INTRODUCTION
In an era of increasing life expectancy, osteoporosis has become a major global health concern. Osteoporosis is a metabolic disease of bone and at least one-third of all postmenopausal females will suffer an osteoporotic fracture in her lifetime. Global estimates postulate that approximately 20 osteoporotic fractures occur every minute and the National Osteoporosis Foundation (NOF) estimates that approximately 54 million Americans suffer from osteoporosis resulting in 2 million fractures annually. Many population-based studies have demonstrated concerning upward trends for increasing incidence of hospitalisations for osteoporotic fractures worldwide.

Vertebral fractures (VFs) account for up to 50% of osteoporotic fractures making them the most common fracture subtype. Incidence of VFs increases with increasing age, with significant heterogeneity between sexes, origin and age of onset. Up to 26% of Scandinavian females will be diagnosed with at least one VF in her lifetime. VFs are a major cause of patient pain, reduced mobility and many patients who have sustained a VF suffer with the psychological fear of isolation and loss of independence. Additionally, sustaining a VF is an independent risk factor for patient mortality, emphasising its significance as a major public health concern. Studies show that patients with previous VFs are five times more likely to obtain an additional VF and are twice as likely to suffer a hip fracture with resulting morbidity and mortality.

Encouragingly, robust evidence has shown that early intervention with pharmacological agents such as bisphosphonates result in a relative risk reduction of up to 0.6 for VFs and up to 0.8 for non-VFs. Therefore, it is vital that VFs are correctly diagnosed so that patients are appropriately investigated and placed on correct medical therapy. Unfortunately, there is a discrepancy between best recommended management and real-life clinical practice with one study concluding that less than one-quarter of patients diagnosed
with an osteoporotic fracture are appropriately investigated and treated for osteoporosis.15

Studies suggest that radiologists have contributed to the current problem due to non-diagnosis or underreporting of VFs.16,17 VFs are evident on various imaging modalities which are performed for alternative clinical indications but are frequently not reported by radiologists.16,19 Use of terminology such as ‘wedging’, ‘endplate compression’ and ‘endplate concavity’ in radiology reports can be confusing and may not be understood to indicate the presence of osteoporotic VF by the reading physician. Non-diagnosis or inappropriate reporting of VFs in this way is a missed opportunity to diagnose osteoporosis, impedes adequate medical treatment and renders the patient vulnerable to sustaining future osteoporotic fractures.15

In the current paper, we discuss radiological assessment of VFs and describe how fractures can be diagnosed on the most commonly used imaging modalities, plain film, MRI, CT and bone scans.

Assessment of fractures
Genant et al devised the semi-quantitative (SQ) method for describing VFs. This method has high inter- and intraobserver agreement, even amongst inexperienced reviewers.20 SQ is relatively straightforward method to grade fractures and avoids otherwise confusing language which may be misinterpreted. First described on lateral radiographs, SQ method employs visual inspection to grade VFs. Grade 0 is normal without loss of vertebral body height. Grade 0.5 are borderline vertebrae. Grade 1 fractures show mild deformity with approximately 20 to 25% loss of height and 10 to 20% reduction in area. Grade two fractures are moderately deformed with 25–40% loss of height and 20–40% loss of area. Grade 3 VFs have lost 40% or more of their height and area.

Recognition of fractures
Imaging modalities
Plain films: For clinically suspected VFs, plain films including anteroposterior (AP) and lateral projections are usually the first line of investigation. The lateral film is particularly useful (Figures 1–2). The radiologist should carefully examine the vertebral body outline, especially the superior and inferior endplates to ensure VFs are not missed. The pedicles should be examined for symmetry on the AP film. Subjectively identifying reduced bone density should heighten the index of suspicion for VFs as these patients are at much greater risk. Dynamic radiographs of the vertebrae can increase the likelihood of correct diagnosis on plain radiography. This method allows the radiologist to compare supine images with lateral sitting radiographs to evaluate for changes in vertebral body height. The sensitivity and specificity of dynamic radiographs for diagnosing acute VFs is 66 and 96%, respectively.21 While moderate and severe VFs are rarely misdiagnosed, there are a number of conditions which can be mistaken for mild VFs leading to overdiagnosis. These include developmental short vertebral height, physiological wedging, Scheuermann’s disease, degenerative scoliosis, Schmorl’s nodes and Cupid’s bow deformity (smooth developmental curvature of the inferior endplate of lumbar vertebra).22 Possible reasons for underdiagnosis of VFs by non-musculoskeletal radiologists include focusing on other acute findings, lacking specialist knowledge about osteoporosis/osteoporotic VFs or simply ignoring osteoporotic VFs completely.23

