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Distribution and sources of bioaccumulative air pollutants at Mezquital Valley, Mexico, as reflected by the atmospheric plant *Tillandsia recurvata* L.

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Abstract. Mezquital Valley (MV), a Mexican wastewaterbased agricultural and industrial region, is a "hot spot" of regulated air pollutants emissions, but the concurrent unregulated ones, like hazardous metals and polycyclic aromatic hydrocarbons (PAH), remain undocumented. A biomonitoring survey with the epiphytic Tillandsia recurvata was conducted there to detect spatial patterns and potential sources of 20 airborne elements and 15 PAH. The natural δ^{13} C and δ^{15} N ratios of this plant helped in source identification. The regional mean concentration of most elements was two (Cr) to over 40 times (Ni, Pb, V) higher than reported for Tillandsia in other countries. Eleven elements, pyrene and chrysene had 18–214% higher mean concentration at the industrial south than at the agricultural north of MV. The total quantified PAH (mean, 572 ng g^{-1} ; range, 143-2568) were composed by medium (65%, phenanthrene to chrysene), low (28%, naphthalene to fluorene) and high molecular weight compounds (7%, Benzo(b)fluoranthene to indeno(1,2,3-cd)pyrene). The δ^{13} C (mean, -14.6%; range, -15.7% to -13.7%) was consistently lower than -15% near the major petroleum combustion sources. The $\delta^{15}N$ (mean, -3.0%; range, -9.9% to 3.3‰) varied from positive at agriculture/industrial areas to negative at rural sites. Factor analysis provided a five-factor solution for 74% of the data variance: 1) crustal rocks, 39.5% (Al, Ba, Cu, Fe, Sr, Ti); 2) soils, 11.3%, contrasting contributions from natural (Mg, Mn, Zn) and saline agriculture soils (Na); 3) cement production and fossil fuel combustion, 9.8% (Ca, Ni, V, chrysene, pyrene); 4) probable agricultural biomass burning, 8.1% (K and benzo(g, h, i)perylene), and

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5) agriculture with wastewater, 5.2% (δ^{15} N and P). These results indicated high deposition of bioaccumulative air pollutants at MV, especially at the industrial area. Since *T. recurvata* reflected the regional differences in exposition, it is recommended as a biomonitor for comparisons within and among countries where it is distributed: southern USA to Argentina.

1 Introduction

Monitoring airborne metals, polycyclic aromatic hydrocarbons (PAH) and other bioaccumulative compounds with living organisms (biomonitoring) is a technique in use and refinement since at least the 1960's. It is a multipurpose environmental tool for exploring pollutant occurrence and dispersion trends at different geographical/time scales, to identify emission sources, estimate atmospheric deposition and relate biological/ecological changes to air pollution (Aboal et al., 2006; Conti and Cecchetti, 2001; Pirintsos and Lopi, 2008; Segala et al., 2008; Wolterbeek, 2002). Since bioaccumulative air pollutants usually travel in particles, biomonitoring is done preferentially with organisms that rely on the atmosphere as primary source of moisture and nutrients, such as lichens, mosses and some vascular plants. This dependence is indicated by positive correlations between the concentration of pollutants in the biomonitor tissues and the amount of atmospheric deposition (Sloff, 1995; St. Clair et al., 2002; Wolterbeek, 2002).

We report results from a biomonitoring survey of airborne metals and PAH at Mezquital Valley (MV) conducted in the context of the MILAGRO 2006 field campaign in Mexico. This region is environmentally better known by severe

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soil and water pollution problems caused by over 100 years of agricultural irrigation with untreated sewage water from Mexico City (Cifuentes et al., 1994; Downs et al., 1999; Friedel et al., 2000; Siebe, 1994; Vázquez-Alarcón et al., 2001). Mezquital Valley is also a major Mexican "hot spot" in emissions of particulate matter, SO₂ and other regulated pollutants by the Tula-Tepeji-Vito industrial corridor located there (SEMARNAT-INE, 2006). Nevertheless, there is very little information on the expected concurrence of unregulated air toxics at MV, such as hazardous metals, PAH and other persistent organic pollutants.

This survey explored MV for spatial deposition trends and potential sources of 20 trace and major elements and 15 hazardous PAH using the "ball moss" (Tillandsia recurvata L.) as a natural receptor. This epiphytic Bromeliaceae is very common at the study region, where it mainly grows on mesquite trees (*Prosopis laevigata* (Willd) M. C. Johnst.). Adult individuals are spherical, ca. 10–12 cm in diameter, easy to recognize in the field and collect by hand. They have a reduced stem and non-functional roots. Most of their biomass is formed by linear leaves profusely covered with absorptive trichomes. This latter morphological feature increases the ability of this plant to capture moisture and particles directly from the surrounding air, which justifies naming it as an "air" or "atmospheric" plant (Schmitt et al., 1989). Tillandsia recurvata has some physiological resistance to high levels of O₃ and SO₂ (Benzig et al., 1992), which may partially explain why is it so abundant at the polluted MV. This and other atmospheric Tillandsia species have been used as in situ air pollution biomonitors in some southern US states and Latin-American countries, where this genus is exclusively distributed (e.g., De Sousa et al., 2007; Husk et al., 2004; Pignata et al., 2002; Pyatt et al., 1999; Schrimpff, 1984; Smodiš et al., 2004; Wannaz et al., 2006a, b). Successful transplanting of *Tillandsia* from rural into urban areas for similar purposes has been reported by Brighina et al. (2002), Figuereido et al. (2007), and Malm et al. (1998).

The natural δ^{13} C and δ^{15} N stable isotope ratios of T. recurvata were determined as additional information to identify regional sources of air pollution. These ratios express the ¹³C to ¹²C and ¹⁵N to ¹⁴N proportions in the samples relative to their proportions in standard materials. They are sensitive to spatial and time variations in the isotopic composition of the C and N sources used by plants (Bukata and Kyser, 2007; Liu et al., 2008; Norra et al., 2005; Skinner et al., 2006; Solga et al., 2006). Since most constitutive plant carbon derives from ¹²CO₂ fixed directly from the atmosphere by photosynthesis, plant δ^{13} C ratios are particularly useful in detecting air pollution by the ¹²C-rich emissions from fossil fuel combustion. These emissions have significantly lower ¹³CO₂: ¹²CO₂ ratios than unpolluted air, as indicated by δ^{13} C values around -25%, -30%, -40%and -8% for coal, petroleum, natural gas and clean air, respectively (Pichlmayer et al., 1998). The mixing of fossil fuel emissions with clean air is thus related to the commonly low air δ^{13} C values at urban/industrial areas compared to rural/remote locations (e.g., -12% vs. -8%, respectively, Boutton, 1991). Because of the increase in the air concentrations of ¹²CO₂ caused by fossil fuel emissions and the preferential photosynthetic fixation of this form of carbon dioxide over ¹³CO₂, plants exposed to such pollution develop lower δ^{13} C values than those growing under clean air. In contrast, the plant $\delta^{15}N$ changes related to air pollution sources cannot be interpreted straightforward because of the variety of N compounds involved, their emission and deposition forms, and the complex isotopic fractionation processes occurring within and between the sources and the plant receptors. The natural δ^{15} N values of land plants range roughly from -7%to +9% (Kelly et al., 2005). They can change because of the isotopic composition of the N sources and the ability of each plant species to discriminate against ¹⁵N. In general, for plants obtaining nutrients exclusively from atmospheric sources, this ratio decreases if the deposited N compounds are dominated by δ^{15} N values lower than theirs. Such compounds are usually reduced N species abundant at, but not restricted to, agriculture/farming areas: e.g., ammonia $(NH_3, \delta^{15}N: -10\% \text{ to } -40\%, \text{ several sources})$. Contrarily, this ratio increases in plants predominantly exposed to compounds with $\delta^{15}N$ values higher than theirs, mainly oxidized N species (NO_x), which are more common to urban/industrial areas because of the heavier use of fossil fuels (Jung et al., 1997; Skinner et al., 2006; Solga et al., 2005). Since agriculture and industry coexist at our study area, we expected the T. recurvata δ^{13} C and δ^{15} N to reflect these land uses.

