Re-analysis of late Quaternary dust mass accumulation rates in Serbia using new luminescence chronology for loess–palaeosol sequence at Surduk

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Loess–palaeosol deposits in the Carpathian Basin preserve some of the longest, most complete terrestrial records of Quaternary climate change in central Europe (Fitzsimmons et al. 2012; Marković et al. 2012, 2015). Palaeoenvironmental interpretations of these sedimentary deposits are most valuable when supported by detailed chronologies. Recent advances in radiocarbon dating of loess, utilizing earthworm granules (Moine et al. 2017) and mollusc shells (Újvári et al. 2014, 2017), are contributing to very high-resolution interpretations of the records. However, the absence of suitable material for radiocarbon dating at some locations, and an upper radiocarbon dating limit of ~50 ka, still present significant limitations to chronology generation. Further, radiocarbon dating provides ages for system closure, i.e. when the exchange of carbon with the environment ceased, which in most cases was when the organism died (Brock Ramsey 2008), rather than an age for the sediment deposition. Therefore, luminescence dating, which directly dates the burial of sediment grains, is often the most suitable geochronological approach in loess research.

Due to a better understanding of electron transfer within the crystal lattice (e.g. Aitken 1985, 1998; Preusser et al. 2008), and athermal stability (Wintle 1973; Murray & Wintle 1999), optically stimulated luminescence (OSL) dating of quartz is usually favoured over the infrared stimulated luminescence (IRSL) of feldspars. However, the use of IRSL signals from feldspars offers an opportunity to date much older material (<500 ka; Thomsen et al. 2011), as quartz reaches saturation at 100–200 Gy (Wintle & Murray 2006; Timar-Gabor et al. 2011). With the dose rates for quartz varying across this area are not consistent in timing or rates. The high-resolution dating strategy identifies a disturbance in sediment deposition that occurred after 45±2 ka and implies that site contains a hiatus. Finally, we show samples that failed routine dose recovery and preheat plateau tests, and had low fast ratios. Supported by bulk sample geochemical analysis it is proposed that a potential abrupt source shift, during the Last Glacial Maximum, may be the cause of the anomalous luminescence behaviour.

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decade, suggesting the need for many previously investigated sites to be revisited.

High-resolution, absolute ages provide a basis for mass accumulation rate (MAR) estimations (Kohfeld & Harrison 2003). MARs can provide an additional proxy for the past environmental change at a site (Újavári et al. 2017), as they ‘normalize’ the magnitude and rate of the changes. This not only allows the opportunity to investigate and correlate changes between different loess sites, but also other dust record archives (Frechen et al. 2003; Újavári et al. 2010; Pigati et al. 2013; Perić et al. 2019). These data sets also feed into global palaeo-dust cycle models, providing information on the global and regional dust fluxes, and interactions between the atmosphere and dust (Mahowald et al. 2006; Muhs 2013; Albani et al. 2015).

We present a new luminescence chronology for the loess–palaeosol profile at Surduk, Serbia. A published chronology from a profile ~2 km from our site (Fuchs et al. 2008) relied on 10 OSL ages determined using an uncorrected IR$_{50}$ signal obtained using an older, multi aliquot additive-dose measurement protocol (MAAD). Subsequently, research by Antoine et al. (2009) supplemented the original chronology with radiocarbon ages and showed that the luminescence chronology was likely underestimating the burial age due to the use of the uncorrected IRSL signal. Here, we present a new chronology based on 13 quartz OSL and 10 polimical pIRIR$_{220}$ ages. Based on this new chronology, MARs are calculated and discussed in the context of regional and global dust records.

Regional setting and site description

Surduk is part of a long stretch of thick loess deposits along the western bank of the Danube in Serbia, with a number of other sites such as Batajnica (Marković et al. 2009) and Stari Slankamen (Marković et al. 2011) previously investigated. The site (latitude 45$^\circ$4′50.24″N, longitude 20$^\circ$18′50.12″E, altitude 107 m a.s.l.) is situated 2 km northwest from a previously studied profile at Surduk, which was investigated for particle size, magnetic susceptibility, total organic carbon, carbonate content, and carbon isotopes (Fuchs et al. 2008; Antoine et al. 2009; Hatté et al. 2013). Due to access constraints the original site, referred to here as Surduk 1, could not be investigated. The new location analysed here is referred to as Surduk 2.

Surduk 2 is located 7 km downstream of the Danube’s confluence with the Tisza River and ~20 km from the Titel Loess Plateau (Fig. 1). The selected loess–palaeosol exposure is located in a gully along a road cutting 200 m from the main river channel. Figure 2 shows a simplified sedimentary succession and sample positions in the investigated Surduk 2 profile. The profile was sampled as two sub-profiles, A and B (Fig. 2), 10 m apart. While fieldwork aimed to ensure the two profiles overlapped, we subsequently suspect that there is a gap of up to 1.5 m between the profiles.

Luminescence dating

Sampling, preparation and measurement

Surduk 2 was sampled for luminescence dating using opaque light tight plastic tubes, hammered into the cleaned loess–palaeosol profile. Seventeen samples were collected at a 30-cm resolution throughout the profile, access permitting. All samples were prepared and analysed at the Oxford Luminescence Dating Laboratory, University of Oxford, under subdued orange-light conditions. To avoid light contamination, sediment from both ends of the sampling tubes was removed and used for particle size and dose rate analysis. All sediment was treated with 32% HCl (~2 h) and 30% H$_2$O$_2$ (up to 2 weeks), to remove carbonates and organic matter, respectively. Sieving and settling was used to isolate 4–11 μm grain sizes, which were divided into two fractions: polimical and quartz. To prepare the quartz enriched fine-grained fraction, samples were additionally treated with H$_5$SiF$_6$ (up to 2 weeks) to remove non-quartz minerals, followed by a brief 32% HCl (~2 h) immersion to dissolve any potential fluorite precipitates. In both polimical and quartz cases, sediment was dispensed onto the surface of 9.7 mm aluminium discs.

Luminescence measurements were made using Riso TL/OSL readers fitted with a calibrated $^{90}$Sr/$^{90}$Y beta source and a bialkali photomultiplier tube. For all quartz analyses, OSL signals were measured in a UV detection window through 7.5 mm U-340 glass filters (Bottjer Jensen et al. 2000) by stimulating with blue-light emitting diodes (470 nm). In the case of polimical aliquots, IRSL signals were detected in the blue-violet region of the electromagnetic spectrum through a combination of Schott BG39/Corning 7-59 filters by stimulation with infra-red light emitting diodes (870 nm).

