Triticale and barley microspore embryogenesis induction requires both reactive oxygen species generation and efficient system of antioxidative defence

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Abstract
The effectiveness of microspore embryogenesis (ME) is determined by a complex network of internal and environmental factors. In the present study on triticale and barley, strong positive correlation (r = 0.85) between the generation of hydrogen peroxide (H2O2) and ME effectiveness confirmed the important role of reactive oxygen species in microspore reprogramming. However, for high effectiveness of ME induction, intensive H2O2 generation had to be associated with high activity of antioxidative enzymes, superoxide dismutase and catalase. The strong seasonal effect on the physiological status of microspores revealed in the study suggests a kind of ‘biological clock’ controlling plant reproduction, crucial for microspore viability and embryogenic potential. Although the effect of various modifications of ME-inducing stress tiller pre-treatment was determined mainly by the physiological condition of microspores, at higher stress intensity positive effects induced by antioxidant molecules—reduced glutathione and its precursor, 1,2-oxothiazolidine-4-carboxylic acid—were observed. High level of variation in the response to ME-inducing stress tiller pre-treatment was also revealed between the two DH lines of triticale and two cultivars of barley and among microspores isolated from subsequently developed spikes.

Key message
Microspore reprogramming towards embryogenic development is initiated by the genotype-specific threshold of reactive oxygen species accumulation. However, high effectiveness of the process requires efficient system of antioxidative defence.

Keywords Antioxidants · Microspore embryogenesis · Oxidative stress · Reactive oxygen species · Hordeum vulgare · ×Triticosecale Wittm

Introduction
The population of isolated and in vitro cultured microspores represents a unique, highly advantageous model for studying the mechanisms which determine the phenomenon of plant cell totipotency, dedifferentiation and reprogramming towards embryogenic development (Testillano 2019). It gives the possibility to monitor the process of embryogenesis continuously, right from the beginning, in a big population of homogeneous cells and to modify a number of environmental and endogenous factors which regulate this process. Several decades of research revealed some important facts but the precise sequence of events and all factors involved in the efficient initiation of microspore embryogenesis (ME) have not been definitely identified yet. The
progress is highly anticipated as better understanding of molecular and physiological background of ME has great breeding importance (Dwivedi et al. 2015), providing the possibility for the development of doubled haploid (DH) procedures for recalcitrant genotypes and its wider incorporation into breeding programmes.

Recent interest focused on the role of redox control in plant growth and development suggests that the signal triggering microspore reprogramming is associated with the generation of reactive oxygen species (ROS). For many years, ROS were mainly considered to be undesirable by-products of aerobic metabolism able to induce serious oxidative damage of cellular components and disrupt metabolic processes, ultimately leading to cell death. However, over the last two decades the research has revealed that starting from fertilization and the first cell division, through shoot, root and flower bud formation up to senescence—all developmental phases are regulated by changes in redox status at organ, tissue, cellular and subcellular levels (Kocsy et al. 2013; Considine and Foyer 2014). Moreover, ROS play a signalling role, informing plants about stress and initiating many defence reactions (Mittler 2017; Schmitt et al. 2014, Noctor et al. 2018; Mhamdi and van Breusegem 2018). Among ROS, hydrogen peroxide (H₂O₂) is particularly interesting, as its ability to diffuse through cell membranes predisposes it to trigger chemical reactions and affect responsive targets, such as metabolites or proteins, inside and outside organelles of its origin. However, other ROS species such as singlet oxygen (¹O₂) and superoxide anion (O₂−) cannot be excluded from the group of potential candidates to play the role of signalling molecules involved in the induction of stress response mechanisms (Schmitt et al. 2014; Dietz et al. 2016; Noctor et al. 2018, Mhamdi and van Breusegem 2018).

ROS generation associated with the process of embryogenesis in in vitro system was first reported by Earnshaw and Johnson (1985, 1987), who examined the effects of antioxidants on the somatic embryo development of wild carrot. The continuation of their research resulted in the conclusion that a reduced environment stimulates cells divisions, whereas a more oxidized environment is necessary for embryo formation and patterning (reviewed in Yeung et al. 2005; Stasolla et al. 2004, 2008).

