Bromeliad catchments as habitats for methanogenesis in tropical rainforest canopies

Shana K. Goffredi 1,*, Gene E. Jang 1, Walter T. Woodside 1 and William Ussler III 2

1 Biology Department, Occidental College, Los Angeles, CA, USA
2 Monterey Bay Aquarium Research Institute, Moss Landing, CA, USA

INTRODUCTION

Because of the significant influence of methane on atmospheric chemistry and its prominent role as a greenhouse gas, it is becoming increasingly important to understand its sources and sinks on Earth (Crutzen, 1991; IPCC, 2001). Methanogenesis is the dominant terminal mineralization process in wetlands and freshwater sediments that experience prolonged flooding and are limited in more energetically favorable electron acceptors (e.g., sulfate, nitrate, and metal oxides; Whalen, 1993; Grosskopf et al., 1998; Galand et al., 2002; Cadillo-Quiroz et al., 2006). Methane emissions from these environments, both natural and man-made, are substantial (~200 Tg year−1) and are estimated to contribute up to 40% of total global CH4 emissions annually (Crutzen, 1991; Grosskopf et al., 1998; Watanabe et al., 1999; IPCC, 2001; Galand et al., 2002). Methane in these ecosystems is produced exclusively by the activity of certain archaea, during the final step in the anaerobic degradation of organic matter.

Methanogenic archaea can use a number of substrates as precursors of methane, with CO2 reduction and acetate fermentation being most important in wetland soils. Although the relative contributions of the two pathways can vary, acetoclastic methanogenesis generally contributes more to total terrestrial CH4 production than hydrogenotrophic CO2 reduction (Jetten et al., 1992; Conrad, 1999; Conrad et al., 2010). Specialist acetotrophic and more versatile methylotrophic (C1) methanogens belong predominantly to the Methanosarcinales, while H2/CO2-utilizing methanogens include members of the Methanocellales and Methanomicrobiales, to name a few. Methane production by these groups can be influenced by many factors, including availability of substrates and electron donors (e.g., H2), which are often formed catabolically by bacteria, quantity and quality of organic carbon, the presence of possible substrate competitors (e.g., bacteria), and environmental conditions such as pH and oxygen levels (Capone et al., 1983; Galand et al., 2002; Juutonen et al., 2005).

There is a general expectation that as the conditions for methanogenesis are discovered to be broader than predicted, so too will the number of habitats known to contribute methane to the atmosphere. Small-scale water catchments such as bromeliad tanks, Heliconia bracts, tree holes, etc., have the capacity to collect water, thereby hosting microbial communities, with the collective potential to possibly influence carbon cycling, when integrated over ecosystem scales. Epiphytic tank-forming bromeliads, in particular, are those that possess foliage arranged in a compact rosette capable of retaining water. They are predominant members of neotropical habitats, including rain and montane cloud forests, and by some estimates may trap as much as 30,000 l of suspended
water in the canopy ha\(^{-1}\) (Sugden and Robins, 1979; Fish, 1983). In this way, bromeliad tanks provide a vast and unique freshwater niche that does not typically occur in other, especially above ground, locations in tropical forests. In fact, in a recent article by Yavitt (2010), the lentic ecosystems found in bromeliads were classified as “cryptic wetlands.” Accumulation of organic matter, such as leaves and carcasses of dead fauna, primarily arthropods, in bromeliad tanks under acidic, submerged conditions, provides the conditions necessary for the creation of an anaerobic microcosm capable of sustained decomposition and remineralization of terrestrial carbon and the potential for production and release of methane.

Martinson et al. (2010) first reported that tank bromeliads are important sources of atmospheric methane. Methane fluxes, measured from bromeliads in the Ecuadorian Andes, were directly correlated with tank diameter and the plant vascular system itself was determined to act as an important conduit for methane release (Martinson et al., 2010). By extrapolation, it was suggested that CH\(_4\) emissions from bromeliad-associated archaea (up to \(\sim\)2 Tg CH\(_4\) year\(^{-1}\)) might possibly explain the anomalously high methane levels observed previously over neotropical forests (Frankenberg et al., 2008; Martinson et al., 2010). Whether or not the contribution of methane by these ecosystems is globally significant, as suggested by Martinson et al. (2010), but downplayed by Yavitt (2010), it is clear that methanogens in bromeliad tank water play a key role in the cycling of carbon in neotropical environments.

Biomass degradation through methanogenesis remains largely unexplored with regard to suspended catchments, such as bromeliad tanks. Within bromeliad tanks, total organic carbon levels are elevated (\(\sim\)46% TOC), relative to nearby soil (4% TOC), the pH is neutral to acidic (6.5–3.5), and oxygen is low (< 1 ppm; Bermudes and Benzing, 1991; Guimaraes-Souza et al., 2006; Goffredi et al., 2011). Additionally, bacteria related to Proteobacteria, Acidobacteria, Bacteroidetes, and Firmicutes, with varying capabilities of organic carbon breakdown, have been recovered from Costa Rican bromeliad tanks (Goffredi et al., 2011). These bacteria presumably contribute to organic matter decomposition, from recalcitrant chitin and cellulose breakdown to the production of volatile fatty acids and hydrogen, which facilitate the final production and evolution of methane gas by archaea. The objective of this study was to provide a molecular and functional assessment of the methanogen communities inhabiting bromeliad tanks in a lowland neotropical rainforest in Costa Rica, with emphasis on community structure and possible influences on methane production potential (e.g., environment and plant morphology). Bromeliads, as well as nearby soil and artificial catchments, were sampled over the course of 18 months, using both 16S rRNA genes, as well as methanogen-specific mcr\(A\) genes, to document variations in methanogen populations among tanks.

