

The complexity-stability hypothesis in plant gas exchange under water deficit

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We hypothesized that more complex, i.e. irregular, temporal dynamics and a more interconnected overall network supports greater stability to gas exchange parameters (herein, CO₂ net assimilation and transpiration) in plants under water deficit. To test this hypothesis two genotypes of *Phaseolus vulgaris* were subjected to a period of absence of irrigation, and subsequent rewatering to achieve recovery. Gas exchanges parameters were measured each 10 s during 6 h to obtain time series to evaluate complexity by Approximate Entropy (ApEn) calculations, and network connectance in each water regime. Notably, the Jalo Precoce genotype showed significantly more stability than the Guarumbé genotype under system perturbation, coincident with greater irregularity in each gas exchange parameter and greater overall connectance for Jalo Precoce. This conclusion is consistent with other observations of greater homeostasis in more complex networks, seen in broad contexts such as cardiac rhythms and respiratory dynamics.

Key words: Approximate Entropy; complexity, network connectance, plant stress theory, photosynthesis, temporal dynamics, water deficit.

A hipótese sobre a relação entre complexidade e estabilidade em trocas gasosas de plantas sob deficiência hídrica: A hipótese testada neste estudo foi a de que dinâmicas temporais mais complexas, isto é, mais irregulares, e uma rede com maior conectância global suportariam maior estabilidade de parâmetros relacionados às trocas gasosas (assimilação líquida de CO₂ e transpiração) em plantas sob deficiência hídrica. Para tanto, dois genótipos de *Phaseolus vulgaris* foram submetidos a um período de suspensão da irrigação e subsequente re-hidratação para avaliar a recuperação das plantas. As trocas gasosas foram medidas em intervalos de 10 segundos durante 6 horas para a obtenção de séries temporais adequadas para os cálculos de complexidade por meio do uso da medida de Entropia Aproximada (ApEn), e conectância da rede de trocas gasosas em cada condição testada. Notavelmente, o genótipo Jalo Precoce mostrou significativa maior estabilidade que o Guarumbé, coincidindo com maior complexidade na dinâmica de cada parâmetro de trocas gasosas e maior conectância global no genótipo Jalo Precoce. Esse resultado foi consistente com outras observações de maior homeostase em redes mais complexas, em um contexto mais amplo, como observado em ritmos cardíacos e dinâmicas respiratórias.

Palavras-chave: Complexidade, conectância em redes, deficiência hídrica, dinâmica temporal, Entropia Aproximada, fotossíntese, teoria de estresse em plantas.

INTRODUCTION

The relationship between complexity and physiological stability has been observed among different kinds of biological systems. A greater irregularity (complexity) of each of cardiac rhythms, respiratory dynamics and blood pressure measurements has been associated with a healthier physiological status in humans. Conversely, and significantly, re-

duction of complexity in these biological parameters has also been associated with the aging process, and with increased risk of morbidity and mortality (Lipsitz and Goldberger, 1992; Pincus, 1995). Analogously in plants, we have previously observed that more complex temporal dynamics in parameters such as photosynthesis, enzymatic reactions and a broad class of fluxes are associated with a greater capacity

of system homeostasis (Hütt and Lüttge, 2002; Souza et al., 2004a). An increase in ecological system complexity has also been related to an increase in its overall stability (van Voris and O'Neill, 1980; Hastings et al., 1993; McCann, 2000).

Biological systems are organized as networks with a variety of interconnection and hierarchical attributes (Kauffman, 1993; Souza and Manzatto, 2000). The existence of significant redundancy (Edelman and Gally, 2001) within the network buffers the primary pathways or mechanisms within biological systems against external perturbations. Systems with sufficient redundancy provide a robustness in performance even when the system suffers an external disturbance, e.g., via transmission across alternate pathways, providing overall stability to the ensemble system (Amzallag, 2001; Edelman and Gally, 2001). Specifically, the quantity and the strength of the connections between network components have been directly correlated with the system stability (Trewavas, 1986; Edelman and Gally, 2001).

In this study, we considered the concept of stability as closely related to the classical concept of homeostasis. Stability is more suitable in studies considering dynamical aspects of biological systems such as herein (Souza and Buckeridge, 2004). However, since the original concept of homeostasis does not explicitly consider the ability of biological systems to dynamically reconfigure under environmental perturbations, authors such as Yates (1994) and Lloyd et al. (2001) have introduced a new and more broadly applicable concept, the homeodynamics concept, to account for such biological dynamical evolution. This revised concept is very interesting and could improve our understanding of plant ecophysiology, pending suitable care in interpretation of results.

Network connectance analysis has been proposed by Amzallag (2001) in plant physiological data, to assess the strength or degree to which specified individual network components are linked in an overall ensemble (see details in M&M). Further studies, such as Prado et al. (2004), Souza et al. (2004b) and Souza et al. (2004c), have considered the hypothesis that 1) systems showing more connected physiological networks are more able to maintain their stability, and 2), under external disturbances, physiological networks tend to maintain or increase the connectance between their parameters as a general response to such environmental variation.