Vertebrae are included on many plain films when there is no clinical suspicion of VF. Examples include abdominal radiographs for patients with abdominal pain or chest radiographs in patients with cardiorespiratory symptoms. Less commonly, the vertebrae are incidentally imaged during barium investigations, interventional, cardiac and fluoroscopic procedures. Even if not
performed to outrule a VF, each imaged vertebra should be carefully evaluated to ensure no underlying occult VF. Despite the obvious opportunity to diagnose VFs in this way, there is a paucity of published literature in the area. The most studied radiographic technique to incidentally diagnose VFs is the chest radiograph. In a large study of over 10,000 post-menopausal females who underwent a lateral chest X-ray, 41% of radiologists who identified a VF failed to document it in the report summary, and only 36% were prescribed bone protection on discharge. In a smaller retrospective review of chest X-rays of post-menopausal females, Gehlbach showed that 14.1% had a moderate or severe VF visible on chest radiograph. Unfortunately, less than one-quarter of visible VFs were referenced in the radiologists’ summary, and only one-seventh of these patients received a discharge diagnosis of VF. As a result, only 18% of patients were discharged with appropriate medical therapy for underlying osteoporosis. The lateral chest radiograph on elderly patients is an opportunity to incidentally diagnose VFs by assessing vertebral bodies and clearly reporting them in the final summary. Despite their importance in initial investigation for suspected VF, many patients with VFs will have no morphological change on plain films. It is important not to dismiss patient symptoms based on normal radiographs since many patients with normal plain films may only have acute changes detectable on MRI. Loss of vertebral height may not be evident at time of acute symptoms but can be evident on subsequent follow-up radiograph.

MRI

MRI is a time-intensive imaging modality with relative contraindications such as claustrophobia, presence of a non-conditional pacemaker and first trimester of pregnancy. MRI has a sensitivity of 100% in detecting spinal trauma and is an excellent method to diagnose and assess VFs. MRI has a sensitivity and specificity of up to 82 and 98%, respectively, for distinguishing osteoporotic VFs from other types of fracture. In addition to identifying a VF, MRI may also diagnose other uncommon causes for back pain such as infection or malignancy, and also allows assessment of spinal ligaments, spinal cord, surrounding cerebrospinal fluid and meninges. The short tau inversion recovery (STIR) sequence is particularly useful as it nullifies marrow fat signal over a large body area such as the entire vertebral column. STIR sequences are helpful to differentiate benign osteoporotic VFs from those caused by malignancy. The presence of marrow oedema recognised as high signal on fluid sensitive STIR or T2 weighted fat-saturated sequences indicates recent fracture. Marrow oedema is absent in a chronic VF. Benign VFs typically are seen as linear low T1 signal. Malignancy or infection in contrast cause diffuse nonlinear replacement of the normal marrow of the vertebra. For every MRI study performed, initial localiser sequences are utilised by radiographers to plan image acquisition. These localisers are obtained from thick slices and are not suitable for diagnostic detail but do represent an opportunity to diagnose a VF when not suspected. The importance of reviewing localisers for fractures with strong interobserver agreement to detect VFs in the thoracic and lumbar spine has been reported. In another study, musculoskeletal radiologists examined 856 localisers of patients undergoing breast MRI. The authors concluded that 8.9% of patients had a VF visible on the MRI localiser, but none were documented in the final report. MRI localisers are a quick and reliable method of diagnosing VFs when not suspected and may negate the necessity for further imaging or using ionising radiation.
CT uses high doses of ionising radiation to acquire images. CT imaging is available 24 h in most tertiary hospitals and offers almost instant acquisition of images. CT has excellent sensitivity and specificity for identifying VFs at 100 and 97%, respectively. CT of the spine may be requested when a VF is clinically suspected and when the radiograph is normal. Of note, a non-displaced fracture accompanied by marked osteopaenia, may not be evident on CT. In patients with known VF, CT can help to provide additional information such as stability of the fracture and protrusion of bone fragments. CT can also aid with clinical decisions such as patient suitability for surgical intervention or vertebroplasty.

