Past fire dynamics inferred from polycyclic aromatic hydrocarbons and monosaccharide anhydrides in a stalagmite from the archaeological site of Mayapan, Mexico

Julia Homann¹, Niklas Karbach¹, Stacy A. Carolin^{2,3}, Daniel H. James², David Hodell², Sebastian F. M. 5 Breitenbach⁴, Ola Kwiecien⁴, Mark Brenner⁵, Carlos Peraza Lope⁶, Thorsten Hoffmann¹

¹Department of Chemistry, Johannes Gutenberg-Universität, Mainz, Germany
 ²Department of Earth Sciences, University of Cambridge, Cambridge, UK
 ³School of Archaeology, University of Oxford, Oxford, UK
 ⁴Department of Geography and Environmental Sciences, Northumbria University, Newcastle upon Tyne, UK
 ⁵Department of Geological Sciences, University of Florida, FL, USA

⁶Instituto Nacional de Antropología e Historia, Centro INAH Yucatán, Mérida, México

Correspondence to: Thorsten Hoffmann (hoffmant@uni-mainz.de)

Abstract

- 15 Speleothems (cave stalagmites) contain inorganic and organic substances that can be used to infer past changes in local and regional paleoenvironmental conditions. Specific biomarkers can be employed to elucidate the history of past fires, caused by interactions among climate, regional hydrology, vegetation, humans, and fire activity. We conducted a simple solid-liquid extraction on pulverised carbonate samples to prepare them for analysis of 16 polycyclic aromatic hydrocarbons (PAHs) and three monosaccharide anhydrides (MAs). The preparation method requires only small samples (0.5-1.0 g); PAHs and MAs were measured by GC-MS and LC-HILIC-MS, respectively. Detection limits range from 0.05-2.1 ng for PAHs and 0.01-0.1 ng for MAs. We applied the method to 10 samples from a ~400-year-old stalagmite from Cenote Ch'en Mul, at Mayapan (Mexico), the largest Postclassic Maya capital of the Yucatán Peninsula. We found a strong correlation (r=0.75, p < 0.05) between</p>
- the major MA (levoglucosan) and non-alkylated PAHs (Σ15). We investigated multiple diagnostic PAH and MA ratios and found that although not all were applicable as paleo-fire proxies, ratios that combine PAHs with MAs are promising tools for
 identifying different fire regimes and inferring the type of fuel burned. In the 1950s and 1960s, levoglucosan and Σ15 concentrations roughly doubled compared to other times in the last 400 years, suggesting greater fire activity at Mayapan during these two decades. The higher concentrations of fire markers may be due to have been a consequence of land clearance

at the site and explorations of the cave by the Carnegie Institution archaeologists.

1 Introduction

30 Speleothems are valuable continental paleoenvironmental archives. They can grow continuously over long time periods (Fairchild and Baker, 2012; Gałuszka et al., 2017) and can provide very high-resolution (sub-annual) proxy time series of past climate and environment (Ridley et al., 2015; Braun et al., 2023). Most can be reliably dated using uranium series isotopes (Mason et al., 2022; Scholz and Hoffmann, 2008). Well-established inorganic proxies such as δ^{18} O and δ^{13} C are increasingly complemented by organic biomarkers that record more specific aspects of the paleoenvironmental conditions (Baker et al.,

- 35 2019; Blyth et al., 2016; Bosle et al., 2014; Heidke et al., 2019; Homann et al., 2022). One important aspect of past environments is the occurrence and dynamics of fires, whether natural or anthropogenic. The development of fire-sensitive proxies in paleoenvironmental archives can help elucidate interactions among climate, regional hydrology, vegetation, and fire activity (Campbell et al., 2023). Two such fire proxies are polycyclic aromatic hydrocarbons (PAHs) and monosaccharide anhydrides (MAs). PAHs are products of incomplete combustion of biomass and petrogenic fuels over a wide temperature 40 range (200-700 °C) (Han et al., 2020; Tobiszewski and Namieśnik, 2012; Yunker et al., 2002b; McGrath et al., 2003).
- -The presence of specific biomarkers is indicative of the fuel source. Retene, for example, is a PAH that is a unique marker for the combustion of gymnosperm biomass (Ramdahl, 1983; Wakeham et al., 1980). MAs are formed only during combustion of biomass at lower temperatures (150-350 °C). The predominant MA is levoglucosan, which is formed during combustion of cellulose (Elias et al., 2001; Simoneit, 2002). Both PAHs and MAs can be present in the gas and the particle phases (Lammel
- 45 et al., 2009; Xie et al., 2014) and have been reported to undergo long-range transport (Luo et al., 2020; Zennaro et al., 2014). Their atmospheric residence times, however, differ widely, ranging from 1-3 hours (gas phase) to 4-5 days (particulate phase) for PAHs and 2-26 days for MAs (Bai et al., 2013; Fraser and Lakshmanan, 2000; Slade and Knopf, 2013). This difference in atmospheric longevity may explain for why Denis et al. (2012) found PAHs to record only very local (≤0.5 km) fires, whereas known fires that occurred 1-2 km from the sampling site were not recorded.
- 50 These different behaviours of PAHs and MAs are the rationale for analysis of both fire proxies in tandem, not only to detect the presence of fire but also to explore changes in fire regime (e.g. fire frequency, intensity, and fuel source). Whereas PAHs and MAs have been measured simultaneously in lake sediment cores (Argiriadis et al., 2018; Battistel et al., 2017; Callegaro et al., 2018), they have not yet been investigated jointly in speleothems, although both have been extracted individually from stalagmites (Argiriadis et al., 2019; Homann et al., 2022; Perrette et al., 2008). Another motivation to survey different markers
- 55 simultaneously is to better understand their transport and incorporation mechanisms in speleothem carbonate. In principle, PAHs and MAs can be transported into a stalagmite either via infiltrating water, and/or via deposited aerosol particles, delivered via cave ventilation or by cave-internal sources, respectively (e.g. campfires, torched, candles, or petroleum lamps). Whereas samples from caves without substantial ventilation (i.e. with no or very narrow entrances) would only archive a dripwater-derived signal, it is likely that in caves with significant substantial ventilation and easy human access, markers would 60 be introduced via both dripping water and externally introduced or cave internally formed aerosol particles.

Here we focus on the extraction of PAHs and MAs from speleothem carbonate. We report results of sequential extraction of 16 PAHs and three MAs from a speleothem and their analysis using GC-MS and LC-HILIC-MS, respectively. We applied the new, simple method to a young (~400-year-old) stalagmite (MAYA-22-7) collected in August 2022 from Cenote Ch'en Mul, Mayapan, <u>YucátanYucatán</u> Peninsula, Mexico. Several smaller stalagmites from the same cave contained charcoal inclusions,
indicating that the studied stalagmite was a promising candidate for this proof-of-concept study. We used the measured PAH

and MA concentrations and selected diagnostic ratios to interpret past aspects of fire dynamics in the cave and surrounding area. Even though some of the biomarkers were below the limit of detection (LOD), the analysed stalagmite samples provided valuable hints about past fire dynamics.

2 Methods

70 2.1 Materials

Ultrapure methanol (MeOH, LC/MS grade) was obtained from Carl Roth, ultrapure dichloromethane (DCM, LC/MS grade, \geq 99.8 %) and water (LC/MS grade) were purchased from Fisher Scientific. Ultrapure acetonitrile (ACN, LC/MS grade) and ammonium formiate (99 %) were obtained from VWR Chemicals. Analytical standards of levoglucosan (1,6-Anhydro- β -D-glucopyranose, 99 %), *p*-terphenyl (\geq 99 %), naphthalene-d₈ (2000 µg mL⁻¹ in DCM), acenaphthene-d₁₀ (2000 µg mL⁻¹ in

- 75 DCM), phenanthrene-d₁₀ (2000 µg mL⁻¹ in DCM), and a standard of 16 PAHs (QTM PAH Mix, 2000 µg mL⁻¹ in DCM), as well as ultrapure ethyl acetate (EA, GC/MS grade) were purchased from Sigma-Aldrich. An analytical standard of ¹³C₆-levoglucosan (98 %) was obtained from Cambridge Isotope Laboratories. An analytical standard of retene (10 µg mL⁻¹ in cyclohexane) was purchased from LGC and analytical standards of mannosan (1,6-Anhydro-β-D-mannopyranose) and galactosan (1,6-Anhydro-β-D-galactopyranose) were obtained from Carbosynth Ltd. Ultrapure water with 18.2 MΩ resistance
- 80 was produced using a Milli-Q water system from Merck Millipore. Solid-phase extraction columns (CHROMABOND SiOH, 3 mL tubes, 45 μm particle size) were purchased from Macherey-Nagel.



