Plant electrophysiological information manifests the composition and nutrient transport characteristics of membrane protein

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Abstract

Background: Almost all life activities of plants are accompanied by electrophysiological information. Plant electrical parameters are considered to be the fastest response to environment.

Results: In this study, the theoretically intrinsic relationships between the clamping force and leaf resistance (R), capacitive reactance (Xc) and inductive reactance (XL) were revealed as 3-parameter exponential decay based on bioenergetics for the first time. The intrinsic resistance (IR), intrinsic capacitive reactance (IXc) and intrinsic inductive reactance (IXL) in plant leaves were monitored via these relationships for the first time, and the nutrient transport capacity (NTC) in plant cells based on IR, IXc and IXL was first defined. The results indicate that IXc and IXL could be used to manifest the composition of surface and binding proteins in cell membrane, plant with high crude proteins and crude ash had higher NTC, and which accurately revealed the nutrient transport strategies in tested plants.

Conclusions: This study highlights that plant electrophysiological information could effectively manifest the composition and nutrient transport characteristics of membrane protein in plant cells.

Keywords: electrophysiological information, bioenergetics, membrane protein composition, nutrient transport.
Almost all life activities in plants, including the metabolism of substances and energy, development, stress resistance and signal transduction, involve charge separation, electron movement, proton and dielectric transport, etc. [1-2]. The electrical properties of plant cells are derived from the cell membrane with a double electric layer, which is two electron density bands approximately 2.5 nm thick on the inside and outside of the membrane and a transparent band approximately 2.5 nm thick in the middle. And membrane lipids and proteins, the mainly compositions of cell membrane, can be regarded as insulating layer, have a high electrical resistivity, enabling the plant cell to store electric charge [3]. Therefore, the electrophysiological information in plants is closely related to the life activities, and the changes of structure, composition and ion permeability in plant cells will inevitably lead to significant changes in electrophysiological information [3-8].

Plant electrical parameters are considered to be the fastest response to environmental stimulus such as drought, salt stimulation, cold stimulation, diseases and insect pests, exogenous force [7-11]. Previously, a traditional approach, the electrical parameters in plants are measured by the insertion of two electrodes into the stem or leaf [12-13]. However, this method is unstability and difficult to manipulate, and the plant electrical signals acquired lacked representativeness, reproducibility and comparability needling injury, as well as different environments, users and other factors. Moreover, the intrinsic or spontaneous electrical parameters in plants are not detected by previously methods. Thus, can the intrinsic relationships between environmental stimulus and electrophysiological parameters be feasible to obtain intrinsic electrical parameters with high reproducibility in plants or evaluate their life phenomena? Can these intrinsic relationships be described by corresponding physical mechanism models?

Generally, a mesophyll cell can be regarded as a concentric sphere capacitor with both inductor and resistor function, and many aligned mesophyll cells make up the leaf capacitor [2, 14]. The ions, ion groups and electric dipoles in mesophyll cells are electrolytes of leaf capacitor and most related to electrophysiological information [15]. Interestingly, Guo et al. [16] reported the capacitance (C) values of maize leaves increased with clamping forces, and manifested clamping forces stimuli changed the electrophysiological information in plant leaves. However, this intrinsic mechanism or relationship between clamping force and the electrophysiological information of plant leaves wasn't revealed. Thus, it is of great practical significance to clarify the intrinsic mechanism between clamping forces and electrophysiological parameters and provide a rapid, accurate and real-time technique for monitoring the physiological state of plant leaves.
Cell is the site of all biochemical reactions, and cell membrane side is an important barrier to ensure a stable environment inside the cell. It has been estimated that 15~30% of the nuclear gene encoded proteins are involved in nutrient transport on the cell membrane, and the energy used by cells in nutrient transport up to two-thirds of the total energy consumed by cells [1]. The nutrient transport capacity of cells is most closely related to the type and quantity of surface and binding proteins in cell membrane, thus, the composition and content of membrane protein can indirectly reflect the nutrient transport capacity of cells. Protein detection methods of biological samples include conventional, electrochemical, molecular biology, electrophoresis and mass spectrometry methods [17]. However, the detection of membrane proteins is limited to single cell or single proteins, and the existing protein detection technology is difficult to accurately evaluate the composition characteristic of cell membrane protein [17-18]. Moreover, the nutrient transport capacity ultimately affects the nutrient use efficiency of plants, and the most commonly used method of plant nutrient use evaluation is the ratio of total nutrient in plants to total input nutrient [19-20]. However, this nutrient use efficiency also does not directly reflect the nutrient transport capacity. To the best of our knowledge, the composition and nutrient transport characteristics of membrane protein has rarely been reported.

