Root proteome alterations in sugarcane promoted by the regrowth cycle in commercial production

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Abstract

The decrease in agricultural productivity in successive cutting of sugarcane plants is associated with several extrinsic and intrinsic factors. However, no studies have focused on the physiological potential of sett roots in successive cuts in sugarcane culture. There have been no proteomic studies on sugarcane sett roots at different stages of cutting. In this study, the UPLC-ESI-TOF-MS system and bioinformatics tools were used to identify proteins of sett roots in the first and fifth cuts of sugarcane cultivar RB966928 in the sprouting stage. Differences in the proteome of sett roots of RB966928 in the first and fifth cuts detected in this study supports the hypothesis that the proteome of sett roots may change after successive cuts in sugarcane culture. A reduction in the number of proteins was observed in the roots of the fifth cut, whereas 34% of proteins, identified exclusively in the first cut, were absent in the fifth cut. Proteome analysis of sett roots in the first and fifth cuts showed that the changes after successive cuts were quantitative (number of proteins) and mainly qualitative. In this study, the detailed list of proteins identified in the first cut but absent in the fifth cut is relevant. The findings of this study may aid further research that employ biotic or abiotic elicitors to induce gene expression of essential proteins absent in sett roots of the fifth cut, and thus increasing the agricultural productivity and longevity of cane fields.

Keywords: Saccharum spp.; longevity; productivity; stress proteins; sett roots proteomic.

Abbreviations: 1D SDS-PAGE - one dimensional sodium dodecyl sulfate polyacrylamide gel electrophoresis; LC-MS/MS - liquid chromatography coupled with sequential mass spectrometry; PPI - protein–protein interaction network; UPLC-ESI-TOF-MS - ultra performance liquid chromatography-electrospray ionization-time of flight-mass spectrometer.

Introduction

Sugarcane (Saccharum spp.) is used for the production of alcohol and sugar worldwide. In addition, the pharmaceutical, cosmetic, and chemical industries are benefited from sugarcane byproducts for the manufacture of various materials and chemical components. Commercial sugarcane is a ratoon crop (Plucknett et al., 1970) depending on the sprouting of axillary buds present in seed-pieces (Ming et al., 2006). An economically viable sugarcane field depends on the high rates of axillary bud sprouting during harvest. Thus, the longevity of the cane field, approximately five cuts for most cultivars in Brazil (Santos and Borêm, 2013), is a determining factor for the success of the culture.

The sprouting of axillary buds in the stem after harvesting is accompanied by forming a root group called sett roots (Glover, 1967; Moore and Botha, 2014). The sett roots are formed in a band of root primordium above the leaf scar at the culm nodes (Van Dillewijn, 1952; Glover, 1967). The sett roots are used for anchoring and sustaining the new sprout, and for soil–plant interaction at the start of axillary bud sprouting (Smith et al., 2005). Sett roots remain for approximately 10 weeks (cultivar-dependent) after spraying, and thereafter replaced by shoot roots (Spaull, 1990). Budding is essential for generating vigorous plants for the next sugarcane production cycle (longevity).

The decrease in agricultural productivity in successive cuttings has been related to several extrinsic and intrinsic factors (Manhães et al., 2015). However, to date, no studies have assessed the physiological potential of sett roots in successive cuts in sugarcane culture. Changes in gene expression of the axillary buds at the fifth cut have been detected in sugarcane cultivars (Benez et al., 2019; Maranho et al., 2019) and may or may not be observed in sett roots after successive cuts. Proteome analysis revealed a reduction of proteins in the axillary buds of the fifth cut in the RB867515 sugarcane cultivar (Maranho et al., 2019). However, there have been no proteomic studies on sugarcane sett roots at different cut stages.

This study hypothesizes that the proteome of sett roots may change after successive cuts in sugarcane culture. This study elucidates the physiological role of sett roots in sugarcane and identifies genes involved in the synthesis of proteins essential for the stability and maintenance of average agricultural productivity in sugarcane fields. The UPLC-ESI-TOF-MS system and bioinformatics tools were used to identify proteins of sett...
roots in the first and fifth cuts of the sugarcane cultivar RB966928 in the sprouting stage.

Results

Root proteome of cultivar RB866928

The complex protein mixture by the TCA/acetone modified method quantified concentrations of 621 µg mL⁻¹ in the set roots of the first cut and concentrations of 924 µg mL⁻¹ in the fifth cut set roots. Pre-fractionation by 1D SDS-PAGE demonstrated good quality extraction (Figure 1), with protein bands ranging from approximately 120 kDa to 22 kDa.

The set root proteome was compounded with 141 proteins, of which 125 and 93 proteins were detected in the first and fifth cuts, respectively. Forty-eight proteins (34.04% of total proteome) were exclusively detected in the first cut (Table S1, markinga – Supplementary material), while only 16 proteins (11.34% of total proteome) were exclusively detected in the fifth cut (Table S2, markingb – Supplementary material). Seventy-seven proteins were identified at both the cutting ages.

Proteins identified in the sugarcane set roots of cultivar RB966928 integrate several diverse cellular processes and components. They are involved in carbon metabolism, lipid metabolism, protein metabolism, response to biotic and abiotic stresses, transcription factors, ribosomal components, and elongation factor, among others.

Differential proteome of first and fifth cut sett roots

The 48 proteins differentially identified in the set roots of first cut seed-pieces, participate in several cellular functions, acting as enzymes, structural proteins, or as proteins and/or polypeptides, regulating transcription and gene regulation.

