Influence of increasing combustion temperature on the AMS $^{14}$C dating of modern crop phytoliths

Jinhui Yin$^{1,2}$, Xue Yang$^1$ & Yonggang Zheng$^1$

$^1$State Key Laboratory of Earthquake Dynamics, Institute of Geology, China Earthquake Administration, Beijing 100029, China, $^2$State Key Laboratory of Loess and Quaternary Geology, Institute of Earth Environment, Chinese Academy of Sciences, Xi’an 710075, China.

Several attempts have been made to directly date phytoliths, but most $^{14}$C results are not consistent with other independent chronologies. Due to the limited dataset, there is not a clear explanation for these discrepancies. Herein, we report the $^{14}$C ages of phytolith-occluded carbon (PhytOC) from contemporary rice and millet crops that were combusted at different temperatures to investigate the relationship between the combustion temperature and resulting $^{14}$C age. Our results show that the $^{14}$C age of PhytOC increases directly with combustion temperature (up to 1100 $^\circ$C) and results in age overestimations of hundreds of years. Considerably older ages are observed at higher temperatures, suggesting that it may be possible to distinguish between two fractions of organic carbon in phytoliths: labile and recalcitrant carbon. These findings challenge the assumption that PhytOC is homogeneous, an assumption made by those who have previously attempted to directly date phytoliths using $^{14}$C.

Rice and millet are two staple food crops that have been cultivated in the Yellow and Yangtze River basins of China since the early Neolithic period$^{1-2}$. There is some uncertainty regarding the origin and spread of these important crops across East Asia, particularly China, due to the lack of adequate evidence from crop remains in archaeological sites$^{3-4}$. In these archaeological sites, with climates ranging from moist and warm to semi-arid, typically only charred plant remains survive for several millennia$^{6-7}$. As an alternative to these scarce, charred plant remains, phytoliths (biogenetic opals formed in plants when roots absorb soluble silica) represent a potentially useful geochronometer$^{8,9}$. When plants die and decay, phytoliths are released into the soil and sediment$^{10,11}$. Phytoliths are extremely durable and can be preserved in large amounts in most archaeological sites, and in some situations, they can form well-defined strata$^{12,13}$.

Organic carbon (PhytOC) is occluded during the formation of a phytolith and sometimes represents up to 2% of the dry weight of a plant$^{14-17}$. The PhytOC contents of millet and rice have been estimated to be 1.36 mg g$^{-1}$ and 2.8 mg g$^{-1}$, respectively$^{18,19}$. The small sample size required by accelerator mass spectrometry (AMS) allows for the analysis of $<500$ µg of carbon; therefore, only a handful of soil can provide a sufficient amount of phytoliths for radiocarbon analysis.

Over the past decades, there have been a few attempts to measure the $^{14}$C age of fossil phytoliths$^{8,10,13,20-22}$. However, most $^{14}$C ages of phytoliths are not consistent with independent chronologies. These discrepancies have been attributed to preferential oxidation$^1$, stratigraphic disturbances$^{20}$, or sample extraction methods$^{23,24}$. Moreover, some new $^{14}$C analyses of PhytOC from phytolith concentrates extracted from living grass or bamboo reported ages of up to several thousand years older$^{25-27}$. A clear explanation for these surprisingly older dates is still lacking due to the limited dataset and exploration of extraction protocols$^{27,28}$.

One hypothesis suggests that there may be two possible components of photosynthetic and recalcitrant organic matter in phytoliths$^{27}$. The exact influence of these two components of PhytOC on $^{14}$C dating needs to be examined in detail because previous attempts to date $^{14}$C PhytOC have focused on the total organic matter within the phytolith, and these attempts have clearly failed. In some cases, the combustion temperature can be used to separate heterogeneous mixtures of labile and refractory carbonaceous components, each of which may have a different apparent radiocarbon age$^{29-31}$. Recently, a more robust phytolith extraction protocol suitable for carbon isotopic analysis was reported$^{32}$, and we use this protocol to further investigate the potential source of PhytOC. In this study, phytoliths were isolated from two different species of modern crops (rice and millet) using a modified version of the recently published protocol$^{32}$ and were then combusted with CuO powder at different temperatures.
ranging from 160°C to greater than 1400°C. We analyse the 14C and δ13C ratios produced by the range of combustion temperatures and compare them with the current atmospheric ratio to investigate the influence of combustion temperature on 14C age. We explore these results in an attempt to delineate between the two hypothesised components of occluded organic carbon (PhytOC).

