Differential Expression of Genes Associated With Degradation Enhancement of Imazethapyr in Barnyardgrass

*(Echinochloa crus-galli)*

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Abstract

The understanding of mechanism of herbicide resistance in weeds is essential for adequate or innovative weed management practices. The aim of this study was to identify and analyze the expression of genes related to degradation enhancement of imazethapyr in barnyardgrass (*Echinochloa crus-galli* L. Beauv.). One susceptible (SUSSP01) and two populations previously identified as resistant to imazethapyr (ARRGR01 and PALMS01) were used. Gene expression of *CYP* and *GST*, the translation initiating factor *eIF4B*, and *ALS* genes were evaluated after imazethapyr spraying. A reference gene stability analysis was carried out, wherein the genes *18S* and *actin* showed to be more stable in response to the population and herbicide treatment. The gene expression analysis was performed by qRT-PCR. There was no difference in the relative expression of the *ALS* gene. The *CYP81A6* and *GSTF1* genes showed higher relative expression in the resistant populations. The *CYP81A6* gene had expression 9.61 and 8.44 higher in the resistant populations ARRGR01 and PALMS01, respectively, in comparison with the untreated susceptible population. The expression of this gene was induced by spraying the herbicide imazethapyr. The *GSTF1* gene showed higher relative expression in PALMS01 population, reaching 12.30 times higher in plants treated with imazethapyr in relation to untreated susceptible population. The expression of *eIF4B* gene in the resistant populations treated with imazethapyr was about six times higher than observed in susceptible population. The high relative expression of *CYP81A6* and *GSTF1* genes indicate the importance of degradation enhancement for the resistance of barnyardgrass to imazethapyr.

**Keywords:** imidazolinones, ALS, CYP, GST, herbicide resistance

1. Introduction

The resistance of weeds to herbicides is related to the occurrence of two primary mechanisms of resistance grouped as target site resistance (TSR) and non target site resistance (NTSR) (Powles & Yu, 2010). The TSR is associated mainly with mutation in the target enzyme encoding gene and increase activity of the target enzyme. NTSR mechanisms encompass the resistance caused by lower uptake and variation of herbicide translocation, rapid necrosis caused by reactive oxygen species, herbicide sequestration in the vacuole, and degradation enhancement of the herbicide (Sammons & Gaines, 2014; Yu & Powles, 2014; Nandula et al., 2015).

The degradation enhancement of herbicides by detoxifying enzymes has been associated with the resistance in several weed species, especially the *Lolium rigidum* Gaud. (Yu, Abdallah, Han, Owen, & Powles, 2009; Busi, Vila-Aiub, & Powles, 2011), *Alopecurus myosuroides* Huds. (Délye, Gardin, Boucansaud, Chauvel, & Petit, 2011) and *Echinochloa phyllopogon* Stapf. (Yasour et al., 2009; Iwakami et al., 2014a). The main enzymes related with herbicide detoxification include cytochrome P450 monooxygenases (known as CYP or P450) and glutathione S-transferases (GST) (Powles & Yu, 2010). The enzymes P450 and GST are fundamental in phases I and II of herbicide metabolism, respectively (Yuan et al., 2007). The resistance due to increased metabolization is particularly alerting, since the same enzyme can detoxify more than one herbicide, leading to cross- and multiple resistance (Beckie & Tardif, 2012).
Barnyardgrass (*Echinochloa crus-galli* L. Beauv.) is a major weed of irrigated rice fields around the world (Chauhan & Johnson, 2011). This weed can cause 21-79% yield rice losses, depending on weed density, rice cultivar and irrigation management (Bajwa et al., 2015), and demands intensive use of different methods of control. Herbicides are the most used method for controlling this weed, and acetolactate synthase (ALS) inhibitors, such as imidazolinones, are widely used. The availability of imidazolinone-resistant rice cultivars (Clearfield® rice) have allowed the repetitive use of these herbicides, selecting resistant barnyardgrass populations. In southern Brazil, the mechanism of resistance of these populations involves both types: mutations in the *ALS* encoding gene and increased metabolism by detoxifying enzymes (Matzenbacher, Bortoly, Kalsing, & Merotto Jr., 2015). In this study the effect of degradation enhancement as the mechanism of resistance was inferred through P450 inhibitors. However, the genetic regulation of this process is unknown in these populations.

