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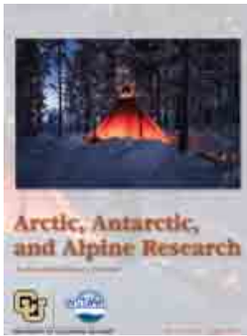
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Terrestrially Derived *n*-Alkane δ D Evidence of Shifting Holocene Paleohydrology in Highland Costa Rica

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Abstract

A previous study of carbon isotopes in the sediments of a glacial lake in Costa Rica led to the hypothesis that changes in the migration of the intertropical convergence zone (ITCZ) over the course of the Holocene significantly affected the hydrology of the surrounding high-elevation páramo ecosystem. This hypothesis was based on millennial-scale changes in terrestrial *n*-alkane carbon isotope ($\delta^{13}\text{C}$) values in a sediment core from Lago de las Morrenas 1, a tarn on the Chirripó massif of the Cordillera de Talamanca. Here we present terrestrial *n*-alkane hydrogen isotope (δ D) data, a more direct proxy of ecosystem drought stress, from the same core. These new data support the previous hypothesis and confirm that the effects of millennial-scale ITCZ dynamics in the circum-Caribbean region were not restricted to tropical lowlands. In southern Central America, these dynamics may have played a fundamental role in millennial-scale fire dynamics in high-elevation páramo ecosystems.

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Introduction

Stable oxygen isotope analyses of carbonates preserved in lacustrine sedimentary sequences have provided a wealth of information regarding paleoclimate variability over a variety of time scales. The development of compound-specific hydrogen isotope (δ D) analysis of biomarkers has widened opportunities to develop paleohydrological records based on lake sediment archives where carbonates are absent or poorly preserved. A widely used class of biomarkers of terrestrial vegetation present in lake sediments are long-chain ($\geq\text{C}_{27}$) *n*-alkanes with odd carbon number chain lengths, which predominantly originate from epicuticular leaf waxes (Eglinton and Hamilton, 1967). Multiple studies have documented the close relationship between terrestrially derived *n*-alkane and meteoric water δ D values along latitudinal (Sachse et al., 2006) and elevational (Bai et al., 2011) transects in modern ecosystems. Furthermore, these studies and others have shown a significant evapotranspiration influence on terrestrial vegetation biomarker δ D values. Over a wide range of modern ecosystems, Sachse et al. (2006) and Polissar and Freeman (2010) documented a consistent enrichment, or apparent fractionation (ϵ_a), in terrestrial *n*-alkane δ D values over values expected based on meteoric water δ D values alone. They attributed this enrichment to soil water evaporation prior to uptake by terrestrial vegetation and biosynthesis into leaf waxes. Transpiration, in contrast, appears to play a minimal role in isotopic enrichment in *n*-alkane δ D values, particularly in grasses (McInerney et al., 2011). Sachse et al. (2006) identified this evaporative enrichment of meteoric waters prior to biosynthesis by terrestrial organisms as a potential proxy for ecosystem evapotranspiration, and Tierny et al. (2008) and Niedermeyer et al. (2010) subsequently employed it as an indicator of paleohydrologic variability on millennial timescales in tropical Africa.

Lane et al. (2011) presented bulk and compound-specific stable carbon isotope ($\delta^{13}\text{C}$) data spanning the Holocene from Lago de las Morrenas 1 (LM1), a glacial lake in highland Costa Rica. The general pattern of high $\delta^{13}\text{C}$ values during the early Holocene,

shifting to more negative $\delta^{13}\text{C}$ values during the middle Holocene, and then returning to relatively higher $\delta^{13}\text{C}$ values during the late Holocene, led Lane et al. to speculate that the $\delta^{13}\text{C}$ *n*-alkane record reflected drought stress in vegetation surrounding the lake and thus may track millennial-scale changes in circum-Caribbean precipitation. In this study, we test this interpretation by analyzing the more direct terrestrial *n*-alkane δ D proxy of ecosystem evapotranspiration in samples from the same LM1 sediment core.

We also compare millennial-scale variations in the LM1 compound-specific δ D values to a high-resolution macroscopic charcoal record from the same lake (League and Horn, 2000) to assess the potential role that precipitation variability may play in fire regimes of high-elevation ecosystems of the neotropics. League and Horn (2000) proposed that millennial-scale variations in charcoal deposition were related to increasing anthropogenic activities and/or precipitation variability over the Holocene. More specifically, League and Horn hypothesized that notably low charcoal influx during the middle Holocene might indicate greater precipitation (lower evaporation/precipitation ratios) between \sim 7000 and 4000 cal yr B.P. However, this hypothesis was speculative because at the time no reliable proxies of precipitation had been established for the LM1 sediment record. Our analysis of the compound-specific δ D proxy, one of the few precipitation proxies that can be extracted from non-carbonate bearing sedimentary records, should provide a test for this hypothesis.

