Defining multiple inhabitations of a cave environment using interdisciplinary archaeometry: the ‘Christmas Cave’ of the Wadi en-Nar/Nahal Qidron, West of the Dead Sea

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Abstract
The present study reports a series of interdisciplinary archaeometrical analyses of objects found in the Christmas Cave, which was discovered by John Allegro and his team in 1960 on the West Bank of the Dead Sea and assumed to be inhabited only in the Chalcolithic era and by Jewish refugees of the second century CE, at the end of the Bar Kokhba Revolt. Like many other Judaean desert caves, there was an abundance of organic material, especially textiles, surviving in the dry environment. In the absence of clear stratigraphy and even a proper publication of the finds, the present study shows how archaeometry can provide important insights. We analysed food crusts on ceramics by gas chromatography with mass spectrometric detection (GC–MS), made petrographic descriptions to estimated provenance of the ceramics, produced new radiocarbon dates from organic material and thermoluminescence (TL) dates from the pottery. It appears from the data that the Christmas Cave has been briefly inhabited or visited intermittently over a very long time, starting ca. 4000 BCE (the Chalcolithic period), and extending all the way to the Medieval period, even though there is also a concentration of dates near the period of the Bar Kokhba Revolt (132–136 CE). We argue, through a detailed analysis of the radiometric and TL-datings and by the artefactual evidence, that there was likely another refuge episode connected with the First Jewish Revolt during which people fled to this cave. However, we see no material connection to Qumran and nearby caves. Overall, our study demonstrates the importance of archaeometric studies in cave environments where stratigraphy is veritably absent.

Keywords: Christmas Cave, Judean Desert, Bar Kokhba Revolt, Radiocarbon dating, TL-dating, Ceramic analyses, Organic residue analyses, Textiles

Introduction
Cave environments are often particularly challenging to archaeologists when it comes to the dating of artefacts found within them. Caves frequently lack significant stratigraphy, one of the cornerstones of defining human occupation, though at times this can be built up by series of rockfalls, debris or even collections of animal droppings. Therefore, when artefacts that do not show clear typological features are found, what appears to be an assemblage from one period can actually come from multiple times, and attest to different events of human activity, and from long term use to sporadic visits. In recent work on Cave 11Q near Qumran, on the coast of the Dead Sea, a range of scientific methods were employed on artefacts alongside classic typological

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analyses, revealing a far more complex picture of human inhabitation of the cave than had hitherto been assumed (see [1]). In the present study, we deploy a range of inter-disciplinary studies in key fields of archaeometry. We consider another cave in the Judaean wilderness, the so-called 'Christmas Cave', in order to probe whether use of these scientific methods can further nuance conclusions about human inhabitation of a cave environment, and the identity of the people who came here. As in other Judean Desert caves, the Christmas Cave is regarded as a ‘time capsule’ preserving various finds from several periods, but with little stratification, and little data from the cave has hitherto been published or studied.

The Christmas Cave: location and excavation
The Christmas Cave was discovered by John Allegro and Howard Stutchbury on Christmas Day of 1960, when they were engaged in excavations at Khirbet Mazin (Fig. 1). After a small initial sounding, it was excavated over two seasons by the British Dead Sea Scrolls Foundation (DSSF), under licence from Jordan, in December 1961 to January 1962 and again in December 1962 to January 1963. While the team's original hope was that this cave could yield more Dead Sea Scrolls, only tiny pieces of one papyrus written in Greek were found, and they never fully published either this or the other artefacts they excavated [2, 3].

The Christmas Cave is located in the eastern segment of Nahal Kidron/Qidron (Wadi an-Nar), near the top of the south-western bank of the wadi, before its last turn eastwards towards the lower canyon, at coordinates 239887/621095 (New Israel Grid), ca. 3 km west of the Dead Sea and 8 km SSW of Qumran. Allegro described the structure of the cave and also reported signs of looting conducted by Bedouins that preceded his visit ([2]: pages 11–13 [3]). Artefacts from the cave were in fact offered to the excavators in due course, and included wooden bowls, spindle whorls, a distaff, combs, kohl container and stick, a seal, beads, an ivory-inlaid box, coins, and an adze handle. The excavations proper brought to light numerous artefacts, amongst them textiles, wood, leather, basketry, metals, ceramics, glass, and flint. A bronze coin minted during the second year of the Bar Kokhba Revolt (132–136 CE), as well as Roman-period pottery, pieces of leather (some from shoes), small fragments of papyrus and a large number of textiles cumulatively corresponding with an interpretation of the cave as a primarily being a place of refuge in the Roman period, likely the second century. At this time, refugees fled to numerous caves of the Judean Desert after the revolt, led by Shimon ben Kosiba (called Bar Kokhba) was brutally quashed by Roman forces.

However, there was also a flint blade and a bone tool assigned to the Chalcolithic period, and the adze handle was also identified as Chalcolithic.

Part of the history of the discovery of the cave and of the excavations was briefly reported by Allegro in his popular book entitled Search in the Desert [2], though Allegro did not present the actual findings and excavations accurately, and only his archives provide the full story (see [3]; these are currently being prepared for publication).

The two seasons of excavations in the cave were never published, owing to political changes. The objects were placed in boxes in the Rockefeller Museum (Palestine Archaeological Museum) close to the storage area of Qumran. After the Six Day war of 1967, the British excavation teams did not further the publication, owing to their close association with the Jordanian royal family. As time passed, the exploration of the Christmas Cave did not attract scholarly attention, and even its location was forgotten due to the lack of geographic references in Allegro’s publication. In the 1980s, the stored boxes were opened by the École Biblique et Archéologique de France (EBAF) and assumed to have come from an unidentified Qumran cave. At this point they were erroneously assigned the prefix QCC, short for ‘Qumran Christmas Cave’. When the study of the material culture of Qumran was resumed in the 1990’s, under the auspices of EBAF, the Christmas Cave was understood as one of the Qumran caves (e.g. [4]). It took some effort before the location of the Christmas Cave was re-traced again in 2007 by the Israel Cave Research Center of the Hebrew University of Jerusalem [5–7]. This renewed salvage investigation included a spelleological and archaeological survey, and brought forward numerous finds, including potsherds, two additional bronze coins (and two more found in small alcoves next to the cave entrance), textile fragments, vegetal food remains, and metal objects.

In tandem, analyses conducted on the finds collected by Allegro began to indicate that the cave experienced human presence in multiple periods, including the Late Chalcolithic, Early Bronze Age I, Early Roman (first century CE), as well as during the Bar Kokhba Revolt [4, 6, 8–12]. Especially telling were the detailed studies devoted to the large collection of textiles [4, 10, 11, 13, 14]. Of the 255 catalogued textiles, 71 linen textiles were attributed to the Chalcolithic period; five pieces were assigned to the Medieval era, together with several modern textiles made of cotton; and the remaining 184 textiles were assigned to the Roman period. The textiles from the Roman period included 113 made of wool, 63 of linen, and 8 of goat-hair [11, 14]. Some of the Roman wool textiles were dyed or decorated with bands or a gamma-shaped design using a
Fig. 1 The Christmas Cave: a map showing the position of the Christmas Cave; b drawing of the morphology of the Christmas Cave.
broad spectrum of colours including red, purple, black, blue, and green [11, 14], similar to other textiles from Roman sites in the Southern Levant (see for example [15, 16]) as well as all the textiles from all the Eastern provinces in the Roman Empire [17, 18]. Belis, Shamir, and Sukenik concluded that since no wool textiles were found in the Qumran Caves [19], and given other technological differences between the two assemblages, it was unlikely that the Christmas Cave inhabitants from the Roman period had any connection with Qumran. On the other hand, the Roman-period Christmas Cave textiles exhibit many similarities to the textile assemblage from the days of the First Jewish Revolt (66–73 CE) found at Masada, as well as to textile collections derived from other cliff caves in the Judean Desert, which served as refuge caves at the end of the Second Jewish Revolt.

While all this then appeared to confirm that the cave was used for refuge in the Bar Kokhba period, the question remained about how to date objects that were not classifiable by stratigraphy or typology to a particular period. How many different periods were represented in the cave?

Clearly, the papyrus pieces would most likely come from the Bar Kokhba period [3: 68–70]. Documents from this time, including some written by Bar Kokhba himself, have been found in the caves of Wadi Murabba’at, Nahal Hever, and Nahal Se’elim ([20]:159–163; [21–25]: Table 3, 156–7). The small pieces of papyrus from the Christmas Cave appear consistent with such written materials, some of which are in Greek. To further define the epochs in which people visited, lived, or took refuge here, we present new investigations of several types of objects excavated in the Christmas Cave which will cast more light on precisely which periods saw occupation, what foodstuffs they ate, and what ceramic technology they possessed. For comparison, and further clarification, we will also qualify to what degree the Christmas Cave saw inhabitation simultaneously with the famous site of Qumran’s main habitation in Periods I and II (from ca. 85 BCE to 68 CE).