Water deficiency is a common environmental factor constraining plants to express their ecophysiological potential. This deficiency leads to decrease in photosynthesis,

although tolerance may vary according to species (Kaiser, 1987; Chaves, 1991; Larcher, 1995; Chaves et al., 2002; Souza et al. 2003).

Souza et al. (2004a) found that sunflower plants, which are more tolerant to water deficit than sugar beet plants, showed more irregular (complex) temporal dynamics in their stomatal conductance runs at different water regimes. It was hypothesized that this complex temporal behavior allowed a more efficient adjustment to conditions of water deficiency. Such hypothesis was also corroborated by van Voris and O'Neill (1980) taking into account the respiration dynamics of a microecological community under different levels of soil toxicity.

In the present context, this study proposes to test the hypothesis that a higher complexity (irregularity) in temporal dynamics of assimilation of CO₂ and transpiration would be more efficient to the maintenance of homeostatic capacity of plants of common bean submitted to water deficit. To test this hypothesis we used two genotypes of *Phaseolus vulgaris* with distinct capacity to resist to water deficiency (Souza et al., 2003). Before submitting plants to a period of absence of irrigation, we measured gas exchanges parameters each 10 s during 6 h to obtain suitable time series in order to evaluate complexity by Approximate Entropy (ApEn) calculations (Pincus, 1991, Souza et al., 2004c). After a 6 day-period of irrigation suspension the plants were rewatered to test their homeostatic capacity.

ApEn assigns a non-negative number to a sequence or time-series, with larger values corresponding to greater apparent process randomness or serial irregularity, and smaller values corresponding to more instances of recognizable features or patterns in the data (Pincus, 1991; Pincus and Goldberger, 1994).

Furthermore, we also assessed the degree of connectance in the networks of gas exchange. These networks were formed by relationships between the physiological variables net CO₂ assimilation, transpiration, stomatal conductance and intracellular concentration of CO₂ (Souza et al. 2004b and c). We utilized this analysis to test the hypothesis that the plants with, *a priori*, higher network connectance would be more efficient to face water deficit.

MATERIAL AND METHODS

Plant material and data acquisition: This study was performed with two genotypes of common bean (*Phaseolus vulgaris* L.), the genotype Jalo Precoce (Type II), from Embrapa Arroz e Feijão, and the landrace Guarumbé from

Foz do Iguacu State Park – PR, Brazil. All seeds were kindly provided by germoplasm bank from Seeds Laboratory at UNOESTE, Presidente Prudente – SP, Brazil.

Plastic pots (5 L) filled with 4 Kg of substrate Plantmax (Eucatex-Agro, Brazil) were used, and irrigated with tap water until saturation. Three seeds of each genotype were sowed in each cup, and only one plant was maintained after complete seedling development.

Sixteen-day old individuals were submitted to water deficiency by suspension of irrigation for 6 days, when the water potential values reached around -1 MPa, the plants were rewatered. The experiment was carried out under greenhouse conditions. The control group was not water limited, being irrigated with 400 mL of water by replicate daily. All plants were supplied with 600 mL of nutrient solution (McCree, 1986) at the 8th and the 16th day after germination.

Water potential of plants under water deficit was measured each two days using dew point microvoltmeter (model HR-33T, Wescor) in a sample chamber (model C-52, Wescor).

Data acquisition was performed according to Souza et al. (2004c). Time series at steady state of gas exchange (net assimilation of CO₂ μmol.m⁻².s⁻¹, *A*, and transpiration mmol H₂O.m⁻².s⁻¹, *E*) were scored each 10 s during 6 h (from 9:00h AM to 3:00h PM) in control, water deficit, and after 36 h recovery treatments. Measurements were performed using an IRGA LI-6400 (LiCor, Lincoln, USA). Data were collected inside the laboratory and environmental conditions were maintained constant within the chamber at the temperature of 30°C, photosynthetic photon flux density (PPFD) of 800 μmol.m⁻².s⁻¹, and 370 ppm of CO₂. The difference of water pressure deficit between leaf and air was maintained at 1 kPa using a dew point generator (LI-610, LiCor, Lincoln, USA). The plants were placed into the laboratory at the experimental conditions one hour before starting the measurements. The same healthy and fully expanded central leaflet was analyzed in one plant per treatment (control, water deficit and recovery). The experiment was repeated three times in August and September 2002 and in February 2003, presenting suitable qualitative reproducibility. The time series shown in Figures 1 and 2 were derived from the third experimental repetition.

The noise in measurements (Pearcy et al., 1989) was evaluated with the empty leaf cuvette. This test showed that the noise due to the LI-6400 was very low (<2 % for net CO₂ assimilation and <0.005 % for stomatal conductance) and thus could be presently ignored. The coefficient of variation

(total CV %, which represents the sum of variations of the air flow rate, CO₂ and water vapor differentials) of the LI-6400 was around 0.1 % during the measurements. Furthermore, the LI-6400 typically takes new measures every 0.75 s, which is suitable for the baseline time interval used in this study (10 s).