Study Site

LM1 (3477 m, 9°29'40"N, 83°29'14"W) is located on the north flank of Cerro Chirripó (3819 m; Fig. 1), the highest peak in the Cordillera de Talamanca and in all of Costa Rica. The lake is one of several dozen glacial lakes on the Chirripó massif and is surrounded by some 5000 ha of neotropical páramo vegetation (Kappelle and Horn, 2005). The Chirripó páramo is dominated by the dwarf bamboo *Chusquea subtessellata* Hitchc., which reaches cover values of 60% or higher over wide areas (Kappelle, 1991)

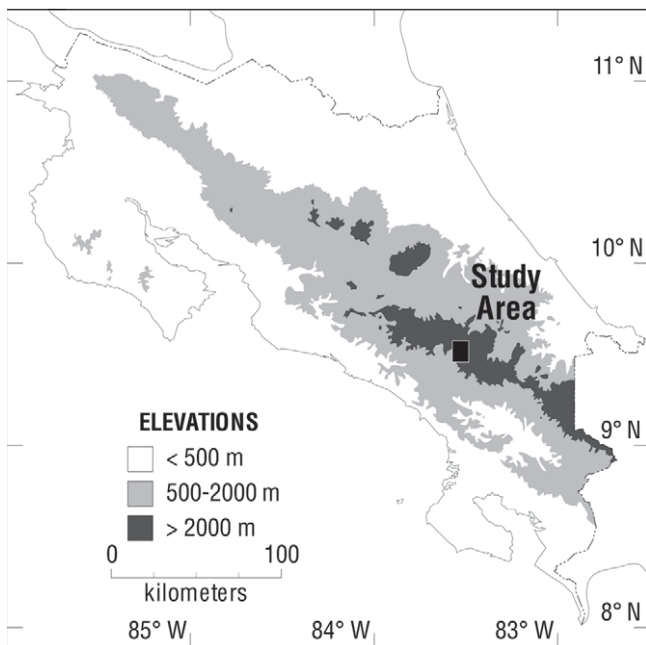


FIGURE 1. Location of the Chirripó massif (“study area”) in the Cordillera de Talamanca of southern Costa Rica, site of Lago de las Morrenas 1.

and is represented in nearly all terrestrial plant communities other than those developed on rock (Chaverri and Cleef, 2005). Precipitation around LM1 is highly seasonal, with the majority of precipitation (~90%) falling between May and November, when the inter-tropical convergence zone (ITCZ) is positioned over the Central American isthmus. The presence of the ITCZ intensifies convective activity along the slopes of the Cordillera de Talamanca and causes a weakening and rise in altitude of the trade wind inversion (TWI), which can sit well below LM1, especially during the boreal winter (Dohrenwend, 1972; Hastenrath, 1991).

Methods

SEDIMENT CORING AND PREVIOUS PROXY ANALYSES

Parallel sediment cores (core 1 and core 2) were collected from LM1 in 1989 from an anchored platform using a square-rod piston corer for the deeper sediments (>1 m) and a PVC pipe fitted with a rubber piston for the near-surface sediments. The sediments of core 1 are the focus of this study and of the bulk and compound-specific carbon isotope analyses carried out by Lane et al. (2011). All other proxy analyses, including pollen, microscopic charcoal, macroscopic charcoal, diatom, and basic geochemical analyses, were conducted using sediments from core 2 (Horn, 1993; Haberyan and Horn, 1999; League and Horn, 2000). The chronology for LM1 core 1 is based on six AMS radiocarbon dates on bulk sediment (Lane et al., 2011). Radiocarbon dates were converted to calibrated years B.P. (where P. = A.D. 1950) using the CALIB 6.0 computer program (Stuiver and Reimer, 1993) and the data set of Reimer et al. (2004). Sedimentation rates were calculated using the weighted means of the probability distributions of the calibrated ages (Telford et al., 2004), and used to linearly interpolate ages for lake sediment horizons located between the positions of radiocarbon-dated materials.

n-ALKANE EXTRACTION AND ISOTOPIC ANALYSIS

Alkanes were extracted from LM1 core 1 sediment samples (1–16 g dry weight) using an ASE 300 accelerated solvent extraction system (ASE 300, Dionex, California, U.S.A.). Samples were flushed with a mixture of 50% dichloromethane and 50% methanol at 125 °C at a pressure of 1500 psi for 10 min. This procedure was repeated three times, yielding a total organic extract that was dried under high purity nitrogen. The aliphatic fraction was then isolated from the total organic extract by solid phase extraction through a silica-gel column with hexane. Straight chain monomers were further isolated from branched and cyclic compounds using urea adduction.

Compound-specific hydrogen isotope analyses were performed on a continuous flow Finnigan MAT Delta-plus XL mass spectrometer interfaced with an Agilent 6890 GC fitted with a DB5-MS silica column (60 m, 0.25 mm i.d., 0.25 μm film thickness) via a Finnigan GC/C III interface equipped with a pyrolysis reactor at 1430 °C. Injection temperatures were 300 °C and the oven temperature program was 60 °C isothermal for 2 min, 15 °C/min to 170 °C, 4 °C/min to 320 °C, and 320 °C isothermal for 30 min. Hydrogen gas and alkane mixtures (provided by A. Schimmelmann, Indiana University) with known hydrogen isotope compositions were used as standards. The Indiana University *n*-alkane mixture was injected following every fourth sample to monitor precision of the instrument. The standard deviation of these standard analyses over the course of this study was less than 5‰. ³H factors were calculated daily using the Isodat software with pulses of increasing reference gas amount. All samples were run in duplicate. Hydrogen isotopic compositions are reported in standard δ-per mil notation relative to Vienna Standard Mean Ocean Water (VSMOW) where:

$$\delta D = 1000 [(R_{\text{sample}}/R_{\text{standard}}) - 1], \quad (1)$$

where $R = {}^2\text{H}/{}^1\text{H}$.