Results

Morphology of phytoliths. The extracted rice straw and millet phytoliths were examined using a scanning electron microscope (SEM) to ensure a lack of visible cellulose adhering to the outside of the phytoliths and to ensure the presence of intact samples with diameters of at least 20 μm (Supplementary Figure S1). When the phytoliths were combusted with CuO to form CO2 gas, a manometer was used to monitor the production of CO2 as the temperature increased from 160°C to 900°C. The combustion profiles of the phytolith samples show that the majority of CO2 is generated in two distinct temperature ranges: 500–600°C and 800–900°C (Supplementary Figure S2). This release pattern can be interpreted in terms of carbon being derived from a mixture of at least two components. As shown in Supplementary Figure S1, when the temperature reaches 900°C, most phytoliths lose their distinct morphological characteristics, and when the samples are heated to 1100°C, the phytoliths disappear.

14C concentrations in the modern atmosphere. The AMS 14C concentrations for modern crops and their phytolith samples, along with the δ13C values and carbon yield rates, are listed in Supplementary Table S1. The 14C concentration in the atmosphere can be estimated from the three living crops. The modern fraction (Fm, the deviation of the 14C/12C ratio of a sample from the 14C/12C ratio of “modern carbon”) values from the direct AMS dating of the plant material for AD 2011 rice straw, AD 2012 rice straw and AD 2012 millet are 1.0312 ± 0.0028, 1.0288 ± 0.0027, and 1.0223 ± 0.0027, respectively. These values are consistent with the atmospheric radiocarbon derived from the bomb radiocarbon curve in the Northern Hemisphere atmosphere22–25, suggesting that the 14C in the plant is in equilibrium with the 14C in the atmosphere.

14C concentrations in phytoliths with increasing combustion temperature. The 14C concentration in an individual phytolith sample was normalised to that in the host crop to more readily identify the deviation between the measured 14C concentration in phytoliths and that measured for the atmosphere. We observe that the AMS 14C concentrations in all 16 phytoliths are lower than those in the atmosphere (Figure 1).

We can confirm a 14C depletion of several percent in PhytOC at combustion temperatures below 1100°C (Supplementary Table S1). For 14C of phytoliths in AD 2011 rice straw, a chi-square test was applied to compare the Fm values of phytoliths combusted at temperatures below 1100°C. The chi-square value of this dataset is 7.04, which is less than 11.07, the critical value for 95% confidence and 5 degrees of freedom, implying that the values are indistinguishable. The weighted mean for the 6 Fm values of phytoliths in AD 2011 rice straw is 0.9393±0.0013. This 14C level is 6% lower than the atmospheric concentration and corresponds to an apparent radiocarbon age of 480 yrs. Similarly, the Fm values of PhytOC from AD 2012 rice straw (C3 plant) and AD 2012 millet (C4 plant) also passed the chi-square test. We obtained weighted means of 0.9684±0.0019 and 0.9577±0.0013, respectively. These results indicate that the deficiency in the 14C content of the two samples is 3% to 4% lower when combusted at temperatures below 1100°C, equivalent to ages of 250 yrs and 320 yrs older than the true ages of the plants, respectively.

In a further experiment, 0.5 g and 0.3 g of phytolith samples were sent to Beta Analytic Inc. to investigate PhytOC during higher-temperature combustions. The phytolith samples were combusted at temperatures greater than 1400°C via the exothermic reaction between tin and oxygen25. The two PhytOC samples combusted at temperatures above 1400°C produced Fm values of 11% and 13% lower than those of the atmosphere, respectively, resulting in age estimates that were ~1000 yrs older than the true ages. Similarly, the average δ13C values of PhytOC in AD 2011 rice straw and in AD 2012 rice straw combusted at temperatures below 1100°C are −33.66‰ and −32.67‰, respectively (Supplementary Table S1). In contrast, when combusted at temperatures over 1400°C, the δ13C values were 1.17‰ and 2.53‰ heavier, respectively, than those of PhytOC combusted at the lower temperature.
Background check. We also performed one $^{14}$C determination on a phytolith assemblage isolated from L1-3 loess to evaluate the background value for this extraction procedure. The OSL age of L1-3 loess is approximately 71 ka BP, which is beyond the limit of the radiocarbon dating method. The $F_m$ value of phytoliths from L1-3 loess is 0.0051 ± 0.0001, corresponding to 42,380 ± 180 yr BP, which is statistically indistinguishable from the background of our AMS line background, which is equivalent to 42,750 ± 190 yr BP.