Figure 1: Scan of stalagmite MAYA-22-7 with samples taken for U-Th dating and biomarker analysis indicated in red and cyan, respectively. For simplicity, rectangles indicate the position of the samples projected <u>onto</u> the central growth axis, whereas samples
 were drilled off-axis, along growth layers. Black <u>vertical</u> lines indicate the growth axis along which grey values were measured. Ages are given in calendar years (C.E. = Common Era) with 95% confidence intervals.

The developed extraction and measurement methods were tested on stalagmite MAYA-22-7, collected in 2022 from Cenote Ch'en Mul at the Postclassic Maya archaeological site of Mayapan in northern Yucatan (Mexico), with permission granted by the Instituto Nacional de Antropología e Historia (INAH). The stalagmite was retrieved from a narrow passage in the southeast sector of the cave (Fig. S1). The growth surface of stalagmite MAYA-22-7 was wet from drip water prior to collection. The stalagmite was cut in half lengthwise, i.e. along the growth axis, and cut again to produce a central working slab and an off-axis working piece, using a Diamond WireTec DSW175 wire saw with a 0.35-mm-diameter diamond-studded steel wire at Northumbria University, UK. Powdered samples for biomarker analysis and U-Th dating were drilled at the University of Cambridge along exposed growth layers on the working half (Fig. 1), using a handheld Dremel drill with a 0.8-mm-diameter stungsten carbide dental drill bit. The bench surface, drill, drill bit, and aluminium tools for powder collection were all sprayed with compressed air and wiped down with a Kimwipe[™] and methanol between sample drilling. Sample pit vertical heights along the growth axis differed depending on the sample pit size, with a range of 3 to 8 mm (Fig. 1). The stalagmite surface was also wiped with a Kimwipe[™] and methanol before each pit was drilled. Powder sample weights ranged from 500 to

1200 mg and were stored in methanol-cleaned glass sample vials with a plastic screw cap.

- 100 The chronology of MAYA-22-7 was determined with a U-Th age model constructed from five U-Th ages drilled from the central working slab (Fig. 1). Sample preparation and analytical chemistry for dating were conducted at the Department of Earth Sciences, University of Oxford. Samples were dissolved in distilled concentrated nitric acid and spiked with a mixed ²²⁹Th-²³⁶U solution. U and Th were separated using column chemistry, following procedures adapted from Edwards et al. (1987). U and Th isotopes were measured using a Nu Plasma II Multi-ion-counting, Multi-collector Inductively Coupled
- 105 Plasma Mass Spectrometer (MC-ICP-MS), following procedures adapted from Hoffman (2008). Ages were calculated iteratively using the ²³⁰Th/²³⁸U age equation (Kaufman and Broecker, 1965), derived in Richards and Dorale (2003). The ²³⁴U and ²³⁰Th half-lives used for age calculations are those reported in Cheng et al. (2013). Age uncertainty was calculated using a Monte Carlo method, which accounts for instrument measurement, chemical blank, and initial detrital ²³⁰Th/²³²Th activity ratio uncertainty. The initial detrital ²³⁰Th/²³²Th activity ratio is set as a uniform distribution of values between 0.1 and 2.0,
- 110 which encompasses the bulk earth value (²³⁰Th/²³²Th activity = 0.82) and the median detritus value determined from a collection of speleothem studies (²³⁰Th/²³²Th activity = 1.5) (Hellstrom, 2006). All U-Th ages are in sequential order, with the corrected youngest date (9.3 mm from stalagmite top) 2001 ± 11 (2σ) CE. The interpolated age model for the stalagmite central growth axis was produced using the OxCal version 4.4 Poisson-process

Deposition model, with the stalagmite vertical growth rate constrained using model inputs $k_0 = 0.1 \text{ mm}^{-1}$ and $\log_{10}(k/k_0) = U(-2, 2)$ (Ramsey, 2008, 2009; Ramsey and Lee, 2013). Because sample pits in this study were drilled off the central axis, visible laminae were tracked to connect the sample pit locations to the central growth axis to determine the mean age range and 95% confidence age range of the sample pits. The OxCal Deposition model input code, an age-depth plot (Fig. S2), U-Th isotope activity ratios and calculated age of U-Th samples (Table S2), as well as ages and their respective errors of

120 2.2 PAH and MA extraction

the biomarker samples (Table S3) are found in the supplement.

The sample preparation procedure is illustrated in Fig. S3. <u>The PAH extraction method was adapted from Shahaoury et al.</u> (2018). <u>Powdered For PAH extraction, powdered</u>-speleothem samples were weighed into baked-out (>8 h at 450 °C) 20 mL glass vials and spiked with 100 μ L of a solution of 100 ng mL⁻¹ naphthalene-d₈, acenaphthene-d₁₀, and phenanthrene-d₁₀ in ethyl acetate (EA). Subsequently, each sample was extracted twice with 5 mL g⁻¹ dichloromethane (DCM) in an ultrasonic

- 125 bath for 45 min. After sonication, samples were allowed to stand for 10-20 minutes to improve phase separation. The supernatant was removed, filtered through a 1 μm glass fibre filter (Macherey Nagel), and loaded onto the preconditioned (3 mL each of EA and DCM) SPE cartridges. The flow-through was collected in baked-out 20 mL glass vials with tapered bottoms. The cartridge was rinsed with 3 mL each of DCM and EA and dried by blowing air through the cartridge. Subsequently, the solution was evaporated under a gentle stream of nitrogen at 30 °C to a volume of approximately 500 μL.
- 130 The walls of the vial were washed with 3 mL EA, the solution was once again evaporated to approximately 500 μ L, and the walls were washed again with another portion of 3 mL EA. Then the sample solution was evaporated to approximately 200 μ L, the walls were washed a last time with 500 μ L EA, and the solution was finally evaporated to a volume of ~100 μ L. This

solution was transferred into a <u>baked_baked_out</u> 250 μL vial. The evaporation vial was washed with 100 μL EA and the solution was added to the sample. To enable volume calculation and corrections during measurement, 2 μL of a solution of *p*-terphenyl (7.5 μg mL⁻¹ in EA) were added. The sample solution was stored in the freezer at -25 °C.

The extracted speleothem carbonate powder was dried in an oven at 50 °C overnight and then spiked with 100 μ L of a solution of 100 ng mL⁻¹ ¹³C₆-levoglucosan in acetonitrile (ACN). The extraction procedure is described in Homann et al. (2022). In brief, two 45 min ultrasonic extractions were performed using 5 mL g⁻¹ methanol (MeOH) as an extraction agent. The supernatant was filtered with a 1 μ m glass fibre fabric filter and evaporated to dryness under a gentle stream of nitrogen at

140 30 °C. The residue was redissolved in ACN/H₂O (95:5) and filtered through a 0.2 μm PA-filter (Altman-Analytik) prior to storage at -25 °C.

2.3 PAH analysis

The analysis was performed using a Thermo Fisher Scientific Orbitrap Exploris GC system. Analytes were separated on a TG-5-SILMS column (30 m, 0.25 mm inner diameter, 0.25 μm film thickness, Thermo Fisher Scientific). A volume of 1 μL was
injected in splitless mode at an injector temperature of 320 °C, a transfer-line temperature of 320 °C, and an oven temperature of 50 °C. The carrier-gas (helium, 5.0, Nippon Gases) flow was set to 1 mL min⁻¹. The initial temperature was held for 2 min, then increased to 160 °C at a rate of 10 °C min⁻¹ (1 min hold), then to 270 °C at 3 °C min⁻¹, then to 300 °C at 30 °C min⁻¹ (5 min hold), and finally to 320 °C at 30 °C min⁻¹ (2.7 min hold). The injector was cleaned after each injection for 5 min at 330 °C with a flow rate of 150 mL min⁻¹. The mass spectrometer was operated in positive electron ionisation mode (EI+) using
selected ion monitoring mode (SIM). The SIM *m/z* ratios and retention times of the respective PAHs are found in <u>Table 1</u><u>Table</u>
Analytes were quantified via the ratio of the peak area of the analyte to the peak area of the internal standard.