Materials

The materials investigated included textiles, reed, dung, plant stones, cereal grain, and ceramics. The 25 samples which have been investigated in the present study are listed in Table 1. In the following table, ‘QCC’ refers to the DSSF materials catalogued by EBAF in the 1980’s, and is not the original listing system made by the British team. The designations “Frumkin” and “CC soil” indicate objects from the 2007 survey. All samples analysed were surface finds.

Methods

Radiocarbon dating

Samples of a variety of materials, rope, wood, charcoal, textile, wool, were dated by radiocarbon. Accelerator Mass Spectroscopy (AMS) dating was performed. The samples underwent standard chemical pre-treatment AAA (Acid–Alkali-Acid), for cleaning and isolation of the datable fraction (see e.g. [26]). Next, the samples were combusted into CO$_2$ by an Elemental Analyzer (EA), coupled on-line with an Isotope Ratio Mass Spectrometer (IRMS). The EA is also used to purify the CO$_2$. In addition, the EA/IRMS system enables precise measurements of the $^{13}$C/$^{12}$C-ratio. Part of the CO$_2$ is reduced to graphite by reacting under excess H$_2$ gas [27]. This graphite was pressed into target holders to be placed in the ion source of the AMS (Accelerator Mass Spectrometer). The AMS measured the $^{14}$C/$^{12}$C-ratio in the graphite.

Most samples were analysed in Groningen (laboratory code GrA). Five samples were analysed in Irvine (laboratory code UCI). The Groningen AMS facility is based on a 2.5 MV Tandetron accelerator [28].

The stable isotope ratio is expressed as a δ-values, defined as the deviation (expressed in per mil) of the rare to abundant isotope ratio from that of a reference material:

$$\delta^{13} = \left( \frac{{^{13}C/^{12}C}_{sample}}{{^{13}C/^{12}C}_{reference}} - 1 \right) \times 1000\%$$

The reference material is belemnite carbonate (V-PDB) [29]. The measured $^{14}$C/$^{12}$C-ratio was converted to a conventional Radiocarbon Age. The convention comprises the use of the Oxalic Acid standard, correction for isotopic fractionation and usage of the conventional half-life [30]. The measured activity ratios are reported in $^{14}a$ (%), 100% being the standard value. The conventional $^{14}$C dates are calculated from these activity ratios and are reported in BP. The $^{14}$C dates (in BP) need to be calibrated into calendar ages. This was done using the OxCal code [31] applying the most recent high-resolution calibration curve IntCal20 [32]. The calibration yields calendar age ranges, which are reported at both the 1σ and 2σ confidence level. All dates are rounded to 5.

Thermoluminescence (TL) dating

TL-measurements were performed in Odense on a DA-12 TL-reader built by Risø National Laboratory in Denmark with the 100–300 µm granulometric fraction of sieved grains using a Single Aliquot Regeneration method adapted from Hong et al. [33], taking the average of four subsamples. No mineral separation has been
done. The calculation of the date required determining the dose received from the environment. This was assumed to originate from three sources: (1) the internal sources—from the four radioactive isotopes present in the samples, \( ^{40} \text{K}, ^{232} \text{Th}, ^{235} \text{U}, \) and \( ^{238} \text{U}; \) (2) the external source—from the same four radioactive isotopes in the surrounding soil; and (3) the cosmic flux. The radioactive isotopes from the samples and the surrounding sediment were measured using LA-ICP-MS (for Si, Th and U) and XRF (for Si and K). The cosmic flux was estimated from the geographical position of the site, its altitude above sea level, the depth of finding and the density of the overlying sediment. The calculation was performed using the “Luminescence” package on R software [34]. The procedure required input of several parameters: (1) the self-shielding was calculated using a measured average density of \( 1.8 \pm 0.3 \text{ g cm}^{-3} \), (2) the grain diameter after sieving was assumed to be \( 200 \pm 100 \mu \text{m} \), (3) the alpha efficiency was assumed to be \( 0.10 \pm 0.02 \) according to Olley et al. [35], and (5) the sediment water content was estimated to be 2 wt%. No HF etching was performed; thus, the alpha particle dose was included in the annual dose rate calculation [36]. These parameters were computed and processed through the AGE software [37] providing the dose rates and the TL-ages.

**Laser ablation inductively coupled plasma mass spectroscopy (LA-ICP-MS)**

Laser ablation (LA) was performed with a CETAC LX5-213 G2 equipped with a NdYAG laser operating at the fifth harmonic at a wavelength of 213 nm. A 25 µm circular aperture was used. The shot frequency was 20 Hz. A line scan was performed with a scan speed of 20 µm s\(^{-1}\) and was ca. 300 s long following a 10 s gas blank. The helium flow was fixed at 600 mL m\(^{-1}\). The laser operations were controlled by the DigiLaz G2 software provided by CETAC. Inductively coupled plasma mass spectrometry (ICP-MS) analyses were carried out using a Bruker Aurora M90 equipped with a frequency matching RF-generator. The basic parameters were as follows: radiofrequency power 1.30 kW; plasma argon gas flow rate 16.5 L min\(^{-1}\); auxiliary gas flow rate 1.65 L min\(^{-1}\); sheath

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**Table 1 Samples analysed in the present study**

| KLR Lab No | Gra Lab No | QUM No | Field no | Sample description | Analyses performed |
|------------|------------|--------|----------|--------------------|--------------------|
| KLR-8008   | 430        | QCC-007| Ceramic with fat | GC–MS, TS, TL, LA-ICP-MS, XRF, FT, SUS, FTIR, XRD |
| KLR-8009   | 426        | QCC-079| Ceramic with fat | GC–MS, TS, TL, LA-ICP-MS, XRF, SUS, FTIR, XRD |
| KLR-8010   | 428        | QCC-264| Ceramic, blackened | TS, TL, LA-ICP-MS, XRF, FT, SUS, FTIR, XRD |
| KLR-8011   | 425        | QCC    | Ceramic, juglet, N trench I | TS, FT, SUS |
| KLR-8012   | 437        | QCC-258| Ceramic, small juglet | GC–MS |
| KLR-8013   | Gra-50530  | 437a   | Dung      | C14 |
| KLR-8014   | Gra-50531  | 427    | III 1A    | Charcoal | C14 |
| KLR-8015   | Gra-50532  | 436    | Wood      | C14 |
| KLR-8016   | QCC test s | Ceramic | GC–MS, TS, LA-ICP-MS, XRF, SUS, FTIR, XRD |
| KLR-8852   | Gra-53282  | C1     | Cereal grain | C14 |
| KLR-8853   | Gra-53271  | C2     | Olive stone | C14 |
| KLR-8854   | Gra-53272  | C3     | Date stone | C14 |
| KLR-8855   | Gra-53273  | C4     | Reed      | C14 |
| KLR-8856   | Gra-53274  | C5     | Textile   | C14 |
| KLR-8857   | Frumkin#5  | Ceramic | TS, TL, LA-ICP-MS, XRF, FT, SUS, FTIR, XRD |
| KLR-8858   | Frumkin#6  | Ceramic | TS, TL, LA-ICP-MS, XRF, FT, SUS, FTIR, XRD |
| KLR-8859   | Frumkin#7  | Ceramic | TS, TL, LA-ICP-MS, XRF, FT, SUS, FTIR, XRD |
| KLR-8860   | Frumkin#11 | Ceramic | TS, TL, LA-ICP-MS, XRF, FT, SUS, FTIR, XRD |
| KLR-8861   | Frumkin#12 | Ceramic | TS, TL, LA-ICP-MS, XRF, FT, SUS, FTIR, XRD |
| KLR-8862   | Frumkin#15 | Ceramic, lamp with soot | TS, TL, LA-ICP-MS, XRF, FT, SUS |
| KLR-8863   | Frumkin#16 | Ceramic | TS, TL, LA-ICP-MS, XRF, FT, SUS |
| KLR-12527  | CC1 soil   | Soil from surface | LA-ICP-MS, XRF |
| KLR-12529  | CC3 soil   | Soil from surface | LA-ICP-MS, XRF |
| KLR-12530  | CC4 soil   | Soil from surface | LA-ICP-MS, XRF |
| KLR-12531  | CC5 soil   | Soil from surface | LA-ICP-MS, XRF |

“QUM” are EBAF and J. Gunneweg numbers. GC–MS Gas Chromatography Mass Spectrometry, C14 Radiocarbon dating, TS Thin Section – petrographic description, FT Firing Temperature determination, TL Thermoluminescence, XRF X-Ray fluorescence, LA-ICP-MS Laser Ablation Inductively Coupled Plasma Mass Spectrometry, SUS magnetic susceptibility, FTIR Fourier Transform Infrared Spectroscopy, XRD X-ray diffraction
gas flow rate 0.18 L min⁻¹. The following isotopes were measured all without skimmer gas: ²⁹Si, ²³²Th, and ²³⁸U. No interference corrections were applied to the selected isotopes. The analysis mode used was peak hopping with 3 points per peak, and the dwell time was 10 ms on ²⁹Si and 100 ms on ²³²Th and ²³⁸U. The quantification was performed with a method similar to that of Golitko and Terrell [38]. An in-house ceramic standard was run before and after batches of three samples in order to monitor the stability of the beam and to act as a standard. The concentration of U and Th were calculated by comparison of the U/Si and Th/Si experimental ratios to the U/Si and Th/Si standard material ratios. A relative error of ca. 10% is estimated from these measurements mostly due to mineral heterogeneity of the samples. For the analyses of other trace elements than U and Th, the same set was used, and the following isotopes were analysed for: ⁵⁵Mn, ⁵⁷Fe, ⁶⁰Co, ⁶⁵Cu, ⁶⁵Zn, ⁷⁵As, ⁸⁵Rb, ⁸⁸Sr, ¹⁰⁷Ag, ¹¹¹Cd, ¹²¹Sb, ²⁰⁵Tl, and ²⁰⁸Pb. Here the analysis mode used was also peak hopping with 3 points per peak. The data were quantified by direct ratios to the count rates of the international standard material NIST612, which was analysed as every fourth sample.

