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Sequential fatty acid analysis of a peat core covering the last two millennia (Tritrivakely lake, Madagascar): Diagenesis appraisal and consequences for palaeoenvironmental reconstruction

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Abstract

Seven samples from a 1 m long peat core, taken from the centre of the Tritrivakely maar lake (Madagascar) and covering the last 2300 yr, were analysed for their fatty acid (FA) content. Solvent extraction, followed by acid hydrolysis and saponification allowed successive release of “free”, H⁺-labile and OH⁻-labile bound FAs. Downcore variations in the concentration of individual compounds released by the different treatments allowed distinction between the following three major types of input: (i) primary production; (ii) secondary production (that most certainly forms as a result of microbial degradation of primary products) immediately underlying the sub-surface; (iii) a notable bacterial stock that appears later and slowly at depth. The latter input is revealed by the regularly increasing concentration of several H⁺-labile compounds with depth. Sedimentary horizons that were sites for good preservation conditions are indicated by high relative amounts of nearly all the compounds, especially those inherited from the primary as well as from the secondary producers (including the primary consumers). In contrast, depth intervals that have been the site for intensive diagenetic alteration contain few compounds, including those inherited from late consumers.

1. Introduction

Among the most common families of biochemical constituents, lipids and especially fatty acids (FAs) have the advantage of comprising a highly diversified group of structures (carbon backbone plus possible functional groups) and a notable resistance to degradation. Consequently, these compounds are frequently analysed in order to gain information on organic matter (OM) sources and diagenetic transformations in a diversity of sediments, mainly from recent marine and lacustrine environments (Volkman et al., 2000, Zegouagh et al., 2000 and Garcette-Lepecq et al., 2004). In this respect, despite the fact that they appear advantageous targets because of their dominant or even exclusive OM content, peat deposits have received relatively little attention (Lehtonen and Ketola, 1990 and Lehtonen and Ketola, 1993). Nevertheless, recent studies on lipid biomarkers in peat have been performed with the aim of reconstructing variations in higher plant inputs as well of deriving new climatic change proxies (Dehmer, 1993, Farrimond and Flanagan, 1995, Ficken et al., 1998, Nott et al., 2000, Duan and Ma, 2001, Pancost et al., 2002, Pancost et al., 2003 and Xie et al., 2004).
When studying FAs organic geochemists classically paid attention either to the “free” fraction or to “bound” compounds commonly released by saponification, without any decisive argument to justify their choice of one or other fraction. It is now well established that more detailed information can be obtained by a study of the various pools of FAs that can be released successively by different chemical treatments (Barnes and Barnes, 1978, Albaigés et al., 1984, Nishimura and Baker, 1987, Amblès et al., 1987, Zegouagh et al., 1996 and Stefanova and Disnar, 2000). Among the “new” treatments, acid hydrolysis has been recognised as being capable of specifically releasing bacterial FAs initially linked to other cell constituents through amide bonds (Rietschel et al., 1972, Mendoza et al., 1987a, Mendoza et al., 1987b, Goossens et al., 1989a and Goossens et al., 1989b). In this paper, we present the results of a three step FA sequential extraction approach carried out on seven samples taken from a one metre peat core representing the uppermost part of the sedimentary fill of the Tritrivakely maar lake in Madagascar. This sequence, which constitutes a ca. 2300 yr long sedimentary record, has previously been the object of a detailed petrographical and geochemical study that allowed distinction of two main accumulation stages during the time in question, the transition from one to the other being most probably driven by a notable regional climate change (Laggoun-Défarge et al., 1999, Bourdon, 1999, Bourdon et al., 1997 and Bourdon et al., 2000). The present work was undertaken to obtain supplementary information on OM inputs and diagenesis through FA analysis. To this end, we applied a new strategy based on sequential FA analysis and comparison of individual component concentration vs. depth.