Data analyses: To assess changes in system network aspects, we evaluated the occurrence of differences in the system modulation of gas exchange when submitted to different leaf water potentials via the concept and measurement of global connectance, *C_g* (Amzallag, 2001). Accordingly, to define connectance, first we specify a collection of paired variables of interest in the network. Next, we utilize the correlation coefficients (*r*) between each paired variable not only to test the significance of the correlation, but also as a measure of the strength of the relationship (connection) between the two parameters, by forming the z-transformation: $z = 0.5 \ln [(1 + |r|)/(1 - |r|)]$. Finally, we define the network global connectance (*C_g*) of the specified collection of paired variables as the average of the absolute z-values calculated above (Amzallag, 2001). In this study, we calculated *C_g* for the following collection of paired variables: *A* (net CO₂ assimilation) x *g_s* (stomatal conductance), *A* x *C_i* (internal CO₂ concentration), *A* x *E* (transpiration), *g_s* x *E*, and *g_s* x *C_i*. This technique was first used in time series by Souza et al. (2004c), providing some interesting and significant findings.

The irregularity of the time-series was assessed by ApEn. Two input parameters, a run length *m* and a tolerance window *r*, must be specified to compute ApEn. This parameter measures the logarithmic likelihood that runs of patterns that are close (within *r*) for *m* contiguous observations remain close (within the same tolerance width *r*) on next incremental comparisons. The opposing extremes are perfectly regular sequences, (e.g., sinusoidal behavior, very low ApEn), and independent sequential processes (very large ApEn). It is imperative to consider ApEn (*m, r*) as a family of parameters; comparisons are intended with fixed *m* and *r*.

When *m* = 1, as is employed herein, we interpret ApEn as a measure of the difference between the probability that runs of length 1 will recur within tolerance *r* and the probability that runs of length 2 will recur to the same tolerance. A high degree of regularity in the data would imply that a given (matched) run of length 1 would often continue with nearly the same third (next) value, producing a low value of ApEn.

ApEn evaluates both dominant and subdominant patterns in data; notably, it will detect changes in underlying episodic behavior not reflected in peak occurrences or amplitudes (Pincus and Keefe, 1992). Additionally, ApEn provides a direct barometer of feedback system change in many coupled systems (Pincus and Keefe, 1992; Pincus, 1994).

ApEn is nearly unaffected by noise of magnitude below r , a *de facto* filter level. ApEn is robust or insensitive to artifacts or outliers: extremely large and small artifacts have small effect on the ApEn calculation, if they occur infrequently. Both these points are evidently useful in clinical and experimental contexts, such as our present setting.

Further technical discussion of mathematical and statistical properties of ApEn, including mesh interplay, relative consistency of (m,r) pair choices, asymptotic normality under general assumptions, and error estimation for general processes can be found elsewhere (Pincus and Huang, 1992; Pincus and Goldberger, 1994). To develop a more intuitive, physiological understanding of the ApEn definition, a multistep description of its typical algorithmic implementation, with figures, is developed in Pincus and Goldberger (1994). Moreover, an extended discussion that covers a broad variety of both methodological aspects, plus considerably more background on the physiological meaning and scope of applicability of ApEn, can be found in Pincus (2000).

For the studies discussed below, ApEn values were calculated with widely established parameter values of $m = 1$, and $r = 20\%$ SD (standard deviation) of the gas exchange time-series (Souza et al., 2004c). *Normalizing* r to each time-series SD in this manner gives ApEn a translation- and scale-invariance (Pincus et al., 1993), in that it remains unchanged under uniform process magnification, reduction, or constant shift higher or lower. Multiple previous studies that included both theoretical analysis (Pincus, 1991; Pincus and Keefe, 1992; Pincus and Goldberger, 1994) and biological applications (Pincus et al., 1993; Pincus et al., 1996; Pincus et al., 1996; Christen et al., 1998; Pincus et al., 1998; Bruhn et al., 2000, Souza et al., 2004c and 2004d) have demonstrated that these input parameters produce good statistical reproducibility for ApEn for time-series of the lengths considered herein. In particular, one ApEn standard deviation ≤ 0.055 under very general conditions for ApEn for time-series of the lengths we analyze herein. Thus ApEn values that differ by 0.15 represent nearly 3 ApEn SDs,

indicating true distinction with error probability nearly $p = 0.001$.

Because the time-series of all physiological variables in the present experiments exhibit some nonstationary aspects, ApEn was applied to the first-differenced gas exchange time-series. This is a standard statistical method to stationarize time-series and is applicable to a very broad class of models (Chatfield, 1989; Willians, 1997); it showed utility in previous analysis of gas exchange dynamics (Souza et al., 2004a and c).