The majority of CTs are performed for clinical indications not specifically related to identification of VFs. Examples include cardiac CT, CT pulmonary angiogram and CT thorax to evaluate for thoracic pathology and CT kidneys, ureters and bladder, CT abdomen/pelvis, CT colonography and CT peripheral angiograms/venograms performed to identify intra-abdominal pathology. Vertebral morphology, particularly on sagittal reformats is well visualised on these CT studies. Modern CT scanners can display vertebrae in the region imaged in excellent bony detail in coronal, sagittal and axial reformats without the requirement for further imaging or radiation exposure to the patient. Of these, the sagittal reconstructions are particularly important to diagnose VFs (Figure 4). Despite the ability to utilise CT to diagnose occult VFs, CT is often not effectively exploited in this way. A New Zealand study retrospectively reviewed sagittal reconstructions of CT abdomen or thorax in patients over 65 years. 22 of 175 patients had a VF visible on sagittal reconstruction, and 77% of these had previously undiagnosed VF. The authors concluded that reviewing reformatted CT of the abdomen and pelvis improved diagnosis of VFs but are frequently not reported—thereby missing an opportunity to treat with appropriate medical therapy to reduce future osteoporotic fractures and associated mortality.

Similar to localisers in MRI, CT scout views are obtained prior to final image acquisition. These use low levels of radiation to acquire two-dimensional images which are used to plan the final CT image. Lateral CT scout views may show fractures not visible on axial CT images. One study of 300 CT scans involving the thoracic and lumbar spines demonstrated the sensitivity and specificity of diagnosing VFs on scout views to be 98.7 and 99.7%, respectively. The authors concluded that scout views should be used to evaluate for VFs on CTs performed for other clinical indications.

Skeletal scintigraphy (bone scans)

$^{99m}$Tc is a radioisotope which can be bound to methylene diphosphonate (MDP) and injected intravenously. The radioisotope travels through the patient’s bloodstream and binds to remodelling bone. 3 h after injection, patients are placed on a gamma camera which identifies bony hotspots where $^{99m}$Tc has accumulated. 80% of VFs are visible as hotspots, usually linear in morphology, at 24 h following injury and almost all return to normal within 2 years.

The major limitation of bone scans is their poor specificity. The most common indication for performing bone scans is to identify osseous metastatic disease in patients with known primary malignancy. However, bone scans are also utilised to identify...
occult fractures or osteomyelitis. Due to their non-specific nature, hotspots can also be caused by degenerative changes. Therefore, a nuclear medicine specialist may have diagnostic difficulty in distinguishing the exact cause of a hotspot. For this reason, bone scans (Figure 5) are often reported in conjunction with other available imaging such as MRI, CT (Figure 6) or plain films.

**DISCUSSION**

Osteoporosis is an increasing public health concern and predisposes patients to VFs. Prompt diagnosis and early intervention with appropriate medical treatment is imperative. The literature shows a failure by radiologists to report incidental VFs, and many which are diagnosed are never transcribed into the patient discharge summary. Untreated and undiagnosed VFs can significantly impact on a patient's quality of life and life expectancy. These patients suffer intolerable pain, lose their independence, suffer psychologically due to fear of isolation, many require polypharmacy for pain control, and all are at substantive risk of future osteoporotic fractures. The mid-thoracic region and thoracolumbar junction are the most commonly affected areas and often lead to spinal kyphotic deformities. Kyphosis predisposes to loss of balance, further muscle fatigue, further degenerative changes at adjacent intervertebral joints, restrictive lung disease, inability to work and loss of earnings.36

Unfortunately, many radiologists use equivocal language such as ‘loss of height’ or ‘wedging’ to describe VFs. This terminology is ambiguous for referring physicians and patients and can contribute to inappropriate management of underlying osteoporosis. An alternative strategy, described by Gehlbach et al is the semi-quantitative method for grading VFs. Even amongst inexperienced observers, the SQ method for grading fractures demonstrates high levels of agreement.16 Clearly stating the existence of an osteoporotic fracture in this way may improve the percentage of patients being discharged on appropriate medical therapy.