Table 1: Trivial-Common name, acronym, sum formula, retention time, m/z value, limit of detection, repeatability, and accuracy of all d-PAHs, internal standards, and PAHs determined by the GC-MS method.

	Analyte	Acronym	Sum	t _R (min)	$m/z [M]^{+}$	LOD	Repeatability	Accuracy
			formula			(ng)	(%)	(%)
Spiked d-PAHs and internal standard	Napthalene-d ₈	d-NAP	$C_{10}D_8$	9.49	136.1128	0.97	4.5	8.3
	Acenaphthene-d ₁₀	d-ACE	$C_{12}D_{10}$	13.67	164.1410^\dagger	1.37	4.6	1.5
	Phenanthrene-d ₁₀	d-PHE	$C_{14}D_{10}$	19.40	188.1410	1.29	11.4	2.6
	p-Terphenyl	PTP	$C_{18}H_{14}$	29.90	230.1096	/	6.6	/
Investigated PAHs	Naphthalene	NAP	$C_{10}H_8$	9.53	128.0626	1.41	5.7	0.6
	Acenaphthylene	ACY	$C_{12}H_8$	13.28	152.0626	0.07	7.0	0.2
	Acenaphthene	ACE	$C_{12}H_{10}$	13.77	154.0783	0.14	5.1	0.3
	Fluorene	FLN	$C_{13}H_{10}$	15.43	166.0783	0.17	25.6	0.1

Phenanthrene	PHE	$C_{14}H_{10}$	19.50	178.0783	2.07	17.6	4.8
Anthracene	ANT	$C_{14}H_{10}$	19.77	178.0783	0.04	3.3	0.1
Fluoranthene	FLT	$C_{16}H_{10}$	26.45	202.0783	0.67	6.2	1.0
Pyrene	PYR	$C_{16}H_{16}$	27.81	202.0777^\dagger	0.34	2.2	0.5
Retene	RET	$C_{18}H_{18}$	30.48	234.1409	0.36	19.4	0.7
Benzo(a)anthracene	BAA	$C_{18}H_{12}$	36.53	228.0939	1.09	3.7	1.2
Chrysene	CHR	$C_{18}H_{12}$	36.75	228.0939	1.14	8.9	0.9
Benzo(b)fluoranthene	BBF	$C_{2}0H_{12}$	44.02	252.0939	1.30	8.7	0.2
Benzo(a)pyrene	BAP	$C_{20}H_{14}$	45.99	252.0939 [†]	0.55	9.6	1.5
Indeno(1,2,3-c,d)pyrene	INP	C22H14	52.00	276.0939†	0.51	10.2	1.1
Dibenz(a,h)anthracene	DBA	C22H14	52.18	278.1096	NA	12.0	1.0
Benzo(g,h,i)perylene	DPE	C22H12	52.73	276.0939	0.47	12.4	1.5

155 [†]: Detection of fragment.

2.4 MA analysis

Analysis of MAs was carried out on a Dionex UltiMate 3000 ultrahigh-performance liquid chromatography system (UHPLC), coupled to a heated electrospray ionisation source (HESI) and a Q Exactive Orbitrap high-resolution mass spectrometer (HRMS) (all by Thermo Fisher Scientific) equipped with an iHILIC-Fusion column, 100 mm x 2.1 mm with 1.8 µm particle

- 160 size (Hilicon). The injection volume was 10 μL. A H₂O/ACN isocratic program was applied at a flow of 0.3 mL min⁻¹ with a run time of 5 min. The eluent composition was 97 % eluent B (100 % ACN) and 3 % eluent A (consisting of 5 mmol L⁻¹ ammonium formate in H₂O). To improve ionisation, a post-column flow of 50 mmol L⁻¹ NH₄OH in MeOH was applied with a flow rate of 0.1 mL min⁻¹. The HESI source was operated in negative mode to form deprotonated molecular ions [M-H]⁻. The HESI probe was heated to 150 °C, the capillary temperature was set to 350 °C, and the spray voltage was -4.0 kV. The
- 165 sheath gas pressure was 60 psi and the auxiliary gas pressure was 20 psi. The mass spectrometer was operated in full scan mode with a resolution of 70,000 and a scan range of *m/z* 80–500. During the expected retention times (<u>Table 2Table 2</u>), the full scan mode was alternated with a targeted MS² mode with a resolution of 17,500. For the MS² mode (i.e. parallel reaction monitoring mode in the software Xcalibur, provided by Thermo Fisher Scientific), higher-energy collisional dissociation (HCD) was used with 35% normalised collision energy (NCE).

170

Table 2: Trivial-Common name, acronym, sum formula, retention time, *m/z* value, limit of detection, repeatability, and accuracy of ¹³C-MAs and MAs determined in the MA LC-MS method.

Analyte	Acronym	Sum formula	t _R (min)	<i>m/z</i> [M-H] ⁻	LOD (ng)	Repeatability (%)	Accuracy (%)
13C6-levoglucosan	¹³ C-LEV	13C6H10O5	3.28	167.0657	NA	11.5	0.1

Mannosan	MAN	$C_6H_{10}O_5$	2.65	161.0455	0.05	4.4	0.3
				129.0193^\dagger			
Galactosan	GAL	C6H10O5	2.77	161.0455	0.01	11.0	0.2
				113.0245^\dagger			
Levoglucosan	LEV	C6H10O5	3.30	161.0455	0.09	6.4	0.1
				113.0245 [†]			
				101.0244^\dagger			

[†]: Fragments resulting from targeted MS².

3 Results and discussion

175 3.1 Method validation

Detection limits (LODs) were calculated using Eq. (1), where B is the blank and SD_B the corresponding standard deviation (Otto, 2014).

 $LOD = B + 3 SD_B , (1)$

Recovery was calculated by spiking samples with known amounts of naphthalene-d₈ (42-102 %), acenaphthene-d₁₀ (41-120 %), phenanthrene-d₁₀ (46-137 %), and ¹³C₆-levoglucosan (62-83 %). The comparatively low <u>PAH</u> recovery rates were likely a consequence of volatilisation during the evaporation step. A comprehensive overview of all recoveries is found in Table S3. The sample set data were not corrected for recovery to acknowledge that the three deuterated PAHs examined do not necessarily reflect the properties of all 16 analysed PAHs. A 25 ng mL⁻¹ (PAHs) or a 2.5 ng mL⁻¹ (MAs) standard was used to calculate the accuracy of the method (percent relative deviation with respect to the standard concentration). Repeatability
was calculated as the standard deviation of nine measurements of 25 ng mL⁻¹ (PAHs) or 2.5 ng mL⁻¹ (MAs) standards. Results are summarised in Table 17able 1 and 2.

3.2 Speleothem MAs and PAHs

Levoglucosan (LEV) and Σ15 (sum of non-alkylated PAHs) concentrations are presented in Fig. 2. Concentrations of individual MAs and PAHs are found in Table S4, whereas calculated sums and ratios are shown in <u>Table 3</u>Table 3. The LEV
and Σ15 concentrations range from 0.6 to 5.7 ng g⁻¹ and 3.8 to 16.9 ng g⁻¹, respectively. The Σ15 concentrations are greater than values presented by Argiriadis et al. (2019) but lower than those reported by Perrette et al. (2008). Σ15 concentrations and to a lesser extent, LEV concentrations, track the colour (grey value) of the stalagmite (Fig. 2Figure 2), suggesting a link between grey scale value and fire activity (soot).