Discussion

If phytoliths are to become a new alternative candidate for $^{14}$C dating for establishing a reliable age control for archaeological sites, the $^{14}$C content in PhytOC must be in equilibrium with the $^{14}$C content in the atmosphere when the organism dies. However, in this study, the most striking feature is that the $^{14}$C contents in all 16 aliquots of phytoliths are lower than that in the atmosphere. These phytoliths were isolated from modern crops using a reliable, published protocol with only slight modifications. PhytOC accounted for 0.06–0.10% and 0.07% of the dry weight of rice and millet phytoliths, respectively (Supplementary Table S1), which is at least 20 times lower than the range of 14–34 mg $^{-1}$ for rice and 25.1 mg $^{-1}$ for millet. These results indicate that the protocols used to extract phytoliths in this study are very harsh and can effectively eliminate possible sources of contamination during processing. We require 1200–2200 mg of phytoliths to produce graphite samples with $\sim$0.8 mg of carbon for AMS $^{14}$C analysis. The precision and reproducibility of our experimental procedure, better than 0.3% for modern samples, could be evaluated by the mean $F_m$ value of the standard material oxalic acid II during AMS $^{14}$C analysis. The background test of the phytolith extraction from L1-3 loess indicates that the protocols in our lab minimise the possibility of introducing contaminant carbon, specifically modern carbon, to the phytolith sample during extraction, graphitisation, and measurement. Moreover, the $^{14}$C overestimated ages (by hundreds of years) of phytolith samples with different weights from each rice crop are consistent when the samples are combusted at temperatures below 1100 $^\circ$C. The average $\delta^{13}$C values of PhytOC also implies that a new carbonaceous compound with a heavier carbon isotope is liberated from phytoliths when combusted at temperatures greater than 1400 $^\circ$C. Regardless, the change in the isotopic composition of PhytOC in rice straw cannot explain the systematic age offset. Therefore, this $^{14}$C depletion in the phytoliths most likely originates from the temperature interval at which the sample itself was combusted. A similar trend has been described in other studies that used the same laboratory protocols. In addition, this problem is not unique to our laboratory protocols because the AMS $^{14}$C dates of phytoliths isolated from mature and recently senesced bamboo leaves using another separation method (microwave digestion) were inexplicably 3.5 ka and 1.9 ka too high, respectively. These results clearly show a consistent shift in the $^{14}$C levels when comparing phytoliths to modern plants regardless of their locations, species, and extraction methods. As mentioned above, for fossil phytoliths, most $^{14}$C ages were inconsistent with expected or independent chronologies. This problem could reveal that, even though the $^{14}$C of the plant is in equilibrium with the atmosphere, the $^{14}$C content of PhytOC is not.

Phytoliths can be changed by heating. Dry ashing has been widely used to recover phytoliths from plants since the beginning of phytolith analysis. This method involves the incineration of plant tissue in muffled furnaces at temperatures of at least 500 $^\circ$C. This technique has been suggested to cause shrinkage, warping and changes to the refractive index of phytoliths. For rice crop phytoliths, it has been found that the original physical characteristics are significantly altered when the extraction temperature exceeds 900 $^\circ$C. Previous studies indicated that the relationship between the $^{14}$C age and combustion temperature can be used to quantify the relative contributions of different pools of organic carbon. For phytoliths in AD 2012 rice straw, the $F_m$ of PhytOC was depleted by approximately 4% when the sample was combusted below 1100 $^\circ$C and was even more depleted (to approximately 12%) when combusted over 1400 $^\circ$C. We observe a similar trend between the combustion temperature and $F_m$ depletion of PhytOC in AD 2011 rice straw, suggesting that a mechanism may govern the $^{14}$C variation with temperature. Our results indicate multiple carbon species, each of which may potentially have a unique $^{14}$C concentration in phytoliths. One possible explanation is that there are two components of organic matter in phytoliths. The low-temperature pool (combusted below 900 $^\circ$C) is associated with relatively labile carbon, whereas the high pool (combusted above 900 $^\circ$C) is commonly associated with recalcitrant carbon.