As previously mentioned, at first ROS generation associated with the initiation of ME was treated as a negative (cytotoxic) but inevitable side-effect of stress treatment. Such approach was in agreement with the observed upregulation of genes and the increase in the activity of antioxidative enzymes—i.e. superoxide dismutase (SOD) and/or catalase (CAT)—as well as the accumulation of low molecular weight antioxidants detected in embryogenic microsperms (Vrinten et al. 1999; Maraschin et al. 2006; Muñoz-Amatriain et al. 2006; Jacquard et al. 2009; Elhiti et al. 2012; Uváčková et al. 2012). Both the above-mentioned enzymes—SOD, which catalyse the dismutation of O²− to H₂O₂, and CAT, which subsequently detoxifies H₂O₂ to O₂ and H₂O—help maintain the redox balance and play a crucial role in cell defence against oxidative stress. Many studies also confirmed that the effectiveness of ME was significantly increased through the application of exogenous ROS-scavenging molecules such as reduced glutathione (GSH), reduced ascorbate (Asc), proline, tocopherol or dimethyl tyrosine (DMT) conjugated peptide (Belmonte et al. 2003; Stasolla et al. 2004; Asif et al. 2013; Hoseini et al. 2014; Sinha and Eudes 2015; Zeng et al. 2017).

The new idea assuming a direct role of ROS in ME initiation can explain why microspore reprogramming can be induced by various stress factors (for example low or high temperature, starvation, osmotic or hormonal shock) as ROS generation is associated with any disturbance in cell metabolism (Mittler 2002). Dual, signalling and toxic effects of ROS overproduction are in agreement with the observations that higher stress intensity increases the effectiveness of ME initiation, but at the same time can reduce cell viability and green plant regeneration ability (Rodriguez-Serrano et al. 2012; Shariatpanahi et al. 2006). Our previous studies with the use of anther and isolated microspore cultures of triticale (Żur et al. 2008, 2009, 2012, 2014, 2019) revealed that various stress factors used for ME initiation enhanced ROS generation and induced genotype-specific changes in the activity of the antioxidative system, which could be associated with the embryogenic potential of the examined cultivar/DH line (Żur et al. 2008, 2009, 2014). We also postulated a dual role of GSH in isolated microspore cultures of triticale, which on the one hand provides direct antioxidative defence and on the other hand, creates a redox status regulating cell proliferation during early embryo-like structure (ELS) development (Żur et al. 2019).

In the present study, we continued to analyse the role of the antioxidative system in the process of microspore reprogramming towards the embryogenic pathway. Our usual model of study—DH lines of winter triticale—was supplemented with two cultivars of barley: cv. Igri combining high embryogenic potential with high green plant regeneration efficiency (Larsen et al. 1991) and cv. Golden Promise, recalcitrant to ME under conventional low temperature treatment (Coronado et al. 2005; Lippmann et al. 2015).

The primary objective of our research was to identify common and specific physiological phenomena associated with ME induction in relation to genetic (different crop species) and environmental (various ME-inducing treatments) variation. The effect of standardly used low temperature tiller pre-treatment (3 weeks at 4 °C; LT) was modified with: (1) GSH, (2) GSH precursor (l-2-oxothiazolidine-4-carboxylic acid, OTC) and (3) buthionine sulfoximine (BSO)—irreversible inhibitor of γ-glutamylcysteine synthetase, the
rate-limiting enzyme for GSH de novo synthesis. We also applied two inhibitors of antioxidative enzymes: diethyl-dithiocarbamate trihydrate (DDC)—Cu²⁺ chelating agent which inhibits SOD activity, and 3-amino-1,2,4-triazole (3-AT) which inhibits CAT activity binding covalently to the heme-containing active centre. All of the used chemicals were analysed with respect to microspore viability, the effectiveness of ME induction and cellular redox status regulated by the balance between ROS generation and antioxidative defence.