The fully expanded leaves, which account for a high proportion of plant biomass, determine and reflect the plant nutrient metabolism. Since the concentration of electrolytes in cells (ions, ion groups and electric dipoles) in leaf cells is directly affected by the nutrient metabolism in plant leaves, and then it is accompanied by vigorously electrical activities. In this study, it was first clarified and constructed the intrinsic mechanisms and physical models between clamping forces and leaf resistance (R), capacitive reactance (Xc) and inductive reactance (XL). Subsequently, the intrinsic electrophysiological parameters in plant leaves were monitored through these mechanism equations. And then the nutrient transport capacity (NTC) in plant leaves in the light of the intrinsic electrophysiological parameters was defined to evaluate the nutrient transport strategies of various tested plants. This study aims to clarify the intrinsic mechanisms among the leaf R, Xc and XL and exogenous stimuli, and provide a novel, feasible technique for real-time monitoring plant nutrient transport.

**Results**

**Intrinsic mechanism relationships of clamping force (F) and leaf R, Xc and XL**
Almost all life activities in plants are closely related to the electrophysiological information. In a mesophyll cell, cell membrane has strict selective permeability to various ions, ion groups and electric dipoles, and the electrolyte solution on both sides of cell membrane forms a specific conductive state. The inside and outside of cell membrane can be simulated as a capacitor, the electrolyte solution on both sides of the membrane is equivalent to the two plates of the capacitor, and cell membrane is equivalent to intermediate medium of the capacitor. Moreover, organelles such as vacuoles and cytoplasm in cells are equivalent to resistors. Thus, mesophyll cell can be regarded as a concentric sphere capacitor with both inductor and resistor functions. The simplified equivalent circuit of mesophyll cell is displayed in Fig. 1.

Fig. 1.

Plant electrophysiological information obtained by the traditional needling method is often less authenticity, reproducibility and comparability due to needling injury, different environments and technicians, and other factors [11]. The ions, ion groups and electric dipoles in the plant leaf were used as electrolytes, and a parallel-plate capacitor sensor could be formed by placing the leaf between the two plates of the parallel-plate capacitor. The leaf R, Xc and XL varied with the ions, ion groups and electric dipoles concentrations in the plant leaf, and different clamping forces which can be regarded as different exogenous stimuli inevitably lead to changes the ions, ion groups and electric dipoles concentrations in plant leaves. To obtain authentic, comparable and reproducible plant electrophysiological data, the intrinsic mechanism models between the clamping force and leaf R, Xc and XL were revealed.

The concentration of the electrolytes determines inside and outside R of the cell membrane. External stimuli change the membrane permeability of the electrolytes and affect their inside and outside concentration of the cell membrane. Under different clamping forces, the membrane permeability of the electrolytes that respond to R in the plant cell membrane changed. According to the bioenergetics, the Nernst equation can be used to quantitatively describe the potential of electrolytes inside and outside of the cell membrane. Thus, the concentration differences in the electrolytes that respond to inside and outside R of the cell membrane obey the Nernst equation and can be expressed as follows:

\[ E - E^0 = \frac{R_0 T}{n R F_0} \ln \frac{C_i}{C_o} \]  

(1)

where \( E \) = the electromotive force (V), \( E^0 \) = the standard electromotive force (V), \( R_0 \) = the gas constant (8.314570 J...
\( K^{-1} \text{ mol}^{-1} \), \( T \) = the thermodynamic temperature (K), \( C_i \) = the concentration of the electrolytes that respond to \( R \) inside the cell membrane (mol L\(^{-1} \)), \( C_o \) = the concentration of the electrolytes that respond to \( R \) outside the cell membrane (mol L\(^{-1} \)), \( F_0 \) = Faraday constant (96485 C mol\(^{-1} \)), and \( n_R \) = the number of transferred electrolytes (mol).

