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A Multi-method Luminescence Dating of the Palaeolithic Sequence of La Ferrassie Based on New Excavations Adjacent to the La Ferrassie 1 and 2 Skeletons

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Abstract

A new interdisciplinary project was initiated to excavate a portion of the Palaeolithic site of La Ferrassie left intact by earlier excavations. One of the aims of this project was to provide chronological information on the succession of Middle and Upper Palaeolithic layers, as well as on the skeletons unearthed by Capitan and Peyrony in the early 1900’s. We report here preliminary results on the lithics, faunal remains, site formation processes, and on the stratigraphic context of the La Ferrassie 1 and 2 skeletons that were found adjacent to our excavations. Finally, results from luminescence dating of the sediments and a preliminary set of radiocarbon ages are presented. Quartz OSL, both at the multi-grain and single-grain levels of analysis, and post-IR IRSL of feldspar at various stimulation temperatures are compared. The quartz/feldspar comparison revealed a bleaching problem for the quartz OSL (and the feldspar pIRIR signals) from Layer 2; as a consequence, the age of this Layer was determined using a minimum age model.

A Mousterian industry with bifaces, at the base of the sequence, has been dated between 91 ± 9 and 44 ± 3 ka. The Ferrassie Mousterian layers are attributed to MIS 3, between 54 ± 3 and 40 ± 2 ka, and thus appear very late in the final Middle Palaeolithic of the region; furthermore, these ages constrain the chronology of the La Ferrassie 1 and 2 skeletons, which have been attributed to one of these Ferrassie Mousterian layers. The Châtelperronian layer is dated to 42 ± 3 ka and the Aurignacian to 37 ± 2 ka. Implications of the ages for the La Ferrassie 1 and 2 skeletons, and for the variability of late Mousterian, are discussed.

Key-words: OSL dating; post-IR IRSL dating; single grain; Middle Palaeolithic; Mousterian.
1. Introduction

For over a century, the Palaeolithic site of La Ferrassie has been the focus of numerous excavations and studies. Located in a small tributary valley of the Vézère River in southwest France (Fig. 1), this site has yielded rich Middle and Upper Palaeolithic lithic assemblages, faunal remains, as well as two nearly complete Neanderthal skeletons and additional partial skeletons. The human remains make La Ferrassie one of the more important sites for the study of Neanderthal morphology and one of the more important data sets when discussing Neanderthal treatment of the dead (Capitan and Peyrony, 1921a, 1922; Peyrony, 1934, 1921; Vandermeersch, 1976; Harrold, 1980; Heim, 1982a, 1982b, 1976, 1968; Chase and Dibble, 1987; Bar Yosef, 1988; Smirnov, 1989; Straus, 1989; Gargett, 1989, 1999; Duday et al., 1990; Lindly and Clark, 1990; Binant, 1991a, 1991b; Defleur, 1993; Riel-Salvatore and Clark, 2001; Pettitt, 2002; Maureille and Vandermeersch, 2007; Langley et al., 2008). The site also gives its name to one variant of the Mousterian rich in scrapers made on Levallois flakes, and the Aurignacian portion of the sequence was originally used to define its stages (Peyrony, 1934; Bordes, 1957, 1961; Sonneville-Bordes, 1960; Delporte, 1984).

Thus there are many reasons why La Ferrassie has been studied in detail. However, aside from some radiocarbon ages completed on Upper Palaeolithic layers from excavations led by Delporte in the 1970s, this key site has remained undated. To address this lack of chronology, we undertook a new interdisciplinary excavation (led by AT) that started in 2010. The first season resulted in the discovery of new deposits in the western portion of the site, very close to the find location of the La Ferrassie 2 skeleton; a great part of the La Ferrassie sequence is represented in these new deposits. Since then, new excavations have been undertaken (i) to re-evaluate the stratigraphic sequence of this area, with a particular emphasis on site formation processes and their implications for the context of the Neanderthal skeletons, (ii) to re-assess the techno-complexes and in particular the eponymous Ferrassie
Mousterian in the current context of Mousterian occupations in SW France (Turq et al., 2008a), and
finally (iii) to determine the chronology of the different occupations.

Here we report results derived from the use of various luminescence dating methods and compare them
with preliminary radiocarbon ages. The results give a coherent chronology that (i) places the base of the
sequence at approximately 90 ka, (ii) shows that the Mousterian with bifaces is older than most similar
industries in southwest France, (iii) demonstrates that the Ferrassie Mousterian is quite late in the
sequence, and (iv) shows that the La Ferrassie 2 skeleton is most likely dated to 43 ± 3 ka and cannot be
older than 54 ± 4 ka, which does not confirm expected older ages (e.g., Turq et al., 2008b).

2. Presentation of the site

The first and most extensive excavations of the so-called Grand Abri or Large Shelter at La Ferrassie
were led by Peyrony and Capitan over a century ago (1934; see also Capitan and Peyrony, 1909; 1910;
1912; 1921; Breuil, 1921). They excavated a series of trenches starting near the road and progressively
working towards the east or deeper into the supposed shelter (see below). They defined an
archaeological sequence of Mousterian with bifaces, Ferrassie Mousterian, Châtelperronian (initially
Lower Perigordian), Aurignacian and Gravettian (initially Upper Perigordian). This work was followed,
from 1968 to 1973, by more limited excavations by Delporte (1984) focused almost exclusively on the
eastern, interior part of the site. As a result of this work, the site today can be divided into two parts: an
inner part to the east where the large section cleaned by Delporte is now preserved, and an outer part
or entrance in the northwest sandwiched between the modern road and rock wall that forms an
enclosure around the inner part (Fig. 2). Prior to the new excavations, it was uncertain whether any
intact deposits remained in this portion of the site, but the general assumption was that it had been
entirely excavated by Peyrony and Capitan and subsequently back filled (Laville, 2007).
In 2010 we opened this part of the site (Fig. 2) and encountered extensive, previously unexcavated, deposits corresponding to the base of La Ferrassie sequence, as previously defined. This area (our Sector I) extends to the underlying bedrock. In the northern portion of the new excavation, we also encountered what appears to be the upper part of the classic sequence. This became Sector II because the stratigraphic link with the base of the sequence was not clear at the time. In 2011 we were able to confirm the presence of a complete sequence in Sector II and began work to link Sectors I and II across the western part of the excavated area where substantial intact deposits remained. This western area became Sector III. In 2012 we expanded work in all three sectors, and as a result we were able to link the separate stratigraphic sequences into a single, unified sequence that is the basis of what is reported here. However, there is still some lateral variation in the upper layers, particularly in Layer 7, which remains to be fully described. Thus for Layer 7 we retain a western and eastern separation (former Sector III and II, respectively).

2.1. Geology of the site

The shelter is part of an extensive tunnel-like karstic system at the intersection of two roughly perpendicular valleys (Fig. S1). Table 1 provides a simplified description of the geology and lithics of the stratigraphic units based on the new excavation (Fig. 3). Correspondences with Peyrony’s original stratigraphy and OSL/radiocarbon sample provenance are also indicated in Table 1. In general, the sediments are fluvially-deposited sands and gravels overlain by slope deposits; Turq et al. (2012; see chapter by Goldberg et al.) studied the nature of these deposits in the current part of the excavation. The transition from fluvial to slope deposits occurs between Layers 2 and 3. Fig. S2 shows the average grain size distribution for the OSL samples from La Ferrassie determined by laser diffraction grain-size analysis, after the >1 mm fraction has been removed. Sand and silt contribute respectively about 75 and 25% of the total, and only minor variations are present from layer to layer.
2.2. Lithics

Assemblage sizes from the new excavations are still small but sufficient to generally confirm the presence of the sequence as defined by Peyrony and Capitan and refined by Delporte. A Middle Palaeolithic industry is present at the base of the sequence, in Layers 1 and 2 (N=682 artefacts > 2.5cm), with discoidal and Levallois technology. A partial biface comes from Layer 1 and a complete biface from Layer 2. The biface, in terms of size and shape, fits within the variability defined for the Mousterian of Acheulian Tradition (MTA). However, aside from bifacial manufacture flakes, no other specifically MTA elements are present. Among retouched tools are scrapers, one pseudo-Levallois point, notches, and denticulates. Overall the assemblages are similar to the Mousterian with bifaces described by Peyrony (1934). The Layer 3 assemblage (N=434) has relatively few Levallois flakes, more notched tools than scrapers, one biface, and some evidence for small flake production in the form of truncated-faceted pieces. Layers 4 and 5 (N=620) are consistent with a Ferrassie type Mousterian, based on a Levallois technology with a predominance of single, double, and convergent scrapers along with truncated-faceted pieces, notches, and denticulates. The Layer 6 assemblage is still quite small (N=164 plus 91 pieces from the 1.5-25 mm fraction); nevertheless, it has yielded artefacts typical of the Châtelperronian (see Chiotti in Turq et al., 2012). The assemblage includes nine retouched bladelets including three truncated bladelets and one Dufour bladelet (sub-type Roc-de-Combe). There are also two Châtelperronian point fragments and one other blade fragment with abrupt retouch that could also be a point fragment. The assemblage also includes one end-scraper and three burins. Layer 7 (N=908 plus 376 pieces from the 1.5-25 mm fraction) is characterized by an Aurignacian lithic production including bladelets produced from carinated scrapers and at least one Dufour bladelet, of the sub-type Roc-de-Combe. Thus far, however, the assemblage does not allow us to determine with certainty which type of Aurignacian is present.
2.3. Faunal remains

The faunal diversity of the current sample is in agreement with what has been previously reported for the sequence (Peyrony, 1930) and further supports the stratigraphic correlation proposed in Table 6. In the older part of the sequence (Layers 1 through 5b), large bovids (aurochs [*Bos primigenius*] and/or bison [*Bison priscus*]) dominate the assemblages, and when these elements can be identified more specifically, they are from bison. Red deer (*Cervus elaphus*) and roe deer (*Capreolus capreolus*) are also common in these older assemblages. A few elements of reindeer (*Rangifer tarandus*) and horse (*Equus simplicidens*) are also found. Specifically, Layer 3 has also yielded remains of woolly mammoth (*Mammuthus primigenius*) and woolly rhinoceros (*Coelodonta antiquitatis*). In Layer 4, the presence of wild boar (*Sus scrofa*) together with red deer and bovids may indicate more temperate conditions, but reindeer are present there, too. Many of the assemblages contain species characteristic of a diversity of habitats: reindeer are usually considered part of a colder, drier, more open environment whereas roe deer are considered part of a warmer, more wooded environment. The topographic relief of the region may have created microhabitats that allowed the persistence of a diversity of species. Time averaging over minor climatic fluctuations may also contribute to the variability. Reindeer clearly dominate the upper part of the sequence (Layers 6 and 7); they are almost exclusive in the large sample from layer 7. Zooarchaeological analyses have demonstrated an abundance of cutmarks, burned bones, and breakage of fresh bones for marrow exploitation throughout the La Ferrassie sequence; bone surfaces are generally well-preserved allowing for sufficient reading of the modifications. Carnivore remains, coprolites, and traces of carnivore activity such as chewed and digested bones are rare. Human activity was the main agent for the accumulation of faunal remains throughout the deposits.

