BP2E-based model for vertebral fracture risk assessment before the clinical benefit can be judged. While the vertebral strength prediction of the FE method used in this study is good for *ex vivo* condition, precision for *in vivo* applications is likely to be more variable. Factors influencing *in vivo* measurements are the soft tissue heterogeneity surrounding the lumbar spine, the variability of *in vivo* positioning and the greater soft tissue volume X-ray absorption. Nonetheless, most of these factors can be controlled using the sagittal and frontal views to assess soft tissue heterogeneity and include it in the calculation of soft tissue attenuation. Variability in patient positioning should not be an issue for 3D reconstruction as the method demonstrated a position precision of maximum 2.1 mm error on spine with severe scoliosis (Cobb's angle > 30°) (Humbert et al., 2009). Moreover the volumetric BMD distribution method takes into account axial rotation as long as the posterior arch does not superimpose the vertebral body on the lateral view, which will automatically overestimate the vBMD distribution as the posterior arch is significantly denser than the vertebral body (Travert et al., 2013). Future work will assess the performance of the BP2E-based FE model in discriminating patients with and without vertebral fracture in a prospective study.

The present study has several limitations to be considered. Possible error sources were the accuracy of the 3D reconstruction, which can affect the vertebral body volume and thus the apparent density, the contribution of the cortical bone layer and, to a lesser extent, the surrounding soft tissues. However, spine 3D reconstruction position precision was quantified to be less than 1.8 mm which should not affect average vBMD distribution (Humbert et al., 2009). Moreover, aBMD accuracy of the EOS® system was determined to be below 5.2 per cent, versus 7.2 per cent for a DXA system in the same conditions (Sapin et al., 2008). Cortical shell was not modeled in either FE models since qCT-scan and X-ray images are not precise enough to measure cortical thickness with pixel sizes being larger than average cortical thickness in a vertebra. The contribution of the cortical shell to vertebral strength is not well established in the literature with cortical shell accounting from 10% to 75% for vertebral strength (Roux et al., 2010; Rockoff et al., 1969; Andresen et al., 1998; McBroom et al., 1985). However trabecular bone appears to play the primary role in determining vertebral body strength (Silva et al., 1997; Hulme et al., 2007; Perilli et al., 2012; Fields et al., 2009). Other limitations are the sample size and the cadaveric nature of the vertebrae. Twenty-eight vertebrae is a reasonable number when looking at correlation however the vertebrae were extracted from 11 donors which makes the data not truly independent. Pseudo-replication was accounted by calculating an adjusted R² coefficient using the average data for each donor. However 11 data points was not enough for correlation analysis and underpowered the study. Given the purpose of the present study is to compare the performance from different modalities in predicting experimental vertebral strength, and therefore using a repeated between-vertebrae design, pseudo-replication should not be a significant issue.

Further limitations are the *in vitro* nature of the study which might not be representative to *in vivo* situation. Therefore these results cannot be directly extrapolated into *in vivo* condition and future work should assess the performance of the BP2E-based FE model in discriminating patients with and without vertebral fracture. Moreover, the loading conditions applied on the vertebrae were designed to replicate anterior wedge-shape fractures but they were not fully representative of *in vivo* loading conditions as adjacent vertebrae and intervertebral disc are major factors to vertebral fractures. Another advantage in using bi-planar X-rays in erect position is to consider patient-specific loading condition *in vivo* and therefore taking into account the patient's posture.

5. Conclusion

In conclusion in this *in vitro* study of lumbar vertebral bodies, it was shown that vertebral strength determined from FE models are strong predictors of experimental failure load, highlighting the capability of bi-planar dual energy (BP2E) X-rays absorptiometry based FE model to predict vertebral body strength. BP2E X-ray absorptiometry allows for fast, low-radiation and minimal cost patient-specific 3D FE model as accurate as qCT-based FE models, currently the gold standard for vertebral strength prediction. These findings support further exploration of the clinical application of BP2E-based FE models in vertebral fracture prediction.

Acknowledgment

The authors would like to thank S. Persohn and M. Jeyasankar for contributing to mechanical testing. The authors would also thank Anabela Darbon, advanced research engineer at EOS Imaging, for EOS® dual energy acquisition and calibration. This work was supported by the Banque Publique d'Investissement through the dexEOS project part of the FUI14. The funding agencies had no role in the design and conduct of the study, in the collection, management, analysis and interpretation of the data, or in the preparation, review, or approval of the manuscript.

Authors'roles

Study design: JC, JMV and WS. Study conduct: JC, JMV, CT, SK, CR and WS. Data collection: JC, JMV and SK. Data analysis: JC, JMV and WS. Data interpretation: JC, SK, CT, CR and WS. Drafting manuscript: JC and WS. Revising manuscript content: JC, JMV, CT, SK, CR and WS. Approval final version of manuscript: JC, JMV, CT, SK, CR and WS. WS takes responsibility for the integrity of the data analysis.

Conflict of interest

The authors have no conflict of interest to declare. Wafa Skalli is the co-inventor of the EOS system without direct financial interest.

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