Results

All *n*-alkanes typically attributed to terrestrial vegetation were present in quantities suitable for compound-specific δD analysis in all but the uppermost sample analyzed (Fig. 2). In this sample, C₃₁ and C₃₃ hydrogen peaks were small and co-eluting with unidentified compounds preventing accurate measurement of the δD compositions (Fig. 2). Shorter-chain alkanes typically attributed to algae (e.g. C₁₇) were insufficiently represented in many samples. The δD values of the C₂₅ and C₂₇ *n*-alkanes show a similar pattern through the sedimentary profile with relatively high δD values at the base of the profile (>425 cm; >7700 cal yr B.P.) that generally decrease towards the middle of the profile (~425–225 cm; ~7700–4200 cal yr B.P.), and then increase at the top of the profile (<225 cm; <4200 cal yr B.P.), with the exception of a slight negative excursion at ~65 cm (~940 cal yr B.P.; Fig. 2; Appendix). The uppermost sample displayed anomalously high δD values for the C₂₅, C₂₇, and C₂₉ *n*-alkanes. The δD values of the C₂₉, C₃₁, and C₃₃ *n*-alkanes show corresponding patterns through the profile, but differ from the C₂₅ and C₂₇ *n*-alkane δD patterns. The C₂₉, C₃₁, and C₃₃ *n*-alkanes display a general pattern of relatively low δD values at the base of the profile (>545 cm; greater than an interpolated age of 11,388 cal yr B.P.), followed by an increase in δD

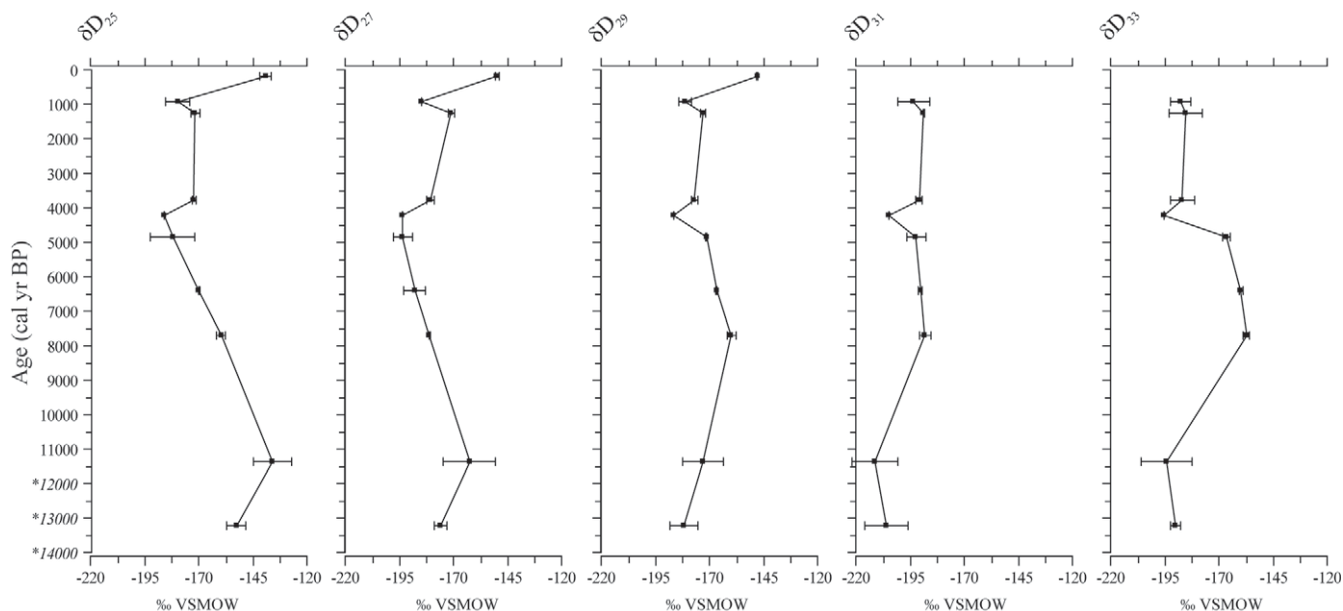


FIGURE 2. Terrestrial *n*-alkane δD data for Lago de las Morrenas 1 core 1. Error bars for the *n*-alkane isotope data represent one standard deviation based on replicate analyses ($n \geq 2$). VSMOW = Vienna Standard Mean Ocean Water.

values, particularly for the C_{33} *n*-alkane (~440 cm; ~7700 cal yr B.P.). The C_{29} , C_{31} , and C_{33} *n*-alkane δD values then display a slight decline upcore until ~190 cm (~3800 cal yr B.P.), above which the values stabilize until ~65 cm (940 cal yr B.P.).

Discussion

n-ALKANE SOURCE AND δD INTERPRETATION

While the C_{29} , C_{31} , and C_{33} *n*-alkanes are typically interpreted as the most reliable biomarkers for terrestrial vegetation, analyses of the *n*-alkane distribution of leaf waxes in modern specimens of *Chusquea* bamboos in the Colombian Andes indicated a disproportionate predominance of C_{25} and C_{27} *n*-alkanes (Boom et al., 2002). This finding, together with the close correspondence of C_{25} and C_{27} δD values through the LM1 sediment profile, lead us to consider that these two *n*-alkanes are originating from the same source and should be representative of the páramo environment surrounding LM1, where *Chusquea subtessellata* dominates in both cover and biomass. The comparatively different δD pattern observed for the longer-chain C_{29} , C_{31} , and C_{33} *n*-alkanes in the LM1 sediments, and their relatively high abundance despite the dominance of *C. subtessellata* in the LM1 watershed (Lane et al., 2011) suggests that these alkanes reflect a different vegetation source. We conjecture that they may derive from montane forest ecosystems at lower elevations, where the waxes are abrasively removed by winds and transported upslope (Schefuß et al., 2003).