A group of proteins related to the chaperone family was evident in the proteome of the set roots during the first cutting. Six heat shock proteins (heat shock cognate 70 kDa protein 2, heat shock cognate 70 kDa protein 1, heat shock cognate 70 kDa protein 3, heat shock cognate protein 80, heat shock protein 90-2, and heat shock protein 82), two chaperonins (chaperonin CPN60-2, mitochondrial precursor, 20 kDa chaperonin, chloroplast precursor), 2 luminal binding proteins (luminal binding protein 5 precursor, luminal binding protein 4 precursor), the chloroplast envelope membrane 70 kDa (chloroplast envelope membrane 70 kDa heat shock-related protein), and two Rubisco subunit binding-protein (Rubisco subunit binding-protein beta subunit, Rubisco subunit binding-protein alpha subunit, chloroplastic) are given in Table S1. Some of these proteins (Rubisco subunit binding-protein beta subunit, luminal binding protein 5, heat shock cognate 70 kDa protein 2, heat shock cognate protein 80, heat shock cognate 70 kDa protein 1, heat shock protein 90-2, heat shock cognate 70 kDa protein 3, chaperonin CPN60-2 mitochondrial precursor, and 20 kDa chaperonin chloroplast precursor) may be observed in Figure 2A (red marking), where the analysis of the PPI network revealed the presence of interactions between them, indicating a biological connection.

Proteins related to stress responses and root growth and development have also been identified only in the set roots of the first cutting, such as phosphoglucomutase, cytoplasmic 1 (EC 3.1.4.4), pyruvate dehydrogenase E1 component beta subunit, mitochondrial (EC 1.2.4.1), phospholipase D 1 precursor (EC 3.1.4.4), and 1-aminocyclopropane-1-carboxylate oxidase (EC 1.14.17.4). Tables 1 and 2 show the main KEGG pathways and UniProt keywords for proteins differentially identified in the set roots of the first cutting of seed pieces obtained by STRING. In the set roots of the fifth cutting, only the proteins 40S ribosomal protein S15, 40S ribosomal protein S4, 40S ribosomal protein S16, and 60S ribosomal protein L2, which were differentially identified, interacted with one another, as may be observed in Figure 2B. The four interacting proteins were ribosomal proteins that were exclusively identified in the roots of the fifth cut (Table S2). Four 14-3-3 family proteins (14-3-3-like protein, 14-3-3-like protein GF14 chi, 14-3-3-like protein 16R, and 14-3-3-like protein E) were also detected in the set roots of the fifth cut. However, the PPI network failed to reveal any evidence of interaction between them (Figure 2B). Table 3 shows the INTERPRO protein domains and features of the five proteins from the PPI network for the proteins differentially identified in the set roots of the fifth cutting.

Discussion

Morphological divergence was not observed in the set roots of RB966928 for the first and fifth cuttings. However, proteome analysis showed a reduction in the number of proteins (25.6% less proteins) in the set roots of the fifth cutting, perhaps related to reduced crop productivity. Essential proteins for photosynthesis, photosynthesis, glycolytic pathway, phytohormone biosynthesis, ribosomal subunits, transcription and elongation factors, protein and amino acid biosynthesis, defense against pathogens, in addition to different types of biotic and abiotic stress, identified in the set roots of the first cut, were different or absent in the roots of the fifth cut.

Proteins related to biotic and abiotic stresses and defense against pathogens commonly described in other plant species were identified in the set roots of the first and fifth cuts in sugarcane. The number of proteins related to biotic and abiotic stress identified in the set roots of the first cut was more than three times greater than the number of proteins related to biotic and abiotic stress identified in the set roots of the fifth cut. Furthermore, the number of proteins related to defense against pathogens identified in the set roots of the first cut was four times greater than the number of proteins produced for defense against pathogens identified in the set roots of the fifth cut. Proteins induced under thermal stress conditions were highlighted in the set roots of the first cut.

Heat shock protein families are necessary to protect cells from heat and various damages by normalizing cellular function during recovery processes (Morimoto et al., 1994; Parsell and Lindquist, 1994; Nové et al., 1996). Seven principal HSP families have been described, based on their molecular masses: Hsp110, Hsp100, Hsp90, Hsp80, Hsp70, Hsp40s, and Hsp (small Hsp) (Gupta et al., 2010; Park and Seo, 2015; Verna et al., 2019). Nine HSP or HSP precursors (Hsp70-2, Hsp70-1, Hsp70-3, Hsp60, Hsp80, Hsp80-B, Hsp90-2, Hsp90-7, and chloroplast envelope membrane 70 kDa heat shock-related protein) were detected in the set roots of the first cut in RB966928. Several studies have indicated that HSPs play an essential role in the survival of plants under thermal stress (Luján et al., 2009; Hu et al., 2010; Li et al., 2014) and greater tolerance to water and salt stress (Augustine et al., 2015).
The chloroplast envelope membrane 70 kDa heat shock-related protein was exclusively detected in the sett roots of the cultivar RB966928 of the first cut forms a complex of Hsp families that are induced under thermal stress conditions. The chaperonin CPN60-2 mitochondrial precursor (HSP60) was also detected exclusively in the sett roots of the first cut. Chaperonins prevent the misfolding of polypeptides and promote proper refolding and assembly of unfolded polypeptides generated under stress conditions in the mitochondrial matrix, which may be induced by a series of biotic and abiotic stresses, such as exposure to cold, ultraviolet light, water deficit, and tissue remodeling (Boston et al., 1996; Hartl et al., 2011). Under field conditions, sugarcane regrowth often occurs at elevated temperatures, causing the accumulation of folded and unfolded proteins. The higher amount of Hsps and SHsps in the sett roots of the first cut may indicate a better response under high thermal stress conditions. Furthermore, the absence of the four HSP70, HSP60, HPS80, and HPS82 in the sett roots of the fifth cut, may reduce the ability of the axillary buds of plants of the fifth cut to detect and respond to thermal and other stresses (abiotic and biotic), and soil-plant interaction, mediated by the sett roots, may alter parameters that reduce the ability of regrowth of the axillary buds.