2.4. Micromorphology and the stratigraphic position of the La Ferrassie 2 foot
An analysis of historical photographs of the Peyrony and Capitan excavations and of the various published and unpublished maps (Laville, 2007) in combination with our own excavations in the same area allow us to place the La Ferrassie 2 skeleton to within ~1-2 m of the stratigraphic section sampled here (Fig. 4c). The observations made by the original excavators and by the members of the scientific commission prior to removal of the skeleton attributed the skeleton to the base of Layer C (our Layer 4) and noted that it was covered by Layer D (our Layer 5). In terms of the spatial association between La Ferrassie 1 and 2, Peyrony clearly thought they were stratigraphically very close. La Ferrassie 1 is described as lying almost on the yellowish layer and sometimes on the reddish layer underneath ("... reposant presque sur la couche jaunâtre et parfois rougeâtre sous-jacente ...") (Peyrony 1934: 24). He describes La Ferrassie 2 as occurring at about the same level and only 50 cm from La Ferrassie 1 ("... sensiblement au même niveau et dans le même axe que le premier squelette, mais en position inverse, puisque les deux têtes n'étaient qu'a 0 m. 50 l'une de l'autre ...") (Peyrony, 1934: 26).

In an attempt to establish/constrain the stratigraphic position of La Ferrassie 2 in the newly excavated sequence, we (AT, PG, and SM) examined the foot of the skeleton stored in the Musée de l'Homme, Paris. A large block of sediment is still attached to the base of the right foot of this specimen, and it was examined along with a complete suite of thin sections collected from of our excavations. The purpose of the investigation was to (i) examine the sediments adhering to the foot to compare them with those from our excavations, and thereby (ii) determine the stratigraphic layer from which it was excavated.

The sediments attached to the foot consist of consolidated (by the Musée) yellow brown silty sands with burned and unburned bone and some lithics scattered throughout (Fig. 4a). When these sediments are compared to those exposed in our sequence (Fig. 4b), we observe a remarkably close correspondence to those of Layer 5; the same components and abundances are observed. Whereas
Layers 4 and 5 are generally similar, the latter is noticeably richer in bone (Fig. 4b), which makes it a closer match to the sediments attached to the hominin specimen. At the same time, we can say that Layer 6 contrasts greatly with Layers 4 and 5 both in terms of the sediments and their cultural content. The sediments preserved under the foot do not match Layer 6 nor do they look like a mix of Layers 5 and 6 as might happen if a burial pit was excavated from 6 into 5 and then refilled. Thus association with Layer 6 can be excluded, and it remains to be determined whether Layer 4 or 5 is the better candidate for the sediment under the skeleton.

The question as to whether the skeletons represent burials, and the definition of the latter, are beyond the scope of this study. Our ongoing study of the processes leading to the formation of the site and the documentation of the original discoveries is expected to shed more light on this question.

3. Material and methods

3.1. Sampling

Given the importance of the site, we decided on a multi-method dating programme. In the almost complete absence of heated stones, only sediments were sampled for luminescence dating, but these covered the entire stratigraphic sequence. Some samples were measured in two different laboratories to provide inter-laboratory comparisons. Since an overlap with the range of radiocarbon dating was considered possible, bone samples were also selected for radiocarbon dating from the top of the Mousterian (Layer 5b) and from Layer 6 (Châtelperonian) in Sector II to provide additional age control.

The two main series of sediment samples were collected from Sectors I and II (Figs. 2, 3; Table 1) in 2010, during the first season of excavation; the samples were collected at night under dim orange lighting. After cleaning the sections, ~1 cm of exposed sediment was removed before actual sampling. At that stage, Sector III was not yet excavated, so the two sampled sections were not linked in stratigraphy.
In the northwest area (Sector II), the Aurignacian, Châtelperronian, and Ferrassie Mousterian Layers were sampled (Layers 4-7 – see Table 1 for the relationship between luminescence samples and archaeological layers). One sample was taken per layer, except for the Mousterian layers; bearing in mind the presence of Neanderthal skeletons in this part of the sequence and the closeness of the La Ferrassie 2 skeleton (~1-2 m east of the sampling location), paired samples were taken from these layers. In this area, only a thin wall of sediments remained from the Peyrony and Capitan excavations. Because of this, the sediment samples were taken close to the bedrock wall (less than 20 cm, possibly 10 cm); this led to additional complexity in the assessment of the gamma dose-rates (see section 4.2 – dose-rates). A further six samples were taken from the trench in Sector I, two from each Mousterian Layer (1, 2 and 3).

The excavations in Sector III in 2012 provided a stratigraphic link between Sectors I and II, and led to the current unified stratigraphy (Fig. 3). It was, therefore, decided to sample Layer 5 in this sector. Finally, another sample, also taken in 2012, comes from Layer 3 at the south end of Sector I. Both samples were collected during the day under opaque black covering. Fig. S3 shows an interactive 3D model of the excavated portion of the site, indicating approximate sampling location (only approximate since the 3D model corresponds to the state of the site two years after the first sampling campaign).

Luminescence dating methods are applicable to different minerals (e.g., quartz and feldspar) measured using different signals and displaying different characteristics in terms of dose-rates, equivalent doses, signal resetting rates when exposed to daylight (hereafter called bleaching rates), etc. They can thus be regarded as complementary tools when combined to study one site (e.g., Richter et al., 2009; Guérin et al., 2012a). For the study of La Ferrassie, we decided to compare ages obtained from both quartz and feldspar. As the bleaching rates of the luminescence signals of these two minerals are very different (by orders of magnitude), the apparent ages can be used to assess the likelihood of age
overestimation resulting from incomplete bleaching at the time of burial (Murray et al., 2012). This is particularly important for slope deposits filling a rockshelter.

3.2. Luminescence

3.2.1. Sample preparation and instrumentation

Chemical preparation

The sediment samples were first split into different portions for grain size analysis, high-resolution gamma spectrometry and mineral separation (performed entirely under controlled low-level yellow light suitable for luminescence samples). Prior to mineral separation the samples were wet sieved to isolate the 180-250 µm sand fractions. This sieving step was undertaken before any chemical treatment and in particular before treatment with hydrochloric acid (HCl), so that any mm to cm scale calcareous aggregates and/or gravels were less likely to release quartz grains of the size range of interest. After sieving, the samples were treated with HCl (10%) to remove carbonates, and then with hydrogen peroxide (H₂O₂) to remove organic contaminants; both treatments were continued until no reaction was visible. The sieved and disaggregated grains were subsequently treated with dilute hydrofluoric acid (HF, 10%) for 30 minutes to clean all grain surfaces and to etch the outer rim of feldspar, and finally rinsed with 10% HCl. Two aqueous solutions of sodium heteropolytungstates (densities 2.58 and 2.62 g.cm⁻³) were used to isolate K-rich feldspar fractions (<2.58 g.cm⁻³) and quartz (>2.62 g.cm⁻³). The quartz fraction was then etched with HF (40%) for 40 minutes to remove the outer portion of the grains affected by alpha irradiation. After etching, any fluoride contaminants were removed by rinsing with 10% HCl. This fraction was then re-sieved at 180 µm for further analysis and in particular for single grain measurements; this latter step removes any small grains resulting for example, from the dissolution of residual feldspar in the quartz-rich fraction.
**Instrumentation**

Grains were mounted in 9 mm base diameter stainless steel cups using silicone oil. Aliquots of ~3 mm in diameter, comprising several hundreds of grains, were measured for both quartz and feldspar extracts. Luminescence measurements were made using Risø TL/OSL DA-15 and DA-20 readers; for quartz, blue (470 nm) light-emitting diodes (LED) were used with 7.5 mm Hoya U-340 detection filters; for feldspar, IR diodes emitting at 875 nm were used in combination with coupled Schott BG39 and Corning 7-59 detection filters (transmission 320–460 nm; Bøtter-Jensen et al., 2003; 2010). Each $^{90}$Sr/$^{90}$Y source was calibrated during the measurement period by measuring several aliquots of calibration quartz dosed with gamma rays (4.81 Gy) from a national secondary-standard $^{137}$Cs source; this calibration has been independently confirmed by Bos et al., 2006.

Single grains of quartz were measured using an automated Risø TL/OSL reader (DA 20) fitted with a single grain attachment (Duller et al., 1999; Bøtter-Jensen et al., 2000). The grains were loaded into aluminium single-grain discs; each disc contains 100 holes 300 µm in diameter and 300 µm deep, drilled on a 10x10 rectangular grid with 600 µm spacing between centres. A green laser (532 nm) was used with a 7.5 mm Hoya U-340 detection filter. To confirm that only one grain was loaded into each hole, the single grain discs were visually inspected using a microscope before measurement. The spatial homogeneity of the $^{90}$Sr/$^{90}$Y beta source (ID 155, produced prior to 2000) used for the single grain measurements presented here was determined using radiochromic film (GAFCHROMIC EBT2) and the coefficient of variation was found to be ~5%, which is unlikely to affect our dose distributions significantly. Using the approach developed by Lapp et al. (2012) to correct for spatial beta source heterogeneity, no significant changes in dose or scatter in our dose distributions could be detected, consistent with the observations of Sim et al. (2014) for the same source. Thus, the dose rate employed in further single grain calculations is based on the average over the whole disc.
Inter laboratory comparisons

Given the importance of the site, it was decided to conduct separate, blind comparisons of equivalent dose measurements on quartz aliquots between two laboratories: the Centre for Research in Physics Applied to Archaeology (CRPAA) in Bordeaux (France) and the Centre for Nuclear Technologies, Danish Technical University (DTU Nutech), Risø Campus (hereafter Risø). Where available, the numbers obtained by both laboratories will be provided.

3.2.2. Dose-rates

Quartz and feldspar grains accumulate dose in sediments because of the exposure to external alpha, beta, gamma and cosmic rays and particles; there are also internal contributions – especially in the case of K-rich feldspar. The internal dose-rate was assumed to be $0.06 \pm 0.03 \text{ Gy.ka}^{-1}$ for quartz (Mejdahl, personal communication to Murray, based on Mejdahl, 1987), and was calculated for feldspar using dose-rate conversion factors for potassium (Guérin et al., 2011) and assuming a potassium content equal to $12.5 \pm 0.5\%$ (Huntley and Baril, 1997). The self-dose fraction of the infinite-matrix dose-rate was taken from Guérin et al. (2012b). The contribution from Rb was calculated according to Readhead (2002) and Huntley and Hancock (2001). Direct potassium measurements of K-rich extracts using a recently-developed XRF system attached to a Risø luminescence reader confirmed the expected high potassium content. Finally, it was assumed that the HF etching had removed the alpha irradiated portions of the quartz and feldspar grains; therefore, no external alpha dose-rate was taken into consideration. Cosmic dose-rates were estimated from the estimated burial depth of the sediments according to Prescott and Hutton (1988) and contributed ~10% of the total dose-rates.