**MATERIALS AND METHODS**

**SAMPLE COLLECTION**

La Selva Biological Station, situated in a wet (4 m annual rainfall) lowland neotropical forest in northern Costa Rica, is located at the confluence of the Sarapiqui and Puerto Viejo rivers in the province of Heredia, Costa Rica (10°26’N, 83°59’W, 37–130 m elevation). The reserve, which covers approx 1600 ha, is home to dozens of bromeliads species, including those within the genera Werauhia (syn. Vriesea), Guzmania, Androlepis, and Aechmea, which were sampled in this study on three separate occasions over 18 months during 2009–2010 (Figure 1). This study included tank water from 106 adult and juvenile bromeliad specimens (Figure 1C), comprising six species; Aechmea mariae-reginae (“Amr”), Aechmea nidi-caulis (“An”), Werauhia gladioliflora (“Wg”), Werauhia kupperiana (“Wk”), Androlepis skinneri (“As”), and Guzmania lingulata (“Gl”), which encompassed a range of tank morphologies (e.g., \(\sim\)3–114 cm plant height) and pH conditions (3.6–6.5; Figure 1D). Additionally, 20 soil samples were collected from near and below sampled bromeliads, as well as eight amber bottles (\(\sim\)100 ml volume) intended to artificially simulate bromeliad tanks, attached to nearby bromeliads for a duration of 6–12 months (Figure 1B). For two bromeliads (Amr1 and Amr51), the pH was artificially depressed, from \(\sim\)6.5 to \(\sim\)4.5 for a total of 62 days, by the frequent addition of 1N hydrochloric acid to the tank (Goffredi et al., 2011). Several bromeliads (and paired soil) were sampled by climber from the canopy (\(\sim\)30 m), but most were within 2 m high on host trees, typically in either man-made or natural clearings. Tank water samples were collected via serological pipette from the bottommost horizon within the tank, with the exception of four bromeliads (with tank depths of \(\sim\)10–15 cm), which were sampled stratigraphically every \(\sim\)3–5 cm. Samples were transported to the

![Figure 1](image-url)
CHEMICAL ANALYSES
Tank pH was measured via hand-held pH electrode (Hanna Instruments HI-98103B) in the field, prior to sampling for DNA analysis. Oxygen was also measured in situ using a colorimetric dissolved oxygen test kit (CHEMetrics, Inc.).

Methane production by bromeliad tank communities was assayed via incubation of tank water (13–30 ml) in gas tight serum vials (total volume 73 ml). Bromeliad tank water for incubation experiments was collected by siphoning through a tube (tygon) inserted, at the other end, into a 100-ml serum vial. Minor amounts of partially decomposed leaf litter and invertebrate remains were entrained in the siphoned water. The serum vials were crimp sealed using a butyl rubber stopper and inverted, at a laboratory temperature of 22˚C in the dark. Over the course of 10–77 days, 10 ml headspace samples were taken via syringe after brief shaking. These samples were taken in the following manner to minimize pressure differences between the interior and exterior of the serum vial. A fully depressed 10-ml syringe with a 22-gauge needle was inserted into the headspace of the serum vial. A second 10-ml syringe with a 22-gauge needle filled with UHP nitrogen gas was inserted into the headspace. Pushing and pulling on the syringe plungers approximately 10–15 times mixed the gases. A 10-ml gas sample was pushed through a low volume magnesium perchlorate drying column prior to injection into the gas chromatograph via a 1-ml sample loop. The addition of nitrogen gas was accounted for when calculating methane concentrations. Headspace methane concentrations were determined using a Shimadzu mini-2 gas chromatograph equipped with a 1/8′′×5-ft stainless steel column packed with Carbosieve G and a flame ionization detector (FID) and operated in isothermal mode (oven temperature 100˚C; detector temperature 110˚C) with a UHP nitrogen carrier gas. Methane gas standards (9.93 and 98.6 ppm CH₄ in helium) were used for calibration. Compound specific isotopic values for methane from three selected samples were measured via gas chromatography-combustion-isotope ratio mass spectrometry using a system consisting of an Agilent 6890 GC, a combustion unit, and a mass spectrometer (Finnigan Delta V plus). Hydrocarbon components were separated by GC, and each individual component slated for isotopic analysis was combusted. The resultant CO₂ was introduced directly into the mass spectrometer, and Finnigan’s Isodat software was utilized for peak detection and quantification (Isotech Laboratories Inc., Champaign, IL, USA).

PHYLOGENETIC ANALYSES
Freshly collected bromeliad tank water (0.5 ml, including debris) was spun at 15,000 × g for 10 min and the resulting pellet was either extracted for total nucleic acids or preserved in 0.5–1.0 ml RNAlater (Ambion, Inc.). DNA was extracted using the Power Soil DNA extraction kit (MoBio Laboratories, Inc., Carlsbad, CA, USA), modified by two initial 5–10 min incubations at 65˚C, one in the presence of solution S1, followed by 5–10 min vortexing. The remainder of the extraction procedure was carried out according to the manufacturer’s instructions, with the exception of a 4˚C incubation in IRS solution (5 min) between solutions S2 and S3 to increase DNA yield and inhibitor removal. SSU (16S) rRNA was amplified by polymerase chain reaction (PCR) from extracted DNA, using the archaea-specific primer pair 1F/1100R (1F, 5′-TCYGKTTGATCCYGSCRGAG-3′; 1100R, 5′-TGGGTCTCGCTCGTTG-3′; Hales et al., 1996), the methanogen-specific primer pair 355F/1068R (355F, 5′-CAGGCCGGAAC ACTTTAC-3′; 1068R, 5′-ATGCTTCACAGTACGAAC-3′; Banning et al., 2005), or a combination of 355F/1100R (Table 1). Thermal cycling conditions were as described, with an annealing temperature of 52˚C, for 25 cycles (Hales et al., 1996; Banning et al., 2005). No difference in relative abundance and

| Sample ID¹ | Catchment pH | Wg37 | Amr34 | Wg104 | Wg104 | An82 | As12 | Wg86 | An91 | Bt15 | Soil13 |
|------------|--------------|------|-------|-------|-------|------|------|------|------|------|--------|
|            |              |      |       |       |       |      |      |      |      |      |        |
| Methanomicrobales | Methanoregula | 14   | –     | 75    | 79    | 77   | 74   | –    | 3    | 12   | 10     |
|             | Methanospririllum | –    | –     | –     | –     | –    | –    | –    | –    | 62   | –      |
| Methanocellales      | Methanocella   | 2    | 32    | –     | –     | 7    | 3    | 21   | 88   | –    | –      |
| Methanosarcinales   | Methanosaeta   | 64   | 24    | 15    | 17    | –    | 23   | 4    | –    | 4    | –      |
|                     | Methanoscarina | 6    | 21    | –     | –     | 3    | 75   | –    | 9    | 73   | –      |
| Thermoplasmatales   | 11             | 12   | 11    | 15    | 10    | 3    | 13   | –    | 9    | 10   | –      |
| Crenarchaeas/Unknown | 3              | 11   | 10    | 4     | 3     | 13   | –    | 9    | 10   | –    | –      |

¹An, Aechmea nudicaulis; Amr, A. mariae-reginae; As, Androlepis skinneri; Wg, Werauhia gladioliflora. Bt15 = an amber bottle (~100 ml volume) intended to artificially simulate a bromeliad tank, attached to Amr1 for a duration of 12 months. Soil13 was a soil sample collected from just below a group of bromeliads.