We still have an incomplete understanding of PhytOC in phytoliths. An early hypothesis suggested that recalcitrant PhytOC (biasing the $^{14}$C results towards age overestimates) could be brought from the soil to the plant by root uptake, but this hypothesis is still under debate. We entertain another hypothesis to explain the discrepancies in the $^{14}$C contents of our measured samples: two possible pools of $^{14}$C exist in the phytoliths, each of which is attributed to a different type of phytolith development. Phytoliths are biogenetic opals that form in the cell walls, cell lumina and intercellular spaces of plants. If biosilicification primarily occurs in the cell walls, hollow forms of phytoliths are formed and often contain cellular organelles such as mitochondria and plastids. However, if biosilicification occurs in cell lumina or between cells, solid forms of phytoliths develop and contain mostly lipids and nucleic acids. For sugar cane, the amount of PhytOC retained within hollow rather than solid phytoliths has been estimated to be 10.12% and 0.15%, respectively. These results suggest that, on average, the PhytOC from hollow phytolith cavities is 50 times more abundant than the PhytOC from solid phytoliths. Similar trends have also been observed in sorghum. We also observe that organic materials are liberated from phytoliths at two distinct stages. For example, rice straw from AD 2011, the carbon yield rate was 0.07%, and the $F_m$ value was 0.9363 ± 0.0038 when combusted below 1100 $^\circ$C (Supplementary Table S1). When combusted above 1400 $^\circ$C, although the carbon yield rates did not change, the $F_m$ value was 0.8724 ± 0.0046. This $F_m$ value, attributed to the recalcitrant fraction of PhytOC, increased the apparent age of phytoliths even though the amount of recalcitrant carbon is minor compared to the total amount of carbon in the samples. The ratio of carbon released below 1100 $^\circ$C to that released above 1100 $^\circ$C during combustion can be estimated using a simple, two end-member mixing model. If one end-member is the $^{14}$C released below 1100 $^\circ$C ($F_m=0.9363 ± 0.0038$ for rice straw from AD 2011), then the other end-member is the $^{14}$C released above 1100 $^\circ$C ($F_m=0$, dead carbon, no $^{14}$C); we calculate the ratio as 13. Similarly, for AD 2012 rice straw, we calculate the ratio as 12. Therefore, it is possible that the low-temperature carbon pool originates from hollow phytoliths formed in porous cell walls during photosynthesis. This carbon pool is more readily available for oxidation and lost from phytoliths at lower temperatures. Conversely, the high-temperature carbon pool may represent solid phytoliths formed in cell lumina. Further investigation of the mechanisms responsible for $^{14}$C differentiation in the two fractions of PhytOC is necessary.

Therefore, we are working toward a new protocol that can successfully remove the recalcitrant carbon from phytoliths such that we can establish a reliable $^{14}$C dating method for phytoliths. Among the many potential uses, a new protocol will allow us to better constrain the timing and origins of agriculture in East Asia.

Methods

Materials. Phytoliths were extracted from modern plants: two rice (Oryza sativa) stems (with leaves) and one millet (Panicum miliaceum). The dried rice straws were harvested from the Hubei province near the Yangtze River (115° E, 30° N) in two successive years: AD 2011 and AD 2012. The millet was obtained from the Nihewan basin (114° E, 40° N) in northwest Beijing in AD 2012. A fossil loess sample was
collected from the Weinan section on the southeastern Chinese Loess Plateau and was dated by optically stimulated luminescence (OSL) to ca. 71 ka BP.

Phytoliths isolated from modern crops and loess. Phytoliths were extracted from modern plants using a protocol involving both a sink–float specific gravity and a wet-diggestion method detailed below (modified from a recently published protocol). Approximately 700 g of each dried crop was used to obtain contemporary phytoliths. The rice and sorghum samples were initially cut into cm-sized pieces and immersed in a distilled H2O ultrasonic bath to remove any matter adhering to their surfaces. The sample was then immersed in 1 N hydrochloric acid (HCl) for 4 h to eliminate carbonatic material and stirred every half hour. The samples were washed three times in distilled H2O after each agitation. The treated samples were dried in an oven at 70°C for three days. A 650 g aliquot of treated rice straws was weighed into a 4000 ml beaker and digested with ~5 ml of concentrated H2SO4 per gram of dry plant material under a fume hood. The digestion lasted for 2 h at 70°C, and the sample then set unheated overnight. The following day, the hot plate was re-heated to 70°C, and hydrocyanic acid (H2CN, 30%) was gradually added to destroy organic material until the liquid was clear. The supernatant liquid was poured off, and the supernatant liquid (with minimal organic matter) was poured off. This process was repeated three times.