Beta dose-rates
Sediment was homogenised by crushing and sealed in plastic boxes containing ~10 g of samples. These sealed samples were then stored for at least three weeks to ensure radon build-up, before measurement using high resolution, low background gamma spectrometry. For each sediment sample, the potassium, uranium and thorium contents were measured; the state of equilibrium of the U-series has been evaluated. For that purpose, two effective U contents were calculated using weighted averages obtained from different gamma lines: the first, based on the top members of the chain, included the rays at 63 keV and 93 keV ($^{234}$Th), 143 keV and 186 keV ($^{235}$U) and 1001 keV ($^{234m}$Pa). The second value was calculated from the lines at 295 keV and 352 keV ($^{214}$Pb), and 609, 1120 and 1764 keV ($^{214}$Bi). For samples FER 7-14, both values agree at one sigma. For the remaining (samples FER 1-6), we calculated extreme scenarios: we calculated the dose-rate using either the top or the bottom chain value as representative of the whole U-series. Given the uranium contribution to the total dose-rates (~20-25 %), the differences between the extreme scenarios resulted in discrepancies of ~5% on average. Given the final uncertainties on the ages, we decided that a detailed analysis of the source of disequilibrium was not necessary. We thus calculated dose-rate values based on the pre- and post-Radon contents in the U decay series, using factors from Guérin et al. (2011).

External beta dose-rates for both quartz and feldspar were calculated from K, U, and Th contents using the factors from Guérin et al. (2011). Grain size attenuation factors were taken from Guérin et al. (2012b). For the effect of moisture on beta dose-rates, values provided by Nathan and Mauz (2008) were used, although they do not include geometric factors and are therefore likely to be underestimates (Guérin et al., 2012b). However, these moisture correction values only apply to the water content of the sediment (Zimmermann, 1971), which typically amounts to ~15% of the total (see below); errors in the moisture correction term will thus result in errors of less than a few % in the dose-rate. The effect of etching on the mean dose-rate was corrected for using values from Nathan (2011). The moisture content of the sediments during burial time must have been different from the values
measured immediately after sampling, because the stratigraphic sections had been exposed to air for several days before sampling and had dried out significantly. Measurements of saturation content yielded maximum values of ~25%, on average. We took 0.6±0.2 as a representative value for the fraction of saturation, leading to a moisture content equal to 15±5% of water content (expressed as mass of water/dry mass). The actual lifetime average water content cannot have been greater than saturation (25%), and it is inconceivable that it could have been (on average) less than 5% (average at the time of sampling). Thus the two standard deviation range of 5-25% covers all possible water content scenarios; the average water content between these two extreme scenarios represents our best guess regarding burial time.

Following Guérin et al. (2012b), the use of the infinite matrix assumption (Aitken, 1985) may be questioned, especially in light of the grain size analysis (Fig. S2) which shows that the sediment samples from La Ferrassie are mainly composed of sands, i.e. of grains of non-negligible dimensions compared to the range of electrons – whether of primary electrons from beta decay or of secondary from photoelectric/Compton interactions of primary photons. The two main dosimetric effects identified by Guérin et al. (2012b) are: (i) the inappropriate use of beta dose-rate attenuation factors (first defined by Mejdahl, 1979) since there is no homogeneous, infinite matrix containing the radioelements and surrounding the dated quartz/feldspar grains; (ii) the unaccounted self-dose to potassium feldspar, which are point like sources of non-negligible size compared to the range of beta particles. It has been shown by Guérin et al. (2012b), using numerical simulations, that for well-sorted sands in which potassium feldspar and quartz grains have the same grain-size distribution, these two effects compensate because the self-dose fraction $\phi$ is the same factor as the one used in beta attenuation calculations (1-$\phi$). A detailed study of these effects for the sediments from La Ferrassie is beyond the scope of this study; however, a few comments are appropriate. Firstly, the presence of potassium feldspar in the sand fraction and in particular in the 180-250 µm fraction used for $D_e$ measurements
indicates that there is at least some self-dose to K-feldspar to compensate for the inappropriate beta attenuation correction. Secondly, the observed bimodal grain size distributions (cf. Fig. S2) complicate the problem, since it is not unlikely that the radioelement contents are different in the two modes – in particular it is well-known that clay fractions are usually more radioactive than sands. Since radionuclide concentrations in these sediment samples are thus likely to be grain size dependent, the two dosimetric effects might not strictly compensate each other. If indeed the silt fraction contained more radioactivity, then there should be no self-dose, and, on average, no attenuation effect (cf. Fig. 3, in Guérin et al., 2012) so our calculated dose-rates would underestimate the true external beta dose-rates received by quartz and feldspar grains. The presence of sand size K-feldspar indicates only that this phenomenon does not fully affect – if it does at all – our samples, so we have adopted a conservative approach and calculated beta dose-rates using the infinite matrix approximation.

**Gamma dose-rates**

Gamma dose-rates were measured *in situ* by burying Al₂O₃:C crystal chips (1 mm thickness, 5 mm in diameter) in the sections in aluminium tubes (length: 30 cm, diameter: 1 cm). The dosimeters were left in place for one year and measured using a Daybreak 2200 OSL reader system (Bortolot, 2000) combining green light stimulation (Nichia NSPG310) with 7.5 mm Hoya U-340 filters, using the same protocol as Richter *et al.* (2010). These measurements are generally representative of present-day gamma dose-rates, because they characterize a volume of ~1 m³ (Guérin and Mercier, 2011) surrounding the measurement point. In contrast, samples analysed in high resolution spectrometry use ~10 g of material; using the resulting K, U, and Th concentrations to derive gamma dose-rates assumes that these 10 g are representative of the ~1000 kg of sediment contributing to the gamma dose-rate. In ‘lumpy’ (Brennan *et al.*, 1997) environments such as a karstic rockshelter, where one avoids taking calcareous lumps in the samples, the situation is clearly problematic since these low radioactivity
minerals will not be measured in gamma spectrometry but will contribute to gamma dose-rates. On the other hand, *in situ* dosimeters do not give any information concerning possible radionuclide disequilibrium, and so cannot indicate whether dose-rates have been constant through time.

Based on these considerations, we distinguish two different situations for the samples from La Ferrassie: samples FER 1-6 were taken from sector II, in the section left by the original excavators, and thus came from a thin layer of sediment covering the bedrock wall; on the other hand, samples FER7-14 were taken from horizons whose lateral extent was greater than the typical range of gamma rays in sediments. As a consequence, the systematic differences between gamma dose-rates measured *in situ* with Al₂O₃:C dosimeters and those calculated from K, U, and Th concentrations (Fig. 5; the factors used to correct gamma dose-rates for the effect of humidity were taken from Guérin and Mercier (2012), taking the average value – 1.15 – for cubic-centred assemblages with an average grain diameter of ~200 µm [*cf. Fig. S2*]) can be explained in two different ways: (i) For samples FER 1-6, the gamma dose-rates measured by dosimeters underestimate the gamma dose-rates received by the samples (by ~35-40%), because the dosimeters were closer to the bedrock wall than the samples. On the other hand, since the K, U, Th values did not include any contribution from the bedrock, these values overestimated the true gamma dose-rates received by the samples. As a consequence, the gamma dose-rates to the samples were taken as the averages between calculated values and those measured *in situ*, and the error was chosen to cover, at two standard deviations, the interval between these two extreme values. (ii) In the case of samples FER 7-14, where the discrepancies (of ~20%, on average) arose because of the absence of calcareous lumps in samples analysed by high resolution gamma spectrometry, *in situ* measurements are considered reliable estimates of gamma dose-rates received by the samples during burial. In addition to these systematic differences, there will also be differences arising from the different locations of the dosimeters and gamma spectrometry samples, but dose-rate discrepancies arising from this origin are assumed to induce random rather than systematic differences. Finally, the present-day
moisture content of 5% is presumed to represent the average water content throughout the burial time of the dosimeters, and gamma dose-rates measured with dosimeters were corrected for this moisture content before application of the correction during sample burial time (WF=15±5%, cf. previous subsection). The association between the dosimeters and the samples, and the gamma dose-rate calculations are shown in Table S2.

3.3. Radiocarbon

Bone samples were pre-treated at the Department of Human Evolution, Max Planck Institute for Evolutionary Anthropology (MPI-EVA), Leipzig, Germany, using the method described in Talamo and Richards (2011). The outer surfaces of the bone samples were first cleaned by a shot blaster, and then 500 mg of bone powder was taken. The samples were then decalcified in 0.5M HCl at room temperature until no CO₂ effervescence was observed, usually for about 4 hours. 0.1M NaOH was added for 30 minutes to remove humics. The NaOH step was followed by a final 0.5M HCl step for 15 minutes. The resulting solid was gelatinized following Longin (1971) at pH3 in a heater block at 75°C for 20h. The gelatine was then filtered in an Eeze-Filter™ (Elkay Laboratory Products (UK) Ltd.) to remove small (<8 µm) particles. The gelatine was then ultra-filtered with Sartorius “Vivaspin 15” 30 KDa ultrafilters (Brown et al., 1988). Prior to use the filter was cleaned to remove carbon-containing humectants (Higham et al., 2006). The samples were finally lyophilized for 48 hours.