Dosimetry

Light-exposed sediment from the sampling tubes was dried, homogenized and used to determine radionuclide concentrations for all samples. Concentrations of uranium, thorium, potassium and rubidium were determined by inductively coupled plasma mass spectrometry (ICP-MS) at the British Geological Survey, Keyworth, and converted to infinite-matrix dose rates using the conversion factors of Guérin et al. (2011). These dose rates were adjusted for alpha efficiency using a-values of 0.04±0.04 for quartz and 0.11±0.02 for polimical fractions (Kreutzer et al. 2014), grain size attenuation for alpha dose rates by Brennan et al. (1991) and Mejdahl (1979) for beta dose rates, and a typical loess moisture content of 15±5% (e.g. Stevens et al. 2011; Schatz et al. 2012; Újavári et al. 2014). Geographical
location, sediment thickness, and altitude were used to calculate the cosmic dose rate (Prescott & Hutton 1994). All dose rates were calculated using the DRAC (v1.2) software (Durcan et al. 2015).

**OSL measurements**

All of the tests and equivalent doses ($D_e$) for quartz aliquots were determined using the single aliquot regenerative dose (SAR) protocol (Murray & Wintle 2000; Wintle & Murray 2006). In all cases, luminescence was measured at 125 °C for 40 s following a preheat, while a 160 °C cut-heat was applied for the test dose. $D_e$s were determined by integrating the first 0.5 s of stimulation and by subtracting the background calculated from the last 20 s of stimulation. Depending on the fit, an exponential or an exponential plus linear function was used to fit the dose-response curves.

**Preheat plateau**

To determine the most appropriate temperature conditions for the quartz SAR protocol, a preheat plateau test for samples SER16/2/1, SER16/2/7 and SER16/2/9 was conducted. Aliquots were bleached twice for 1000 s with blue light stimulation at 50 °C with a 10 000 s pause in between, before administering a dose approximating a natural one: ~65, ~149 and ~119 Gy, respectively. Three aliquots for each of the six tested temperatures were measured using the SAR protocol described above. Figure 3 shows the results of the quartz preheat plateau. Samples SER16/2/1 and SER16/2/9 show a similar pattern with a drop in the ability to recover the given dose at intermediate temperatures. While a number of temperatures appear suitable, further dose recovery tests at 200 and at 260 °C showed that the latter recovers the dose more accurately. On that basis, a 260 °C preheat temperature was selected for all further tests and $D_e$ determination. In contrast, sample SER16/2/7 performed poorly across the preheat temperature range (Fig. 3) and did not recover the administered dose. Additionally, initial $D_e$ tests showed that all measured discs failed the recycling ratio, suggesting $D_e$ determination using the SAR protocol is not appropriate for this sample. Therefore, no further measurements using the quartz OSL signal were made for this sample.
Dose recovery

To examine the suitability of the selected preheat temperature, and therefore the SAR protocol’s ability to recover a laboratory administered dose, dose recovery tests were performed. Three samples (12 aliquots from each) that capture the age spread of the profile were selected. As in the case of the preheat plateau, aliquots were bleached twice and dosed with levels approximating those naturally occurring: ~65 Gy (SER16/2/1), ~69 Gy (SER16/2/3) and ~92 Gy (SER16/2/9). Almost all discs recovered the administered dose within uncertainty (Fig. 4), with an average measured to given dose ratio of 0.94/0.06 (n = 36, Table 1), demonstrating the suitability of the quartz SAR protocol to recover a laboratory administered dose.

Quartz signal assessment and provenance

The quartz OSL signal comprises a number of discrete components, fast, medium and slow (e.g. Smith & Rhodes 1994; Bailey et al. 1997). Signal from the fast component is preferred for dating because it bleaches rapidly in nature and is thermally stable (Wintle & Murray 2006). To quantitatively test for the dominance of the fast component in the initial part of the OSL signals, the fast ratio (Durcan & Duller 2011) was calculated with the R calc_FastRatio script (King et al. 2019). In addition, continuous-wave OSL signals were fitted with the sum of exponentials using the fit_CW-Curve script (Kreutzer 2019) to assess any variability in the number of components identified, and the relative contribution of signal from the various components to the total OSL signal.

The majority of measured quartz signals decayed rapidly to background levels within a few seconds (Fig. 5A), and qualitatively, this suggests dominance of the fast component. This qualitative assessment is confirmed by the fast ratio, calculated for both natural and regenerated signals, of selected representative samples from Surduk 2A and Surduk 2B (SER16/2/1, SER16/2/2, SER16/2/3, SER16/2/4, SER16/2/7, SER16/2/8, SER16/2/9, SER16/2/23 and SER16/2/20). Averages above 20 were calculated for all samples, except for the sample SER16/2/8. However, there was a lot of variability between individual discs especially in the samples from Surduk 2B, e.g. for SER16/2/23 fast values range between 8.3/2.9 and 357.9/530.5. Sample SER16/2/8 had a notably less bright OSL signals and decayed more slowly than other samples (Fig. 5D), with 13 out of 15 signals displaying a fast ratio below 20 (FR average = 13.1±2.3). This suggests that signal from the fast component does not dominate the initial part of the OSL signal, which may result in erroneous D_S determination (Durcan & Duller 2011). On this basis, quartz D_S for this sample are considered unreliable.

The application of routinely applied tests such as preheat plateau, dose recovery, and the fast ratio provides grounds for signal rejection, but does not provide insights into the cause of the signal problems. One avenue to explore is sediment geochemical compo-
sition to investigate any potential causes of this abrupt shift in quartz behaviour. While much work has been undertaken to establish links between provenance and luminescence dating (cf. Gray et al. 2019), only a few studies have looked at geochemical properties for comparison (Götte & Ramseyer 2012; Stevens et al. 2013; Rodrigues et al. 2019).