The internal energy of the electromotive force can be converted into pressure work, and they have a direct relationship, \( PV = aE \), that is:

\[
PV = aE = aE^0 + \frac{aR_0T}{nRF_0} \ln \frac{Q_i}{Q_o} \quad (2)
\]

where \( P \) = the pressure intensity on the leaf cells (Pa), \( a \) = the energy conversion coefficient of the electromotive force, and \( V \) = the cell volume (m\(^3\)). \( P = \frac{F}{S} \), where \( F \) = the clamping force (N) and \( S \) = the effective area of the electrode plate (m\(^2\)). \( F \) can be calculated by the gravity formula:

\[
F = (M + m)g \quad (3)
\]

where \( M \) = the iron block mass (kg), \( m \) = the mass of the plastic rod and the plate electrode (kg), and \( g \) = 9.8 N/kg.

For mesophyll cells, the sum of \( C_o \) and \( C_i \) is certain. \( C_i \) is directly proportional to the conductivity of the electrolytes that respond to \( R \), and the conductivity is the reciprocal of \( R \). Hence, \( C_i/C_o \) can be expressed as \( \frac{C_i}{C_o} = f_0 \frac{f_0}{C_T R - f_0} \), where \( f_0 \) = the ratio coefficient of the conversion between \( C_i \) and \( R \), and \( C_T = C_o + C_i \). Therefore, formula (2) was transformed into formula (4):

\[
\frac{V}{S} F = aE^0 + \frac{aR_0T}{nRF_0} \ln \frac{C_T R - f_0}{f_0} \quad (4)
\]

Formula (4) was rewritten:

\[
\frac{aR_0T}{nRF_0} \ln \frac{C_T R - f_0}{f_0} = aE^0 - \frac{V}{S} F \quad (5)
\]

and

\[
\ln \frac{C_T R - f_0}{f_0} = \frac{nRF_0E^0}{S R_0T} \frac{V}{S} aR_0T F \quad (6)
\]

Formula (6) takes the exponents of both sides:

\[
\frac{C_T R - f_0}{f_0} = e^{\frac{nRF_0E^0}{S R_0T}} \left( \frac{V}{S} aR_0T F \right) \quad (7)
\]

Further:
\[ R = \frac{f_0}{C_T} + \frac{f_0}{C_T} e^{\frac{n_R F_0^0}{R_0^T}} e\left(\frac{-V n_R F_0^0}{a R_0^T} \right) \]  

(8)

Because \( d = \frac{V}{a} \), formula (8) was transformed into:

\[ R = \frac{f_0}{C_T} + \frac{f_0}{C_T} e^{\frac{n_R F_0^0}{R_0^T}} e\left(\frac{-d n_R F_0^0}{a R_0^T} \right) \]  

(9)

For the same leaf tested in the same environment, the \( d, a, E^0, R_0, T, n_R, F_0, C_T, \) and \( f_0 \) of formula (9) are constant.

Let \( y_0 = \frac{f_0}{C_T} \), \( k_1 = \frac{f_0}{C_T} e^{\frac{n_R F_0^0}{R_0^T}} \), \( b_1 = \frac{d n_R F_0^0}{a R_0^T} \), and the intrinsic mechanism relationships of leaf \( R \) and \( F \) was:

\[ R = y_0 + k_1 e^{-b_1 F} \]  

(10)

where \( y_0, k_1 \) and \( b_1 \) are model parameters.

When \( F=0 \), the intrinsic resistance (IR) of the plant leaves could be obtained:

\[ IR = y_0 + k_1 \]  

(11)

With the same \( R \), the intrinsic mechanism relationships of leaf \( X_c \) and \( F \) was revealed (Additional file 1):

\[ X_c = p_0 + k_2 e^{-b_2 F} \]  

(12)

where \( p_0, k_2 \) and \( b_2 \) are model parameters.

When \( F=0 \), the intrinsic capacitive reactance (IXc) of plant leaves could be calculated as:

\[ IX_c = p_0 + k_2 \]  

(13)

With the same \( R \), the intrinsic mechanism relationships of leaf \( X_L \) and \( F \) was revealed (Additional file 1):

\[ X_L = q_0 + k_3 e^{-b_3 F} \]  

(14)

where \( q_0, k_3 \) and \( b_3 \) are model parameters.