A data matrix with 50 MV sites and 38 chemical variables was reduced by standard univariate statistical techniques. Exploratory factor analysis (FA) of selected variables and mapping allowed us to detect the major regional emission sources. As far as we know, this is the first report on the bioaccumulative air pollution of MV using biomonitoring techniques.

2 Materials and methods

2.1 Study area

Mezquital Valley is located in Hidalgo State, Mexico, ca. 60 km NW of Mexico City (Fig. 1). It is a semiarid region (ca. 2429 km²) with about 500 000 inhabitants distributed in medium-size towns (e.g., Tula and Tepeji del Río) and many small villages. The mean annual temperature and precipitation range from 16.5°C and 432 mm at the north portion of the valley, which is mainly dedicated to agriculture, to 17.2°C and 647 mm at the industrialized south. The mean elevation at the agriculture flatlands is 2000 m a.m.s.l., and up to 2400 m at the surrounding mountains. The original valley's vegetation was a xerophytic shrubland with "mesquite" tree as dominant element. This vegetation disappeared at the

agricultural areas and is now restricted to the foothills and highlands. Over 90 000 ha of MV are currently dedicated to agriculture with raw sewage water from Mexico City. This is the major cause of chemical and biological contamination of the regional soils (Siebe and Cifuentes, 1995). The Tepeji-Tula-Atitalaquia industrial corridor has about 140 industries, including a petroleum refinery (325 000 barrels day⁻¹); a 1500 MW electricity power plant fueled with residual fuel oil and, secondarily, with natural gas, and over 20 cement plants. That power plant is the largest emitter of air toxics among similar Mexican facilities (CEPAL-SEMARNAT, 2004). The regional cement production relies on raw limestone material obtained nearby by dusty quarries. Cement is processed using residual fuel oil, petroleum coke and other materials, like used tires and industrial wastes. Other MV industries include non-ferrous metal manufacturing, textile, chemical, processed food, and disposal, recycling or incineration of waste materials. Cabrera et al. (2003) estimated the MV emissions of total suspended particles (TSP) from industrial and mobile sources in 21538 ton year⁻¹. This is a rather large amount representing ca. 80% of the same emissions at the much larger and higher populated Mexico City $(27308 \text{ ton year}^{-1}, \text{SMA}, 2006)$. The contribution from natural and agricultural sources to the MV air pollution is still unknown. Two major regional natural/crustal sources of particulate material are the Cretaceous limestone (El Doctor Formation) that covers about 60% of MV, mainly exposed at the E-SE sector (Silva-Mora, 1997), and Lower Tertiary igneous rocks, roughly located at the valley's west half.

2.2 Sampling and sample preparation

Tillandsia recurvata was sampled at 50 sites throughout MV in the late spring and early summer, 2006 (Fig. 1). Most sites were at mountain foothills where mesquite trees, from which the biomonitor was obtained, are more common. In a few sites lacking this tree, sampling was done on alternative trees and cacti. The mean elevation of the sampling sites was 2200 m (min, 1879 m; max, 2435 m). Three composite samples per site were collected from tree branches at ≥ 1 m from the ground, each composed by six to eight "ball moss" individuals from different trees. Cross contamination between samples was avoided by wearing new dust-free latex gloves per sample. Samples were taken to the laboratory in air-dried condition in brown paper bags, and stored at ambient temperature until processing (late September-October 2006). They were manually cleaned prior to chemical analyses by removing dead plant parts as well as insects, feathers and spider webs, among other materials extraneous to the samples, but were not subjected to further cleaning procedures, as recommended for air pollution biomonitoring with plants (Rossini Oliva and Raitio, 2003). The samples were then subdivided into three portions for metal, PAH and isotopic analyses, and stored in polyethylene bags at -40° C until analyses. Since T. recurvata is perennial, potential age-related variability in

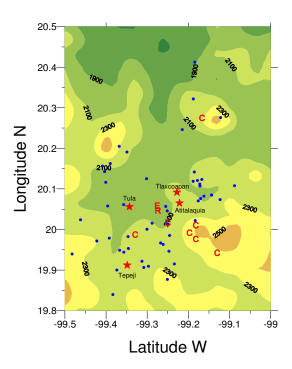


Fig. 1. Mezquital Valley study area. Symbols: dots, sampling sites; stars, main urban settlements; E, electricity power plant; R, petroleum refinery; C, cement plants. The isoline numbers indicate topographic elevation (m).

pollutant bioaccumulation was minimized by analyzing only the newest 3–4 pairs of leaves per shoot. Thus, the time period of environmental exposition was assumed to be similar among sites, probably representing one or two years.

2.2.1 Metals

The glass and Teflon material for sample digestion was cleaned by immersion in 10% HNO₃ (24 h) followed by profuse rinsing with deionized water. The plant samples were oven-dried to constant weight (70°C, three days) and ground to fine powder with agate mortar and pestle. A fraction of the powder (0.15–0.2 g per sample) was digested with 6.0 ml HNO₃ (65%, Merck), 4.0 ml HCl (38%, JT Baker, Ultrex II), and 0.2 ml HF (48%, JT Baker, Ultrex II) in a microwave oven (Anton Paar, Multiwave 3000) using the "pine needles" program: phase I (power, 1400 W; ramp, 10 min; hold, 10 min; fan 1), phase 2 (power, 0 W; hold, 30 min; fan 2). Digestion temperature was 180°C. The digests were filtered with Whatman 4 paper and brought to 25 ml with deionized water.

The elements were determined by inductively coupled plasma/optical emission spectrometry (ICP-OES, Perkin-Elmer, Optima 3200 DV) following the USEPA 6010C method (EPA, 2000). The sample injection flow was 1.0 ml min⁻¹. Calibration curves were prepared with diluted element standard solutions (High Purity). QC repeatability

was checked up by injecting element mixtures (Ultra Scientific) every 10–15 samples. The percent recovery of seven certified elements in seven equally processed samples of the NIST 1575a standard reference material (trace elements in pine needles) was (mean \pm coefficient of variation): Al (102.2 \pm 2.0), Ca (113.5 \pm 10.1), Fe (117.3 \pm 7.5), Mg (84.8 \pm 13.9), P (96.5 \pm 2.0), Zn (83.0 \pm 5.1). The recovery of Ba was low (35.8 \pm 6.4%), but was very good for In (103.7 \pm 2.9%), which was added as surrogate from a 5.0 ppm standard solution.

2.2.2 PAH

The samples were organically extracted in microwave oven and analyzed by high performance liquid chromatography (HPLC) for the 15 PAH listed in Table 1. All glassware and Teflon materials for sample preparation were cleaned with liquid detergent (Liqui Nox), running water and consecutive rinsing with bidistilled water, acetone and dichloromethane (DCM). Samples (5 to 8 g, air-dry weight) were frozen with liquid nitrogen and ground to powder with ceramic mortar and pestle. A fraction of the powder (0.25–0.5 g) was ovendried (90°C, 24h) for dry weight determination and about 2.0 g were extracted in 10-sample batches with 30 ml DCM in microwave oven (CEM, model MarsX). The oven was programmed according to USEPA 3546 method: potency, 1200 W; pressure, 100 psi; temperature, 115°C; time, 15 min; total extraction time, 30 min; cooling, 2 h. Each extraction batch included one pure DCM blank and one positive laboratory control sample (LCS). The LC samples were added 0.5 g of dehydrated and deactivated Cromosorb (60/80 mesh, Supelco, Inc; 400°C for 4h) as a surrogate for the vegetal matrix and 200 μ l of standard mixture containing 5 μ g ml⁻¹ of each target PAH (Chemservice). The extracts were filtered through Millipore membrane and concentrated to 1.0 ml under a gentle stream of ultrapure nitrogen supplied with a nitroevaporator (8158, N EVAP111, Organomation Associates, Inc). The concentrates were cleaned with chromatographic glass columns (40 cm length×1.5 cm ID) filled with deactivated (400°C for 4h) alumina-silica-gel (USEPA 3610B and 3630C). The columns were packed from bottom to top with 1.0 cm silanized glass fiber (Alltech) humidified with DCM, 10 g alumina (Baker, Inc.), 3.0 g silica gel (60/200 mesh, Mallincroft) and 2.0 g anhydrous sodium sulfate (JT Baker) dissolved in DCM. Extra DCM (30 ml) was added to the columns for final PAH separation. Eluates were captured in glass vials and concentrated to 1.0 ml under ultrapure nitrogen. They were then changed into acetonitrile (4.0 ml, Burdick & Jackson), filtered with 0.2 μ m acrodiscs (Pall Gelman Laboratory), reconcentrated to 0.5 ml with ultrapure nitrogen and stored at -40° C in amber glass vials until analyses.

