Seed moisture content can be used to accelerate dormancy release during after-ripening of *Urochloa humidicola* cv. Llanero spikelets

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**ABSTRACT** Increased demand for livestock products is leading to research on more productive pastures adapted to different environments. *Urochloa humidicola* is a species adapted to low-fertility conditions and to environments occasionally subjected to temporary flooding and with highly dormant seeds at harvest. Mechanical and sulphuric acid (H₂SO₄) scarification are used to release these seeds from dormancy; although, there are several side effects. Hydrogen peroxide (H₂O₂) is linked to dormancy release in many crop species. The objective of this research was to characterize the dormancy release and H₂O₂ accumulation during one year of storage in dormant spikelets of *U. humidicola* cv. Llanero equilibrated to 4.5% and 50% relative humidity (RH) and stored at 20 ºC, during which time the seeds had moisture contents (MCs) of 0.03 and 0.1 g H₂O g⁻¹ dry weight (DW), respectively. The seeds were evaluated initially and at 3, 6 and 12 months; the seeds from spikelets or just caryopses were evaluated. Germination and dormancy percentage, H₂O₂ content, superoxide dismutase (SOD) and peroxidase (PRX) activity were determined. The germination and H₂O₂ content increased mainly in the caryopses, and SOD activity increased. Moreover, dormancy and PRX activity decreased during storage. For the first time, it was shown, in *U. humidicola* cv. Llanero that dormancy release during storage is caused mainly by the accumulation of reactive oxygen species (ROS) in the spikelets even under ultra-dry storage conditions (4.5% RH), and this process could be accelerated by seed storage at a MC of 0.1 g H₂O g⁻¹ DW at 20 ºC.

**Key words:** Brachiaria, germination, oxidative window, ultra-dry storage, ROS.

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**Uso do teor de água em sementes para acelerar a quebra de dormência pós-colheita de caryopses de *Urochloa humidicola* cv. Llanero**

**RESUMO:** O aumento da demanda de produtos carnêos tem levado à pesquisa de pastagens mais adaptadas para diferentes ambientes. *Urochloa humidicola* é uma espécie adaptada a solos de baixa fertilidade e sujeitos a alagamentos ocasionais e que apresentam um alto grau de dormência por ocasião da colheita. Escarificação mecânica e por ácido sulfúrico são normalmente usadas para quebrar a dormência nesta espécie, porém com diversos efeitos colaterais. Peróxido de hidrogênio (H₂O₂) está ligado à quebra de dormência em diversas culturas. O objetivo deste trabalho foi caracterizar a perda de dormência e o acúmulo de H₂O₂, durante um ano de armazenamento em espiguetas dormentes de *U. humidicola* cv. Llanero, equilibradas a 4,5% e 50% de umidade relativa (UR) e estocadas a 20 ºC; com um conteúdo de água de 0,03 e 0,1 g H₂O g⁻¹ matéria seca (MS), respectivamente. As sementes foram avaliadas inicialmente (tempo zero) e aos três, seis e 12 meses; como espiguetas ou caryopses. Foram avaliadas as percentagens de germinação e dormência, teor de H₂O₂, e as atividades de superóxido dismutase (SOD) e peroxidase (PRX). A germinação e o H₂O₂ aumentaram, nas caryopses, bem como a atividade de SOD. A dormência e a atividade de PRX diminuíram durante o armazenamento. Pela primeira vez, demonstrou-se em *U. humidicola* cv. Llanero, que a liberação da dormência durante o armazenamento é causado principalmente pelo acúmulo de espécies reativas de oxigênio (ROS) em espiguetas mesmo em condições de armazenamento ultra seco (4,5% UR), e que este processo pode ser acelerado por armazenamento de sementes com teor de água de 0,1 g H₂O g⁻¹ MS a 20 ºC.