Our original interpretation, based on carbon isotopes, was that upslope transport was likely an insignificant source of *n*-alkanes to the LM1 sediments, and probably did not strongly affect stable isotope data (Lane et al., 2011). However, the opposite trends in δD values of the shorter-chain (C_{25} and C_{27}) and longer-chain (C_{29} , C_{31} , and C_{33}) *n*-alkanes in our new data set indicate that the plants producing these biomarkers are not accessing the same water

sources or originating from the same locale. Garcin et al. (2012) found a similar discrepancy in δD values of sedimentary C_{29} and C_{31} alkanes collected along an elevational and latitudinal transect in Cameroon. Garcin et al. were unable to definitively explain this discrepancy, but proposed the possibility that plants producing an abundance of C_{31} alkanes were accessing evaporated lake waters while plants producing an abundance of C_{29} alkanes were more directly accessing meteoric water prior to significant evaporation.

We similarly consider the most parsimonious explanation for the diverging temporal trends in LM1 δD data to be that plants are accessing water of differing δD composition, especially during the early to middle Holocene. However, it is unlikely that such divergent trends in δD values could be produced wholly within the páramo ecosystem surrounding LM1 because of the predominance of shallow, poorly developed soils and high daily solar radiation under cloud-free conditions, which combine to make the high-altitude páramo ecosystem particularly prone to soil water evaporation prior to incorporation by plants (Polissar and Freeman, 2010). Thus, we propose that the differing trends in δD values of shorter-chain and longer-chain *n*-alkanes are the result of significant elevational differences in precipitation and subsequent evaporative enrichment of these meteoric waters prior to uptake by plants. We propose that it is these elevational differences that ultimately produced differing δD temporal trends of *n*-alkanes sourced from lower-elevation forest communities versus those produced in the high-elevation páramo ecosystem. With this in mind, we consider temporal variations in the δD of C_{25} and C_{27} *n*-alkanes isolated from the sediments of LM1 to primarily reflect either changes in the δD composition of meteoric waters entering the Chirripó páramo ecosystem or variations in soil water evaporation in the páramo. Conversely, the δD values of C_{29} , C_{31} , and C_{33} *n*-alkanes more likely represent the δD composition of meteoric waters entering lower-elevation montane forests and any subsequent evaporative enrichment of deuterium prior to uptake by vegetation.

Prior studies have established an important life form effect on ecosystem *n*-alkane δ D signatures (Liu et al., 2006; Smith and Freeman, 2006; Polissar and Freeman, 2010), but available paleoecological data indicate that the páramo ecosystem surrounding LM1 was established soon after deglaciation of the valley \sim 10,000 cal yr B.P. (Horn, 1993), making life form an unlikely driver of the millennial-scale δ D dynamics. Compound-specific $\delta^{13}\text{C}$ data from LM1 indicate a potentially higher predominance of C_4 photosynthetic plants in the late Pleistocene and early Holocene, but the Chirripó páramo ecosystem likely included *C. subtessellata* as a significant component of the landscape at that time and throughout the postglacial period (Horn, 1993; Lane et al., 2011).

The most likely causes of millennial-scale variations in the δ D composition of meteoric waters entering the Chirripó páramo ecosystem prior to soil water evaporation are amount or altitude effects. The amount effect has been documented as the primary driver of meteoric water isotopic composition along the Central American isthmus, with higher rainfall totals yielding more negative $\delta^{18}\text{O}$ and δ D values (Dansgaard, 1964; Lachniet and Patterson, 2002). δ D values of meteoric waters also show an inverse relationship with elevation (Lachniet and Patterson, 2002), a phenomenon primarily driven by changes in atmospheric temperature. While we can assume no significant change in the elevation of LM1 over the time span of this study, we must consider the possibility that isotherms may have shifted up- or downslope in the Cordillera de Talamanca during the Holocene. Available paleoclimate data indicate temperature depressions of 7–8 °C during the Last Glacial Maximum in the highlands of Costa Rica (Hooghiemstra et al., 1992; Orvis and Horn, 2000), but similar reconstructions have not been possible during the Holocene due to a lack of suitable sites and proxies. δ D values display a positive relationship with temperature change, with decreasing temperatures causing a decrease in δ D values of meteoric waters and vice versa (Dansgaard, 1964). Fossil pollen data from LM1 show no clear evidence of shifting treelines during the Holocene (Horn, 1993), leading us to assume that any temperature effects on the δ D composition of precipitation are relatively minor through the Holocene.