²The archaea-specific primer pair was 1F/1100R (1F, 5′-TCYGKTTGATCCYGSCRGAG-3′; 1100R, 5′-TGGGTCTCGCTCGTTG-3′; Hales et al., 1996).

³The methanogen-specific primer pair was 355F/1068R (355F, 5′-CAGGCCGGAACACTTTAC-3′; 1068R, 5′-ATGCTTCACAGTACGAAC-3′; Banning et al., 2005).

nm = not measured.
types of methanogens was detected between those recovered using the two main primer pairs (see Wö104; Table 1).

Presence/absence of methanogens was judged by electrophoretic separation and visualization of PCR products produced by the 35S5F/1068R primer set. For seven bromeliads, one artificial catchment, and one soil sample, PCR-amplified bacterial 16S rRNA genes were used for clone library construction. PCR products were pooled in triplicate prior to ligation. Transformants (30–48 clones analyzed for each bromeliad library; Table 1) were screened directly for the presence of inserts using M13F/R vector primers (8 min initial denaturation). M13 amplicons were cleaned prior to sequencing with MultiScreen HTS plates (Millipore Corporation, Bedford, MA, USA). Sequencing reactions were performed using the Genome Lab DTCS Quick Start Kit (Beckman Coulter, Fullerton, CA, USA), precipitated according to the manufacturer's instructions, and run on a CEQ 8800 Genetic Analysis System (Beckman Coulter, Fullerton, CA, USA). Representative ribotypes, based on >97% sequence similarity, were selected for near full-length sequencing (Figure 4).

Sequences were assembled and edited using Sequencher v4.10.1 (Gene Codes Corporation). Initial sequence homology searches were performed using BLASTn (NCBI) and the Ribosomal Database Project classifier. Our 16S rRNA sequences along with additional sequences obtained from GenBank were compiled in ARB, after initial alignment using the SILVA Aligner function, with subsequent manual refinements (Ludwig et al., 2004; Preusse et al., 2007). For near full-length representatives and closest relatives, neighbor-joining (NJ) analysis was conducted with Felsenstein distance correction. In some cases, partial sequences recovered in our study were added to the NJ tree in ARB via parsimony insertion within a tree of longer sequences. NJ analysis was performed with 2000 bootstrap replicates to assign confidence levels to nodes, shown in Figure 4, if >70% confidence (PAUP*4.0b10; Swoford, 1998). Sequences obtained in this study have been deposited in the GenBank database under accession numbers JN810747–JN810784 (archaeal 16S rRNA ribotypes) and JN810785–JN810789 (mcrA genes).

**QUANTITATIVE PCR**

A SYBR green I assay was used to quantify total copy numbers of methyl coenzyme M reductase A (mcrA) genes present in bromeliad tank genomic DNA extracts. Methyl coenzyme M reductase is the key enzyme, unique among methanogenic archaea, which catalyzes the reduction of methyl coenzyme M, the final reductase is the key enzyme, unique among methanogenic archaea, in bromeliad tank genomic DNA extracts. Methyl coenzyme M reductase gene (mcrA) genes present were amplified and sequenced, as described earlier (Table A1 in Appendix). Serial dilutions of the plasmids were used in the optimization of primer concentrations, as well as generation of a standard curve for cycle threshold (Ct) versus gene copy number. The Ct values from these plasmids were converted into copy numbers with consideration of the plasmid preparation nucleic acid concentration, length of the plasmid (3956 bp) and gene insert length (489 bp), average molecular weight of a double stranded DNA molecule (660 g mole−1), and Avogadro's number, resulting in a DNA mass conversion of 1.096 × 10−21 g bp−1. The slope of the standard curve [regression line of Ct versus log(DNA)] was used to assess amplification efficiency and was empirically determined to be 3.3 for the mcrA primer set. Quantitative PCR amplifications (20 μl), performed in duplicate, contained 417 nmol of primer mcrA-mlas and 278 nmol of primer mcrA-rev, and 1× PCR buffer containing MgCl2, dNTPs, and AmpliTaq Gold in the SYBR green master mix (Power SYBR® Green PCR Master Mix; Applied Biosystems, Foster City, CA, USA). Purified DNA from bromeliad tank water was added as template (~1–6 ng ml−1), in replicate dilution series. DNA concentrations for samples and standards were measured using the Quant-it PicoGreen dsDNA assay kit (Invitrogen) and a DTX880 multimode detector and workstation (Beckman Coulter, Brea, CA, USA). Lambda-DNA was used as a standard, as per the manufacturer’s instructions. QPCR amplifications were carried out using an ABI Prism 7500 Sequence Detection System (Applied Biosystems). Cycling parameters included an initial denaturation of 95°C for 3 min, followed by 50 cycles of denaturation at 95°C (15 s) and 60 s of annealing/extension at 60°C. A dissociation curve from each reaction was examined to further ensure proper target sequence amplification (i.e., Tm > 80°C). Given the uncertainty of using plasmids as standards for absolute copy numbers (Hou et al., 2010), mcrA values are shown as relative in Figure 3.

**TERMINAL RESTRICTION FRAGMENT LENGTH POLYMORPHISM AND DIVERSITY**

Methanogen-specific terminal restriction fragment length polymorphism (T-RFLP) profiling was used to characterize the relative proportions, and corresponding diversity, of methanogen groups associated with bromeliad catchments. This approach has been used previously to quantify major methanogen phylotypes from acidic peatlands, a minerotrophic fen, and brackish lake sediments (Banning et al., 2005; Cadillo-Quiroz et al., 2006, 2008). A subset of 25 bromeliads, encompassing a range of morphological and physical–chemical parameters, was selected for T-RFLP fingerprinting. 16S rRNA genes from purified DNA samples, as described above, were PCR-amplified using methanogen-specific primers 355F/1100R [the latter was fluorescently labeled at position 1100 (E. coli numbering) with Well RED dye D4, Sigma-Proligo, St. Louis, MO, USA; Cadillo-Quiroz et al., 2006], using the conditions described above for unlabelled PCR amplifications. For each sample, duplicate PCR amplifications were performed and pooled (300–500 ng total) prior to digestion with Hhal/Sau961 (for 8 h at 37°C, with addition of bovine serum albumin; New England Biolabs, Beverly, MA, USA). Fluorescently labeled fragments were separated by capillary electrophoresis and analyzed on a CEQ 8800 Genetic Analysis System (Beckman Coulter, Fullerton,
CA, USA). Fragment sizes were parsed by separation of >3 bp, and relative abundances were estimated using the CEQ 8800 Fragment Analysis software.