**Palavras-chave:** Brachiaria, germinação, janela oxidativa, armazenamento ultra seco, ROS.

**INTRODUCTION**

The increasing demand for livestock products is driven by population growth, increased richness and urbanization, especially in developing countries. This global demand is projected to increase by approximately 70% by 2050 to meet an estimated population of 9.6 billion. The larger part of this demand will be achieved through intensive livestock production (concentrated animal feeding operations) and traditional systems (animals raised in pastures) (FAO, 2018).

In a survey about changes in natural cover and land use in Brazil (IBGE, 2020), from 2010 to 2014, the area occupied by natural and non-natural pastures decreased, but even though beef production and the number of animals per unit area in Brazil increased during that time, Brazil has the second largest cattle herd (232 million heads); as such, Brazil has again attained the position of the highest global exporter of in natura beef (UDSA, 2018).
The increase in animal production in Brazil is due to technological advances related to genetics, animal nutrition and pasture quality. Brazilian pastures with *Urochloa* (syn. *Brachiaria*) spp. have been increasing in area, mainly because of the robustness and adaptability of these species to a range of both climate and soil conditions (CARDOSO et al., 2014).

Dormancy is defined as a temporary failure of a seed to complete germination under favourable conditions, allowing dispersal in time and space (BEWLEY et al., 2013). *Urochloa* seeds are characterized as exhibiting non-deep physiological dormancy (PD; BASKIN & BASKIN, 2014); this is a problem for the seed industry, as they needed to be maintained under storage until dormancy is released and for initial pasture establishment, which is impaired by delays in seedling emergence, leading to problems of weed infestation (MARTINS & SILVA, 2003).

One of the causes of PD in Poaceae species is the impermeability of the seed coat to oxygen, as noted for seeds of *Elymus nutans*, *Lolium perenne*, *Leymus chinenesis*, *Avena sativa* (LV et al., 2017), *Triticum vulgare*, *Oryza sativa*, *Zea mays*, *Panicum miliaceum*, *Setaria italica* and the *Sorghum bicolor* x *S. sudanensis* hybrid (SUN et al., 2018).

Many studies have been conducted to increase the germination of forage grasses either by chemicals such as sulphuric acid (H2SO4; MESCHEDE et al., 2004, USBERTI & MARTINS, 2007, COSTA et al., 2011) or by mechanical scarification, both of which are commonly used methods applied to seeds with gas-impermeable seed coats, such as those of *Urochloa* (BRASIL 2009). H2SO4 easily degrades residue (ALVAREZ et al., 2007). Thus, it is necessary to neutralize residual H2SO4 to avoid environmental problems; moreover, H2SO4 has negative effects on the germination of *U. humidicola* (COSTA et al., 2011; USBERTI & MARTINS, 2007). H2SO4 also requires an excessive amount of care to circumvent risks for workers and for the environment as an adequate destination of its residue (ALVAREZ et al., 2007).

Seed germination is associated with water absorption, adequate temperature and oxygen supply (BEWLEY et al., 2013). It is divided into three imbibition phases: Phase I, which occurs because of water potential differences; Phase II, which involves the slowing of absorption, during which time several metabolic reactions occur, leading to the emergence of the primary root; and Phase III, which involves the resumption of water absorption due to the growth of the embryonic axis, degradation of reserve substances and growth of the seedling (BEWLEY et al., 2013). However, dormant seeds do not reach Phase III, even when adequate conditions are supplied. *Urochloa* seed dormancy can be released by exogenous application of gibberellic acid (GA) (SILVA et al., 2013; SORIGOTTI et al., 2016). Dormancy release occurs when there is a decrease in abscisic acid (ABA) levels and an increase in GA levels, as this hormone promotes the synthesis of α-amylase, which is involved in the germination process (DIAZ-VIVANCOS et al., 2013).