**Material and methods**

**Plant material and growth conditions**

The research was done into two DH lines of winter triticale (× *Triticosecale* Wittm.) derived from the F₁ generation of a cross between German inbred line Saka 3006 and Polish cv. Modus (responsive DH28 and recalcitrant DH19) and two barley (*Hordeum vulgare* L.) cultivars (responsive winter cv. Igri and recalcitrant spring cv. Golden Promise). The seeds of DH lines of triticale were obtained from the State Breeding Institute at the University of Hohenheim (Germany), whereas the seeds of barley cultivars were obtained from Leibniz Institute of Plant Genetics and Crop Plant Research (Gatersleben, Germany).

Kernels of triticale and winter barley cv. Igri were germinated for 2 days at 20 °C in perlite pre-soaked with Hoagland’s salt solution (HS; according to Wędzony 2003) and vernalized for 7 weeks at 4 °C and 8 h/16 h (day/night) photoperiod. Kernels of spring cv. Golden Promise were germinated under the same conditions and kept at 4 °C for 2 weeks. Cold-treated seedlings were planted into pots containing a mixture of soil and sand (3/1; v/v), and grown in a glasshouse at 20 °C and 16 h/8 h photoperiod. Additionalillumination with the irradiance of 400 μmol m⁻² s⁻¹ was supplied by high pressure sodium (HPS) lamps SON-T + AGRO (Philips, Brussels, Belgium) during unfavorable growth conditions.

A series of four experiments were carried out over the course of two years. The experiments were conducted over different growing seasons, with microspore isolation starting in May/July (Exp. 1), November (Exp. 2), April (in the case of cv. Igri prolonged to June; Exp. 3), and March (Exp. 4).

**Tiller pre-treatment**

Tillers were harvested at the phase when the majority of microspores were at mid- to late uni-nucleate stage of development, placed in HS medium and stored for 3 weeks at 4 °C in the dark. In Exps. 1 and 3, 4 days before microspore isolation, a part of tillers was transferred to fresh HS medium supplemented with 0.3 mmol dm⁻³ GSH (LT + GSH), 0.3 mol dm⁻³ mannitol (LT + MAN) or both GSH and mannitol (LT + MAN + GSH). The control tillers were transferred to fresh HS medium (LT).

In Exp. 2, the scheme of the tiller pre-treatment was similar, but HS medium was supplemented with: (1) diethyl-dithiocarbamate trihydrate (Cat. No. 228680, Sigma-Aldrich) (LT + DDC); (2) 3-amino-1,2,4-triazole (Cat. No. A8056, Sigma-Aldrich) (LT + 3-AT), (3) buthionine sulfoximine (Cat. No. B2515, Sigma-Aldrich) (LT + BSO), and (4) 1-2-oxothiazolidine-4-carboxylic acid (Cat. No. O6254, Sigma-Aldrich) (LT + OTC). All the chemicals were applied at 0.25 mmol dm⁻³ concentration selected based on the literature data. The control tillers were transferred to fresh HS medium (LT).

In Exp. 4, the effects of two substances which produced the most interesting results in Exp. 3—OTC and DDC—were analysed again with the use of the same experimental scheme.

**Microspore isolation and culture**

Depending on plant growing season and plant species, 20–40 and 20–80 spikes were used for one isolation procedure for triticale and barley, respectively.

The procedure of triticale microspore isolation was described in detail in Żur et al. (2019). The isolation of barley microspores was performed according to the same method with the exception of the concentration of MAN, which was increased to 0.4 mol dm⁻³. The total number of collected microspores was estimated using a Neubauer counting chamber, then the suspension was sampled for in vitro culture and biochemical analysis. Induction media 190-2 (Zhuang and Xu 1983) modified according to Pauk et al. (2003) and KBP (Kumlehn et al. 2006) were used for isolated microspore cultures of triticale and barley, respectively.

Microspore suspensions with the final density of approximately 7×10⁴ microspores per ml (mcs ml⁻¹) for triticale and 10⁵ mcs ml⁻¹ for barley were transferred to 15×60 mm Petri dishes. Microspores of triticale were co-cultured with isolated ovaries dissected simultaneously with microspores (10 per 1 ml of suspension). All cultures were incubated in the dark at 26 °C.