RESULTS

Effects of water deficiency on gas exchange: The period of suspension of irrigation imposed to both genotypes significant ($p < 0.05$) reductions in the gas exchange parameters A , E and g_s (table 1) and in the leaf water potential (table 2). Under control condition, both genotypes presented similar average rates of CO_2 assimilation. However, mean values of g_s and E were higher ($p < 0.05$) in the genotype Guarumbé (table 1), accordingly, water use efficiency (WUE) in the genotype Jalo Precoce was superior (table 2). In spite of the fact that E and g_s in Guarumbé have been relatively more affected by water deficiency than in Jalo Precoce, higher absolute values ($p < 0.05$) under water deficit were maintained in Guarumbé (table 1), still allowing a slight higher WUE in Jalo Precoce.

Thirty six hours after recovering, mean values of A were not completely restored to control levels in both genotypes, although they were higher ($p < 0.05$) than values during water deficiency (table 1). Jalo Precoce presented a complete recovery of E and g_s values, while Guarumbé still presented lower values in these parameters than under the control conditions (table 1). This likely explains lower values of WUE after recovery in Jalo Precoce (3.74) than in Guarumbé (4.75).

It is notable that leaf water potential has not returned to initial values in both genotypes after recovery, markedly in Jalo Precoce (Table 2). Nevertheless, Jalo Precoce exhibited more substantial and significant ($p < 0.05$) recovery in A than Guarumbé. While the latter recovered less than 50% of CO_2 net assimilation, the former showed a recovery of approximately 65%.

Effects of water deficiency on gas exchange networks: For each treatment, the coefficients of correlation (r) were calculated for each pair of parameters: A and g_s , A and C_i , A and E , g_s and E , g_s and C_i . In Jalo Precoce, all pairs of parameters were closely correlated ($p < 0.001$), especially the

pairs $g_s \times E$ and $g_s \times C_i$ ($p < 0.0001$). Conversely, in Guarumbé g_s showed no significant correlation with A ($p > 0.9$) and E ($p > 0.1$), suggesting an absence of a straightforward control of stomata on A and E , under irrigated condition. Since the relation between $A \times C_i$ and $g_s \times C_i$ was significant ($p < 0.001$), the stomatal regulation of A probably occurred through C_i (table 3). Under the water deficit condition, all parameters of Guarumbé were significantly correlated, especially $g_s \times A$ and $g_s \times E$ ($p < 0.0001$), suggesting that water limitation induced a strong stomatal control on A and E . Moreover, although there was variation in the extent of correlation r among tested pairs, all correlations between measured variables for Jalo Precoce remained significant

($p < 0.001$). The $A \times C_i$ relationship showed a negative correlation, which is expected because of the low stomatal and mesophyll conductance under water deficit (Flexas et al., 2004). Thus, while intercellular CO_2 is used as substrate for photosynthesis, supporting net photosynthesis, CO_2 uptake limited by low g_s causes a decrease in C_i . This also could help to explain the low correlation between g_s - C_i . As well, highly significant and strongly positive correlations among all pairs were observed during the recovery period. Guarumbé maintained a significant large positive correlation between g_s and E , and between g_s and C_i during recovery ($p < 0.001$); the other three pairwise comparisons, A - g_s , A - C_i and A - E now showed inverse (negative) correlations (table 3).

Table 1. Mean values and standard deviations of gas exchange parameters (net assimilation of CO_2 $\mu\text{mol}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$, A , transpiration $\text{mmol H}_2\text{O}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$, E , and stomatal conductance $\text{mol H}_2\text{O}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$, g_s , intercellular CO_2 concentration, $\mu\text{mol}\cdot\text{L}^{-1}$, C_i) measured each 10 s during 6 h in two genotypes of *P. vulgaris* under conditions of irrigation (control), water deficit, and after 36 h of recovering.

	Jalo Precoce			Guarumbé		
	control	deficit	recovery	control	deficit	recovery
A	15.18±1.56	2.88±1.04	9.84±1.69	15.50±0.54	4.43±0.98	7.45±1.66
E	2.19±0.51	0.46±0.08	2.63±0.94	5.76±0.71	0.80±0.16	1.57±0.19
g_s	0.130±0.040	0.029±0.007	0.190±0.110	1.066±0.141	0.043±0.014	0.061±0.014
C_i	183.78±21.94	196.16±13.61	221.21±37.17	270.04±10.51	172.50±44.71	155.78±63.14

Table 2. Values of leaf water potential (ψ , MPa) and water use efficiency (WUE, $\mu\text{mol CO}_2\cdot\text{mmol}^{-1} \text{H}_2\text{O}$) of *P. vulgaris* under conditions of irrigation (control), water deficit, and after 36 h of recovering.

