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Leaf wax biomarkers of a high-mountain lake area in western iberia—Implications for environmental reconstructions

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Leaf wax *n*-alkane biomarkers are widely used to infer past vegetation dynamics and hydroclimate changes. The use of these compounds strongly relies on the characterization of modern plants. However, few studies have explored leaf waxes of modern plants and their application to reconstructing climate and environmental changes in the Iberian Peninsula, a region known for its high vulnerability to climate change. In this study, we characterize the distributions and compound-specific isotopic compositions of the leaf waxes of dominant plants in the vegetation cover, soil, and surface sediment of the Lake Peixão area, a high-mountain glacial lake in Serra da Estrela (central Portugal). Our results show that the modern oro-Mediterranean (subalpine) vegetation of the study area is dominated by C3 grasses/herbs and shrubs that preferentially produce long-chain leaf waxes ($\geq C_{27}$). The C_{31} *n*-alkane display the overall highest concentration, produced by some grasses and shrubs, but especially Erica sp (heather), which is highlighted as a major source for the total n-alkane pool in the lake sediments. C₂₉ is the second-most abundant and the most equally produced *n*-alkane of the vegetation cover; C₂₅ and C₂₇ homologs are mainly associated with aquatic-related grasses/herbs, while C33 and C35 are particularly linked to cold-drought tolerant Juniperus sp. shrubs. Shrubs show higher but proportional values than grasses/herbs in the isotopic space, suggesting a directly proportional physiological adaptation of the two ecological forms to the prevailing climatic and environmental factors of the study area. C₂₉ is pointed as the most representative (or less plant-biased) leaf wax *n*-alkane in the lake sediments. Thus, δD of C_{29} *n*-alkane is interpreted as a robust terrestrial hydrological indicator (δD_{terr}), which signal is believed to be strongly influenced by the mean air temperature and/or precipitation amount. Despite the sparse vegetation and small catchment area, the apparent hydrogen fractionation factor, determined from δD_{terr} of the lake surface sediment, is in line with the modeled global mean values for the latitude of the study area. The different molecular and compound-specific signatures of the studied oro-Mediterranean species have the potential to support future interpretations of leaf wax biomarkers in the Iberian Peninsula.

KEYWORDS

leaf wax *n*-alkanes, terrestrial biomarkers, stable isotopes, western iberia, high-mountain lake



1 Introduction

Reconstruction of past climate and environmental conditions requires a thorough characterization and understanding of how modern ecosystems produce physical, chemical, or biological indicators, known as proxies, in response to specific climatic and environmental conditions. An ecosystem that promptly responds to climate change is often referred to as a "climatic sensor" (Evans et al., 2013). An excellent example of this type of sensor is the vegetation cover of the Iberian high mountains, known as sensitive ecosystems and natural laboratories (Toro et al., 2006). The lakes of these high mountains can act as sentinels of climate and environment changes and are capable of archiving evidence of the temporal evolution of the surrounding landscape in their sediments (Adrian et al., 2009). The Iberian Peninsula (IP) displays a wide range of climatic conditions due to its mid-latitude location, between the Eastern North Atlantic and the western Mediterranean regions, its complex orography, and the interplay between sub-tropical and sub-polar air masses. These features make IP bioclimatic diverse and particularly sensitive to climate changes and a recognized "hot spot" for the study of past climate, especially in quaternary studies (e.g., Sánchez-López et al., 2016; Abrantes et al., 2017; Baldini et al., 2019; Thatcher et al., 2020).



FIGURE 1

Morphological map of the Iberian Peninsula. The red star indicates the Lake Peixão location (Serra da Estrela, Portugal). Csa: Hot summer Mediterranean climate; Csb: Mild summer Mediterranean climate; Cfb: Oceanic. The dark blue arrows represent the moist rich prevailing westerlies from the Atlantic Ocean entering Iberia. Sources: European map from shadedrelief. com. River network from the European Environment Agency (EEA). Topography and bathymetry, sourced from portal.emodnetbathymetry.eu, overlapped with the simplified Köppen-Geiger climate classification for the Iberian Peninsula (1971–2000) (adapted from the Iberian Climate Atlas, 2011).



FIGURE 2

The Lake Peixão area. The light-blue dashed line represents the upslope watershed contour obtained from the elevation model of Google Earth Pro. The enumerated triangles represent sites of sample collection: 1–*Agrostis* sp.; 2–*Antinoria* sp.; 3–*Juncus* sp.; 4–*Cytisus* sp.; 5–*Juniperus* sp. and respective litter; 6–*Erica* sp. and respective litter; 8–Soil sample. Colors represent orange–grasses; light blue–aquatic grass; light green–soft rush; dark green–shrub; brown–soil. The eye-view symbol point to the view seen in the respective photos (A,B), and (C). Satellite imagery from Google Earth (date: 26 May 2019).