Eleven proteins related to biotic and abiotic stress (20 kDa chaperonin/chloroplast precursor, BiP4, BiP5, vacuolar ATP synthase catalytic subunit A, isoflavone reductase homolog, Ras-related protein Rab7A, seed lipoygenase-2, glutathione reductase cystolic, flavonol synthase, aminopeptidase/chloroplast precursor, and 1-aminoacyclopropane-1-carboxylate oxidase) were identified in the sett roots of the first cut; however, they were absent in the sett roots of the fifth cut. The 20 kDa chaperonin, a chloroplast precursor (CPN20), shows to function as a co-chaperonin and CPN60 to promote proper refolding and assembly of unfolded polypeptides generated under stress. A review by Zhao and Liu (2018) highlighted the importance of the structure and function of the chaperonin-chloroplast system because chloroplasts have been suggested as sensors for various environmental stresses. Two BiP proteins (BiP4 and BiP5), detected in the sett roots of the sugarcane cultivar RB966928 of the first cut, are molecular chaperones of the endoplasmic reticulum that acts on protein folding and maturation (Carvalho et al., 2014). In the absence of BiP, many secretory pathway proteins do not assume their active conformation (Pobre et al., 2019). Several studies have demonstrated the role of BiP proteins in plant protection under various stress conditions (Alvim et al., 2001; Wang et al., 2005; Costa et al., 2018; Valente et al., 2009; Reis et al., 2011). The vacuolar ATP synthase catalytic subunit A (V-ATPases) proteins hydrolyze ATP to ADP and Pi coupled with H+ transport (Toei et al., 2010). The transcript levels of V-ATPases are regulated by environmental stresses such as salinity, heat, and drought responses that help plants maintain their water balance (Okamoto and Futai, 2018). The isoflavone reductase homolog (IFR; EC 1.3.1.1-4) catalyzes a stereospecific NADPH-dependent reduction to (3R)-isoflavonone (Wang et al., 2006), and a greater expression of IFR has been shown after induction by UV, wounding, pathogen infection, and cold (Lucheta et al., 2007), indicating the role of IFR in responding to stressful situations. The Ras-related protein Rab7A is localized mainly to late endosomes (Bucci et al., 2000; Vieira et al., 2003; Saito-Nakano et al., 2007). The Rab7 gene expression in rice is differentially regulated by environmental stimuli, such as cold, NaCl, dehydration, and abscisic acid (Nahm et al., 2003). Guerra and Bucci (2016) reported the main functions of the small GTPase Rab7, highlighting that the central role of Rab7 in correct cargo selection, biogenesis, positioning, motility, and functioning of lysosomes, phagolysosomes, and autolysosomes is fundamental for many cellular processes. The seed lipoygenase-2 (LOX; EC 1.13.11.12) catalyzes the addition of molecular oxygen to polyunsaturated fatty acids to produce unsaturated fatty acid hydroperoxide. A review by Porta and Rocha-Sosa (2002) reported recent advances in the role of LOXs in the physiology of plants. Glutathione reductase cytosolic (GR; EC 1.6.4.2) play an important role in protecting cells against oxidative damage (Trivedi et al., 2013). GR overexpression studies in various plants have revealed that it protects the cell from abiotic stress induced by oxidative damage. Flavonol synthase (FLS; EC 1.14.11.-) catalyzes the early step in the flavonoid biosynthesis pathway, leading to the production of flavonals and anthocyanins against oxidative damage (Jin et al., 2005). The 1-aminoacyclopropane-1-carboxylate oxidase (ACO; EC 1.14.17.4), detected only in the sett roots of the first cut, is related to biosynthesis of the ethylene phytohormone and in the plant’s defense against biotic and abiotic stresses (Van de Poel et al., 2015). The absence of ACO in the sett roots of the fifth cut suggests a loss of capacity by roots to respond to biotic and abiotic stresses, which may restrict the budding process of axillary buds at advanced cutting ages.

The protein cell division cycle protein 48 homolog, ubiquitin-activating enzyme 1, proteasome subunit beta type 1, gluconendo-1,3-beta-glucosidase GlI precursor, and trypsin inhibitor B, exclusively identified in the sett roots of the first cut, are reported to be essential against pathogens. The cell division cycle protein 48 homolog and ubiquitin-activating enzyme E1 in the sett roots of the first cut, although missing in the sett roots of the fifth cut, suggest that defensive responses to pathogens in the fifth cut may be deficient, causing plants to be more exposed to attacks by pathogenic microorganisms. The cell division cycle protein 48 homolog (product of Cdc48 gene) is an essential protein for plant cell cycle progression, and it is a component of protein quality control in plant immunity against pathogen infections (Chisholm et al., 2006; Jones and Dangl, 2006; Dangl et al., 2013; Bégue et al., 2018). The proteasome subunit beta type 1 (EC 3.4.25.1) is a core component of the 20S proteasome involved in several aspects of plant immunity (Ustán et al., 2016). The Glucon endo-1,3-β-glucosidase GlI precursor degrades fungal cell walls; therefore, it may inhibit pathogen growth by directly acting as an elicitor of defense reactions (Shetty et al., 2009). The application of purified β-1,3-glucan, isolated from the cell walls of the pathogen, has been shown to protect a susceptible wheat cultivar from disease development, accompanied by accumulation of pathogenesis-related proteins and callose. Trypsin inhibitor B is important in homeostasis, and changes in protease function leading to pathogenesis. It also maintains physiological homeostasis and helps the plant’s innate defense machinery (Arnaiz et al., 2018).