Notably, the mechanisms underlying primary seed dormancy and its regulation after ripening are poorly understood (CHAHTANE et al., 2017). In dry (0.1 g H2O g−1 dry weight (DW)) or ultra-dry (0.03 g H2O g−1 DW) sunflower seeds, dormancy release is not associated with complex events ranging from gene transcription to post-translational modification of proteins because, in this state, the water is tightly linked to macromolecules, preventing the synthesis of new biomolecules (MEIMOUN et al., 2014). Therefore, the reactions triggering the breaking of dormancy of dry seeds are not associated with reactions driven by enzymes but are instead due to oxidative processes associated with reactive oxygen species (ROS).

ROS, such as superoxide anions (•O2−), hydrogen peroxide (H2O2), singlet oxygen (1O2) and hydroxyl radicals (•OH), are the principal triggers of many reactions in cells. They are produced mainly in the respiratory pathway and not only are involved in oxidative damage but also act as signalling molecules. ROS are also involved in many events in cells: elongation, cell death, peroxidation, membrane rupture, lignification and protection against herbivores (FOYER et al. 2017; FOYER & NOCTOR, 2016; MITTLER, 2017). In dry storage, ROS can weaken seed dormancy, and during this process, the seeds age. The enzymes superoxide dismutase (SOD), catalase (CAT) and ascorbate peroxidase (APX) compose the antioxidant system, which works to remove ROS and protect cells from oxidative damage (CHEN & ARORA, 2011).

Oxidative stress occurs when the amount of ROS produced is greater than the ability of the organism to eliminate them. ROS production was initially considered the production of only toxic molecules because of the impairment of primary metabolism resulted. However, there are reports in which seed germination is linked to internal ROS...
contents and the activities of the antioxidant system (BAILLY & KRANNER, 2011; DIAZ-VIVANCOS et al., 2013; GOMES & GARCIA, 2013; EL-MAAROUF-BOUTEAU et al., 2015; FOYER & NOCTOR, 2016; FOYER et al., 2017; MITTLER, 2017); thus, these molecules also act as signalling molecules in biological processes such as seed germination (BAILLY, 2004). According to BAILLY et al. (2008), germination occurs within an “oxidative window”; if the ROS levels are above this window, the seeds are undergoing oxidative damage, leading to deterioration and loss of germination, but below this window, the amount of ROS is too low to allow germination (BAILLY et al., 2008; FOYER et al., 2017; KUMAR et al., 2015; MITTLER, 2017).

ROS production is beneficial in the process of cell wall loosening because the increase in and release of ROS during germination coincides with the period in which the expanding embryo penetrates the seed coat and becomes exposed to the environment (SCHOPFER et al., 2001). During dry storage (0.03 g to 0.1 g H₂O g⁻¹ DW), the accumulation of ROS acts as a signal that regulates cellular activity during the post-maturation period (ORACZ et al., 2007), and breaking dormancy is related to changes in gene expression, which were shown to have been caused by the accumulation of ROS in sunflower seeds (EL-MAAROUF-BOUTEAU et al. 2007), but this variation in gene expression does not happen during the after-ripening period because of the low water content when the cytoplasm reaches the glass state (BEWLEY et al., 2013; MEIMOUN et al., 2014). At this moisture content (MC), the growth of microorganisms is prevented only because of chemical reactions (Amadori-Mailard reactions and those involving ROS) happen (BEWLEY et al., 2013).

The accumulation of H₂O₂ in many ways is directly related to decreasing amounts of ABA, which can lead to seed germination (BAILLY et al., 2008; EL-MAAROUF-BOUTEAU et al., 2007). The internal content of H₂O₂ is related to many phenomena in plants (SIEGEL, 1993) and germination, and exogenous H₂O₂ application was shown to induce germination of dormant Arabidopsis seeds by triggering ABA catabolism and by stimulating GA biosynthesis (LIU et al., 2010) and shown to induce seed germination in wild almond (ZEINALABEDINI et al., 2009), pea (BARBA-ESPÍN et al. 2010, 2011) and barley (BAHIN et al., 2011). This was confirmed by the use of diphenyliodonium (DPI), which decreases the production of O₂⁻ and; consequently, H₂O₂, increasing seed dormancy, while the application of exogenous H₂O₂ was shown to reverse the effects of DPI in Arabidopsis completely (LIU et al., 2010).