Evaporative enrichment of deuterium in soils is considered to be the primary driver of variations in the apparent fractionation (ϵ_a) between meteoric water δ D and terrestrial *n*-alkane δ D in páramo ecosystems (Polissar and Freeman, 2010). The term ϵ_a accounts for the fractionation effects on meteoric waters during biosynthesis and soil water evapotranspiration. Thus, ϵ_a increases significantly with increased aridity as soil water is evaporatively enriched in deuterium pushing it closer to the δ D composition of meteoric waters (Polissar and Freeman, 2010). Polissar and Freeman (2010) reported ϵ_a values from approximately -90 to -110‰ in modern páramo ecosystems in Venezuela. Discounting the anomalous δ D value of the uppermost sample in the LM1 record, and assuming an average C_{27} δ D value of -175‰ for C_{27} *n*-alkanes in the LM1 profile (Fig. 2; Appendix) and a meteoric water δ D composition of about -90‰ (Lachniet and Patterson, 2002), our estimated ϵ_a (-85‰) falls close to the high end of this range. With these variables in mind, we consider the most likely drivers of Holocene variation in C_{25} and C_{27} δ D in the LM1 record to be amount effects and/or evaporative enrichment of soil water that increase ϵ_a values. In either case, increased aridity should increase C_{25} and C_{27} δ D values, while an increase in precipitation should decrease C_{25} and C_{27} δ D values.

The anomalously high δ D values in all of the measured *n*-alkanes in the uppermost sample are difficult to explain (Fig. 2). We find it unlikely that such a high δ D value could have resulted from hydrological changes alone, as no other proxies indicate such severe shifts. Lichtfouse et al. (1994) documented the potential algal origin of some long-chain alkanes following low-temperature diagenesis in sedimentary systems that could explain this anomaly if algal organisms were imparting a very different fractionation on meteoric waters. A wide range of fractionation factors has been reported for algae under similar growth conditions (Zhang and Sachs, 2007). Alternatively, low-temperature diagenetic processes could explain the decreased relative abundance of C_{31} and C_{33} *n*-alkanes in the uppermost sample, although we find it puzzling that this effect would only be obvious in our uppermost sample, when the older sediments have had more time to undergo diagenesis.

Discounting the anomalous upper sample, the pattern of C_{25} and C_{27} δ D variability through the Holocene corresponds well with the terrestrial *n*-alkane $\delta^{13}\text{C}$ record presented by Lane et al. (2011; Fig. 3). This result supports the hypothesis that the observed variations in terrestrial *n*-alkane $\delta^{13}\text{C}$ are a signal of shifting ecosystem drought stress during the Holocene. The overall pattern of relatively high δ D values during the early Holocene, trending toward more negative values during the middle Holocene, and back to more positive values during the late Holocene, indicates millennial-scale shifts in Holocene aridity around LM1—specifically, relatively arid conditions during the early and late Holocene, interrupted by a relatively moist middle Holocene. Our reconstruction from isotopic analyses of the LM1 sediments shows similarities to the Holocene titanium record from Cariaco Basin sediments, which reflects precipitation in adjacent northern South America (Fig. 3) and has been interpreted as a record of millennial-scale ITCZ dynamics in response to the orbital precession cycle (Haug et al., 2001). The Cariaco Basin titanium record indicates a more rapid transition to mesic conditions during the early Holocene than the LM1 δ D record, but the general trend toward more mesic conditions during the middle Holocene is apparent in both records. The slower response of the LM1 record to changes in ITCZ dynamics may be related to more rapid early Holocene warming of northern South America (Cariaco Basin sediment source area) compared to the Central American isthmus, which is surrounded by large water bodies. The high thermal inertia of the Pacific Ocean and the Caribbean Sea relative to the South American landmass may have restricted migration of the ITCZ over the isthmus until sea surface temperatures warmed sufficiently. Similar restrictions in northward ITCZ mobility occur today during El Niño events due to relatively cool sea surface temperatures near the Central American isthmus (Giannini et al., 2001). The tight linkage between modern ITCZ migration and precipitation around LM1 supports the hypothesis that shifting ITCZ migrational dynamics are the primary mechanism responsible for shifting paleohydrology in the circum-Caribbean region during the Holocene.

On Cerro Chirripó, precipitation is tightly linked to ITCZ dynamics via the trade wind inversion (TWI; Lane et al., 2011). When the ITCZ is displaced southwards, convective activity in the region decreases, and the TWI intensifies and lowers, often to an elevation below the highest peaks of the Cordillera de Talamanca (Dohren-

