

Short communication

The elusive quantification of nitrogen fixation in xeric shrubs: The case of *Adesmia volckmanni*, a Patagonian leguminous shrubR.A. Golluscio^{a, b, *}, Regina Irueta^a, P.A. Cipriotti^{b, c}^a Department of Animal Production, School of Agriculture, University of Buenos Aires, Av. San Martín 4453, 1417, Buenos Aires, Argentina^b IFEVA (UBA-CONICET), Argentina^c Department of Quantitative Methods and Information Systems, School of Agriculture, University of Buenos Aires, Av. San Martín 4453, 1417, Buenos Aires, Argentina

ARTICLE INFO

Article history:

Received 12 November 2013

Received in revised form

13 July 2014

Accepted 16 July 2014

Available online 5 August 2014

Keywords:

Desertification

¹⁵N natural abundance

N cycle

Nodule-leaf ^δ¹⁵N difference

Patagonian steppes

Watering responses

ABSTRACT

The comparison of ¹⁵N natural abundance ($\delta^{15}\text{N}$) between fixing and non-fixing reference species, is the most feasible method for nitrogen fixation studies in xeric shrubs. However, it assumes that both species use the same sources of soil N, the $\delta^{15}\text{N}$ of such source is clearly different from zero, and the difference between the $\delta^{15}\text{N}$ of both species is $\geq 5\text{‰}$. On the other hand, the analysis of the difference between the $\delta^{15}\text{N}$ of nodules and leaves ($\Delta\delta^{15}\text{N}_{n-l}$) does not require a reference species. To evaluate the effect of water availability on nitrogen fixation, we analyzed the variation of $\delta^{15}\text{N}$ and $\Delta\delta^{15}\text{N}_{n-l}$ in watered and unwatered plants of *Adesmia volckmanni* Phil. (Fabaceae) and *Mulinum spinosum* (Cav.) Pers. (Apiaceae), formerly proposed as non-fixing reference species. However, we found that *M. spinosum* was not valid as reference species to study N fixation in *Adesmia volckmanni* because its $\delta^{15}\text{N}$ was closer to zero than that of the presumably fixing species, suggesting they use different soil N sources. In addition, water availability did not affect biological fixation after watering because both leaf $\delta^{15}\text{N}$ and $\Delta\delta^{15}\text{N}_{n-l}$ of *A. volckmanni* did not change either as soil dried up along the growing season nor in response to watering. However, the presence of nodules itself, the $\delta^{15}\text{N}$ closer to zero than in a former dry year, and the $\Delta\delta^{15}\text{N}_{n-l}$ ($+1.23\text{‰}$) suggest that biological fixation could have occurred before watering, probably during early spring.

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1. Introduction

Quantification of nitrogen fixation of xeric shrubs is very difficult. The three main available procedures are the acetylene reduction assay, the ¹⁵N dilution method, and the ¹⁵N natural abundance method (Virginia et al., 1989). The acetylene reduction assay is accepted as the only method measuring potential nitrogenase activity, closely related to potential N fixation ability. However, it requires the use of live nodules, which stay alive a very short time once detached from roots, and in the case of xeric shrubs are scarce and very deeply located in soil. The ¹⁵N dilution method is the only method measuring the proportion of plant N obtained by fixation. However, it requires the use of expensive marked N in the vast and deep soil volume occupied by shrub root systems. In

addition it must be applied with water, therefore confusing the measurement of N fixation itself with the effect of water addition on N fixation. Natural abundance of ¹⁵N ($\delta^{15}\text{N}$) of fixing plants was proved to be more similar to that of atmosphere (atmospheric $\delta^{15}\text{N} = 0$) as more active is fixation (Dawson et al., 2002). As a consequence, the comparison between the leaf $\delta^{15}\text{N}$ of “fixing” and “non-fixing plants” may be a valid indirect evidence of N fixation (Dawson et al., 2002). However, this approach relies on two basic assumptions: both species use the same sources of soil N, and the $\delta^{15}\text{N}$ of such source is clearly different from zero (Högberg, 1997). In addition, the method relies on a $\geq 5\text{‰}$ difference in the $\delta^{15}\text{N}$ values of both species, under active fixation (Högberg, 1997).