The extracts were analyzed with a liquid chromatograph (Agilent HP, 1100 series) equipped with Nucleosil column (Macherey-Nagel, 265 mm, 100-5 C18 PAH), automatic sample injector, and DAD and fluorescence detectors. The

Table 1. Target PAH and HPLC analytical parameters.

РАН	FLD λex-λem ¹	RT^2	QL^3
	(nm)	(min)	$(\mu \text{g ml}^{-1})$
Naphthalene, NAP	275–350	4.29	0.03
Acenaphthylene, ACY	275-350	4.95	0.03
Acenaphthene, ACE	275-350	6.75	0.01
Fluorene, FLN	275-350	7.30	0.01
Phenanthrene, PHE	275-350	8.59	0.01
Anthracene, ANT	275-450	10.45	0.01
Fluoranthene, FLT	275-450	11.94	0.01
Pyrene, PYR	275-410	13.14	0.03
Benzo(a)anthracene, BaA	275-410	17.29	0.03
Chrysene, CHR	275-410	18.47	0.01
Benzo(b)fluoranthene, BbF	275-410	21.32	0.01
Benzo(k)fluoranthene, B b F	275-410	23.02	0.03
Benzo(a)pyrene, BaP	275-410	24.28	0.03
Benzo (g, h, i) perylene, B g P	275-410	27.56	0.01
Indeno($1,2,3$ - cd)pyrene, Icd P	275–495	29.01	0.03

¹ Fluorescence detector excitation (ex) and emission (em) wave length; ² retention time; ³ Quantification limit.

analyses were done in 10-sample batches plus one blank and one positive control. A calibration curve per batch was developed with seven concentrations (0.0625 to $5.0 \,\mu \mathrm{g} \,\mathrm{ml}^{-1}$) of stock acetonitrile solution containing $5 \mu g \text{ ml}^{-1}$ of each target PAH. Samples were added 150 μ l of 4,4'-difluorobifenil (Chemservice) solution in acetonitrile $(4 \mu g \text{ ml}^{-1})$ as internal standard. Solvent A (methanol, 50%; acetonitrile, 25%; HPLC water, 25%) was injected from 0 to 3 min and gradually exchanged by solvent B (acetonitrile) from 3 to 21 min. Only this latter solvent was added from 21 to 35 min. The injection volume was 5 μ l; flux, 0.3 ml min⁻¹; column temperature, 25°C; wavelength (FLD: 275–495 nm, Table 1; DAD: 230 nm). The total time of analysis per sample extract was 30 min. The method validation parameters were: linearity ($R^2 > .98$); accuracy and precision (RSD<3%); detection limit $(0.004 \,\mu\mathrm{g\,ml^{-1}})$ and quantification limits, depending on the compound, 0.01 to 0.03 μ g ml⁻¹ (Table 1).

2.2.3 δ^{13} C and δ^{15} N

Air-dried samples (200–500 mg) were ground with cleaned ceramic mortar and pestle under liquid nitrogen, and dried at 80°C for 24 h. The δ^{13} C and δ^{15} N ratios were determined with a dual carbon and nitrogen analyzer coupled to a continuous flow isotope ratio mass spectrometer (Europa Scientific). Equipment precision was 0.1% for C and 0.2% for N. The isotopic composition is defined by:

$$\delta^{13}$$
C or δ^{15} N(‰) = $(R_{\text{sample}}/R_{\text{standard}} - 1) \times 1000$

where R is the sample or standard ratio of the heavy to the light isotope: $^{13}\text{C}/^{12}\text{C}$ and $^{15}\text{N}/^{14}\text{N}$. The V-PDB (Viena Pee Dee Belemnite) and atmospheric nitrogen were used as

standards to determine δ^{13} C and δ^{15} N, respectively (Mariotti, 1974; Coplen, 1995).

2.3 Statistics

The raw site-pollutant data base was regionally summarized by simple central tendency and dispersion statistics. The geographical distribution of pollutants was explored by mapping with a geostatistical gridding method for irregularly spaced data (Surfer, Ver. 7.05, Kriging method) and comparing concentrations between and among MV areas with ttests or ANOVA, as appropriate. Potential emission sources were explored by factor analyses (FA) with principal component extraction and normalized varimax rotation (Statistica, Ver. 6.0). Since 50 sampling sites was a small number of cases, FA was used conservatively. Only 20 pollutants were included in FA to keep a 2.5 site to variable ratio. The standardized median site values were used for FA. Pollutants excluded from this analysis had one or more of the following problems: they were not detected or below quantification levels at ≥ 8 sampling sites (i.e., 16% of sites: Pb, Sb, BaA, BbF, BkF, BaP and IcdP); high skewness (>2) and/or kurtosis (>7); strong deviation from normality even after log transformation (Shapiro-Wilk test); no significant correlation with any other pollutant, because they tended to form factors with single pollutants (e.g., Cd and Cr), or contrarily, very high correlation (r>0.9), which may lead to multicollinearity. All low and some medium molecular weight PAH, which predominantly disperse in the gas phase, were also excluded from FA (NAP, ACY, ACE, FLN, PHE, ANT and FLT). Finally, the pollutant selection for FA was aided by considering element signal to noise ratios (SNR), which relate the regional to the site variability, as defined by Wolterbeek et al. (1996), and the element enrichment factors (EF) calculated as

$$EF = (X_s/Al_s)/(X_r/Al_r)$$

where X_s and Al_s are the sample element and aluminum concentrations (ppm), respectively; X_r and Al_r are the element and aluminum concentrations (ppm) in the main crustal rocks of MV: limestone, according to Lozano and Bernal (2005), and acidic igneous rocks, using averaged data from an internet data base by Surendra (2001).

3 Results and discussion

Mezquital Valley was biomonitored with *T. recurvata* to detect the regional dispersion trends and potential emission sources of airborne metals and PAH.

3.1 Metals

Table 2 summarizes the regional element concentrations in the biomonitor. Calcium was the most abundant element (regional mean, 1.3% on dry weight basis; min, 0.5%; max,

Table 2. Regional element concentrations in *T. recurvata* (mg kg⁻¹).