The recovered phytoliths were dried and retained. After the phytoliths were recovered, they were washed and reheated at 70°C for 2 h in concentrated HNO3 and NaClO2 to ensure that any organic material clinging to the outer surfaces was removed. The sample set unheated overnight, and the liquid was decanted. This step was repeated 4 times to maximize the oxidation of organic material. The phytoliths were rinsed 3 times with distilled water and dried. Subsequently, they were immersed in 0.001 M NaOH solution and heated at 70°C for 15 min to remove any alkaline-soluble forms of organic matter. The isolated phytoliths were washed again with concentrated HNO3 and rinsed 3 times with distilled water. Finally, phytolith concentrates were dried at 70°C, weighed and observed with an optical microscope.

The background extraction method for phytoliths in loess is similar to the aforementioned method for modern plants. Sand grains larger than phytoliths were first removed by wet-sieving through an 80-mesh screen and then added to 1 N HCl to eliminate carbonates. Subsequently, two 50-litre settling containers were used to separate the clays and organic colloids. The sedimentation-decanting procedure was repeated several times until the water was clear. The sample then was dried at 70°C. The subsequent steps were the same as for the modern plants, but the extraction is usually repeated twice to increase phytolith recovery.

Cellulose extracted from modern crops. The radiocarbon content of cellulose separated from modern crops was used to represent the atmospheric δ13C values of crops in China. Chinese Sci. Bull. 56, 3451–3456 (2011).

Li, Z., Song, Z., Parr, J. F. & Wang, H. Occluded C in rice phytoliths: implications for radiocarbon dating. Radiocarbon 52, 275–281 (2009).

Prior, C. A., Carter, J. & Rieser, U. Are phytolith radiocarbon dates reliable? The 10th International Conference on Accelerator Mass Spectrometry, Berkeley, USA. (http://ilni.confex.com/ilni/ams10/techprogram/P1592.HTM) (2005, September 25–30).

Carter, J. A. Atmospheric carbon isotope signatures in phytolith-occluded carbon. Quatern. Int. 193, 20–29 (2009).

Santos, G. M. et al. The phytolith 14C puzzle: a tale of background determinations and accuracy tests. Radiocarbon 52, 113–128 (2010).

Santos, G. M. et al. Possible source of ancient carbon in phytolith concentrates from harvested grasses. Biogeosciences 9, 1873–1884 (2012).

Sullivan, L. A. & Parr, J. F. Comment on: “Possible source of ancient carbon in phytolith concentrates from harvested grasses” by G. M. Santos et al. Biogeosciences 9, 2273–2275 (2012).