A number of imaging techniques performed for various clinical indications may show VFs in their field of view. However, there is underreporting of VFs which are clearly visible on lateral chest radiographs, MRI localisers and CT scout views. Failure to request bony sagittal reformats from CT scans often renders VFs invisible, even to experienced musculoskeletal radiologists. The term ‘inattentional blindness’ refers to an inability to notice unexpected events when immersed in an alternative task. In one experiment, 83% of expert radiologists failed to recognise a gorilla drawn onto a stack of CT images when they were focusing on finding pulmonary nodules.37 Another phenomenon coined ‘satisfaction of search’ refers to a relative difficulty in identifying further pathological findings following identification of another significant abnormality.38 Both are relevant to radiologists when searching for potentially life-threatening pathology on X-ray, MRI or CT and VFs can easily be overlooked.

Dedicated education programmes delivered to radiologists and internal medical physicians may help to improve the diagnosis and management of VFs. In one study, recognition of VFs amongst internists almost doubled from 22 to 43% following provision of basic lectures, posters and flyers. The same study demonstrated a significant increase in patients discharged on osteoporosis treatment from 11 to 40%.39 In another study, there was a marked improvement in the ability of a radiology resident to correctly identify VFs after undergoing specific teaching.40

**CONCLUSION**

In conclusion, VFs are a major health concern in an era of aging population. Many factors have contributed to poor diagnosis and treatment of VFs. When identified, ambiguous
Figure 5. There are foci of increased radioisotope uptake in the mid-thoracic spine on bone scan.

Figure 6. Sclerotic pathological wedge compression fractures of T6 and T7 secondary to metastatic disease on sagittal reconstruction of staging CT thorax in patient with primary non-small cell lung cancer correlating with uptake in bone scan in Figure 5.
terminology should be avoided and SQ method employed. Irrespective of clinical indication or imaging modality, a high index of suspicion for VFs should be employed at all times. Basic education programmes delivered to radiologists and internists have been shown to improve diagnosis and treatment of VFs.

REFERENCES

1. Zhao JG, Zeng XT, Wang J, Liu L. Association between calcium or vitamin D supplementation and fracture incidence in community-dwelling older adults: a systematic review and meta-analysis. JAMA 2017; 318: 2466–62. https://doi.org/10.1001/jama.2017.19344

2. Chrischilles EA, Cooper C, Lane AW, Riggs BL. Perspective. How many women have osteoporosis? J Bone Miner Res 1992; 7: 1005–10. https://doi.org/10.1002/jbmr.5650070902

3. (N.d.). Foundation no.National Osteoporosis Foundation Website. Available from: https://www.nof.org/preventing-fractures/general-facts/2019

4. Glinkowski W, Narloch J, Krasuski K, M, Schaufler T, Senn C, et al. Fracture incidence of living with an osteoporotic vertebral compression fracture. Osteoporos Int 2016; 27: 1279–36. https://doi.org/10.1007/s00198-015-3445-y

5. Kelly MA, McGowan B, McKenna MJ, Bennett K, Carey JJ, Whelan B, et al. Emerging trends in hospitalisation for fragility fractures in ireland. Ir J Med Sci 2018; 187: 601–8. https://doi.org/10.1007/s11845-018-1743-z

6. Lippuner K, Popp AW, Schwab P, Gitlin M, Schaufler T, Senn C, et al. Fracture hospitalizations between years 2000 and 2007 in switzerland: a trend analysis. Osteoporos Int 2011; 22: 2487–97. https://doi.org/10.1007/s10994-010-1487-8

7. Schousboe JT. Epidemiology of vertebral fractures. J Clin Densitom 2016; 19; S1094-6950(15)00168-7: 8–22. https://doi.org/10.1016/j.jocd.2015.08.004

8. Ballane G, Cauley JA, Luckey MM, El-Hajj Fuleihan G. Worldwide prevalence and incidence of osteoporotic vertebral fractures. Osteoporos Int May 2017; 28: 1531–39. https://doi.org/10.1007/s00198-017-3909-3

9. Arima K, Abe Y, Nishimura T, Okabe T, Tomita Y, Mizukami S, et al. Association of vertebral compression fractures with physical performance measures among community-dwelling japanese women aged 40 years and older. BMC Musculoskelet Disord April 28, 2017; 18(1): 176. https://doi.org/10.1186/s12891-017-1531-3