Petrographic descriptions
Thin sections of 30 μm thickness were manufactured from the ceramic samples in the Catamarca laboratory. They were polished with silicon carbide 80, 180, 400 and 600, photographed and described petrographically. Crossed polarized light (XPL) and plane polarized light (PPL) observations were done in a Leitz Ortholux polarizing microscope at 40X-100X. JMicrovision® software was used to make all the measurements and the point-counting analysis in each thin-section. A total of 300 points were recorded for each thin-section.

Magnetic susceptibility and TL-sensitivity measurements
The method of characterising clay sediments has proven effective in studies of provenancing archaeological ceramics, pot shards, and red brick. Magnetic susceptibility was measured using ca. 1 g samples with a KLY-2 susceptibility-metre capable of reaching ca. 1 × 10⁻⁶ SI-units. The thermoluminescence (TL) sensitivity was measured on four 10 mg aliquots of the crushed and sieved grain size fraction 100–300 μm. For the TL-measurements a TL/OSL-system TL-DA-12 manufactured at Riso National Laboratory, Denmark was used. The palaeosignals were erased by heating the samples to 400 °C. Subsequently the samples were irradiated for 60 s under a 0.8 GBq ⁹⁰Sr-source. The samples were then annealed for 30 s at 200 °C, and the TL-signal measured from 202 to 235 °C and integrated in order to yield the TL-sensitivity. Drift in photodetector high tension, optical transmission and other system parameters were monitored by measuring four aliquots of a standard sample daily. The method and its application are described in [39–47].

Firing temperature determination
The maximum firing temperature was determined at University of Southern Denmark in the Odense laboratory by stepwise reheating as described in Rasmussen et al. [48]. This method was chosen over other methods such as e.g. the thermal expansion measurement method (see e.g. Tite [49]) as it is generally producing more reliable results. The samples were heated at 20 degrees steps starting from 200 °C until 1060 °C. The heating time was 24 h for the lowest temperatures, gradually decreased to 1 h with raising temperature. Upon cooling after each heating step, the samples were weighed and measured in quadruple for magnetic susceptibility on a KLY-2 Kappebridge manufactured by Geofyzika, Brno, with a detection limit of 10⁻⁶ SI-units. The samples were all a factor of ca. 1000 times higher than the detection limit. A sample with known susceptibility was measured daily for absolute calibration. The empty sample holder was measured before and after each session and the average background susceptibility was subtracted from the actual measurements. The uncertainty was calculated from the four repeated measurements, between which the sample was turned in the sample holder. The reported uncertainties therefore to a large degree reflects the anisotropy of the sample. The firing temperature was read off a graph showing the squared first derivative of the susceptibility as a function of the reheating temperature [48].

Micro-X-ray fluorescence (µ-XRF)
The embedded and polished samples were subjected to analysis by µ-XRF for the quantification of K and Si. An ARTAX-800 µ-XRF situated at University of Southern Denmark and manufactured by Bruker-Nano was used and operated with a high tension of 50 kV and a current 600 μA. Absolute calibration of the concentrations has been performed by the DCCR-method (Direct Calibration from Count Rates) provided by the Bruker software using the standard reference material NIST-2711.

Fourier Transform Infrared spectroscopy (FTIR)
The samples were analysed at University of Southern Denmark using a FTIR instrument (Agilent Technology, Cary 630) with a diamond crystal ATR accessory. Spectra were collected from 32 co-added scans in the spectral range 4000–650 cm⁻¹ with a resolution of 8 cm⁻¹. The background was measured with 32 scans. The instrument was controlled through MicroLab software and data were processed with Spectragraphy software (v.1.2.14). The second derivative of the spectra were calculated. This is
a widely used and useful method in FTIR studies for the identification of peaks barely visible in the raw spectra [51–56].

X-ray diffraction (XRD)
The XRD analyses were performed University of Southern Denmark using a PANalytical X’Pert PRO MPD system (PW3050/60) diffractometer with Cu Kα radiation as the source (λ = 1.54 Å) and a PIXcel3D detector. The X-ray generator was set to an acceleration voltage of 45 kV and a filament emission to 40 mA. All measurements were performed in continuous mode with a 2Θ angle step size of 0.026° and a scan speed of 0.022° s⁻¹ in a range from 5° to 90° 2Θ. The qualitative analysis was performed using Highscore Plus software linked to the ICDD PDF-2 database.

Gas Chromatography Mass Spectroscopy (GC–MS)
GC–MS analysis was performed at the University of Pisa after extraction of ground fragments of food crusts adhered to the internal surface of four ceramic vessels (KLR8008, KLR8009, KLR8012, and KLR8016) with CH₂Cl₂/MeOH. The extracts were dried and submitted to alkaline hydrolysis following a procedure described in the literature [57]). Particularly, the neutral (unsaponifiable) and the acidic fractions were obtained by extraction with hexane and, after acidification, with diethyl ether. Each fraction was derivatised with a silylating agent, N,O-Bis(trimethylsilyl)trifluoroacetamide (BSTFA), and separately analysed by GC–MS.

Results
Radiocarbon dating
The radiocarbon dates acquired in this study are listed in Table 2. The columns are the laboratory identification numbers, the δ¹³C values, the ¹⁴C activities and measurement uncertainties, the Carbon content (C%) of the analysed sample materials, the δ¹³C values, the ¹⁴C ages in BP and the calibrated age ranges in CE/BCE, both 1σ and 2σ. The ¹⁴C dates and calibrated dates are rounded to 5 years. Multiple calibrated ranges are taken together.

**TL-datings**
The Si, K, U, and Th concentrations and the TL-dates are listed in Table 3. Both 1σ and 2σ uncertainty ranges are listed, and both intervals are used in the discussion below. Results for 4 soil samples later procured by Amos Frumkin are listed below the samples in Table 3 (one additional soil sample failed to yield usable results for Si and was therefore discarded). Below this is listed the average soil composition together with a 1σ uncertainty calculated as 10% of the concentration values. The latter is a best estimate, based mainly on an assessment of the inhomogeneity of the soil samples. The radiation contribution from the soil is applied as half the calculated average soil composition, because the ceramic samples were assumed to have been residing on the floor of the cave, not dug down into the chalkstone.

Petrographic and mineralogic analysis of the ceramic samples
The results of the XRD and FTIR analysis (Table 4) confirmed the mineralogical observations and brought some complementary information. FTIR second-derivative spectra and XRD diffractograms, as well as the results of the firing temperature determinations are available in Additional file 1.

Secondary calcite was identified in some sherds, and here the combination of petrography and the results of the FTIR show the presence of primary calcite (KLR-8810, KLR-8859, KLR-8861, KLR-8862, and KLR-8863). In the felsic minerals, XRD and FTIR revealed the presence of quartz and several potassium and plagioclase series feldspars. The XRD patterns point to the presence

**Table 2 Results of the radiocarbon dating**

| KLR lab No | GrA lab No | δ¹³C (%o) | Sample material | C% | ¹⁴a (%) | σ ¹⁴a | Age (BP) | Calibrated age range (1σ) | Calibrated age range (2σ) |
|------------|------------|-----------|-----------------|----|---------|-------|----------|--------------------------|--------------------------|
| 8853       | 53271      | -20.94    | Olive stone     | 48.8| 92.21   | 0.38  | 650±35   | CE1290–1390               | CE1280–1400               |
| 8854       | 53272      | -23.41    | Date stone      | 46.0| 79.71   | 0.34  | 1820±35  | CE170–325                 | CE125–330                 |
| 8013       | 50530      | -24.45    | Macro remains   | 35.1| 79.42   | 0.38  | 1850±40  | CE130–240                 | CE80–325                 |
| 8856       | 53274      | -25.11    | Textile         | 39.9| 78.82   | 0.33  | 1910±35  | CE80–205                  | CE25–220                 |
| 8855       | 53273      | -23.99    | Reed            | 43.4| 78.44   | 0.33  | 1950±35  | CE25–125                  | BCE40–CE205              |
| 8015       | 50532      | -24.78    | Wood            | 48.4| 78.30   | 0.37  | 1965±40  | BCE5–CE120                | BCE50–CE205              |
| 8014       | 50531      | -22.11    | Charcoal        | 75.6| 52.57   | 0.28  | 5170±40  | BCE405–3950               | BCE4160–3805             |
| 8852       | 53282      | -22.63    | Cereal grain    | 35.0| 51.86   | 0.25  | 5275±40  | BCE4230–3995              | BCE4240–3985             |

The columns show KLR and GrA laboratory code numbers, δ¹³C in ‰, sample material, ¹⁴C activity ratio corrected for isotope fractionation in %, its standard deviation, ¹⁴C age in BP including its standard deviation, and calibrated age ranges in BCE at 1σ and 2σ confidence intervals. The ¹⁴C dates are calibrated with the latest calibration curve IntCal20 [32]. See text for details.
of a moderate amount of potassium feldspars, including three sherds with sanidine (KLR-8859, KLR-8860, and KLR-8861), while the FTIR data give more information about the potassium feldspars, showing the presence in most of the sherds of orthoclase and sanidine, and, in some sherds, microcline. Concerning the plagioclase series, the presence of two specific feldspars was observed: anorthite and albite, anorthite being the most common plagioclase mineral. Various amounts of gehlenite, wollastonite, and diopside have been detected in all the sherds. These neo-formation minerals have been detected by both instrumental techniques, but the FTIR second-derivative method appears to be more sensitive than the XRD. Iron oxides such as hematite were identified in at least four of the sherds (KLR-8008, KLR-8009, KLR-8858, and KLR-8861).