Plants produce a wide range of organic compounds, including leaf wax biomarkers, which are used as climate and environmental proxies (Peters et al., 2005; Castañeda and Schouten, 2011; Berke, 2018). However, the use of these compounds requires a comprehensive characterization of modern plants composition and understanding of their respond to the prevailing climate and environment factors (e.g., Bush and McInerney, 2013; Howard et al., 2018; Liu et al., 2018; Wang et al., 2018). The characterization of modern leaf wax compositions in the IP is still understudied (Ferreira et al., 2007; Ortiz et al., 2010, 2011, 2016; López-Días et al., 2013), despite the increasing number of past reconstructions using these compounds in both marine and lacustrine records (e.g., Jambrina-Enríquez et al., 2016; Taylor et al., 2018; Schirrmacher et al., 2020; Toney et al., 2020). Here, we focus on leaf wax *n*-alkanes, *n*-alkyl lipids with the general formula of $C_n H_{2n+2}$ (where *n* is the carbon number in the chemical chain), synthesized by plants to act as a protective barrier from the external environment and prevent water loss (Eglinton and Hamilton, 1967; Cranwell et al., 1987; Post-Beittenmiller, 1996; Shepherd and Griffiths, 2006). The high resistance to degradation and strong isotopic relationship of *n*-alkanes with vegetation type and water sources make compound-specific stable isotope analysis (CSIA) a valuable tool to infer past organic matter sources and hydrologic changes (e.g., Leider et al., 2013; Curtin et al., 2019; Hahn et al., 2021; Imfeld et al., 2022). Temporal variations in the stable carbon isotope ratio ($^{13}\mathrm{C}/$ ¹²C) of *n*-alkanes ($\delta^{13}C_{n-alk}$) in sedimentary records generally reflect changes in vegetation sources and water use efficiency of a catchment area (e.g., Freeman and Colarusso, 2001; Schefuß et al., 2003; Rommerskirchen et al., 2006; Diefendorf and Freimuth, 2017). The hydrogen isotopic ratios (D/H) of leaf wax *n*-alkanes (δD_{n-alk}) ultimately reflect the isotopic composition of precipitation. However, this signal results from a net fractionation process involving isotopic signatures of groundwater, lake water, soil, plant transpiration, and biosynthetic fractionation (see Sachse et al., 2012 for review).

In this work, we aim to fill the gap in studies of modern plants and their relationship as precursors of hydrocarbon biomarkers in IP. For that, we explored leaf wax *n*-alkanes of predominant species in the vegetation cover and surface sediment of Lake Peixão. This study area, located on top of the Serra da Estrela Mountain, lies between the Atlantic and Mediterranean bioclimatic zones, being considered a sensitive place to explore past hydrological changes and is in one of the most vulnerable regions to current climate changes (Allan et al., 2021). These results will add new insights about leaf wax biomarkers in IP and ultimately contribute to more reliable and robust interpretations of their sedimentary records in this key-region for reconstructing past climate and environmental changes.

2 Materials and methods

2.1 Study area

The study area is located in the Serra da Estrela Mountain, a NE–SW-directed intraplate mountain range in the westernmost sector of the Iberian Central System (Figure 1), central Portugal. This alpine range of the granitic basement (Ribeiro et al., 1990)

comprises the highest point (1,993 m above mean sea level, amsl) and the largest nature conservation area of continental Portugal. Here, a remarkable and unique set of glacial landforms and deposits (e.g., see Vieira, 2004, 2008) promoted the recognition of this region as a UNESCO Global Geopark. One of these glacial features is the study site, Lake Peixão (Figure 1) (40°20'35"N; 7°36'19"W), a relatively small (0.015 km²) and subcircular lake (ca. 140 m diameter) at 1,677 m amsl. This alpine lake has a glacial origin, formed after the melting of the last icesheets ca. 14.7 \pm 0.32 ka cal BP, which the small catchment area (ca. 0.30 km²) is characterized by enclosed and steep granitic rocks (Figure 2) [Hernández et al., (2022) (in review)]. Few and small ephemeral streams flow into the lake, which has slightly acidic, oligotrophic, and monomitic waters (Boavida and Gliwicz, 1996). A spill point in the southern part can control the maximum lake level, draining into the Candeira valley (Figure 2). The landscape near the lake is characterized by sparse but well-constrained oro-Mediterranean vegetation patches dominated by shrubs such as Erica spp., Juniperus communis, Genista spp., Cytisus spp., and Calluna vulgaris (Connor et al., 2021), and grasslands mainly composed by Agrostis delicatula and Nardus stricta. Soils are rare and poorly developed; aquatic plants are sparse and mainly located near ponds and the lake, while trees are absent.

In general, Serra da Estrela region is characterized by a Mediterranean climate, with warm/dry summers and cold/wet winters, and is marked by strong vertical gradients in temperature and precipitation (Vieira et al., 2005; Mora, 2010). The ensemble historical climate of the region, from 1971 to 2000 [see http://portaldoclima.pt/en (Accessed 29 March 2022)], shows that the warmest and driest months are July and August, with mean values of approximately 20°C and 17 mm of rainfall. The coldest and wettest months are December and January, with a mean temperature of approximately 5°C and 150 mm precipitation, where precipitation often falls in the form of snow on the top of the mountain (Carreira et al., 2009, 2011).

The spatial variability of precipitation in this mountain range is highly controlled by altitude and slope orientation, with the basal areas receiving approximately half of the precipitation amount of the mountain summit, which can reach values of approximately 2,500 mm/year (Mora, 2010). The mountain range roughly defines a boundary between two Mediterranean climates (following the Köppen-Geiger climate classification system), one to the north characterized by mild summers (csb) and the other to the south characterized by hot and dry summers (csa) (Figure 1). The moisture-rich westerlies are predominant during winters and show a gradual D-depletion from the coast inland (Carreira et al., 2009). The atmospheric moisture sources of this part of the IP are almost exclusively derived from the prevailing westerly Atlantic air masses (Carreira et al., 2009; Cortesi et al., 2013; Thatcher et al., 2020), contrasting with other regions, such as the southeastern, which can also be significantly influenced by Mediterranean sources (Cortesi et al., 2013; Taylor et al., 2018; Schirrmacher et al., 2020; Toney et al., 2020). The high-altitude setting of Serra da Estrela results in temperature being an important driver of moisture condensation and the δ D value of precipitation (δ D_{prc}) (Carreira et al., 2009; Oliveira and Lima, 2010). This is reflected by a strong relationship between mean values of δ^{18} O and δ D vs. temperature of monthly rainfall for Portugal's mainland (Carreira et al., 2009), which is also observed in other Iberian mountains (Giménez et al., 2021). The combination of the settings mentioned above and climatic and environmental factors makes Serra da Estrela one of the most biodiverse in Europe (Jansen, 2011) and a prime location to investigate the relationship between climate and ecology (Connor et al., 2021).