The involvement of *CYP* and *GST* genes has been associated with herbicide resistance in weeds and selectivity in crops. The *CYP81A* is one of the most important gene subfamilies associated with the metabolism of herbicides in plants. The *CYP81A6* gene has already been related to the metabolism of bensulfuron-methyl and sulfonylureas in rice (Pan et al., 2006; Liu et al., 2012). Similarly, a *CYP81* gene is involved in the resistance of *A. japonicus* to the herbicide fenoxaprop-P-ethyl (Chen, Xu, Zhang, Bai, & Dong, 2018). In *E. phyllopoogon*, the higher expressions of the *CYP81A12* and *CYP81A21* genes are associated with resistance to the *ALS*-inhibiting herbicides penoxsulam and bensulfuron-methyl (Iwakami et al., 2014a). In this species, previous spraying of bispyribac-sodium, another ALS inhibitor, induced the expression of *CYP71AK2* and *CYP72A234* genes in plants resistant to these herbicides (Iwakami et al., 2014b). In rice and *Arabidopsis thaliana*, the *CYP72A31* gene confers tolerance to bispyribac-sodium (Saika et al., 2014). *GST* genes encode enzymes that catalyze conjugation reactions of herbicides with more soluble molecules, decreasing the phytotoxicity of the compounds (Yuan et al., 2007). These enzymes also have other functions in herbicide detoxification, such as peroxidase activity and stress signaling (Dixon et al., 2002; Powles & Yu, 2010; Cummins et al., 2013). The *GSTF1* gene conferred greater tolerance to the herbicides chlorotoluron and fenoxaprop-P-ethyl in *A. myosuroides* and *L. rigidum*, causing multiple resistance (Cummins et al., 2013). In rice, overexpression of the *GSTL1* or *GSTL2* genes led to greater tolerance to chlorsulfuron and glyphosate (Hu, Qv, Xiao, & Huang, 2009; Hu, 2014). Likewise, the greater expression of the *GST1* gene has been related with resistance of *E. crus-galli* plants to quinclorac (Li et al., 2013).

The qRT-PCR (*quantitative reverse transcription-polymerase chain reaction*) is useful to access the gene expression, which is associated with enzyme activity. In this procedure it is necessary to analyze the stability of reference genes used for calculating the relative expression of the target gene. A number of studies demonstrate that the expression of reference genes widely used in this type of study can vary considerably with experimental conditions, tissues and species (Thellin et al., 1999; Stürzenbaum & Kille, 2001). The analysis of the stability in the expression of reference genes in weeds responding to herbicide stress has already been performed in *A. myosuroides* for the acetyl CoA carboxylase (ACCase) inhibitors herbicides (Petit, Permin, Heydel, & Délye, 2012), and in *L. rigidum* for ALS inhibitors herbicides (Duhoux & Délye, 2013). In these studies, the most stable reference gene for *A. myosuroides* and *L. rigidum* were genes coding for tubulin and CAP proteins, respectively. The stability of reference genes for barnyardgrass under stress by herbicides has not been investigated yet.

The objectives of this study were to evaluate the stability of reference genes and the expression of *CYP* and *GST* genes in imidazolinone-susceptible and -resistant barnyardgrass populations exposed to imazethapyr.

### 2. Method

#### 2.1 Plant Material

The source of susceptible population was at Engenheiro Coelho, SP (SUSSP01) and the resistant populations were at Arroio Grande-RS (ARRGR01) and Palmares do Sul-RS (PALM01). Barnyardgrass resistant populations were collected from paddy fields of South Brazil, where escapees of control with imidazolinone herbicides had occurred and historic use of Clearfield®-rice cultivars was known. The susceptible population SUSSP01 was originally from an area where no herbicides had been applied before and efficient control had been obtained during previous pot-studies using imazethapyr. The PALM01 population is resistant to imidazolinones due to the mutation Ser653Asn in *ALS*, but enhanced metabolism was also identified through P450 inhibitors (Dalazen, Pisoni, Raffaeli, & Merotto Jr., in press). Meanwhile, in the population AARRGR01 there was absence of *ALS*-gene mutation associated with herbicide resistance (Matzenbacher et al., 2015).