McGeen, J. et al. Stepped-combustion 14C dating approach: A comparison with established techniques. Radiocarbon 43, 259–261 (2001).
33. Levin, I. et al. Observations and modelling of the global distribution and long-term trend of atmospheric \(^{14}\text{CO}_2\). *Tellus B* 62B, 26–46 (2010).
34. Graven, H. D., Guilderson, T. P. & Keeling, R. F. Observations of radiocarbon in \(^{14}\text{CO}_2\) at La Jolla, California, USA 1992–2007: Analysis of the long-term trend. *J. Geophys. Res.* 117 (2012).
35. Hua, Q., Barbetti, M. & Rakowski, A. Z. Atmospheric Radiocarbon for the Period 1950–2010. *Radiocarbon* 55, 2059–2072 (2013).
36. Kang, S., Lu, Y. & Wang, X. Closely-spaced recuperated OSL dating of the last interglacial paleosol in the southeastern margin of the Chinese Loess Plateau. *Quat. Geochronol.* 6, 490–491 (2011).
37. McClaran, M. P. & Umlauf, M. Desert grassland dynamics estimated from carbon isotopes in grass phytoliths and soil organic matter. *J. Veg. Sci.* 11, 71–76 (2000).
38. Rovner, I. [Plant opal phytolith analysis: major advances in archaeobotanical research] *Advances in archaeological method and theory* [Schiffer, M. B. (ed.)] 225–266 (Academic Press, New York, 1983).
39. Elbaum, R., Weiner, S., Albert, R. M. & Elbaum, M. Detection of burning of plant materials in the archaeological record by changes in the refractive indices of siliceous phytoliths. *J. Archaeol. Sci.* 30, 217–226 (2003).
40. Parr, J. F., Lentfer, C. J. & Boyd, W. E. A comparative analysis of wet and dry ashing techniques for the extraction of phytoliths from plant material. *J. Archaeol. Sci.* 28, 875–886 (2001).
41. Wu, Y., Wang, C. & Hill, D. V. The transformation of phytolith morphology as the result of their exposure to high temperature. *Microsc. Res. Techniq.* 75, 852–855 (2012).
42. Pronin, J. et al. [Individual characterization of phytoliths: experimental approach and consequences on paleoenvironment understanding] *Phytoliths: Applications in Earth Science and Human History* [Meunier, J. D. & Colin, F. (eds.)] 329–341 (A.A. Balkema Publishers, Lisse, 2001).
43. Smith, F. A. & Anderson, K. B. [Characterization of organic compounds in phytoliths: Improving the resolving power of phytolith \(\delta^{13}\text{C}\) as a tool for palaeoecological reconstruction of C3 and C4 grasses] *Phytoliths: applications in earth sciences and human history* [Meunier, J. D. & Colin, F. (eds.)] 317–327 (A.A. Balkema Publishers, Lisse, 2001).
44. Perry, C. C., Williams, R. J. P. & Fry, S. C. Cell wall biosynthesis during silicification of grass hairs. *J. Plant Physiol.* 126, 437–448 (1987).
45. Elbaum, R., Melamed-Bessudo, C., Tuross, N., Levy, A. A. & Weiner, S. New methods to isolate organic materials from silicified phytoliths reveal fragmented glycoproteins but no DNA. *Quatern. Int.* 193, 11–19 (2009).
46. Évett, R. Interactive comment on “Comment on: "Possible source of ancient carbon in phytolith concentrates from harvested grasses" by G. M. Santos et al.(2012)” by L. A. Sullivan and J. F. Parr. http://www.biogeosciences-discuss.net/9/C6286/2012/ (2012) Date of access: 5 December 2012.
47. Song, Z., Wang, H., Strong, P. J., Li Z. & Jiang, P. Plant impact on the coupled terrestrial biogeochemical cycles of silicon and carbon: Implications for biogeochemical carbon sequestration. *Earth-Sci. Rev.* 115, 319–331 (2012).
48. Piperno, D. R. *Phytolith analysis: an archaeological and geological perspective.* (Academic Press, San Diego, 1988).
49. Hodson, M. Interactive comment on “Comment on: "Possible source of ancient carbon in phytolith concentrates from harvested grasses" by G. M. Santos et al. (2012)” by L. A. Sullivan and J. F. Parr. http://www.biogeosciences-discuss.net/9/C5612/2012/ (2012) Date of access: 12 November 2012.
50. Parr, J. F. & Sullivan, L. A. Comparison of two methods for the isolation of phytolith occluded carbon from plant material. *Plant Soil* 374, 45–53 (2014).
51. Wang, Y. & Liu, H. *The study of phytolith and its application* (in Chinese). (China Ocean Press, Beijing, 1993).
52. Liu, W., An, Z., Zhou, W., M., J. & Cai, D. Carbon isotope and C/N ratios of suspended matter in rivers: an indicator of seasonal change in C4/C3 vegetation. *Appl. Geochem.* 18, 1241–1249 (2003).
53. Vogel, J. S., Southon, J. R., Nelson, D. E. & Brown, T. A. Performance of catalytically condensed carbon for use in accelerator mass spectrometry. *Nucl. Instrum. Meth.* B 5, 289–293 (1984).
54. Liu, K. et al. A new compact AMS system at Peking University. *Nucl. Instrum. Meth.* B 259, 23–26 (2007).

Acknowledgments

This work was jointly supported by State Key Laboratory of Earthquake Dynamics (Project No. LED2013A08) and the fundamental scientific research special project of Institute of Geology, China Earthquake Administration (No: IGCEA-1116), and State Key Laboratory of Loess and Quaternary Geology, Institute of Earth Environment, CAS(SKLQG0707). We are especially indebted to the technical support of the staff in PKUAMS center for their assistance and patient accommodation of our many AMS measurement requests.

Author contributions

J. Y. conceived the study, phytolith extraction, data interpretation, and wrote the manuscript. X. Y. performed the \(^{14}\text{C}\) graphite and discussion of the results. Y. Z. assigned sample collection and sample pre-clean.

Additional information

Supplementary information accompanies this paper at http://www.nature.com/scientificreports/

Competing financial interests: The authors declare no competing financial interests.

How to cite this article: Yin, J., Yang, X. & Zheng, Y. Influence of increasing combustion temperature on the AMS \(^{14}\text{C}\) dating of modern crop phytoliths. *Sci. Rep.* 4, 6511; DOI:10.1038/srep06511 (2014).

This work is licensed under a Creative Commons Attribution-NonCommercial-NoDerivs 4.0 International License. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder in order to reproduce the material. To view a copy of this license, visit http://creativecommons.org/licenses/by-nc-nd/4.0/