Bulk sample geochemical data from ICP analyses were examined in more detail. Trace elements, normalized relative to the upper continental crust (UCC), are shown in Fig. 6A. Negative anomalies relative to the UCC can be seen for Rb, Ba and Sr for all the Surduk 2 samples, which is likely related to the removal of these elements as part of the weathering process and/or low concentrations of feldspars in the samples. The relatively high enrichment of transitional elements (V, Cr and Ni; Fig. 6A) suggests a mafic character of the source rocks (Gasparon et al. 1993; Bracciali et al. 2007), especially in samples SER16/2/7, SER16/2/8 and SER16/2/9. Further, sample SER16/2/8 is also strongly enriched in all light rare earth elements (LREE) especially Eu, and has one of the higher Eu anomalies (0.77, Table S1). The Eu anomaly, calculated by Eu/Eu* (where Eu* is (SmN*GdN)0.5), provides a measure of Eu3+ fractionation from Eu3+ relative to the neighbouring elements in chondrite normalized rare earth elements (REE). In a sedimentary setting, the calculated values are indicative of source rock types (Kasanzu et al. 2008), with a negative anomaly associated with felsic rocks (feldspar bearing rocks), and no anomaly with mafic inputs (Cullers 1994; Gao & Wedepohl 1995; Gallet et al. 1998). The negative Eu/Eu* values suggest felsic inputs in all samples, although samples with the highest values (e.g. SER16/2/8) indicate incorporation of more mafic material. Thus, these samples might have slightly different source rocks, or additions from another source that is affecting the bulk sample geochemical signals. While links between provenance change and quartz signal characteristics are not clear yet, a mechanism could be envisaged where non sensitised quartz grains from smaller, proximal source rocks are added (Pietsch et al. 2008; Fitzsimmons 2011).

It is beyond the scope of this paper to explore the properties (e.g. components, sensitivity) of the luminescence signal in quartz, and how these are influenced by changes in geochemistry of the sediment or the provenance. However, this section shows the potential for linking geochemical properties with luminescence properties and highlights the need for further investigation.

On the basis of preheat plateau tests, dose recovery tests, the fast ratio and bulk sample geochemistry, quartz OSL signals are considered suitable for dating for most of the samples in Surduk 2A, where calculated Des are less than 100 Gy, and are not in saturation. Therefore, ages from Surduk 2A are based on quartz signals, apart from samples SER16/2/7 and SER16/2/8, which are excluded from further analyses due to the poor preheat test and recycling ratio performances (SER16/2/7), and non-dominance of the fast component (SER16/2/8). In section Surduk 2B, quartz De values exceed 175 Gy for all samples (Table 2). Multiple studies have suggested that quartz OSL ages become less reliable when the De is in the 100–200 Gy bracket (e.g. Buylaert et al. 2012) due to signal saturation, and the large asymmetric uncer-
tainties that result from interpolating onto the asymptote of the dose-response curve (e.g. Fig. 5A). Therefore, for samples with D_{s} above 100 Gy and SER16/2/7, the D_{e} was determined using 4–11 μm polymineral fraction and the pIRIR signal.

pIR-IRSL measurements

D_{s}s from polymineral aliquots were calculated using a modified pIRIR SAR protocol (after Thomsen et al. 2008; Buylaert et al. 2012). In all cases, after a preheat, IR_{50} and pIRIR_{elv} signals were measured for 200 s. The same sequence was followed for the test dose measurement with the addition of a 290 °C ‘hot-bleach’ (for 200 s) at the end of each cycle. Signals from the first 2 s of stimulation were integrated, with the background taken from the final 100 s of measurements subtracted. An exponential function was used to fit dose-response curves.

Preheat temperature selection

A number of loess luminescence studies from the Carpathian Basin have reported problems with dose recovery when using an elevated temperature pIRIR protocol (Stevens et al. 2011; Schatz et al. 2012; Murray et al. 2014; Újvári et al. 2014), especially the 320 °C preheat – 290 °C stimulation temperature combination. Further, research by Roberts (2012) demonstrated the influence of preheat temperature in the pIRIR protocol on the final D_{e} value, and therefore the importance of a preheat plateau test for feldspar and polymineral samples. The impact of temperature on D_{s}s was tested through preheat plateau (Fig. 7). Six preheat and elevated pIRIR temperature combinations (three aliquots for each) were analysed, with the pIRIR measurement temperature ~25–30 °C lower than the preheat temperature. For preheats above 280 °C, higher D_{s}s were measured with the pIRIR_{290} protocol (preheat of 320 °C) producing D_{s}s almost twice as large (274±76 Gy) as those in the 225–280 °C temperature range (average 153±5 Gy). These results support observations of Murray et al. (2009) and Roberts (2012) who also noted changes in D_{s}s with an increased preheat temperature, which is most likely resulting from sensitivity change occurring between Ln and Tn cycles, which cannot be measured and therefore accounted for (Roberts 2012). The overlapping D_{e} values for 225–280 °C preheats suggests the preheat plateau is achieved for these temperatures. Therefore, two of the lower temperatures, 220 and 250 °C, were selected for further dose recovery testing.

Residuals

Buylaert et al. (2011) and Sohbati et al. (2012) indicated that parts of the feldspar signal may be unbleachable, which may affect the ability to recover given doses. Further, if these ‘residual’ components are large enough they could have an effect on D_{e} calculation, and therefore age. While many regional loess studies have reported
minimal unbleachable components (Schatz et al. 2012; Murray et al. 2014; Ujvári et al. 2014), some have shown quite large residual signals e.g. 15–20 Gy (Austria; Thiel et al. 2011) and up to 40 Gy (in some Serbian samples; Stevens et al. 2011), demonstrating the importance of testing for this component.