2. Materials and methods

2.1. Setting and sampling

The site and sedimentary section have been described by Bourdon et al. (2000). Tritrivakely Lake (19°47 min S, 46°55 min E, 1778 m) is located in the central part of the volcanic Ankaratra Plateau (Madagascar). It is presently a maar about 600 m wide and 50 m deep. As revealed by piston core examination, sedimentary infill represents a more than 40 m thick accumulation of lacustrine and paludal sediments that probably record the last climatic cycle (Gasse et al., 1994, Sifeddine et al., 1995 and Gasse and Van Campo, 1998). The uppermost 3 m of sediment consist of peat accumulated during the late Holocene (Burney, 1987 and Gasse et al., 1994).

The regional climate, controlled by latitude and topography, is of the mountain tropical type (mean annual precipitation about 1600 mm/yr; mean annual temperature about 16 °C) with a warm and wet summer and a dry and cool winter (Chaperon et al., 1993). The lake has no surface outlet. Changes in water level are, therefore, directly controlled by the precipitation/evaporation ratio, with considerable variation (e.g., the lake was almost dry in Nov. 1992, while the maximum water level reached 0.6 m in Jan. 1992 and up to 2 m after an extremely rainy season in May 1994). The water is fresh, oligotrophic (conductivity about 20.6 μS cm⁻¹), slightly acidic (pH 5.5–6.3) and has a ¹⁴C activity of 98.14%, corresponding to an apparent age of about 1000 yr (Gasse et al., 1994). The water body is presently colonised by aquatic and emergent plants. The flora consists of Cyperaceae (Cyperus madagascariensis dominant, and Cyperus papyrus and Heliocharis equisetina) with some Poaceae species like Leersia hexandra. Rice and corn are cropped on the slopes of the crater.
The present study is concerned with the uppermost metre of peat which has accumulated since 2300 yr BP in the maar (Bourdon et al., 1997). The core was taken in 1994 near the centre of the marsh. The sections analysed (1–4 cm thick) were taken at 0–1, 2–5, 9–12, 19–22, 30–34, 59–60 and 96–99 cm depth. C and N determination was performed using combustion with a Leco® CNS 2000 analyser on an aliquot of the crushed and dried samples.

2.2. Sequential lipid extraction

Sequential lipid extraction was performed according to the protocol applied by Stefanova and Disnar (2000). Briefly, a “reversed” sequential extraction scheme was applied, where acid attack preceeded saponification.

2.2.1. Isolation of “free” extractable lipids

The lyophilised samples (1–2 g) were soaked overnight in acetone–pentane (1:1), with a 1:50 w/v sample:solvent ratio. The mixture was sonicated (2 × 5 min) at ambient temperature. The extracts were separated by filtration and the residues were carefully washed with acetone–pentane. The extract and the washes were combined and the solvent was removed under vacuum.

2.2.2. Isolation of H⁺-labile lipids

The solid residues from the previous step were transferred to 25 ml Pyrex® tubes. Methanol (10 ml) and 3 N HCl (10 ml) were added. The tubes were tightly closed under vacuum and left overnight in an oven at 80 ± 2 °C.

The insoluble residues were isolated by filtration and washed with distilled water. The filtrates were combined, concentrated under reduced pressure to about 10 ml and extracted with diethyl ether (2 × 20 ml). The soluble portions were dried with anhydrous Na₂SO₄, filtered and finally concentrated for chromatographic separation.

2.2.3. Isolation of OH⁻-labile lipids

The solid residues from the previous treatment were transferred to Pyrex® tubes and were saponified with 1.5 N NaOH. After addition of the NaOH solution, the tubes were shaken vigorously, closed and left for 48 h in an oven at 55 °C. After cooling, the reaction products were neutralised with dilute HCl. The insoluble residue was separated using filtration and washed with water and diethyl ether. The organic phase was separated from the aqueous solution. The latter was extracted with diethyl ether until extracts were colourless. Then, the organic phases were combined, dried and concentrated under reduced pressure.