The seed dormancy overcoming was performed by immersing the seeds in KNO3 solution (0.2%) at a temperature of 25 °C until germination (radicle emission), which occurred approximately four days after immersion. The seeds were then rinsed in distilled water and placed in Petri dishes incubated at 25 °C until the
emergence of the first leaf. Then, seedlings were transplanted into 200 ml pots filled with a mixture of ultisol and organic compound at a ratio of 10:1, and the mineral fertilizer (05-20-20 NPK complex) at 2.5 g kg⁻¹. Plants were maintained in greenhouse with temperatures ranging from 25 to 27 °C, relative humidity of about 70% and photoperiod of 14/10 hours (day/night).

The herbicide imazethapyr was sprayed at the label dose of 106 g ha⁻¹ plus adjuvant (Dash 0.5% v/v). The spray treatments were applied in plants with three to four leaves stage using an automated spray chamber with nozzle TJ8002E, spray pressure of 2.89 bar and speed of 1.16 m s⁻¹, resulting in a spray volume of 200 L ha⁻¹.

Leaves of barnyardgrass plants treated and untreated with imazethapyr were analyzed. The samples were collected before (T0, untreated control) and 24 hours after herbicide spraying (T24). Three biological replicates were used. The samples were collected in a 1.5 mL microtube and immediately conditioned in liquid nitrogen (LN₂). The samples were stored in an ultrafreezer (-85 °C) until the RNA extractions.

2.2 RNA Extraction and cDNA Synthesis

RNA extraction was performed using the Trizol® method (Invitrogen). The RNA was quantified in a spectrophotometer (Genesys 2™, Thermo Spectronic) at a wavelength of 260 nm and diluted in RNAse free water at 1 μg μL⁻¹. Each sample was purified with DNase® I (Invitrogen) in an amount of 1 μg of total RNA, as per the manufacturer's recommendation. The next step consisted of obtaining the cDNA strand from RNA through SuperScrip® III reverse transcriptase (Invitrogen) in the amount of 1 μg of RNA using polidT primers. 

2.3 Candidate Genes and Primer Design

The reference genes (Table B1) evaluated for stability analysis were selected from the study of Duhoux and Délye (2013), including actin, CAP (catabolite activator protein), EF1 (elongation factor 1), rubisco, ubiquitin, 18S (18S ribosomal RNA) and 28S (28S ribosomal RNA). The candidate genes CYP and GST were chosen based on a large literature review (Table B2). The analyzed genes were CYP81A6, CYP81A12, CYP81A21, CYP71C30, CYP71AK2, CYP72A254, CYP72A3, GSTF1, and GSTL1. In addition to these genes, the expression of ALS and eIF4B (translation initiator factor) was considered. The eIF4B gene is present in the genus Echinochloa (Iwakami et al., 2014b) and its involvement in the detoxification of xenobiotics in other eukaryotic organisms has been reported (Kim et al., 2011).

The primers sequences were designed by the program Primer3Plus (http://www.bioinformatics.nl/cgi-bin/primer3plus/primer3plus.cgi) from the most conserved region of each sequence obtained in the Genbank (http://www.ncbi.nlm.nih.gov/genbank). For each gene, at least three primer pairs were designed based on annealing temperature around 60 °C, size close to 20 bp and expected fragment size around 100 bp.

2.4 qRT-PCR Analysis

The obtained cDNA was amplified by real-time PCR using the SYBR Green® kit. qRT-PCR analysis was performed with the 7300 Real-Time PCR System (Applied Biosystems) on 96-well plates PCR-96M2-HS-C® (Axygen) with a sealer MicroAmp® Optical Adhesive Film (Applied Biosystems).