The objective of this research was to characterize the dormancy release and H₂O₂ accumulation in dormant spikelets of U. humidicola cv. Llanero equilibrated to conditions of 4.5% and 50% relative humidity (RH) and stored at 20 °C for one year.

**MATERIALS AND METHODS**

**Seed selection and conditioning**

This experiment was conducted with recently harvested pure seeds of Urochloa humidicola cv. Llanero whose caryopses were more than two-thirds filled. Seeds were selected by air flow in a South Dakota (DeLeo, Porto Alegre/RS) seed blower and were 90% viable according to the tetrazolium test (TZ), and 24% of seeds germinated.

The selected seed lot was split into two portions. The portions were placed into paper bags that were equilibrated over white silica gel (with orange silica used as an indicator), providing 4.5% RH, and over a lithium chloride solution (36.4 g of LiCl in 100 mL of H₂O), providing 50% RH (HAY et al., 2008). The MC was determined gravimetrically one week after the seeds were equilibrated; for this, one gram of seeds per treatment (two replicates each) was placed in an oven without air flow at 105 ± 3 °C for 24 h (BRASIL, 2009). The equilibrated seeds were maintained in two hermetic plastic boxes with a rubber seal (28.5 x 10.0 x 21.5 cm) during the storage period of 12 months at 20 ± 3 °C with silica gel or lithium chloride, during which time the MC of the seeds was maintained at 0.03 g or 0.10 g H₂O g⁻¹ DW. The RH was checked using a WP4T Dew Point Potentiometer (Decagon Devices Inc., Pullman, WA, USA) that measures the water potential (MPa), which was subsequently converted to RH (%).

The seeds were sampled at 90-day intervals throughout one year for evaluation. At each time point the seeds from spikelets were used (the caryopses coated with a lemma, palea and lower glumes) or only the caryopses.

**Seed viability**

Seed viability was evaluated by the TZ test according to the methods of (CUSTÓDIO et al., 2012), with 4 subsamples of 25 seeds from each period. Seeds (from the spikelets or just the caryopses) were imbibed for 16 h on moistened paper (the proportion of moisture was 2.5 times the mass of the dried paper) and then manually cut (longitudinally);
half of the seeds were immersed in a solution of 0.1% 2,3,5-triphenyl tetrazolium chloride for 5 h at 40 °C in the dark. Afterward, the solution was discarded, the seeds were rinsed, and the embryos were considered viable when the vital parts were coloured red. Results were expressed as the percentage of viable seeds.

**Germination**

The germination test was performed for four subsamples of 25 seeds from each period. The spikelets and caryopses were sown onto two paper towels within transparent plastic boxes, which were subsequently kept in a germinator whose temperature alternated between 15 and 35 °C, with an eight-hour photoperiod coinciding with the higher temperature (BRASIL, 2009). Germination was measured every two days for 21 days after sowing, and the seeds were considered germinated when the primary root protrusion reached 2 mm.

At the end of the germination test, the remaining seeds were subjected to a TZ test, as we sought to determine the percentage of dormant (coloured by TZ) and dead seeds (uncoloured by TZ). Owing to the discrepancy between results of the initial TZ coloured seeds and the results of the germination test, we calculated the average of these two results via the formula $D = \frac{D_{EG} + (1 - Tz)}{2}$, where $D$ is the percentage of dormant seeds (%), $D_{EG}$ is the number of viable seeds coloured by TZ after the germination test, $Tz$ is the percentage of seeds coloured by TZ (%) and $G$ (%) is the result of the germination test.

During each period, a portion of spikelets and caryopses were subjected to biochemical analysis for quantification of $\text{H}_2\text{O}_2$, guaiacol peroxidase (PRX) activity and SOD activity.