4. Results

4.1. OSL - Multi-grain quartz (3 mm aliquots)

Signal characteristics and measurement protocol

The OSL signal from the La Ferrassie quartz is dominated by the fast component (see comparison with the signal from heated calibration quartz in Fig. S4a). The net signal intensity used in
further calculations was derived from the sum of the OSL in the first 0.8 s of stimulation minus a background signal. An early background subtraction method (Cunningham and Wallinga, 2010) was used to remove any contribution from the medium component of the OSL signal; the background intensity was taken as the sum of the OSL from the interval 0.8 to 3.2 s of stimulation, and then time averaged. It should be noted here that for sample FER 1, a late background subtraction was also tested (using the last 8 s of stimulation). With the early background subtraction, the average equivalent dose for this sample is $70.7 \pm 2.0 \text{ Gy}$. With a late background subtraction, this average equivalent dose becomes $70.5 \pm 1.9 \text{ Gy}$. This indicates that for quartz samples from La Ferrassie, whose OSL signal is dominated by the fast component, early and late background subtraction lead to indistinguishable results. SAR sensitivity-corrected dose response curves (Murray and Wintle, 2000; 2003) were constructed (Fig. S4b) using at least four regenerative doses bracketing the natural $D_e$. The effect of IR stimulation at $125^\circ\text{C}$ of varying duration (from 0 up to 320 s) was tested on sample FER 7 (Fig. S4c) and showed no dependency of $D_e$ as a function of IR stimulation duration. Furthermore, the IRSL signal intensity was, on average, less than 1% than OSL signals. This indicates that the quartz fraction is not contaminated significantly by feldspar and we can thus use a standard SAR protocol without prior IR stimulation. At the end of the SAR cycle, the first regeneration dose was repeated twice to obtain a recycling ratio and an IR depletion ratio (Duller, 2003). Both these ratios were within $\pm 15\%$ of unity for all aliquots, and no correlation between equivalent dose and IR depletion ratio was observed, confirming that the observed $D_e$ was insensitive to IR stimulation, and thus presumably not influenced by signals from feldspar. Recuperation was checked using a zero dose cycle after measurement of the largest regeneration dose and showed no significant inter-cycle signal transfer. All aliquots (between 16 and 32, depending on the variability in $D_e$ measurements to obtain comparable relative uncertainties) were thus used in the calculations of the unweighted average equivalent doses (see also discussion in section 4.3 about single grain data analysis). The SAR protocol was applied with various preheat conditions and Fig. S4d shows a preheat
plateau obtained for sample FER 7; each point corresponds to an average of 6 aliquots, except for the preheat temperature of 220°C (2 high dose outliers were removed). The preheat temperature was held for 10 s prior to L_i measurements. The cutheat temperature (applied before reading the T_i signal, used to monitor sensitivity changes) was 40°C lower than the preheat temperature. A dose recovery test, using a preheat at 240°C for 10 s and a cutheat of 200 °C, was performed on six aliquots from 10 different samples covering the entire sequence; the given dose (~60 Gy) was delivered to the aliquots in the reader after exposure to a solar simulator (Hönle SOL 2) for 3 hours. The average measured/given dose recovery ratio was 0.98±0.01. In Bordeaux, dose recovery tests were independently performed on 4 different samples (FER1, 3, 7 and 8) with the same parameters; the average measured to given dose ratio was 0.99 ± 0.01. Fig. S5 shows a frequency histogram of dose recovery ratios obtained for individual aliquots (n=60), indicating that our SAR protocol is able to accurately measure a laboratory dose given before any thermal treatment of the sample (see Table S1, SOM, for the measurement protocol used to estimate equivalent doses). The quartz samples from La Ferrassie thus met all laboratory-based criteria for a reliable estimate of equivalent dose. The uncertainties on individual equivalent doses have been assigned using the Analyst software (Duller, 2007) and include contributions from counting statistics (assuming Poisson statistics) and curve fitting errors.

**Data interpretation**

In Table 2 we summarize the relevant dose-rate data and in Table 3 the equivalent doses and resulting multi-grain aliquot quartz ages for all samples. Fig. 6 shows the relationship between archaeological layer and age. Four of the fourteen samples were measured both at Risø (Fig. 6, filled black squares) and in Bordeaux (Fig. 6, filled red circles) to ensure that our age estimates are not affected by inter-laboratory systematic errors. These paired ages agree well with each other (three pairs are consistent within one standard error and the fourth is consistent within two standard errors); we
thus consider it unlikely that our ages suffer from significant systematic off-sets that could, for instance,

arise from different calibrations of laboratory dose-rates.

There is a general increase of age with burial depth, or more precisely when moving downwards
in the stratigraphy (i.e., taking into account the tilting of the layers as evidenced by the fabrics – see
chapter by McPherron in Turq et al., 2012). A noticeable exception occurs for sample FER 2 (Layer 6),
which appears to be too old; for this sample we have an age inversion that is significant at two standard
errors.

Based on observations of sediment thin sections, post-depositional mixing does not appear to
have affected the integrity of the different layers. Furthermore, we do not consider it likely that this age
inversion is caused by beta dose-rate heterogeneity because neither the mineralogy nor the grain size
distributions changed significantly between Layers 5, 6 and 7. On the other hand, a consideration of the
process of sediment accumulation through slope transport and deposition mechanisms suggest that
incomplete bleaching may be a problem for these samples, i.e. the OSL signal may not always have been
adequately reset prior to sediment deposition. At this stage the most likely explanation for the age
inversion is incomplete bleaching which at least has affected Layer 6 more than the other layers. This is
possibly caused by the fact that Layer 6 as exposed today is quite sandy and non-calcareous. Field
evidence shows the presence of decalcified bedrock among the sand, demonstrating that much of the
sand is derived from the direct decomposition of the bedrock and was likely little exposed to ambient
light before burial.

One approach to investigating the presence of incomplete bleaching is to compare the OSL ages
obtained from quartz with IRSL ages obtained from feldspar. Given the different bleaching rates of these
two minerals, such a comparison enables us to make qualitative statements about the likelihood of the
presence of incomplete bleaching (Murray et al., 2012, and references therein). The quartz OSL signal is
much more rapidly reset by sunlight than the conventional IRSL signal from feldspar (~1 order of magnitude difference; e.g. Godfrey-Smith et al., 1988; Thomsen et al., 2008; Buylaert et al., 2012), which means that if the feldspar IRSL ages are comparable to the quartz OSL ages, then both signals are likely to have been sufficiently reset at deposition. If the feldspar ages are significantly greater than the quartz ones, then it indicates that incomplete bleaching may be a problem, at least for feldspar IRSL signals.

4.2. Feldspar

Feldspar as a natural luminescence dosimeter has several advantages compared to quartz (e.g., sensitivity and age range) but suffers from an athermal signal instability, known as anomalous fading (Wintle, 1973; Spooner, 1994). The conventional IRSL signal (measured close to room temperature, usually 50°C, and here identified as $\text{IR}_{50}$) typically fades by 3-5% per decade of time (Huntley and Lamothe, 2001). However, recent studies have demonstrated the potential of elevated temperature post IR-IRSL protocols (Thomsen et al., 2008; Buylaert et al., 2009), in which a further IR stimulation is made at some elevated temperature after an $\text{IR}_{50}$ stimulation, to overcome this problem. In particular, the post-IR IRSL signal at 290°C ($\text{pIRIR}_{290}$, Thiel et al., 2011) has been shown to be essentially non-fading by experiments on known age samples; this avoids the need for any inherently model-dependent fading corrections (Buylaert et al., 2012). Moreover, Buylaert et al. (2012) have shown that this signal is, in comparison with quartz OSL, very slowly reset by exposure to sunlight – probably orders of magnitude slower than is quartz. As a consequence, $\text{pIRIR}_{290}$ ages can only be greater than, or at best equal to, quartz ages. In the latter case this indicates good bleaching of both signals at the time of deposition. The situation where $\text{pIRIR}_{290}$ ages are greater than quartz ages indicates poor bleaching of the $\text{pIRIR}_{290}$ signal, but does not necessarily imply poor bleaching of the quartz signal (Murray et al., 2012) - or indeed of the $\text{IR}_{50}$ feldspar signal.
In this study we have measured K-rich feldspar extracts, not with the intention of providing independently-determined luminescence ages, but as an indicator of the degree of bleaching of quartz. This avoids the need for an accurate estimate of field fading rates and unbleachable residual signals as well as the debate on the applicability of fading correction models to samples > 50 ka (Huntley and Lamothe, 2001; Morthekai et al., 2008). Murray et al. (2012) have pointed out that because of the differential bleaching rates of quartz and the IRSL and pIRIR signals from feldspar even an incompletely bleached feldspar signal may indicate a well-bleached quartz signal.

Here we have used a SAR based protocol (see Table S1), with 200 s IRSL stimulations. The net signals were calculated using the initial 2 s of the pIRIR signals after subtracting a background obtained from the last 40 s of stimulation. We have derived pIRIR$\text{290}$ ages for seven of our samples and the results are given in Table 3 and Fig. 7a. All pIRIR$\text{290}$ ages are significantly larger than the corresponding quartz ages; indeed the pIRIR$\text{290}$ signal for FER 12 is so close to saturation that a dose could not be determined ("sat" in Table 3). The average ratio of pIRIR$\text{290}$ age to multi-grain quartz age is $2.3 \pm 0.3$ (n=7, Fig. 7c, open symbols). Such a large difference in this age range can almost certainly be attributed to incomplete bleaching of at least the pIRIR$\text{290}$ signal. Indeed, the measured excess doses for our samples, defined as the post IR-IRSL dose on top of the expected dose based on the quartz age – range from 61 to 490 Gy (Table 3). As a consequence we conclude that for all the measured samples, the pIRIR$\text{290}$ signal was probably not completely reset at the time of sediment deposition (although of course this does not necessarily indicate poor bleaching of the quartz signal).

Although the pIRIR$\text{290}$ signals may not have been completely bleached, the same does not necessarily apply to lower temperature pIRIR signals. Both theoretical models (Jain and Ankjærgaard, 2011) and experimental studies (e.g., Thomsen et al., 2008; Reimann et al., 2011; Reimann and Tsukamoto, 2012) indicate that when the stimulation temperature is decreased it becomes easier to
reset the signal under daylight (although the fading rate also tends to increase). So, by decreasing pIRIR stimulation temperature below 290°C, we would expect a decrease in equivalent dose because of better signal bleaching but also because of an increase in fading rate.

Figure 8 shows the equivalent doses obtained for sample FER 6, measured using different post IR-IRSL stimulation temperatures (see Table S1); the preheat temperature was systematically 30°C higher than the stimulation temperature, to minimise the risk of any isothermal thermoluminescence signal contributing to the pIRIR signals. Each point is the average of three aliquots. It appears that the equivalent dose increases with stimulation temperature by a factor of ~2 (corresponding to ~90 Gy) between 130 and 290°C. This is too large to be explained in terms of differential fading; the fading rate is confidently expected to be lower than that for IR at 50°C signals, which typically underestimates ages for well-bleached samples by ~30% (e.g., Buylaert et al., 2011; Huntley and Lamothe, 2001). We have measured the corresponding fading rate for the pIRIR signal for three different samples (FER 1, 3 and 6; 6 aliquots each); individual values are indistinguishable and the average fading rate is 2.05 ± 0.05 % per decade. Thus, we attribute this difference in D_e between low and high-temperature pIRIR signals to be mainly due to differential bleaching rates (perhaps arising in part from thermal transfer), although it must be acknowledged that there may also be a contribution from inaccurate measurement. The latter can be identified by e.g., a dose recovery experiment (see below). We also note that the equivalent dose is less sensitive to temperature change at low temperatures (Fig. 8); this supports the possibility that thermal transfer is playing a role.