The reactions were carried out in a final volume of 20 μL, consisting of 10 μL of the cDNA sample (diluted 1:100) in Mili-Q water; 10 μL of the constituents of the reaction composed of 2 μL 10X buffer, 0.5 μL dNTPs (10 μM of each nucleotide), 1.2 μL of MgCl₂ solution (50 mM), 2 μL of SYBR Green® (Invitrogen) diluted 1:100 (prepared at the time of use from diluted 100X solution), 0.2 μL of ROX Reference Dye, 0.1 μL Taq Platinum® (Invitrogen) and 0.4 μL of the combination of forward and reverse primers.

The amplification steps included an initial cycle of 95 °C for 5 min, followed by a 40-cycle sequence: started at 94 °C for 15 sec, 60 °C for 10 sec, 72 °C for 15 sec and 60 °C for 35 sec, and a final denaturation cycle of 95 °C for 15 sec, 60 °C for 60 sec, 95 °C for 15 sec and 60 °C for 15 sec.

2.5 Reference Genes Stability Analysis

Analysis of the stability of the reference genes was performed from the Ct values obtained in the qRT-PCR reaction (Wang, Ma, Huang, & Zhang, 2015). The software RefFinder (http://fulxie.0fees.us/?ckattempt=1) was used, which is based on the algorithms geNorm, Norm finder, BestKeeper and Delta Ct method. The stability coefficients (SC) were classified in each of the algorithms and, finally, a comprehensive classification of all the algorithms was determined by the program RefFinder. A lower SC value indicate greater the stability of the reference gene. The two genes with lower stability coefficients were used as reference genes in calculations of relative expression of genes related to the herbicide imazethapyr.
2.6 Analysis of Gene Expression

The Ct values mean, the standard deviation, and the confidence interval per treatment were calculated. The relative quantification was performed by adjusting the curves by analyzing the efficiency of the PCR using the LinRegPCR software (version 12.2), which analyzes the exponential amplification curve. Values of R > 0.99, with efficiency between 1.8 and 2, and numbers of points greater than 4 were accepted (Tuomi, Voorbraak, Jones, & Ruijter, 2010).

The relative expression was calculated using the ΔCt method (Dussault & Pouliot, 2006), by the equation,

\[ \Delta \Delta Ct = (Ct_{target} - Ct_{reference}) - (Ct_{calibrator} - Ct_{reference}) \]

(1)

where, ΔCt is the relative expression of the gene, and the application of the result in \(2^{\Delta \Delta Ct}\) gives the variation dimension. The Ctreference value was determined by the average of the reference genes that presented lower coefficients in the stability analysis performed in the RefFinder computational program as described above.

2.7 Conventional PCR Analysis

Polymerase Chain Reaction (PCR) was carried out using Thermal Cycler (Life Science Research, Bio-Rad). Each reaction was set up in 30 μl volume, containing 50 ng genomic DNA, 0.2 mM dNTPs, 0.35 U Taq DNA polymerase (Invitrogen), 0.15 μM primers (forward and reverse) (Table B2), 1.3% of DMSO 100%, 3 mM of MgCl2, 1X PCR buffer and water to complete the total volume. The reactions were carried out using the following cycling parameters: 94 °C for 3 min, followed by 35 cycles of 94 °C for 30 sec, 57 °C for 30 sec, 72 °C for 60 sec, and 72 °C for 5 min. PCR products were separated by electrophoresis in 2.0% agarose gels and product size was identified using 100 bp DNA Ladder (Invitrogen). Subsequently, the PCR products were visualized using the UV transilluminator L-PIX IMAGE Release 2.6 (Locus Bioteologica).

The expression was evaluated in the imidazolinone-susceptible (SUSSP01) and in the -resistant (ARRGR01) populations, which does not have mutation in ALS gene associated with herbicide resistance. Gene expression was evaluated in untreated leaves (T0) and leaves collected 24 h after treatment with imazethapyr (T24), as described in section 2.1.