Statistical analyses were performed using the JMP statistical software (version 4.04; SAS Institute, Cary, NC, USA). Diversity estimates of T-RFLP data were performed using Primer v6 (Clarke, 1993; Clarke and Gorley, 2006). Hierarchical cluster analysis, using Bray–Curtis similarity resemblance, Euclidean distance, and no transformation of the data, was also performed using Primer v6.

RESULTS

METHANE PRODUCTION IN BROMELIAD TANK INCUBATIONS

Methane production potential was assayed via incubation of bromeliad tank water in gas tight serum vials. Methane increased within a few days and reached a plateau, by day ~30, at ~900 nmol ml⁻¹ CH₄ in the headspace (= ~22000 ppm CH₄; Figure 2). This plateau was likely due to constraints within the incubation vials as methane production resumed following a purge of the headspace, at day 76, with nitrogen (Figure 2). Methane production rates varied initially between 7 and 38 nmol CH₄ ml⁻¹ tank water⁻¹ day⁻¹, and increased to 100–300 nmol CH₄ ml⁻¹ day⁻¹ immediately following the purge. Additionally, δ¹³C stable isotopic signatures of the generated methane, taken at day 76, were −56.5 and −58.4‰, indicative of a combination of both hydrogenotrophic and aceticlastic methanogenesis. The δ¹³CH₄ value of an additional sample (Wg104), not monitored for methane production, was −62.5‰.

MEASURES OF, AND INFLUENCES ON, MCRA GENE COPY NUMBER

The methanogenic potential of the tank community was examined by QPCR assay of the methyl coenzyme M reductase A (mcrA) gene. Relative mcrA copy number ranged from 1.6 × 10⁶ to 7.0 × 10⁷ ng⁻¹ DNA extract, however, actual copy numbers (and corresponding methanogen abundance) may have been overestimated due to use of plasmids as a standard curve for QPCR analysis (Hou et al., 2010). Nevertheless, there was a positive correlation between mcrA copy number and plant height, suggesting a greater capacity for methanogenesis in larger tanks (Figure 3, R² = 0.61, P = 0.0009). For seven clonal specimens of A. nidi-caulis (ranging from 2 to 7 cm tank depth and 4–20 cm estimated plant height), quantitative assessment of mcrA revealed the same trend of increased copy number in larger plants, and no detection of mcrA genes in the smallest individuals (Figure 3, inset). It should be noted that this relationship did not hold for plants with heights of >54 cm. Although they did possess mcrA genes (data not shown), greater stratification, and a sample volume of ~10–15 ml, likely resulted in an inability to quantitatively sample larger tanks. Additionally, QPCR analysis on four of the artificial catchments did not render detectable levels of the mcrA gene. Two artificial catchments, however, did possess methanogens, albeit at lower relative levels (40% of signal, as compared to the bromeliads with which they were paired).

PHYLGENETIC CHARACTERIZATION OF METHANOCENOCIC ASSEMBLAGES IN BROMELIAD TANKS

Archaeal community composition in bromeliad catchments was analyzed by cloning and sequencing of 16S rRNA genes. Archaea were dominated by methanogens (77–90% of recovered ribotypes) and community structure, although variable, revealed a predominance of various hydrogenotrophic methanogenic orders (e.g., Methanomicrobiales and Methanocellales), as well as both general and specific acetotrophs (Methanosarcinales; Table 1; Figure 4). Very few non-methanogen taxa were detected, using archaeal domain primers, but additional minor archaeal groups included possible members of Thermoplasmatales/Candidate Division II, and uncultured crenarchaeae. Notably, the “methanogen-specific” primer set not only amplified the 16S rRNA genes from other Eur-yarchaeota, perhaps not surprisingly, but also Crenarchaeota in proportions not much different from the archaeal primers, which themselves, are somewhat biased toward Eur-yarchaeota. Many

![Figure 2](https://www.frontiersin.org)

**Figure 2** | Methane production by bromeliad catchment water. Methane increased within a few days and reached a plateau, by day ~30, at ~900 nmol ml⁻¹ CH₄ in the headspace (= ~22000 ppm CH₄). Following a purge of the headspace with nitrogen, at day 76, methane production resumed, and was measured once more at day 192. Methane production rates varied initially between 7 and 38 nmol CH₄ ml⁻¹ tank water⁻¹ day⁻¹, and increased to 100–300 nmol CH₄ ml⁻¹ day⁻¹ immediately following the purge (days 77–78). Additionally, the δ¹³C isotopic signatures of the generated methane, taken at day 76 prior to the purge, were −56.5 and −58.4‰, for Wg57 and As12, closed and open circles, respectively.
euryarchaeal ribotypes, using both archaeal- and methanogen-specific primers, were closely related (based on 97–100% similarity in partial 16S rRNA) to cultured representatives, including *Methanoregula boonei*, isolated from an acidic peat bog (Bräuer et al., 2006, 2011), *Methanocella paludicola*, isolated from rice paddy soil (Sakai et al., 2008), *Methanosarcinaarkeri*, isolated from rice paddies and an underground gas storage facility (Joulian et al., 1998; Tarasov et al., 2011), and *Methanoseta concilii* isolated from an anaerobic sewage sludge fermenter (Patel, 1984). Others were related to ribotypes associated with uncultured groups previously recovered from a range of environments, including estuaries, landfills, deep aquifers, fens, and bromeliads. In fact, several ribotypes recovered in the present study were most closely related to those recovered from Ecuadorian bromeliads (Goffredi et al., 2010).

The predominant group of methanogens recovered in clone libraries varied between bromeliads. For example, recovered ribotypes associated with *Methanocella* were numerically dominant (46 and 97% of methanogen 16S rRNA ribotypes recovered) in the tanks of two bromeliads (*Amr34* and *Amr91*, respectively), *Methanoregula*-associated ribotypes were dominant (75 and 82%) in two other bromeliads (*Wg104* and *An82*, respectively), while *Methanosaeta* and *Methanosarcina*-associated ribotypes each dominated one bromeliad, as assayed via clone library construction (72 and 75% for *Wg37* and *Wg86*, respectively; Table 1; Figure 4). Both the *Methanosaeta* and *Methanocella*-associated clusters, in particular, appeared to be bromeliad-specific, in that ribotypes were most closely related to each other than to any other sequences deposited in the databases (Figure 4).