	Jalo Precoce			Guarumbé		
	control	deficit	recovery	control	deficit	recovery
Ψ	-0.5±0.04	-1.0±0.14	-0.7±0.08	-0.5±0.08	-1.1±0.10	-0.6±0.1
WUE	6.88	6.26	3.74	2.69	5.54	4.75

Table 3. Values of correlation coefficients (r) for pairs of gas exchange parameters in two genotypes of *P. vulgaris* under conditions of irrigation (control), water deficiency, and after 36 h of recovery. These values were calculated by linear regression of the complete time series data. Except for $A \times g_s$ ($p = 0.909$) and $g_s \times E$ ($p = 0.194$) at control condition in Guarumbé, all others r values showed $p < 0.001$.

	Jalo Precoce			Guarumbé		
	control	deficit	recovery	control	deficit	recovery
A - g_s	0.593	0.929	0.726	-0.003	0.775	-0.366
A - C_i	0.510	-0.122	0.728	0.846	0.270	-0.405
A - E	0.597	0.873	0.793	0.808	0.729	-0.279
g_s - E	0.977	0.965	0.954	-0.031	0.947	0.672
g_s - C_i	0.841	0.139	0.688	-0.177	0.714	0.792
C_g	1.077	1.055	1.129	0.515	0.987	0.598

The r values were transformed into z values (see Material and Methods) for calculation of global connectance (C_g), which is the average of z values for each pair of parameters in each treatment. According to Gardner and Ashby (1970), the connectance quantifies the level of linkage within a system. Under the control condition, C_g in Jalo Precoce was 52 % higher than in Guarumbé. However, under water deficiency C_g of Guarumbé increased 47.8 % compared to control, while C_g value in Jalo Precoce remained practically unchanged, but still greater than in Guarumbé (table 3). Finally, after the recovery period, C_g in Guarumbé decreased 39 %, reaching a value close to the baseline control condition. In considerable contrast, C_g in Jalo Precoce remained very stable from baseline through deficit to recovery, with a slight final stage increase of 7 % (table 3). These results suggest that water deficiency had substantially greater effect on overall gas exchange network linkages in Guarumbé than in Jalo Precoce.

Steady-state temporal dynamics in A and E runs: Although all data have been measured under constant environmental conditions, different temporal dynamic patterns can be observed (figures 1 and 2). Both genotypes presented oscillations in A and E runs during the measurements period. Oscillations in A and E were synchronized, i.e., there was an agreement between valleys and peaks for these parameters throughout measurements, as may be observed by the correlation coefficients between A and E , except for Jalo Precoce under control conditions and Guarumbé after recovery (table 3). However, while Guarumbé presented only a single oscillation period of 33 min, Jalo Precoce showed two frequencies, a longer one of 33 min, and a short one of 15 min. There also was a greater variation in the amplitude of oscillations in Jalo Precoce runs (figure 1).

According to visual comparison of the presented dynamics (figures 1 and 2), it can be observed that under water deficiency, both genotypes lost their primary oscillatory behavior, and only Jalo Precoce tended to recover it after rewatering.

Values of ApEn for time series at steady state of A and E under the control condition showed that complexity (irregularity) in both series were greater for Jalo Precoce (table 4). This tendency of higher values of ApEn for Jalo Precoce were observed in other repetitions of the experiment (data not shown), confirming a greater irregularity in temporal dynamics for A and E in this genotype. This difference in values of ApEn between the genotypes was

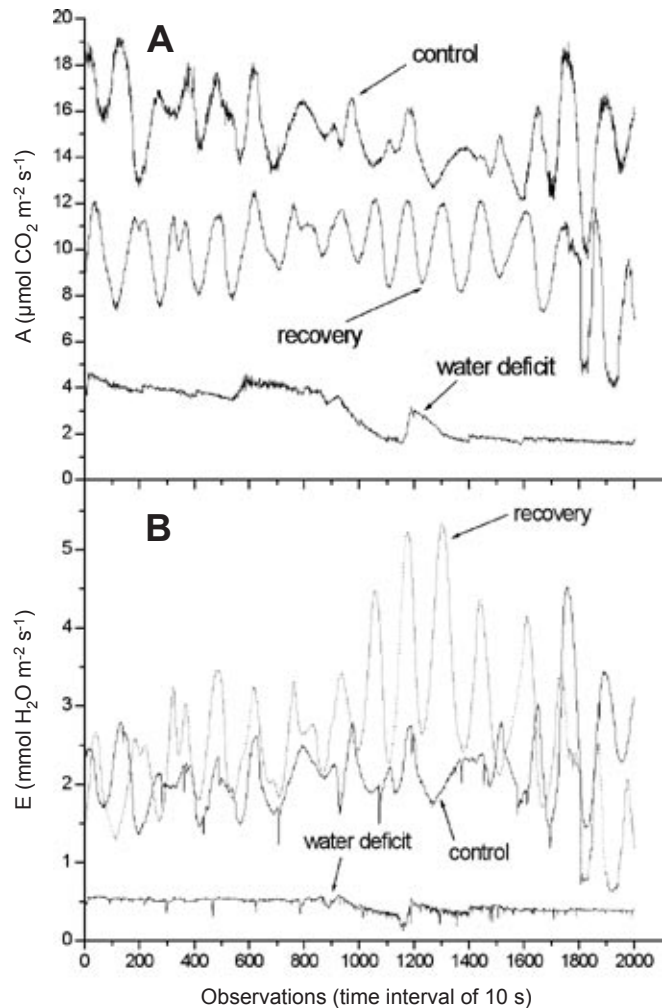


Figure 1. Temporal dynamics of A) net CO₂ assimilation, A , and B) transpiration, E , for Jalo Precoce under conditions of irrigation (control), water deficiency, and after 36 h of recuperation. The series were evaluated in similar healthy totally expanded leaves and data were taken each 10 s during 6 h.