A dose recovery ratio using the pIRIR signal was determined by adding a laboratory dose of 103 Gy on top of the natural dose for 6 aliquots of sample FER 6; after subtracting the natural D_e (111 ± 4 Gy) the measured to given dose ratio was 1.02 ± 0.05, confirming the ability of our measurement protocol to accurately determine a laboratory dose given prior to any thermal treatment; we assume
this conclusion also applies to natural doses. We used the $D_e$ determined with a pIRIR stimulation temperature of 160°C (111±4 Gy) to calculate an age (Table 3) of 50 ± 3 ka (without correcting for fading or subtracting any residual); this is consistent with the quartz age of 54 ± 4 ka. This agreement further supports the interpretation of Fig. 8 as a bleaching/thermal transfer plateau; the older ages for higher post IR-IRSL stimulation temperatures result from incomplete bleaching, but low temperature signals are better bleached (cf. Fig. 7b, c). The dashed line in Fig. 8 shows predicted equivalent dose derived by multiplying the quartz age by the dose-rate to feldspar; this is the dose expected if the feldspar signal does not fade, was well-bleached at deposition and does not preserve any significant unbleachable residual signal. The possible dose plateau at low stimulation temperatures is ~9% lower than that expected (rather than the ~20% lower expected from fading). This may indicate either (i) some residual incomplete bleaching, (ii) inaccurate laboratory estimation of field fading rates (e.g., Wallinga et al., 2007; Reimann et al., 2011), or (iii) the presence of an unbleachable/difficult-to-bleach residual dose. On the latter point, several studies (Madsen et al. 2011; Reimann et al. 2011; Reimann and Tsukamoto 2012) have reported very low residual doses (≤ 1 Gy) for low temperature pIRIR signals (pIRIR$_{150}$ and pIRIR$_{180}$). Furthermore, Kars et al. (2014) have shown that experimental determination of such pIRIR residual doses in the laboratory should be treated with caution. In the absence of modern analogues and because we do not intend to use feldspar pIRIR signals for producing an absolute chronology, we have decided to neglect potential residual, unbleachable dose. The average pIRIR$_{160}$/quartz age ratio (Table 3, Fig. 7c) is 1.08±0.05 (n=8) for samples between FER 1 and FER 8 in the stratigraphy, but excluding the outlier FER 2, for which the ratio is 1.97±0.23. Since the bleaching of feldspar signals occurs so much more slowly than that of quartz, we deduce that the conclusion above that the quartz signals in FER 6 were well-bleached applies also to all these samples (except FER 2). Because the pIRIR$_{160}$ to quartz age ratios increase below FER 8 (although not to the value recorded by FER 2, Fig. 7c), we cannot with confidence apply this conclusion to the deeper samples.
In sample FER 2 there is (i) an age inversion observed with multi-grain quartz aliquots, and (ii) an unusually large ratio of the pIRIR$_{160}$ to quartz age. Based on these observations, we suggest that the quartz luminescence signals of FER 2, from Châtelperronian Layer 6, are probably inadequately bleached. This is supported by the micromorphological observations (section 4.1), regarding the nature of Layer 6 and is investigated further in the following.

4.3. Single grains of quartz

4.3.1. Rationale

If a sample is significantly affected by incomplete bleaching, it is very likely that large multi-grain aliquot measurements will result in an overestimate of deposition age, because each aliquot is likely to contain both well-bleached and poorly bleached grains. One approach to determining a more accurate age for the depositional event is to measure individual grains and apply a minimum age model in an attempt to identify those grains most likely to have been well-bleached at burial. As the quartz OSL signal is most readily reset by exposure to light, the likelihood of identifying a well-bleached grain population in an incompletely bleached sample is greater for quartz than for feldspar.

As was discussed above (section 4.2) the comparison of quartz and feldspar multi-grain ages suggested that the quartz signals from samples FER 1, 3, 14, 4, 5, 6, 13 and 8 were all well-bleached, but that the quartz signals from sample FER 2 from the Châtelperronian Layer 6 may not be. Based on the quartz and feldspar comparison alone, we were not able to conclude whether the quartz signals from samples FER 7, 10, 9, 11 and 12 were well-bleached or not. To address these issues in more detail we have measured quartz single grains for six samples (FER 1, 2, 3, 4, 7 and 9) to allow us to compare single grain ages with the other ages and assess the applied methodology more thoroughly. In particular, obtaining a reliable OSL age for sample FER 2 requires single grain measurements and subsequent minimum age modelling to resolve insufficient bleaching.
4.3.2. Measurements and rejection criteria

The SAR protocol (Table S1) employed in these single grain measurements included a post-IR green signal (1 s stimulation with green laser at 125 °C following a 40 s stimulation with IR diodes at 125 °C; UV detection). The same preheat and cutheat temperatures as for the multi-grain measurements were used. The net signal used in $D_e$ calculations was derived from the sum of the OSL in the first 0.05 s of stimulation minus a background signal (time average of the last 0.2 s). Dose response curves were measured up to 200 Gy for each individual grain. These were fitted with a single saturating exponential function of the form $y = A (1 - \exp \left( \frac{D}{D_0} \right))$ where $y$ is the sensitivity corrected OSL signal, $D$ is the dose, $A$ is the saturated level of the sensitivity-corrected OSL signal and $D_0$ is the dose at which $y$ is at 63 % of the saturation level.

The dose estimates from individual grains were accepted based on their sensitivity by selecting only those grains for which the uncertainty on the first test dose signal was less than 20%. Thomsen et al. (submitted) have recently tested the effect of different commonly-employed single grain rejection criteria (e.g., Jacobs et al., 2006; Thomsen et al., 2012) on different samples. They observed that neither the average $D_e$ nor the central dose or the overdispersion from the Central Age Model (CAM; Galbraith et al., 1999) vary significantly when grains are rejected based on recycling ratios, recuperation, or IR sensitivity although this process results in a rejection of a great number of grains. We have investigated the effect of these rejection criteria on all analysed samples presumed to have been well-bleached at deposition (cf. Table S3: samples FER 1, 3, 4, 7 and 9). Since these criteria result in the rejection of a significant fraction of grains (26 % on average) for no significant effect on $D_e$ and OD values (and for no apparent improvement in reliability), these rejection criteria are not employed further in this study.

However, it should also be noted that between 8 and 22% of the otherwise accepted single grain analyses were deemed to be indistinguishable from saturation. Here we define saturation as a $L_N/T_N$
ratio (L/T ratio for the natural dose cycle) greater than or indistinguishable from the saturation level (A) of the laboratory dose response curve. Rejecting a significant fraction of measured grains because of saturation is common, and there is no accepted unbiased method for dealing with such analyses; all reported studies in the literature arbitrarily discard the results from such grains (e.g., Yoshida et al., 2000; Stone and Bailey, 2012; Thomsen et al., 2012); this must lead to a bias, but to an unknown degree. Thomsen et al. (submitted) have recently proposed an additional selection criterion which may provide an unbiased method of rejecting many of such saturated grains. This criterion is based on the ability of a grain to measure the dose of interest, i.e. based on the curvature of the dose response curve (the $D_0$ value). Philosophically, this is no different from choosing to use feldspar rather than quartz, because feldspar has a larger $D_0$ than quartz. For instance in the probably well-bleached sample FER 1, it seems very likely that the equivalent dose is ~70 Gy. However, for this sample, analysis of single grain dose response curves shows that there are grains with $D_0$ values of <20 Gy (see Fig. S6). Such grains would be close to 98% of saturation for a burial dose of 70 Gy. Assuming a typical measurement uncertainty of >5% implies that a significant fraction of measurements would be greater than or indistinguishable from saturation and so would be rejected. As a result, the burial dose based on the remaining accepted grains would be underestimated. By only accepting grains with a large $D_0$ value the risk of such bias is reduced, although not all saturated grains are removed (see Table S4). We argue that such a selection criterion must introduce less bias than that consisting of rejecting grains because $D_e$ is greater than 2 $D_0$ (as suggested by Wintle and Murray, 2006); indeed, such a rejection is $D_e$ dependent and, as such, must lead to some bias. In this study, we begin by only accepting grains with $D_0$>100 Gy. Application of this criterion reduces the number of grains appearing to be in saturation and systematically increases the burial dose for presumably well bleached samples by 11 ± 3% (n=6). The effect of this selection criterion on the dose distributions is shown in Fig. S7 for samples FER 1, 2 and 3 (radial plots are also provided, see Fig. S8) and the implications for age determination are discussed in section 4.5. However, as this $D_0$
criterion has not yet been used extensively in single grain OSL dating, we also report the results obtained without this additional criterion to enable a direct comparison with previously published single grain studies (values in brackets in Table 4).

4.3.3. Data analysis

Our single grain age determinations make use of methodological results, described elsewhere, concerning dose-rate distributions and their effect on equivalent dose population (Guérin et al., 2015); here we briefly outline the approach and describe the models used, based on the key concept of overdispersion defined as that part of the dispersion in results that cannot be explained by measurement uncertainties.

Three sources of dispersion contribute to the scatter in natural $D_e$ distributions: (i) measurement uncertainties (including a 2% uncertainty per OSL measurement arising from instrument reproducibility; see, e.g., Thomsen et al., 2005; Jacobs et al., 2006); (ii) intrinsic overdispersion, defined as that part of the dispersion in doses observed when measuring a known dose given in the laboratory (e.g., in a dose recovery experiment) that cannot be explained by measurement uncertainty; and finally (iii) extrinsic overdispersion, defined as that part of the dispersion in equivalent doses measured using natural samples that cannot be explained by (i) and (ii). The relative contributions of these sources of dispersion, as well as their characteristics, need to be assessed as part of selecting the most appropriate age model.

Analytical errors on individual dose estimates are, on average, ~20%. Fig. S9 shows the relationship between these errors (absolute and relative) and dose estimates. The absolute errors tend to increase as the dose increases, although the correlation is weak. A plot of relative errors against dose shows an even weaker trend, if any. We deduce that our analytical uncertainties tend to have multiplicative error properties, rather than additive (see e.g., Galbraith and Roberts, 2012). Such error properties make models such as the CAM, where the central dose is calculated using a weighted
geometric mean, more appropriate than models such as the CAM$_{UL}$, where the central dose is estimated using a weighted arithmetic mean. Models such as the latter are suited for distributions where additive error properties explain the dispersion in $D_e$ measurements (e.g. Galbraith and Roberts, 2012).

We carried out single-grain dose recovery experiments using portions of samples FER 3 and FER 9 that were bleached for 3h in a solar simulator (Hönle SOL 2 at a distance of ~80 cm) prior to being given gamma doses of 57.6 and 95.4 Gy. The relative intrinsic overdispersion (OD) values for these single dose populations, determined by the CAM were, on average, 12%. To investigate the nature of this intrinsic overdispersion, we analysed the standardised residuals from both the CAM and the CAM$_{UL}$ (see, e.g., Galbraith and Roberts, 2012). In these models, either the same relative or the same absolute error is added in quadrature to analytical uncertainties for each dose estimate. Fig. S10 shows quantile-quantile plots of standardised residuals from both the CAM and the CAM$_{UL}$ applied to the 57.6 Gy dose recovery $D_e$ population from sample FER 3. The CAM seems to provide a better fit to the data than the CAM$_{UL}$, presumably because the dispersion in estimated $D_e$ values is dominated by errors with multiplicative rather than additive properties. We thus consider the CAM best suited to interpret dose recovery tests. The average dose recovery ratio ($1.05 \pm 0.03$; see Table 4) cannot be distinguished from unity at two standard errors; our SAR protocol was thus deemed suitable for all samples.