Age (Year CE)*	2010- 2015	2000- 2010	1995- 2000	1945- 1985	1945- 1965	1920- 1935	1890- 1905	1855- 1875	1760- 1790	1640- 1650
Sample #	4	3	5	2	6	8	9	1	10	11
$\Sigma 15^{\dagger}$	6.44	6.2	9.95	14.7	15.4	3.8	11.9	6.7	10.3	5.4
LMW [‡]	6	5.6	9	13.8	14.5	3.7	11.3	6	9.7	5.04
HMW^{\Diamond}	0.4	0.5	0.9	0.9	0.9	0.02	0.6	0.7	0.5	0.3
LMW/HMW	13.6	10.4	10.33	14.9	15.8	175.5	19.3	8.5	18.9	14.7
PHE/ANT	74.7	4	43	3.4	NA	NA	NA	4.8	NA	NA
ANT/(ANT+PHE)	0.01	0.2	0.02	0.2	NA	NA	NA	0.2	NA	NA
RET/(RET+PHE+ANT)	NA	NA	0.16	NA	NA	NA	0.1	NA	NA	NA
$LEV/(LEV+\Sigma15)$	0.5	0.2	0.4	0.4	0.5	0.6	0.3	0.3	0.4	0.5
LEV/MAN	5.9	9.7	8	9.5	18.8	7.4	NA	9.1	8.8	6
LEV/(MAN+GAL)	3.3	4.7	4.6	4.8	5.9	3.7	18.6	5.4	4.7	3.1

195 Table 3: Calculated sums and diagnostic ratios of the 10 samples from stalagmite MAYA-22-7 from Cenote Ch'en Mul, Mayapan.

*: Start and end dates have errors of ± 6-20 years (95 % confidence interval) †: Sum of non-alkylated PAHs; ‡: Sum of two and three-ring

PAHs; ⁶: Sum of four and five-ring PAHs. Errors of sums and diagnostic ratios are reported in Table S5.



200 Figure 2: Concentrations of levoglucosan (dark blue) and Σ15 (sum of non-alkylated PAHs; orange) in the 10 analysed samples from stalagmite MAYA-22-7. Symbols are centred on the middle of their respective sample pit (Fig. 1). Error bars in the x-axis reflect the

Kommentiert [HJ1]: Updated Fig. 2 includes ages in green and explanation of grey value scale

lengths of the sample pits. Error bars on the y-axis represent one standard deviation and may be smaller than the symbols. The grey line shows the grey value of stalagmite MAYA-22-7, measured along the growth axis. <u>A value of 0 = black and 250 = white. Green 205 shading represent the width of the sample pits and their respective U-Th dates are given at the top.</u>

205 Overall, a strong correlation (r = 0.75, p < 0.05) between LEV and Σ15 is observed (Figure 3Fig. 3). This correlation changes only slightly (r=0.76, p < 0.05) when solely low-molecular-weight PAHs (LMW, sum of two- and three-ring PAHs) are considered. This is consistent with the findings of Battistel et al. (2017) and suggests a common origin for LEV and smaller PAHs. The three most abundant PAHs (phenanthrene (PHE), naphthalene (NAP), and fluorene (FLN)) constitute an average of 82 ±7 % of Σ15 in the analysed samples. However, PHE, the most abundant PAH, does not show a significant correlation
210 with LEV (r=0.60, p < 0.1, Figure 3Fig. 3, green circles). This is likely caused by different sources of PHE and LEV; it is possible that PHE is generated by higher-intensity fires than required to produce LEV.



Figure 3: Correlation of levoglucosan and PAHs in the 10 samples analysed from stalagmite MAYA-22-7. Blue squares: Σ15 (sum of non-alkylated PAHs); beige triangles: LMW (sum of two and three-ring PAHs); green circles: PHE. Error bars represent one standard deviation and may be smaller than the symbols.

Most high molecular weight PAHs (HMW, four- and five-ring PAHs) are below LOD in the analysed sample set. This is consistent with the results of Argiriadis et al. (2019) and Perrette et al. (2008) and is likely a consequence of a filtering by the overlying soil and epikarst, as discussed by Perrette et al. (2013). They found that HMW PAHs accumulate in soils and only LMW PAHs are transferred into the ground water under steady state conditions. No literature addresses the inclusion of airborne PAHs into speleothems or whether fractionation between HMW and LMW PAHs might be expected. However, it is plausible that LMW PAHs are more readily dissolved in the water-film that covers actively growing speleothems, than are HMW PAHs. This is supported by the respective n-octanol/water partition coefficient (*K*_{OW}) and air/water partition coefficient (*K*_{AW}) values, according to which, for instance, NAP is 1-2 orders of magnitude more likely to enter the aqueous phase than BAA (Lu et al., 2008; Rayne and Forest, 2016). HMW PAHs adsorbed to particles could also be incorporated into the growing speleothem. However, Dickson et al. (2023) demonstrated that microscopic particles are situated on the flanks of a stalagmite

rather than at the centre. As the samples analysed in this study were drilled along the growth axis, particulate PAHs would very likely not be encountered.

Ratio	Values	Interpretation	Reference	Medium applied to
LMW/HMW	< 1	Pyrogenic	Zhang et al. (2008)	Water
	> 1	Petrogenic		
PHE/ANT	< 10	Pyrogenic	Zhang et al. (2008)	Water
	> 15	Petrogenic		
ANT/(PHE+ANT)	< 0.1	Petrogenic	Yunker et al. (2002a)	River sediment
	> 0.1	Pyrogenic		
RET/(RET+CHR)	0.15-0.50	Petrol combustion	Yan et al. (2005)	Lake sediment
	0.30-0.45	Coal combustion		
	0.83-0.96	Softwood combustion		
RET/(RET+PHE+ANT)	> 0.1	Gymnosperm combustion	Karp et al. (2020)	Aerosols
	< 0.1	Angiosperm combustion		
$LEV/(LEV{+}\Sigma15)^{\dagger}$	< 0.5	High-intensity fires	Ruan et al. (2020)	Marine Sediment
	0.5	Boundary		
	> 0.5	Low-intensity fires		
LEV/MAN	0.5-14	Softwoods	Fabbri et al. (2009) and	Aerosols; Chars
	3-32	Hardwoods	references therein; Kuo	
	4-55	Grasses	et al. (2011)	
	30-90	Lignite		
LEV/(MAN+GAL)	0.5-6	Softwoods	Fabbri et al. (2009) and	Aerosols; Chars
	1.5-10	Hardwoods	references therein; Kuo	
	4-55	Grasses	et al. (2011)	
	30-90	Lignite		

Table 4: Typically reported values of diagnostic ratios of PAHs and MAs, and respective interpretations.

230 \dagger : LEV and Σ 15 are normalised to the highest value of the series.

The lack of HMW PAHs means that some PAH ratios normally used to characterise PAH profiles (<u>Table 4</u>Table 4) may not apply to speleothems. An example of such a ratio is LMW/HMW; for directly deposited samples such as aerosols or sediments, a ratio > 1 indicates a petrogenic origin of PAHs (Soclo et al., 2000; Zhang et al., 2008). The artificially lowered HMW levels
235 observed in speleothems results in very high LMW/HMW ratios (<u>Table 3</u>Table 3), rendering this ratio insensitive to the origin of the PAHs. More promising diagnostic metrics in speleothem analysis could be individual PAHs. For this purpose, we studied the ratios of two or more compounds with similar molecular formulas but different molecular structures, assuming that the

compounds have comparable physicochemical properties (i.e. volatility, solubility, atmospheric lifetime, adsorption, and transport mechanisms) but different predominant origins (Yunker et al., 2002b). At high temperatures, a less stable, "kinetic"

- 240 isomer is formed, which has more aromatic rings and less alkyl substitutes compared to the product at lower combustion temperatures or of petrogenic origin (Han et al., 2020; McGrath et al., 2003; Tobiszewski and Namieśnik, 2012). Phenanthrene (PHE) and anthracene (ANT) form one pair, where PHE is the thermodynamic and ANT the kinetic isomer (Soclo et al., 2000). The resulting PHE/ANT and ANT/(ANT+PHE) ratios can also be used to distinguish between pyrogenic (PHE/ANT < 10; ANT/(ANT+PHE) > 0.1) and petrogenic (> 10; < 0.1) origins of the PAHs and appear to apply to our stalagmite samples. The</p>
- 245 PHE-ANT ratios of samples #1-3 indicate pyrogenic origin; for the remainder of the samples, ANT is very close to or below the LOD. Therefore, we cannot conclude with certainty a petrogenic origin of the PAHs in samples #4-11. Retene (RET) is an alkylated PAH and a molecular marker of gymnosperm combustion (Muri et al., 2003; Ramdahl, 1983;