Ele- ment	N	n	Mean	Median	Range	CV (%)	SNR	<i>p</i> (<i>r</i>)	p(log)
Al	150	50	4155	3969	290-11 900	41.1	4.5		>.05
Ba	150	50	38.1	33.2	8.9-131	51.0	4.1		
Ca	149	50	12556	10 190	2883-46780	55.2	7.3		>.05
Cd	150	50	2.2	1.9	0.9-6.5	42.7	2.1		>.05
Cr	139	49	6.1	4.5	0.8-40.4	96.3	6.1		
Cu	150	50	7.0	5.9	2.0-48.4	71.6	3.5		>.05
Fe	149	50	1579	1444	401-4368	41.1	3.5		
K	150	50	7568	7559	1398-16380	33.3	2.5	>.05	
Li	150	50	18.1	17.0	6.0-34.0	24.8	3.2		>.05
Mg	150	50	2906	2815	868-6285	33.8	2.7		>.05
Mn	149	50	80.3	75.2	14.2-327.4	39.4	4.3		>.05
Mo	145	47	4.0	4.0	0.3 - 28.6	76.6	5.5		
Na	149	50	3236	3046	424-10 280	57.6	6.1	>.05	
Ni	123	50	15.9	14.0	1.6-44.2	60.4	4.5		>.05
P	150	50	501	486	163-1642	45.1	3.8		>.05
Pb	86	31	33.1	28.6	1.4-226	92.0	3.6		
Sb	25	9	4.0	2.9	0.2 - 25.4	124.3	3.6		
Sr	148	50	30.5	28.4	1.9-83.2	44.8	2.8		>.05
Ti	150	50	196	177	58.0-532	42.8	2.8		
V	150	50	43.4	29.7	6.5-220	86.8	5.5		>.05
Zn	150	50	41.9	36.4	3.7–192	57.0	4.0		

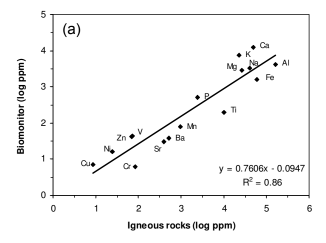
N, number of samples; n, number of sampling sites with quantifiable amount of elements; CV, coefficient of variation; SNR, mean signal to noise ratio; p(r) and $p(\log)$, probability associated to Shapiro-Wilks normality test for raw and \log_{10} transformed data, respectively; empty cells, non-normal distribution.

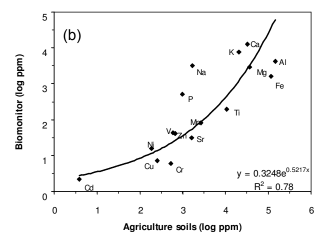
4.5%). This indicated high regional exposure to limestone dust at MV, which is exacerbated by the cement industry. Predominance of Ca in lichen biomonitors growing near a cement plant was recently reported by Branquinho et al. (2008). They considered this element as the best indicator for cement-dust. Other major geological elements followed Ca in abundance (%): K(0.76) > Al(0.42) > Na(0.34) > Mg(0.29) > Fe(0.17) > P(0.05) > Ti(0.02) > Mn(0.008).

The trace elements constituted individually less than 0.005% of the *T. recurvata* dry biomass: V(0.0044) > Zn(0.0042) > Ba(0.0038) > Pb(0.0034) > Sr(0.0031) > Li(0.0018) > Ni(0.0016) > Cu(0.0007) > Cr(0.0006) > Sb(0.0005) > Mo(0.0004) > Cd(0.0002). The prevalence of V, an element typically abundant in crude oils (Bairwise, 1990), can be attributable to emissions from the major industrial users of petroleum fuels at MV. This will be detailed later.

At the gross level of data reduction in Table 2, the element concentrations in T. recurvata reflected the composition of the MV acidic crustal rocks and agriculture soils (Fig. 2). According to this figure, there was a closer chemical similarity between the biomonitor and the igneous rocks (R^2 =0.86) than for soils (R^2 =0.78) and limestone (R^2 =0.64). Thus, a high proportion of the spatial element variability in the biomonitor could be expected to derive from these sources.

Most measured elements had higher mean concentrations than reported for other *Tillandsia* biomonitors in other US and Latin American areas. This is shown in Table 3 by the element concentration ratios between MV and other countries





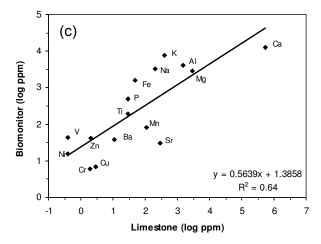


Fig. 2. Elements in *T. recurvata* and crustal/soil sources at Mezquital Valley (log₁₀ transformed mean values, ppm=mg kg⁻¹). Geochemical data: acidic igneous rocks, calculated from Verma-Surendra (2001); agricultural soils, median values among Flores-Delgadillo et al. (1992), Hernández-Silva et al. (1994), Huerta et al. (2002), Lucho-Constantino et al. (2005), and Siebe (1994); limestone (Lozano and Bernal, 2005).

(MV:OT). This ratio was ≥ 2 in 25 out of 44 cases in Table 3, which implies >100% higher bioaccumulation by the Mexican *Tillandsia*. The difference was larger for anthropogenic elements such as Ni, Pb and V, whose respective concentrations at MV were 79, 66 and up to 43 times higher than measured in Tillandsia biomonitors from Argentina (Wannaz et al., 2006a, b). Assuming similar interspecies ability to capture and retain airborne metals, such differences indicate that regional atmospheric deposition levels at MV are at least as high as indicated by those ratios. The MV:OT ratio was <1.0 only in 11 cases; i.e., lower element concentration at MV (Table 3). Most of them involved Cu and Zn biomonitored within urban/industrial areas at São Paulo, Brazil (Figuereido et al., 2007), Cali and Medellin, Colombia (Schrimpff, 1984). This is attributable to the fewer rural sites included in these studies, which are usually less exposed to anthropogenic emissions and tend to lower the regional mean estimates.

3.2 PAH

Table 4 summarizes the regional concentration and variability of PAH in the biomonitor. The sum of quantified PAH per site ranged from 143 to 2568 ng g^{-1} , with regional mean and median equal to 572 and 439 ng g⁻¹, respectively, indicating a data distribution skewed to the right. Apart from PHE and CHR, which were lognormally distributed, the rest of PAH had neither normal nor lognormal distributions. The regional variability of PAH depended on the compound, ranging from 58% (ACY) to 165% (FLN). The medium molecular weight PAH (MMW: PHE, ANT, FLT, PYR, BaA and CHR) was the most abundant group, representing 64.5% of the total measured PAH. The low molecular weight PAH (LMW: NAP, ACY, ACE and FLN) constituted 28.1%, and the high molecular weight (HMW: BbF, BkF, BaP, BghiP and IcdP) only 7.3%. The relatively low amount of the most hazardous PAH group (HMW) deserves further research to be explained. Hypothetically, it may be attributed to photodegradation enhanced by the high solar irradiation rates received in this tropical semiarid region and the scarce protection from it provided by the very open local vegetation.

The most frequent PAH were FLT, FLN, PHE and PYR, which were present at all sampled sites, followed by CHR, NAP and BghiP, which were missed only at one or two sites. Most HMW PAH had low to very low frequencies. For instance, BaP had quantifiable levels at only nine sites, and BaA, BkF and IcdP at the two sites closest to the petroleum refinery and the electrical power plant (Fig. 1). The most abundant compounds were FLT (24.2% of total quantified PAH, regional average), PHE (17.1%) and NAP (10.4%).

Table 5 compares the mean total PAH in *T. recurvata* at MV with other biomonitoring studies using *Tillandsia*. Such comparisons are, however, limited by the several factors affecting the total sum of PAH of each study; e.g., the number and type of measured compounds, the biomonitor's ability to

Table 3. Elements in other *Tillandsia* biomonitors from other countries (OT) respect to *T. recurvata* at Mezquital Valley, Mexico (MV). Units: mg kg⁻¹; values in parentheses, MV:OT ratio.