Table 1. Residual, dose recovery and fading results for selected samples. Fading values and errors reported as means calculated based on three aliquots. * = residual subtracted dose recovery ratio.

| Sample       | Quartz Dose recovery* | Residual dose (Gy) | Polymineral Dose recovery* | IR50 g-value (%) | pIRIR225 g-value (%) |
|--------------|-----------------------|--------------------|---------------------------|------------------|----------------------|
| SER16/2/1    | 0.96±0.06             | –                  | –                         | –                | –                    |
| SER16/2/3    | 0.91±0.06             | –                  | –                         | –                | –                    |
| SER16/2/7    | 0.94±0.06             | 1.10±0.16          | 1.08±0.04                 | 13.0±0.9         | 2.1±1.1              |
| SER16/2/9    | 0.94±0.06             | 1.25±0.31          | 0.87±0.05                 | 16.3±0.9         | 0.9±1.1              |
| SER16/2/10   | –                     | 2.07±0.17          | 0.87±0.04                 | 10.2±0.9         | 2.5±1.2              |
| SER16/2/12   | –                     | 1.23±0.16          | –                         | 5.4±1.2          | 1.8±1.0              |
| SER16/2/17   | –                     | 1.23±0.16          | –                         | 10.2±0.9         | 2.5±1.2              |
| SER16/2/23   | –                     | 1.23±0.16          | –                         | 10.2±0.9         | 2.5±1.2              |
| Average      | 0.94±0.06             | 1.41±0.20          | 0.92±0.04                 | 11.37±0.99       | 1.87±1.11            |

Fig. 5. A. Decay and dose-response curves for quartz (SER16/2/4). B. Decay curve for a typical quartz disc from sample SER16/2/7 with fitted components; insert shows component contributions to the total curve and residuals from the component fitting. C. Fast ratio from the natural signal measurement as a function of fast ratio from the nearest regeneration point for sample SER16/2/7. D. Decay curve for typical quartz disc from sample SER16/2/8 with fitted components; insert shows component contributions to the total curve and residuals from the component fitting. All fast ratios are presented with absolute error values.
To test the size of the residual component in the Surduk 2 samples, six aliquots from four samples, SER16/2/9, SER16/2/10, SER16/2/17, SER16/2/23, were bleached in daylight for 7 days (during May in the UK), and measured using the pIRIR protocol. Of the 24 aliquots measured, 18 passed the recycling ratio, but none passed a 5% recuperation ratio. As the initial assessment of the signal suggested small natural signals, all recuperation values were converted into absolute values (in Gy) to test for absolute size.

The average residual component measured is $1.41 \pm 0.20$ Gy ($1.10 \pm 0.16$–$2.07 \pm 0.17$ Gy range; Table 1), equivalent to $\sim 0.4$ ka (when calculated using the average polymineral dose rate). These results show that the unbleachable component has a negligible impact on the final ages. It is not possible to compare these values with the previous Surduk 1 study as the residual test was not performed (Fuchs et al. 2008). Previously reported Serbian values range between 1 and 40 Gy (Stevens et al. 2011; Murray et al. 2014; Böskén et al. 2017), and along with our results, demonstrate that residual components are sample and site specific, and should be routinely assessed.

**Dose recovery**

As with quartz, the suitability of the pIRIR protocol was tested using a dose recovery test. First, two pIRIR protocols, pIRIR$_{200}$ (225 °C preheat) and pIRIR$_{225}$ (250 °C), were tested using sample SER16/2/9. Six aliquots were bleached in daylight for 7 days (during May in the UK), irradiated with a laboratory beta-dose approximating their natural one, and measured using the two protocols. Figure 8 shows that the pIRIR$_{225}$ protocol struggled to recover the given dose with an average given to measured ratio of 1.38 ± 0.06. Aliquots measured using the pIRIR$_{200}$ protocol recovered the laboratory dose, giving an average ratio of 1.08 ± 0.04. Subtracting the average residual value from the measured equivalent dose prior to the calculation of the dose recovery takes the ratios down to 1.36 ± 0.06 for pIRIR$_{225}$, while the pIRIR$_{200}$ remains unchanged due to the very small size of residual dose. These results show that the lower preheat temperature better recovers a known dose for this sample.

Following selection of the pIRIR$_{200}$ protocol an additional 12 aliquots per sample (SER16/2/10 and

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*Fig. 6. Bulk sample trace elements UCC-normalized spider diagrams for Surduk 2. A. Large-ion lithophile elements (LILE) and high-field-strength elements (HFSE) elements. B. Light earth elements (LREE) and heavy earth elements (HREE).*
Table 2. Samples, depths, radionuclide concentrations, dose rates, $D_e$ values and the final luminescence ages obtained from the 4–11 μm quartz (OSL protocol) and polymineral fractions (pIRIR$_{220}$ protocol). Values are presented to two decimal places, and all calculations were made prior to rounding. Based on the results of the fading correction no fading corrections have been made and therefore the pIRIR$_{220}$ age reflects the uncorrected age. * = overdispersion.