The objective of this work is to demonstrate the existence of N fixation in a leguminous shrub of the Patagonian steppe. In a previous work, we compared the leaf $\delta^{15}\text{N}$ of two conspicuous shrub species of the Patagonian steppes, a presumably N-fixing leguminous (*Adesmia volckmanni* Phil., Fabaceae) and a non-fixing umbelliferous (*Mulinum spinosum* (Cav.) Pers., Apiaceae) (Golluscio et al., 2006). Because of the high similarity of their root profiles and phenological patterns, we assumed that both species would access

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the same fractions of soil mineral N (Golluscio et al., 2005). However, the $\delta^{15}\text{N}$ of *Adesmia volckmanni* was similar to that of *M. spinosum* plants growing together in a xeric steppe ($\sim -1.5\%$ for both species), but was nearer to zero for *A. volckmanni* plants growing close to a wetland, and significantly different to the $\delta^{15}\text{N}$ of co-existing *M. spinosum* plants. As a consequence, we hypothesized that fixation in *A. volckmanni* would depend on water availability (Golluscio et al., 2006). The logical prediction from this hypothesis would be that biological fixation of *A. volckmanni* must increase with watering, and must decrease along the growing season, as water availability decreases (Sala et al., 1989). However, this may be an oversimplified hypothesis due to several logical and methodological problems. First, increasing water availability could also increase mineralization, nitrification, and soil N availability, then decreasing N fixation. Second, Patagonian shrubs usually show little response to watering, because they explore deep, humid soil layers (Golluscio et al., 1998, 2009). Third, even if watering triggers a N fixation pulse in *A. volckmanni*, it would not cause a sufficient $\delta^{15}\text{N}$ difference with respect to the reference species *M. spinosum*. Fourth, in spite of their phenology and root-structure similarities, both species may use different sources of edaphic N.

To deal with these problems, we not only compared the $\delta^{15}\text{N}$ of both watered and unwatered plants of both species but also analyzed the difference between the $\delta^{15}\text{N}$ of *A. volckmanni* nodules and leaves. This isotopic index may be useful to qualitatively evaluate biological N fixation, and has the advantage of being independent of the existence of a valid reference species. In N-fixing legume species, nodules are enriched in ^{15}N (i.e. its $\delta^{15}\text{N}$ is higher than that of their soil source) while leaf $\delta^{15}\text{N}$ is lower than that of nodules (Shearer et al., 1982), and the ^{15}N enrichment of nodules increases with fixation activity (Khadka and Tatsumi, 2006a). This pattern would respond to the discrimination against ^{15}N during the process of transference of fixed N from nodules to other sites, and also during the subsequent allocation and circulation of fixed N from shoot to root (Khadka and Tatsumi, 2006a). As a consequence, the difference between the $\delta^{15}\text{N}$ of nodules and leaves ($\Delta\delta^{15}\text{N}_{n-l}$) increases proportionally to biological fixation (Wanek and Arndt, 2002). This hypothesis was successfully tested for annual legume crops as soybean (Khadka and Tatsumi, 2006a; Schweiger et al., 2012; Wanek and Arndt, 2002), bean and cowpea (Khadka and Tatsumi, 2006a), but also for perennial legumes, as *Lespedeza cuneata* (Khadka and Tatsumi, 2006b). Although we do not know the exact relationship between $\Delta\delta^{15}\text{N}_{n-l}$ and N fixation for *A. volckmanni*, we will adopt $\Delta\delta^{15}\text{N}_{n-l}$ to qualitatively estimate N fixation. We expect this approach may be useful for other researchers aiming to assess N fixation in shrubs growing in arid environments.