When \( F=0 \), the intrinsic inductive reactance (IXL) of plant leaves could be calculated as:

\[ IX_L = q_0 + k_3 \]  

(15)

**Determination of the nutrient transport parameters**
The IR of the plant leaves is calculated according to formula (16):

\[
\frac{1}{\text{IR}} = \frac{1}{\text{IR}_1} + \frac{1}{\text{IR}_2} + \frac{1}{\text{IR}_3} + \ldots + \frac{1}{\text{IR}_n}
\]  

(16)

It is assumed that the membrane inside and outside resistance of each cell is equal, then IR$_1$, IR$_2$, IR$_3$, … IR$_n$ can represent intrinsic resistance of each unit cell membrane. It is assumed that the intrinsic resistance of each cell membrane is equal, that is IR$_1$=IR$_2$=IR$_3$=…=IR$_n$=IR$_0$. Thus, the IR of the plant leaves was obtained:

\[
\frac{1}{\text{IR}} = \frac{n}{\text{IR}_0}
\]  

(17)

Due to membrane resistance is most closely related to proteins and lipids of cell membrane, then n can be characterized as the amount of proteins and lipids that induce membrane resistance in plant leaves.

The IXc of the plant leaves is calculated according to formula (18):

\[
\frac{1}{\text{IXc}} = \frac{1}{\text{IXc}_1} + \frac{1}{\text{IXc}_2} + \frac{1}{\text{IXc}_3} + \ldots + \frac{1}{\text{IXc}_p}
\]  

(18)

It is assumed that the membrane inside and outside capacitive resistance of each cell is equal, then IXc$_1$, IXc$_2$, IXc$_3$, … IXc$_p$ can represent intrinsic capacitive resistance of each unit cell membrane. Similarly, it is assumed that the intrinsic capacitive resistance of each cell membrane is equal, that is IXc$_1$=IXc$_2$=IXc$_3$=…=IXc$_p$=IXc$_0$. Thus, the IXc of the plant leaves was obtained:

\[
\frac{1}{\text{IXc}} = \frac{p}{\text{IXc}_0}
\]  

(19)

Due to membrane capacitive resistance is most closely related to surface proteins of cell membrane, then IXc or p can be characterized as the amount of surface proteins that induce membrane capacitive resistance in plant leaves.

Clearly, IXc is inversely proportional to p. The lower IXc, the more surface proteins.

The IXL of the plant leaves is calculated according to formula (20):

\[
\frac{1}{\text{IXL}} = \frac{1}{\text{IXL}_1} + \frac{1}{\text{IXL}_2} + \frac{1}{\text{IXL}_3} + \ldots + \frac{1}{\text{IXL}_q}
\]  

(20)

It is assumed that the membrane inside and outside inductive resistance of each cell is equal, then IXL$_1$, IXL$_2$, IXL$_3$, … IXL$_q$ can represent intrinsic inductive resistance of each unit cell membrane. Similarly, it is assumed that
the intrinsic inductive resistance of each cell membrane is equal, that is $I_{XL_1}=I_{XL_2}=I_{XL_3}=\ldots=I_{XL_q}=I_{XL_0}$. Thus, the $I_{XL}$ of the plant leaves was obtained:

$$\frac{1}{I_{XL}} = \frac{q}{I_{XL_0}}$$  \hspace{1cm} (21)$$

Due to membrane inductive resistance is most closely related to binding proteins of cell membrane, then $I_{XL}$ or $q$ can be characterized as the amount of binding proteins that induce membrane inductive resistance in plant leaves. Same, $I_{XL}$ is inversely proportional to $q$. The lower $I_{XL}$, the more binding proteins.

The cell membrane proteins are most closely related to the nutrient transport, thus, the nutrient transport capacity (NTC) could be represented by formula (22):

$$NTC = \frac{I_{Xc}+I_{XL}}{I_{IR}}$$  \hspace{1cm} (22)$$

**Electrophysiological information and nutrient transport** of *B. papyrifera* grow in two habitats

The fitting equation parameters of between clamping force and leaf $R$, $Xc$, and $XL$ of *B. papyrifera* grown in agricultural and moderately rocky desertification soils are shown in Table 1, Fig. 2 randomly lists the fitting curves for 1-4 leaf of *B. papyrifera* in agricultural soil. The correlation coefficients ($R^2$) of the fitting equations of $R$-$F$, $Xc$-$F$, and $XL$-$F$ for nine leaves of *B. papyrifera* grown in agricultural and moderately rocky desertification soils were 0.9044~0.9929, 0.9033~0.9910 and 0.9085~0.9895, and 0.9722~0.9976, 0.9910~0.9986 and 0.9862~0.9976, respectively. Moreover, all the $P$ values of the fitting equation parameters were lower than 0.0001. These results show that the relationships of between clamping force and leaf $R$, $Xc$, and $XL$ display good correlations, and highlight that the intrinsic mechanism relationships of those are authentic existence.