The absence of clay minerals (such as illite, smectite, and kaolinite) in the XRD and FTIR analyses indicated that the pottery was fired above 500–550 °C. As most of the pottery analysed is Ca-based ceramics, the presence of several neo-formation mineral phases in the

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### Table 3  Results of the TL-dating of the ceramic samples

| Lab No  | Code       | Field no | Si wt% | K wt% | U µg/g | Th µg/g | Date BCE/CE | 1σ BCE/CE | 1σ interval BCE/CE | 2σ interval BCE/CE |
|---------|------------|----------|--------|-------|--------|---------|-------------|-----------|-------------------|-------------------|
| KLR-8008 | QCC-430    | QCC-007  | 25.1   | 1.14  | 2.32   | 4.92    | 450 CE     | 100       | 350–550 CE        | 250–650 CE        |
| KLR-8009 | QCC-426    | QCC-079  | 30.4   | 1.71  | 1.84   | 6.33    | 334 CE     | 106       | 228–440 CE        | 122–546 CE        |
| KLR-8010 | QCC-428    | QCC-264  | 8.63   | 1.91  | 1.53   | 1.86    | 3208 BCE   | 354       | 3563–2854 BCE     | 3916–2500 BCE     |
| KLR-8016 | ?          | ?        | 24.5   | 1.58  | 1.43   | 5.35    | 413 BCE   | 155       | 568–258 BCE       | 723–103 BCE       |
| KLR-8857 | Frumkin#5  | Frumkin#5 | 29.3   | 1.51  | 2.15   | 8.60    | 59 BCE    | 127       | 186 BCE – 70 CE   | 313 BCE–195 CE    |
| KLR-8858 | Frumkin#6  | Frumkin#6 | 31.8   | 1.58  | 2.99   | 10.9    | 240 CE    | 110       | 130–350 CE        | 20–460 CE         |
| KLR-8859 | Frumkin#7  | Frumkin#7 | 19.0   | 3.55  | 1.87   | 4.05    | 476 CE    | 107       | 369–583 CE        | 262–690 CE        |
| KLR-8860 | Frumkin#11 | Frumkin#11| 22.1   | 4.17  | 2.37   | 5.85    | 698 BCE   | 206       | 904–492 BCE       | 1110–286 BCE      |
| KLR-8861 | Frumkin#12 | Frumkin#12| 17.3   | 4.14  | 2.46   | 4.02    | 92 BCE    | 142       | 234 BCE – 50 CE   | 376 BCE–192 CE    |
| KLR-8862 | Frumkin#15 | Frumkin#15| 10.6   | 2.67  | 3.72   | 3.98    | 145 BCE   | 151       | 296 BCE – 6 CE    | 447 BCE–157 CE    |
| KLR-8863 | Frumkin#16 | Frumkin#16| 22.2   | 1.15  | 2.24   | 6.68    | 272 BCE   | 180       | 456–92 BCE        | 632 BCE–88 CE     |
| KLR-12527| CC1 soil   |          | 3.99   | 2.87  | 1.085  | 0.495   |            |           |                   |                   |
| KLR-12529| CC3 soil   |          | 0.165  | 1.93  | 0.098  | 0.021   |            |           |                   |                   |
| KLR-12530| CC4 soil   |          | 0.325  | 1.49  | 0.476  | 0.049   |            |           |                   |                   |
| KLR-12531| CCS soil   |          | 0.230  | 3.38  | 0.268  | 0.046   |            |           |                   |                   |
| Average  | soil       |          | 1.18   | 2.42  | 0.482  | 0.153   |            |           |                   |                   |
| 1σ (10%) |           |          | 0.12   | 0.242 | 0.048  | 0.015   |            |           |                   |                   |

A sample called CC2-soil was available but did not contain quantifiable amounts of Si by XRF and were consequently left out.

### Table 4  Mineral phases identified by XRD and FTIR analyses

| Samples   | XRD | FT-IR |
|-----------|-----|-------|
| Samples   | Qz  | Cal  | K-fs | An | Al  | Hem | Grt | Gh | Di | Wo | Qz  | Cal | Cal 2nd | An | Al  | Or | Mc | Sn | Gh | Di | Wo |
| KLR-8008  | xxxx | –    | xx   | X  | –   | –   | –   | –  | –  | –  | xxxx | x    | x    | x    | x  | x  | –  | x  | x  | x  | x  | x  |
| KLR-8009  | xxxx | –    | xx   | xx | xx  | –   | –   | –  | –  | –  | xxxx | xx   | x    | x    | x  | x  | –  | x  | x  | x  | x  | x  |
| KLR-8010  | xxxx | xx   | xx   | XX | x   | –   | –   | –  | x  | –  | xx   | xxxx | –    | xx   | x   | x  | x  | x  | x  | x  | x  | x  |
| KLR-8016  | xxxx | xx   | xx   | xx | –   | xx  | –   | x  | –  | x  | xxxx | x    | x    | x    | x  | x  | –  | x  | x  | x  | x  | x  |
| KLR-8857  | xxxx | xx   | xx   | xx | x   | –   | x   | –  | –  | –  | xxxx | xx   | xx   | xx   | xx | x  | –  | x  | x  | x  | x  | x  |
| KLR-8858  | xxxx | x    | xx   | xx | xx  | –   | –   | –  | –  | –  | xxxx | xx   | x    | x    | x  | x  | –  | x  | x  | x  | x  | x  |
| KLR-8859  | xxxx | xx   | xx   | x  | –   | xx  | –   | x  | –  | x  | xxxx | xx   | xx   | xx   | x  | x  | –  | x  | x  | x  | x  | x  |
| KLR-8860  | xxxx | xx   | xx   | –  | x   | xxx  | –   | x  | –  | x  | xxxx | xx   | x    | x    | x  | x  | –  | x  | x  | x  | x  | x  |
| KLR-8861  | xxxx | xxxx | xx   | –  | –   | –   | –   | x  | –  | x  | xxxx | xx   | xx   | x    | x  | x  | –  | x  | x  | x  | x  | x  |
fabrics was expected. The presence of wollastonite, a neo-formed mineral at the carbonate-quartz interface, indicates firing temperatures as high as 850–900°C or above (sherds KLR-8009, KLR-8010, KLR-8016, KLR-8858, and KLR-8860). The presence of diopside and gehlenite indicate temperatures mostly above 750–800 °C, and gehlenite mostly is a good indicator of temperatures above 800 °C, although it starts to nucleate at 650°C. The presence of microcline together with orthoclase in one sherd suggest firing temperatures not above 750 °C (KLR-8857). The absence of relevant peaks of calcite in the XRD patterns in some of the sherds and the presence of neo-formation mineral phases suggest firing temperatures in the range of 900–950°C (KLR-8860 and KLR-8858) (see Additional file 1).

The ceramic samples are mostly characterized by presence of very fine or fine quartz-grained fabrics. The matrices are generally isotropic (i.e., not optically active) and most of them are formed by aggregates and very fine-grained quartz (Fig. 2 and Additional file 1). Only one sherd (KLR-8859) presented an anisotropic matrix formed by mica (biotite) in addition to very fine-grained quartz. Most of the sherds have an average of 14.4% inclusions, 78.8% matrix and 6.8% voids. The common primary minerals identified are quartz (and polycrystalline quartz), plagioclase feldspar, and calcite (mostly secondary). Accessory minerals identified are pyroxene, amphibole (brown hornblende), and some opaque minerals. Additionally, some rock fragments (mostly igneous rock fragments, granite) were identified at trace level. One exception was sample KLR-8810, which features abundant large granite fragments. Sample KLR-8858 also presents fragments of limestone (sedimentary rock) at trace levels. Generally, mineral inclusions and voids are evenly distributed. According to the mineral assemblages recorded by optical microscopy on thin-sections, the investigated sherds can be classified into 7 petrographical fabrics:

- Fabric 1: KLR-8008 (rounded quartz, probably coming from fluvial deposits).
- Fabric 2: KLR-8010 (a very distinctive sherd, with mineralogy characterized by very fine/fine quartz inclusions + large igneous rock fragments of granite type).
- Fabric 3: KLR-8858 (high percentage of argillaceous inclusions + very fine quartz).
- Fabric 4: KLR-8857, and KLR-8009 (very fine quartz + plagioclase with a good distribution and a selection of temper).
- Fabric 5: KLR-8860 (very fine low sphericity quartz + plagioclase; presence of grog).
- Fabric 6: KLR-8861 and KLR-8863 (high percentage of primary and secondary calcite). KLR-8859 (very fine quartz + argillaceous inclusions + grog).
- Fabric 7: KLR-8862 (high percentage of primary and secondary calcite + argillaceous inclusions).