Among the higher number of proteins related to biotic and abiotic stresses and defense against pathogens in the sett roots of the first cut, only one protein of defense against pathogens (α-amylase inhibitors), and six proteins related to biotic and abiotic stresses different from those detected in sett roots of the first cut (spermidine synthases, 265 protease regulatory subunit 6A homolog, nitrilase 4, glycine-rich RNA-
binding proteins, profilin 6, and α-amylase inhibitor) were identified in the sett roots of the fifth cut from R8966928. Proteins related to biotic and abiotic stresses and defense against pathogens originally synthesized in the sett roots of the first cut were absent at the fifth cut. In contrast, other biotic and abiotic stresses were evident at advanced cutting ages.

Spermidine syntheses (SPDS; EC 2.5.1.16) are involved in normal cellular metabolism and stress response signaling regulators in stress-signaling pathways, leading to stress tolerance mechanisms in Arabidopsis thaliana plants under stress conditions (Kasukabe et al., 2004; Sekula and Dauter, 2019). Polyamines have been implicated in a wide range of biological processes in plant growth and development, including senescence, environmental stress, and infection by fungi and viruses (Kaur-Sawhney et al., 2003; Pál et al., 2018).

The 26S protease regulatory subunit 6A homolog is involved in the ATP-dependent degradation of ubiquitinated proteins in the protein catabolic process (Marino et al., 2012; Dudler, 2013). The 26S protease regulatory subunit 6A in the roots of the fifth cut may be another indication for response to stress situations due to infection by pathogens detected at this cut-off age, where defective or exogenous proteins from pathogens may be degraded by proteosomes.

Nitrilase 4 (NIT; EC 3.5.5.1) is an isozyme of the nitrilase superfamily that catalyzes the hydrolysis of nitrile compounds to carboxylic acids and ammonia. Nitrilases are tissue-specific (reviewed by O’Reilly and Turner, 2003; Howden and Preston, 2009). NIT4 activity (detected in the sett roots of the fifth cut in this study) is described as higher in senescent leaves of A. thaliana than in non-senescent ones (Piotrowski et al., 2001). The authors suggested that the upregulation of NIT4 activity during senescence may be linked to increased ethylene biosynthesis and cyanide production.

α-Amylase inhibitors (α-AIs) are part of the natural defense mechanisms against pests by interfering with their digestion process. They may also provide access to new pest management strategies (Kaur et al., 2014). Several important roles of glycine-rich RNA-binding proteins (GR-RBPs) in response to abiotic stresses in actively proliferating organs, such as young plants, root tips, and flowers, were reviewed by Wang et al. (2018). Profilin 6 represents a group of proteins (profilins) that are key regulators of actin cytoskeleton dynamics through their interaction with monomeric actin (G-actin). Interaction of profilins with phosphatidyl inositol-4,5-bisphosphate, a major component of cell-signaling transduction pathways, is essential for integrating stress responses through cytoskeleton rearrangement, in addition to processes such as cell movement and cytokinesis driven by actin polymerization dynamics (Wilke and Otto, 2003).

Quantitative differences were observed in the expression of genes involved in the synthesis of stress proteins identified in the sett roots of the first and fifth cuts of R8966928. However, the most striking difference in the proteome of the sett root in the first and fifth cuts was related to the essential proteins for photosynthesis, photorespiration, glycolytic pathway, control of gene expression, and protein and amino acid biosynthesis to plant growth and development exclusively observed in the sett roots of the first cut. The triosephosphate isomerase, ribulose 1,5-bisphosphate carboxylase/oxygenase (RuBisCO) subunit binding protein α-subunit, RuBisCO subunit binding protein β-subunit and transketolase 10 are essential proteins for photosynthetic processes detected in the sett roots of the first cut although absent in the sett roots of the fifth cut. Triosephosphate isomerase (TI; EC 5.3.1.1) is present in several metabolic pathways involved in glycolysis, gluconeogenesis, and the Calvin cycle. Chen and Thelen (2010) showed that a reduction in gene expression and plastid TPI activity (pdTPI) resulted in a stunted plant that failed to reach reproductive maturity. The RuBisCO (EC 4.1.1.39) subunit binding protein α-subunit and β-subunit are recognized as chaperonin 60, which is intimately involved in the assembly of the RuBisCo enzyme (Zhao and Liu, 2018). Transketolase (TKT; EC 2.2.1.1) is integral to the Calvin cycle and the oxidative pentose phosphate pathway of higher-plant chloroplasts (Khozaee et al., 2015).