Out of 20 nearby soil samples (both ground level and canopy height, ~33 m), only one sample (soil13) produced a positive signal for methanogens. Methanogen ribotypes recovered from this soil sample were generally distinct from those recovered from nearby bromeliads, and were, instead, closely related to *Methanosarcina vacuolata* from sludge and *Methanosarcina mazei* from rice paddy soil (97–99% similarity in 16S rRNA; Table 1; Figure 4; Deppenmeier et al., 2002; Liu et al., 2009). Conversely, ribotypes associated with *Methanoregula* were recovered predominantly from a nearby bromeliad (data not shown). Additional methanogenic ribotypes in soil included those found in meadow soil and aquifers (Figure 4).

Artificial catchments (amber bottles), intended to simulate bromeliad tanks, were also analyzed for methanogen presence. Methanogens were generally not detected within the contents of these artificial catchments, suspended in the canopy near bromeliads, even though they had been in place, collecting debris, for 6–12 months. Despite holding similar volumes of water (~100 ml), oxygen levels in bottles were generally elevated (6–8 ppm O2), compared to <1 ppm O2 in the bottommost depths of paired bromeliad tanks. Two artificial catchments did show evidence of methanogens. Visual examination of these two bottles revealed that they contained the largest amount of organic debris. Moreover, they had the lowest pH values of all bottles measured (5.1 versus 6.6 average pH, comparatively). Unfortunately, oxygen levels were not measured in these specific bottles. Time in the field did not appear to matter, as both bottles that assayed positive for methanogens were in place for only 6 months, compared to others in place for up 12 months. The community composition within one bottle (‘Btl5,’ Table 1) was investigated further and
FIGURE 4 | Phylogenetic relationships among archaea associated with Costa Rican bromeliads, a nearby soil sample, and an artificial catchment, relative to selected cultured and environmental sequences in public databases. Relationships are based on sequence divergence within the 16S rRNA gene, with emphasis on methanogenic archaea, as well as those related to the Thermoplasmatales/Candidate Division II, and an uncultured member of the Korarchaeota (DQ465910) as an out group (not shown). The symbols at the nodes represent bootstrap values from a Kimura-2 parameter neighbor-joining analysis obtained from 2000 replicate samplings (open symbol = 70–80%, closed symbol = 80+ % bootstrap support). Sequences obtained in this study have been deposited in the GenBank database under accession numbers JN810747–JN810784.
determined to be distinct from nearby bromeliads, with ribotypes most closely related (97% similarity) to *Methanospirillum hungatei*, isolated from sewage sludge (Ferry et al., 1974), as the dominant methanogen (75% of known methanogen 16S rRNA ribotypes; Figure 4). Additional close relatives (99% similarity) included uncultured representatives isolated from minerotrophic fens and sludge, including a few *Methanoregula*-associated ribotypes (Figure 4; Cadillo-Quiroz et al., 2008; Rivière et al., 2009).

Ribotypes tentatively characterized as *Thermoplasmatales*-associated were recovered in significant numbers from two bromeliads (~12% of the recovered ribotypes; *Amn*34 and *Wg*37) and one artificial catchment (at least 17%; B1); Table 1). *Thermoplasmatales* do occur in low temperature soils (Horn et al., 2003; Juottonen et al., 2005), however, nearest neighbors in other studies (based on 16S rRNA) have been classified as distinct from *Thermoplasmatales* and may, in fact, represent new members within the Candidate Division II (Watanabe et al., 2002; Liu et al., 2009). Whether or not they play a role in methane metabolism is currently unknown.

The alpha subunit of the methyl coenzyme M reductase gene (*mcrA*) was also cloned and sequenced, using the QPCR primers described above, from the tank water of three bromeliads (Table A1 in Appendix). Based on recovered *mcrA* gene sequences, the dominant methanogens were closely related (86–96% amino acid similarity) to members of both the *Methanosarcinaceae* and *Methanomicrobiales*, many of which were previously recovered from similar habitats (e.g., rice paddy soil, acidic peat bogs, and wastewater sludge; Table A1 in Appendix).

PRESENCE AND ABSENCE OF METHANOGENS IN TANK-FORMING BROMELIADS

Tank water samples from 106 specimens, comprising 6 bromeliad species (including *A. mariae-reginae*, *A. nudicaulis*, *W. gladioliflora*, and *A. skinneri*, as noted in the above section, and *W. kupperiana* and *G. linguata*), were extracted for examination of methanogen presence by PCR. Methanogen-specific 16S rRNA genes were recovered for 75 of the 106 bromeliad tanks sampled and included all species, which ranged in plant height from 20 to 114 cm and tank pH of 3.6–6.5. Of the 31 tanks that were negative, or below the detection limit for methanogens, 10 were juvenile plants (less than 5 cm height, with small tank volumes of ~2–8 ml), 10 were small adult plants (~<20 cm plant height and <5–7 cm tank depth), 3 were *Guzmania* species, which typically have very shallow tanks, leaving only eight that were within all specifications of those that did possess methanogens. Seven clonal, rhizomatous specimens of *A. nudicaulis*, ranging from 2 to 7 cm tank depth (~4–20 cm estimated plant height for this species in general) were compared and found to possess methanogens only when tank depth exceeded ~4 cm (see also *mcrA* results; Figure 3).

Four bromeliads (with tank depths of ~10–15 cm) were sampled stratigraphically with three tank horizons sampled every ~3–5 cm depth. In all cases, only the deepest, bottommost layer tested positive for methanogens (data not shown). Strong pH and oxygen gradients were measured in the deeper horizons as well, with pH depressed by 0.2–0.6 pH units and oxygen dramatically reduced (<1 ppm O₂ at depths of 6–7 cm, compared to 2–8 ppm O₂ within 1 cm of the surface of the tank, n = 3; Walter T. Woodside, personal observation).