Trace elements and magnetic susceptibility versus TL-sensitivity

The trace elements compositions determined by LA-ICP-MS, the magnetic susceptibility, and the TL-sensitivity results are all listed in Table 5.

In the magnetic susceptibility versus TL-sensitivity cross plot (Fig. 3), the ceramics studied here are compared to 12 ceramic sherds from Qumran [42]. More than half of the samples from Qumran are gathered in the top-left corner of the plot with high to moderate magnetic susceptibility and low TL-sensitivity. The sherds from the Christmas Cave are more widely spread on the plot. Most of them seem to exhibit a higher TL-sensitivity than the Qumran ceramics indicating a difference in the raw materials used for the fabrication of the ceramics in Qumran and in those found in the Christmas Cave.

To achieve a better distinction between the samples, the trace elements data were processed using multivariate technique. Because of the of the low number of sample and the higher number of variables (p > n), the use of Sparse PCA (SPCA) is more suited than regular PCA and eases the interpretation of the cross plot. The cross plot (Fig. 4) was computed with the R package “SparseBiplots” [58] using the LASSO method [59] with λ = 0.22. The results show that the 52% of the variation is explained by PC1 (principal component 1) and 22% is explained by PC2. There are strong contributions of Co, Fe, Sb, and Ni aligned with PC1, whereas Zn, Ag, Pb, and Mn are aligned with the PC2. Arsenic, Cd, Ti, and Sr are contributing to both PC1 and PC2. Six samples are located on the left side of the biplot with negative coordinates on PC1 (KLR-8857, KLR-8859, KLR-8860, KLR-8861, KLR-8862, and KLR-8863). The three samples KLR-8008, KLR-8010, and KLR-8858 are separate from these with positive PC1 values. The comparison between the chemical results and the petrological fabrics is consistent even if several petrological fabrics are gathered in the same area to the left in the SPCA cross plot.

Analyses of organic residues

The results of the GC–MS analyses of organic material associated with the pottery fragments are shown in Fig. 5. Peak assignments are listed in Tables 6 and 7. The results highlight a complex pattern of mono- and di-carboxylic fatty acids (Fig. 5a and c) in all the four
Fig. 2 Microphotographs of six sherds in crossed polarized light (XPL) showing the main mineral and rock fragment inclusions for six of the seven defined petrographic groups (40X)
Table 5  Results of the trace element concentrations determined by LA-ICP-MS

| Mineral | Mn  | Fe  | Co  | Ni  | Cu  | Zn  | As  | Rb  | Sr  | Ag  | Cd  | Sb  | Tl  | Pb  | SUS | 1σ  | TL-235 | 1σ  |
|---------|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|--------|------|
| KLR-8008 | 1146 | 4.35 | 27.8 | 144.9 | 46.4 | 245.8 | 13.4 | 49.3 | 252 | 2.09 | 0.528 | 1.97 | 0.623 | 353 | 10.0 | 0.01 | 1.30 | 0.45 |
| KLR-8010 | 678  | 6.19 | 22.3 | 124  | 99.8 | nd  | 195 | 53.9 | 673 | 0.259 | 1.71  | 1.24 | 3.25  | 6.05 | 7.34 | 0.07 | 0.550 | 0.13 |
| KLR-8857 | 787  | 4.44 | 19.4 | 59.3 | 29.5 | 90.9 | 112 | 51.0 | 109 | 0.367 | 0.116 | 0.401 | 0.304 | 12.0 | 25.8 | 0.02 | 1.13 | 1.40 |
| KLR-8858 | 1751 | 6.37 | 38.0 | 87.0 | 44.0 | 124  | 605 | 81.2 | 148 | 0.533 | 0.165 | 0.912 | 0.558 | 19.7 | 26.1 | 0.13 | 6.28 | 0.74 |
| KLR-8859 | 562  | 4.77 | 19.2 | 53.4 | 29.7 | 101  | 128 | 96.2 | 373 | 0.307 | 0.446 | 0.600 | 0.686 | 17.3 | 7.13 | 0.05 | 0.740 | 0.31 |
| KLR-8860 | 486  | 4.49 | 19.1 | 43.1 | 23.5 | 112  | 15.3 | 93.6 | 194 | 0.194 | 0.220 | 0.588 | 0.509 | 15.0 | 32.5 | 0.86 | 0.950 | 0.40 |
| KLR-8861 | 197  | 2.32 | 9.33 | 20.6 | 14.7 | 64.3 | 6.97 | 62.6 | 167 | 0.270 | 0.107 | 0.298 | 0.329 | 7.86 | 3.69 | 0.02 | 0.330 | 0.41 |
| KLR-8862 | 465  | 2.98 | 13.8 | 46.7 | 31.8 | 168  | 7.59 | 71.5 | 270 | 0.271 | 0.097 | 0.442 | 0.0333 | 13.2 | 17.1 | 0.01 | 2.03 | 0.74 |
| KLR-8863 | 451  | 3.69 | 16.2 | 33.3 | 21.2 | 96.5 | 13.3 | 96.6 | 254 | 0.350 | 0.151 | 0.499 | 0.557 | 17.0 | 14.2 | 0.02 | 0.520 | 0.15 |

*Iron is given in wt%, the rest of the trace elements in µg g⁻¹, the relative standard deviation is ca. 15%. Also listed is magnetic susceptibility (SUS), which is measured in 10⁻³ SI units, and the TL-sensitivity (TL-235) measured in arbitrary units as described in [41].*
samples analysed by GC–MS: glycerolipids are the main component of the vessel lipids. The use of calibration curves for the linear fatty acids allowed us to perform an estimation of the lipid content, which was above 50 μg for samples KLR-8009 and KLR-8012 and above 120 μg for sample KLR-8008 and KLR-8016 (the quantitation limit, based on procedural blanks, is ca. 3 μg). In particular, the acidic fractions of the saponified extracts were showed to contain even and odd chain saturated linear monocarboxylic fatty acids, with 8–24 carbon atoms in the acyl chain, with hexadecanoic acid (palmitic acid, peak #30) and octadecanoic acid (stearic acid, peak #41) as the most abundant. The fatty acid profiles also featured variable amounts of the unsaturated cis and trans octadecenoic acid isomers (oleic and elaidic acids, peaks #39 and 40), particularly abundant in samples KLR-8012 and KLR-8016.

The neutral fraction of sample KLR-8008 features a relevant amount of long chain ketones (Fig. 5b, peaks #26, 29–39 in the neutral fraction). Finally, some plasticizers (i.e., phthalates) were also detected.

Table 8 summarizes the identified compounds and materials for all the four samples analysed.

**Discussion**

**Dating and occupational phases**

The collective set of radiocarbon dates and TL-dates from the present study and previous studies shows that the Christmas Cave has been temporarily used in several episodes during the last 6500 years (Figs. 6 and 7; Table 9).

A single early fifth millennium BCE radiocarbon date (GrA-24265) was made on wood. However, it is undetermined if it should be taken to represent an early
Chalcolithic use of the cave, or whether the wood was re-used at a later time. In any case, no clear archaeological evidence for an early fifth millennium phase was attested in the material record otherwise from the cave, or elsewhere in the Judean Desert caves to date ([60]: 172), which strongly suggests that old wood was used by the occupants at an unknown time thereafter.

The cave was certainly occupied in the Late Chalcolithic, during the last quarter of the fifth millennium BCE. This is indicated by three radiocarbon dates from a cereal grain and two pieces of charcoal (KLR-8852/GrA-53282, KLR-5464/GrA-24261, KLR-8014/GrA-50531, see Tables 2 and 9). This is in accordance with the archaeological evidence in the form of a small yet typical Late Chalcolithic pottery assemblage ([60]: 381–384, Fig. 33.4). It is likely that some of the 71 linen fragments from the DSSF excavation classified by Shamir and Suknik as “Chalcolithic” [11, 14] also belong to this phase according to the splicing technique that were observed in those textiles ([61]: 34).

A second late prehistoric habitation phase is attested by three radiocarbon dates and one TL-date from pieces of textile and a ceramic fragment (samples KLR-10430/GrA-65501, UCI-79817, UCI-79815, and KLR-8010). The three radiocarbon dates are clustered around 3630–3530 BCE, a time span currently associated with the Early Bronze Age 1A in the conventional South Levantine chronology (e.g., [62, 63]: 26–43). Activity during this time span is virtually unknown from the Judean Desert [64], with only one other radiocarbon date thus far obtained from the region (the latest date from the “Cave of the Warrior” in Wadi al-Makkukh; [65]: sample RT-1943). There has not yet been found independent archaeologically dated materials from the Christmas Cave from Early Bronze Age 1A, although it is possible that some of the textiles previously classified as “Chalcolithic” may belong to this phase.