The aminomethyltransferase mitochondrial precursor, glycine dehydrogenase mitochondrial, serine hydroxymethyltransferase mitochondrial, malate synthase glyoxysomal, and aconitate hydratase cytoplasmic are essential proteins for photorespiration processes detected in the sett roots of the first cut; however, they are absent in the sett roots of the fifth cut. The aminomethyltransferase mitochondrial precursor (AMT; EC 2.1.2.10) protein is one of the four enzymes that work in the glycine cleavage system, producing methyl groups transferred to other key molecules, such as purines and methionine. The glycine cleavage system is required for photorespiration in C3 plants (Timm et al., 2015). Glycine dehydrogenase B, mitochondrial (GDC; EC 1.4.4.2), is also an integral part of the photorespiratory system involved in the glycine catabolic process (Palmieri et al., 2010). Serine hydroxymethyltransferase mitochondrial (SHMT; EC 2.1.2.1) functions in the photorespiratory pathway by catalyzing serine and glycine interconversion. Glycine and serine are intermediates in the photorespiration of glycerate to 3-phosphoglycerate. Moreno et al. (2005) observed that, in addition to acting as an essential protein for photorespiration, SHMT is involved in controlling cell damage caused by abiotic stress, such as high light, salt rates, and also in the defense response of plants. The malate synthase glyoxysomal (MS; EC 2.3.3.9) enzyme is specific to the glyoxylate cycle responsible for malate synthesis from acetyl-CoA and glycolate in the glyoxysome. In addition to gluconeogenesis, the glyoxylate cycle is also important for replenishing succinate in the TCA cycle, the major cellular machinery for oxidative metabolism and energy production (Pua et al., 2003). The cytoplasmic aconitate hydratase (cACO; EC 4.2.1.3) is involved in the reversible conversion of citrate to isocitrate, crucial to the Krebs cycle, which plays a key role in aerobic respiration. Carrari et al. (2003) have shown that a reduction in aconitase activity causes a stunted phenotype in the early stages of tomato (Lycopersicon pennellii) development plants.

Phosphoglucomutase and fructokinases are essential proteins in the glycolytic pathway detected in the sett roots of the first cut and absent in the sett roots of the fifth cut. Essential proteins for glycolytic via and respiration, such as the phosphoglucomutase (cPGM; EC 5.4.2.2) isozymes identified in the sett roots of the first cut and absent in the roots of the fifth cut, catalyze the reversible interconversion of glucose 6-phosphate and glucose 1-phosphate. The absence of cPGM in the sett roots of the fifth cut may be related to metabolic changes in the radicular cells and probable deficit in the conduction of water and nutrients to the axillary bud, which results in a lower sprouting rate, less
Table 1. KEGG Pathways for proteins differently identified in the sett roots of the first cutting of sugarcane seed-pieces obtained by STRING 11.0 database.

| KEGG Pathways | Description                                     | Count in gene set | False discovery rate |
|---------------|-------------------------------------------------|-------------------|----------------------|
| cmo01110      | Biosynthesis of secondary metabolites           | 15 of 958         | 1.05e-08             |
| cmo01200      | Carbon metabolism                               | 9 of 229          | 1.79e-08             |
| cmo01100      | Metabolic pathways                              | 18 of 1685        | 1.79e-08             |
| cmo01230      | Biosynthesis of amino acids                     | 8 of 201          | 7.84e-08             |
| cmo00630      | Glyoxylate and dicarboxylic metabolism          | 6 of 69           | 7.84e-08             |
| cmo04141      | Protein processing in endoplasmic reticulum     | 6 of 168          | 8.61e-06             |
| cmo00710      | Carbon fixation in photosynthetic organisms     | 4 of 64           | 6.23e-05             |
| cmo00260      | Glycine, serine and threonine metabolism        | 4 of 72           | 8.49e-05             |
| cmo01210      | 2-Oxocarboxylic acid metabolism                 | 3 of 44           | 0.00054              |
| cmo00670      | One carbon pool by folate                       | 2 of 18           | 0.0031               |
| cmo00010      | Glycolysis / Gluconeogenesis                     | 3 of 111          | 0.0059               |
| cmo00500      | Starch and sucrose metabolism                   | 3 of 116          | 0.0061               |
| cmo04144      | Endocytosis                                      | 3 of 145          | 0.0105               |
| cmo00020      | Citrate cycle (TCA cycle)                       | 2 of 44           | 0.0116               |
| cmo00030      | Pentose phosphate pathway                       | 2 of 47           | 0.0122               |
| cmo00051      | Fructose and mannose metabolism                 | 2 of 51           | 0.0134               |
| cmo00270      | Cysteine and methionine metabolism              | 2 of 84           | 0.0320               |
| cmo03018      | RNA degradation                                 | 2 of 94           | 0.0372               |

Figure 1. 1D SDS-PAGE with the protein profile from sett roots of the sugarcane cultivar RB966928. A - Roots of first cut; B - Roots of fifth cut; L - Ladder. Red arrows indicate differential bands for the cut ages.

Table 2. UniProt Keywords for proteins differently identified in the sett roots of the first cutting of sugarcane seed-pieces obtained by STRING 11.0 database.

| UniProt Keywords | Description                 | Count in gene set | False discovery rate |
|------------------|----------------------------|-------------------|----------------------|
| KW-0346          | Stress response             | 5 of 52           | 6.39e-06             |
| KW-0809          | Transit peptide             | 3 of 26           | 0.00086              |
| KW-0554          | One-carbon metabolism       | 2 of 14           | 0.0092               |
| KW-0496          | Mitochondrion               | 3 of 70           | 0.0092               |
| KW-0489          | Methyltransferase           | 4 of 254          | 0.0189               |
| KW-0067          | ATP-binding                 | 7 of 851          | 0.0189               |
| KW-0032          | Aminotransferase            | 2 of 38           | 0.0255               |
| KW-0460          | Magnesium                   | 3 of 161          | 0.0314               |
| KW-0663          | Pyridoxal phosphate         | 2 of 55           | 0.0381               |
| KW-0560          | Oxidoreductase              | 5 of 607          | 0.0433               |
Figure 2. Protein-protein analysis by STRING 11.0. (A) STRING PPI network of some differential proteins identified in sett roots of first cut sugarcane. Network contains 95 edges (vs. 34 expected edges); enrichment p-value < 1.0e-16. (B) STRING PPI network of some differential proteins identified in sett roots of fifth cut sugarcane. PPI network contains 15 nodes with 6 edges (vs. 4 expected edges); enrichment p-value < 0.199. Minimum required interaction score was set at 0.400 (medium confidence) for both analyses.