**Hydrogen peroxide quantification**

To quantify the amount of $\text{H}_2\text{O}_2$, four seed subsamples (1 g each) were frozen in liquid nitrogen, macerated in 3.5 mL of cold 5% trichloroacetic acid (TCA), and then centrifuged at 14,000 rpm for 5 minutes at 4 °C. After centrifugation, 800 μL of the supernatant was collected, and 200 μL of 10 mM ferrous ammonium sulphate and 100 μL of 2.5 mM potassium thiocyanate were added. Measurements were performed via a spectrophotometer at 480 nm. The amount of $\text{H}_2\text{O}_2$ was expressed in micromoles per gram of fresh mass according to the modified methodology of SAGISAKA (1976).

**Enzyme extraction**

Four subsamples of approximately 0.2 g of dry seeds from each treatment were frozen in liquid nitrogen and then ground with a mortar and pestle. Afterward, 0.1 M sodium phosphate buffer (pH 7.8) containing 0.4 g polyvinylpyrrolidone and consisting of 2 mM dithiothreitol, 0.1 mM EDTA and 1.25 mM PEG 4000 at 4 °C was added, after which the contents were mixed thoroughly. Dry seeds were used to measure the enzyme activity present in the coats (palea and lemmas) and in the caryopses in the absence of any de novo synthesis. Extracts were centrifuged at 12,000 g for 20 minutes, and the supernatant was divided into four aliquots, one of which was used for protein quantitation (BRADFORD, 1976). The remaining aliquots were immediately analysed or stored at -80 °C and then used to measure enzyme activity.

**Superoxide dismutase (SOD, EC.1.15.11)**

SOD activity was measured according to the method described by MORIYA et al. (2015). One unit of SOD (mg protein$^{-1}$) was defined as the enzyme activity capable of inhibiting the photoreduction of nitro blue tetrazolium (NBT) to blue formazan by 50%.

Fifty microlitres of the extract was added to 0.1 M phosphate buffer (pH 7.8) added of 1.3 μM riboflavin, 13 mM methionine and 63 μM NBT. Tubes were incubated at 25 °C for 15 minutes under fluorescent light, after which the absorbance was measured at 560 nm. Tubes containing the same mixture and 50 μL of the extraction buffer and without sample extracts were used as controls. The SOD activity data were normalized to the protein content according to the BRADFORD (1976) method and expressed in units of SOD per milligram of protein (LEI et al., 2005).

**Peroxidase (PRX, EC. 1.11.1.7)**

Four 100 mg subsamples of seeds were frozen in liquid nitrogen, crushed with a mortar with a pestle, homogenized in 3 mL of 0.01 mol L$^{-1}$ sodium phosphate buffer (pH 6.0) and centrifuged at 12,000 rpm for 25 minutes at 4 °C. The PRX activity was measured by the addition of one microgram of protein adjusted to 100 μL of the supernatant in 2.9 mL of 0.01 mol L$^{-1}$ phosphate buffer (pH 6.0) (ARAÚJO et al., 2005) added of 13 mM guaiacol and 5 mM H$\text{O}_2$ (MORIYA et al., 2015). The mixture was incubated for 20 minutes at 30 °C, after which its absorbance was measured via a spectrophotometer at 470 nm. The enzyme activity was expressed as A$\text{A}_{470}$ nm per minute per milligram of fresh sample.

**Statistical analysis**

The experiment was conducted in accordance with a completely randomized design.
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RESULTS

The initial germination of the spikelets was 24% but increased with increasing storage duration and RH. The spikelets stored with greater water content achieved greater germination faster than those stored with lower water content. Compared with the spikelets, the caryopses exhibited a 141.7% increase in initial germination, showing that the physical covers of the caryopses constitute a barrier to germination. With respect to the caryopses germination, no difference was detected between the storage environments, and the maximum germination was reached at six months of storage (Table 1). The maximum germination of the spikelets was achieved after 12 months of storage, regardless of the storage RH, but, the dormancy release was faster in the moister environment than under dry conditions. The maximum germination point of the caryopses – nine months – was lower than that of the spikelets, and the difference was attributed to removal of the covers; the wounds resulting from these injuries could serve as gateways for pathogens and might have compromised germination (Table 1).