Table 4. Calculated mean values of Approximate Entropy (ApEn) of steady state time series of net CO₂ assimilation (A) and transpiration (E) in two genotypes of *P. vulgaris* under control, water deficit and recovery treatments. (ApEn parameters $m = 1$ and $r = 20$ % of SD).

	Jalo Precoce	Guarumbé
	<i>A</i>	
Control	2.041	1.414
Water deficit	1.884	1.945
Recovery	1.744	1.740
	<i>E</i>	
Control	1.395	1.195
Water deficit	1.397	2.437
Recovery	1.467	1.425

most pronounced in A , and statistically significant ($p < 0.05$), based on all experimental repetitions. Although for each experiment repetition the values of ApEn in the E time series for Jalo Precoce were always higher than for Guarumbé, we did not quite achieve statistical significance here. As previously suggested by Souza et al. (2004c) in a temporal dynamic study of gas exchange with *Citrus sinensis* under situation of different VPD , the variable A seemed to be the more sensitive of the two in exhibiting dynamical differences due to environmental effects in gas exchange time series.

While the runs became more irregular under water deficit in Guarumbé, the runs of genotype Jalo Precoce remained approximately the same, tending towards a lower irregularity. After the recovery period the complexity of all time series tended to be lower than under water deficit, except for transpiration in Jalo Precoce (table 4). These trends were also observed in temporal dynamics of sap flow in tropical tree species under different water regimes (Souza et al., 2004d).

DISCUSSION

Water content has a pronounced effect on photosynthesis. Water deficiency leads invariably to a decrease in photosynthetic rate, although levels of tolerance can vary for different plant species (Kaiser, 1987; Chaves, 1991; Larcher, 1995; Chaves et al., 2002). Among the factors that contribute to this photosynthesis reduction we could consider stomatal closure in direct response to a reduction of leaf water potential. A reduction of stomatal conductance limits intercellular CO_2 reposition, reducing C_i , consequently constraining net CO_2 assimilation (Jones, 1998; Cornic, 2000; Chaves et al., 2002). As a result of stomatal closure, loss of water to atmosphere is also reduced, contributing to the maintenance of plant water potential (Jones, 1998; Chaves et al., 2002). Under moderate water restriction, with a partial stomatal closure, photosynthesis is less affected than transpiration, increasing WUE (A/E). This WUE increase was clearly observed in Guarumbé under water deficiency (table 2). Nevertheless, WUE decreases quickly as stomata are almost closed (Chaves, 1991; Larcher, 1995; Nobel, 1999).

Thus, under water deficiency, stomata assume a central role in gas exchange control (Chaves, 1991; Jones 1998). Therefore, we assumed herein that values of C_g might be especially dependent on g_s . In addition to the direct effect of g_s over A and E , there may exist multiple feedback processes acting directly or indirectly on g_s . Moreover, environmental factors also can influence any of these variables and their

feedback linkages (Farquar and Sharkey, 1982; Zeiger et al., 1987; Jones, 1998). Accordingly, we proposed a diagram to represent gas exchanges regulation in this study, based upon Jones (1998) (figure 3). A greater stomatal control is a desirable character in plants in order to tolerate water deficiencies, allowing a suitable adjust in gas exchange, resulting in a faster photosynthesis recovering after rewatering (Kramer and Boyer, 1995).

The results of gas exchange network analyses (table 3) suggest that a higher C_g in Jalo Precoce, at initial irrigated condition, may facilitate system homeostasis, compared with the Guarumbé genotype. We have suggested that higher values of C_g provides a better system adjustment under environment perturbations, permitting the system to adjust itself faster in response to environmental stimulus. Additionally, environmental restrictions may canalize the system, reinforcing linkages between network components in the face of a particular environment condition (stress), thus maintaining its homeostasis (Souza et al., 2004b,c). Both these two behaviors were observed in this study. Although Guarumbé increased C_g under water deficiency, Jalo Precoce maintained high values of C_g in all experimental states, supporting our homeostatic thesis. Similar results to the present analysis were obtained in gas exchange responses to water deficiency in two tropical tree species (Souza et al., 2004b). In that study, the species with higher C_g values did not exhibit significant physiological change in this measure under water deficit, whereas the other species, with lower values of C_g , presented a limited and diminished recuperation after the water deficit condition.