Wakeham et al., 1980). Unfortunately, RET was below the LOD for all but two samples; nevertheless, RET might provide a promising marker of fire dynamics in speleothems if it is present in sufficient concentrations. Literature suggests that

- 250 gymnosperm vegetation is not common in the study area (Contreras-Medina et al., 2007; Douglas et al., 2012; Etnoflora Yucatanense, 1994; Tellez et al., 2020). Nevertheless, RET might provide a promising marker of fire dynamics in speleothems if it is present in sufficient concentrations. RET is also used in various diagnostic ratios. For instance, in combination with chrysene (CHR), where RET/(RET + CHR) helps to determine fuel type (Yan et al., 2005). Because CHR is one of the HMW PAHs, it is below the LOD for all samples in this study, but this ratio could potentially be utilised in other samples. In addition,
- Karp et al. (2020) suggested the use of a ratio that includes three 3-ring PAHs, RET, PHE, and ANT, where RET/(RET+PHE+ANT) is used to distinguish between gymnosperm (softwood, > 0.1) and angiosperm (hardwood, < 0.1) combustion. As mentioned above, in our sample set, RET is above the LOD in only two samples. For one of these (1995-2000 CE) the ratio points towards gymnosperm fuel₁ whereas the result for the other sample (1890-1905 CE) falls right on the boundary between angiosperm and gymnosperm fuel. This ratio could be particularly useful in combination with the MAs,
- 260 which can also be used to determine fuel type.



265

Pigne 4. (a) Lev MAA' (dark blue) and LEV (MAA' $(3AL)^{-1}$ (b) and SAL) (b) and SAL in the barnet of sample analysed rioh statighter MAA' A 22-7. Coloured bars on the right indicate the burned fuel based on the LEV (MAA' $(3AL)^{-1}$ ratio according to Fabbri et al. (2009) and Kuo et al. (2011). Green: grasses; grey: hardwoods; blue: softwoods. Age range listed is the OxCal modelled mean age at the top and bottom of each sample pit. Start and end dates have errors of \pm 6-20 years (95 % confidence interval; see Table S2). Error bars represent one standard deviation. (b) LEV (LEV $\pm \Sigma 15$)⁻¹ ratios of the 10 samples analysed. The dashed line represents the boundary between MA-dominated fire regimes (low intensity) and PAH-dominated fire regimes (high intensity). Age range listed is the OxCal modelled mean age at the top and bottom of each sample pit. Start and end dates have errors of \pm 6-20 years (95 % confidence interval; see Table S2). Error bars represent one standard deviation.

- MA ratios, LEV MAN⁻¹ and LEV (MAN + GAL)⁻¹, respectively, have been shown to capture the type of biomass burned (Fabbri et al., 2009 and references therein; Kuo et al., 2011). According to Fabbri et al. (2009), the use of LEV (MAN + GAL)⁻¹ is even more sensitive and discriminating. The ratios in our study are shown in Figure 4Fig. 4 (a) and indicate combustion of predominantly mixed hardwoods and softwoods. Kuo et al. (2011), however, showed that high combustion temperatures may shift the MA ratios to higher values because hemicellulose is less thermally stable than cellulose. This means that MAN and
- 275 GAL are not emitted at higher temperatures, whereas LEV can still be formed. This could be the case for our sample from

Kommentiert [HJ2]: Updated Fig. 4 (b) does not have a reversed y-Axis anymore.

1890-1905 CE (light orange in Fig. 4 (a)), in which no MAN could be detected and the resulting LEV (MAN + GAL)⁻¹ is very high. Therefore, it might be useful to combine the MA ratios with an indicator of fire intensity. Ruan et al. (2020) suggested the use of LEV (LEV + $\Sigma 15$)⁻¹, with both LEV and $\Sigma 15$ normalised to the highest value in the series. Values > 0.5 indicate a predominance of PAHs and high fire intensity, whereas values < 0.5 indicate low intensity fires with dominant MA emission. Fig. 4 (b) shows two distinct peaks in fire intensity (i.e. low LEV contribution) in the 1855-1875 CE and 2000-2010 CE samples. The 1890-1905 CE sample mentioned above also shows high-high-intensity fires, supporting the interpretation that the LEV (MAN + GAL)⁻¹ ratio may have been artificially increased. The 1920-1935 CE sample is the only one that clearly indicates low fire intensity, whereas the remaining samples fall into the high-fire-intensity fire regime. We note however, the relatively large analytical uncertainties make a definitive interpretation difficult.

285 **3.3 Lev LEV** and PAHs as indicators of fire activity at Cenote Ch'en Mul

280

We applied our results to infer past fire activity in and around Cenote Ch'en Mul. Mayapan was the last capital of the Late Postclassic Maya period and was abandoned in the mid-15th century (Kennett et al., 2022; Milbrath and Peraza Lope, 2003). Its ruins, including the round temple and the Kukulcan temple, were first reported to the English-English-speaking world in 1841 by John Stephens. Although preliminary excavations were conducted in the early 20th century, major archaeological work

- 290 only took place in the 1950s under the auspices the Carnegie Institution (Milbrath and Peraza Lope, 2003). These archaeological activities are relevant to the present study, because human-induced fires related to the excavations (land-clearing, use of kerosene lamps or open fires, etc.) produced LEV and PAH that should be recorded in our dataset. Activities like water collection or ritual incense burning have a very long history in Yucatan and are still conducted to this day. Any open flame combustion thus adds to the overall PAH and MA load in the cave, and potentially in speleothems. The signal observed
- 295 in a stalagmite represents the integral of all accumulated PAHs and MAs, and in-depth research is required to differentiate the relative contribution of each pathway.

Stalagmite MAYA-22-7 grew fast (growth rate of 200-300 μ m/year before the dark layer at a distance of 15-23 mm from the top, and 500-600 μ m/year after that dark layer), and covers only the last ~400 years, i.e. the period after the site was abandoned by the Postclassic Maya. The fire history inferred from the stalagmite is likely related to both natural fires in the low-stature,

- 300 thorny, deciduous tropical forest that characterised the area, and land-clearing activities that preceeded archaeological excavation at the site. Early and middle 20th century archaeological efforts likely used kerosene lamps for work in Cenote Ch'en Mul. Parts of the Mayapan site might have been cleared of vegetation using fire, although it is difficult to find evidence for this practice in archaeological field reports. The Carnegie excavations might have used both kerosene and battery powered lights. Either way, it is plausible that LEV and PAHs from human-induced fires might be recorded in the stalagmite.
- 305 This is in accordance with our findings, in that the samples covering the period 1945-1985 show the highest LEV and PAH concentrations in the whole sample set (Fig. 2). Finding elevated PAH and MA concentrations in speleothem carbonate that dates to a time of higher human-induced fire activity is encouraging and suggests that speleothems are sensitive recorders of combustion products. Further work should be directed towards i) deciphering transport mechanisms and potential scavenging

effects in the soil and epikarst, ii) detailed understanding of incorporation pathways into carbonate, and iii) refinement of 310 methods to delineate dripwater and aerosol derived fire proxies.

4 Conclusions and outlook

We used a sequential extraction preparation method for analysis of 16 PAHs and three MAs in samples from a stalagmite collected at the Postclassic Maya archaeological site of Mayapan, Yucatán, Mexico. Sample preparation involved solid-liquid extraction of pulverised carbonate samples. To our knowledge, this is one of only a few methods for analysis of PAHs in

315 speleothems and the only one that also enables analysis of MAs. The new method requires relatively small sample amounts (0.5-1.0 g) and simple instrumentation, i.e. an ultrasonic bath and a sample concentrator. PAHs and MAs were measured by GC-MS and LC-HILIC-MS, respectively.

We applied the method to 10 samples from the MAYA-22-7 stalagmite, collected at Cenote Ch'en Mul, directly below the site of Mayapan. We used concentrations and selected ratios of PAHs and MAs described in the literature to explore fire history in this area of the Yucatán Peninsula. We found a positive correlation between the major MA, levoglucosan, and non-alkylated

PAHs (Σ15), or two- and three-ring PAHs (LMW), (r = 0.75, p < 0.05, and r = 0.76, p < 0.05, respectively). Not all diagnostic PAH ratios were applicable, but the ratios combining PAHs with MAs appear to have promise for discriminating among different fire regimes and inferring the source of burned fuel. Furthermore, retene (RET) can be used as a molecular marker for gymnosperm combustion, if present in high enough concentrations. <u>However, it has to be noted that the HMW PAHs</u>
were often below LOD, probably because of a filtering effect of the overlying soil. This limited our ability to fully exploit the data set to interpret past fire dynamics at Mayapan.

data set to interpret past fire dynamics at Mayapan.