2. Materials and methods

We performed a manipulative experiment from November 2006 to March 2007 within a 1 ha plot closed to the access of big herbivores since 1983, located at the experimental farm of INTA (Instituto Nacional de Tecnología Agropecuaria) at Río Mayo, SW of Chubut province, Argentina ($45^{\circ}41' \text{ S}$, $70^{\circ} 16' \text{ W}$). The sampling area is representative of the Patagonian semiarid shrub-grass steppes. Soils have a 45–60 cm deep upper layer, sandy with high gravel content, over a calcareous layer with even higher gravel content (Paruelo et al., 1988). Mean annual temperature is 8.4°C , with 14°C in the hottest month (January) and 2°C in the coldest one (July) (Golluscio et al., 1998). Mean annual precipitation is 150 mm, 70% falling in autumn-winter, the water recharge season (Jobbágy and Sala, 2000; Sala et al., 1989). Most water used by plants during the experiment fell during 2006, a humid year 24% over the historical mean (186 mm, concentrated between May and

September). As usual, precipitations during the first three months of 2007 were scarce (6.1 mm between January and March 2007).

We irrigated plants with a unique water pulse at the beginning of growing season (November 7), until a 1 m deep soil cylinder of $R + 0.5 \text{ m}$ of radius ($R = \text{mean radius of the individual shrub}$) attained field capacity (see water retention curves in Paruelo et al., 1988). We compared irrigated vs. non-irrigated plants growing in the field, focusing on (a) the variation of the leaf natural abundance of ^{15}N ($\delta^{15}\text{N}$) along the growing season between *Adesmia volckmanni*, presumably “fixing” species, and *M. spinosum*, utilized as reference “non-fixing” species (Golluscio et al., 2006), and (b) the difference in $\delta^{15}\text{N}$ between nodules and leaves of the fixing species. We also measured leaf water potential and nitrogen content of both species, nitrogen content in *A. volckmanni* nodules, and $\delta^{15}\text{N}$, N content, and gravimetric water content at five soil depths (5, 15, 30, 45 and 60 cm) under plants of both species. We sampled all variables at four sampling dates during the growing season (Southern Hemisphere: December 6, January 7, February 7, and March 7).

The sampling units were 24 plant pairs, each one composed of one individual of each species, separated $\leq 1.5 \text{ m}$. At the time of watering, we randomly defined 6 pairs to be sampled at each of four sampling dates, and half of these pairs to be watered. At each sampling date, on the six randomly chosen pairs, we measured leaf water potential with a BioControl model 6 (Argentina) pressure bombe (Scholander et al., 1965). We also sampled leaves of both species to measure their $\delta^{15}\text{N}$ and N content, and extracted one 1 m deep soil parallelepiped ($0.8 \times 0.8 \text{ m}$ section), with one vertical face below the canopy of each individual. At five depths of both faces we gathered the soil samples, on which we measured all the above mentioned soil variables. In addition, on the face below *A. volckmanni*, we harvested nodules to measure their $\delta^{15}\text{N}$ and N content.

We performed the analyses of soil and plant N content and $\delta^{15}\text{N}$ (standard: atmospheric N_2) at CATNAS (Centro de Aplicaciones de Tecnología Nuclear en Agricultura Sostenible) of University of the Republic, Uruguay, using a Finnigan MAT Delta Plus XL mass spectrometer. The precision of the device, calculated as the standard deviations of the repeated measurements made on laboratory standards were 0.1% and 0.1‰ for plant samples, and 0.2% and 0.2‰ for soil samples (for N content and $\delta^{15}\text{N}$ respectively). Since both species produce green leaves at the same time of the year and did not retain green leaves of previous years (Golluscio et al., 2005), the leaves of both species had essentially the same age at each sampling date. Soil and leaf samples were oven dried at 60°C for 48 h, then ground with mortar and pestle until samples could pass through a $40 \mu\text{m}$ mesh.