3. Results and Discussion

3.1 Reference Genes Stability Analysis

The expression levels of the reference genes in all samples (barnyardgrass population x herbicide treatment) evaluated by the Ct values from qRT-PCR reactions are presented in Figure 1. A lower Ct value indicates a greater transcription of a particular gene. The mean Ct values for the seven reference genes evaluated ranged from 23.35 to 32.75. The gene with the highest number of transcripts was rubisco, followed by 18S, 28S, EF1, actin, CAP and ubiquitin.

![Figure 1. Ct values of reference genes in leaves of barnyardgrass (Echinochloa crus-galli) susceptible and resistant to imidazolinones evaluated before and 24 hours after spraying of imazethapyr (vertical bars indicate the standard deviation)](image-url)
Five algorithms were used to determine the stability of the reference genes. The Delta Ct method determines the stability of the genes according to the variation of the Ct value (Silver et al., 2006). In the geNorm method, the stability coefficient is called the M-value, which is calculated according to the variation of Ct eliminating the two least stable genes in the first round of the calculation. In this analysis, the lower the M-value the greater the stability of the reference gene (Vandesompele et al., 2002). The NormFinder method calculates the stability coefficient S and the standard deviation (SD). Reference genes are considerable stable when both S and SD values are low (Andersen, Jensen, & Orntoft, 2004). In the BestKeeper method the variation of the Ct values and the standard deviation (SD) of each gene are used for the calculation of stability. Genes with SD < 1 are considered stable and ordered based on the correlation between the Ct value and the geometric mean of the Ct values of all values with SD < 1 (BestKeeper index). The candidate genes with the highest correlation with the BestKeeper index are considered to be the most stable (Pfaffl, Tichopad, Prgomet, & Neuvians, 2004). Finally, the RefFinder method uses an integrated way the other methods of analysis, classifying them comprehensively. Based on the ordering of the methods described above, the RefFinder method assigns an appropriate weight to an individual gene and calculates the geometric mean of its weights for the comprehensive classification of candidate reference genes (Xie, Xiao, Chen, Xu, & Zhang, 2012). The stability analysis of the reference genes indicated low variations of the classification and stability among the methods used (Table 1 and Appendix A).

<table>
<thead>
<tr>
<th>Method</th>
<th>Ranking order (better-good-average)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Delta CT</td>
<td>18S Act Ubi EF1 28S CAP Rub</td>
</tr>
<tr>
<td>BestKeeper</td>
<td>18S Act 28S Ubi EF1 CAP Rub</td>
</tr>
<tr>
<td>NormFinder</td>
<td>Act 18S Ubi EF1 28S CAP Rub</td>
</tr>
<tr>
<td>geNorm</td>
<td>18S/Act Ubi 28S EF1 CAP Rub</td>
</tr>
<tr>
<td>RefFinder</td>
<td>18S Act Ubi 28S EF1 CAP Rub</td>
</tr>
</tbody>
</table>

Note. 18S: 18S ribosomal RNA; Act: actin; Ubi: ubiquitin; 28S: 28S ribosomal RNA; EF1: elongation factor 1; CAP: catabolite activator protein; Rub: Rubisco.

3.2 Expression Analysis of the ALS Gene

The relative expression of the ALS gene was not altered by the evaluated treatments (Figure 2). The expression in both resistant populations ARRGR01 and PALMS01 was similar to that observed in SUSSP01 susceptible population. Spraying of imazethapyr had no effect on the relative expression of the ALS gene. This indicated that the greater expression or the greater number of copies of the ALS gene is not a mechanism of resistance to imazethapyr in these populations of barnyardgrass.
3.3 CYP and GST Genes Expression