Table 3. INTERPRO Protein Domains and Features for proteins identified in the sett roots of sugarcane obtained by STRING 11.0 database.

| INTERPRO Protein Domains and Features |
|--------------------------------------|
| Domain | Description | Count in gene set | False discovery rate |
|-------|-------------|-------------------|---------------------|
| IPR036815 | 14-3-3 domain superfamily | 4 of 25 | 2.42e-08 |
| IPR023410 | 14-3-3 domain | 4 of 25 | 2.42e-08 |
| IPR000308 | 14-3-3 protein | 4 of 25 | 2.42e-08 |
| IPR023409 | 14-3-3 protein, conserved site | 3 of 22 | 1.77e-06 |
| IPR014722 | Ribosomal protein L2, domain 2 | 2 of 36 | 0.0012 |

Figure 3. Seed-pieces with individualized axillary buds and sett roots developed from the node’s root primordia. A – Seed-piece of first cut; B - Seed-piece of fifth cut.
robust plants, and decreased agricultural productivity of the sugarcane plantation (Malinova et al., 2014). Fructokinases (FRKs; EC 2.7.1.4) are important enzymes that catalyze the key metabolic step of fructose phosphorylation (Riggs et al., 2017) to enter metabolism and plant development (review by Stein and Granot, 2018).

5-adenosylmethionine synthetase 1 (SAM; EC 2.5.1.6) is an essential protein that controls gene expression detected in the sett roots of the first cut and absent in the sett roots of the fifth cut. Gong et al. (2014) showed that overexpression of the SAM gene in tomato (Lycopersicon esculentum) provided a stronger root system, exhibited a significant increase in tolerance to alkali stress and maintained nutrient balance, higher photosynthetic capacity, and lower oxidative stress.

Proteins involved in ribosomal subunit formation (60S ribosomal protein L12 and 60S ribosomal protein L7), elongation factors (elongation factor 2) for protein biosynthesis, and proteins involved in amino acid biosynthesis (5-methyltetrahydropteroyltriglutamate-homocysteine methyltransferase, ketol-acid reductoisomerase chloroplastic precursor, aspartate aminotransferase cytoplasmic isoform 1, and alanine aminotransferase 2) were exclusively identified in the sett roots of the first cut. Elongation factor 2 (eEF-2) is originally a GTP-binding protein involved in the translation process for polypeptide chains and protein synthesis, and eEF-2 is also associated with plant cold (Guo et al., 2002; Shi et al., 2019).

The enzyme 5-methyltetrahydropteroyltriglutamate-homocysteine methyltransferase (MET6; EC 2.1.1.14) catalyzes the reaction of the last step in L-methionine biosynthesis (Eichel et al., 1995). The reaction catalyzed by Ketol-acid reductoisomerase (KARI; EC 1.1.1.86) consists of the biosynthesis of branched-chain amino acids (Lee et al., 2005). Aspartate aminotransferase cytoplasmic isoform 1 (AAT2; EC 2.6.1.1) catalyzes the reversible reaction of transamination between aspartate and 2-oxoglutarate to generate glutamate and oxaloacetate, regulating the synthesis of aspartate or catabolism. Alanine aminotransferase 2 (AlaAT; EC 2.6.1.2) catalyzes the reversible conversion of pyruvate and glutamate into alanine and α-oxoglutarate, regulating the synthesis of aspartate or catabolism. Alanine aminotransferase 2 (AlaAT; EC 2.6.1.2) catalyzes the reversible conversion of pyruvate and glutamate into alanine and α-oxoglutarate. Studies have shown that AlaAT transcript levels could be induced by hypoxic conditions (Ricout et al., 2006; Miyashita et al., 2007) and also by light and N (Xu et al., 2017), regulating shoot root allocation.

Although different proteins involved in phytohormone biosynthesis, such as phospholipase D1 precursor, 1-aminocyclopropane-1-carboxylate oxidase, and pyruvate dehydrogenase E1 component β-subunit, were identified in the sett roots of the first cut, only the 14-3-3-like protein family was detected in the sett roots of the fifth cut. The Phospholipase D1 precursor (PLD 1; EC 3.1.4.4), exclusively detected in the sett roots of the first cut, cleaves the phosphodiester bond in the phospholipid, yielding phosphatidic acid (PA) and a free head group (Wang, 2000; Chen et al., 2011). Phospholipases may also bind and modulate other membrane regulatory proteins, including the subunits of G-protein complexes and sphingosine kinase (Pandey, 2016). Therefore, the maintenance of homeostasis and membrane stability is partly ensured by phospholipases (Lanteri et al., 2008; Takáč et al., 2019). According to Takáč et al. (2019), the various mechanisms of action of phospholipases have suggested that these enzymes are involved in plant responses to drought, cold, heat, salt, osmotic pressure, heavy metal stress, phosphorus deficiency, and pathogens. The absence of PLD 1 in the sett roots of the fifth cut may imply changes in the various mechanisms of action described for phospholipases and, consequently, reduced growth and development of axillary buds.