T-RFLP ANALYSIS OF METHANOGENIC ASSEMBLAGE STRUCTURE AND DIVERSITY

For a subset of bromeliads (n = 25) that encompassed a range of morphologies and environmental conditions, T-RFLP fingerprinting was used to compare the diversity and relative ratios of associated methanogen-specific groups (Figure 5; Table A2 in Appendix). These bromeliads spanned a range of plant heights from 21 to 114 cm, photosynthetically active radiation (PAR) of 33–1282 μmol m⁻² s⁻¹, position off ground of 0.5–34 m, 2–51 mg ml⁻¹ debris within the catchment fluid, and catchment water pH of 3.5–6.5 (including two bromeliads that underwent a manual pH adjustment from 4.5 to 6.5). Fragments (T-RFs) were assigned to particular phylogenetic groups, based on in silico analysis and direct comparison of T-RFs from methanogen clones recovered from bromeliad tank 16S rRNA libraries, as well as previous data from Cadillo-Quiroz et al., 2006, including: *Methanosarcinaceae* (~87, 142, 271 bp; *Methanococaceae* – 90, 119, 399 bp; *Methanomicrobiales*/Fen Cluster (group E2) or *Methanocellales* (Rice Cluster 1) – 232 bp; *Methanocellaclae* (group E1) – 291 bp; *Methanocellaclae* (RC1) – 363 bp. Although Cadillo-Quiroz et al. (2006) distinguished T-RF 230 bp (RC1) from 233 bp (Fen Cluster group E2), we have conservatively combined them, as did Martinson et al. (2010), as a representation of H₂-consuming methanogens. Certain T-RF’s could not be phylogenetically resolved (e.g., 75–79, 135, and 143–150 bp). These unknown fragments typically represented <20% of the total (average 9.8 ± 1.6%), with the exception of five samples (cluster B, described below, Figure 5), for which the unknowns represented 30–49% of the community. In general, T-RFLP analysis resolved the dominant methanogen-affiliated taxa and supported the proportions of methanogens recovered (within ~10%) in the clone libraries. For *Wg*104 for example, the *Methanomicrobiales*/Fen Cluster represented ~78% of the clone library (Methanoregula spp. specifically) versus ~69% via T-RFLP analysis (T-RF 233 specifically). Likewise, the clone library for *Wg*104 recovered 17% *Methanosaeta*-associated relatives, whereas the T-RFLP analysis showed ~26% *Methanosaeta* (Table 1; Figure 5).

Hierarchical analysis revealed three main clusters of samples, with regard to diversity and relative ratios of associated methanogens, based on T-RF abundances, in bromeliad catchment water (>50% community similarity, Figure 5). Group A, which consisted of only specimens of *A. mariae-reginae*, was primarily distinguished by a high abundance of *Methanosaetaceae* (38.1 ± 8.8% average) and very few *Methanomicrobiales* (3.1 ± 0.8%, n = 3). Group B showed the highest abundance (40.6 ± 3.1%, n = 5) of the putative unknown methanogen ribotypes T-RF 143 bp. Group C, comprised of subgroups C1 and C2, had a high abundance of *Methanomicrobiales* (group E2)/*Methanocellae* (56.5 ± 3.4%, n = 16; Figure 5). G171 was distinct from the main groups, likely due to the very low diversity and dominance by *Methanomicrobiales*/*Methanocellae* (92% of T-RFs; Figure 5). Groups B and C1 also had moderately high abundances (~26 ± 3% average) of *Methanosarcinaceae*. Diversity indices ranged from 1.36 ± 0.08 (H′, Shannon index), 0.66 ± 0.03
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FIGURE 5 | Hierarchical cluster analysis of methanogen community composition, as analyzed by T-RFLP fingerprinting, for the catchment water of 25 bromeliad individuals, comprising 6 species. Fragments (T-RFs) were assigned to particular phylogenetic groups, including: Methanosarcinaceae, Methanosaetaceae, Methanomicrobiales/Fen Cluster (group E2) or Methanocellales (Rice Cluster I), Methanomicrobiales/Fen Cluster (group E1), and Methanocellales (RCI) specifically. Hierarchical analysis revealed three main clusters of samples (A–C), with regard to diversity and relative ratios of associated methanogens, shown at left, shaded in relation to % abundance (performed using Primer v6, Bray-Curtis similarity resemblance, Euclidean distance, and no transformation of the data). “M-micro-E2” or “E1” = Methanomicrobiales/Fen Cluster group E2 or E1, respectively.

DISCUSSION

Plants within the family Bromeliaceae are known for their capacity for extreme epiphytism, sometimes growing on bare rock or suspended from vines. Tank-forming bromeliads, in particular, possess foliage arranged in a compact rosette capable of retaining water. In lieu of uptake via root systems, these plants are thought to rely on the catchments and products of decomposition of impounded material (litter and animals) for water and

(1H′, Simpson index), 0.77 ± 0.02 (J′, Pielou’s evenness) for all bromeliads measured (n = 25; Table A2 in Appendix; Figure 5). Bromeliads that underwent manual perturbation of tank pH (manual adjustment from pH 4.5 to 6.5 for bromeliads Amr1 and Amr51) had the highest measures of methanogen diversity (2.01, 0.83, and 0.87, for H′, I'H', and J', respectively; Table A2 in Appendix). The heterogeneous nature of methanogen community structure was similar to that observed in Martinson et al. (2010), and did not appear to correlate with any of the in situ parameters measured, including PAR, amount of solid debris within the tank, position off ground, specific location within the forest, or host plant species. Two notable exceptions included Group A, which were all A. mariae-reginae with high tank pH [>6.5, either naturally (Amr106) or experimentally induced (Amr1 and Amr51)] and Group B, which only included bromeliads collected from sites high in the canopy (11–34 m estimated height off ground).
As a consequence, the suspended water within bromeliad tanks provides a unique niche in that water-saturated, acidic, anaerobic conditions do not typically occur in other locations in the canopy of tropical forests. These catchments are an ideal site for microbial decomposition, and possibly important habitats with regard to global biogeochemical cycling, including CO2 and CH4 efflux and organic carbon storage.

**METHANE PRODUCTION POTENTIAL IN BROMELIAD TANKS**

*In vitro* CH4 production potential measured for bromeliad tank water was comparable to methane production rates in ecosystems known to emit large quantities of methane. Methane production by bromeliad tanks varied between ~12 and 72 nmol CH4 ml−1 day−1, as compared to rice paddies (43–212 nmol CH4 g−1 day−1; Watanabe et al., 2009) and certain peatland ecosystems (13–47 nmol CH4 g−1 day−1; Basilio et al., 2003). Methane dynamics in water-saturated ecosystems are complex and additional parameters must be known, including diffusion, conductance by plants, and oxidation by methanotrophs in overlying aerobic layers, in order to accurately extrapolate bromeliad tank water production rates to field emission rates. We have not identified aerobic methanotrophs in bromeliad tank water, nor have we successfully amplified the key gene involved in aerobic methane oxidation (pmoA; Goffredi unpublished observation; Goffredi et al., 2011). To provide an estimate of net emission rate by tank bromeliads, we assume that aerobic methane oxidation is insignificant relative to methane production rate, and that methane transfer across the tank water (or plant tissue)-atmosphere interface is by diffusion and is in steady-state. Therefore, assuming an average bromeliad tank volume of 100 ml and a bromeliad density of 25,000–175,000 ha−1 (Sugden and Robins, 1979; Martinson et al., 2010), potential emission rates are of the order ~10–500 moles CH4 ha−1 year−1. These estimated emission rates are comparable to those measured for montane bromeliads (82 moles CH4 ha−1 year−1; Martinson et al., 2010) and plant-based pasture production (~190 moles CH4 ha−1 year−1; Parsons et al., 2006), but still fall short of the high, yet variable, values noted for acidic peatlands in general (72–210,000 moles CH4 ha−1 year−1 which depend primarily on moisture regime; Svensson and Rosswall, 1984; Galand et al., 2002; Horn et al., 2003).