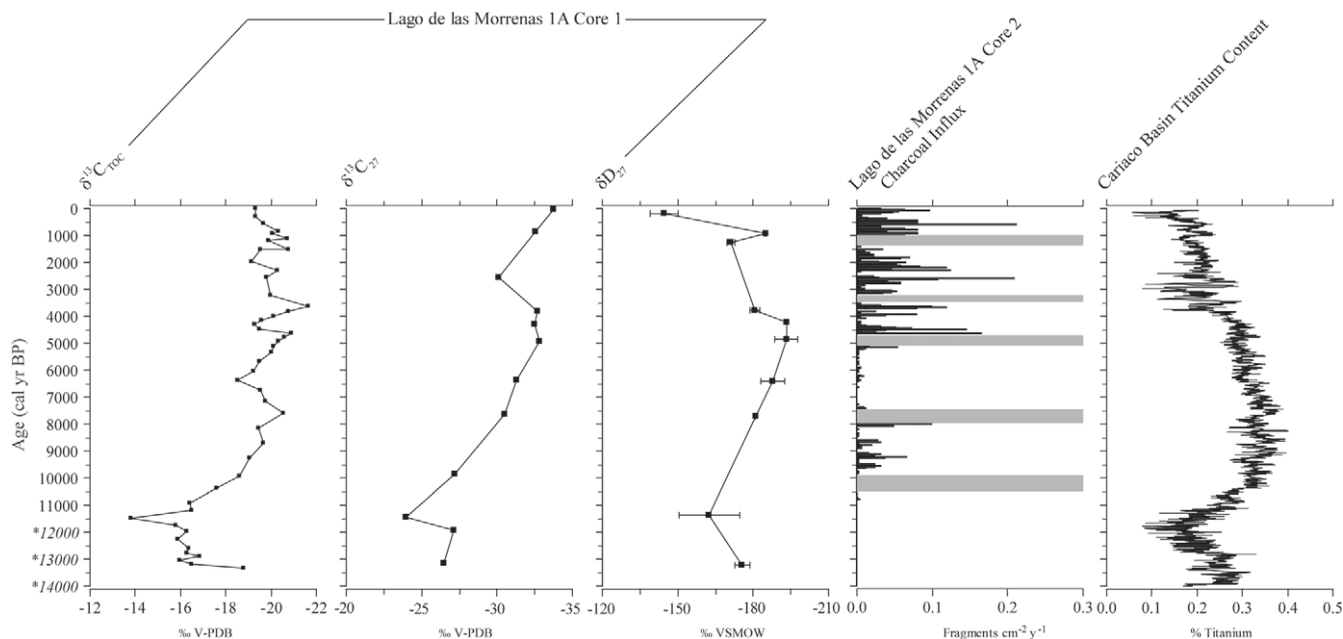


FIGURE 3. Comparison of Lago de las Morrenas 1 core 1 stable isotope data (bulk sedimentary $\delta^{13}\text{C}$, C_{27} *n*-alkane $\delta^{13}\text{C}$, and C_{27} *n*-alkane δD) to Lago de las Morrenas 1 core 2 macroscopic charcoal influx data (League and Horn, 2000), and the Cariaco Basin titanium record (Ocean Drilling Program Site 1002; Haug et al., 2001). The age model for the Cariaco Basin data has been adjusted slightly such that B.P. refers to years before A.D. 1950. Dates extrapolated from the lowest date (11,230 cal yr B.P.) are italicized and marked with asterisks; these should be regarded as estimates for graphing purposes only (see Lane et al., 2011). V-PDB = Vienna–Peedee Belemnite, VSMOW = Vienna Standard Mean Ocean Water.

wend, 1972; Stadtmüller, 1987; Hastenrath, 1991). This inhibits cloud formation and precipitation on the high mountain tops, thereby increasing E/P ratios in the lakes and surrounding watersheds. Conversely, northward migration of the ITCZ promotes convective activity in the region, and weakens and raises the elevation of the TWI, allowing for cloud formation and precipitation on Cerro Chirripó and the other high peaks of the Cordillera de Talamanca.

If ITCZ and TWI dynamics are the primary controls on millennial-scale precipitation dynamics on Cerro Chirripó, this could explain the predominantly inverse relationship between the LMI C_{25} and C_{27} δD record and the C_{29} , C_{31} , and C_{33} δD record (Figs. 2 and 4). A more southerly mean annual position of the ITCZ and a related lowering of the mean annual height of the TWI during the early and late Holocene could lower mean annual cloud heights and lifting condensation level (LCL). This would increase aridity in páramo ecosystems, but could leave subalpine and montane forests at lower elevations relatively unaffected, as these forests would still lie within, or below, the cloud belt. Limited available climate data from the Monteverde Cloud Forest of Costa Rica indicate relatively stable monthly average relative humidity (always >77%; Pounds et al., 1999; Johnson et al., 2005), even during periods of intensified TWIs that can cause relative humidity on the high peaks to drop well below 50% (Dohrenwend, 1972).

This altitudinal humidity and precipitation pattern would explain the relative stability of the C_{29} and C_{31} biomarker δD records that we interpret as having a primarily montane or subalpine source when compared to the C_{25} and C_{27} biomarker δD records that we interpret as primarily originating from the high-elevation páramo ecosystem (Fig. 2). This could also explain why C_{29} , C_{31} , and C_{33}

δD values increase between ~ 7700 and 4200 cal yr B.P. while C_{25} and C_{27} δD values are becoming more negative. If the ITCZ were located in a more northerly mean annual position during this time, the TWI would have been positioned, on average, at a higher elevation, promoting cloud formation and precipitation in the páramos and perhaps increasing water stress at lower elevations via an upward vertical displacement of the LCL and associated clouds that typically bathe montane and subalpine cloud forests year-round. Pollen evidence from the La Chonta bog at 2600 m elevation in the Cordillera de Talamanca (Hooghiemstra et al., 1992) suggests shifts in forest composition that may be associated with early to middle Holocene aridity, perhaps resulting from an increase in mean annual TWI altitude. Existing climate models support this hypothesis as they indicate higher LCLs and reduced cloud contact with tropical cloud forests with increased temperatures (Still et al., 1999). This was likely the situation in southern Central America during the early to middle Holocene, when northern hemisphere solar insolation was significantly higher than southern hemisphere solar insolation (Voorhies and Metcalfe, 2007) and the ITCZ was located in a more northerly mean annual position (Haug et al., 2001). Simultaneous evidence of increased aridity in pollen records from relatively low- (~ 1360 m) and high-elevation (~ 3280 m) sites located near the lower and upper boundary of modern cloud forest distributions in the Peruvian Andes (Bush et al., 2004, 2005) may indicate a narrowed altitudinal range of these specialized ecosystems during the early to middle Holocene in South America. This pattern would be consistent with decreased ITCZ-sourced rainfall and related evapotranspiration rates at low elevations that can significantly impact cloud formation (Lawton et al., 2001) and lowered LCLs as a result of a lower mean annual TWI altitude.