The 1-aminocyclopropane-1-carboxylate oxidase (ACO; EC 1.14.17.4) is related to the biosynthesis of ethylene phytohormones. It is an important regulator of many developmental and physiological processes, such as seed dormancy, germination, vegetative growth, flowering, climacteric fruit ripening, and senescence (Houben and Van de Poel, 2019). The pyruvate dehydrogenase E1 component β-subunit (PDH E1-β; EC 1.2.4.1) is important for regulating the distribution of PIN proteins in the cytoplasmic membrane of cells. PIN proteins are integral membrane proteins involved in transporting across membranes (Guan et al., 2019; Ohbayashi et al., 2019). Low expression of the gene for PDH E1B inhibited the transport of auxin, which is crucial for developing roots, and other organs mediated by auxin. Subsequently, the non-detection of PDG E1B in the roots of the fifth cut may be related to defective transport of auxins, with a consequent reduction in enzymatic activities, respiration, and amino acid metabolism of the sett roots.

The 14-3-3-like protein family represents 25% of the proteins exclusively detected in the sett roots of the fifth cut from RB966928 (14-3-3-like protein 14, 3-3-like protein GF14 chi, 14-3-3-like protein 16R, and 14-3-3-like protein E). Camoni et al. (2018) highlighted the role of 14-3-3 proteins in regulating hormonal signaling, biosynthesis, and transport. The 14-3-3-like proteins were also detected exclusively in the axillary buds of the fifth cut of sugarcane cultivar RR867515 (Maranho et al., 2019). It has been reported that 14-3-3-like proteins in A. thaliana may function as positive regulators of the phytohormone brassinosteroids, while 14-3-3 isoforms have different regulatory properties for brassinosteroid signaling (Lee et al., 2020). In contrast, in A. thaliana transgenic plants overexpressing the TAI4-3-3 gene (from Triticum aestivum), 14-3-3-like proteins are growth inhibitors that provide plants with delayed growth rates, shorter primary roots, and delayed flowering (Jing et al., 2013). The 14-3-3-like proteins were exclusively found in the sett roots of the fifth cut from RB966928 may indicate an incidence of stress that activates genes for 14-3-3-like proteins that act on signaling routes in the sett root cells. The large expression of genes for this group of proteins may inhibit the complete development of the sett roots of the fifth-cut specimens. The 14-3-3-like proteins, exclusively in the sett roots of the fifth cut, and the absence or reduction of other proteins not detected at this cut-off age may cause deficient soil-plant communication, interfering with the sprouting of axillary buds.

Ribosomal proteins (40S and 60S) represent another 25% of the proteins exclusively detected in the sett roots of the fifth cutting from RB966928. The 40S and 60S ribosomal proteins are essential for the organization of the ribosomal small subunit assembly and ribosomal larger subunit assembly, respectively, during ribosome biogenesis (Carroll, 2013; Sáez-Vázquez and Delseny, 2019). Several studies have shown that plant-pathogen interactions in the early stages cause transcriptional reprogramming, coupled to translational regulation. In current study, the authors suspect that the expressionive proportion of ribosomal proteins in the sett roots of the fifth cut from RB966928 may be related to plant responses to biotic stress.

Differences in the proteome of the sett roots from RB966928 in the first and fifth cuts detected in current study supports
the hypothesis that the proteome of sett roots may change after successive cuts in sugarcane culture. A reduction in the number of proteins (25.6%) was observed in the roots of the fifth cut; however, 34% of the proteins identified exclusively in the first cut were absent in the fifth cut. Proteome analysis of sett roots in the first and fifth cuts showed that the changes after successive cuts were mainly qualitative. Essential proteins for the promotion of plant growth and development involved with photosynthesis, photorespiration, glycolytic pathway, control of gene expression, and protein and amino acid biosynthesis were not detected in the roots of the fifth cut. In addition, while stress proteins exclusive to the first cut roots are involved in protecting and guaranteeing the plant development processes, stress proteins exclusively identified in the fifth cut may act as growth inhibitors, providing plants with delayed growth. The absence of essential proteins to promote plant growth and development and stress proteins involved in protecting and guaranteeing the development processes in sett roots of the fifth cut from RB966928 may reduce budding of the axillary buds. The reduction in agricultural productivity of RB966928 in the fifth cut was approximately 51% (information by farmers). The reduction in the budding of axillary buds may generate less vigorous plants for the next sugarcane productive cycle and compromise the longevity of cane fields. The development of strategies to induce the synthesis of proteins involved in photosynthesis, photorespiration, glycolysis, control of gene expression, protein bioactivity, and amino acids (undetected in the roots of the fifth cut) may be an alternative to increase the longevity of the cane fields of the cultivar RB9662869. The detailed list of proteins identified in the first cut but absent in the fifth cut reported in the present study is relevant because it opens a perspective for further researches that employ biotic or abiotic elicitors to induce gene expression. Application of biotic or abiotic elicitors may be a promising strategy to induce gene expression of the essential proteins absent in the sett roots of the fifth cut, increasing the agricultural productivity and longevity of cane fields.