10. Svensson HK, Olofsson EH, Karlsson J, Hansson T, Olsson LE. A painful, never ending story: older women’s experiences of living with an osteoporotic vertebral compression fracture. Osteoporos Int 2016; 27: 1279–36. https://doi.org/10.1007/s00198-015-3445-y

11. Cauley JA, Thompson DE, Ensrud KC, Scott JC, Black D. Risk of mortality following clinical fractures. Osteoporos Int 2000; 11: 556–61. https://doi.org/10.1007/s001980010075

12. Delmas PD, Genant HK, Crans GG, Stock JL, Wong M, Siris E, et al. Severity of prevalent vertebral fractures and the risk of subsequent vertebral and nonvertebral fractures: results from the more trial. Bone 2003; 33: 522–32. https://doi.org/10.1016/s8756-3282(03)00241-2

13. Klotzbuecher CM, Ross PD, Landsman PB, Abbott TA 3rd, Berger M. Patients with prior fractures have an increased risk of future fractures: a summary of the literature and statistical synthesis. J Bone Miner Res 2000; 15: 721–39. https://doi.org/10.1359/jbmr.2000.15.4.721

14. Crandall CJ, Newberry SJ, Diamant A, Lim HC, Whitehouse RW. Under-reporting of vertebral fractures is a worldwide problem: the impact study. J Bone Miner Res 2005; 20: 557–63. https://doi.org/10.1359/jbmr.041214

15. Freedman KB, Kaplan FS, Billsker WB, Strom BL, Lowe RA. Treatment of osteoporosis: are physicians missing an opportunity? J Bone Joint Surg Am 2000; 82: 1063–70. https://doi.org/10.2106/00004632-200008000-00001

16. Gehlbach SH, Bigelow C, Heimsdottir M, May S, Walker M, Kirkwood JR. Recognition of vertebral fracture in a clinical setting. Osteoporos Int 2000; 11: 577–82. https://doi.org/10.1007/s001980070078

17. Delmas PD, van de Lameriget L, Watts NB, Eastell R, Genant H, Grauer A, et al. Underdiagnosis of vertebral fractures is a worldwide problem: the impact study. J Bone Miner Res 2005; 20: 557–63. https://doi.org/10.1359/jbmr.041214

18. Bartalena T, Rinaldi MF, Modolon C, Braccioli L, Sverzellati N, Rossi G, et al. Incidental vertebral compression fractures in imaging studies: lessons not learned by radiologists. World J Radiol 2010; 2: 399–404. https://doi.org/10.4329/wjr.v2i110.399

19. Williams AL, Al-Busaidi A, Sparrow PJ, Adams JE, Whitehouse RW. Under-reporting of osteoporotic vertebral fractures on computed tomography. Eur J Radiol 2009; 69: 179–83. https://doi.org/10.1016/j.ejrad.2007.08.028

20. Genant HK, Wu CY, van Kuijk C, Nevitt MC. Vertebral fracture assessment using a semiquantitative technique. J Bone Miner Res 1993; 8: 1137–48. https://doi.org/10.1002/jbmr.5650080915

21. Niimi R, Kono T, Nishihara A, Hasegawa M, Matsumine A, Kono T, et al. Efficacy of the dynamic radiographs for diagnosing acute osteoporotic vertebral fractures. Osteoporos Int 2014; 25: 605–12. https://doi.org/10.1007/s00198-013-2456-9

22. Griffith JF. Identifying osteoporotic vertebral fracture. Quant Imaging Med Surg 2015; 5: 592–602. https://doi.org/10.3978/j.issn.2223-4292.2015.08.01

23. Li Y, Yan L, Cai S, Wang P, Zhiang H, Yu H. The prevalence and under-diagnosis of vertebral fractures on chest radiograph. BMC Musculoskelet Disord 2018; 19: 235. https://doi.org/10.1186/s12891-018-2171-y

24. Morris CA, Carrino JA, Lang P, Solomon DH. Incidental vertebral fractures on chest radiographs. J Gen Intern Med 2006; 21: 352–56. https://doi.org/10.1111/j.1525-1497.2006.00395.x

25. Adams JE. Opportunistic identification of vertebral fractures. J Clin Densitom 2016; 19; S1094-6950(15)00174-2: 54–62. https://doi.org/10.1016/j.jocd.2015.08.010