| Sample     | Depth (m) | Unit | U (ppm ±1σ) | Th (ppm ±1σ) | Rb (ppm ±1σ) | K (% ±1σ) | Quartz | Total dose rate (Gy ka$^{-1}$) | OD %* | $D_e$ (Gy) | OSL age (ka) | Total dose rate (Gy ka$^{-1}$) | Polyminal | OD %* | $D_e$ (Gy) | pIRIR$_{220}$ age (ka) |
|------------|-----------|------|--------------|--------------|--------------|-----------|--------|--------------------------------|-------|-------------|-------------|--------------------------------|-----------|-------|-------------|------------------------|
| SER16/2/1  | 1.5       | II   | 1.91±0.19    | 9.06±0.91    | 74.64±7.46   | 1.88±0.19  | 0.18±0.02 | 3.19±0.22                     | 9     | 64.52±1.88  | 20.23±1.49   | –                                 | –         | –     | –            | –                      |
| SER16/2/2  | 1.8       | II   | 1.86±0.19    | 9.29±0.93    | 76.73±7.67   | 2.08±0.21  | 0.17±0.02 | 3.36±0.23                     | 1.5   | 67.69±0.93  | 20.13±1.40   | –                                 | –         | –     | –            | –                      |
| SER16/2/3  | 2.1       | II   | 2.54±0.25    | 9.20±0.92    | 78.64±7.86   | 2.27±0.23  | 0.16±0.02 | 3.72±0.26                     | 3.3   | 70.13±1.25  | 18.85±1.33   | –                                 | –         | –     | –            | –                      |
| SER16/2/4  | 2.4       | II   | 2.21±0.22    | 10.02±1.0    | 83.43±8.34   | 2.47±0.25  | 0.16±0.02 | 3.87±0.27                     | 5.6   | 86.43±1.90  | 22.35±1.61   | –                                 | –         | –     | –            | –                      |
| SER16/2/5  | 2.7       | II   | 2.10±0.21    | 10.02±1.0    | 87.78±8.78   | 2.26±0.23  | 0.15±0.02 | 3.64±0.25                     | 4.4   | 82.05±2.23  | 22.52±1.66   | –                                 | –         | –     | –            | –                      |
| SER16/2/6  | 3.2       | II   | 2.06±0.21    | 10.19±1.02   | 94.94±9.49   | 2.27±0.23  | 0.15±0.02 | 3.65±0.25                     | 7.5   | 85.27±2.07  | 23.37±1.70   | –                                 | –         | –     | –            | –                      |
| SER16/2/7  | 3.3       | II   | 2.04±0.20    | 10.26±1.03   | 95.80±9.58   | 2.38±0.24  | 0.14±0.01 | –                               | –     | –            | –            | 4.40±0.28                   | 2.6       | 99.24±1.39 | 22.57±1.46   | –                      |
| SER16/2/8  | 3.6       | II   | 2.14±0.21    | 11.82±1.18   | 100.03±1.00  | 2.43±0.24  | 0.14±0.01 | 3.95±0.27                     | 5.4   | 84.58±1.96  | 21.43±1.57   | –                                 | –         | –     | –            | –                      |
| SER16/2/9  | 3.9       | II   | 2.08±0.21    | 11.08±1.11   | 95.52±9.55   | 2.37±0.24  | 0.13±0.01 | 3.81±0.26                     | 7.4   | 92.73±2.36  | 24.35±1.79   | –                                 | –         | –     | –            | –                      |
| SER16/2/10 | 4.4       | IV   | 1.86±0.19    | 9.79±0.98    | 81.13±8.11   | 1.92±0.19  | 0.14±0.01 | 3.23±0.22                     | 0     | 175.66±3.63 | 54.45±3.92   | 3.84±0.24                   | 9         | 174.34±5.87 | 45.46±3.26   | –                      |
| SER16/2/11 | 4.8       | IV   | 1.89±0.19    | 9.47±0.95    | 78.49±7.85   | 1.83±0.18  | 0.13±0.01 | 3.20±0.22                     | 17    | 120.61±7.50 | 38.60±3.58   | 3.73±0.24                   | 3.6       | 176.16±2.80 | 47.29±3.09   | –                      |
| SER16/2/12 | 5.2       | IV   | 1.81±0.18    | 9.67±0.97    | 83.85±8.39   | 1.93±0.19  | 0.13±0.01 | 3.20±0.22                     | 4.6   | 169.94±3.89 | 53.09±3.87   | 3.80±0.24                   | 4         | 179.77±3.29 | 47.33±3.12   | –                      |
| SER16/2/13 | 5.5       | IV   | 1.98±0.20    | 9.73±0.97    | 76.73±7.67   | 1.80±0.18  | 0.12±0.01 | –                               | –     | 3.24±0.26   | 48.94±3.24   | –                                 | 0         | 187.56±2.46 | 49.94±3.24   | –                      |
| SER16/2/14 | 5.8       | IV   | 1.98±0.20    | 9.42±0.94    | 80.88±8.09   | 1.82±0.18  | 0.12±0.01 | –                               | –     | 3.48±0.23   | 49.77±3.22   | –                                 | 1.3       | 186.19±2.45 | 49.77±3.22   | –                      |
| SER16/2/20 | 6.1       | IV   | 1.92±0.19    | 9.41±0.94    | 78.42±7.84   | 1.80±0.18  | 0.12±0.01 | 3.09±0.21                     | 0     | 200.13±3.94 | 64.83±4.67   | 3.69±0.23                   | 0         | 191.85±2.56 | 51.99±3.37   | –                      |
| SER16/2/21 | 6.4       | IV   | 1.89±0.19    | 10.11±1.01   | 85.83±8.58   | 1.90±0.19  | 0.11±0.01 | –                               | –     | 3.85±0.25   | 50.95±3.32   | –                                 | 2         | 196.20±2.64 | 50.95±3.32   | –                      |
| SER16/2/22 | 7.2       | V    | 1.99±0.20    | 10.48±1.05   | 88.84±8.88   | 1.85±0.19  | 0.11±0.01 | 3.23±0.23                     | 4     | 193.33±5.03 | 59.78±4.46   | 3.89±0.25                   | 0         | 199.32±2.75 | 51.30±3.36   | –                      |
SER16/2/17) were analysed following the same preparation and measurement protocol. The average measured to given dose ratio for these samples was 0.92 ± 0.04 \((n = 18)\), suggesting a satisfactory pIRIR protocol for \(D_e\) measurement.

**Laboratory fading**

Laboratory fading rates \((g^2/\text{days})\) were tested for the pIRIR\(_{200}\) signal, using the same protocol used for \(D_e\) measurements. Following Auclair et al. (2003) three aliquots per sample were irradiated with a fixed dose \((-90 \text{ Gy})\), a test dose \((-40 \text{ Gy})\) and measured following a series of pauses \((0, 1, 10, 100 \text{ and } 1000 \text{ h})\). The mean fading rates for the IR\(_{50}\) and pIRIR\(_{200}\) signals are 11.37 ± 0.99%/decade and 1.87 ± 1.11%/decade \((n = 15, \text{ five samples})\), respectively. The range of the fading rates for the IR\(_{50}\) signal varies between 0.43 ± 0.94 and 21.53 ± 0.74%/decade, whereas the pIRIR\(_{200}\) results range from 0.27 ± 0.92 to 3.46 ± 1.23%/decade (Fig. 9, Table 1). These results demonstrate that IR\(_{50}\) signals undergo high rates of fading at Surduk 2. In contrast, fading rates for the pIRIR\(_{220}\) are sufficiently low that they do not require a fading correction, following the arguments of Buylaert et al. (2012) and Thiel et al. (2011).

**\(D_e\) and age calculation**

All measured aliquots \((n = 345)\) were screened using signal recuperation \((<5\%)\), a recycling ratio (Murray & Wintle 2000), and additionally for quartz, an OSL IR depletion ratio (Duller 2003). In total, 24 signals were rejected due to a failed recycling ratio, and six signals failed recuperation. All but five quartz signals passed the IR depletion ratio. Overdispersion (Table 2) was found to be below 10% (except for the quartz sample SER16/2/23) and therefore all equivalent doses were calculated using the central age model (CAM; Galbraith et al. 1999). CAM \(D_e\)'s were divided by the environmental dose rate to derive the age. As discussed, \(D_e\)'s were calculated from quartz for Surduk 2A samples, with the exception of SER16/2/7 and SER16/2/8. For all other samples, polymineral pIRIR signals were used.