2.2.4. Fatty acid analysis

All the lipid extracts were separated into neutral and acidic portions using column chromatography on KOH-impregnated silica gel columns (McCarthy and Duthie, 1962). The acidic fractions, eluted by diethyl ether–HCOOH (9:1, v/v) were concentrated and esterified with diazomethane (Lombardi, 1990). Tetradecene
(cis + trans) was added as an internal standard prior to gas chromatography (GC) and combined gas chromatography–mass spectrometry (GC–MS).

The methylated fractions were analysed using a Perkin–Elmer XL Autosystem® gas chromatograph equipped with a CP sil 5 CB column (0.22 mm × 30 m; 0.25 μm film thickness), a flame ionisation detector (300 °C) and a split/splitless capillary injector maintained at 300 °C and used in the splitless mode (valve reopened one min after injection). After 1 min hold at 50 °C, the oven temperature was increased from 50 to 120 °C at 30 °C/min and then from 120° to 300° at 5°/min; this final temperature was maintained for a period of 1 h. The carrier gas was helium.

GC–MS analysis was performed with a Varian 300 chromatograph connected to an Ion Trap ITD 800 (Finnigan Mat) by a 2 m capillary interface heated to 300 °C. Other GC conditions were identical to those described above. Operating conditions for the ion trap were: temperature, 220 °C; ionisation energy, 80 eV; 1 scan/2 s from 50 to 500 amu.

3. Results and discussion

3.1. Bulk composition of samples and acid fractions

The total organic carbon (TOC), between 43.1% and 49.6% over the whole section, expresses a paucity of mineral matter. More important variations affect the N content, since the C/N ratio value reaching 19.6 in the upper level, remains between 12 and 15 below this. The three FA fractions were named as follows: “free”, for directly solvent-extracted fractions; “H⁺-labile” and “OH⁻-labile” for the fractions released after acid treatment and saponification, respectively. A general description of the FA distributions in the different fractions is given in Table 1. The changes in total FA content with depth in the three fractions are shown in Fig. 1. These profiles are further discussed.
Table 1. Acids series distributions: carbon range, dominant compound and maximal content (μg/g Corg)

As illustrated in Fig. 2 for the upper- (0–2 cm), mid- (30–34 cm) and lower core (96–99 cm) samples, respectively, n-saturated acids and especially n-C16:0 and n-C18:0 strongly dominate compound distributions. The contribution of different series of acids to the “free”, H⁺-labile and OH⁻-labile acid fractions and their changes with burial depth are shown in Fig. 3. The main observation is that FA indicators of high biological activity (namely, branched and 3-OH FAs) were released after the acidic and basic treatments.

Hopanoic acids were represented by only one compound. This was tentatively identified as a C24 17,21-seco-hopanoic acid from the spectrum that showed a molecular ion at 374 amu, loss of a COOCH₃ group (M⁺ – 59; m/z 315; 100%) and intensive ions at m/z 191 and m/z 177. The presence of a seco-acid in recent sediments is not surprising. In the Miocene Clarkia deposit, Huang et al. (1996) identified a suite of seco-terpenoic acids having ursane, oleanane and friedelane skeletons, all attributed to higher plants on the basis of carbon isotopic data. In our study, the possible seco-hopanoic acid could derive from a previous hopanoic acid by microbial reworking. It is, however, slightly suspicious that the quantity of this acid appears neither influenced by extraction treatment nor by diagenesis (i.e., depth) since, except at the surface where it is present in slightly higher amounts (ca. 13–16 μg/g Corg), it represented ~2% of all extracts, at all levels (Table 1).
Fig. 2. Distributions of $n$-fatty acids ($\mu$g/g C$_{org}$) in “free” and “bound” acid fractions of upper-, mid- and downcore samples.
3.2. FA variation with depth

The following discussion on the biological origin of FAs and their variation with depth is focussed mostly on the analysis of the individual concentration profiles. Information deduced from their structure is often disputable because of the general multiplicity of possible source organisms.
3.2.1. Variation in total quantities of “free” and bound FAs