The expression of the genes CYP81A12, CYP81A21, CYP71C30, CYP71AK2, CYP72A254, CYP72A31, GST1 and GSTL1 was not significant different among the populations evaluated and neither due to imazethapyr spraying (data not shown). Otherwise, meanwhile the CYP81A6 and the GSTF1 presented higher expression in these conditions. The CYP81A6 gene showed higher expression in both resistant populations, especially when the plants were treated with imazethapyr (Figures 3 and 6). The relative expression of the gene CYP81A6 for resistant population ARRGR01 treated with the herbicide imazethapyr (T24) was 9.61 times greater in comparison to the untreated susceptible plants (T0). For the PALMS01 resistant population, the relative expression was 8.44 times higher when the plants were sprayed with imazethapyr (T24). The expression of the CYP81A6 gene, even in the absence of herbicide, was 4.96 fold higher in the PALMS01 population than the untreated susceptible population. For the resistant population ARRGR01, the relative expression of this gene in untreated plants was 1.67. Thus, in addition to the expression of the CYP81A6 gene in the resistant populations, the spraying of the herbicide imazethapyr induced greater expression of this gene in resistant plants.
The relationship of the *CYP81A6* gene to herbicide detoxification has already been reported in crops and weeds. Some poaceous species, such as rice (*O. sativa* L.) and wheat (*Triticum aestivum* L.) are naturally tolerant to bentazon herbicides (FSII inhibitor) and to sulfonyleureas (ALS inhibitors). The selectivity of these herbicides on these crops is attributed to the greater expression of the *CYP81A6* gene, responsible for the phase I of the detoxification of these herbicides in the plants (Pan et al., 2006; Zhang et al., 2007). In addition, the insertion of this gene into non-poaceous species such as *A. thaliana* L. and *Nicotiana tabacum* L. conferred tolerance to these herbicides (C. Liu, S. Liu, F. Wang, Y. Wang, & K. Liu, 2012). In contrast, the silencing of this gene by means of interference RNA made rice plants, previously tolerant, sensitive to the herbicide bentazon, proving the function of this gene in the process of detoxification of herbicides in plants (Liu et al., 2012). In the present study, spraying of the herbicide imazethapyr induced *CYP81A6* gene expression in both resistant populations evaluated (Figure 3). Similar results were observed in rice, in which both bentazon and metsulfuron-methyl spraying caused increased *CYP81A6* gene expression when compared to untreated plants (Lu et al., 2015).

The *GSTF1* gene showed highest expression in the PALMS01 resistant population in leaves collected 24 hours after imazethapyr spraying (Figure 4). In plants of the ARRG01 resistant population treated with imazethapyr (T24), the expression of this gene was 6.41 times higher than the untreated susceptible population SUSSP01 (T0) (Figures 4 and 6). However, in the absence of the herbicide (T0), the relative expression was similar to that observed in the susceptible population. In the PALMS01 population, the relative expression of the *GSTF1* gene was higher in both untreated (T0) and treated (T24) plants. In untreated plants (T0), the relative expression was 10.24 times higher than the susceptible population, whereas in treated plants (T24), the expression was 12.30 times greater. It is also observed that, even in susceptible plants, there was greater expression of this gene in plants treated with imazethapyr (T24) in comparison with the untreated control plants.

![Figure 4. Relative expression of the GSTF1 gene in imidazolinone-susceptible (SUSSP01) and -resistant (ARRRG01 and PALMS01) barnyardgrass (*Echinochloa crus-galli*), in response to spraying of imazethapyr. T0; Untreated leaves; T24: leaves collected 24 hours after spraying of imazethapyr. Vertical bars indicate the confidence interval (α = 0.05)](image)

The GST enzymes are involved in phase II of the herbicide metabolism in plants. In this phase occurs the conjugation of the herbicide molecule with glutathione, after being activated in phase I (Yuan, Tranel, & Stewart Jr., 2007). However, these enzymes may also present other functions in plants, such as peroxidase activity and signaling for the production of secondary metabolites in the stress protection (Dixon, Skipsey, & Edwards, 2010; Powles & Yu, 2010). In *A. myosurioides*, the *GSTF1* gene showed low herbicide detoxifying activity, however, it was highly active as a glutathione peroxidase, catalyzing the reduction of organic hydroperoxides (Cummins et al., 1999). The presence of the same gene in *A. thaliana* resulted in multiple resistance to herbicides due to increased activity of peroxidase enzymes and increased accumulation of antioxidant compounds (glutathione, flavonoids and anthocyanins) (Cummins et al., 2013). Although there is little information on the formation of reactive oxygen species by ALS-inhibiting herbicides, some studies have demonstrated increase of the levels of certain antioxidant enzymes due to the presence of these herbicides (Wang et al., 2006; Wang, Zhou, & Ren, 2009).