The tank rosette itself may be important for the maintenance of the methanogenic microcosm. Stable, strong gradients, with regard to micro-limnological parameters (O2, pH, Corg) may, in part, be related to tank size and shape. Methane release from Ecuadorian bromeliads correlated, exponentially, with tank diameter (Martinson et al., 2010). Similarly, we observed a relationship between mcrA copy number and plant height, suggesting an influential role of plant (and consequently tank) height on methane production capacity, and possibly methanogen community structure. Taller, deeper tanks have time to collect more debris, assuming tank size is positively correlated with plant lifespan. Larger tanks may also experience less mixing and decreased susceptibility to desiccation (Benzing et al., 1972), thereby sustaining methanogen-favorable conditions. Similarly, oxygen levels appear to have some effect on methanogens, which were not observed in either the uppermost horizons within bromeliad tanks or throughout the water column in artificial catchments (i.e., amber bottles), both of which had elevated O2 levels (2–8 ppm O2), compared to the bottommost methanogen-dominated horizons within bromeliad tanks (<1 ppm O2). This implies that either tank shape, or other plant-related parameter, is required for establishing methanogen-favorable conditions (i.e., low oxygen, production of fermentation products).

**METHANOGENIC COMMUNITIES ASSOCIATED WITH BROMELIAD TANKS**

Molecular analysis of the archael communities within bromeliad tanks revealed a dominance of methanogens (77–90% of the recovered ribotypes) and demonstrated the presence of members of both the Methanomicrobiales and Methanosarcinales, known to use H2/CO2, as well as the Methanosarcinales, which are versatile, but known commonly to be acetaticlastic. Similar substrate heterogeneity, in terms of methane production, has been observed for methanogenic communities found in other freshwater environments, including fen soil, acidic peat bogs, and the Florida Everglades (Kotsyurbenko et al., 2004; Smith et al., 2007; Watanabe et al., 2009; Wüst et al., 2009; Martinson et al., 2010; Borrell et al., 2011; Kanokratana et al., 2011). For Ecuadorian bromeliads specifically, there was a similarly strong presence of ribotypes related to Methanosaeta, Methanocella, and what was called the Fen Cluster (or Methanomicrobiales), presumably Methanoregula (Martinson et al., 2010). Interestingly, the Methanobacteria, which represented ~30% of the ribotypes recovered from montane bromeliad tanks (Martinson et al., 2010), were not observed in the lowland forest bromeliads of the present study. Although many parameters are required to determine more certainly the relative contributions of hydrogenotrophic versus acetaticlastic methanogenesis in bromeliad catchments (Conrad, 1999, 2005; Whiticar, 1999; Kotsyurbenko et al., 2004), isotopic measures of methane in two tank microcosms reflected an environment in which methane was generally produced from both processes (δ13CH4 = approx. ~57‰ for Wg37 and As12), and one for which perhaps a slight shift in balance toward H2/CO2-based CH4 production was suggested (~62.5% δ13CH4 for Wg104). This assumption is consistent with clone library results from Wg37 and As12, which both showed ~ 30:70 ratios of the two major methanogenic guilds, whereas Methanoregula was the dominant methanogen in Wg104 (~80%; Table 1; Figure 2).

The relative contribution of acetate and H2/CO2 to CH4 production can vary within ecosystems (Conrad, 1999, 2005) and, among many influences on community composition, it appears that pH may play a role. In this study, individual bromeliads with lower tank pH values (~<6.1) were overwhelmingly dominated by H2/CO2-utilizing methanogens, including those most closely related to Methanoregula (Fen Cluster group E2), and Methanocella (Rice Cluster I), both found previously to be abundant in similarly organic-rich, acidic habitats (Ramakrishnan et al., 2001; Galand et al., 2003; Sizova et al., 2003; Cadillo-Quiroz et al., 2006, 2008; Wüst et al., 2009). These two groups are tolerant of low pH and are known to use H2 as their primary electron donor (Sakai et al., 2008; Bräuer et al., 2011), and, in the case of Rice Cluster I, possess a genome specifically suggestive of a H2/CO2-dependent lifestyle (Erkel et al., 2006). Indeed, for methanogenic
The more versatile Methanosarcina, on the other hand, were only dominant in one bromeliad, by library construction, and occurred in moderate abundances (∼26% average) in nine other bromeliad tanks, based on T-RFLP analysis. In the only other previous study of methanogenic communities within bromeliad catchments (from the high elevation Ecuadorian Andes), Methanosarcina were not observed (Martinson et al., 2010). Importantly, Methanosarcina are common in many soil habitats, including flooded rice paddies (Kudo et al., 1997; Ramakrishnan et al., 2001; Yuan et al., 2011), and we found them to also be the dominant methanogen in the only soil sample (out of 20) that showed positive presence of methanogens.

CONCLUSION
Epiphytic, tank-forming bromeliads are predominant members of neotropical habitats, including cloud and rainforests, and collectively suspend large amounts of water in the canopy, thus representing a large environmental resource. Our results indicate that in stratified bromeliad catchments, methane production occurs and that members of the Methanomicrobiales, Methanocellales, and Methanosarcinales (Methanosetaeaceae in particular) are common. This, plus laboratory incubation of tank waters, provides compelling evidence of bromeliads as an important source of methane, produced by the activities of both hydrogenotrophic and aceticlastic methanogens during the final steps in the decomposition of organic matter. Although at this point linkages cannot be confidently drawn between environmental parameters, bromeliad methanogen community structure, and CH4 production dynamics, it is likely that a combination of factors, including substrate quality and availability, degree of anoxia and desiccation, tank pH, and tank morphology are most influential in regulating production and release of methane to the atmosphere above neotropical rainforests.