In the natural samples, the measured OD values range between 29±3% (FER 1) to 51±5% (FER 2). The average OD is 35.0±1.7% (n=5) excluding FER 2 which is likely to be incompletely bleached. From the quartz and feldspar age comparison, we concluded that the quartz in sample FER 1 to FER 8 (but not including FER 2) were probably well-bleached at deposition, but we were unable to demonstrate that quartz from samples FER 7 and below were well-bleached. However, the OD values determined from the single grain measurements of FER 7 (35±3%) and FER 9 (39±3%) are completely consistent with the ODs
determined from the well-bleached samples (FER 1, 3 and 4), we now conclude that the quartz signals from these samples were also probably well-bleached at deposition.

The fact that the average OD of 35% for natural, well-bleached distributions is significantly greater than the average OD of 12% determined in the dose recovery experiments suggests that the main source of dispersion in our well-bleached natural samples is extrinsic. We also applied the CAM and the CAM\textsubscript{UL} to our natural D\textsubscript{e} distributions, and Fig. S11 shows a quantile-quantile plot of the standardised residuals from the two models applied to the natural D\textsubscript{e} population from sample FER 3. Neither of these two models provides a good fit of the data; this suggests that the dispersion in natural D\textsubscript{e} distributions cannot simply be explained by additional uncertainties with either multiplicative or additive properties.

Given the important contribution of potassium beta dose-rates to total dose-rates, skewed, log-normal distributions can result from the presence of K-feldspar in sediments (e.g., Brennan et al., 1997; Mayya et al., 2006), especially when the average K-content is relatively low (~0.55 – 1 %, cf. Table 2). In addition, our samples include a strong sand component with average grain sizes of several 100 µm, and so the inter-grain distance is not negligible compared to the average range of beta particles in sediments. As a result dose-rate distributions are likely to be an important contribution to origins of the spread in measured D\textsubscript{e} values; we assume that it is the dominant source of extrinsic dispersion in our natural, well-bleached D\textsubscript{e} distributions (samples FER 1, 3, 4, 7 and 9). Whatever the nature of the single-grain dose-rate distribution, it is important to note here that any dose-rate measurements (direct or indirect) provide average dose-rates (arithmetic mean). In cases where dose-rate distributions are an important source of spread in D\textsubscript{e} distributions, it is likely that the CAM will underestimate the central dose value; i.e. the dose that must be divided by the measured/calculated average dose-rate (arithmetic mean, which is independent of the nature of the dose-rate distributions) to give an accurate age. As a
result a CAM dose will underestimate the age, because (i) the CAM provides geometric means
(systematically lower than arithmetic means, for any distribution) of \( D_e \) values and (ii) the weighting of
individual estimates by the CAM is dominated by sources of dispersion (in this case, those from
dispersions in dose-rate) that have nothing to do with measurement uncertainties (the bigger the
extrinsic overdispersion, the closer the CAM result will be to the geometric mean of the dose
distribution, irrespective of measurement uncertainties – see last column of Table 4). As a consequence,
the presence of two sources of dispersion, each of a different nature argues against the use of models
based on either weighted geometric means such as the CAM (because they are not suited to
distributions where dose-rate variations explain a large part of the dispersion) or on weighted arithmetic
means such as the CAM\(_{ul}\) or the IEU models (Thomsen et al., 2007), because they are not suited to
distributions displaying multiplicative error properties.

For these reasons, in our \( D_e \) calculations, we consider it more appropriate to ignore our
individual error estimates on \( D_e \) values derived from individual grains (because they only explain a minor
part of the dispersion in our natural distributions); we calculate unweighted, arithmetic means of
individual \( D_e \) estimates (see Thomsen et al., submitted). All single grain ages are listed in Table 3; there is
agreement within two standard deviations with the multi-grain results for the well-bleached samples
FER 1, 3, 7 and 9 and within three standard deviations for FER 4.

4.3.4. Minimum age modelling

A minimum age model was used to interpret the dose distribution of sample FER 2 (Fig. S7);
poor-bleaching in this sample appears to have resulted in an OD value (51 ± 5 %) considerably greater
than those obtained for the well-bleached samples (29 ± 3 % for FER 1 and 35 ± 3 % for FER 3; see Table
4) directly above and below the sampling location. Furthermore, after selection of grains based on \( D_0 \)
values, for sample FER 2, 9 of the otherwise accepted grains (i.e. 9% of the grains) appeared to be in
saturation after selection of grains having $D_0 > 100$ Gy; this fraction of such rejected grains was much higher than for samples 1, 3 and 4 and further indicates the possibility of poor bleaching (see Table 4).

Given the discussion above on various central age models, there is no obvious model to use for sample FER 2; even the well-bleached portion of the grains from this sample will have a significant overdispersion dominated by extrinsic factors such as dose-rate dispersion. In our case, samples FER 1 and 3 bracket sample FER 2 in stratigraphy. Moreover, the age range is similar since the quartz multi-grain ages range from $36.7 \pm 2.0$ ka (FER 1) to $42.2 \pm 2.8$ ka (FER 3). The OD values measured on these well-bleached samples (presumed to arise solely from intrinsic factors and dose-rate heterogeneities) were very similar (32 % on average).

Thus, we have applied the 3 parameter MAM (Galbraith et al., 1999) to the distribution of $D_e$ values from FER 2, after selection based on $D_0$ values and with an input OD value of 32%. However, the MAM does not fit the data very well as indicated by the log likelihood profiles (Fig. S1). The log likelihood profiles are not symmetrical bell-shaped curves with well-defined 68% and 95% confidence intervals, which can be taken as evidence that the resulting OSL age is unreliable (Arnold et al., 2009).

Nevertheless, the MAM burial dose estimate of $76.3 \pm 6.7$ Gy (at 68% CI) corresponds to an age of $41.0 \pm 3.9$ ka, which is consistent with the chrono-stratigraphy and with other reported ages for Châtelperronian occupations in southwest France (e.g., Hublin et al., 2012).

We have also applied the IEU minimum age model to the same data set. This model has been tested against independent age control, and here we use an a value of 0.32 (this study) and a typical b value of 0.02 Gy (e.g., Thomsen et al., 2007; Sim et al., 2014; Medaldea et al., 2014). The resulting burial dose for FER 2 is $78.4 \pm 3.5$ Gy (using 76 grains, i.e. 84% of the total); this gives an age of $42.2 \pm 2.9$ ka, which is in agreement with the MAM age and the chrono-stratigraphy.

4.4. Summary of luminescence data
A comparison between quartz and various pIRIR signals has enabled us to identify one poorly-bleached sample (FER 2) and distinguish it from other surrounding samples. By using pIRIR signals as bleaching proxies in combination with single grain measurements we confirmed that for the remaining samples, the quartz ages obtained from conventional multi-grain measurements were unlikely to be systematically in error because of incomplete bleaching. However, sample FER 2 was probably poorly bleached (which confirms field observations of the corresponding sediment), and required a combination of single grain analysis and minimum age modelling to identify and evaluate the $D_e$ of well-bleached grains. As a result, three datasets have been used to derive ages: multi-grain quartz OSL data from both Risø and Bordeaux laboratories, and single grain quartz OSL data from the Risø laboratory (Fig. 9). The resulting dose estimates were indistinguishable from those samples (FER 1, FER3 and FER 7) for which all three data sets were available, and so a weighted average quartz age has been calculated for each of these samples (Table 3). These ages provide the basis for the discussion in Section 5 of the implications of the chronology of the different occupation Layers of La Ferrassie (Table 6).

4.5. Radiocarbon ages and overall chronology

Collagen yield ranges between 2.5 and 4.4 % for the samples listed in Table 5; C:N ratios are well within the accepted range (Van Klinken 1999). The one sample from the Châtelperronian Layer 6 is dated to between 42.4 and 41.3 ka cal BP (95% confidence interval - CI), whereas three samples from Layer 5b provided ages between 44.4 and 47.3 ka cal BP (lower and higher limits of the 95% CI from the three samples).

4.6. Single grain OSL data analysis in view of radiocarbon results

Given these radiocarbon results, we can now discuss the appropriateness of our single grain OSL analysis. We first consider the choice between the CAM, the CAM$_{UL}$, the IEU and the unweighted average of $D_e$ values in providing the best estimate of the burial dose. For sample FER 3 (the only well-
bleached sample for which a comparison with radiocarbon is available) the single grain CAM age is 38.1 ± 2.2 ka, the single grain $\text{CAM}_{\text{UL}}$ age 37.8 ± 2.2 ka, and the IEU age 38.4 ± 2.6 ka. None of these are consistent (95% confidence) with the $^{14}$C age range. In contrast our preferred unweighted average of $D_e$ values gives an age of 40.9 ± 3.1 ka (see Table 3), which is consistent (95% confidence) with the radiocarbon ages. This supports our contention that none of the more statistically robust models (CAM, $\text{CAM}_{\text{UL}}$, IEU) offer an appropriate treatment of distributions where both multiplicative error properties and important dose-rate (i.e., true dose) variations contribute to the scatter in measured $D_e$ values.

Even though we have no radiocarbon age to compare with the single grain OSL age of sample FER 1, it is interesting to note that the ages obtained with the CAM (36.6 ± 2.0 ka), the $\text{CAM}_{\text{UL}}$ (35.6 ± 1.9 ka) and the IEU (35.7 ± 2.1 ka) are systematically younger than that based on the age derived from the unweighted arithmetic mean of $D_e$ values (38.4 ± 2.5 ka).

The minimum ages obtained for sample FER 2 using the MAM or the IEU cannot be distinguished (95% confidence) from the $^{14}$C age range of 42.40 to 41.25 cal BP ka (95 % CI). It is difficult to choose which minimum age model provides the most accurate age; however, because (i) the likelihood profiles obtained with the MAM (Fig. S12) do not appear very satisfactory, (ii) the age uncertainty obtained with the IEU is smaller than with the MAM, and (iii) the IEU age is closer to the $^{14}$C age, in Table 3 the single grain age of FER 2 is that obtained with the IEU. The excellent agreement between the IEU age and the $^{14}$C available leads us to formulate the hypothesis that for this particular sample, poor-bleaching is the dominant factor of dispersion in measured $D_e$ values. Even though the IEU does not seem to handle the dispersion factors for well-bleached samples, it seems to give a reliable age estimate for the poorly-bleached sample FER 2.