Starting from the sixth week of culture, ELS of triticale over 1 mm in size were transferred to 0.6% agar solidified regeneration medium 190-2 (Zhuang and Xu 1983) supplemented with 0.5 mg dm⁻³ kinetin, 0.5 mg dm⁻³ NAA and 30 g dm⁻³ sucrose, pH 6.0.

ELS of barley were transferred onto KBP4P medium for two weeks, then to K4NBT according to the procedure described by Otto et al. (2015). All cultures were incubated
at 26 °C, with 16 h/8 h (day/night) photoperiod in dim light—at about 30 μmol (hv) m⁻² s⁻¹ (PAR) during the first week, then transferred to 80–100 μmol (hv) m⁻² s⁻¹ (PAR).

**Measurements of cytological parameters associated with microspore embryogenesis initiation and further stages of embryogenic development**

The number of isolated microspores received per one spike of the donor plant was considered as ‘the microspore yield'. Microspore viability was determined through fluorochromatic reaction to fluorescein diacetate (FDA; 0.01%; λ<sub>Ex</sub> = 465 nm, λ<sub>Em</sub> = 515 nm, green fluorescence; Heslop-Harrison and Heslop-Harrison 1970). The samples were examined with Nikon Eclipse E600 epifluorescence microscope equipped with Zyla 4.2 (Andor) camera. Photos were taken using NIS-Elements AR 4.00 software. At least 500 microspores from 10 fields of view (magnification ×10) for each individual sample were evaluated. Microspore samples were standardly collected at the end of the isolation procedure. Only in Exp. 2, the samples were collected twice: (1) after the first centrifugation from crude sediment of homogenised spikes, and (2) at the end of the isolation, after the washing procedure separating somatic tissue debris and dead or damaged microspores from viable cells.

Progressive stages of ME were monitored at five time points—on the isolation day (0 day) and on the 8th, 14th, 28th and 42th day of in vitro culture. Observations were made at least for 10 fields of view per one Petri dish using Nikon Eclipse TS100 inverted microscope and processed by DS-Ri1 digital camera and NIS-Elements AR 4.0 software. The number of ELS over 1 mm in size was assessed after 6 weeks of in vitro culture and calculated per one spike of the donor plant (ELS per spike).

**Measurements of antioxidative enzymes activities**

The activities of antioxidative enzymes were determined in freshly isolated microspores after 1 h incubation at 26 °C in the dark. The samples of microspores suspended in the medium were centrifuged for 3 min at 500 rpm at room temperature. Collected pellet was homogenized by vortexing (90 s) with glass grinding balls in 50 mM potassium phosphate buffer (PPB, pH 7.5), supplemented with 1 mM EDTA, in the proportion of 3 ml buffer for 1 g microspores. The homogenate was centrifuged at 12,000 rpm for 3 min. Protein content was determined in the supernatant according to Bradford (1976) using bovine serum albumin (BSA) as the standard. The activity of superoxide dismutase (SOD, EC 1.15.11) was registered at λ = 550 nm using the modified method of McCord and Fridovich (1969), where one unit of SOD activity was expressed as the amount of the enzyme required to cause 50% inhibition of cytochrome c in a coupled system, using xanthine and xanthine oxidase at pH 7.8 and 25 °C in a 1.0 ml reaction volume. Xanthine oxidase was used in the concentration which produced an initial (uninhibited) ΔA 550 nm of 0.025 ± 0.005 min⁻¹. The activity of catalase (CAT, EC 1.11.1.6) was determined at λ = 240 nm according to Aebi (1984). The reaction mixture contained 50 mM PPB buffer (pH 7.0), 0.1 mM EDTA, 0.04% (v/v) H₂O₂ and 35 μl enzyme extract in 0.7 ml total reaction volume. A unit of enzyme activity (U) was defined as a decrease in absorbance at 25 °C equal to 0.0145 (consumption of 1 μM of H₂O₂).

All measurements were carried out using a ULTRASPEC 2100 PRO (GE Healthcare Life Sciences) UV-Vis spectrophotometer in at least three biological replicates.