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Table A1 | Summary of archaeal methyl coenzyme M reductase (mcrA) clone library results from the tank water of three bromeliads specimens.

| Sequence i.d. | Closest cultured relative     | % Similarity nucleotides (BLASTn) | Closest environmental relative (ace #/environment) | % Similarity amino acid (BLASTx) |
|---------------|-------------------------------|-----------------------------------|----------------------------------------------------|---------------------------------|
| An9_mcrA_D4   | Methanoregula boonei          | 97                                | CAH68747 acidic peat bog                            | 95                              |
| An9_mcrA_A6   | Methanosarcina mazei          | 84                                | CBL29136 rice paddy soil                            | 93                              |
| An9_mcrA_B6   | Methanothermobacter wolfeii    | 86                                | ACD35158 biogas plant                               | 90                              |
| An22_mcrA_C9  | Methanoregula boonei          | 97                                | CAH68747 acidic peat bog                            | 96                              |
| Wg37_mcrA_E9  | Methanoseta concilii          | 89                                | CBA18220 Amazon lake sediment                       | 86                              |

An, Aechmea nudicaulis; Wg, Werauhia gladioliflora.

Table A2 | Bromeliads used in T-RFLP analysis, including catchment pH, plant diameter (cm), photosynthetically active radiation (μmol m⁻² s⁻¹) exposure at the time of sample collection, position off ground (m), group-specific abundances (%) based on T-RFLP analysis, and corresponding diversity indices (Shannon index H', Pielou's evenness J', and Simpson's index 1/λ').

| ID | Species¹       | pH     | Size² | PAR  | POG ³ | M-saeta | M-sarc | E2   | E1 | other | H' | J' | 1/λ' |
|----|----------------|--------|-------|------|-------|---------|--------|------|----|-------|----|----|-----|
| 106| Amr            | 6.50   | nm    | nm   | nm    | 47      | 3      | 5    | 45 | 0     | 1.01 | 0.63 | 0.58 |
| 1  | Amr            | 6.40   | 145   | nm   | 1.2   | 21      | 20     | 12   | 34 | 13    | 1.96 | 0.85 | 0.82 |
| 51 | An             | 6.35   | 110   | nm   | 0.3   | 47      | 13     | 11   | 14 | 15    | 2.06 | 0.89 | 0.84 |
| 91 | An             | 4.65   | nm    | 62   | 11    | 0       | 31     | 20   | 0  | 49    | 1.75 | 0.98 | 0.82 |
| 96 | Wg             | 4.95   | nm    | 48   | 10    | 5       | 17     | 22   | 26 | 30    | 1.87 | 0.90 | 0.83 |
| 61 | Wg             | 4.27   | 30    | 114  | 34    | 11      | 25     | 20   | 0  | 44    | 1.94 | 0.93 | 0.84 |
| 82 | An             | 5.14   | nm    | 64   | 32    | 0       | 27     | 33   | 0  | 40    | 1.58 | 0.81 | 0.76 |
| 83 | An             | 5.12   | nm    | 89   | 32    | 0       | 32     | 28   | 0  | 40    | 1.68 | 0.86 | 0.78 |
| 71 | Gl             | 5.60   | 31    | 147  | 34    | 0       | 0      | 92   | 0  | 8     | 0.87 | 0.79 | 0.52 |
| 23 | An             | 4.95   | 24    | nm   | 2     | 0       | 10     | 69   | 0  | 21    | 0.89 | 0.64 | 0.48 |
| 54 | Wg             | 4.20   | 116   | 200  | 0.5   | 6       | 1      | 73   | 0  | 20    | 1.02 | 0.52 | 0.45 |
| 70 | Gl             | 4.96   | 30    | 92   | 34    | 5       | 6      | 71   | 0  | 19    | 1.15 | 0.64 | 0.53 |
| 99 | Wg             | 4.83   | nm    | nm   | 18    | 22      | 60     | 0    | 0  | 0     | 1.02 | 0.74 | 0.57 |
| 88 | Wg             | 4.50   | 102   | 200  | 0.5   | 29      | 0      | 67   | 0  | 3     | 0.74 | 0.67 | 0.46 |
| 89 | Wg             | 3.65   | 36    | 730  | 0.3   | 14      | 4      | 77   | 0  | 4     | 0.75 | 0.54 | 0.38 |
| 58 | Wg             | 4.30   | 120   | 200  | 0.5   | 22      | 0      | 55   | 22 | 0     | 1.20 | 0.87 | 0.66 |
| 25 | Wg             | 4.08   | 167   | nm   | 0     | 41      | 4      | 50   | 0  | 5     | 1.29 | 0.66 | 0.64 |
| 27 | Wk             | 6.08   | 99    | nm   | 0     | 20      | 10     | 55   | 0  | 15    | 1.47 | 0.67 | 0.65 |
| 104| Wg             | 5.10   | nm    | nm   | 26    | 0       | 68     | 0    | 6  | 0     | 0.78 | 0.71 | 0.46 |
| 92 | An             | nm     | nm    | 62   | 11    | 9       | 30     | 36   | 8  | 16    | 1.99 | 0.83 | 0.81 |
| 95 | An             | 4.82   | nm    | 62   | 11    | 0       | 32     | 47   | 0  | 21    | 1.17 | 0.85 | 0.65 |
| 57 | Wg             | 5.60   | 110   | 200  | 0.5   | 11      | 23     | 55   | 0  | 11    | 1.49 | 0.83 | 0.72 |
| 60 | An             | 4.32   | 56    | 33   | 34    | 22      | 14     | 61   | 0  | 2     | 1.50 | 0.77 | 0.69 |
| 55 | Wg             | 5.38   | nm    | 107  | 0.5   | 27      | 18     | 45   | 0  | 10    | 1.42 | 0.88 | 0.72 |
| 87 | Wg             | 5.30   | 116   | 200  | 0.5   | 18      | 36     | 40   | 0  | 6     | 1.41 | 0.79 | 0.71 |

¹An, Aechmea nudicaulis; Amr, A. mariae-reginae; As, Androlepis skinneri; Wg, Werauhia gladioliflora; Wk, Werauhia kupperiana; Gl, Guzmania lingulata.
²In some cases, plant diameter was estimated from longest leaf length (using a conversion of ×1.7).
³A position off ground (POG) of 0 m indicates that the plant had established itself in soil, presumably after falling from a tree.
⁴E2 and E1 correspond to Methanomicrobiales/Fen Cluster, group E2 or E1, respectively.
nn = not measured.