To measure gravimetric soil water content, immediately following gathering, we placed $< 2 \text{ mm}$ sieved soil samples within double polyethylene bags to avoid water losses. We kept samples at air temperature during field work and afterwards in refrigerator at 5°C until processing. Then, we weighed mass of samples, before and after dry them in oven at 105°C until constant weight, and calculated gravimetric water content as the mass ratio between water and dry soil.

We analyzed the results by Analysis of Variance (ANOVA) for linear mixed models. For plant variables (leaf water potential, $\delta^{15}\text{N}$ and N content of leaves and nodules), we used a two way array (4 dates \times 2 watering levels), with plots splitted by species (2 levels: *M. spinosum* or *A. volckmanni*, except in the case of nodules, which only existed below *A. volckmanni*). To analyze soil variables ($\delta^{15}\text{N}$, N content and gravimetric water content), we used the same statistical layout (4 dates \times 2 watering levels), with plots splitted by species (2), but assessing the depth effect (5 depths) below the respective plant species. To do this, we incorporated into the linear model a simple correlation structure associated to each plot and species. All studied effects were fixed, except for plots. In all cases

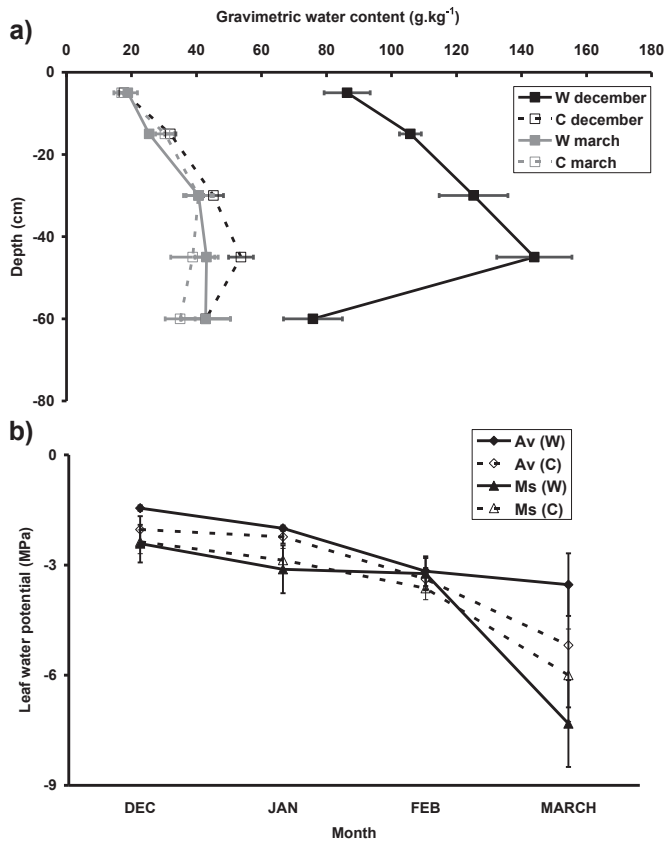


Fig. 1. (a) Gravimetric soil water content (GWC) at 5, 15, 30, 45 and 60 cm of depth for watered (black symbols and full lines) and control (white symbols and dotted lines) plots in the two extreme measurements of December (black color) and March (grey color). Values are average of soils below *Adesmia volckmanni* and *Mulinum spinosum* because we did not find significant species effects. Horizontal lines are standard errors. (b) Leaf water potential of *Adesmia volckmanni* (rhombi) and *Mulinum spinosum* (triangles), watered (black symbols and full lines) and control (white symbols and dotted lines) plants. Vertical lines are standard errors.

we used a $\alpha = 0.01$ significance level to characterize treatments effects. We used the angular arc sin \sqrt{p} transformation to analyze percentage data ($p = \text{percentage}/100$). All statistical analyses were performed with the software *Infostat* (Córdoba University,

Argentina) in combination with the package *nlme* from R environment (Pinheiro et al., 2014).