**Age-depth-model and MARs**

All final luminescence ages were re-calculated using Bayesian and inverse modelling (Zeeden et al. 2018), which separates systematic and random errors of luminescence ages. This model creates probability density functions for both types of error but models only the random component of the uncertainty, making this approach better suited for the analysis of luminescence ages than other Bayesian models such as OxCAL (Bronk Ramsey 1995). Due to the lack of overlap and to uncertainty regarding the sediment thickness, two separate models were created: one for Surduk 2A and the second for Surduk 2B. These age-depth models provide a basis for calculating sediment accumulation rates, as well as a better visual representation of the age distribution throughout the profile.

Loess–palaeosol sequences can preserve information about dust fluxes, yet to date no sites from Europe have
been included in the global palaeodust cycle modelling efforts (Albani et al. 2015), mostly resulting from the lack of high-resolution chronologies. To aid those efforts, a number of loess studies (Újaví et al. 2015; Stevens et al. 2016; Perić et al. 2019) have converted age-depth models into MARs. Here, modelled luminescence ages

![Figure 8](image8.png)

**Fig. 8.** Polyminal dose recovery for samples SER16/2/9, SER16/2/10 and SER16/2/17. Note dose recovery tests for the pIRIR$_{220}$ and pIRIR$_{225}$ protocols were carried out using sample SER16/2/9. Dotted lines mark ±10% from unity. All samples had a residual dose subtracted prior to the calculation.

![Figure 9](image9.png)

**Fig. 9.** Calculated fading rates for the IR$_{50}$ and pIRIR$_{200}$ luminescence signals for all aliquots.
were used to calculate MARs (Kohfeld & Harrison 2003) using the following equation

$$\text{MAR} (g m^{-2} a^{-1}) = SR \times f_{\text{eol}} \times \rho_{\text{dry}} \quad (1)$$

where SR is the sedimentation rate (m a^{-1}), $f_{\text{eol}}$ is the fraction of the sediment that is aeolian in origin, and $\rho_{\text{dry}}$ is the bulk density of dry sediment (g m^{-3}). As the sediment is interpreted as loess and therefore aeolian in origin, $f_{\text{eol}} = 1$. A bulk density value of 1.5 g cm^{-3} was used in all calculations based on the reported estimates of loess bulk density in Hungary and Serbia (Újvári et al. 2010; Perić et al. 2019). Finally, in desert research it has been shown that using only mean ages to construct sedimentation rates for sand dune accumulation may provide an inaccurate picture of aeolian and climatic history (Leighton et al. 2014), therefore for the first time in loess research two MARs are calculated based on mean and minimal sedimentation rates. Mean sedimentation rates are calculated as $\text{SR}_{\text{mean}} = x_1 - x_2/\gamma_1 - \gamma_2$, where $x_{1,2}$ are two depths and $\gamma_{1,2}$ corresponding ages in the profile. The $\text{SR}_{\text{min}} = x_1 - x_2/((\gamma_1 + \sigma_1) - (\gamma_2 - \sigma_2))$ also includes age errors for each relative age.

To ensure a meaningful and fair comparison of MARs between loess studies, only sites with a high-resolution, absolute chronology were selected. Further, sites where the luminescence chronology used an IR_{50} signal that had not been tested for fading were also excluded. Therefore, three sites, Crvenka (Stevens et al. 2011), Dunaszécső (Újvári et al. 2017) and Titel (Perić et al. 2019), were selected for MAR assessment.

**Palaeoenvironmental results and discussion**

**A new chronology and sedimentation rates for the Surduk 2 loess-palaeosol sequence**

Unmodelled quartz and polymineral pIRIR_{200} luminescence ages for Surduk 2 are presented in Table 2. The preferred (see Quartz quartz signal assessment and provenance discussion) unmodelled and modelled ages are presented in Fig. 10 and Table S2. Sixteen ages span from the top of the transitional palaeosol (Unit III) to 1.5 m below the modern-day surface. All but one of the ages come from loess Unit II, with the youngest age 19.03±1.03 ka. The bottom age, for the transitional zone between palaeosol and loess, places its burial at 52.55±2.71 ka.

The ages for both parts of the profile form two stratigraphically consistent depositional groups. The first (Surduk 2A) covers 1.5 m within the top ~4 m and represents the period between 19.03±1.03 and 24.62±1.31 ka. The second group (Surduk 2B) consists of eight samples, spread over 2.8 m, with ages between 44.99±2.20 and 52.55±2.71 ka. These two groups are separated by a sampling gap that is estimated to be between 1 and 1.5 m. The comparison of ages above and below shows relatively rapid accumulation rates and suggests that within that packet of sediment a change occurred. Without further high-resolution chronological investigation at this stage, it is not possible to definitively determine whether this gap represents a period of sediment accumulation lows, driven by environmental or climatic factors, or if this is an erosion boundary between different periods of dust accumulation. However, it is most likely that an erosional boundary is present above L1LL2, supporting a previously argued notion that loess sequences are neither continuous nor homogenous, and are likely to preserve disturbances in deposition (Stevens et al. 2008, 2018).

**Comparison with other sites and implications**

The oldest age obtained from Surduk 2B comes from the transitional palaeosol-loess zone near the bottom of the profile and dates sediment burial to 52.55±2.71 ka. The calcification and compaction of the palaeosol below precluded collection of samples and determination of the beginning of deposition for this part of the profile/sequence. The comparison of the basal parts between Surduk 1 and 2 shows a large age discrepancy (Fig. 10). Based on stratigraphical position, the bottom palaeosol at Surduk 2B may correspond either with the ‘Basal soil complex’ or the ‘Middle soil complex’ at Surduk 1, and therefore be interpreted as either S1 or L1SS2 palaeosol.