Our analyses suggest that within the analysed time series, the period 1945-1985 stands out in terms of high fire activity. We tentatively interpret this result to indicate clearing of the site prior to and during mid-20th-century archaeological excavation. A longer record of the two biochemical proxies from older stalagmites from Che'Mul Cave could document the history of

330 cave use by the Maya for ceremonial and practical purposes - burning undoubtedly occurred within the cave (e.g. torches or fire for lighting).

5 Future Work

320

Future work should be directed towards i) refinement of methods to delineate dripwater- and aerosol-derived fire proxies, ii) deciphering transport mechanisms and potential scavenging effects in the soil and epikarst, and iii) interplay with other proxies.
 The PAH extraction method should be improved to optimise recovery and reduce its variance. One possible approach is to perform a systematic solvent screening to improve the extraction efficiency. Alternatively, the extraction method could be modified, e.g. by utilisation of microwave extraction or accelerated solvent extraction. Another route is the addition of a keeper

(e.g. nonane) during evaporation, to reduce losses to volatilisation, similar to the method described by Wietzoreck et al. (2022). To correct the data for recovery, additional deuterated PAH standards could be added.

- 340 In addition, monitoring rainfall and dripwater at the sample site would be useful to track changes in PAH and MA patterns and to gain a deeper understanding of site-specific transport mechanisms. It would also be beneficial to apply our method to other speleothems from the same cave and other sites, to test whether our results are generally applicable. For future work, thethe two biochemical proxies studied here should be combined with other, established proxies such as carbon and oxygen isotope ratios as well as trace elements described by Campbell et al. (2023)_s to gain further insights into the interplay among fire,
- 345 hydrology, vegetation, and human activities. In addition, monitoring rainfall and dripwater at the sample site would be useful to track changes in PAH and MA patterns and to gain a deeper understanding of site-specific transport mechanisms. It would also be beneficial to apply our method to other speleothems from the same cave and other sites, to test whether our results are generally applicable. Finally, other source-specific PAH species could be added to enhance the interpretation of the PAH patterns. Potentially suitable PAHs include 1,2- and 1,7-dimethylphenanthrene (1,2- and 1,7-DMP). Kappenberg et al. (2019)
- 350 suggested that their ratio could be used as an alternative to distinguish among different types of burned biomass.

Data availability. All data from our study are presented numerically in the paper and in the Supplement.

- Author contributions. JH and TH designed the study. JH and NK conducted the method development, sample preparation, and data analysis for PAHs and MAs. CL and MB enabled sampling in Cenote Ch'en Mul. DJ, OK, DH, and SB collected stalagmite MAYA-22-7. OK and SB surveyed Cenote Ch'en Mul and prepared the cave map. SC and DJ subsampled the stalagmite for analysis and constructed the age model. All co-authors participated in discussions, interpretation of the data, and writing of the manuscript.
- 360 **Competing interests.** The contact author has declared that none of the authors has any competing interests.
- Acknowledgements. The authors thank <u>Drs.</u> Susan Milbrath and <u>Marylin Marilyn</u> Masson for insights <u>oninto</u> the history of the archaeological excavations. JH gratefully acknowledges financial support from the Max Planck Graduate Center Mainz. Some of the analytical developments included here were developed with financial support from DFG Project HO 1748/20-1
- 365 (Organic trace analysis of atmospheric marker substances in ice cores). DH acknowledges financial support from the Leverhulme Trust Award RPG-2019-228.

References

Argiriadis, E., Battistel, D., McWethy, D. B., Vecchiato, M., Kirchgeorg, T., Kehrwald, N. M., Whitlock, C., Wilmshurst, J.

- 370 M., and Barbante, C.: Lake sediment fecal and biomass burning biomarkers provide direct evidence for prehistoric human-lit fires in New Zealand, Sci Rep, 8, 12113, https://doi.org/10.1038/s41598-018-30606-3, 2018.
 - Argiriadis, E., Denniston, R. F., and Barbante, C.: Improved Polycyclic Aromatic Hydrocarbon and n-Alkane Determination in Speleothems through Cleanroom Sample Processing, Anal. Chem., 91, 7007–7011, https://doi.org/10.1021/acs.analchem.9b00767, 2019.
- 375 Bai, J., Sun, X., Zhang, C., Xu, Y., and Qi, C.: The OH-initiated atmospheric reaction mechanism and kinetics for levoglucosan emitted in biomass burning, Chemosphere, 93, 2004–2010, https://doi.org/10.1016/j.chemosphere.2013.07.021, 2013.
 - Baker, A., Blyth, A. J., Jex, C. N., Mcdonald, J. A., Woltering, M., and Khan, S. J.: Glycerol dialkyl glycerol tetraethers (GDGT) distributions from soil to cave: Refining the speleothem paleothermometer, Organic Geochemistry, 136,
- 380 103890, https://doi.org/10.1016/j.orggeochem.2019.06.011, 2019.
- Battistel, D., Argiriadis, E., Kehrwald, N., Spigariol, M., Russell, J. M., and Barbante, C.: Fire and human record at Lake Victoria, East Africa, during the Early Iron Age: Did humans or climate cause massive ecosystem changes?, The Holocene, 27, 997–1007, https://doi.org/10.1177/0959683616678466, 2017.
- Blyth, A. J., Hartland, A., and Baker, A.: Organic proxies in speleothems New developments, advantages and limitations,
- 385 Quaternary Science Reviews, 149, 1–17, https://doi.org/10.1016/j.quascirev.2016.07.001, 2016.
- Bosle, J. M., Mischel, S. A., Schulze, A.-L., Scholz, D., and Hoffmann, T.: Quantification of low molecular weight fatty acids in cave drip water and speleothems using HPLC-ESI-IT/MS -- development and validation of a selective method, Analytical and bioanalytical chemistry, 406, 3167–3177, https://doi.org/10.1007/s00216-014-7743-6, 2014.

Braun, T., Breitenbach, S. F. M., Skiba, V., Lechleitner, F. A., Ray, E. E., Baldini, L. M., Polyak, V. J., Baldini, J. U. L.,

- 390 Kennett, D. J., Prufer, K. M., and Marwan, N.: Decline in seasonal predictability potentially destabilized Classic Maya societies, Commun Earth Environ, 4, 1–12, https://doi.org/10.1038/s43247-023-00717-5, 2023.
 - Callegaro, A., Battistel, D., Kehrwald, N. M., Matsubara Pereira, F., Kirchgeorg, T., Del Villoslada Hidalgo, M. C., Bird, B. W., and Barbante, C.: Fire, vegetation, and Holocene climate in a southeastern Tibetan lake: a multi-biomarker reconstruction from Paru Co, Clim. Past, 14, 1543–1563, https://doi.org/10.5194/cp-14-1543-2018, 2018.
- 395 Campbell, M., McDonough, L., Treble, P. C., Baker, A., Kosarac, N., Coleborn, K., Wynn, P. M., and Schmitt, A. K.: A Review of Speleothems as Archives for Paleofire Proxies, With Australian Case Studies, Reviews of Geophysics, 61, https://doi.org/10.1029/2022RG000790, 2023.
 - Cheng, H., Lawrence Edwards, R., Shen, C.-C., Polyak, V. J., Asmerom, Y., Woodhead, J., Hellstrom, J., Wang, Y., Kong, X., Spötl, C., Wang, X., and Calvin Alexander, E.: Improvements in 230Th dating, 230Th and 234U half-life values, and