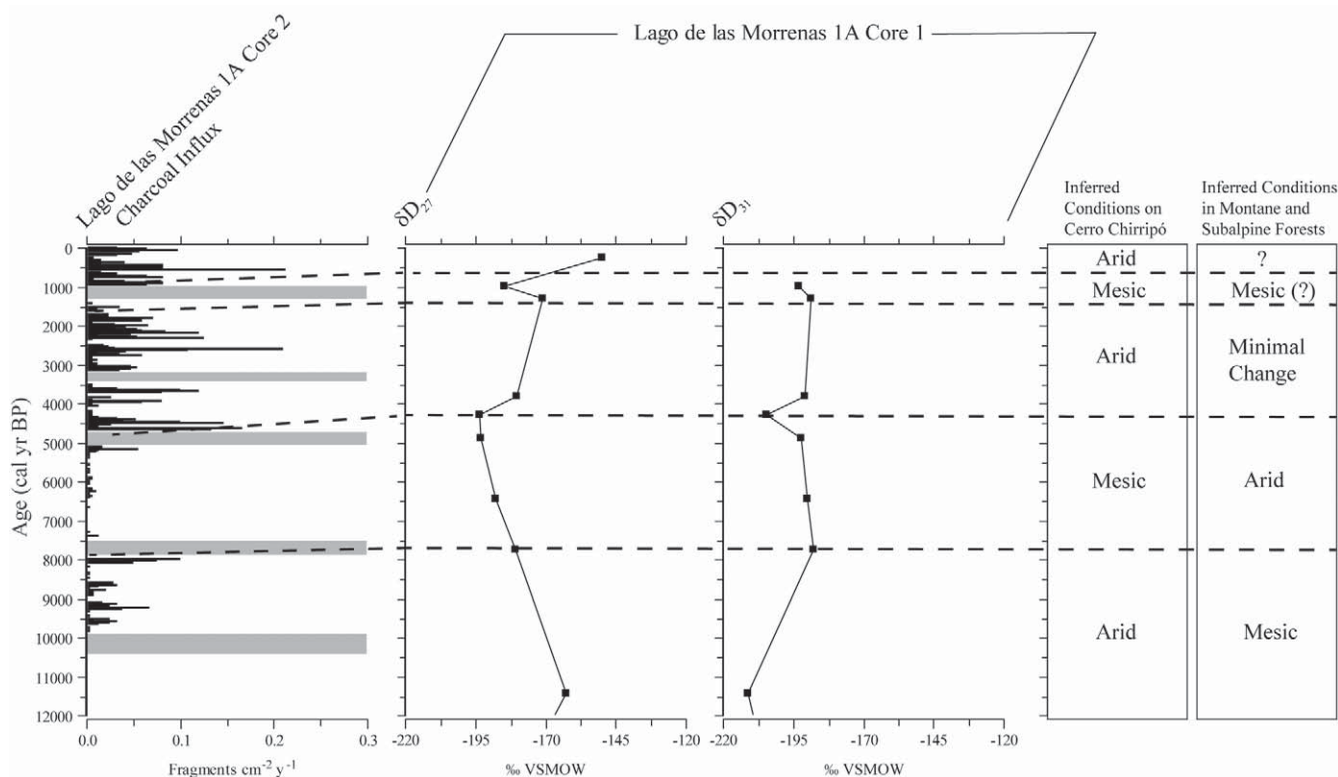


FIGURE 4. Comparison of $\delta^{13}\text{C}$ and C_{27} *n*-alkane δD isotope data with Lago de las Morrenas 1 core 2 macroscopic charcoal data (League and Horn, 2000), C_{27} *n*-alkane δD values, and C_{31} *n*-alkane δD values. Inferred climate conditions on Cerro Chirripó are based on relative changes in C_{27} *n*-alkane δD values. Inferred climate conditions at lower elevations (montane and subalpine forests below the páramo) are based on relative changes in C_{31} *n*-alkane δD values. The terms “mesic” and “arid” are relative and refer to temporal changes in *n*-alkane δD values compared to the preceding time period. VSMOW = Vienna Standard Mean Ocean Water.

MILLENNIAL-SCALE PRECIPITATION VARIABILITY AND FIRE DYNAMICS IN THE LM1 WATERSHED

The C_{27} δD record supports the interpretations of League and Horn (2000), who speculated that millennial-scale patterns of macroscopic charcoal deposition in LM1 core 2 were related to Holocene precipitation variability, with wetter conditions leading to fewer fires and lower charcoal influx. The lowest C_{25} and C_{27} δD values in LM1, between ~ 7700 and 4200 cal yr B.P., coincide with very low macroscopic charcoal influx, supporting the hypothesis that relatively moist conditions during the middle Holocene suppressed fires on the Chirripó massif (Fig. 4). Mayle and Power (2012) documented a coincident mid-Holocene rise in fire activity in the Amazon basin that they associate with increased aridity. This pattern of a wetter Central American isthmus and a drier Amazon basin would be expected if the ITCZ were located in a more northerly mean annual position during this time. Conversely, the sudden rise in C_{27} δD values after ~ 4200 cal yr B.P. coincides with a steep increase in macroscopic charcoal influx, suggesting that increased regional aridity led to increased fire activity around LM1. This rise in charcoal ~ 4200 cal yr B.P. also coincides with a sudden decrease in mean Ti concentrations in the Cariaco Basin, interpreted as decreased ITCZ-related precipitation in northern South America, further indicating a tight linkage between ITCZ dynamics and fire dynamics in the Chirripó páramo.