The results presented herein (table 1) showed that Jalo Precoce had a greater stomatal control on gas exchanges, reaching smaller values of g_s than Guarumbé in water deficit condition. The greater stomatal control of Jalo Precoce is supported by higher values of r observed in the relationships between $A \times g_s$ and $E \times g_s$ (table 2).

The hypothesis in this study was that more irregular (complex) temporal dynamics and a more interconnected overall network could support higher stability to CO_2 net assimilation and to transpiration in plants under water deficit.

Although both genotypes exhibited apparently oscillatory behavior (figures 1 and 2), ApEn results showed consistent differences between E and A dynamics for both genotypes. Moreover, results of ApEn calculations (table 4) could be directly related to physiological parameters that showed higher recovery after water deficiency, i.e., transpiration and CO_2 net assimilation in the Jalo Precoce

genotype. This relation found herein and observations from Souza et al. (2004c) on gas exchanges temporal dynamics of *C. sinensis* under different VPD situations support the hypothesis that systems with more irregular temporal dynamics may have greater tolerance to environmental perturbations, thus maintaining greater system stability (homeostasis). Therefore, just as for heart rate, respiration and blood pressure (Lipsitz and Goldberger, 1992; Pincus, 1995), more robust systems had higher ApEn values, and were more stable, fluid or less affected by external perturbations. Similarly here, Jalo Precoce showed higher baseline ApEn than Guarumbé genotype, and again was more stable under perturbation.

Furthermore, the overall variation in ApEn values ($T_{\text{var}}\%$), over the three treatment conditions, calculated as: $T_{\text{var}}\% = \frac{\text{max of the mean ApEn values} - \text{min of the mean ApEn values}}{\text{average of the three ApEn values}} \times 100$, showed that the runs of Jalo Precoce were more stable than in Guarumbé. Consideration of the time series of net CO_2 assimilation, while $T_{\text{var}}\%$ in Jalo was 15.7%, in Guarumbé $T_{\text{var}}\%$ was 31.2%. This difference was even profoundly more significant for the transpiration time series. $T_{\text{var}}\%$ in Jalo was nearly a factor of 15 smaller (more stable) than that for Guarumbé, respectively, 5.0% compared to 73.7%. These results support the hypothesis that the genotype that showed a more complex baseline can retain its stability (homeostasis) with greater efficiency.

Plants, as biological systems, are dynamic systems, thermodynamically in non equilibrium. This implies that those systems, especially their physiological processes, evolve during time, presenting a variability of dynamic behaviors. Such behaviors may be significantly influenced by environment stimuli (Hütt and Lüttge, 2002; Souza et al., 2004a). Complex temporal dynamics can be more efficient in supporting environmental perturbations due to its greater intrinsic flexibility of responses, which can then more effectively stabilize the system, i.e., preserve homeostasis. Van Voris and O'Neill (1980) verified that a greater complexity in respiratory metabolism in a micro-ecosystem was associated with the ability to support higher doses of heavy metals. Møller et al. (1998) in a study of chaotic dynamics in peroxidase catalyzed reactions found that such irregular dynamics improved system stability. Finally, in studies with stomatal conductance temporal dynamics Souza et al. (2004a) observed that plants with higher complexity in g_s time series were more able to recover gas exchanges status after period of water deficit.

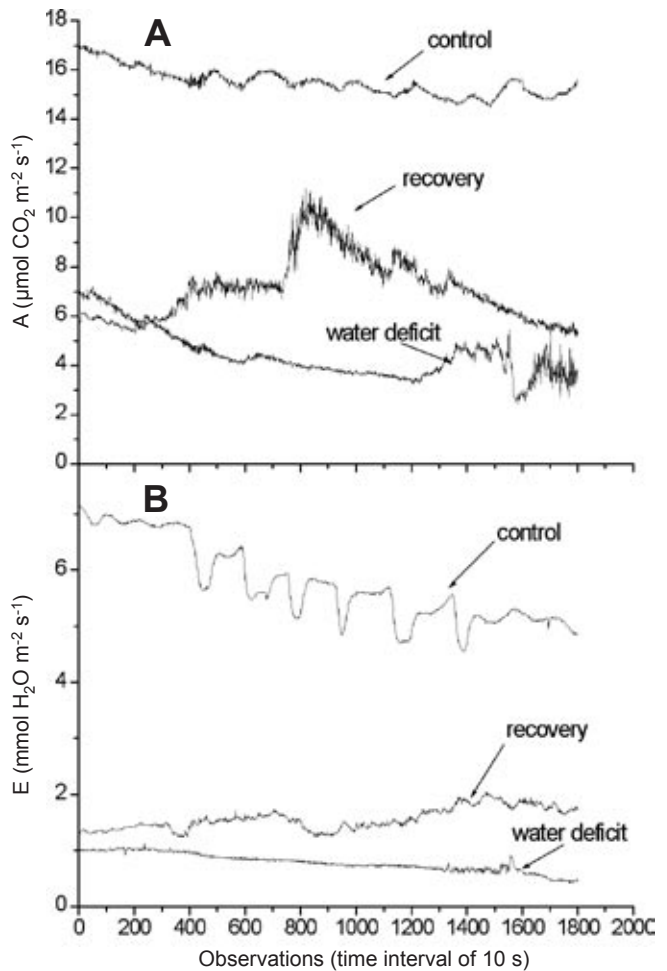


Figure 2. Temporal dynamics of A) net CO_2 assimilation, A , and B) transpiration, E , for Guarumbé under conditions of irrigation (control), water deficiency, and after 36 h of recuperation. The series were evaluated in similar healthy totally expanded leaves and data were taken each 10 s during 6 h.