A decrease in dormancy was achieved after six months of spikelets storage, independent of the RH of the storage environment (Table 2). The dormancy decreased until the third month, when the coats were removed and when the caryopses were isolated. Afterward, there was no difference between them, independent of the use of caryopses or spikelets (Table 2). This finding seems to be associated with an increase in the endogenous content of H$_2$O$_2$ (Table 3), while the loss of dormancy with the removal of the coverings did not alter the levels of H$_2$O$_2$ (Table 3).

The SOD activity was initially greater in the spikelets than in the caryopses, but after three months of storage, the removal of the coats of the caryopses increased the SOD activity, which stayed high until the end of the experiment. After six months, samples from the ultra-dry storage conditions (4.5% RH) presented high activity (Table 4).

The PRX activity was greater in the spikelets than in the caryopses at the first evaluation (0 storage period). However, in the other periods, no difference was observed between the spikelets and the caryopses. The PRX activity was high for up to 3 months of storage, but it decreased after that (Table 5). A portion of the PRX activity was seemingly initially linked to the seed coat, as shown in Table 5 (month zero).

Antioxidant metabolism, represented by SOD and PRX activity, decreased in the spikelets during the storage period (Tables 4 and 5), and for the caryopses, this also happened, albeit just for PRX. However, the SOD activity in the caryopses increased over time (Table 5).

The maintenance of seeds during after-ripening generates ROS, either by ageing or by stress conditions such as high RH and high temperature. In this paper, seeds were stored in two different environments, one moister (50% RH) than the other (4.5% RH), at the same temperature. In both cases, the generation of ROS could be related to a gradual increase in germination (Table 1) by dormancy release (Table 2) and an increase in H$_2$O$_2$ content (Table 3).

After 12 months, the maximum values of germination (Table 1) and the minimum dormancy

| Seed treatment     | 0  | 3  | 6  | 12 |
|--------------------|----|----|----|----|
|                   | Germination (%) |                |    |    |
| Spikelet 4.5% RH   | 24.0 ±2.3 bC   | 46.0 ±8.9 bB   | 88.0±4.9 aA | 92.0±4.0 aA |
| Spikelet 50% RH    | 24.0 ±2.3 bC   | 76.0 ±4.0 aB   | 79.0±5.7 aB | 95.0± 2.5aA |
| Caryopses 4.5% RH  | 58.0 ±4.2 aB   | 69.0 ±8.5 aB   | 89.0±4.1 aA | 77.0±5.2 bA |
| Caryopses 50% RH   | 58.0 ±4.2 aB   | 79.0±2.5 aA    | 85.0±3.4 aA | 74.0±3.5 bA |

Means followed by equal letters, lowercase in the column and upper case in the row, do not differ by Scott-Knott test (P<0.05).
(Table 2) were recorded for all seed treatments. For spikelets stored at 50% RH, there were a faster lack of dormancy (Table 2) and a faster accumulation of H$_2$O$_2$, differing from those observed in the other treatments after 6 months of storage (Table 3).

When the activity of PRX decreased during storage (Table 5), there was a natural increase in the amount of H$_2$O$_2$ (Table 3).

Germination and dormancy had a strong coefficient of determination under all conditions (Table 6), but there were strong negative correlations (Table 6) in the three situations, with spikelets at 4.5% RH and spikelets and caryopses at 50% RH. In the case of germination and H$_2$O$_2$, there were moderate correlations for spikelets but not for caryopses, and in the case of germination and SOD, there were moderate correlations for spikelets and a strong correlation for caryopses under both storage conditions. Between germination and PRX, there were moderate correlations for the spikelets but not for the caryopses. At 4.5% RH, there was a weak correlation between dormancy and H$_2$O$_2$ for the spikelets and a strong correlation at 50% RH. No correlations were detected between dormancy and SOD at 4.5% RH either for the spikelets or caryopses, but moderate correlations were detected for the spikelets and caryopses stored at 50% RH. With respect to dormancy and PRX, there was a strong correlation for the spikelets stored at 4.5% RH and moderate correlations for other combinations.