The loess deposition on top of the ‘Basal soil complex’ at Surduk 1 was dated to 82.6±9.0 ka (Fig. 10). Due to the blanket nature of loess deposition, it is highly unlikely that there is a ‘real’ ~30-ka discrepancy between Surduk 2 and Surduk 1 deposits, especially given that they are located so closely together. The luminescence investigation at Surduk 1 did not test the IR_{50} signal for fading and ages were not corrected (Fuchs et al. 2008). Fading tests at Surduk 2 show that the polymineral IR_{50} signal is athermal unstable (Fig. 9). Similar fading behaviour can be expected from Surduk 1, and it is likely that Surduk 1’s chronology underestimates the true ages. Therefore, the age of 82.6±9.0 ka for the ‘Basal soil complex’ at Surduk 1 is likely to be too young. However, without fading rate data for those samples, combined with the old age of the samples (and therefore the difficulties in correcting the non-linear part of the dose-response curve), it is not reasonable to speculate upon the degree of under-estimation of each age at Surduk 1 in Fuchs et al. (2008). Nonetheless taking the fading rates broadly into consideration and the depths of the units at both profiles, it is likely that the Unit VI palaeosol at Surduk 2B corresponds with the ‘Middle soil complex’ at Surduk 1, dated to 36.3±4.1 ka (Fuchs et al. 2008).

Nevertheless, if the correlation based on stratigraphy was correct, it would suggest favourable pedogenic conditions at Surduk 2B persisting until 52.55±2.71
Fig. 10. Schematic stratigraphical diagram for Surduk 1 (Fuchs et al. 2008; Antoine et al. 2009) and Surduk 2. For the Surduk 2 quartz OSL (marked by *), polymineral pIRIR220 and Bayesian modelled ages (bold) are presented.
ka, and no loess deposits corresponding to the initial climate deterioration seen across the region and associated with MIS 3 (Bokhorst et al. 2011; Panagiotopoulos et al. 2014; Wegwerth et al. 2016). This could be a result of repeated erosional events at Surduk 2, possible due to the geomorphic position of the site. Alternatively, a number of loess profiles in central Europe e.g. Stalač (Bosken et al. 2017), Suttő (Novothny et al. 2009), and Stari Slankamen (Schmidt et al. 2010), indicate persistent pedogenesis until the MIS 3 stage, which may have also occurred at Surduk. However, without further geochronological investigation at Surduk 1, it is not possible to determine if the stratigraphic discrepancy between Surduk 1 and 2 palaeosols represents: multiple erosional events at Surduk 2; differences in pedogenic conditions; or a chronological discrepancy.

Following the transitional unit, primary loess deposition began at Surduk 2B after 52.55±2.71 ka and lasted at least until 44.99±2.20 ka (Fig. 10). At the beginning of this period, the mass accumulation rates were the highest in the profile, 1288 g m⁻² a⁻¹ (mean; minimum = 163 g m⁻² a⁻¹), after which they decreased in a step-like manner all the way up through Unit IV (Fig. 11). However, there are two brief periods of increased accumulation between 49.93±2.29 and 49.17±2.37 ka (mean 591 g m⁻² a⁻¹ and min. 174 g m⁻² a⁻¹) and 48.00±2.02 and 46.84±2.21 ka (mean 514 g m⁻² a⁻¹ and min. 173 g m⁻² a⁻¹). This indicates temporary environmental deterioration or increased supply of material during the early MIS 3. While a decrease in MAR values is also seen in Crvenka over this period (Stevens et al. 2011), the lack of a temporary increase could be attributed to the lower chronological resolution of the study or the site’s position. We propose that the increase in accumulation was locally driven and probably related to the site’s proximity to the Danube and sediment source. Further, it is important to note that while mean rates show a relatively large change in the MAR values, e.g. from 607 to 479 g m⁻² a⁻¹, the difference in minimum rates is very small (176 and 171 g m⁻² a⁻¹). Seeing that these rates are almost three times lower and do not really show corresponding trends, this highlights that care must be taken when interpreting MARs, especially when only mean rates are presented.

A hiatus in luminescence ages is observed between 44.99±2.20 and 24.62±1.31 ka. At present, it is not possible to determine if the stratigraphic discrepancy between Surduk 1 and 2 palaeosols represents: multiple erosional events at Surduk 2; differences in pedogenic conditions; or a chronological discrepancy.

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colder temperatures, increased precipitation, and changes in vegetation cover (Kageyama et al. 2006; Sümenegi et al. 2013, 2019; Zech et al. 2013). This can be seen represented at other sites such as Dunaszékeső (Újvári et al. 2017) or Titel (Perić et al. 2019) by an increase in MAR values, which is expected as the ice sheets grow to their maximum extent. Taking this regional picture into consideration, we suggest that Surduk 2 contains a sedimentological hiatus, and if a palaeosol was present, it has likely been eroded. It is not possible to determine how much sediment has been removed as only the remaining features are datable. However, a further higher resolution chronological investigation should be able to identify when exactly the gap occurs.

The top part of Unit II preserves ~5.5 ka of dust accumulation and represents the prelude to, and the main part of the Last Glacial Maximum (LGM). While not particularly thick in comparison to other loess sections recording this period of accumulation, it still indicates progressive and relatively rapid sediment accumulation from 24.62±1.31 ka (mean 744 g m⁻² a⁻¹; Fig. 11). This period of deposition also corresponds to a potential shift in provenance. Bulk sample geochemical analyses (Fig. 6) suggest sediment at that time had slightly more mafic and/or less felsic source rock inputs. The analysis did not imply complete change in provenance but rather the gap occurs.

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Following this period at 23.42±1.07 ka (Fig. 11), MARs rose to the second highest values for Surduk 2 (mean 1058 g m⁻² a⁻¹), indicating increased environmental instability. High MARs did not persist for long and began to drop after 22.99±1.01 ka and continued to fall until 20.28±0.92 ka (mean 333 g m⁻² a⁻¹). This was followed by a temporary rise to the second highest rate at Surduk 2A (mean 1036 g m⁻² a⁻¹ and min. 207 g m⁻² a⁻¹) that lasted until 19.85±0.81 ka. These results indicate that even during the LGM, which is thought to be the period of highest sediment accumulation, the deposition was not constant.

The values at Surduk 2 are as much as five times greater than those shown for this period at the nearby Titel Plateau (Perić et al. 2019), which are exceeded even by the conservative minimum values of 207 g m⁻² a⁻¹. While the Surduk 1 profile does not fully capture a chronology for this period, it still suggests increased wind speeds or delivery of coarser material to the system as a higher proportion of sands and reduction of clay is noted (Antoine et al. 2009), further supporting a lack of vegetation and cold/windy conditions.