Element	MV	Argentina ¹	Argentina ²	Brazil ³	Colombia ⁴	Colombia ⁵	US ⁶
Al	4155			1112 (3.7)			
Ba	38.1			36.8 (1)			
Ca	12556			3312 (3.8)			4160 (3.0)
Cd	2.2				0.4 (5.3)	0.82 (2.6)	
Cr	6.1			2.2 (2.8)			
Cu	7.0	0.9 (7.8)	0.9 (7.0)	19.3 (0.4)	52.7 (0.1)	30.5 (0.2)	7.1 (1.0)
Fe	1579	223 (7.1)	257 (6.1)	1144 (1.4)			575 (2.7)
K	7568			5890 (1.3)			2550 (3.0)
Mg	2906			2025 (1.4)			414 (7.0)
Mn	80.3	22.6 (3.6)	22.3 (3.6)	174 (0.5)			52.8 (1.5)
Na	3236			1923 (1.7)			
Ni	15.9	0.2 (79.4)	0.2 (79.4)		11.9 (1.3)	52.8 (0.3)	
Pb	33.1	0.5 (66.2)	0.5 (66.2)		40.6 (0.8)	26.3 (1.3)	
Sb	4.0		0.5 (8.0)				
V	43.4	1.0 (43.4)	1.1 (39.5)	2.7 (16.1)			
Zn	41.9	4.0 (10.5)	3.8 (11.0)	111 (0.4)	119 (0.4)	379 (0.1)	73.5 (0.6)

¹ Wannaz et al. (2006a), *T. capillaris*; ² Wannaz et al. (2006b), *T. permutata*; ³ Figuereido et al. (2007), *T. usneoides*, average value for the two-months transplanting period presenting the maximum concentration, excluding the control site, ^{4,5} Schrimpff (1984), Cali and Medellín cities, respectively; ⁶ Husk et al. (2004), *T. usneoides* in Florida, samples recollected in 1998. Empty cells: data not reported.

capture and retain them, and design aspects of the biomonitoring studies; for instance, number and type of sampling sites, length of exposure and sampling strategy (in situ vs. transplanted samples). No difference was clear between the total PAH at MV and one heavily polluted site at Mexico City measured by Hwang et al. (2003) with pine needles, a morphologically and physiologically different receptor. The mean total PAH reported for T. usneoides from polluted sites in Río de Janeiro (De Sousa et al., 2007) doubled the mean value at MV. Nevertheless, the most polluted MV sites had similar or slightly different total PAH (1284-1748 ng g⁻¹) than the polluted sites at Rio de Janeiro, and no MV site had lower total PAH than unpolluted Brazilian sites (93 to 175 ng g^{-1}). Levels of total PAH at MV were 70% to 140% higher than recorded for two different Tillandsia species transplanted from Costa Rica into downtown Florence, Italy, for eight months (Brighina et al., 2002).

3.3 δ^{13} C and δ^{15} N

The spatial patterns of the δ^{13} C and δ^{15} N in *T. recurvata* reflected quite well the main land uses at MV. Interestingly, these ratios showed that this epiphyte is naturally less discriminatory against 13 C and more discriminatory against 15 N than most terrestrial plants.

The regional δ^{13} C ratio (mean, -14.6%; min, -15.7%; max, -13.7%) was similar to known values for this species (-15.3% to -13.2%, Martin, 1994) and within range for *Tillandsia* with crassulacean acid metabolism (CAM; δ^{13} C>-20%, Crayn et al., 2004); i.e., plants which fix CO₂

Table 4. Regional PAH concentrations (ng g^{-1}) in *T. recurvata*.

HAP	N	n	Mean	Median	Range	CV
						(%)
NAP	141	49	82.3	49.6	6.0-607	122.8
ACY	85	40	88.6	78.8	10.2-292	61.9
ACE	51	32	19.6	13.4	1.1 - 87.0	95.6
FLN	129	50	37.6	21.4	7.2 - 386	134.6
PHE	150	50	136	114	9.9-624	60.3
ANT	92	42	9.0	7.1	1.7-27.0	63.0
FLT	150	50	192	72.2	19.3-1674	164.9
PYR	148	50	25.9	22.1	0.03-158	88.0
BaA	2	2	12.8	12.8	0.2 - 25.4	139.7
CHR	127	49	22.5	18.2	1.9-92.9	72.6
BbF	75	40	18.3	13.2	0.1 - 131	118.0
BkF	2	2	13.5	13.5	1.7-25.3	123.6
BaP	12	9	41.7	39.9	2.4-87.6	58.2
BghiP	112	48	28.1	22.5	4.0-134	73.7
I <i>cd</i> P	2	2	67.0	67.0	22.2-112	94.4
LMW	150	50	164	128	12.4-738	81.2
MMW	150	50	376	261	74.1-2131	96.8
HMW	122	50	42.5	32.8	0.1 - 250	85.0
$\Sigma HAPs$	150	50	572	439	142.6–2568	76.7

N, number of samples; n, number of sampling sites with quantifiable amount of PAH; CV, coefficient of variation.

Table 5. Mean PAH concentration (ng g⁻¹) in *T. recurvata* at Mezquital Valley (MV) and *Tillandsia* in other countries (OT). Values in parentheses are the MV:OT ratio.

HAPs	MV	Mexico City ¹	Sao Paulo, Brazil ²		Florence, Italy ³		
			Polluted sites	Unpolluted sites	T. caput- medusae	T. bulbosa	
NAP	82.3		70.7 (1.2)	11.4 (7.2)			
ACY	88.6	8.3 (10.7)	10.2 (8.7)	1.4 (63.3)	23.4 (3.8)	55.1 (1.6)	
ACE	19.6	1.7(11.5)	13.7 (1.4)	8.6 (2.3)	53.6 (0.4)	53.9 (0.4)	
FLN	37.6		24.6 (1.5)	9.5 (4.0)	4.0 (9.4)	112.1 (0.3)	
PHE	136	15.8 (8.6)	198.2 (0.7)	18.6 (7.3)			
ANT	9.0	2.39 (3.8)	23.3 (0.4)	1.5 (6.0)	6.9 (1.3)	14.0 (0.6)	
FLT	192	66.9 (2.9)	211.5 (0.9)	20.1 (9.6)			
PYR	25.9	118 (0.2)	145.1 (0.2)	14.2 (1.8)			
BaA	12.8	11.6 (1.1)	41.9 (0.3)	3.9 (3.3)	24.5 (0.5)	10.8 (1.2)	
CHR	22.5	49.8 (0.5)	119.4 (0.2)	18.3 (1.2)	34.9 (0.6)	18.5 (1.2)	
Bb+kF	31.8	47.2 (0.7)	111.6 (0.3)	15.9 (2.0)			
BaP	41.7	9.2 (4.5)	72.7 (0.6)	4.0 (10.4)	8.26 (5.0)	35.0 (1.2)	
BghiP	28.1	39.7 (0.7)	29.3 (1.0)	5.0 (5.6)	28.2 (1.0)	, ,	
IdcP	67.0	28.2 (2.4)	31.1 (2.2)	3.9 (17.2)	33.2 (2.0)	12.4 (5.4)	
Σ PAH	572	563 (1.0)	1067 (0.5)	138 (4.1)	238 (2.4)	330 (1.7)	

¹ Hwang et al. (2003): measurements in *Pinus maximartinezii* needles at one site with vehicle/industrial emissions. ² After de Souza Pereira et al. (2007): average summer and winter measurements in *T. usneoides*. ³ Brighigna et al. (2002): eight-month transplanting study at an urban site. Empty cells: data not reported.

predominantly at night as a water-saving adaption to arid environs (Pierce et al., 2002). This ratio was the most homogeneous of the measured variables. It varied only 2‰ throughout MV (CV, 9.0%). Its less negative values occurred at the predominantly rural NW sector of the valley (δ^{13} C = $-14.3\% \pm 0.13$, mean \pm standard error), being on average 0.6‰ higher than at the industrialized SE sector ($-14.9\% \pm 0.12$; p<0.006 from one-way ANOVA by sectors). This suggests that *T. recurvata* has a "background" δ^{13} C signature similar to values recorded at the NW sector of MV (-14.3%, on average).