Approximately two thirds of the radiocarbon and TL dates fall between 45 BCE and 320 CE (2σ). A single date from an olive stone (KLR-8853/GrA53271) is medieval, 1280–1395 CE, which falls in the early Mamluk period and testifies to a visitor from this time. A single TL-date (KLR-8860, Table 3) falls in the late Iron Age, or First Temple Period, around 700 BCE.
The calibrated date intervals (1σ and 2σ) between 500 BCE and 500 CE are depicted in Fig. 7, which also shows three rectangles indicating the main habitation period for the Qumran settlement (ca. 100 BCE to 68 CE; ignoring the Roman occupation after 68 CE), the First Jewish Revolt (66–73 CE), and the Second Jewish Revolt or the reign of Bar Kokhba (132–136 CE).

There are six radiocarbon dates with 1σ ranges in-between the two revolts, i.e.

- from 73 to 132 CE. However, if the 2σ ranges are considered, multiple radiometric and TL dates overlap with both the Qumran Main Habitation period and the Second Jewish Revolt. So, the temporal resolution in the dates available is not sufficient to resolve with certainty if the Christmas Cave was visited during both or only one of the revolts, or indeed in the time between the revolts.

Pottery from the Hasmonean era (2nd to first century BCE) may be linked with the nearby harbour of Khirbet Mazin, at the mouth of the Wadi Qidron, which was established in the time of Alexander Jannaeus (ca. 85 BCE) and continued to be in use through the Early Roman period (see [66]: 77–78; [67]: 18–29; [68]). The Hasmoneans also re-used old Iron Age sites in the nearby Buqe’ah (see [69]: 145). North of the Lower Qidron valley there was a track leading from the Dead Sea through the wilderness to Bethlehem and Jerusalem, thus travellers in peacetime could also have made use of the caves temporarily.

The TL-dates of ceramic fragments from the 4th to sixth centuries are consistent with what we know of

| Peak # | Retention time (min) | Peak assignment |
|--------|---------------------|-----------------|
| 1      | 8.0                 | Benzoic acid    |
| 2      | 8.3                 | Octanoic acid   |
| 3      | 8.8                 | Tridecane       |
| 4      | 9.6                 | Nonanoic acid   |
| 5      | 10.2                | Tetradecane     |
| 6      | 10.9                | Decanoic acid   |
| 7      | 11.4                | Heptanedioc acid, 1-methyl ester |
| 8      | 11.7                | Hexanedioc acid |
| 9      | 12.1                | Undecanoic acid |
| 10     | 12.7                | Hexadecane (IS1) |
| 11     | 12.6                | Octanedioc acid, 1-methyl ester |
| 12     | 12.8                | Heptanedioc acid |
| 13     | 13.2                | Dodecanedioc acid |
| 14     | 13.7                | Octanedioc acid, 1-methyl ester |
| 15     | 13.7                | Octanedioc acid |
| *      | 13.9                | Phthalic acid   |
| 16     | 14.3                | Tridecanedioic acid (IS2) |
| *      | 15.1                | Terephthalic acid |
| 17     | 14.9                | Nonanedioc acid |
| 18     | 15.0                | ω-hydroxy-decanoic acid |
| 19     | 15.1                | Tetradecanoic acid (br) |
| 20     | 15.6                | Tetradecanoic acid |
| *      | 15.9                | Phthalate (contaminant) |
| 21     | 16.3                | Dodecanedioc acid |
| 22     | 16.6                | Pentadecanoic acid (br) |
| 23     | 16.8                | Pentadecanoic acid (br) |
| 24     | 17.4                | Hexadecanoic acid, methyl ester |
| 25     | 17.3                | Pentadecanoic acid |
| 26     | 18.0                | Undecanedioc acid |
| 27     | 18.3                | Hexanedioc acid (br) |
| 28     | 18.8                | (Z)-9-hexadecenoic acid |
| 29     | 19.0                | (E)-9-hexadecenoic acid |
| 30     | 18.9                | Hexanedioc acid |
| 31     | 19.5                | Dodecanedioc acid |
| 32     | 19.8                | Heptadecanoic acid (br) |
| 33     | 19.9                | Heptadecanoic acid (br) |
| 34     | 20.4                | Octadecanoic acid, methyl ester |
| 35     | 20.3                | Heptadecanoic acid |
| 36     | 20.7                | Tridecanedioic acid |
| 37     | 21.0                | Octadecanoic acid (br) |
| 38     | 21.4                | 9,12-octadecadienoic acid |
| 39     | 21.1                | (Z)-9-octadecenoic acid |
| 40     | 21.7                | (E)-9-octadecenoic acid |
| 41     | 21.5                | Octadecanoic acid |
| 42     | 21.9                | Tetradecanedioc acid |
| 43     | 22.2                | Nonadecanoic acid (br) |
| 44     | 22.5                | Hydroxy-carboxylic FA-TMS |
| 45     | 22.5                | Nonadecanoic acid |

All organic acids and alcohols are detected as their TMS esters and ethers, respectively.

The calibrated date intervals (1σ and 2σ) between 500 BCE and 500 CE are depicted in Fig. 7, which also shows three rectangles indicating the main habitation period for the Qumran settlement (ca. 100 BCE to 68 CE; ignoring the Roman occupation after 68 CE), the First Jewish Revolt (66–73 CE), and the Second Jewish Revolt or the reign of Bar Kokhba (132–136 CE).

There are six radiocarbon dates with 1σ ranges in-between the two revolts, i.e., from 73 to 132 CE. However, if the 2σ ranges are considered, multiple radiometric and TL dates overlap with both the Qumran Main Habitation period and the Second Jewish Revolt. So, the temporal resolution in the dates available is not sufficient to resolve with certainty if the Christmas Cave was visited during both or only one of the revolts, or indeed in the time between the revolts.

Pottery from the Hasmonean era (2nd to first century BCE) may be linked with the nearby harbour of Khirbet Mazin, at the mouth of the Wadi Qidron, which was established in the time of Alexander Jannaeus (ca. 85 BCE) and continued to be in use through the Early Roman period (see [66]: 77–78; [67]: 18–29; [68]). The Hasmoneans also re-used old Iron Age sites in the nearby Buqe’ah (see [69]: 145). North of the Lower Qidron valley there was a track leading from the Dead Sea through the wilderness to Bethlehem and Jerusalem, thus travellers in peacetime could also have made use of the caves temporarily.

The TL-dates of ceramic fragments from the 4th to sixth centuries are consistent with what we know of
Byzantine hermits who lived in desert caves during this time (KLR-8008, 8009, 8858, and 8859). The Byzantine sample KLR-8008 is special, because it is a sample containing thermally altered fats, it has a specific fabric type (1), and it is the sample with the highest percentage of voids and the highest percentage of quartz. Also, it plots distinctly different from the Qumran samples in the provenancing plot (Fig. 4).

Apart from the one Bronze Age TL-date, the other TL-dates are rather uniformly spread out from 700 BCE to 475 CE. Three TL-dates are overlapping the main habitation period of Qumran (ca. 100 BCE to 68 CE) with the 1σ interval (KLR-8857, 8861, and 8862). One TL-date overlaps the Second Revolt period in the 1σ interval (KLR-8858).

### The ceramics

It is evident from the archaeological evidence of the site that none of the ceramics found in the Christmas Cave were produced close by. The ceramic samples exhibit traits that can trace their place of origin, although they cannot be provenanced to any specific location, at least not from the present study. The analyses of the ceramics show some differences in the resources used for their manufacture, which again might indicate differences in location of origin of the people seeking shelter or refuge in the cave.

From a technological point of view, the pottery found in the Christmas Cave was made using different recipes by ancient potters. Two main *momentums* can be defined. The oldest sherds (KLR-8010, 8860, 8861, 8862, and 8863) are characterized by ceramic pastes with poorly sorted minerals and very low grain selection; low firing temperatures are very characteristic in these early shards as indicated by the presence of primary calcite. Surprisingly, the presence of grog which was found in only one shard (KLR-8860) indicates something more than routine recipes for the making of the pottery. This practice should be fully investigated further in the future. One of the older sherds, KLR-8010, was found to have a relative high firing temperature. The younger sherds (KLR-8857, 8858, 8008, and 8859) are characterized by well sorted mineral grains and an excellent grain selection used to elaborate the ceramic pastes. Most of them are very fine quartz tempered with low percentages of voids, indicating a very elaborated recipe to make the vessels.

Summarizing the inorganic measurements of the ceramic material it appears that there are probably four different sources for the raw materials of the ceramics. Ceramics were fired at temperatures well above 600°C, reaching 800–850°C for some of them. Several neo-formation mineral phases were identified by XRD and FTIR. The results of the LA-ICP-MS and the mineralogy analysis seen through the SPCA analysis (Fig. 4) suggest that the samples KLR-8008, KLR-8010 and KLR-8858 are possibly from three different locations. The six samples KLR-8857, KLR-8859, KLR-8860, KLR-8861, KLR-8862, and KLR-8863 group together which might indicate that they share a common origin. The three samples KLR-8861, 8862 and 8857 are probably contemporary with Qumran.