The quantities of FAs represent the sum of the compounds identified using GC–MS and quantified using GC. Their variation with depth is presented in Fig. 1. The first notable point is the relatively high amount of “free” FAs in the surface sample. This demonstrates two essential facts: first, rapid hydrolysis of the ester (and amide) bonds of the various combinations under which FAs exist in living organisms and second, a temporary accumulation of “free” FAs at the surface of the sediment column. This second point further implies a faster release of FAs from their original combinations than their consumption by microorganisms. This latter phenomenon is nevertheless very rapid as proven by the abrupt decrease recorded by “free” FA concentrations between the surface and the immediately underlying level. Despite the relative speed of this consumption process, notable quantities of free FAs still exist at depth. These form a nearly continuous background all along the section. The stock that occurs at depth might either represent a transitory pool of compounds progressively liberated by hydrolysis of esters and/or amides and waiting to be degraded, or to a stock of FAs freed rapidly during early diagenesis and preserved as a result of the rapid disappearance of any noticeable biological activity at depth.

The initially bound acids released by H+ and by OH− treatments display different profiles but show notable FA contents at almost all sedimentary levels. The persistence, if not the increase in the amounts of bound acids all along the profile, is consistent with an absence of degradation of bound (as well as of “free”) acids at depth, or more precisely below 10 cm. These various considerations suggest that, once past the phase of intensive degradation at the surface, the FAs are likely to preserve information on their environment of deposition during further burial. Thus, the relatively high amounts of “free” and OH−-labile FAs at 30–34 cm and 96–99 cm depth either denote particularly important inputs of these compounds and/or particularly favourable preservation conditions during deposition and early diagenesis. For the sample taken at 33–34 cm, the former hypothesis is supported by independent observations revealing that the corresponding sediment was deposited under well oxygenated waters that were the site of a rather important production of diatoms and of the alga Botryococcus braunii (Bourdon, 1999 and Bourdon et al., 2000).

3.2.2. “Free” FA variation with depth

The “free” FA fraction is dominated by low molecular weight linear saturated acids, i.e., n-C16 and n-C18 (Fig. 2 and Fig. 3). To a lesser extent these compounds are accompanied by their unsaturated homologues, n-C16:1 and n-C18:1 (Fig. 4a and b). All these compounds are abundant in all living organisms (Matsuda and Koyama, 1977) but most particularly in phytoplankton (Cranwell, 1978, Cranwell, 1982 and Robinson et al., 1984) and even in bacteria (Holton and Blecker, 1970). They are accompanied by a relatively minor contribution from even numbered higher molecular weight saturated homologues, n-C20+ that, in most cases are dominated by the n-C24 component (Fig. 2). The latter are particularly abundant in terrestrial higher plants (Cranwell, 1978 and Cranwell, 1982). The “free” fractions also contain small proportions of C15 and C16 branched iso- and anteiso-compounds, namely i-C15, ai-C15, i-C16 and ai-C16, that are usually considered as being derived from bacteria (Poupet, 1989 and Volkman et al., 1998).
Fig. 4. (a,b) Downcore correlations of some fatty acids (μg/g Corg) in the “free” acid fraction.
At the peat surface, the amounts of linear saturated FAs record a much stronger decrease (Fig. 4a) than their unsaturated counterparts despite the fact that the former are usually considered as more refractory with respect to biodegradation. This contradictory observation might tentatively be explained by a comparable rate of release of both types of compounds from their original lipid combinations, followed by a preferential consumption of the “free” unsaturated FAs that are more easily assimilated than their saturated counterparts. The lower abundance of the transitory stock of the latter necessarily entails a lower decrease in their amount during early diagenesis [i.e., between the surface (0–1 cm) and the immediately underlying sub-surface level (2–5 cm depth)].