2.2 Material

A total of seven samples of dominant vegetation (four grass/herbs and three shrubs), three vegetation litters (one grass/herbs and two shrubs), a soil sample collected from the Lake Peixão catchment area (Figure 2), and a lake surface sediment. Modern plant samples were collected in September 2020; each plant sample was composed of several branches or leaves from different plants near the same collecting spot. These samples were wrapped in aluminum foil and sealed in an airtight bag. The vegetation litter samples were collected beneath the respective plant to ensure their source, and the soil sample comprises the top 2 cm of one of the few places in the lake's catchment where soils were available (Figure 2). Both litter and soil samples were collected directly with precombusted jar glasses. All samples were stored in a cool bag, directly transported to the Portuguese Institute of Sea and Atmosphere (IPMA, Portugal), and stored in a cool room (~5°C) until analysis. The lake surface sediment sample correspond to the 0 cm-1 cm depth of 8.63 m long core retrieved from the center of the Lake Peixão in June 2019 (PEX19-01) using a UWITEC® Piston Corer. Based on ²¹⁰Pb measurements, this surface sediment sample comprises 11 ± 4 years [Hernández et al. (2022) in review].

2.3 *n*-alkane analyses and characterization

All samples were freeze-dried, homogenized, and analyzed in the Biogeochemistry laboratory at IPMA. The vegetation samples were composed of leaves and stems from seven samples. Depending on the vegetation type, wood and stems were removed as much as possible. The samples were cut into small pieces with clean metal scissors. The three samples of the vegetation litter, the soil sample, and the lake surface sediment sample presented minimal clastic material (silt and clays), and no sieving was needed. Before extraction, 1,000 ng of an internal standard solution (hexatriacontane, tetracontane, and nonadecanol-1-ol) was added to approximately 2.5 g of the sample. The lipid compounds were extracted three times using dichloromethane in an ultrasonic bath.

The total lipid extract (TLE) was hydrolyzed with potassium hydroxide in methanol (KOH/MeOH 6%, ~12 h, at room temperature). The neutral fraction containing n-alkanes was recovered with n-hexane by liquid-liquid extraction and cleaned with ultrapure water to remove traces of KOH. The *n*-alkanes were fractionated from the neutral extracts using 5 cm of deactivated silica gel (0.040 μ m–0.063 μ m mesh) and AgNO₃silica gel Pasteur pipette columns with n-hexane as eluent. The *n*-alkane extracts were dried and diluted with toluene, and 1 µl was injected into a gas chromatography-flame ionization detector (GC-FID; Varian Model 3,800) equipped with a 1,079 programmable temperature vaporizing injector (PTV) for cold on-column with a CPSIL-5 CB column coated with 100% dimethylsiloxane capillary column (50 m \times 0.32 mm \times 0.12 µm film coating) connected to a fused silica precolumn (5 m \times 0.25 mm \times 0.25 µm). The oven temperature was programmed from 90° (holding time of 1 min) to 170°C at 20°C/min, then to 280°C at 6°C/min (holding time 25 min), and finally, to 315°C at 10°C/min (holding time of 12 min). The injector was programmed from 90°C (holding time of 0.5 min) to 310°C at 200°C/min. Samples were injected (1 μ l) in split mode using H₂ as the carrier gas (~2.5 ml/min). The temperature of the detector was constant at 320 $^\circ\mathrm{C}$ with 35 ml/min of H₂ flow, 300 ml/min of airflow, and 30 ml/min of N2 flow. The total run time for each sample was 63.83 min.

The *n*-alkanes were identified by comparing the retention times to an external standard mixture (*n*-alkanes ranging from C_{17} - C_{36}). The *n*-alkane concentrations (ng/g, dry weight sample) were determined based on the internal standard *n*-hexatriacontane (C_{36}).

Several indices have been developed to characterize the n-alkane distributions of samples (see Li et al., 2020). The most widely used are (a) the average chain length (ACL, Poynter, 1989); (b) the carbon preference index (CPI, Bray and Evans, 1961; Marzi et al., 1993), and (c) the proportion of aquatic plants (P_{aq}, Ficken et al., 2000).

(a) The ACL is used to identify the preferentially produced *n*-alkane length chains in a sample (Poynter, 1989) based on a weighted average of the *n*-alkane concentrations from C₂₇ to C₃₃, calculated as:

$$ACL_{27-33} = odd \sum_{n=27}^{33} (Cn \times n) / \sum_{n=27}^{33} (Cn)$$
(1)

where C_n is the concentration of a given *n*-alkane and *n* is the number of carbons in its chain.