The documentation of the relationship between millennial-scale precipitation and fire dynamics in the Chirripó páramo helps fill a gap in the understanding of neotropical páramo environments. While modern fire regimes are strongly shaped by humans, charcoal in ancient sediments shows that fire was present in neotropical páramos long before human settlement (Horn and Kappelle, 2009). Fires require dry fuels to spread, and, whether set by humans or nature, depend on short-term climate and weather. The relationships between human activity, climate, and fire are complex and are only beginning to be documented in neotropical highlands (Asbjornsen and Wickel, 2009; Horn and Kappelle, 2009). Resolving links between climate and fire over millennial time scales, as we have done for the Chirripó páramo with hydrogen isotope and macroscopic charcoal analysis, may provide key information for modeling and managing potential changes in fire susceptibility or frequency in páramo habitats under changing climate.

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Appendix

Average δD and $\delta^{13}\text{C}$ values of duplicate analyses of Lago de las Morrenas 1A sedimentary *n*-alkanes typically attributed to emergent or terrestrial plants. Depths are average depths of the sub-sampled sediment, and ages are based on linear interpolation of weighted calibrated ages bracketing the sample position.

Depth (cm)	Age (cal yr B.P.)	δD (‰)											
		C_{23}	σ	C_{25}	σ	C_{27}	σ	C_{29}	σ	C_{31}	σ	C_{33}	σ
16.00	201.28	-147.85	2.05	-139.07	2.74	-149.81	0.70	-147.92	0.46	n.d.	-	n.d.	-
65.00	937.15	-179.64	0.91	-179.44	5.57	-184.86	0.11	-181.16	2.71	-193.24	14.96	-187.56	4.67
85.00	1257.15	-177.57	6.59	-171.62	2.19	-171.09	1.48	-172.97	1.32	-188.63	0.15	-185.11	7.73
190.00	3772.70	-173.38	0.75	-172.33	0.90	-180.55	1.94	-176.95	1.57	-190.78	1.54	-186.99	5.61
240.00	4234.26	-184.62	0.27	-185.72	0.20	-193.53	0.30	-186.35	0.70	-204.58	0.60	-195.35	0.27
302.00	4854.96	-147.69	11.57	-181.95	10.27	-193.27	4.49	-171.43	0.32	-192.28	4.40	-166.52	1.96
390.00	6404.93	-137.11	1.22	-169.83	0.21	-187.93	4.82	-166.61	0.38	-190.04	0.89	-159.80	0.79
440.00	7709.45	-126.54	1.89	-159.70	2.11	-180.89	0.27	-159.74	2.05	-187.97	2.50	-157.28	1.68
545.50	11,383.49	-161.99	6.14	-135.79	8.94	-162.54	12.15	-173.19	9.40	-211.18	10.82	-194.27	11.84
598.50	13,228.60	-172.34	5.76	-152.41	4.36	-175.70	3.04	-181.89	6.31	-205.75	10.13	-190.11	2.23

Depth (cm)	Age (cal yr B.P.)	$\delta^{13}\text{C}$ (‰)											
		C_{23}	σ	C_{25}	σ	C_{27}	σ	C_{29}	σ	C_{31}	σ	C_{33}	σ
3.47	52.04	-34.24	0.30	-31.68	0.72	-33.07	1.64	-31.90	0.50	-31.39	2.48	-28.06	0.91
57.00	856.01	-33.30	0.47	-31.08	1.23	-31.67	0.96	-32.17	1.28	-31.13	0.70	-27.29	0.54
136.29	2571.17	-33.08	0.54	-29.56	1.29	-29.57	0.05	-30.97	1.04	-29.90	1.17	-27.95	0.04
194.50	3817.74	-34.20	0.53	-31.15	0.40	-32.20	0.14	-32.50	1.12	-30.69	0.03	-31.76	0.60
243.25	4305.79	-34.15	0.23	-32.50	0.13	-32.00	0.76	-32.98	0.55	-31.86	0.82	-31.35	0.86
305.00	4924.00	-36.44	0.99	-31.45	0.71	-32.55	0.58	-33.39	0.68	-31.95	0.15	-31.94	0.48
386.25	6360.32	-33.31	1.39	-30.18	0.14	-30.54	0.06	-32.31	1.63	-30.95	0.66	-26.99	0.53
436.25	7618.29	-34.11	0.76	-29.36	0.09	-30.26	0.21	-31.72	0.28	-30.98	0.44	-31.29	0.60
500.25	9846.66	-34.85	2.05	-30.56	0.20	-26.96	0.11	-29.69	0.42	-28.79	1.90	-31.14	1.47
546.00	11,439.60	-21.21	2.35	-24.06	0.54	-23.29	0.10	-29.03	0.79	-29.23	0.99	-31.11	0.02
560.00	11,927.10	-22.75	0.83	-26.12	1.21	-27.08	0.86	-31.02	0.75	-31.84	1.35	-32.09	0.64
595.25	13,154.40	-23.86	0.31	-26.29	0.08	-25.97	0.80	-29.84	0.95	-31.54	0.04	-31.59	0.34