The δ^{13} C values of *T. recurvata* decreased from the rural periphery of MV to the core industrial area, from values \geq -14.5% to \leq -15% (Fig. 3a). This negative shift reflected the increased emissions of ¹³C-depleted compounds from industrial fossil fuel combustion. The crude oils processed there have an average δ^{13} C = -27‰, as inferred from available data for Mexican petroleum (e.g., Macko and Parker, 1983; Prinzhofer et al., 2000); i.e., 12.4% more negative than the mean value for T. recurvata. The effect from petroleum sources was further recognized by the regional correlation of δ^{13} C with the Ni/V ratio (r=0.39, p=0.01), which was even closer (r=0.70, p=0.001) when considering only the sampling sites along a belt transect in the predominant wind direction (NE-SW). Since Ni and V are the predominant trace metals in crude oils and keep relatively constant mutual proportions, their ratios (Ni/V or V/Ni) are used for geochemical classification of crude oils (Bairwise, 1990) and as environmental indicators for petroleum sources (Ganor et al., 1988). Figure 4 illustrates the parallel pattern followed by δ^{13} C and Ni/V along a NE-SW transect, including a sharp decline at the closest downwind sites from the main petroleum sources (km 0). The regional mean Ni/V for *T. recurvata* was 0.49 ± 0.29 (n=50). The lower (<0.20) and higher values (>0.80) of this ratio occurred, respectively, near and far from the petroleum facilities. The Mexican crude oils used at MV have Ni/V values ranging from 0.07 to 0.24, and 0.17 on average, calculated from data by Castro and Vazquez (2009) and Espinosa Pena et al. (1996). Thus, the biomonitor tended to acquire Ni/V values similar to crude oil at the sites more exposed to petroleum combustion emissions (Fig. 4).

The spatial pattern of $\delta^{15}N$ reflected clearly the agriculture, industrial and rural land uses at MV. The regional mean was -3.0% (min, -9.9%; max, 3.3%), which represented a 6‰ to 9.6‰ ¹⁵N enrichment respect to other Mexican atmospheric *Tillandsia* from less polluted habitats (δ^{15} N = -9.0% to -12.6%, Hietz and Wanek, 2003; Hietz et al., 1999). The lowest, probably "background" signature for T. recurvata, occurred at the rural SW corner of MV (δ^{15} N \approx -7.0%). Since the $\delta^{15}N$ of land plants range roughly from -7% to +9% (Kelly et al., 2005), this *Tillandsia* belongs to the most discriminative plants against ¹⁵N. This suggests that it could rely on biologically usable atmospheric compounds with low δ^{15} N, such as gaseous NH₃ and its derivative aerosol NH₄ (Krupa, 2003). Another contributing factor may be a symbiotic association with bacteria able to convert atmospheric ¹⁴N₂ into usable salts for *T. recurvata*. Puente and Bashan (1994) documented the occurrence of such bacteria in Mexican T. recurvata, but their function needs to be proven.

The T. recurvata $\delta^{15}N$ pattern at MV changed from positive (>3.0%) to negative (<-6.0%) along the predominant wind direction (NE-SW, Fig. 3b). The most positive to slightly negative values occurred at the NNE sector of the valley, near agriculture and industrial/urban areas, whereas the most negative were recorded at the farther WSW rural sites. Similarly high to low plant $\delta^{15}N$ spatial contrasts have been documented for other industrial/urban or agriculture areas respect to rural/remote locations (Gerdol et al., 2002; Jung et al., 1997; Pearson et al., 2000; Solga et al., 2005). Such contrasts are attributed in general to the higher N emission/deposition by human activities in the former areas, an explanation that finds support from the positive correlations often observed between the plant $\delta^{15}N$ and the total N deposition (e.g., Pardo et al., 2007; Skinner et al., 2006). Two major, probably supplementary, hypotheses have been proposed about what causes the higher plant $\delta^{15}N$ at the urban/industrial areas: a) increased exposure to some oxidized N species (NO_x) from fossil fuel combustion, and b) fractionating losses of the lighter ¹⁴N compounds from the plants by leaching or nitrification processes with subsequent concentration of ¹⁵N in the remaining N pools (Jung et al., 1997). Solga et al. (2005) reported a δ^{15} N spatial difference between mosses near agriculture areas (lower values)

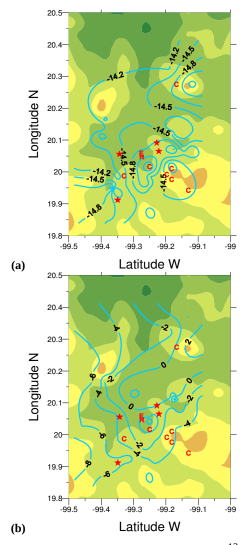


Fig. 3. Spatial distribution of the natural *T. recurvata* δ^{13} C (a) and δ^{15} N (b) isotopic ratios (per mil, ‰) at Mezquital Valley.

and rural environs (higher values) negatively correlated with the $NH_4^+:NO_3^-$ ratio of the bulk N deposition. This implies that increasing the proportion of NO_3^- and compounds with high $\delta^{15}N$, like N_2O (+6.7 to +7.0%, Brenninkmeijer et al., 2003), would shift positively this plant ratio. The same effect would result by lowering the proportion of NH_4^+ and other ^{15}N -depleted compounds.

Although there may be a variety of N atmospheric compounds contributing to the *T. recurvata* $\delta^{15}N$ spatial pattern, specifying which of them accounted for it was beyond the reach of this survey. It can be, however, hypothesized that the higher $\delta^{15}N$ values near the agriculture areas may be related to the heavy use of untreated wastewater for irrigation. There are no $\delta^{15}N$ data for this water, but it is reasonable to assume high positive values, as reported elsewhere for wastewater ($\delta^{15}N$ = +10% to +22%). Soils and plants exposed to such ^{15}N -enriched water tend to acquire high $\delta^{15}N$

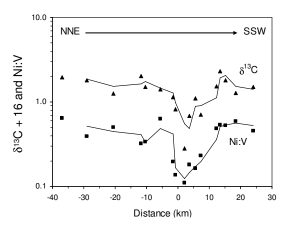


Fig. 4. Median δ^{13} C (triangles) and Ni:V ratios (squares) in *T. recurvata* at sites along a transect oriented with the predominant wind direction at Mezquital Valley (arrow). The main petroleum combustion sources are located at km 0. The δ^{13} C values were added 16 to allow the logarithmic transformation of the y-axis. The fitting lines are two-period running means.

values (Cole et al., 2004; Wigand et al., 2007). This is most likely the case at MV, where agriculture soils have a net positive N balance despite the substantial loss that may occur by volatilization from the open wastewater channels and during irrigation (Siebe, 1998). However, it can be hypothesized that gaseous and particulate emissions of compounds with high $\delta^{15}N$ (e.g., N_2O and NO_3^- , respectively) from those channels and soils are large enough to convert the naturally negative δ^{15} N of *T. recurvata* into positive values. Recent, spatially limited measurements (5 ha) on the emissions of N2O from local agriculture soils showed rates varying from $0.01-0.04 \,\mathrm{mg}\,\mathrm{m}^{-2}\,\mathrm{h}^{-1}$ before irrigation with wastewater to $3.4 \,\mathrm{mg}\,\mathrm{m}^{-2}\,\mathrm{h}^{-1}$ after irrigation, which are similar to fluxes from artificially fertilized soils (González-Méndez, et al. 2008). Other regional activities, like burning agriculture debris, may also contribute with high ¹⁵N emissions.

The $\delta^{15}N$ values at the core industrial area were similar to those at agriculture sites (0% to +3%). This could be due both to in situ NO_x emissions from fossil fuel combustion sources (signaled with C, E and R in Fig. 3b) and to downwind transport of ^{15}N -enriched compounds from the main agriculture areas located immediately to the north of the industrial park.

3.4 Distribution of pollutants and identification of major emission sources

Some insights on the element spatial dispersion at MV can be derived from their regional variability in the biomonitor (Table 2). The average coefficient of variation was 58% and ranged from 25% (Li) to 124% (Sb). The elements with lower variability, such as the same Li and K (33%) were in general more homogeneously distributed throughout the region than those more variable (e.g., V, 87%, Cr, 96%). The

former elements are more likely to be emitted by widely distributed sources; e.g., soil or crustal rocks, whereas the latter ones may derive from localized emission sources.