The second analytical approach that can be tested by comparison with $^{14}$C is the use of a selection criterion based on the curvature parameter ($D_0$) of laboratory dose-response curves, and in
particular the choice of a value for this parameter which selects only those grains that are able to record the full range of doses recorded in the sediment. Table S4 lists the results obtained for two of the three uppermost samples in the stratigraphy: FER 2 and 3 (unweighted averages of $D_0$ values and corresponding ages for sample FER 3; IEU doses and ages for poorly-bleached sample FER 2). These are the only two samples with independent $^{14}C$ age control. As expected there is a systematic increase in age for both samples (FER 2: 18%; FER 3: 8%) using the $D_0$>100 Gy criterion. However, the equivalent dose does not depend significantly on $D_0$ values > 60 Gy. Furthermore, for all $D_0$ values between 80 and 120 Gy, the single-grain OSL ages for samples FER 2 and 3 are in agreement with the available radiocarbon ages (one exception to that rule is for $D_0$>120 Gy, where the age of FER 3 is not consistent with the greatest radiocarbon age of sample S-EVA 26506, but is consistent with the other two radiocarbon ages from Layer 5b). We conclude that (i) without applying this $D_0$ criterion the single grain quartz OSL ages would underestimate the expected age based on $^{14}C$ and (ii) this statement is independent of the chosen value of $D_0$ between 80 and 120 Gy. This supports our a priori argument that the $D_0$>100 Gy selection criterion is necessary and provides an unbiased approach to reject most of the saturated grains. This also seems to confirm that these low $D_0$ grains are not capable of recording the dose range required to calculate an accurate age for these samples.

5. Discussion of the ages and implications

The OSL data suggest the following chronology for the La Ferrassie sequence: Layer 1 was deposited during MIS 5 (most likely 5b), around 90 ka (even though the overestimation of this age by pIRIR$_{160}$ signals leaves room for an underestimation of the deposition age by quartz OSL, arising from saturation effects in the quartz OSL signals), and Layer 2 during MIS 4 with ages ranging from 74 ± 5 to 62 ± 4 ka. The ice segregation and cryoturbation patterns identified in Layer 2 and at the top of Layer 1 seem consistent with the deposition of Layer 2 during MIS 4. Layer 3 was deposited during MIS 3
between 57 ± 3 and 44 ± 3. From a chronological point of view, Layer 4 is indistinguishable with ages of
54 ± 4 and 44 ± 3 ka. Layer 5 yielded three statistically indistinguishable OSL ages with an average of 43 ±
3 ka, which is consistent with the radiocarbon results (from 47.3 to 44.4 ka BP). Application of a
minimum age model to the single-grain quartz OSL dose distribution gives an age of 42 ± 3 ka for Layer
6; this OSL age is indistinguishable from the radiocarbon age of between 42.4 and 41.3 ka cal BP (95% CI)
from the same layer and also compares well with radiocarbon ages obtained recently for the same
industry from the Grotte du Renne in Arcy-sur-Cure and from Saint-Césaire (Hublin et al., 2012; see also
Mercier et al., 1991). The agreement between single-grain OSL and radiocarbon data for this layer
indicates, a posteriori, that there was, indeed, most likely a portion of the quartz grains from sample FER
2 which were sufficiently exposed to sunlight to have their OSL reset signal prior to deposition. Finally,
Layer 7 is dated to 37 ± 2 ka. This age also agrees well with the Aurignacian attribution of this layer
(Banks et al., 2013, and references therein). Taxa associated with cold climate conditions (reindeer,
woolly mammoth, or woolly rhinoceros) occur throughout the sequence (with the exception of Layer 4)
and are dominant in Layers 6 and 7 (reindeer only). Thus it could be that Layers 6 and 7 correspond to
one of the cold events of MIS 3. Taxa associated with warmer environments (especially roe deer) are
found in Layers 5b and below. Layer 2, associated with MIS 4, preserves a significant number of taxa
adapted to temperate conditions (in particular, higher abundance of roe deer over reindeer) – which can
be linked with, e.g., the Roc de Marsal sequence where the fauna attributed via TL and OSL dating to
MIS 4 also shows diversity in taxa (Guérin et al., 2012). Further refinement of the chronology of the
upper part of the sequence using radiocarbon is ongoing. These data are all summarized in Fig. 9 and
Table 6.

These ages add to the growing database for late Middle Paleolithic sites in southwest France
(e.g. Guibert et al., 2008) and to the long-debated question of whether lithic variability in these sites
follows chronological patterns (Mellars 1969). The variability organized into the so-called Mousterian
facies (e.g. Ferrassie Mousterian, MTA, etc.) consists of three main axes: (i) whether blank production is dominated by Levallois techniques or Quina techniques, (ii) how common scrapers are among the retouched tools; and (iii) the presence or lack of, typically, cordiform bifaces. Prior to the development of absolute dating techniques applicable to this time period, Mellars (1969, 1996) noted that (a) in stratigraphic successions scraper-rich industries (Ferrassie and Quina Mousterian) occur below the MTA, (b) the scraper-rich industries which Levallois technology (Ferrassie Mousterian) gives way to later Quina techniques, and (c) MTA assemblages are at the top of the sequence and are frequently immediately under the Châtelperonian (further suggesting that MTA represents the final Mousterian). However, more recent efforts to re-evaluate existing ages and to re-date the sites of southwest France have not fully supported this model and have instead shown that scraper rich industries (Quina Mousterian) overlap – within uncertainties – with those with few scrapers (Denticulate Mousterian) and that bifacial industries (MTA) overlap with both of these (Guibert et al., 2008; Richter et al., 2013a).

The base of the La Ferrassie stratigraphic sequence does not contradict the Mellars’ stratigraphic model because the bifacial assemblages found there under the Ferrassie Mousterian have not been previously considered either Acheulian or MTA (cf. Mellars, 1969:167). Typically in southwest France the Acheulian is older and the MTA is younger (Soressi et al., 2007; Guibert et al., 2008; Vieillevigne et al., 2008; McPherron et al., 2012; Richter et al., 2013) than the ages reported here. However, aside from these two techno-complexes, bifaces occur at a larger geographical scale throughout the Middle Palaeolithic (Monnier, 2006; Ruebens, 2012). This is especially true of northern France where bifacial industries tend to date to MIS 5 (see review in Ruebens, 2012). Thus the La Ferrassie ages from the lower levels suggest more variability in the late Middle Palaeolithic of South West France than the Mousterian facies framework considers. A more precise characterization of the La Ferrassie biface assemblages and their affinities with other known bifacial industries is work ongoing.
The age of the Ferrassie Mousterian levels at La Ferrassie raises even more questions. Given that the overlying Châtelperronian is increasingly well constrained elsewhere to no earlier than approximately 45 ka (Hublin et al., 2012), it seems that these Ferrassie layers at La Ferrassie are indeed quite late in the Middle Palaeolithic of southwest France. Scraper-rich industries based on Levallois technology have remained outside recent dating efforts (e.g. Guibert et al., 2008) probably because they are relatively rare in southern France and virtually all examples of it come from relatively old excavations with sometimes questionable stratigraphic contexts. Examples include Abri Chadourne (Bordes et al., 1954), Caminade (Sonneville-Bordes and Montureux, 1955), Pech de Bourre (Peyrony, 1942), Roc en Pail (Gruet, 1969; 1984), and Petit Puymoyen (Favraud, 1908). There are also four examples of Ferrassie Mousterian layers from Bordes’ excavations at Combe Grenal. That said, the new results from the eponymous site of La Ferrassie suggest that the layers in question are chronologically indistinguishable from what has been previously reported for Denticulate Mousterian, MTA, and for most of Quina Mousterian (Soressi et al., 2007; Guibert et al., 2008; Vieilllevigne et al., 2008; McPherron et al., 2012; Richter et al., 2013a, b), meaning that a scraper-rich industry with Levallois technology is also part of late Middle Palaeolithic variability. Further, the age of Layer 5 of La Ferrassie is younger than the ages determined for the Quina Layer SW-US22 from Chez Pinaud-Jonzac (TL on heated flints: 73 ± 8 ka, Richter et al., 2013b). Thus, in view of these radiometric dating results, it seems that the hypothesis of a regional stratigraphy where Quina Mousterian always precedes Ferrassie Mousterian does not hold against chronological information.

One alternative possibility is that the Mousterian in Layers 4 and 5 at La Ferrassie – thus far recognised as Ferrassie Mousterian – is not actually a Ferrassie Mousterian. Older collections such as that from La Ferrassie can have significant excavator bias, meaning that particular pieces were preferentially saved. At Combe-Capelle Bas, for example, new excavations (Dibble and Lenoir 1995:168-170) showed that the 1930s collections favoured scrapers over notches and denticulates. The same
could have easily happened at La Ferrassie. Our collections are at present limited, and so a better understanding of the significance of the young age of these deposits for models of Mousterian stone tool variability will have to wait for a more extensive excavation to generate a sufficient sample for a proper characterization of the lithics (currently < 1 m² has been excavated).

Finally, the ages reported here provide a chronological context for the La Ferrassie 2 skeleton. According to the original excavators and our recent analysis of the sediments attached to the foot in comparison with those from our stratigraphic section, the LF 2 skeleton – and most likely LF 1, which was found 50 cm east of La Ferrassie 2 (cf. section 2.4) – was found in one of the Ferrassie Mousterian layers; whether it was in our Layer 4 or 5 remains unclear at this stage, but this still provides chronological information. The oldest OSL age for Layer 4 provides a Terminus Post Quem: the LF 2 skeleton is not older than 54 ± 4 ka and can now be placed in the final Mousterian of southwest France, most likely around 44 ± 3 (top of Layer 4) or 43 ± 3 (Average of individual ages from Layer 5).

It is likely that this age also applies to the La Ferrassie 1 skeleton. The age of the remaining skeletal material uncovered at the site, however, remains unknown. Though the archaeological sequences of the east section preserved today at La Ferrassie and our own sequence are the same and although Peyrony and Capitan reported a consistent stratigraphy across the site, it is clear from our on-going analysis that there were two different sources for the sediments in the western versus the eastern portions of the entire cave complex, i.e. for the sediments of our excavations at the entrance to La Ferrassie, and for those of Peyrony and Delporte further inside the cave. Thus, until strong arguments can be made to place each of these Neanderthals in one depositional context or the other, it would be premature to discuss their possible age.