### Table 7

| Peak # | Retention time (min) | Peak assignment               |
|--------|----------------------|--------------------------------|
| 1      | 8.8                  | Tridecane                      |
| 2      | 10.2                 | Tetradecane                    |
| 3      | 12.2                 | Dodecanol                      |
| 4      | 12.7                 | Hexadecane (IS1)               |
| 5      | 13.4                 | 1-tridecanol                   |
| 6      | 13.8                 | Heptadecane                    |
| 7      | 14.3                 | Tridecanoic acid (IS2)         |
| 8      | 14.5                 | 1-tetradecanol                 |
| 9      | 15.6                 | 1-pentadecanol                 |
| 10     | 15.9                 | Phthalate (contaminant)        |
| 11     | 17.5                 | 1-hexadecanol                  |
| 12     | 18.9                 | Hexadecanoic acid              |
| 13     | 19.1                 | 1-heptadecanol                 |
| 14     | 20.4                 | 1-oktadecanol                  |
| 15     | 21.5                 | Octadecanoic acid              |
| 16     | 21.6                 | 1-nonadecanol                  |
| 17     | 22.1                 | Tricosane                      |
| 18     | 22.6                 | 1-eicosanol                    |
| 19     | 23.6                 | 1-eneicosanol                  |
| 20     | 24                   | Pentacosane                    |
| 21     | 24.3                 | Phthalate (contaminant)        |
| 22     | 24.4                 | 1-docosanol                    |
| 23     | 25.7                 | Heptacosane                    |
| 24     | 26.1                 | 1-tetracosanol                 |
| 25     | 27.9                 | Octacosane                     |
| 26     | 28.4                 | 1-hexacosanol                  |
| 27     | 30.6                 | Nonacosan-14-one               |
| 28     | 32.4                 | Cholesterol                    |
| 29     | 32.7                 | Triacantan-15-one              |
| 30     | 35.2                 | Hentriacontan-16-one           |
| 31     | 38.3                 | Dotriacontan-16-one            |
| 32     | 40.6                 | Tritriacantanone               |
| 33     | 42.3                 | Triacconten-16-one             |
| 34     | 41.4                 | Triaccontan-16-one             |
| 35     | 45.1                 | Tetracontanone                 |
| 36     | 45.7                 | Tetracontanone                 |
| 37     | 47.1                 | Tetracontan-16-one             |
| 38     | 52                   | Pentatricontan-18-one          |
| 39     | 53.4                 | Pentatricontan-18-one          |

*All organic acids and alcohols are detected as their TMS esters and ethers, respectively*
but they plot in an area clearly distinct from the Qumran samples in the SPCA plot (Fig. 4). The two oldest samples KLR-8860 and KLR-8863 are the only ones placed inside the Qumran ellipsis but predate the Qumran habitation phase beginning ca. 85 BCE.

The organic residues

The organic residues were analysed in four ceramic samples, two sherds TL-dated to the Byzantine period, one sherd dated 568–258 BCE, and one undated sherd. The samples contained glycerolipids as the main component, in relevant amounts. The presence in all samples of dicarboxylic acids (e.g., nonanedioic acid, peak #17), of hydroxy-acids, and the co-presence of unsaturated cis and trans octadecenoic acid isomers and of 10-oxo-octadecanoic acid (#49) may indicate the oxidation undergone by the material during heating (cooking) processes in presence of oxygen, but they can also result from degradation processes occurring during burial, ascribable to microbial metabolism.

| Sample         | Identified compounds                                         | Identified material                                                                 |
|----------------|--------------------------------------------------------------|-------------------------------------------------------------------------------------|
| KLR-8008       | Mono-, hydroxy- and di-carboxylic fatty acids (linear and branched). Long chain linear and branched alcohols and alkanes/alkenes. Long chain ketones | Oxidised glycerolipids of animal and plant origin, plant waxes. Lipid Material submitted to heating |
| KLR-8009       | Mono-, hydroxy- and di-carboxylic fatty acids (linear and branched). Long chain linear and branched alcohols and alkanes/alkenes | Oxidised glycerolipids of animal and plant origin, plant waxes |
| KLR-8012, KLR-8016 | Mono-, hydroxy- and di-carboxylic fatty acids (linear and branched) | Oxidised glycerolipids of animal and plant origin, plant waxes |

Fig. 6 The radiocarbon dates and TL-dates reported in this study and reported in the literature showing the age range from 5000 BCE to 1400 CE. The TL-dates are shown with 1σ uncertainties; the radiocarbon dates with the probability distributions and 1σ bars below. Note that there is a gap in the calendar axis between 2500 and 1000 BCE, during which time interval there are no dates.
The observed profiles of the above-mentioned fatty acids are non-specific, only allowing us to hypothesize the presence of plant lipids or animal fat, or a mixture of both. The presence of plant lipids is sustained by the detection in the neutral fraction of long chain alcohols and hydrocarbons (alkanes and alkenes), particularly abundant in samples KLR-8008 and present in traces in KLR-8009 and KLR-8012: these can derive from plant epicuticular waxes extracted from vegetables [70, 71] possibly cooked in the potteries. A predominance of odd over even chain length alkanes would have further supported the occurrence of cuticular waxes, however, this was not observed in this case, leaving the discussion open for the origin of these species in the neutral fractions. Interestingly, the same sample (KLR-8008) contains a relevant amount of long chain ketones in the neutral fraction (Fig. 5b, peaks #26, 29–39 in the neutral fraction), whose presence is consistent with heating process undergone by the lipids during cooking [72, 73].

Other molecular markers mentioned in the literature for the assessment of food residue include isoprenoic fatty acids, ω-(o-alkylpenyl)alkanoic acids, β-sitosterol and polyunsaturated fatty acids (PUFAs). None of these species was detected in the chromatograms. Conversely, the concomitant presence of non-negligible amounts of branched and odd-chain fatty acids (mainly C₁₅ and C₁₇) and of cholesterol (Fig. 5d, peak #62) in samples KLR-8009 and KLR-8012 demonstrate the simultaneous presence of lipids of animal origin [74]. These odd-branched chain fatty acids are common in ruminant fats but may also have a bacterial origin [75].

Finally, the presence of a relatively high amount of contaminants such as phthalates can be ascribed to the plastic zip bags which were employed for sample shipment. Nonetheless, their presence did not affect the interpretation of the chromatographic profiles and did not interfere with the identification of the lipids of plant and animal origin.

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### Table 9  Radiocarbon dates of materials from the Christmas Cave reported in other studies

| AMS lab No | KLR lab No | Sample material | Age (BP) | δ¹³C (‰) | % C | Calibrated age range (1σ) | Calibrated age range (2σ) | Ref. |
|------------|------------|-----------------|---------|---------|-----|--------------------------|--------------------------|------|
| Gra-17427  | 3334       | Green wool QUM543 QCC230 (Old 528) | 1905 ± 30 | −20.70 | 44.1 | CE 80–205                 | CE 30–220                 | A    |
| Gra-24260  | 5462       | Red wool QUM544 QCC184 (old 530) | 1915 ± 30 | −19.05 | 44.8 | CE 75–200                 | CE 25–215                 | A    |
| Gra-17423  | 3336       | Charcoal QUM545 QCC174 CC III cave 14 wood | 5230 ± 45 | −21.71 | 69.1 | BCE 4215–3975           | BCE 4230–3960             | A    |
| Gra-24261  | 5464       | Charred wood QUM546 QCC174, Tr III w end of tree amongst rock formation | 5845 ± 45 | −9.22  | 42.0 | BCE 4785–4620           | BCE 4830–4550             | A    |
| Gra-17419  | 3333       | Textile QUM527 QCC230 | 1850 ± 40 | −19.65 | 42.9 | CE 130–235               | CE 80–325                 | A    |
| Gra-65501  | 10,430     | Textile, linen, IAA-585785, QCC585 QCC785 | 4745 ± 35 | −24.32 | 43.1 | BCE 3630–3385           | 3635–3380                 | B    |
| UCI-79816  | 1870 ± 15  | Fabric (flax) | n/a     | CE 130–210 | CE 125–220 |
| UCI-79814  | 1950 ± 15  | Rope (flax) | n/a     | CE 30–115  | CE 25–120   |
| UCI-79813  | 1900 ± 15  | Rope (flax, hemp) | n/a | CE 120–205 | CE 80–210 |
| UCI-79817  | 4760 ± 15  | Fabric (flax, hemp) | n/a | BCE 3625–3530 | BCE 3630–3520 |
| UCI-79815  | 4770 ± 15  | Fabric (flax, hemp) | n/a | BCE 3630–3530 | BCE 3630–3525 |

The ¹⁴C dates are recalibrated with the latest calibration curve IntCal20 [32]. See text for details. References: A: Rasmussen et al. [9]; B: Taylor et al. [88]; C: Murphy et al. [122].
not interfere with the determination of other organic molecules.