Materials and methods

Plant material – sett roots of cultivar RB966828

Sugarcane stalks of the 10-month-old cultivar RB966928 in the first and fifth cuts were collected from farms of Nova Aralo Industrial (20°53'29.82"S, 50°26'55.25"W) in the state of São Paulo, Brazil. The sugarcane plants were cultivated in dystrophic Argisol-type soil with low water availability and medium cation exchange capacity. Based on leaf count by the Kuijper system, described by Van Dillewijn (1952), only auxiliary buds from the fourth to the ninth nodes were used to avoid the greater influence of auxins in the axillary buds near the stem apex. Auxiliary buds from plant cane (first cut) and plants of the fifth cut (fourth ratoon) were individualized and planted in vermiculite to initiate sprouting and sett root development, in labeled 10-L trays, with a spacing of 3 cm between samples. They were irrigated every two days. Sprouting occurred in a greenhouse at 22°C after 5 days. The sett roots of each cutting age (first and fifth; Figure 3) were excised with a scalpel, instantly frozen in liquid nitrogen, and stored in an ultra-freezer at −80°C until use. At each cutting age (first and fifth cut), the sett roots were taken from three plants (three plants from the first cut and three plants from the fifth cut; biological triplicate) and divided into three aliquots (technical triplicates).

Extraction, quantification, and protein digestion

Sett roots (200 mg) extracted from the first and fifth cutting stages were used in triplicate from each biological replicate. Total proteins were extracted using the modified TCA-acetone method for sugarcane axillary buds (Maranho et al., 2018). Proteins were quantified by fluorimetry (Qubit Fluorometer 1.0; Invitrogen, Carlsbad, CA, USA) with a Qubit Protein Assay Kit (Invitrogen, Carlsbad, CA, USA). The complex mixture of proteins was pre-fractionated by 1D SDS-PAGE (sodium dodecyl sulfate polyacrylamide gel electrophoresis), with 12% and 5% visualization and stacking gel, respectively. Electrophoresis lasted ~3 h at 200 V. Gel was then stained with 0.1% Coomassie Brilliant Blue R-250. The SDS-PAGE verified previously differences (Figure 1, red arrows) in protein expression in each sample and assessed protein extraction quality.

Tryptic digestion was performed following the method described by Villén and Gygi (2008). Disulfide bonds were reduced by incubation in dithiothreitol (DTT) solution (DTT 5 mM in 50 mM NH₄HCO₃ - ammonium bicarbonate), Cysteine residues were alkylated by incubation in an alkylation solution (14 mM iodoacetamide and 50 mM NH₄HCO₃). Proteins were digested with Trypsin/Lys-C Mix (Promega, Madison WI, USA) at a final concentration of 20 ng µL⁻¹ for 16 h at 37°C.

UPLC separation, MS analysis, and Bioinformatics

The protein digest was analyzed with an ultra-high performance liquid chromatograph ACQUITY UPLC M-Class System (Waters™, Milford MA USA), coupled to a time-of-flight high-resolution mass spectrometer (Xevo G2-XS TOF, Waters™) equipped with an electrospray ionization source. Chromatographic separation was performed using an Acquity UPLC® M-Class HSS T3 column packed with 1.8 µm particle size, 300 µm x 150 mm column (Waters™, UK), at a flow rate of 6 µL min⁻¹. The gradient mixture of solvents A (H₂O with 0.1% formic acid; v/v) and B (acetonitrile with 0.1% formic acid, v/v) was as follows: 3% B 0–1 min, 40% B 1–80 min, 97% B 80–90 min, maintained at 97% B 90–97 min, 3% B 97–100 min, and maintained at 3% B 100–103 min at 40°C. Capillary voltage was operated in positive mode, set at a 3.0 capillary, sampling cone 40 V, and desolvation gas 600 L h⁻¹/400°C. Data were collected from m/z 50 to 2000 using MSe acquisition, scan time = 0.5, and ramp collision energy 15–45 V.

After LC-MS/MS, raw data files were processed and analyzed with ProteinLynx Global ServerTM 3.0.3, and the sequences were searched against the Viridiplantae and Saccharum taxonomy in the UniProtKB database (downloaded in April 2019). The following parameters were used for database searches: cleavage specificity, trypsin with one missed cleavage allowed, Min Fragment Ion Matches per Peptide = 2, Min Fragment Ion Matches per Protein = 5, Min Peptide Matches Per Protein = 1, Fixed Modifier Reagent: Carbamidomethyl C; Variable Modifier Reagents: Oxidation M; FDR: 5%.

KEEG Pathways, Uniprot Keywords, and INTERPRO protein domains and features enrichment analysis were performed using the multiple protein search interface of the STRING database (version 11.0) (Szklarczyk et al., 2019) to evaluate known and predicted protein relationships. Differential proteins in the sett roots of the sugarcane plant and of the fourth ratoon were compared against Cucumis melo and Musa acuminata protein databases, respectively, for protein–
protein interaction network (PPI) analysis. *Cucumis melo* and *Musa acuminata* databases were selected due to more annotations of their biological processes and molecular functions in the STRING Consortium database and the lack of a *Saccharum* protein database.

All MS data were available via the jPOST Repository Database and ProteomeXchange with identifiers JPST001038 and PXDO23098, respectively.

**Conclusion**

The absence of essential proteins to promote plant growth and development (proteins involved in photosynthesis, photorespiration, glycolytic pathway, control of gene expression, protein and amino acid biosynthesis), and stress proteins involved in protecting and guaranteeing the development processes in sett roots of the fifth cut may reduce the budding of the axillary buds. The reduction in the budding of axillary buds may generate less vigorous plants for the next sugarcane productive cycle and compromise the longevity of cane fields. The absence of essential proteins in sett roots compromises the main function of these roots to provide water and nutrients to the young developing shoots.

**Supplementary Material for Publication**

Link MS data deposited in jPOST repository.
https://repository.jpostdb.org/ Acession: JPST001038.
Link MS data deposited in ProteomeXchange.
http://www.proteomexchange.org/ Acession: PXDO23098.

**Conflict of Interest**

The authors declare no conflict of interest.

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