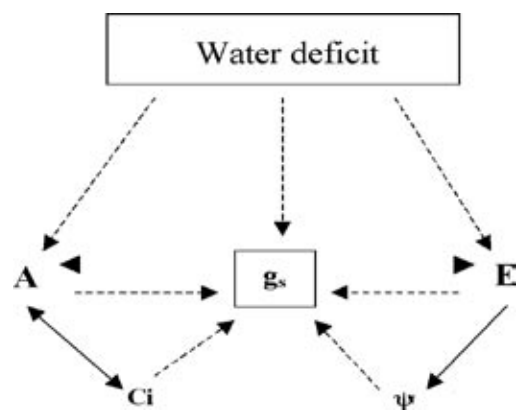


Figure 3. Schematic representation of gas exchange relationships under water deficit. A = CO_2 net assimilation, E = transpiration, g_s = stomatal conductance, C_i = intercellular CO_2 concentration, ψ = leaf water potential. Adapted from Jones (1998).

Generally, studies of temporal dynamics on plant physiology, particularly on gas exchanges, are restricted to the rhythms of circadian cycles (Fredeen et al., 1991; Hennessey and Field, 1991; Hennessey et al., 1993; Webb, 1998; Rasher et al., 2001). Nevertheless, variable stomatal conductance behavior across a smaller time scale (minutes or seconds) has been observed (Cardon et al., 1994; Haefner et al., 1997; Mott and Buckley, 1998 and 2000; Souza et al., 2004a). The phase, amplitude and period of stomatal oscillations can substantially vary among patches in a single leaf (Cardon et al., 1994; Mott and Buckley, 1998). This variation of temporal behavior among patches could reflect in a higher or lower irregularity of stomatal conductance dynamics measured by IRGAs (Infra red gas analyzers), which often sample a leaf area containing many patches (Souza et al., 2004a). Various mechanisms have been proposed to explain oscillatory behavior of stomata. Among the most important ones, we especially note those based upon hydraulic mechanisms of interaction between epidermis cells and the adjacent stomata (Sharpe et al., 1987; Haefner et al., 1997), and the influence of oscillations in calcium intracellular concentrations on stomatal aperture (Allen et al., 2001; Evans et al., 2001; Ng and McAinsh, 2003). As both mechanisms presented could be influenced by environmental changes, the former influenced by humidity (Haefner et al., 1997; Mott and Buckley, 1998) and the latter by different environmental factors, since calcium has been considered as an important secondary messenger in response to environmental perturbations (Reddy, 2001), it might be expected that temporal dynamics of stomatal conductance and, consequently, leaf gas exchanges, were influenced by other external stimuli.

Concluding remarks: The physiological studies discussed herein clarify that common bean gas exchanges can exhibit well-defined, non-random network structure, as well as subtle, yet consistent irregularities in individual variable dynamics. Accordingly, given the absence of external influences, above we inferred intrinsic differences between plant genotypes, under several different physiological states, on the basis of both network and individual variable analysis. The assessment of both connectance (C_g) and sequential irregularity ($ApEn$) was quite useful in evaluating endogenous processes of physiological regulation. Thus, these two entirely distinct and complementary statistical perspectives to investigate plant stability showed corroborative findings supporting our homeostatic hypothesis.

We proposed that a greater system connectance can support a greater complexity in the temporal dynamics of plants, since systems with a high degree of internal connections could more rapidly make fine tuning adjustments. Moreover, our results support the hypothesis of Souza et al. (2004b) that an increase in connectance (C_g values) indicates plant responses to environmental perturbations, improving their resistance, and that systems with a high degree of C_g would be more tolerant to environmental restrictions. In this context, we regard resistance as the capacity of the plants to oppose environmental perturbation due to modifications in their regulation network, and tolerance as the capacity of plants to support environmental changes without significant alterations to their system. Thus, recalling the results of the connectance analyses, Jalo Precoce was the more tolerant genotype, while Guarumbé showed more predominant resistance. This conceptual ecophysiological framework is consistent with the themes of a more systemic stress framework proposed by Souza and Cardoso (2003).

Finally, the results presented herein are somewhat specific to our experimental model, and should be corroborated with subsequent experiments incorporating different sets of environmental variables as well as different species, to verify the robustness and generality of the stability-complexity relation hypothesis.

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