The uppermost sample at Surduk 2, located 1.5 m below the modern land surface, indicates loess deposition at 19.03±1.03 ka (Fig. 10). It is expected that during deglaciation and a transition to the Holocene, sediment availability declined and depositional rates decreased as vegetation became more established. The transition to the Holocene palaeosol is not strongly marked or visible. A lack of thick well-developed Holocene soil, as well as a thicker deglaciation loess unit, is also seen at Surduk 1. Moreover, considering human activities on the modern land surface at both localities, it is likely that Holocene soil (S0) has been removed as part of agricultural processes.

**Comparison with Greenland and implications for MAR interpretations**

A number of studies have attempted to link central European loess deposits with Greenland dust records based on geochemistry (Újvári et al. 2015), MARs (Újvári et al. 2017), particle size (e.g. Rousseau et al. 2002; Antoine et al. 2009) and models (e.g. Sima et al. 2009) to show loess contributions to the global dust record. Figure 11 shows MARs from four sites around the southern part of the Carpathian Basin plotted against dust records from Greenland (Rasmussen et al. 2014).

Firstly, and most notably, it can be observed that peaks and troughs in accumulation do not always overlap. While some broad similarities are seen, records often conflict with each other, e.g. Surduk 2 MARs show increased accumulation followed by a decrease at around 23 ka; however, at the same time at Crvenka rates fall dramatically and stabilize for several thousands of years. It must be pointed out that with higher dating resolution less of the sedimentary sequence is averaged, hence records from sites such as Dunaszékeső show much more detail and variability by comparison. Even if broad trends can be seen there appears to be Carpathian Basin wide heterogeneity, with no obvious trends in accumulation seen over relatively short distances. This suggests that loess does not develop as a blanket feature at the same time across the landscape, as sediment is not transported to the sites uniformly. The comparison between sites in the Carpathian Basin thus suggests that accumulation at each site is likely driven by short-distance transport and local factors, such as sediment availability on flood-plains.

When further compared with the dust record from Greenland (Fig. 11) the picture is quite complex. The MAR patterns seen in the Carpathian Basin do not always mirror trends in Greenland. For example, during the drop in accumulation between 51.61±2.42 and 50.87±2.37 ka at Surduk 2 (shift from 1288 to 607 g m⁻² a⁻¹) values in Greenland remained relatively constant. At the same time while Surduk 2 notes high rates of accumulation the dust flux in Greenland is low. However, the drop-rise pattern between 50–47 ka at Surduk 2 approximates the pattern seen in Greenland. In both cases, increases in MARs at Surduk 2 precede changes in Greenland, although dust flux ceased at the same time.
The same pattern is also observed for peaks at 20.28±0.92 ka at Surduk 2A and smaller peak at ~28 ka at Dunaszékcső (Ujvári et al. 2017). It could be therefore argued that for some instances the record from the Carpathian Basin is preceding the changes that are eventually picked up in Greenland.

Lastly, it is important to note how much variability is seen between the mean and minimum MAR rates. Calculating minimum MAR values takes into account the uncertainty of ages and enables a fuller exploration of the geomorphological signal. For the most part, the same trends are seen between minimum and mean values but the discrepancy between the two values can reach almost two orders of magnitude (e.g. Dunaszékcső). This approach is not taken to discredit the mean values, but to highlight that the true MAR values are somewhere in between the minimum and maximum (which is often impossible to calculate). Therefore, while the trends can be explored when using mean values alone, interpretation of values where only mean values are presented should be taken with caution, especially when comparing with other dust records, such as Greenland. Finally, this point is particularly pertinent to the debate surrounding the contributions of European loess deposits to the dust recorded in Greenland (Sima et al. 2009; Albani et al. 2015; Ujvári et al. 2015, 2017). It is unlikely that a sole site was contributing greatly, but the lack of overlapping trends between sites and the discrepancy between minimum and mean rates suggest that European loess contributions are periodic, and secondary. However, more sites with higher resolution chronologies are needed to unpack this complex dust record, and its contribution to global dust archives.

Conclusions

This study presents a new high-resolution chronology for the loess–palaeosol sequence at Surduk 2, based on 13 quartz OSL and 10 palynometric pIRIR$_{2\alpha}$ ages, that provides a basis for a broad review of the palaeoenvironmental record preserved at this site. The modelled luminescence ages based on both protocols bracket the primary loess deposition between 52.55±2.71 and 19.03±1.03 ka. The results, however, suggest that this record is not continuous and contains a hiatus that occurred after 44.99±2.20 ka and prior to 24.62±1.31 ka. The bulk sample elemental analysis identified a potential abrupt but small provenance shift that occurred between 24.62±1.31 and 23.42±1.07 ka. Further, the high-resolution chronology provides a basis for calculation of MARs for Surduk 2 and comparison with other sites from the Carpathian Basin, Crvenka (Stevens et al. 2011), Dunaszékcső (Ujvári et al. 2017), Titel (Perić et al. 2019) and Greenland (Rasmussen et al. 2014). The analysis demonstrates that sites experienced increased periods of accumulation at different times, showing that sites’ depositional histories are not uniform, despite their relatively close locations. Therefore, preserved loess sequences likely depend on a site’s geomorphological position and palaeotopography, which control the appearance and intensity of erosional and pedogenic processes (Vandenberghe et al. 2014; Marković et al. 2018). While European loess contributions to Greenland are still debated, these results suggest that they do not play a major role. Finally, we suggest that additional chronological analysis is needed to identify the exact position of the hiatus and to determine how long this site was stabilized for prior to 53 ka and whether it experienced pedogenically favourable conditions continuously since the last interglacial.

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Author contributions. – SMB assisted KF with sample collection. All laboratory analysis, including sediment dating, writing, and figure and table preparation was conducted by KF. JAD assisted with luminescence experiments planning, and discussion of results. JAD, DSGT, ILM and SMB provided comments prior to paper submission.

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Supporting Information

Additional Supporting Information may be found in the online version of this article at http://www.boreas.dk.

**Table S1.** Raw bulk sample geochemical composition for loess–paleosol samples from Surduk 2 profile.

**Table S2.** The unmodelled and modelled ages for loess-paleosol samples from Surduk 2 profile.