3. Results

Gravimetric soil water content increased from the upper layer to the 45 cm deep layer, and then decreased at 60 cm depth (Fig. 1a). This pattern was similar below both species at all depths and dates (Table 1), and increased in response to watering at every depth (Fig. 1a). However, the watering effect tended to disappear towards the end of the growing season (Fig. 1a; $T \times D$ and $T \times W$ interactions, in Table 1). Soil watering did not generate any change in leaf water potential of both species (Fig. 1b; Table 1). As a consequence of soil desiccation from December to March (Fig. 1a), leaf water potential decreased along the growing season in both species (Fig. 1b). Although soil water content did not differ between species at any depth, *A. volckmanni* showed higher leaf water potential than *M. spinosum* (Fig. 1b).

Foliar $\delta^{15}\text{N}$ was <0 (Fig. 2a), and did not change with watering nor along the growing season in both species (Table 1). Contrarily to expected, the non fixing species, *M. spinosum*, showed values higher, and closer to 0, than the presumably fixing one, *A. volckmanni* (Fig. 2a, Table 1). On the other hand, *A. volckmanni* showed, as usual in leguminous species, higher foliar N concentration than *M. spinosum*. In addition, the response of foliar N concentration to wetting, and their variation along the growing season, was different for the two species ($S \times T$ and $S \times W$ interactions; Table 1). The foliar N concentration of *A. volckmanni* showed a marginally significant increase in response to watering ($p = 0.07$; Fig. 2a) but did not show any pattern along the growing season, while that of *M. spinosum* showed a clear decrease along the growing season but did not change with watering (data not shown).

The $\delta^{15}\text{N}$ and N concentration of *A. volckmanni* nodules did not show any trend along the growing season or in response to watering (Table 1). Both variables showed higher values than those recorded at leaves (mean $\delta^{15}\text{N} = 0.23\text{‰}$ and mean N = 2.22%; Fig. 2a), and $\delta^{15}\text{N}$ was not statistically different from zero. The difference between the $\delta^{15}\text{N}$ of nodules and leaves averaged 1.23‰, and did not change with watering or along the growing season.

The natural abundance of ^{15}N of soil was >0 in all cases and was even higher than that of nodules (overall mean = 6.2‰; Fig. 2b). Soil $\delta^{15}\text{N}$ did not change along the growing season (Table 1), was higher in lower than in upper soil layers, and increased in response

Table 1
Statistical significance of the effects of watering (W), time (T), depth (D) and species (S) on soil and plant variables (D excluded for plant variables, and S excluded for nodule $\delta^{15}\text{N}$). We report Snedecor's F from a two way ANOVA (time \times watering) with plots splitted according a two way array (species \times depth) for the soil variables or a one way array (species) for the plant variables. Soil and plant N content (%) and Gravimetric Water Content (%) data were subjected to the angular transformation.***: $p < 0.001$; **: $0.001 < p < 0.01$; *: $0.01 < p < 0.05$ (in bold).

	Plant variables					Soil variables		
	ψ leaf (Mpa)	Leaf $\delta^{15}\text{N}$ (‰)	Leaf N content (%)	Nodule $\delta^{15}\text{N}$ (‰)	Nodule N content (%)	$\delta^{15}\text{N}$ (‰)	N content (%)	GSWC (%)
Species	7.8**	8.41**	9.47**			1.24	0.2	0.03
Time	465.7***	0.94	14.8**	1.55	0.23	1.04	0.39	45.7***
Watering	6.8	0.48	2.73	0.02	1.17	5.37*	0.35	85.13***
Depth						8.42***	11.02**	26.84***
S \times T	1.65	0.50	3.32*			0.66	1.10	1.16
S \times W	1.86	0.04	4.19*			3.39	1.61	2.32
S \times D						1.84	0.44	0.73
T \times W	0.04	1.47	0.55	0.76	0.34	0.87	0.51	24.99***
T \times D						2.19	0.95	3.85***
W \times D						1.68	1.28	1.30
S \times T \times W	0.92	0.55	0.07			0.002	0.006	0.6
S \times T \times D						2.55*	0.20	0.49
S \times W \times D						2.57*	3.82**	1.58
T \times W \times D						0.46	2.62*	1.92*
S \times T \times W \times D						0.31	0.25	0.83