(b) The CPI is the concentration weighted ratio of odd to even *n*-alkane chain lengths, which is normally used to examine

the general source of sedimentary biomarkers and/or degradation processes. It can be calculated as follows:

$$CPI_{27-33} = 1/2 [odd \sum_{n=27}^{33} (Cn) / even \sum_{n=26}^{32} (Cn) + (odd \sum_{n=27}^{33} (Cn) / even \sum_{n=28}^{34} (Cn)]$$
(2)

where C_n is the concentration of a given *n*-alkane.

(c) The P_{aq} commonly used to evaluate the relative contributions of *n*-alkanes from aquatic versus terrestrial plants in the lake sediments: $P_{aq}=(C_{23} + C_{25})/(C_{23} + C_{25} + C_{29} + C_{31})$, where C_{23} and C_{25} are considered from aquatic-related plant origins, whereas C_{29} and C_{31} are from terrestrial plants (Ficken et al., 2000).

CSIA was conducted on the *n*-alkanes when amounts were allowed by gas chromatography-isotope ratio mass spectrometry (GC-IRMS) at the MARUM Center for Marine Environmental Science, University of Bremen (Germany).

The $\delta^{13}C_{n-alk}$ values were measured with a Thermo Trace GC Ultra coupled to a Finnigan MAT 252 mass spectrometer. The n-alkane extracts were injected in splitless mode before being oxidized to CO₂ by a combustion reactor at 1,000°C. The instrument was equipped with an HP-5ms capillary column $(30 \text{ m} \times 0.25 \text{ mm} \times 0.25 \text{ µm} \text{ film coating})$; helium was the carrier gas (purity 99.999%) at a constant flow of 1.5 ml/min. The produced CO₂ was then injected into the mass spectrometer for analysis. The injector temperature was programmed to be constant at 250°C. The GC temperature was set from 120°C (hold time of 3 min) to 320°C at a heating rate of 5°C/min (hold time of 15 min). The $\delta^{13}C_{n-alk}$ values of the individual compounds were calibrated by injecting pulses of CO₂ from an external reference gas that was automatically introduced into IRMS at the beginning and end of each analysis. The $\delta^{13}C$ values are reported in ‰ relative to the Vienna Pee Dee Belemnite (VPDB) standard and represent an average of duplicates with a standard deviation less than 0.5‰ whenever n-alkane amounts allowed for multiple runs.

Standard δD_{n-alk} was performed on a Thermo Trace GC equipped with an HP-5ms column (30 m × 0.25 mm × 0.10 µm film coating) coupled to a Thermo Fisher MAT 253 (IRMS) *via* a pyrolysis reactor (operated at 1,420°C). The GC oven program was similar to the conditions used to analyze the carbon isotopic composition. The injection of known isotopic composition standards after every six measurements and the daily determination of the H³⁺ factor using H₂ reference gas were used to control measurement accuracy. The δD_{n-alk} was calibrated against an external H₂ reference gas and reported

Sample	Ecological form	Environment	(n-alkane	(n-alkanes ng/g)															(<i>n</i> -alkanes ng/g)	<i>n</i> -alkane indices						
			Sample type**	C ₁₇	C ₁₈	C ₁₉	C ₂₀	C ₂₁	C ₂₂	C ₂₃	C ₂₄	C ₂₅	C ₂₆	C ₂₇	C ₂₈	C ₂₉	C ₃₀	C ₃₁	C ₃₂	C ₃₃	C ₃₄	C ₃₅	Total n-alk***	ACL ₂₇₋₃₃	CPI ₂₇₋₃₃	P _{aq}
Agrostis sp.	grass/herb	terrestrial	Veg	11	326	65	193	354	145	2,857	1912	45,038	4,765	96,331	15,640	130,298	9,927	112,330	5,719	15,198	833	563	4.43×10^{5}	29.3	10.4	0.2
Nardus sp.	grass/herb	terrestrial	Veg	155	286	5	200	39	102	910	948	14,346	3,420	19,565	8,361	106,318	7,281	307,444	4,367	56,622	184	1,332	5.32×105^{5}	30.6	22.6	0.0
Antinoria sp.	grass/herb	aquatic-related	Veg	128	266	159	250	1,523	1,163	5,795	4,868	33,311	2,985	18,591	2074	16,629	1,349	25,081	1,054	22,256	186	1703	1.39×10^5	30.2	14.4	0.5
Juncus sp.	grass/herb	aquatic-related	Veg	28	218	263	89	830	690	3,685	1,222	2,905	1892	12,554	1,040	5,212	239	2,775	203	300	78	168	3.44×10^4	28.1	9.8	0.5
Erica sp.	shrub	terrestrial	Veg	45	204	188	201	4,360	1,018	6,152	1,411	12,809	3,831	84,604	13,728	311,759	42,497	659,377	41,027	178,062	2,669	845	1.36×10^{6}	30.5	12.3	0.0
Juniperus sp.	shrub	terrestrial	Veg	5	7	0	33	4	78	275	103	413	171	1,225	453	6,556	1,196	33,999	15,430	373,934	23,653	88,742	5.46×10^{5}	32.8	17.2	0.0
Cytisus sp.	shrub	terrestrial	Veg	4	322	32	206	255	681	39,449	3,532	24,616	757	6,830	756	13,283	1,447	18,675	369	538	55	80	1.12×10^{5}	29.7	13.4	0.7
Nardus sp. (L.)	grass/herb	terrestrial	Veg. litter	28	63	2	30	43	56	100	58	253	50	504	160	3,635	430	17,934	579	7,780	113	402	3.22×10^{4}	31.2	23.9	0.0
Erica sp. (L.)	shrub	terrestrial	Veg. litter	61	134	8	59	50	48	95	67	207	65	590	142	2,365	365	11,105	690	5,552	86	137	2.18×10^4	31.2	15.4	0.0
Juniperus sp. (L.)	shrub	terrestrial	Veg. litter	25	156	0	61	318	113	430	80	664	153	2,755	519	10,323	1870	37,055	6,652	118,922	3,845	15,050	1.99×10^{5}	32.2	15.8	0.0
Soil sample	-	terrestrial	soil	1	3	8	5	27	29	57	19	105	30	312	81	1,117	103	1,561	124	790	33	46	4.45×10^3	30.5	11.2	0.1
Lake surface sediment	-	lake	sediment	392	421	1,161	695	2,968	1,504	5,482	2,702	8,899	4,231	15,116	3,975	22,677	3,251	37,111	3,069	19,609	841	1921	1.36×10^5	30.3	7.5	0.2