The concentration profiles of the “free” branched FAs thought to be derived from bacteria, show clear similarities, the profiles of i-C15 and ai-C15 being even strictly comparable with high compound amounts at ca. 32 and 98 cm depth (Fig. 4b). In contrast to i-C15 and ai-C15, i-C16 and ai-C16 are both present at the surface but do not maximise at the same depth intervals. All these observations indicate that there are at least three major sources (or levels of major production) for the four branched compounds: at the surface (i- and ai-C16), at ca. 20 cm (i-C16) and at ca. 32 cm (ai-C16, i-C15 and ai-C15). Thus, the organisms that synthesised the ai-C16 obviously thrived at ca. 32 cm with those that synthesised the i-C15 and ai-C15, whereas the conditions that prevailed during deposition of the sedimentary material presently at ca. 98 cm depth were unfavourable for the ai-C16 producer(s).

Whereas all the already considered “free” FAs seem to have given rise to (an) accumulation(s) under specific medium conditions, the saturated acid n-C24 (taken as representative of all the even numbered n-C20+ species) records first a drastic lowering in concentration just below the surface, then shows a downward regularly increasing, but progressively flattening concentration curve. This increase might be due to a progressive release of n-C24 by hydrolysis or even to its synthesis (very unlikely since this would entail a compound classically attributed to higher plants also being produced by microorganisms). The high abundance of n-C24 FA in the bound form at the base of the section does not allow us to favour either of these two hypotheses.

3.2.3. OH⁻-labile FA evolution with depth

In contrast to the order of the extraction steps, we comment first on the evolution of the FAs liberated by classical saponification and supposed to arise from widespread lipid esters, then, we examine the data on FAs obtained by acid hydrolysis which are supposed to be specific for amide-bound compounds of bacterial origin (Mendoza et al., 1987a and Mendoza et al., 1987b).

All the major compounds that are present in the “free” state, namely the n-C16 and n-C18 FAs, their monounsaturated counterparts n-C16:1 and n-C18:1 and their higher molecular weight homologues such as n-C24, also exist in the fraction released by saponification (Fig. 5a). The high abundance of all these compounds at the sediment surface is fully consistent with an inheritance from organisms that grew in the marsh. As already assumed in the previous section, the important decrease in the abundance of these compounds between the surface sample and the immediately underlying level certainly results from microbial attack. At greater depth, the presence
of saturated und unsaturated linear FAs in rather high concentration (especially at ca. 98 cm) reveals episodes of high production and/or preservation of these compounds.
Fig. 5. (a,b) Downcore correlations of some fatty acids (μg/g Corg) in OH⁻-labile acid fraction.

Following the same interpretation as above, the strong decrease recorded by the branched acid \( i-C_{16} \) immediately below the surface (Fig. 5a) suggests it was also inherited from a primary producer, or more precisely from bacteria that thrived in the marsh waters, since branched FAs are generally supposed to be specific to such organisms (Poupet, 1989 and Volkman et al., 1998). The profile of this \( i-C_{16} \) compound parallels that of the \( n-C_{24} \) species. In other respects, it must be stressed that the concentration profile of the \( n-C_{24} \) released by saponification is totally independent of that of the same compound in the free state (Fig. 4a). The absence of such a relationship makes the question of the origin of the “free” \( n-C_{24} \) component even more puzzling.

The compounds that appear just below the surface (i.e., at 2–5 cm depth; Fig. 5b) are especially those which are usually considered as deriving from bacteria, such as the \( i-C_{15}, ai-C_{15} \) and 3-OH-C\(_{14} \) acids. These three compounds show similar profiles, with high values at three different levels that strongly suggest that they have a common
origin or simply that they derive from microorganisms that prosper under similar conditions.

The di-C_{16} and di-C_{18} αω-dicarboxylic acids have nearly identical profiles (Fig. 5b) as would be reasonably expected for compounds that most probably have the same source, namely higher plant cuticles (Kolattukudy et al., 1976). These diacids appear in the sub-surface (i.e., at ca. 5 cm) but disappear almost completely afterwards, maybe as a result of rapid degradation. Nevertheless, they are both abundant at 98 cm depth, thus probably providing additional evidence that this level has been the site of good organic matter preservation (Bourdon, 1999).