**DISCUSSION**

The removal of the covers was related to dormancy release, but it was not enough to eliminate the dormancy completely or to increase germination to the maximum in this study, even though the covers were impermeable to gas (LV et al., 2017; SUN et al., 2018); the after-ripening process during storage was also related to this.

The maximum values of germination observed after 12 months might have occurred because seeds with greater moisture have a greater respiration rate than do relatively drier seeds, which means that the former would accumulate more ROS, such as H$_2$O$_2$, which is a trigger for many reactions,
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including dormancy release (EL-MAAROUF-BOUTEAU & BAILLY, 2008, EL-MAAROUF-BOUTEAU et al., 2015, MITTLER, 2017).

The H$_2$O$_2$ increase during storage was an effect of a relatively high ROS production and was related to a decrease in the scavenger (PRX) activity during storage, and as a result, a dormancy release occurred during the storage period (after-ripening), followed by the start of the decrease in germination, such as that which occurred for the caryopses in both treatments; this imbalance of the antioxidant system and ROS production is a characterization of oxidative stress (EL-MAAROUF-BOUTEAU & BAILLY, 2008; MITTLER et al., 2017).  

ROS are related to the balance of hormones, mainly ABA (EL-MAAROUF-BOUTEAU & BAILLY, 2008; BAHIN et al., 2011; BAILLY & KRANNER, 2011; BARBA-ESPÍN et al., 2011; DIAZ-VIVANCOS et al., 2013; EL-MAAROUF-BOUTEAU et al., 2015); consequently, ABA is involved in dormancy because during dormancy, the ABA pathway is active and inhibits germination. High amounts of ABA can maintain high levels of scavenger enzymes, resulting in low amounts of ROS during the soaking of dormant seeds (EL-MAAROUF-BOUTEAU & BAILLY, 2008).

According to EL-MAAROUF-BOUTEAU & BAILLY (2008) and a review written by KUMAR et al. (2015), there is a certain amount of ROS needed for germination, which the authors refer to as an “oxidative window” for germination; below this amount, dormancy is maintained, and above it, seeds will begin to deteriorate. This is closely related to the resistance phase of stress cited by KRANNER et al. (2010).

SOD plays a key role in the antioxidant defence system because it converts •O$_2^-$ into H$_2$O$_2$ and water, thereby affording protection against the deleterious effects of the superoxide radical (•O$_2^-$) (MITTLER et al., 2004; GILL et al., 2012). In the present study, SOD activity increased during storage with high seed moisture and with the removal of

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**Table 4** - Superoxide dismutase activity (SOD) in spikelets or caryopses of *U. humidicola* cv. Llanero seeds during 12 months in two conditions of storage.

| Seed treatment | Storage (months) | SOD units g fresh weight$^{-1}$ |
|----------------|------------------|-------------------------------|
| Spikelet 4.5% RH | 0                | 13.638±0.5 aA                 |
|                | 3                | 8.618±0.6 bB                  |
|                | 6                | 14.714±0.7 aA                 |
|                | 12               | 8.593±0.5 cB                  |
| Spikelet 50% RH | 0                | 13.638±0.5 aA                 |
|                | 3                | 9.466±1.7 bB                  |
|                | 6                | 8.464±0.4 cB                  |
|                | 12               | 8.623±0.9 cB                  |
| Caryopses 4.5% RH | 0            | 7.025±0.2 bC                   |
|                | 3                | 19.459±0.6 aA                 |
|                | 6                | 15.070±0.3 aB                  |
|                | 12               | 19.655±1.3 aA                  |
| Caryopses 50% RH | 0            | 7.025±0.2 bC                   |
|                | 3                | 17.851±1.5 aA                 |
|                | 6                | 11.040±0.3 bB                  |
|                | 12               | 16.471±0.7 bA                  |

Means followed by equal letters, lowercase in the column and upper case in the row, do not differ by Scott-Knott test (P<0.05).