**Table 1**

**Fig. 2.**

The intrinsic electrophysiological information and the nutrient transport capacity of *B. papyrifera* in two conditions were successful monitored using the corresponding equation parameters. As shown in Table 2, the leaf IR,
IXc and IXL of *B. papyrifera* in the agricultural soil are significantly \((p < 0.01)\) lower than those of that in the moderately rocky desertification soil. Theoretically, the lower IXc and IXL, the more surface and binding proteins.

Actually, crude protein of *B. papyrifera* in the agricultural soil are significant \((p < 0.05)\) higher than those of that in the moderately rocky desertification soil, which is in good agreement with IXc and IXL. Moreover, for the same plant, the leaf IXc is lower than IXL which shows that binding proteins is more than surface proteins. As displayed in Table 2, the NTC and crude ash of *B. papyrifera* in the agricultural soil are significantly \((p < 0.01)\) higher than those of that in the moderately rocky desertification soil. The results showed that *B. papyrifera* in the agricultural soil grow well under the high nutrient (crude ash) conditions, and cell membrane proteins (crude protein) were relatively much which supported it higher NTC as compared to that in the moderately rocky desertification soil.

Table 2

**Electrophysiological information and nutrient transport** of the herbaceous and woody plants

As illustrated in Table 3, the IR, IXc and IXL of different plants are obviously different, the IXc is lower than IXL in same plant. For the same species plants in the same growth habitat, the NTC, crude protein and crude ash of *R. chinensis* are significantly \((p < 0.01)\) higher than those of *T. sinensis*, and those of *I. batatas* were significantly \((p < 0.05)\) higher than those of *S. scandens*. The results showed that the higher crude protein and crude ash in same species plants, the higher NTC.

Table 3

**Electrophysiological information and nutrient transport** of *S. tuberosum* and *C. annuum*

As shown in Table 4, the leaf IR, IXc and IXL of *S. tuberosum* are significantly \((p < 0.01)\) lower than those of *C. annuum* in the same growth habitat, while NTC, crude protein and crude ash are higher. And IXc is lower than IXL in same plant. The results showed that *S. tuberosum* with high membrane protein (crude protein) and nutrient (crude ash) contents promote the efficient transport and utilization of nutrients by its membrane proteins, which made it had higher nutrient transport capacity (NTC).


Almost all life activities in plants involve charge separation, electron movement, proton and dielectric transport, etc. In mesophyll cells, cells and organelles are both surrounded by the cell membrane composed of 50% lipids, 40% proteins and 2~10% sugars [1-2]. Membrane lipids and membrane proteins can be regarded as insulating layer, have a high electrical resistivity, enabling the plant cell to store electric charge [3]. Surface (or peripheral) proteins account for 20~30% of membrane proteins, bind to lipids on both sides of the membrane with charged amino acids or groups, and binding (or intrinsic) proteins account for 70~80% of membrane proteins, bind to lipids through hydrophobic hydroxyl groups in the membrane [1-2]. Surface proteins affect the capacitive reactance and capacitance, while binding proteins affect the inductive reactance and inductance. Therefore, the mesophyll cells can be regarded as a concentric sphere capacitor with both inductor and resistor function, and the ions, ion groups and electric dipoles are equivalent to electrolytes of capacitor [2, 14-15].

When plant leaves are subjected to clamping force stimuli (or environmental stresses), the cell membrane permeability of leaves changes instantly, and then the concentration of the ions, ion groups and electric dipoles inevitably changes, resulting in the changes of the leaf R, Xc and XL. Nernst equation can quantitatively describe the potential formed by ions between systems A and B, and it can theoretically also be used to quantitatively describe the diffusion potential of the electrolytes inside and outside of the cell membrane. Based on this fact, the R, Xc or XL of plant leaves were successfully obtained via the theoretically intrinsic relationships between clamping force and leaf R, Xc or XL were revealed for the first time. The results show that the relationships of between clamping force and leaf R, Xc, and XL displayed good correlations, and highlight that the aforementioned intrinsic mechanism are authentic existence. Generally, the intrinsic or spontaneous electrophysiological information in plants are not detectable [11]. In this study, the IR, IXc and IXL of plant leaves were successfully obtained via the theoretically intrinsic relationships between clamping force and leaf R, Xc and XL for the first time, which overcome the lack of representativeness, stability and reproducibility of the traditional needing approach.