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3.4 Expression Analysis of the eIF4B Gene

In addition to the greater expression of the CYP81A6 and GSTF1 genes, the greater expression of the eIF4B translation initiator factor (Figures 5 and 6) was also observed. The relative expression of this gene in the ARRGR01 resistant population was 2.51 times higher in untreated (T0) and 6.46 fold higher in treated (T24) leaves with imazethapyr. In the PALMS01 resistant population, relative expression was 2.30 times higher in untreated (T0) and 5.96 fold higher in treated (T24) leaves with imazethapyr. In the SUSSP01 susceptible population, the relative expression of this gene was higher in leaves treated with the herbicide in comparison with the untreated plants. For all populations, spraying of imazethapyr induced expression of the eIF4B gene. The basal expression (untreated plants) was higher in the resistant populations compared to the susceptible population.

![Graph showing relative expression of eIF4B gene](image)

Figure 5. Relative expression of the eIF4B gene in imidazolinone-susceptible (SUSSP01) and -resistant (ARRGR01 and PALMS01) barnyardgrass (Echinochloa crus-galli), in response to spraying of imazethapyr. T0; Untreated leaves; T24: leaves collected 24 hours after spraying of imazethapyr. Vertical bars indicate the confidence interval (α = 0.05).

The eIF4B translation initiation factor, along with other proteins that also act as translation initiators, is responsible for the process of recognition of mRNA by the ribosomes during the process of protein synthesis (Spriggs, Bushell, & Willis, 2010). This protein presents a helicase function, responsible for the unwinding of some mRNA in the 5'-UTR region and initial codon (AUG) exposure for the initiation of translation in ribosomes (Shahbazian et al., 2010). The differential expression of this translation-initiating factor is related to the occurrence of stresses in plants, being important in post-transcriptional gene regulation in eukaryotes (Sonenberg & Hinnebusch, 2009). In addition, the higher expression of this gene is directly related to the lower efficiency of chemotherapy in the treatment of cancer cells, and there is a high correlation between the greater expression of the eIF4B gene and the higher production of proteins responsible for cancerous diseases (Dzgen, Barron, Natarajan, Widlund, & Rheinwald, 2013). Thus, depending on the structure of the mRNA in the 5'UTR region, the higher expression of the gene eIF4B may culminate in the higher content of detoxifying enzymes, such as P450 and GST. Therefore, the greater expression of this gene and, consequently, the greater production of this protein, can contribute to the regulation of the translation of the mRNA of enzymes P450, regulating its production and the capacity to metabolize the herbicides in resistant plants. Although this mechanism has already been reported in cases of chemotherapy for the treatment of cancer in humans (Kim et al., 2011; Degen et al., 2013), this is the first report of the possible involvement of this protein on herbicide resistance in weeds.
Figure 6. Expression of CYP81A6 (A), GSTF1 (B), and eIF4B (C) genes in imidazolinone-susceptible (SUSSP01) and -resistant (ARRGR01) barnyardgrass (Echinochloa crus-galli), in response to imazethapyr spray. T0: untreated leaves; T24: leaves collected 24 hours after the herbicide treatment.

Stability analysis of reference genes expression demonstrated that the 18S and actin were the most stable in imidazolinone-susceptible and -resistant barnyardgrass leaves, in the presence or absence of the stress caused by the herbicide imazethapyr. Expression analysis of the ALS gene demonstrated that the amount of transcripts in all populations evaluated was similar among population and imazethapyr treatment. Thus, it can be stated that the greater expression of the ALS gene is not the cause of the resistance of the populations evaluated in this study (Figure 2).