TABLE 1 – n-Alkane concentrations* and indices (ACL₂₇₋₃₃, CPI₂₇₋₃₃, Paq) of plant leaves and stems and soil and surface sediment samples from the Lake Peixão (Serra da Estrela, Portugal).

*Highest concentrations values in bold; **Veg. meaning vegetation; ***sum of *n*-alkane concentrations from C₁₇ to C₃₅.



n-alkane distributions and compound-specific isotopic values ($C_{25}-C_{35}$) of the dominant plants, vegetation litter (L), soil and surface sediment samples of the Lake Peixão area. The whiskers correspond to 1 SD, where multiple measurements on the respective compound were performed. *Normalization against the total n-alkane concentration of each sample.



FIGURE 4

ACL₂₇₋₃₃ vs. total *n*-alkane concentrations (ng/g, in logarithmic scale) of the two dominant ecological forms of the Lake Peixão catchment. The overall trend is represented by a logarithm regression line in gray. Green triangles represent shrubs; colored circles represent grasses/herbs, aquatic-related plants in blue, and nonaquatic plants in yellowish.



in ‰ Vienna Standard Mean Ocean Water (VSMOW). When *n*-alkane amounts allowed multiple runs, the samples were run in duplicate, and the reported values represent the mean with a standard deviation less than 3‰.

The apparent fractionation (ε_{app}) between δD of leaf wax *n*-alkanes (δD_{n-alk}) and precipitation (δD_{prc}) was calculated using the equation described in Sessions et al. (1999): $\varepsilon_{n-alk/}$ _{water} = 1,000 [(δD_{n-alk} +1,000)/(δD_{prc} +1,000)] -1, in %. The δD_{prc} used as a reference for the mean annual precipitation was estimated using the Online Isotopes in Precipitation Calculator (OIPC3.1; Bowen and Revenaugh, 2003) by constraining the central altitude, longitude, and elevation of the lake catchment area. This modeled δD_{prc} value was then compared with the hydrogen isotopic value of nearby spring water.

3 Results

3.1 Distribution and concentration of *n*-alkanes

The modern vegetation samples exhibit the typical strong odd over even chain length predominance, with a wide range of total *n*-alkane concentration (sum of concentrations from C_{17} to C_{35} *n*-alkane), varying from 3.4×10^4 to 1.4×10^6 ng/g (dry weight) (Table 1). Most of the analyzed plants show a predominance of long-chain n-alkanes ($\geq C_{27}$), particularly C_{31} and C_{29} homologs (Figure 3).

Principal component analysis (PCA) of the *n*-alkanes (concentration and isotopic compositions) performed in

PAST4.07 free software (Hammer et al., 2001) (Supplementary Figure S1) suggests that samples can be grouped according to the ecological forms: grasses/herbs and shrubs (see Figure 3). On the other hand, for ACL_{27-33} (U, p = 0.12) and CPI_{27-33} (U, p = 0.35) the Mann-Whitney U tests, performed in STATISTICA 7.0, suggests no statistical difference in between the two forms (Supplementary Figure S2). In addition, grasses/herbs and shrubs appear to share a generally positive trend between ACL₂₇₋₃₃ and total *n*-alkane concentration, however under low statistical significance (Figure 4). Nardus sp. and Agrostis sp. presented the highest n-alkane concentrations in this ecological form, preferentially producing C₃₁ and C₂₉, respectively (Table 1). Antinoria sp (aquatic-related grass) exhibits proportional amounts of C25, C31, C33, and C27 homologs (decrescent in this order) (Figure 3). Juncus sp (herb) has the lowest total n-alkane concentration and a maximum at C27. Erica sp. is highlighted as having the highest total n-alkane concentration of among all the studied plants (Table 1). These species preferentially produce C₃₁, twenty times more than the other investigated shrubs. On the other hand, Cvtisus sp. is the shrub with the lowest total n-alkane concentration but produces considerably more mid-chains than the other shrubs, particularly the C_{23} homolog (Figure 3; Table 1).

The decrease in *n*-alkane concentration from vegetation to the respective litter samples was significant, although the dominant homolog is preserved from its vegetation source (Table 1). For example, Juniperus sp. displays a decrease of 68% in the dominant homolog (C33), Nardus sp. a decrease of 96% in $C_{31,}$ while ${\it Erica}$ sp. a decrease of 97% in $C_{31}.$ The soil sample exhibits a strong odd over even chain length dominated by long-chain n-alkanes, resulting in ACL₂₇₋₃₃ and CPI₂₇₋₃₃ values of 30.5 and 11.2, respectively (Table 1). The total concentration of odd C_{27} — C_{33} chains is 3.8 × 10³ ng/g (dry weight) and accounts for ~85% of the total n-alkane concentration. The lake surface sediment and soil samples share the same dominance and long-chain n-alkane distribution (Table 1) ($C_{31} > C_{29} > C_{33} > C_{27} > C_{25} > C_{35}$) and similar ACL₂₇₋₃₃ (30.3). However, the lake sediments have a considerably larger concentration (thirty times more) of *n*-alkanes, lower CPI_{27-33} (7.5), and higher P_{aq} (0.19) than the soil sample (Table 1).