The obtained results therefore sustain the hypothesis of the occurrence of a mixture of different fats and oils, with variable degree of degradation and oxidation, deriving from the use of the vessels as cooking pots for different commodities containing plant and animal lipids.

**Who visited the Christmas Cave and when?**

The Christmas Cave is one of the major refuge caves in the northern Judean Desert. Its small, hidden entrance, the wide and comfortable inner passages, and its relatively easy access, made it a preferred temporary shelter in several periods, most notably during the times of Jewish rebellions against Roman rule in Judaea-Palaestina in the first and second centuries CE. The relatively large amount of material and ecofactual assemblages testify to its regional importance. It should be noted that no other large caves are known anywhere along the course of Nahal Qidron, a fact that also singles out the Christmas Cave as the most important natural hideout in this drainage basin. The availability of seasonal waterholes in the valley below the cave, and the easy access granted through the chain of desert trails, also made it a favourable hideout.

The archaeological finds discovered in the Christmas Cave indicate that human activity in the cave began during the Late Chalcolithic period. The nature of these late prehistoric occurrences is unclear, also in terms of the present findings, but it has been suggested that the Judean Desert cliff caves were used as temporary refuge places at times of societal unrest [60, 64, 76]. The mineralogy of the ceramic sherd KLR-8010 is characterized by a different composition than the rest of the pottery from later periods. Its fabric has large granitic inclusions, very poorly sorted felsic minerals, calcareous rock fragments, and the presence of Ca-based silicates as shown by XRD and FTIR indicates a high and oxidizing firing temperature (Additional file 1) (also see [77] for a description of the different Chalcolithic petrographic groups). On the other hand, the chemical data highlight that this sherd presents a very distinctive geochemical signature, different from the rest of the analysed potsherds (Fig. 4).

Following a period without occupation according to the dates available, activity was next seen again in the Early Bronze Age stage IA. This is supported by three radiocarbon dates and possibly one TL-date (KLR-10430/GrA-24261, UCI-79817, UCI-79815, and KLR-8010). From an archaeological perspective, however, the Early Bronze Age IA activity seems to be an isolated phenomenon in the regional perspective.

Considering the series of radiocarbon- and TL-dates and two coins from the days of Agrippa I and Pontius Pilate, both found during the recent survey of the cave [6], it is likely that the Christmas Cave served as a refuge during the First Jewish Revolt (ca. 68 CE), a time when Josephus specifically indicates that people fled from Jericho to the hills ahead of Vespasian’s army (War 4.449–51). It seems possible also that the cave served refugees fleeing from Jerusalem and its hinterland, located less than a day’s walk upstream. Several villages located on the desert margins east of Jerusalem are known to have existed till at least the First Revolt, for example Bethphage (Beit Pagge) and Bethany (Beit ‘Anya), as well as other sites in their vicinity, including Bethlehem ([78]: 96–110, 310). These villages and others are possible candidates for being the home settlements of refugees fleeing to the Christmas Cave. However, the provenances of the ceramics point to 3 to 4 groups of clay resources, where likely only one can originate from this area. The results obtained in the present study are in agreement with a larger technological and provenance study developed for Qumran and Jericho pottery [79]. This later study established that most of Qumran pottery was made of pure clay containing fragments of shales, and none of this pottery, especially the “scroll jars”, was assigned to the Wadi Qumran clay banks.

It should be remembered that ca. 4 km downstream from the Christmas Cave, north of the outlet of Nahal Qidron Valley into the Dead Sea lay the harbour of Khirbet Mazin, established in the Hasmonean era, likely under Alexander Jannaeus (ca. 85 BCE), and occupied through the Roman period. In addition, several complexes known as “cell sites” yield evidence for activity during the early years of the First Revolt. The main cluster of remains has been found on the southern bank of the canyon at a site called “Qidron South”, where dozens of coins dated to the 2nd year of the revolt (ca. 68 CE) were found in the 1993 excavations ([80]; see also [81, 82]). Further numismatic evidence was collected from the area of the “Cave of the Coin”, ca. 2 km south of the Qidron Outlet [83]. In 68 CE Vespasian’s troops conquered Jericho and the Qumran area thereafter creating a camp at Jericho and occupying the site of Qumran (the Period III occupation, see map of the Roman military movements Fig. 5.1 in [84]; see also [85]). Notwithstanding the geographic proximity, is seems possible that the Qidron Outlet cluster was operated by rebels during the early stages of the First Revolt, possibly as part of a guerrilla warfare intended to interrupt the conquest of the western shore of the Dead Sea (e.g., Ein Gedi) by Roman army units, while the Christmas Cave possibly served refugees fleeing from the Judean Highlands and elsewhere (whether this activity should be interpreted to have been limited to the early rather than the more advanced stages of the revolt is difficult to decide on the basis of the current
evidence, even if one would take into account Josephus’s statement that later on all escape from Jerusalem was cut off; War 4.476). This is, of course, only a hypothesis which is consistent with our observations; it does require further testing.

The Bar Kokhba coin found in the British excavations, as well as the looted coins purchased by this team, and the Roman denarius from the days of Nerva found in a nearby cave in the new survey [6], all indicate that by the end of the Bar Kokhba Revolt the Christmas Cave served once again for temporary refuge. Several radiocarbon dates, on a variety of artefacts and ecofacts, indicate that during the Second Revolt the occupation of the cave was more prolonged or more frequent than during the First Revolt. To be as precise as we can, it is safe to say that the oldest half of the dates are consistent with the First Revolt at the 2σ level; for the Second Revolt, this is the case for practically all dates, which otherwise cover the range from roughly 50 BCE to 350 CE. The origin of the refugees in this case is less clear, especially owing to the lack of archaeological data regarding the Jerusalem eastern hinterland at that time. Jerusalem itself was emptied from its Jewish population during the First Revolt. The Christmas Cave could have served refugees from Jericho, which continued to have a Jewish population, and also from Bethbassi, Herodion, and the Bethlehem region, or the Judean Highlands and wilderness, as most refuge caves opening into the main valleys draining the eastern flank of Judea towards the Dead Sea (see [6] for a detailed reconstruction).

It is unlikely that any occupants of the cave came from Qumran. Our analysis of the ceramic material exhibits 3–4 different sources of the clay. In addition, the ceramic material reflects various technological production methods. It seems that only two of the ceramic sherds could be derived from the Qumran area. These two sherds are, however, older than the main habitation period in Qumran. Three other sherds are TL-dated in accordance with the Qumran settlement, but their chemistry and mineralogy are different from ceramic materials from Qumran indicating that they do not originate from the Qumran area. The organic analyses of the sherds show traces of decayed organic material stemming from both plants and animal fat, including the effect of cooking the food stuff.

According to Allegro’s testimony ([2]: 121–122), his archives and the meagre remains found in the recent survey ([6]: 47), the TL-dates and a single radiocarbon date, the Christmas Cave was also ephemerally used during the Byzantine and Islamic eras. It is possible that the single date from the Mamluk period in the present study (GrA-53271/KLR-8858) is related to Islamic-period pottery collected by Allegro’s DSSF team. The phenomenon of returning to refuge cliff caves in later periods is attested also in other instances in the Judean Desert, e.g., in the Murabba’at caves located ca. 12 km south of the Christmas Cave [76, 86, 87]. The extensive use of caves, rock shelters and built structures in the Lower Nahal Qidron Area for seasonal retreats of monks during the Byzantine and Early Islamic periods [82] may also account for the scarce evidence from the Christmas Cave during these periods.

Conclusions

New radiocarbon dates and TL-dates are presented, which collectively demonstrate intermittent occupation of the Christmas Cave for more than 6000 years. There is a concentration of evidence that coheres with the impression that this was a refuge cave of the Bar Kokhba Revolt. However, it seems there were multiple other visits to this cave.

In terms of chronology there is an overlap between the occupation of the Qumran settlement and some use of the Christmas Cave, in the first century BCE and first century CE. However, apart from a chronological correlation there is nothing in the data that indicates a closer connection between the people who used the Christmas Cave and the occupants of Qumran in the main phase of habitation. This supports the general notion of dichotomy between the phenomenon of Qumran and related caves and that of the refuge cliff caves distributed in other parts of the Judean Desert, to where rebels and refugees from Judea fled in times of stress, i.e., during the First and Second Jewish Revolts. The new data presented in this study confirm that there is no material reason to connect the Christmas Cave with Qumran.

In conclusion, the new evidence presented in this study is in agreement with the textile evidence, which also indicates an origin different than Qumran. This study tentatively suggests a connection with villages from the Judean Highlands east of Jerusalem as a possible place of origin for the refugee occupants of the Christmas Cave.

Overall, in deploying multiple scientific methods we can see how it is possible to define the dating of human habitation of the cave with much more accuracy, thus allowing for a more varied and expansive understanding of how the cave was used in different times. While materials are likely concentrated in the mid-second century, the cave clearly attracted diverse people over the millennia. We have also been able to test how the people who came here may have related to other places occupied concurrently. Further comparative studies using such methods would undoubtedly lead to clearer specificity of such relationships and occupations.
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