The CYP81A6 and GSTF1 genes showed higher relative expression in resistant populations. The CYP81A6 gene presented expression 9.61 and 8.44 times higher in the ARRG01 and PALMS01 populations, respectively. Expression of this gene was induced by the spraying of the herbicide imazethapyr. The GSTF1 gene presented higher relative expression in the PALMS01 population, being 12.3 times higher in plants treated with imazethapyr. In addition to these genes, the greater expression of the eIF4B translation initiator factor may be involved in the resistance of these populations to the herbicide imazethapyr. The results demonstrate that degradation enhancement by P450 and GST enzymes is involved in the resistance mechanism of these populations. This type of NTSR is particularly important because it has evolved in multiple- or cross-resistance. In these cases, the simple alteration of the chemical group or herbicide mechanism of action may not be efficacious for the management of resistant weeds. Also, the occurrence of resistance mechanisms not yet described in weeds, related to the greater activity of proteins responsible for the recognition of mRNA in the protein synthesis, require the accomplishment of more detailed studies.

The present study points out the importance of the genes CYP81A6 and GSTF1 in the degradation enhancement of imazethapyr in barnyardgras. Moreover, it is reported for the first time that the translation initiator factor eIF4B was related with the herbicide resistance. These results contribute for the improving the knowledge about the genetic regulation of herbicide NTSR in weeds and could be used for developing the P450s pathway associated with the herbicida detoxification.

References


Appendix A

Ranking order and stability coefficients of candidate reference genes in leaves of imidazolinone-resistant and -susceptible barnyardgrass (Echinochloa crus-galli), in response to treatment with imazethapyr.

![Graph showing ranking order and stability coefficients of candidate reference genes in leaves of imidazolinone-resistant and -susceptible barnyardgrass.]

Note. A lower stability coefficient value indicates greater reference gene stability.
Appendix B

Genes and primers analyzed in the research

Table B1. Reference genes and primers sequences used for the qRT-PCR analysis

<table>
<thead>
<tr>
<th>Gene†</th>
<th>Forward primer sequence</th>
<th>Reverse primer sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Act</td>
<td>ttacaaggtagttagtgag</td>
<td>acacaggacacaaaccaacta</td>
</tr>
<tr>
<td>CAP</td>
<td>ctccagggaagactgtgaag</td>
<td>cttgagaagcccaaatctaa</td>
</tr>
<tr>
<td>EF1</td>
<td>caactctgtactcatacaaa</td>
<td>gtaacagtgagccctgttga</td>
</tr>
<tr>
<td>Rub</td>
<td>ggagatgtaaaccagactgta</td>
<td>gttgtcctagctacagtgaga</td>
</tr>
<tr>
<td>Ubi</td>
<td>caagaagaagctacacaaag</td>
<td>gaccttgataactccagag</td>
</tr>
<tr>
<td>18S</td>
<td>gtagcggaggatttaggttc</td>
<td>tgtgaggtgtggtaatttg</td>
</tr>
<tr>
<td>28S</td>
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<td>tgtgaaaatgtaaagggc</td>
</tr>
</tbody>
</table>

Note. † Act: actin; CAP: catabolite activator protein; EF1: elongation factor 1; Rub: Rubisco; Ubi: ubiquitin; 18S: 18S ribosomal RNA; 28S: 28 S ribosomal RNA.

Table B2. DNA sequences of the primers used for real-time RT-PCR analysis of CYP and GST genes

<table>
<thead>
<tr>
<th>Gene</th>
<th>NCBI number</th>
<th>Forward primer sequence</th>
<th>Reverse primer sequence</th>
<th>Reference</th>
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<tr>
<td>CYP81A6†</td>
<td>DQ341412</td>
<td>gaacccggaggtttaaaca</td>
<td>ggcgaagatgetctctctct</td>
<td>Zhang et al., 2007</td>
</tr>
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<td>aacacccagtctctctgtaa</td>
<td>atctgtcctgacacctcgc</td>
<td>Liu et al., 2012b</td>
</tr>
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<td>caagttctgcatcacaacccga</td>
<td>Iwakami et al., 2014a</td>
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<td>Fischer et al., 2001</td>
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<td>Iwakami et al., 2014b</td>
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<td>Matzenbacher et al., 2015</td>
</tr>
</tbody>
</table>

Note. † Primers sequences designed from NCBI number using the software Primer3Plus. §Primer sequence available in the reference.

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