3.2 Compound-specific $\delta^{13}C_{n-alk}$ (odd $C_{25}-C_{33}$)

The $\delta^{13}C_{n-alk}$ values of the dominant plants of the Lake Peixão catchment area show an overall range of -37 to -29%(Supplementary Table S1), with minor differences between homologs but considerable interspecific variability (Figure 3). In general, longer carbon chains show the lowest $\delta^{13}C$ values, mainly C_{29} and C_{31} homologs (Figure 3). Grasses/herbs display the most negative δ^{13} C values, ranging from -37 to -30‰. From those, *Juncus* sp. showed the highest values (-35 to -30‰) (Supplementary Table S1). The $\delta^{13}C_{n-alk}$ of shrubs varies from -34 to -29‰. *Juniperus* sp. and *Erica* sp. exhibit particularly narrow δ^{13} C ranges, the latter ranging from -32 to -30‰, whereas *Cytisus* sp. has a broader range (-34 to -29‰) (Supplementary Table S1). The δ^{13} C values of C_{29} in the two dominant ecological forms in the study area showed clear and opposite linear relationships with ACL₂₇₋₃₃ (Figure 5) and CPI (Supplementary Figure S3). The $\delta^{13}C_{n-alk}$ values of vegetation litter vary from -35 to -30‰, generally displaying similar values to their respective vegetation sources (Supplementary Table S1). The lake surface sediment shows similar $\delta^{13}C_{n-alk}$ values among homologs (-33 to -32‰), with the lowest values of C_{29} and C_{31} (Figure 3).

3.3 Compound-specific δD_{n-alk} (odd $C_{25}-C_{33}$)

The δD_{n-alk} of the analyzed vegetation shows a wide range of values, from -236 to -119%, with, grasses displaying lower δD_{n-alk} values than shrubs (Supplementary Table S1). There is a remarkable difference between nonaquatic and aquatic-related grasses/herbs, with the latter showing higher δD_{n-alk} values than their terrestrial counterparts. For example, for C277, the terrestrial grasses Agrostis sp. and Nardus sp. display values of ca. -210‰, while Antinoria sp. and Juncus sp. show values of ca. -180‰ (Figure 3). In addition, Cytisus sp. showed higher and more variable values among homologs (-146 to -119‰) than plants such as Erica sp (-161 to -151‰) (Figure 3). Litter of Nardus sp. exhibits higher δD_{n-alk} values than its source; in C_{29} , this relative increase corresponds to ca. 18%, from -236 to -193‰. This contrasts with shrub litter, which shows lower values than their sources (Figure 3). The δD values of C_{31} in *Erica* sp. represent a decrease of 18%, from -155 to -183‰, and in Juniperus sp., a similar decrease from -137 to -157‰ (Figure 3; Supplementary Table S1). The δD_{n-alk} from lake surface sediment displays values ranging from –189 to –161‰, with $C_{\rm 25}$ and $C_{\rm 27}$ homologs showing the highest (similar) values (Figure 3). In contrast, longer chains, particularly C₃₁, displayed the lowest values.

3.4 Apparent hydrogen isotope fractionation

The modern δD of the mean annual precipitation (δD_{MAP}) estimated using OIPC3.1 for the study area is $-56 \pm 5\%$, a value similar to the -53% mean value for the wetter months during the main growing season (from March to May) and within the range of the near-annual mean value of $-51 \pm 2\%$ from the Fonte dos Perús spring, which is ca. 1 km of the study area but in the opposite façade of the mountain (Carvalho, 2013).



The specific fractionation factors ($\varepsilon_{n-alk/MAP}$) determined for the modern vegetation range from $-191 \pm 17\%$ to $-67 \pm 6\%$ (Table 1) values constrained by a common δD_{MAP} value of the study area; error is given by the standard deviation of measurements in *n*-alkanes and δD_{MAP} . Therefore, Agrostis sp. and Nardus sp. display mean ε_{app} values (from C₂₅ to C₃₃) of $-158 \pm 32\%$ and $-177 \pm 35\%$, respectively, whereas the aquatic-related counterparts, Antinoria sp. and Juncus sp., show higher values of $-138 \pm 28\%$ and $-127 \pm 23\%$, respectively. C₃₁ was the only compound present in sufficient quantities in all three shrub species for reliable $\delta^{13}C/\delta D$ measurements; thus, using this component as a reference, Erica sp. shows the highest fractionation ($\varepsilon_{C31/MAP} = -105 \pm 9\%$), while *Cytisus* sp. the lowest ($\epsilon_{C31/MAP} = -77 \pm 7\%$). The surface sediment of the lake also shows an increase in $\varepsilon_{n-alk/MAP}$ from $C_{29} > C_{33} > C_{31}$, ranging from $-129 \pm 12\%$ to $-141 \pm 13\%$, and C₂₅ and C₂₇ homologs display the lowest $\varepsilon_{n-alk/MAP}$, approximately $-112 \pm 10\%$.