3.2.4. H⁺-labile FA evolution with depth

The dominant compounds in this fraction are the lower molecular weight saturated acids, n-C_{16} and n-C_{18}, as well as the unsaturated n-C_{18:1} acid. The profiles of these three compounds appear relatively well correlated, but with slightly higher values for the n-C_{18} species at ca. 32 cm than at ca. 98 cm (Fig. 6a). These three profiles can probably be explained by the fact that all these very common FAs can be derived from multiple sources. The n-C_{24} and n-C_{16:1} profiles (Fig. 6b) show first a strong decrease attributable to the rapid degradation of a stock inherited from primary producers and second a rather regular increase with increasing depth that leads us to hypothesise an in situ production of these compounds at depth. The other components of the fraction, i.e., mostly the C_{15} and C_{16} branched acids and the 3-OH-C_{16}FA, are more abundant at the sub-surface than at the surface, consistent with a secondary (microbial) origin. The notable decline that immediately follows their production stage can be interpreted as consequential to the immediate degradation of a notable part or even the totality of the primary stock. Below, i.e., beyond the first 10 cm, the profiles of the i-C_{16}, 3-OH-C_{16} and di-C_{16} compounds resemble those of the n-C_{16}, n-C_{18} and n-C_{18:1} acids discussed above, but with a more pronounced increase with increasing depth (Fig. 6b). In fact, these profiles, which seem to tend to an asymptote at depth, might well evoke an efficient but progressively declining in situ (i.e., bacterial) production. The putative bacteria might be dead or alive, and in the second case might be active or only subsisting in a latent state (Given et al., 1983 and Chasar et al., 2000).
Fig. 6. (a,b) Downcore correlations of some fatty acids (μg/g C<sub>org</sub>) in H<sup>+</sup>-labile acid fraction.
The profiles of the $i\text{-C}_{15}$ and $ai\text{-C}_{15}$ acids, which are well correlated (Fig. 6b), denote a production just below the surface and additional episodes of compound production or rather good preservation at 20 and 60 cm. This pattern, just the reverse of the one produced by the same acids released by saponification, suggests a multiplicity of bacterial sources that compete one with an other depending on conditions (and/or timing of diagenesis).

4. General discussion

In the following, the possibility of incorporation of some FAs into newly formed compounds, namely humic substances, is not discussed. Such a possibility should not be omitted but first, it can be considered as a kind of preservation mechanism and second, at least for the less common compounds, it is contradicted by a complete lack of FA at some levels, in one or several fractions.

As assumed in the previous sections, the most reliable indications of inherited FAs should be found in the $OH^-$-labile fraction since this one should comprise all the compounds freed from different kinds of esters that constitute the most important lipid fractions of living organisms. A major difference between all the compound concentration profiles drawn for that fraction can, at least to a first indication, be seen as typical evidence of primary or secondary production. Thus, compounds present in high proportion at the surface (e.g., $n\text{-C}_{16}$ and $i\text{-C}_{16}$; Fig. 5a) can be considered as being inherited from the primary production. In contrast, the compounds that are nearly totally absent at the surface but which are abundant in the immediately underlying level might be revealing an in situ secondary production (namely 3-OH-$C_{14}$, $i\text{-}$ and $ai\text{-C}_{15}$; Fig. 5b).