In looking for general spatial patterns at MV, the predominantly agricultural north was compared to the industrial south using the median site concentrations and straight t-tests. Eleven elements (Al, Ba, Ca, Fe, Mg, Mo, Ni, Sr, Ti, V and Zn) plus CHR and PYR had 18 to 214% higher concentrations at the industrial area (p < 0.05, Table 6). None was significantly more concentrated at the agricultural north. Such spatial difference is partially illustrated in Fig. 5. This figure plots the site normalized median concentration of Ca, CHR, PYR and V within a belt transect oriented along the predominant wind direction (NNE to SSW), including the main types of particle emitters in the region. At the industrial area $(\geq 0 \text{ km in Fig. 5})$, those pollutants had site concentrations up to 5.2 (Ca), 4.2 (CHR), 4.8 (PYR) and 29 (V) times higher than at the farthest north, rural location (Mixquiahuala), located 37 km from the industrial area (Fig. 5). At sites near or within the main north agriculture area (-2 to -30 km in)Fig. 5), most elements and PAH had intermediate concentrations.

Figure 6 illustrates four selected distribution patterns of pollutants as mapped with the median site data. Vanadium had a well-defined concentration peak (>175 mg kg⁻¹) located 3.7 km to the south of the petroleum refinery and the power plant (Fig. 6a). Its concentration dropped centrifugally from there, but at faster rate toward the north than to the south, indicating air transport in this latter direction. The V concentration increased again southward (ca., 12 km from the main peak) probably due to further fossil fuel emissions from cement industries, which are also heavy users of residual petroleum and other materials (petcoke, natural gas, vehicle tires and a variety of residual materials).

Chromium had also a single concentration peak (>20 mg kg⁻¹, Fig. 6b). In contrast with V, this was located at the north of the valley; i.e., upwind from the industrial park. At the rest of the sampling areas, the biomonitor accumulated rather similar Cr levels ($\approx 5 \text{ mg kg}^{-1}$). Two potential sources causing that Cr peak are nearby polluted agriculture soils and limestone dust from quarrying.

Calcium (Fig. 6c) had two high concentration spots, one at the SE ($>36\,000\,\mathrm{mg\,kg^{-1}}$) and another at the NE ($>25\,000\,\mathrm{mg\,kg^{-1}}$) of the study region, coinciding with two major limestone areas where mining for cement materials occurs. In general, this element was more concentrated in the biomonitor at the East half of MV were limestone rocks predominate.

Figure 6d shows the distribution of total PAH. The highest concentrations were recorded at the south of MV, mainly to the WSW of the core industrial area. The lowest levels occurred at rural NW sector, upwind respect to the urban/industrial areas. The tree concentration peaks in Fig. 6d were determined by the most abundant MMW (FLT and

Table 6. Mean and standard deviation of elements $(mg kg^{-1})$ and PAH $(ng g^{-1})$ in *T. recurvata* at north and south Mezquital Valley.

Pollutant	North	South	D^1
Al	3724±919	4625±2004	24.2
Ba	31.0 ± 6.6	43.5 ± 18.2	40.3
Ca	9453 ± 4074	15 577±7079	64.8
Cd	2.1 ± 0.8	2.1 ± 0.6	
Cr	5.9 ± 6.1	4.7 ± 2.2	
Cu	6.1 ± 2	6.7 ± 2.9	
Fe	1283 ± 278	1848 ± 702	44.0
K	7078 ± 2058	7786 ± 1478	
Li	17.0 ± 3.2	18.7 ± 3.8	
Mg	2647 ± 693	3111 ± 971	17.5
Mn	77.3 ± 18.0	79.4 ± 23.7	
Mo	3.2 ± 1.7	4.8 ± 3.3	50.0
Na	3215 ± 1414	2815 ± 1421	
Ni	11.2 ± 5.0	20.8 ± 10.3	85.7
P	478 ± 131	505 ± 235	
Sr	26.8 ± 7.2	34.8 ± 14.3	29.9
Ti	164 ± 35	230 ± 93	40.2
V	21.0 ± 7.0	65.9 ± 38.9	213.8
Zn	31.4 ± 6.9	52.0 ± 27.8	65.6
NAP	55.5 ± 39.2	107 ± 144	
FLN	26.9 ± 25.1	46.9 ± 73.3	
PHE	113±55	144 ± 69	
FTN	136±26 6	159 ± 171	
PYR	16.1 ± 8.6	28.8 ± 11.9	78.9
CRY	14.8 ± 7.2	27.3 ± 16.8	84.5
BGP	25.5 ± 12.0	28.0 ± 15.6	

¹ D, significant South – North percent difference (Student t, p < 0.05).

PHE) and LMW (NAP) PAH, probably reflecting vehicular emissions from the closest urban settlements.

Prior to factor analysis, the element signal-to-noise ratio (SNR) and enrichment factors (EF) were considered as criteria for variable selection (see Sect. 2.3). The average SNR was 4.1 (min, 2.1, Cd; max, 7.3, Ca, Table 2), a good ratio in signal information for a biomonitoring survey. Comparatively, Wolterbeek et al. (1996) reported average SNR values from 1.3 to 3.4 for lichens, mosses and tree bark biomonitors in Europe.

Figure 7 shows the mean element EFs relative to the Al concentrations in the regional limestone and igneous rocks. It includes only elements quantified in both rock types. Enrichment values close to 1.0 are indicative of crustal origin. Higher values, subjectively ≥5, indicate enrichment by anthropogenic emissions. Since K, Ni, P, V and Zn were more enriched than expected from both rock types, they are likely to derive from industrial (Ni, V and Zn) and, tentatively, agricultural sources (K and P). In contrast, Na was enriched only respect to limestone; whereas Ca and Cu were enriched respect to igneous rocks. The rest of elements in Fig. 7, Ba,

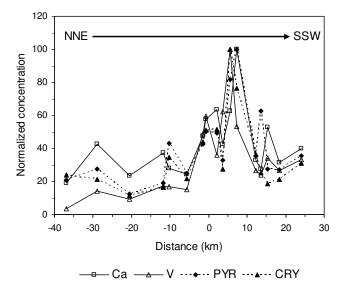


Fig. 5. Normalized median site concentration of two metals and two PAH in *T. recurvata* along a NNE-SSW transect at Mezquital Valley. The 100 value represents the maximum site concentration: Ca, $32\,600\,\mathrm{mg\,kg^{-1}}$; CRY, $82.1\,\mathrm{ng\,g^{-1}}$; PYR, $65.9\,\mathrm{ng\,g^{-1}}$; V, $197.2\,\mathrm{mg\,kg^{-1}}$. The border between the main agricultural and industrial areas is roughly located at km 0.

Cr, Mg, Mn, Sr and Ti, probably originated from crustal/soil sources. The natural emission of some elements can be enhanced by activities like mining. This could be the case for Ca and Cr, whose highest site concentrations occurred near limestone mining areas.

Table 7 shows a factor analysis solution for the 50 MV sites × 20 pollutants standardized data. Five factors (F1 to F5) or potential sources with eigenvalues > 1.0 accounted for 74% of the data variance. Most of it (39.5%) was associated to F1, which was loaded by Ti >Ba >Fe >Al >Sr >Cu (loadings > 0.6), and to some extent by Ca (0.49). Such composition indicates crustal sources, including both igneous rocks and limestone (Ca). According to the site scores from FA (not shown), this factor was geographically centered at the MV agriculture/industrial frontier, 1.5 km from the refinery in NE direction, where the first four elements (Ti to Al) peaked in concentration. This suggests soil dust traveling N to S from the agriculture areas as the element vehicle. Since most F1 elements are considerably more concentrated in the local igneous rocks than in limestone (igneous to limestone concentration ratios: Al, 106:1; Ba, 48:1; Cu, 3:1; Fe, 1201:1; Ti 340:1, with data by Lozano and Bernal, 2005; Surendra, 2001), this factor appears more related to soil sources composed predominantly by igneous parental material.

Factor 2 (11.3% of variance) was loaded (>0.60) by Mn>Mg>Zn, and slightly by Cu (0.44). Interestingly, Na had a negative loading (-0.71) on this factor. This reflected

Table 7. Factor analysis of pollutants in *T. recurvata* (n=50).