4 Discussion

4.1 Leaf wax *n*-alkane biomarkers of modern plants

The small and well-constrained catchment area of the Lake Peixão presents ideal characteristics for assessing the vegetation cover inputs into lake sediments. The C_{31} and C_{29} are the predominant *n*-alkanes of the modern plants in the vegetation cover. Grasses/herbs and shrubs display similar ACL_{27-33} and CPI_{27-33} values (Supplementary Figure S2) that can suggest traits of a common physiological adaptation of this vegetation type, but also that the application of such indicators alone may be questionable to differentiate between the two terrestrial forms. This common physiological response may also be reflected in the suggested positive trend between ACL_{27-33} and *n*-alkane concentration (Figure 4); however, the limited dataset appears to constraint the statistical significance of this hypothesis, which will be likely addressed in future studies.

The high amount of *n*-alkanes produced by Erica sp. suggests that these plants are significant sources of leaf waxes in the study area, particularly the C₃₁ *n*-alkane. These findings agree with other modern vegetation studies in the IP (Ferreira et al., 2007; Ortiz et al., 2016), where C_{31} *n*-alkane is pointed out as a possible indicator of heather inputs (Loidi et al., 2007; Ortiz et al., 2016). Our results also suggest that Juniperus sp. is another important source of long-chain *n*-alkanes ($\geq C_{33}$), accounting almost exclusively for the production of C35 in the vegetation cover (Figure 3; Table 1). These observations align with the results obtained for other species of Juniperus sp. in central Spain (Schäfer et al., 2016a), suggesting that these especially longchains compounds can be used as an ecological proxy and indicators of arid and cold conditions (Thomas et al., 2007; Dasgupta et al., 2022). In Iberian heathlands, the relative increase in grasses/herbs over shrubs can reflect an increase in water availability, as they are often in contact with mires (Loidi et al., 2007). In the study area, the aquatic-related plants are restricted to the grass/herb ecological form, such as Antinoria sp. and Juncus sp., which are relatively abundant in small ponds and streams. The n-alkane analyses of these species suggest C27 and C25 as indicators of aquatic or aquatic-related plants matter and water availability in the lake Peixão ecosystem.

The relatively high contribution of mid-chain compounds of some shrubs, such as Cytisus sp., (Figure 3; Table 1), highlights that caution is needed when applying distributions n-alkane, such as Paq, to infer past aquatic inputs or lake levels in downcore paleoreconstructions (Xie et al., 2020). However, the relative deviation in the n-alkane distribution of Cytisus sp. relative to other shrubs is not reflected in the isotopic space ($\delta^{13}C_{n-alk}$ and δD_{n-alk} (Figures 3, 6), which encourages the application of CSIA to better assess vegetation sources in lake sediments. In fact, the two ecological forms are well-separated in the isotopic space (Figure 6), where the observed linear proportionality may result from the inferred common physiological response to the shared climatic and environmental conditions of the study area, i.e., precipitation, temperature, humidity, and water source. For example, the lower δD_{n-alk} of grasses/herbs may imply lower transpiration rates and lower water loss than shrubs, but on the other hand, its lower $\delta^{13}C_{n-alk}$ values may also imply that grasses/herbs have less water use efficiency than shrubs (Hou et al., 2007; Gao et al., 2014). While the similar ACL₂₇₋₃₃/CPI₂₇₋₃₃, along with the proportionally distinct isotopic

values, are inferred to be the reason for the opposite trends between the two ecological forms in ACL/CPI vs. $\delta^{13}Cn_{-alk}$ (e.g., Figure 5; Supplementary Figure S3).

4.2 The source of Lake Peixão *n*-alkanes and its implications for paleo-reconstructions

The strong predominance of odd-over-even carbon chain compounds, high ACL₂₇₋₃₃, and low P_{aq} values (Table 1) in the surface sediment of Lake Peixão suggests that the *n*-alkanes were primarily derived from terrestrial plants from the catchment area. Additionally the high CPI₂₇₋₃₃ values (> 5) in the lake sediments suggest good preservation of the *n*-alkane content which gives confidence in the use of leaf waxes record in the lake sediments (Pancost and Boot, 2004).

Our results show that from plants to vegetation litter, and lake surface sediment the isotopic signal is well preserved despite important changes in *n*-alkane concentrations (Table 1). Moreover, the relative proportion of dominant chains (Figure 3) is also maintained The ACL₂₇₋₃₃ and *n*-alkane distribution of modern soil and the lake surface sediment remain identical even though the latter reflects higher contribution of aquatic or aquatic-related plants (Figure 3; Table 1). On the other hand, the CPI₂₇₋₃₃ offset between plants, vegetation litter, and the lake surface sediment reflects the complex plant-to-sediment transfer processes, which involve a series of admixtures of different plant inputs and biodegradation; for instance, by microbes (Zech et al., 2010; Schäfer et al., 2016; Stout, 2020; Thomas et al., 2021).