All the compounds considered as issuing from the primary producers are also present in high relative proportions in the “free” fraction, including $i\text{-C}_{16}$ and even its $ai$ isomer (Fig. 4b) that was undetectable in the $OH^-$-labile fraction. This is fully consistent with the release of “free” FAs presumably by (microbial) ester and/or amide bond hydrolysis in the very early stages of diagenesis. The only compounds attributed to secondary production in the $OH^-$-labile fraction and that also exist in the “free” fraction are the $i$- and $ai\text{-C}_{15}$ acids. However, these compounds were not observed in the free state at the subsurface as in $OH^-$-labile fraction but only at depth (i.e., 32 and 98 cm). This might indicate a rather slow release of these compounds from their original combinations, i.e., in a more general way, much slower hydrolysis occurring in the sediment than in the surface waters as a result of much reduced microbial activity. This is in a certain way confirmed by the absence from the “free” fraction of other compounds that were also found at the subsurface in the $OH^-$-labile fraction and that were thus supposed to be derived from secondary production, namely OH-FAs and di-acids. The polyfunctional nature of these compounds might at least partly explain their rather more difficult release than for simple FAs.

The abundance of all primary as well as secondary products at ca. 98 cm provides evidence for very good OM preservation conditions during the deposition of this section. In contrast, the sample from ca. 32 cm depth is only rich in secondary products, i.e., $i$- and $ai\text{-C}_{15}$, plus 3-OH-$C_{14}$. In fact, it has previously been shown (Bourdon, 1999 and Laggoun-Défarge et al., 1999) that this level was particularly rich in mucilage, an extracellular polysaccharidic secretion (EPS) that can be produced
not only by plant roots, but also by bacteria and fungi (Leppard et al., 1977, Decho, 1990 and Défarge et al., 1996). A notable fungal activity could be consistent with (i) the absence of di-acids and 3-OH-C_{16} of supposed bacterial origin and (ii) notable lignin degradation (Bourdon, 1999 and Bourdon et al., 2000).

Except for the presence of minor amounts n-C_{24}, di-C_{18} and i-C_{16}, the general paucity of FAs in the OH−-labile fraction of the sample taken at ca. 22 cm suggests that it has been the site of more extensive degradation than at 32 cm. Even more extensive degradation might have occurred at 10 and 60 cm where almost all compounds are missing from the “free” (Fig. 4a and b) as well as from the OH−-labile fraction (Fig. 5a and b), with only one notable exception for the n-C_{24}FA (and its C_{20+} homologues). For this compound, a significant release at the surface (certainly followed by rapid consumption) is followed by what seems to be a very slow release with increasing depth. In the absence of a much more plausible explanation this regular pattern was previously attributed (Section 3.2.2) to possible in situ microbial production.

Acid hydrolysis is supposed to preferentially cleave the amide bonds that are important in bacterial lipids. Several of the compounds released in this way, and especially most of the more abundant ones, show a more or less regular increase with depth, namely n-C_{16}, n-C_{18}, i-C_{16}, 3-OH-C_{16} and di-C_{16} (Fig. 6a). Irregularities in this trend can be attributed to the influence of non-microbial sources. For example for the n-C_{16} and n-C_{18}, the most ubiquitous of the considered compounds, the high proportions in the uppermost levels can simply be explained by their partial release from lipid esters abundant in all organisms, especially higher plants. A comparable explanation might hold for i-C_{16}, 3-OH-C_{16} and di-C_{16} acids (Fig. 6a) but with sources other than the previous compounds since, contrary to these latter ones, they are more abundant in the subsurface sample than in the surface one. The notable decrease in their quantity recorded below the subsurface level at ca. 2 cm depth and the onset of a regular increase from 10 cm depth, suggests that they are produced during two successive stages and at different rates, i.e., rapidly during very early diagenesis (at about 2 cm) and slowly more deeply. The question as to whether these two production episodes are due to the same or different organisms remains unclear. However, the progressive flattening in the compound concentration vs. depth curves indicates that the inferred late bacterial production slows down progressively with increasing time.