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**Table 5** - Guaiacol peroxidase (PRX) activity in spikelets or caryopses of *U. humidicola* cv. Llanero seeds during 12 months in two conditions of storage.

| Seed treatment | Storage (months) | PRX (A470nm min$^{-1}$ mg$^{-1}$ fresh weight) |
|----------------|------------------|-----------------------------------------------|
| Spikelet 4.5% RH | 0            | 0.874±0.09 bA                               |
|                | 3                | 0.937±0.04 aA                              |
|                | 6                | 0.226±0.01 aB                              |
|                | 12               | 0.208±0.01 aB                              |
| Spikelet 50% RH | 0            | 0.874±0.09 bA                               |
|                | 3                | 0.938±0.07 aA                              |
|                | 6                | 0.230±0.02 aB                              |
|                | 12               | 0.232±0.02 aB                              |
| Caryopses 4.5% RH | 0            | 0.673±0.04 aB                               |
|                | 3                | 1.005±0.03 aA                              |
|                | 6                | 0.213±0.02 aC                              |
|                | 12               | 0.240±0.02 aC                              |
| Caryopses 50% RH | 0            | 0.673±0.04 aB                               |
|                | 3                | 0.964±0.05 aA                              |
|                | 6                | 0.243±0.01 aC                              |
|                | 12               | 0.224±0.01 aC                              |

Means followed by equal letters, lowercase in the column and upper case in the row, do not differ by Scott-Knott test (P<0.05).
lines overexpressing the 1-Cys PER1 protein from barley (*Hordeum vulgare*) or presenting reduced AtPER1 accumulation because of antisense or RNA interference, there was a correlation between PRX levels and the duration needed to trigger germination, as reported by HASLEKAS et al. (2003). The increase in H₂O₂ content per se is a signal of dormancy release that differs between dormant and non-dormant seeds, and H₂O₂ increases with imbibition (EL-MAAROUF-BOUTEAU et al., 2007).

Conversely, compared with those immersed in water, pea seeds immersed in H₂O₂ for 24 h (BARBA-ESPÍN ET AL. 2012) presented an increase in the percentage of germination and seedling growth, and results were dose dependent; e.g., until a certain concentration, there was positive input, and afterward, a decline in germination was observed, which fits perfectly with the oxidative window (EL-MAAROUF-BOUTEAU & BAILLY 2008). Response of pea seeds to increasing doses of H₂O₂ included an increase in H₂O₂-induced cell death because it activates ABA-degrading enzymes (BARBA-ESPÍN et al. 2010, 2011), which are involved in signalling, plant development, elongation and cell division, as well as in the activity of APX, POX and ascorbate oxidase (AAO), reducing the levels of ABA and cytokinin (BARBA-ESPÍN et al., 2010).

Storage in an environment of relatively high moisture (50% RH) increased the dormancy release
in *U. humidicola* cv. Llanero seeds, and this method is safer than scarification with H₂SO₄ and better than mechanical scarification, for which germinability starts to decrease after nine months. Conversely, storage of very dry seeds maintains dormancy and quality. H₂O₂ determination may help to decide what storage time is adequate to remove and sell seeds.

**CONCLUSION**

It was demonstrated, for the first time, that dormancy release, in *U. humidicola* cv. Llanero, during storage was caused mainly by the accumulation of ROS in the spikelets even under ultra-dry storage conditions (4.5% RH), and this process could be accelerated by seed storage at an MC of 0.1 g H₂O g⁻¹ DW at 20 °C.

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**DECLARATION OF CONFLICTS OF INTERESTS**

The authors declare no conflict of interest. The founding sponsors had no role in the design of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript, and in the decision to publish the results.

**AUTHORS’ CONTRIBUTIONS**

All authors contributed equally for the conception and writing of the manuscript. All authors critically revised the manuscript and approved the final version.

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