The $\delta^{13}C_{n-alk}$ values (ca. -33‰) of lake surface sediment suggest a predominance of C3 plants (Figure 3) and reinforce shrubs as a major source of waxes in the study area (Figure 3; Table 1). The two dominant ecological forms produce significant and relatively equivalent amounts of C29, showing proportional values of carbon and hydrogen fractionation that may result from a consistent physiological response of plants to the same environment and climatic factors. Thus, we expect C₂₉ to provide a more holistic overview of the n-alkane composition of the terrestrial vegetation cover in the Lake Peixão sediments compared with other long-chain compounds, such as C₃₁, that appear to be particularly biased towards species like Erica. This hypothesis supports global compilations that report C₂₉ as a widely produced terrestrial n-alkane strongly and consistently associated with regional mean δD_{prc} (Sachse et al., 2012; Liu and Liu, 2019; McFarlin et al., 2019). Moreover, the modern $\varepsilon C_{29}/_{MAP}$ value (-129 ± 12‰) of lake surface sediment is remarkably consistent with the global modeled values by Liu and An (2019) and McFarlin et al. (2019) for the latitude of the study area. Therefore, we suggest the δD of C_{29} as a robust indicator of terrestrial hydrology, i.e., δD_{terr} , in paleostudies in both sediments of Lake Peixão and downstream areas.

e and *n*-alkane isotopic signatures. tor in higher $\delta^{13}C_{n-alk}$ and δD_{n-alk}

The strong relationship between temperature and n precipitation is an important paleo-hydrological factor in h the study area since plants mostly synthesize their n-alkanes during early growing season or when water is available (Freimuth et al., 2017; Diefendorf et al., 2021), n which in the IP occurs during spring-early summer and the during autumn-winter (Pérez Latorre and Cabezudo, 2002; E García-Alix et al., 2021). Therefore, the downcore temporal variability in the values of δD_{terr} in the Lake Peixão sediments al is expected to mainly reflect bi-seasonal to annual changes in mean air temperature and/or precipitation amount. In the study site, lower temperatures often imply an increase in al condensation, resulting in an inherent coupled effect of

precipitation amount (amount effect) that may be imprinted in the δD_{terr} . Hence, low (high) δD_{terr} values in downcore reconstructions are expected to reflect a decrease (increase) in temperatures and/or increase (decrease) in precipitation (snow or rainfall).

Moreover, based on the ecology of modern plants and the compositions of different *n*-alkanes analyzed, we expected the δD values of C25 and C27 in the lake sediments to better reflect changes in lake water sources. The scarcity of true aquatic plants in the study area makes it challenging to confirm this hypothesis. Aquatic plants typically show higher $\delta^{13}C_{n-alk}$ values than their terrestrial counterparts (e.g., Jiménez-Moreno et al., 2013; Hockun et al., 2016; Ortiz et al., 2021), which is not evident in the lake surface sediment nor in the analyzed plants, only Juncus sp. displaying relatively higher $\delta^{13}C_{n-alk}$ than their ecological counterparts (Figure 3; Table 1). The lack of evidence of less δ^{13} C depletion in mid-chains *n*-alkanes in the lake surface sediment supports terrestrial plant dominance over aquatic plant inputs in modern lake settings. This suggests that caution needs to be taken when using mid-chain n-alkanes without compound-specific $\delta^{13}C$ analyses in sedimentary archives to infer past changes in aquatic plant inputs and lake waters (Hockun et al., 2016; McFarlin et al., 2019). Moreover, it also highlights the importance of using $\delta^{13}C_{n-alk}$ over other indicators, such as P_{aq}, to better constrain aquatic inputs in the lake sediments. Beyond that, the distinct values of $\epsilon_{\rm app}$ between modern plants and lake surface sediment underline the importance of constraining the vegetation source when using specific fractionation factors in sediments (Yang et al., 2011).

5 Conclusion

Our results show that the oro-Mediterranean vegetation cover of the Lake Peixão area preferentially produces longchain *n*-alkanes ($\geq C_{27}$) that control the *n*-alkane content in the lake. This vegetation is dominated by C₃ grasses/herbs and shrubs, two ecological forms that display similar ACL₂₇₋₃₃ and CPI₂₇₋₃₃ values but are clearly distinguished by their

n-alkane isotopic signatures. Shrubs show proportionally higher $\delta^{13}C_{n-alk}$ and δD_{n-alk} values than grasses/herbs, suggesting a congruent physiological response to the common climatic and environmental conditions. The n-alkane C31 is the predominant leaf wax of the study area, produced by some grasses and shrubs, particularly by Erica sp. (heather), a shrub that is stands out as a major source for the total n-alkane pool. The C29 is the second-most abundant and the more equally produced leaf *n*-alkane of the two dominant ecological forms. The *n*-alkanes C₃₃ and C₃₅, although less abundant than other long chains homologs, are also important in the study area since they are particularly associated with cold and drought-tolerant vegetation such as Juniperus sp. Finally, C₂₇ and C₂₅ n-alkanes are mainly linked to aquatic-related grasses/herbs and are inferred to better reflect changes in the water availability of the study area and the isotopic compositions of lake water. However, the aquatic origin of these homologs should be ensured in downcore reconstructions, for instance, performing compound-specific carbon isotopic analysis. The use δD of C_{29} in the lake sediments (δD_{terr}) is interpreted as a robust terrestrial hydrology indicator. Despite the small number of plant species and different ε_{app} among modern plants, the integrated ϵ_{app} value of Lake Peixão's climatic sensor, determined from δD_{terr} ($\epsilon_{C29/MAP} = -129 \pm 12\%$), is in line with the projected values of global compilation studies. Overall, this dataset establishes a baseline for interpreting the leaf wax biomarkers in the region and Lake Peixão sediments, which is a valuable archive to reconstruct regional past hydroclimate changes in the region.

Data availability statement

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding authors.

Author contributions

RS collected the data. RS, ES, LC, and TR produced the data. RS wrote the manuscript. DO, AH, and AR contributed to improving the manuscript. All authors contributed to the discussion and interpretation of the data and provided comments and suggestions to the manuscript.

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Supplementary material

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