26. Terakado A, Orita S, Inage K, Kubota G, Kanzaki T, Mori H, et al. A clinical prospective observational cohort study on the prevalence and primary diagnostic accuracy of occult vertebral fractures in aged women with acute lower back pain using magnetic resonance imaging. Pain Res Manag 2017; 2017: 9265259. https://doi.org/10.1155/2017/9265259

27. Abdel-Wanis ME, Solymy MM, Hasan NMA. Sensitivity, specificity and accuracy of magnetic resonance imaging for differentiating vertebral compression fractures caused by malignancy, osteoporosis, and infections. J Orthop Surg (Hong Kong) 2011; 19: 145–50. https://doi.org/10.1177/230949901101900203

28. Ogura A, Hayakawa K, Maeda F, Saeki F, Syukutani A, Shibutani S, et al. Differential diagnosis of vertebral compression fracture using in-phase/opposed-phase and short
ti inversion recovery imaging. *Acta Radiol* 2012; 53: 450–55. https://doi.org/10.1258/ar.2012.110524

29. Kaniewska M, de Beus JM, Alhheim F, Mameghani A, Eid K, Kubik-Huch RA, et al. Whole spine localizers of magnetic resonance imaging detect unexpected vertebral fractures. *Acta Radiol* 2019; 60: 742–48. https://doi.org/10.1177/0284185118796673

30. Bazzocchi A, Spinnato P, Garzillo G, Ciccarese F, Albisinni U, Mignani S, et al. Detection of incidental vertebral fractures in breast imaging: the potential role of mr localisers. *Acta Radiol* 2012; 22: 2617–23. https://doi.org/10.1007/s00330-012-2521-3

31. Berry GE, Adams S, Harris MB, Boles CA, McKernan MG, Collinson F, et al. Are plain radiographs of the spine necessary during evaluation after blunt trauma? accuracy of screening torso computed tomography in thoracic/lumbar spine fracture diagnosis. *J Trauma* 2005; 59: 1410–13. https://doi.org/10.1097/01.ta.0000197279.97113.0e

32. Müller D, Bauer JS, Zeile M, Rummeny EJ, Link TM. Significance of sagittal reformations in routine thoracic and abdominal multislice ct studies for detecting osteoporotic fractures and other spine abnormalities. *Eur Radiol* 2008; 18: 1696–1702. https://doi.org/10.1007/s00330-008-0920-2

33. Chan PL, Reddy T, Milne D, Bolland MJ. Incidental vertebral fractures on computed tomography. *N Z Med J* 2012; 125: 45–50.

34. Bazzocchi A, Fuzzi F, Garzillo G, Diano D, Rimondi E, Merlino B, et al. Reliability and accuracy of scout ct in the detection of vertebral fractures. *Br J Radiol* 2013; 86(1032): 20130373. https://doi.org/10.1259/bjr.20130373

35. Matin P. The appearance of bone scans following fractures, including immediate and long-term studies. *J Nucl Med* 1979; 20: 1227–31.

36. Al-Sari UA, Tobias J, Clark E. Health-related quality of life in older people with osteoporotic vertebral fractures: a systematic review and meta-analysis. *Osteoporos Int* October 2016; 27: 2891–2900. https://doi.org/10.1007/s00198-016-3648-x

37. Drew T, Võ ML-H, Wolfe JM. The invisible gorilla strikes again: sustained inattentional blindness in expert observers. *Psychol Sci* 2013; 24: 1848–53. https://doi.org/10.1177/0956797613479386

38. Berbaum KS, Franken EA Jr, Dorfman DD, Miller EM, Caldwell RT, Kuehn DM, et al. Role of faulty visual search in the satisfaction of search effect in chest radiography. *Acad Radiol* 1998; 5: 9–19. https://doi.org/10.1016/s1076-6332(98)80006-8

39. Caser P, Uebelhart B, Gaspoz JM, Ferrari S, Louis-Simonet M, Rizzoli R. Targeted education improves the very low recognition of vertebral fractures and osteoporosis management by general internists. *Osteoporos Int* 2006; 17: 965–70. https://doi.org/10.1007/s00198-005-0064-z

40. Gruber M, Dingcs J, Müller D, Baum T, Rummeny EJ, Bauer J. Impact of specific training in detecting osteoporotic vertebral fractures on routine chest radiographs. *Rofo* 2013; 185: 1074–80. https://doi.org/10.1055/s-0033-1335230