Once the stage of surface release is past, the n-C_{24} and n-C_{16:1} acids provide a completely different pattern from the previous compounds since the amounts of these two in the H^{+}-labile fraction seem to increase progressively with depth (Fig. 6b). The interpretation of such a pattern is not straightforward. It can tentatively be assumed that it could result from a competition between the release of the two considered compounds from combined state and their subsequent consumption, the latter proceeding at a faster rate in the upper levels than at depth, consistent with a progressive decline in microbial activity with time. Even if the assumed pattern is not so clear for the n-C_{24} component, the proposed interpretation is strengthened by the indication of a notable but declining production of this compound at depth, as provided by the “free” fraction (Fig. 4a). In contrast to the already considered compounds in the H^{+}-labile fraction, the i-C_{15}, ai-C_{15}, 3-OH-C_{14} and ai-C_{16} acids display very irregular patterns (Fig. 6b) The 3-OH-C_{14} acid is present in notable proportion at the surface but also at 10 cm where the absence of any compound in
the “free” as well as in the OH−-labile fraction suggests extensive degradation. If the profile of the ai-C_{16} acid is little informative, it is worth mentioning that the i- and ai-C_{15} acids are relatively abundant at 2 cm depth, as in the OH−-labile fraction, but below this, have completely opposite patterns, i.e., high levels at ca. 22 and 60 cm (Fig. 6b) against maxima at 32 and 98 cm in the OH−-labile fraction (Fig. 5b). These observations provide clear evidence that these two branched isomers certainly exist under different combinations in microorganisms and additionally that these organisms (whatever they are) do not produce the i- and ai-C_{16} higher homologues.

Following the hypothesis of a late and slow third production, if the profile of the 3-OH-C_{16} FA at depth is considered as a bacterial growth curve, the “half-growth depth” occurs at about 20 cm, thus entailing a half-growth time of about 500 yr if the accumulation rate remained constant during deposition of the section presently between 0 and 32 cm. If this interpretation is correct, it remains to be seen whether such a late bacterial growth is common in peat deposits or is due to local (normal or fortuitous) environmental factors. The hypothesis of an important and widespread microbial production at depth in peat deposits is supported by the recent determination of more than 20 times higher amounts of amino acids at depth in a Swiss peat than at the surface (Comont and Gautret; personal communication).

5. Conclusions

1. High amounts of “free” and OH−-labile FAs (saturated and unsaturated n-FAs, plus i-C_{16}) at the surface of the peat sequence and low amounts in the immediately underlying level indicate rapid hydrolysis of lipid esters inherited from the primary production, followed by slightly slower consumption.

2. A whole range of OH−-labile FAs (diacids, i- and ai-C_{15}, 3-OH-C_{14}) that are almost or even totally absent from the peat surface, but are abundant in the immediate sub-surface before decreasing just below, denote in situ secondary production and subsequent decay. All these compounds are totally missing from the “free” fraction of the surface sample.

3. Once past the upper levels where FA dynamics are very active in response to primary inputs, secondary production and active diagenesis, most of the H+ -labile FAs attest to an important but late in situ microbial production that slows down progressively with increasing depth. The latter can be considered as a third production.

4. In contrast, compounds such as i-C_{15} and ai-C_{15} acids also released by acid hydrolysis and which are abundant in the immediate sub-surface but vary very irregularly at depth, might be inherited from consumers acting as secondary producers. In fact, these compounds maximise at levels (ca. 30 and 60 cm) containing low amounts of “free” and OH−-labile FAs normally inherited from the primary producers.

5. Good OM preservation conditions (e.g., at ca. 98 cm depth) are indicated by high relative amounts of nearly all compounds, especially in the “free” and OH−-labile fraction (i.e., inherited from the primary as well as from the secondary producers, consumers included).
6. The comparison of compound concentration profiles permits recognition of whether compounds are related to each other or not, i.e., whether they have the same source (e.g., i- and ai-C15), or are independent.

7. Active molecular dynamics during diagenesis do not allow the drawing of reliable geochemical interpretation for isolated recent geological samples (sediments or soils) after FA analysis. In other words, as formulated in a previous study on lake sediments (Stefanova and Disnar, 2000), extensive diagenesis can provoke the disappearance of most fatty acids and thus nearly all information on the contributing organisms except the latest consumers-producers, i.e., methanotrophs.

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