Currently, the detection of membrane proteins is limited to single cell or single proteins, and the existing protein detection technology is hardly evaluate the composition characteristic of cell membrane protein [17-18]. The results
in this study showed that IXc and IXL could be used to manifest the composition of surface and binding proteins in cell membrane, that was, the lower IXc and IXL, the more surface and binding proteins. This is closely related to the fact that the high content of membrane proteins promoted the nutrient elements to pass through cell membrane more smoothly, thus made the cell membrane resistivity lower. In this study, plant with high crude proteins had relatively lower IR, IXc and IXL, which strongly supported the feasibility of using IXc and IXL to characterize the composition characteristic of membrane proteins. This study found that a phenomenon was common in the all tested plants, that was, the IXc was lower than IXL in same plant. This result perfectly proves the life fact that binding proteins is more than surface proteins in cell membrane [1-2].

Due to the poor nutritional environments, plants in rocky desertification soils are more vulnerable to low nutrient stress than those in cultivated soils [22-24]. The results showed that B. papyrifera in the agricultural soil grow well under the high nutrient (or crude ash) conditions, and cell membrane protein (or crude protein) content were higher which supported it higher nutrient transport capacity as compared to that in the moderately rocky desertification soil. The monitoring of the transport capacity of plant nutrients has rarely been reported in previous studies. In this study, the nutrient transport capacity (NTC) was defined based on IR, IXc and IXL for the first time. The results showed that the higher crude protein and crude ash in all tested plants, the higher NTC. The possible reason is that plant with high membrane protein (crude protein) and nutrient (crude ash) contents promoted the efficient transport and utilization of nutrients by its membrane proteins, which made it had well nutrient transport capacity. Overall, NTC commendably reflected the nutrient transport strategies in various tested plant, and could monitor the nutrient transport status of plants in real time. Additionally, the novel nutrient parameter were obtained by the intrinsic electrophysiological information in plants, which had well authenticity, stability, comparability and reproducibility. This study highlights that IR, IXc and IXL of plant electrophysiological information could effectively manifest the composition and nutrient transport characteristics of membrane protein in plant cells.

Conclusion

The present work provided a novel method based on plant electrophysiological information for accurately manifest the composition and nutrient transport characteristics of membrane protein in plant cells. The theoretically intrinsic relationships among the leaf R, Xc, XL and clamping force were first revealed on the basis of Nernst equation, and the IR, IXc and IXL of the intrinsic electrophysiological parameters in plant leaves were monitored via
these relationships for the first time and used to manifest the composition characteristic of cell membrane proteins. NTC was firstly defined based on IR, IXc and IXL which accurately revealed and reflected the nutrient transport strategies in tested plants.

Materials and methods

Experimental materials

The two Broussonetia papyrifera grown in the agricultural and moderate rocky desertification soil in Puding county, Guizhou Province (26°37′ N, 105°77′ E). Rhus chinensis Mill. and Toona sinensis grown in the moderate rocky desertification soil in Puding county, and Ipomoea batatas (L.) Lam. and Senecio scandens Buch.-Ham. ex D. grown in the cultivated soil in Puding county. Solanum tuberosum L. and Capsicum annuum L. were grown in the potted agricultural soil of Guizhou vocational college of agriculture in Qingzhen county, Guizhou Province (26°58′ N, 106°43′ E). The average annual temperature, sunshine hours and precipitation in Puding and Qingzhen counties were 15.1 and 14.1°C, 1164.9 and 1128.2 hours and 1378.2 and 1180.9 mm, respectively. The growth age, habitat information, measurement conditions and sampling weather of all tested plants are shown in Table 5. The fully expanded leaves of fresh branch as experimental materials were measured. First, the fully expanded leaves were taken from the third, fourth, and fifth leaf positions of each branch, and the fresh leaves were immediately soaked in water for 30 min. Then, the water on the surface of the leaves was removed. Three branches of each plant were measured. The tested leaves were sampled and measured at 8~10 a.m. on sunny days, and the measurement temperature was room temperature (25.0±2.0 °C).