- 400 U–Th isotopic measurements by multi-collector inductively coupled plasma mass spectrometry, Earth and Planetary Science Letters, 371-372, 82–91, https://doi.org/10.1016/j.epsl.2013.04.006, 2013.
 - Contreras-Medina, R., Vega, I. L., and Morrone, J. J.: Gymnosperms and cladistic biogeography of the Mexican Transition Zone, Taxon, 56, 905–916, https://doi.org/10.2307/25065872, 2007.
 - Denis, E. H., Toney, J. L., Tarozo, R., Scott Anderson, R., Roach, L. D., and Huang, Y.: Polycyclic aromatic hydrocarbons
- 405 (PAHs) in lake sediments record historic fire events: Validation using HPLC-fluorescence detection, Organic Geochemistry, 45, 7–17, https://doi.org/10.1016/j.orggeochem.2012.01.005, 2012.
 - Dickson, B., Sniderman, J. K., Korasidis, V. A., and Woodhead, J.: The distribution of fossil pollen and charcoal in stalagmites, Quat. res., 1–13, https://doi.org/10.1017/qua.2023.11, 2023.
 - Douglas, P. M., Pagani, M., Brenner, M., Hodell, D. A., and Curtis, J. H.: Aridity and vegetation composition are important
- 410 determinants of leaf-wax δD values in southeastern Mexico and Central America, Geochimica et Cosmochimica Acta,
 97, 24–45, https://doi.org/10.1016/j.gca.2012.09.005, 2012.
 - Edwards, R. L., Chen, J. H., Ku, T. L., and Wasserburg, G. J.: Precise timing of the last interglacial period from mass spectrometric determination of thorium-230 in corals, Science (New York, N.Y.), 236, 1547–1553, https://doi.org/10.1126/science.236.4808.1547, 1987.
- 415 Elias, V. O., Simoneit, B. R., Cordeiro, R. C., and Turcq, B.: Evaluating levoglucosan as an indicator of biomass burning in Carajás, amazônia: a comparison to the charcoal record, Geochimica et Cosmochimica Acta, 65, 267–272, https://doi.org/10.1016/S0016-7037(00)00522-6, 2001.

Etnoflora Yucatanense, Xalapa, Veracruz, México, 169 pp., 1994.

Fabbri, D., Torri, C., Simoneit, B. R., Marynowski, L., Rushdi, A. I., and Fabiańska, M. J.: Levoglucosan and other cellulose

420 and lignin markers in emissions from burning of Miocene lignites, Atmospheric Environment, 43, 2286–2295, https://doi.org/10.1016/j.atmosenv.2009.01.030, 2009.

Fairchild, I. J. and Baker, A.: Speleothem Science, John Wiley & Sons, Ltd, Chichester, UK, 2012.

- Fraser, M. P. and Lakshmanan, K.: Using Levoglucosan as a Molecular Marker for the Long-Range Transport of Biomass Combustion Aerosols, Environ. Sci. Technol., 34, 4560–4564, https://doi.org/10.1021/es9912291, 2000.
- 425 Gałuszka, A., Migaszewski, Z. M., and Namieśnik, J.: The role of analytical chemistry in the study of the Anthropocene, TrAC Trends in Analytical Chemistry, 97, 146–152, https://doi.org/10.1016/j.trac.2017.08.017, 2017.
 - Han, Y., Chen, Y., Feng, Y., Song, W., Cao, F., Zhang, Y., Li, Q., Yang, X., and Chen, J.: Different formation mechanisms of PAH during wood and coal combustion under different temperatures, Atmospheric Environment, 222, 117084, https://doi.org/10.1016/j.atmosenv.2019.117084, 2020.
- 430 Heidke, I., Scholz, D., and Hoffmann, T.: Lignin oxidation products as a potential proxy for vegetation and environmental changes in speleothems and cave drip water – a first record from the Herbstlabyrinth, central Germany, Clim. Past, 15, 1025–1037, https://doi.org/10.5194/cp-15-1025-2019, 2019.

Hellstrom, J.: U-Th dating of speleothems with high initial 230Th using stratigraphical constraint, Quaternary Geochronology, 1, 289-295, https://doi.org/10.1016/j.quageo.2007.01.004, 2006.

- 435 Hoffmann, D. L.: 230Th isotope measurements of femtogram quantities for U-series dating using multi ion counting (MIC) MC-ICPMS, International Journal of Mass Spectrometry, 275, 75-79, https://doi.org/10.1016/j.ijms.2008.05.033, 2008. Homann, J., Oster, J. L., Wet, C. B. de, Breitenbach, S. F. M., and Hoffmann, T.: Linked fire activity and climate whiplash in California during the early Holocene, Nat Commun, 13, 1-9, https://doi.org/10.1038/s41467-022-34950-x, 2022. Kappenberg, A., Braun, M., Amelung, W., and Lehndorff, E.: Fire condensates and charcoals: Chemical composition and
- 440
- fuel source identification, Organic Geochemistry, 130, 43-50, https://doi.org/10.1016/j.orggeochem.2019.01.009, 2019. Karp, A. T., Holman, A. I., Hopper, P., Grice, K., and Freeman, K. H.: Fire distinguishers: Refined interpretations of polycyclic aromatic hydrocarbons for paleo-applications, Geochimica et Cosmochimica Acta, 289, 93-113, https://doi.org/10.1016/j.gca.2020.08.024, 2020.
- Kaufman, A. and Broecker, W.: Comparison of Th 230 and C 14 ages for carbonate materials from lakes Lahontan and Bonneville, J. Geophys, Res., 70, 4039-4054, https://doi.org/10.1029/JZ070i016p04039, 1965. 445
- Kennett, D. J., Masson, M., Lope, C. P., Serafin, S., George, R. J., Spencer, T. C., Hoggarth, J. A., Culleton, B. J., Harper, T. K., Prufer, K. M., Milbrath, S., Russell, B. W., González, E. U., McCool, W. C., Aquino, V. V., Paris, E. H., Curtis, J. H., Marwan, N., Zhang, M., Asmerom, Y., Polyak, V. J., Carolin, S. A., James, D. H., Mason, A. J., Henderson, G. M., Brenner, M., Baldini, J. U. L., Breitenbach, S. F. M., and Hodell, D. A.: Drought-Induced Civil Conflict Among the
- 450 Ancient Maya, Nat Commun, 13, 3911, https://doi.org/10.1038/s41467-022-31522-x, 2022.
- Kuo, L.-J., Louchouarn, P., and Herbert, B. E.: Influence of combustion conditions on yields of solvent-extractable anhydrosugars and lignin phenols in chars: implications for characterizations of biomass combustion residues, Chemosphere, 85, 797-805, https://doi.org/10.1016/j.chemosphere.2011.06.074, 2011.
- Lammel, G., Sehili, A. M., Bond, T. C., Feichter, J., and Grassl, H.: Gas/particle partitioning and global distribution of
- 455 polycyclic aromatic hydrocarbons -- a modelling approach, Chemosphere, 76, 98-106, https://doi.org/10.1016/j.chemosphere.2009.02.017, 2009.
 - Lu, G.-N., Tao, X.-Q., Dang, Z., Yi, X.-Y., and Yang, C.: Estimation of n-octanol/water partition coefficients of polycyclic aromatic hydrocarbons by quantum chemical descriptors, Open Chemistry, 6, 310-318, https://doi.org/10.2478/s11532-008-0010-y, 2008.
- 460 Luo, J., Han, Y., Zhao, Y., Huang, Y., Liu, X., Tao, S., Liu, J., Huang, T., Wang, L., Chen, K., and Ma, J.: Effect of northern boreal forest fires on PAH fluctuations across the arctic, Environmental pollution (Barking, Essex 1987), 261, 114186, https://doi.org/10.1016/j.envpol.2020.114186, 2020.
- Mason, A. J., Vaks, A., Breitenbach, S. F. M., Hooker, J. N., and Henderson, G. M.: A simplified isotope dilution approach for the U-Pb dating of speleogenic and other low-232Th carbonates by multi-collector ICP-MS, Geochronology, 4, 33-54, https://doi.org/10.5194/gchron-4-33-2022, 2022. 465

McGrath, T. E., Chan, W., and Hajaligol, M. R.: Low temperature mechanism for the formation of polycyclic aromatic hydrocarbons from the pyrolysis of cellulose, Journal of Analytical and Applied Pyrolysis, 66, 51–70, https://doi.org/10.1016/S0165-2370(02)00105-5, 2003.

Milbrath, S. and Peraza Lope, C.: REVISITING MAYAPAN: Mexico's last Maya capital, Ancient Mesoam, 14, 1-46,

470 https://doi.org/10.1017/S0956536103132178, 2003.