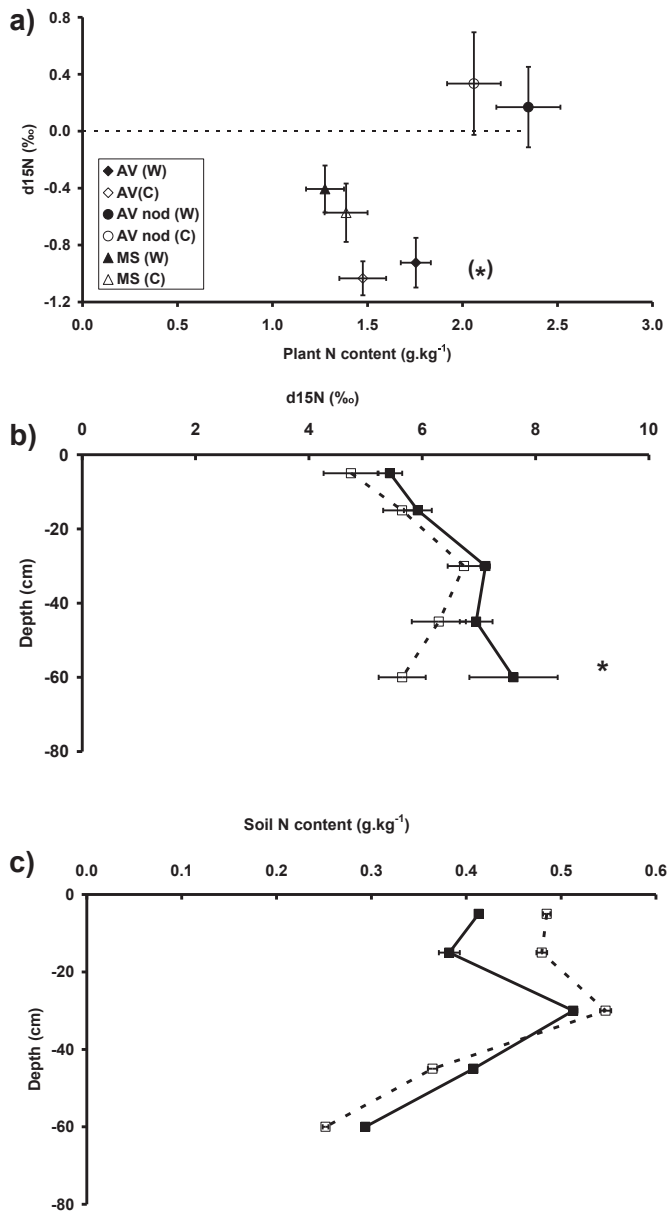


Fig. 2. (a) Natural abundance of ^{15}N ($\delta^{15}\text{N}$) in leaves of *Mulinum spinosum* (triangles) and *Adesmia volckmanni* (rhombi), and in nodules of *A. volckmanni* (circles) as a function of their leaf N content in watered (black symbols and full lines) and control (white symbols and dotted lines) plants. Vertical and horizontal lines are the respective standard errors. (b) Soil natural abundance of ^{15}N ($\delta^{15}\text{N}$), (c) and soil N concentration along the soil profile below watered (black symbols and full lines) and control (white symbols and dotted lines) plants. Values are average of soils below *Adesmia volckmanni* and *Mulinum spinosum* and all sampling dates because we did not find significant species or date effects. Horizontal lines are standard errors. Asterisks indicate significant differences between watered and control plots (within each depth in panels b to d) (LSD test): **: $p < 0.01$; *: $0.01 < p < 0.05$; (*): $0.05 < p < 0.10$.

to watering, mainly in the deepest layer (Fig. 2b). As soil $\delta^{15}\text{N}$, soil N concentration did not change along the growing season, but opposite to $\delta^{15}\text{N}$, it was higher in upper than in lower layers (Fig. 2c).