Table 5

Leaf electrophysiological parameters and crude ash measurement

The fully expanded leaves from the third, fourth, and fifth leaf positions of three branches in plants were measured.
The leaf electrophysiological parameters were measured using a LCR-6300 tester (Gwinstek, Taiwan, China) with a frequency and voltage of 3 kHz and 1.5 V, respectively, as described by Zhang et al. [24]. Every mesophyll cell can be regarded as a concentric sphere capacitor, many aligned mesophyll cells make up the leaf capacitor, the parallel connection modes of LCR is thus applied. Firstly, the leaf was put between the two electrodes of a self-made parallel-plate capacitor with a diameter of 7 mm (Fig. 3). And then leaf capacitance (C), impedance (Z) and R at different clamping forces were continuously collected by adding the same quality iron blocks, and recorded 11-13 data each clamping force. Finally, leaf Xc and XL were respectively obtained according to formula (23) and (24):

\[ Xc = \frac{1}{2\pi fC} \quad (23) \]

\[ \frac{1}{-XL} = \frac{1}{Z} - \frac{1}{R} - \frac{1}{Xc} \quad (24) \]

where Xc= capacitive reactance, \( \pi = 3.1416 \), f= frequency, C= physiological capacitance, XL= inductive reactance, Z= impedance, R= resistance.

For crude ash measurement, the three tested leaves of each branch were rinsed with distilled water, dried in the shade, and then dried at low temperature, smashed and mixed. Crude protein and ash of samples were determined as described by Rayees et al. [25].

**Data analyses**

The data were analyzed using SPSS 18.0 (SPSS Inc., Chicago, IL, USA). A one-way analysis of variance followed by Duncan’s test was performed.

**List of abbreviations**

C: capacitance, Z: impedance, R: resistance, Xc: capacitive reactance, XL: inductive reactance, IR: intrinsic resistance, IXc: intrinsic capacitive reactance, IXL: intrinsic inductive reactance, NTC: nutrient transport capacity.
Supplementary information accompanies this paper at Additional file 1. Construction of the relationship models of clamping force (F) and leaf Xc, XL.

Additional file 2. Raw data

Ethics approval and consent to participate
Not applicable.

Consent for publication
Not applicable.

Availability of data and materials
The datasets generated and/or analysed during the current study are available in this published article and its supplementary information files.

Competing interests
The authors declare that they have no conflicts of interest.

Funding
We thank the National Natural Science Foundation of China (No. U1612441-2), the Key Technologies Research and Development Program of China (No. 2016YFC0502607-02, 2016YFC0502602-5), the Science and technology innovation talent project of Guizhou Province [No. (2015)4035], and the scientific and technological achievement
transformation project of Guizhou Province [No. (2017)4124] for supporting this research.

Authors’ contributions

YYW constructed conception. YYW and CZ designed research. CZ, YS and LF performed research. CZ and DX analyzed data. CZ and YYW wrote the paper. All authors read and approved the final manuscript.

Acknowledgements

We would like to thank the comprehensive experimental station of Karst Ecology in Puding, Guizhou Province, Chinese Academy of Sciences for providing the necessary support for this study, such as plant materials and corresponding premises, etc.

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Figure captions

Fig. 1. Simplified equivalent circuit of cells. $Z=\text{impedance}$, $C_m=\text{capacitance of membrane}$, $R_m=\text{resistance of membrane}$, $X_{cm}=\text{capacitive reactance of membrane}$, $X_{Lm}=\text{inductive reactance of membrane}$, $R_o=\text{resistance of membrane outside}$, $R_i=\text{resistance of membrane inside}$

Fig. 2. Fitting equations of the relationship between $R$ (a), $X_c$ (b), $X_L$ (c) of the fourth expanded leaf of the first branch of $B. papyrifera$ grown in the agricultural soils and champing force (F)

Fig. 3. The experimental setup used in the study and a schematic diagram of the parallel-plate capacitor. 1= holder (315 mm of height), 2= cystosepiment (32 mm of diameter), 3= plate electrode (7 mm of diameter), 4= electrical conductor, 5= iron block, 6= plastic rod (295 mm of height), 7= bench holdfast (130 mm of length).

Table captions

Table 1 The fitting equation parameters of $B. papyrifera$ in two habitats

Table 2 The nutrient transport parameters of $B. papyrifera$ in two habitats

Table 3 The nutrient transport parameters of four plants

Table 4 The nutrient transport parameters of $S. tuberosum$ and $C. annuum$

Table 5 Growth age, habitat information, measuring conditions and sampling weather of all tested plants