6. Conclusion
Luminescence dating methods have been applied to sediment samples from the entire stratigraphic sequence of La Ferrassie. The comparison between quartz OSL and feldspar post-IR IRSL measurements, at various stimulation temperatures, on multi-grain aliquots has indicated that the quartz OSL signal from all samples but one (FER 2, from the Châtelperronian layer) was well-bleached at the time of deposition and thus can provide reliable chronological data. Single grain OSL measurements were undertaken on sample FER 2, and on adjacent well-bleached samples, to investigate the sources of dispersion in the equivalent dose distributions. Finally, a preliminary set of radiocarbon ages has shown excellent agreement with OSL ages for the youngest Ferrassie Mousterian in the sequence and for the Châtelperronian layer, thus confirming the accuracy of our OSL age modelling at La Ferrassie (and the relative inaccuracy of models not taking dose-rate variations into account, at least for well-bleached samples).

The presence of bifaces in a Mousterian industry, at the base of the sequence, has been dated to the MIS 5 (most likely 5b), the MIS 4 and the MIS 3, which in the first two cases is rather unusual for cave sites in southwest France and in particular older than dated occurrences of MTA in the area. The Ferrassie Mousterian layers have been dated to the MIS 3 (between 54 ± 4 and 40 ± 2 ka) and thus appear quite late in the Middle Palaeolithic of the region; these chronological results both suggest additional complexity in late Mousterian variability. The La Ferrassie 1 and 2 skeletons, which have been attributed to one of these Ferrassie Mousterian layers, are most likely dated to ~43-45 ka and are no older than 54 ± 4 ka. The upper part of the Ferrassie Mousterian layer is chronologically indistinguishable from the Châtelperronian layer (~42 ± 3 ka), which seems to be contemporaneous of other Châtelperronian assemblages in the region. Finally, the Aurignacian Layer is dated to 37 ± 2 ka.

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Table captions

Table 1. Tentative interpretation of geological history of La Ferrassie entrance section. Archaeological layers are indicated, together with the correspondence with Peyrony’s stratigraphy and the lithics industry. The list of sediment samples collected for luminescence dating also appears. The sectors refer to the site map (Fig. 2).

Table 2. Dose-rate information for the luminescence samples. K, U and Th contents have been determined by high-resolution gamma spectrometry. Beta and gamma dose-rates have been corrected for the effect of humidity in the sediments. Etching has been taken into account for the beta dose-rates. See text for details.

Table 3. Equivalent doses and ages for the luminescence samples. As explained in the text, not all ‘ages’ should be considered for the chronology: pIRIR ages in particular have here been used as bleaching proxies for the sediments. For clarity, the preferred ages have been highlighted in bold and an average quartz age, which should be the working value, has been calculated for each sample as the average between multi-grain aliquots (including measurements performed in Bordeaux) and single grain values. FER 2 is an exception to that rule: because of poor-bleaching, only the modelled single grain age (calculated using the IEU, cf. section 4.3) is considered reliable and thus appears in the average column.
For sample FER 12, the $D_e$ values determined using the pIRIR$_{160}$ signal systematically lie above $2D_0$ (but below the saturation level); these doses are reported as minimum values.

Table 4. Single grain OSL data. “N” is the number of measured grains; “$S_T < 20\%$” is the number of grains for which the uncertainty on the first (‘natural’) test dose response is less than 20%; “sat. (%)” is the percentage of grains that are rejected due to saturation. “n” is the number of grains used in $D_e$ calculations, i.e. grains with $S_T < 20\%$, $D_0>100$ Gy and not in saturation. “CAM $D_e$” is the weighted dose calculated using the Central Age Model (Galbraith et al., 1999) and “OD” is the corresponding relative overdispersion. “DR” is the dose recovery ratio, i.e. the measured to given dose ratio, calculated using the CAM (see text for details). “Avg. $D_e$” is the arithmetic (unweighted) average of equivalent dose estimates used in the age calculation. In all columns, the values quoted in brackets are derived from datasets containing grains with low $D_0$ values (i.e. $D_0<100$ Gy; see text for details). The values given for FER 2* have been obtained with the IEU model (for this line, the value indicated in the “Avg. $D_e$” column is the dose determined with the IEU). The last column gives the CAM to unweighted average dose ratio, which seems to increase when the CAM overdispersion is increased.

Table 5. AMS radiocarbon dating results of 4 samples from La Ferrassie. Isotopic values, C:N ratios, amount of collagen extracted (%Coll) refer to the >30 kDa fraction. The $\delta^{13}C$ values are reported relative to the vPDB standard and $\delta^{15}N$ values are reported relative to the AIR standard. Ages have been calibrated using OxCal 4.2 (Bronk Ramsey and Lee, 2013) using the international calibration curve IntCal 13 (Reimer et al., 2013). These ages can thus be directly compared with the luminescence ages.

Table 6. Summary of relevant ages, including radiocarbon data, presented together with available data on geology, lithics, hominin remains, and fauna analysis, and in connection with marine isotope stages.
Figure captions

Fig. 1. Regional map, indicating the location of La Ferrassie and other important Palaeolithic sites in the Dordogne region of France.

Fig. 2. Site map with newly excavated sectors and the location of luminescence samples (blue squares).

Fig. 3. Stratigraphic section of La Ferrassie. OSL samples are indicated as blue squares and Al$_2$O$_3$:C dosimeters as red circles.

Fig. 4. Connection between the foot of the La Ferrassie 2 individual and our stratigraphy. (a) Foot as preserved, with sediments attached to it, in the Musée de l’Homme (Paris). (b) View of the sampled section for micromorphology analysis, and corresponding thin sections. The best match between the sediments attached to the foot and the thin sections place the LF 2 individual in Layer 5. (c) Approximate find location (horizontal only) for the skeleton of LF 2, according to historic photographs of the excavations carried out by Peyrony and Capitan.

Fig. 5. Comparison between *in situ* measurements (red filled squares) and infinite matrix gamma dose-rates (blue open circles) as deduced from K, U and Th contents using the conversion factors from Guérin et al. (2011). All values have been corrected for moisture content following Guérin and Mercier (2012). The origin of the discrepancies and their consequences on dating results are discussed in the main text. See Fig. 3 for sample and dosimeter location.

Fig. 6. OSL ages from quartz multi-grain aliquots measured in Risø (black filled squares) and Bordeaux (red filled circles).

Fig. 7. Comparison between quartz multi-grain OSL (measured in Risø: black filled squares; in Bordeaux: red filled circles) and feldspar pIRIR ages. (a) pIRIR$_{290}$, open circles; (b) pIRIR$_{160}$, open squares. For sample FER 12, the $D_e$ values determined using the pIRIR$_{160}$ signal systematically lie above 2 $D_0$ (but below the
saturation level) and are thus not plotted. (c) Ratio of pIRIR to OSL ages (filled squares: pIRIR\textsubscript{160} to OSL; open squares: pIRIR\textsubscript{290} to OSL).

**Fig. 8.** pIRIR equivalent doses for samples FER 6, as a function of stimulation temperature (the preheat temperature was in all cases 30°C higher than the stimulation temperature). The dashed line indicates the expected equivalent dose based on measurements of quartz multi-grain aliquots, assuming a non-fading pIRIR signal and no residual signal at deposition. The equivalent dose increases with stimulation temperature, and the low-temperature flat region seems to indicate a bleaching plateau.

**Fig. 9.** Summary of reliable age estimates: all OSL ages obtained from quartz multi-grain aliquots measured in Bordeaux (red circles) and Risø (black squares) – except for the poorly-bleached sample FER 2 – as well as from single grains (black stars) plotted as a function of Archaeological Layers. These ages correspond to the bold values in Table 3 and are the ones considered in the discussion.
<table>
<thead>
<tr>
<th>Layer</th>
<th>Peyrony 1934</th>
<th>Industry</th>
<th>Description</th>
<th>Comments</th>
<th>OSL Sample</th>
<th>(^{14}C) Samples</th>
<th>Sector</th>
</tr>
</thead>
</table>
| 8     | I            | sterile           | • Light reddish rounded and angular gravel that continues to close to the top of the profile.  
• Connection with platform above the site next to upper cave remains to be determined | Derived from flat area in front of upper cave                                              |            |                     |        |
| 7b    | H            | Aurignacian       | • Compact, dark yellow brown sandy silt with abundant lithics and bone      | • Dry fall talus cone whose origin is a platform emanating from upper cave but whose apex was in the area near the road above the trench. | FER1       |                     | II     |
| 7a    | G            | G                 | • Compact, dark yellow brown sandy silt with abundant lithics and bone      | • Idem                                                                                      |            |                     |        |
| 6     | E            | Châtelperronian   | • Compact massive reddish brown silty fine sand                             | • Idem                                                                                      | FER 2      | S-EVA 26510         | II     |
| 5b    | D            | Ferrassie Mousterian | • compact bedded yellowish red brown silty sand with abundant ~cm-sized fragments of lithics and bone  
• Somewhat richer in lithics and less sandy than 5a                                  | • Idem                                                                                      | FER 3      | S-EVA 26506-26507-26508 | II, III |
| 5a    | D            | Ferrassie Mousterian | • compact bedded yellowish red brown silty sand with abundant ~cm-sized     | • Idem                                                                                      | FER 4      | FER 14              | II, III |
|   |  |   | fragments of lithics and bone  
• Somewhat less rich in lithics and sandier than 5b. |
|---|---|---|---|
| 4 | C | Ferrassie Mousterian | Massive, compact silty medium sand with relatively abundant cm-sized pieces of bone and flint, which are larger than those in unit 5  
• Truncates layer 3 in Sq. I4  
• Large roof fall blocks collapsed during the middle of Unit 4 time  
• Upper half of Unit 4 is banked up against the roof fall and is generally horizontal |
|   |   |   | Large collapse of roof after initial accumulation of Unit 4  
• First major inputs from human occupation  
FER 5  
FER 6 |
| 3 |   | Bifaces + Large flakes | Poorly sorted light brown silty sand with pebbles and cm-sized fragments of Mn-stained, angular burned bone fragments.  
• Inclined to SSE  
• Increasingly stony and flint- and bone-rich to north |
|   |   |   | Mudflow derived from NW  
• Source area is no longer visible  
• Roof beginning to collapse  
FER 7  
FER 8  
FER 13 |
| 2 | B | Bifaces + large flakes | Cemented stony sand with lenses of coarser limestone, which climb upward in the profile; some well-rounded cm-sized gravel  
• Increasingly limestone rich toward the wall  
• Unconformity between 1 and 2  
• During excavation, the base |
|   |   |   | They are inclined from NW to SE thus pointing to a source of this sediment in the direction of the road to the NW  
• The original source material was removed during construction of the  
FER 9  
FER 10 |
of this unit is inclined to NW
• 2-shots on bones also show orientation in this direction
• Deposits are subhorizontal but contact dips to the S

road
• Likely solifluction deposit synchronously affected by cryoturbation that also affected the top of Unit 1

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<th>Bifaces + large flakes</th>
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<td>• Locally red sand (with <em>éboulis</em>) locally laminated and with pockets and concentrations of iron pisolites</td>
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Bedrock

Table 6
Figure 1
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Figure 3

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Figure 5
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