	F1	F2	F3	F4	F5
Ti	0.89	0.07	0.25	0.16	0.17
Ba	0.87	0.26	0.14	-0.07	0.17
Fe	0.86	0.08	0.32	0.14	0.19
Al	0.78	-0.30	0.30	0.08	0.22
Sr	0.72	0.26	0.25	-0.15	0.21
Cu	0.61	0.44	0.09	0.27	0.05
Mn	0.25	0.70	-0.14	0.18	-0.18
Mg	0.35	0.68	0.13	0.24	0.21
Zn	0.24	0.66	0.57	0.08	0.19
Na	0.22	-0.71	-0.08	0.37	0.06
PYR	0.17	0.05	0.81	0.12	0.03
V	0.33	0.19	0.79	-0.06	0.29
CRY	0.23	0.24	0.78	0.05	0.34
Ni	0.12	-0.28	0.73	-0.04	-0.26
Ca	0.49	0.04	0.62	0.05	0.33
BgP	-0.20	0.13	0.14	0.72	-0.15
K	0.31	-0.02	-0.02	0.71	0.04
$\delta^{15}N$	0.26	-0.19	-0.05	-0.2	0.69
P	0.42	0.27	0.17	0.08	0.67
δ^{13} C	-0.08	-0.01	-0.19	-0.01	-0.73
% variance	39.5	11.3	9.8	8.1	5.2

a major spatial contrast between Na, with higher concentrations (4000 to 5000 mg kg⁻¹) within or near agriculture areas (Fig. 8); whereas the other elements were more concentrated at the farther rural areas to the south (Mn>100 mg kg⁻¹) and SE of MV (Mg>4000 mg kg⁻¹). Although Zn had high levels at some industrial sites ($>60 \,\mathrm{mg \, kg^{-1}}$), it kept similar concentrations at the SE rural area, resembling the Mg pattern. This factor may be also related to soil sources, but distinguishing saline agriculture soils at the MV lowlands (Na) from natural soils (Mg, Mn and Zn). Sodium is the main cation in irrigation wastewater at MV (Cervantes-Medel and Armienta, 2004; Jimenez and Chávez, 2004), which is attributed transport by wastewater from the saline areas near Mexico City (Siebe, 1998). However, potential inputs from other sources for the elements loading on this factor cannot be excluded. For instance, industrial particulate matter traveling N to S (Cu and Zn) and gasoline vehicle emissions (Mn) from a major line source located at the S limit of MV, the Mexico City-Querétaro highway.

Factor 3 (F3) accounted for 9.8% of the variance and was significantly loaded (>0.60) by PYR >V>CRY>Ni>Ca and Zn (0.57). It is clearly an anthropogenic factor with pollutants from fossil fuel combustion and cement production (Ca). The presence of Ca on this factor is explained by the spatial concurrence of the main petroleum combustion sources and the major limestone quarries and cement plants, which are also heavy users of fossil fuels, at the SSE sector of MV. The negative loading of δ^{13} C on this factor added

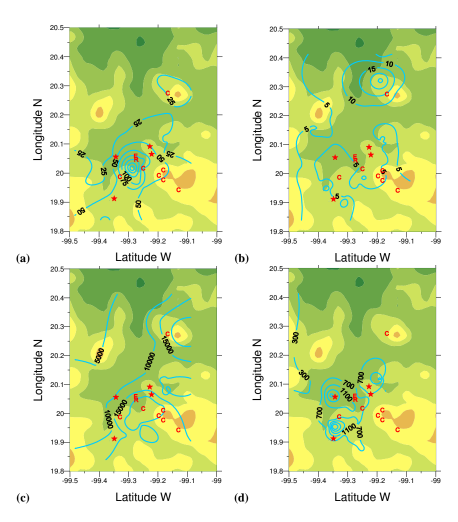


Fig. 6. Spatial distribution of V (a), Cr (b), Ca (c) and total PAH (d) in *T. recurvata* at Mezquital Valley. Units: mg kg⁻¹.

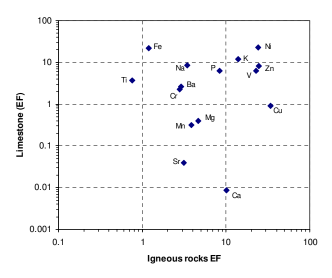


Fig. 7. Mean element enrichment factors in T. recurvata at Mezquital Valley.

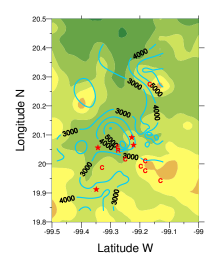


Fig. 8. Spatial distribution of Na in *T. recurvata* at Mezquital Valley. Units: $mg kg^{-1}$.

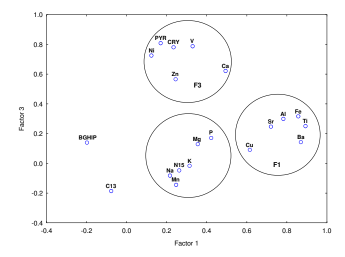


Fig. 9. Factor 3 vs. factor 1 plot. F1 includes elements from mainly crustal origin; F3, elements and PAH from petroleum fuel combustion and cement production (Ca). The third circle includes elements related to agriculture soil sources (F2, F4 and F5, not shown).

further evidence to associate it with petroleum combustion. Figure 9 illustrates how these pollutants separate in the factor space from the predominantly crustal/soil elements (Al, Ba, Cu, Fe, Sr and Ti).

Factor 4 (F4) had 8.1% of the variance associated with it. It was loaded by K and benzo(g, h, i) perylene, deriving tentatively from biomass burning, which may include agriculture debris in the flatlands and forest fires in the highlands at the W side of MV.

Factor 5 (F5) accounted only by 5.2% of the variance. It was loaded by the δ^{15} N isotope ratio and P. This factor may reflect agriculture activities with wastewater, characterized by large emissions of N compounds, mainly NH₃, NH₄⁺ and NO₃⁻ (Cervantes-Medel and Armienta, 2004; Jimenez and Chávez, 2004) and P (Ortega-Larrocea et al., 2001). Phosphorus may also derive from pesticides used in agriculture, which often contain this element. The negative loading of δ^{13} C on this factor added evidence on the regional difference between agriculture and industrial (fossil fuel combustion) land uses.

Although FA did not extracted pure factors/sources, it acceptably distinguished contributions from the most relevant sources of particulate matter into the MV air. Most element variability in the biomonitor, 64% out of 74% total explained variance, appeared associated with crustal/soil sources and agriculture activities (F1+F2+F4+F5), and only 10% (F3) was attributable to emissions from industrial fossil fuel combustion, including power generation, petroleum refining and cement production. Increasing the number of sampling sites, especially around some major fixed sources, such as the power plant and cement factories, may allows us to differentiate their particular contributions to the load of bioaccumulative atmospheric pollutants in the study region.

4 Conclusions

This survey produced previously unknown information on the regional levels, sources and dispersion of airborne metals and PAH for Mezquital Valley, Mexico. Tillandsia recurvata was confirmed as a good natural receptor for those pollutants. The site to site concentration variability, as indicated by high signal-to-noise ratios for most metals (>3.0), reflected the differential long-term exposition to most measured pollutants, especially to the anthropogenic ones. Some anthropogenic metals (e.g., Ni, Pb and V) had particularly high concentrations respect to values for similar biomonitors in other countries, indicating that atmospheric deposition should be several times higher at this Mexican region. The levels of PAH were more similar to values for Tillandsia biomonitors, with a few exceptions (e.g., ACY and IcdP). The southern industrial portion of MV was more polluted than the agricultural north, as implied by the higher concentrations of 11 elements and two PAH. From the chemical correlation between the biomonitor and the regional rocks, results from factor analysis and the geographic patterns of pollutants, it is concluded that crustal/soil sources had a higher relative impact than agriculture and industry on the spatial variability of pollutants. The high "background" δ^{13} C and low δ^{15} N of *T. recurvata* make this epiphyte a sensitive biomonitor for ¹³C-depleted and ¹⁵N-enriched sources. It may be thus recommendable to use it as a common natural receptor for comparative studies within and among Latin American countries and the southern US states, where it is naturally distributed. This would minimize potential variability from using different species.

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