4. Discussion

This experiment questions the validity of *M. spinosum* as reference species to study biological N fixation in *A. volckmanni*, even being the species with the most similar root profiles and

phenological patterns (Golluscio et al., 2005, 2006), because the $\delta^{15}\text{N}$ of *M. spinosum* was higher, and closer to zero, than that of *A. volckmanni* (Fig. 2a). Even if the fraction of the leaf N obtained by *A. volckmanni* via biological fixation would have been minimal, its $\delta^{15}\text{N}$ would have been closer to zero than that of *M. spinosum* if both species would have utilized exactly the same sources of mineral soil N. The pattern we found would indicate that both species utilize different sources of mineral soil N. As leaf $\delta^{15}\text{N}$ of both species was lower than that of overall soil, it may be concluded that they use mineral N sources recently mineralized but not subjected to gaseous losses (Högberg, 1997). The deep root systems of both species (Golluscio et al., 2006) support the virtual absence of gaseous losses in the layers they explore. On the other hand, Gherardi Arbizu et al. (2013) demonstrated they prefer to absorb nitrate over ammonium, the former more frequent than the latter in deep layers because of their differential mobility in soil. As, in absence of gaseous losses, nitrate is more ^{15}N depleted than ammonium (Aranibar et al., 2008), the lower $\delta^{15}\text{N}$ of *A. volckmanni* respect to *M. spinosum* suggest that *A. volckmanni* absorb mineral N with lower ammonium/nitrate ratio than *M. spinosum*. This is consistent with the fact that root systems of *A. volckmanni* are slightly deeper than those of *M. spinosum* (Golluscio et al., 2006).

As demonstrated before, Patagonian shrubs showed little physiological responses to watering (Golluscio et al., 1998, 2009). In the present experiment, *M. spinosum* did not show any watering response, while *A. volckmanni* showed a little increment of leaf N concentration (Figs. 1 and 2). The virtual constancy of leaf ^{15}N natural abundance ($\delta^{15}\text{N}$) in response to watering and along the growing season (Fig. 2a and Table 1), as soil water content decreased (Fig. 1), suggests that *A. volckmanni* would not have obtained significant amounts of atmospheric N via biological fixation after the experimental watering. The existence of nodules may not be *per se* a sufficient evidence of N-fixation as perennial nodules have been reported in other harsh environments of the world, as the arctic tundra, which are not active during most of the year (Schulman et al., 1988). However, although the $\delta^{15}\text{N}$ of *A. volckmanni* was lower than that of air (Fig. 1a), it was 0.5‰ closer to zero in this humid year (186 mm, 24% above historic mean) than in the dry year of the former experiment (134 mm, 10% below historic mean; Golluscio et al., 2006). Furthermore, the 1.23‰ difference between the $\delta^{15}\text{N}$ of nodules and that of leaves, independent of watering, suggests that *A. volckmanni* plants must have obtained atmospheric N during the life of nodules (Wanek and Arndt, 2002). Probably N fixation may have taken place before watering, at the early spring following winter soil water recharge season. We expect our approach will encourage other researchers to develop specific models allowing the use of $\Delta\delta^{15}\text{N}_{n-1}$ to estimate N fixation in xeric shrubs.

Acknowledgments

We thank the unvaluable field aid of Natalie Dudinszky, Guillermo García Martínez, Fernando Cavagnaro, Magalí Valenta, and Paula Leva, the critical reading of Olga Correa, Agustín Grimoldi, Amy T. Austin, Marcela Gally, Roberto Fernández Aldúncin, Cristina Armas and an anonymous reviewer, the logistical facilities provided by INTA, and the financial support of Agencia Nacional de Promoción de la Ciencia y la Tecnología (PICT 0463), and University of Buenos Aires (UBA G044).

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