Muri, G., Wakeham, S. G., and Faganeli, J.: Polycyclic aromatic hydrocarbons and black carbon in sediments of a remote alpine Lake (Lake Planina, northwest Slovenia), Environ Toxicol Chem, 22, 1009–1016, https://doi.org/10.1002/etc.5620220508, 2003.

Otto, M.: Analytische Chemie, 4., überarb. und erg. Aufl., 1. Nachdruck, Bachelor, Wiley-VCH, Weinheim, 674 pp., 2014.

475 Perrette, Y., Poulenard, J., Durand, A., Quiers, M., Malet, E., Fanget, B., and Naffrechoux, E.: Atmospheric sources and soil filtering of PAH content in karst seepage waters, Organic Geochemistry, 65, 37–45, https://doi.org/10.1016/j.orggeochem.2013.10.005, 2013.

Perrette, Y., Poulenard, J., Saber, A.-I., Fanget, B., Guittonneau, S., Ghaleb, B., and Garaudee, S.: Polycyclic Aromatic Hydrocarbons in stalagmites: Occurrence and use for analyzing past environments, Chemical Geology, 251, 67–76,

480 https://doi.org/10.1016/j.chemgeo.2008.02.013, 2008.

- Ramdahl, T.: Retene—a molecular marker of wood combustion in ambient air, Nature, 306, 580–582, https://doi.org/10.1038/306580a0, 1983.
- Ramsey, C. B.: Bayesian Analysis of Radiocarbon Dates, Radiocarbon, 51, 337–360, https://doi.org/10.1017/S0033822200033865, 2009.
- 485 Ramsey, C. B.: Deposition models for chronological records, Quaternary Science Reviews, 27, 42–60, https://doi.org/10.1016/j.quascirev.2007.01.019, 2008.

Ramsey, C. B. and Lee, S.: Recent and Planned Developments of the Program OxCal, Radiocarbon, 55, 720–730, https://doi.org/10.1017/S0033822200057878, 2013.

Rayne, S. and Forest, K.: Air-water partition coefficients for a suite of polycyclic aromatic and other C10 through C20

- 490 unsaturated hydrocarbons, Journal of environmental science and health. Part A, Toxic/hazardous substances & environmental engineering, 51, 938–953, https://doi.org/10.1080/10934529.2016.1191812, 2016.
 - Richards, D. A. and Dorale, J. A.: Uranium-series Chronology and Environmental Applications of Speleothems, Reviews in Mineralogy and Geochemistry, 52, 407–460, https://doi.org/10.2113/0520407, 2003.

Ridley, H. E., Asmerom, Y., Baldini, J. U. L., Breitenbach, S. F. M., Aquino, V. V., Prufer, K. M., Culleton, B. J., Polyak,

495 V., Lechleitner, F. A., Kennett, D. J., Zhang, M., Marwan, N., Macpherson, C. G., Baldini, L. M., Xiao, T., Peterkin, J. L., Awe, J., and Haug, G. H.: Aerosol forcing of the position of the intertropical convergence zone since ad 1550, Nat. Geosci., 8, 195–200, https://doi.org/10.1038/ngeo2353, 2015.

- Ruan, Y., Mohtadi, M., Dupont, L. M., Hebbeln, D., Kaars, S., Hopmans, E. C., Schouten, S., Hyer, E. J., and Schefuß, E.: Interaction of Fire, Vegetation, and Climate in Tropical Ecosystems: A Multiproxy Study Over the Past 22,000 Years,
- 500 Global Biogeochem. Cycles, 34, https://doi.org/10.1029/2020GB006677, 2020.
 - Scholz, D. and Hoffmann, D.: 230Th/U-dating of fossil corals and speleothems, Quaternary Science Journal, 52–76, https://doi.org/10.23689/fidgeo-1056, 2008.
 - Simoneit, B. R.: Biomass burning a review of organic tracers for smoke from incomplete combustion, Applied Geochemistry, 17, 129–162, https://doi.org/10.1016/S0883-2927(01)00061-0, 2002.
- 505 Slade, J. H. and Knopf, D. A.: Heterogeneous OH oxidation of biomass burning organic aerosol surrogate compounds: assessment of volatilisation products and the role of OH concentration on the reactive uptake kinetics, Physical chemistry chemical physics PCCP, 15, 5898–5915, https://doi.org/10.1039/c3cp44695f, 2013.
 - Soclo, H., Garrigues, P., and Ewald, M.: Origin of Polycyclic Aromatic Hydrocarbons (PAHs) in Coastal Marine Sediments: Case Studies in Cotonou (Benin) and Aquitaine (France) Areas, Marine Pollution Bulletin, 40, 387–396,
- 510 https://doi.org/10.1016/S0025-326X(99)00200-3, 2000.
 Tellez, O., Mattana, E., Diazgranados, M., Kühn, N., Castillo-Lorenzo, E., Lira, R., Montes-Leyva, L., Rodriguez, I., Flores
 Ortiz, C. M., Way, M., Dávila, P., and Ulian, T.: Native trees of Mexico: diversity, distribution, uses and conservation,
 PeerJ, 8, e9898, https://doi.org/10.7717/peerj.98988, 2020.
 - Tobiszewski, M. and Namieśnik, J.: PAH diagnostic ratios for the identification of pollution emission sources,
- 515 Environmental pollution (Barking, Essex 1987), 162, 110–119, https://doi.org/10.1016/j.envpol.2011.10.025, 2012.
 Wakeham, S. G., Schaffner, C., and Giger, W.: Poly cyclic aromatic hydrocarbons in Recent lake sediments—II.
 Compounds derived from biogenic precursors during early diagenesis, Geochimica et Cosmochimica Acta, 44, 415–429, https://doi.org/10.1016/0016-7037(80)90041-1, 1980.
 - Wietzoreck, M., Bandowe, B. A. M., Hofman, J., Martiník, J., Nežiková, B., Kukučka, P., Přibylová, P., and Lammel, G.:
- 520 Nitro- and oxy-PAHs in grassland soils from decade-long sampling in central Europe, Environmental geochemistry and health, 44, 2743–2765, https://doi.org/10.1007/s10653-021-01066-y, 2022.
 - Xie, M., Hannigan, M. P., and Barsanti, K. C.: Gas/particle partitioning of 2-methyltetrols and levoglucosan at an urban site in Denver, Environ. Sci. Technol., 48, 2835–2842, https://doi.org/10.1021/es405356n, 2014.
 - Yan, B., Abrajano, T. A., Bopp, R. F., Chaky, D. A., Benedict, L. A., and Chillrud, S. N.: Molecular tracers of saturated and
- 525 polycyclic aromatic hydrocarbon inputs into Central Park Lake, New York City, Environ. Sci. Technol., 39, 7012–7019, https://doi.org/10.1021/es0506105, 2005.
 - Yunker, M. B., Backus, S. M., Graf Pannatier, E., Jeffries, D. S., and Macdonald, R. W.: Sources and Significance of Alkane and PAH Hydrocarbons in Canadian Arctic Rivers, Estuarine, Coastal and Shelf Science, 55, 1–31, https://doi.org/10.1006/ecss.2001.0880, 2002a.

530 Yunker, M. B., Macdonald, R. W., Vingarzan, R., Mitchell, R. H., Goyette, D., and Sylvestre, S.: PAHs in the Fraser River basin: a critical appraisal of PAH ratios as indicators of PAH source and composition, Organic Geochemistry, 33, 489– 515, https://doi.org/10.1016/S0146-6380(02)00002-5, 2002b.

Zennaro, P., Kehrwald, N., McConnell, J. R., Schüpbach, S., Maselli, O. J., Marlon, J., Vallelonga, P., Leuenberger, D., Zangrando, R., Spolaor, A., Borrotti, M., Barbaro, E., Gambaro, A., and Barbante, C.: Fire in ice: two millennia of

535 boreal forest fire history from the Greenland NEEM ice core, Clim. Past, 10, 1905–1924, https://doi.org/10.5194/cp-10-1905-2014, 2014.

Zhang, W., Zhang, S., Wan, C., Yue, D., Ye, Y., and Wang, X.: Source diagnostics of polycyclic aromatic hydrocarbons in urban road runoff, dust, rain and canopy throughfall, Environmental Pollution, 153, 594–601, https://doi.org/10.1016/j.envpol.2007.09.004, 2008.

540