**Determination of endogenous H₂O₂ content**

The concentration of H₂O₂ in microspores was measured using the Amplex Red Hydrogen Peroxide/Peroxidase Assay Kit by Invitrogen (Waltham, Massachusetts, USA). After isolation, triticale and barley microspores were incubated in the induction media for 1 h at 26 °C in the dark. Then, microspores were washed with 50 mM Tris HCl with 1 mM EDTA buffer (pH 7.4) and homogenised in the ice-cold phosphate buffered saline (PBS, provided in the kit) at the ratio 1:5 (w/v). The obtained homogenates were centrifuged (10,000 × g, 4 min at 4 °C) and H₂O₂ concentration was measured colorimetrically in the supernatant according to the manufacturer’s protocol using a microplate reader (Synergy 2; BioTek, Winooski, Vermont, USA) under λ<sub>Ex</sub> = 530 nm and λ<sub>Em</sub> = 590 nm. The standard curve in the range 0.05–1 μg dm⁻³ was obtained using different concentrations of H₂O₂ provided in the kit.

**Statistical analysis**

The effect of normally distributed variables was examined by one-factor analysis of variance (ANOVA), after which post-hoc comparison was conducted using Duncan’s multiple range test (p ≤ 0.05). Data expressed as percentages were first transformed using arcsine transformation. Pearson correlation coefficients (r) were used to analyze associations between variables. All statistical analyses were performed using STATISTICA version 10.0 package (StatSoft, USA, 2011).
Results

The influence of genotype and environmental factors on the microspore yield, viability and the effectiveness of ME induced by low-temperature (control) tiller pre-treatment

Significant variation among the studied plant materials with respect to their embryogenic potential was fully confirmed in the conducted experiments. Several parameters given below distinctly differentiated the isolated microspores—both common and distinct for the examined plant species.

The first visible difference among the studied plant materials was the yield of isolated microspores received after the standardly used LT tiller pre-treatment (21 days at 4 °C, control), calculated as the mean number of microspores isolated per individual spike. Comparing the two studied cereal species, triticale was characterized by a significantly higher microspore yield (Tables 1 and 2) with the mean number of ca. 111,000 microspores per individual spike in comparison with 18,000 characteristic for barley. No significant difference was detected between DH lines of triticale, while 1.3–2.6–fold higher number of microspores was received from a spike of cv. Igri in comparison with cv. Golden Promise (Tables 1 and 2).

The viability of cells measured after LT treatment in crude microspore suspensions (estimated in Exp. 2) varied significantly from 24% to 37% (Supplementary Source 1) but was not associated with embryogenic potential. The washing procedure based on density gradient centrifugation resulted in a mean two-fold increase in the number of viable cells in comparison to crude microspore suspensions. Moreover, after the washing procedure a significantly higher microspore viability was detected in the responsive line DH28 and cv. Igri in comparison with the recalcitrant DH19 and cv. Golden Promise.

For the majority of the studied genotypes, the frequency of viable microspores in suspensions isolated from LT pretreated tillers depended significantly on the donor plant’s growing season (Fig. 1). Both winter triticale DH lines

Table 1  Experiment 1 and 3. The effect of the donor plant’s growing season and tillers pre-treatment on microspore yield [microspores per spike] and the effectiveness of ME induction [ELS per spike] in isolated microspore cultures of triticale and barley. Data represent means from 3 to 5 biological replications (isolations) ± SE. Statistical analysis was performed separately for each DH line/cultivar. Data marked with different letters differ significantly according to the Duncan test (p ≤ 0.05); NS—variation not significant

| Plant material | Treatment                  | Experiment 1A |          |          | Experiment 3B |          |          |
|----------------|---------------------------|---------------|----------|----------|---------------|----------|----------|
|                |                           | Microspores per spike [× 10³] ± SE | % Control | ELS per spike [× 10³] ± SE | % Control |          |
|                |                           |               |          |          |               |          |          |
| DH28           | LT (control)              | 116 ± 12NS    | 100      | 32 ± 7b  | 91 ± 10NS     | 100      | 144 ± 31a|
|                | LT + GSH                  | 94 ± 17       | 79       | 123 ± 24a| 72 ± 17       | 79       | 11 ± 2b  |
|                | LT + MAN                  | 87 ± 7        | 70       | 8 ± 3b   | 79 ± 15       | 86       | 9 ± 3b   |
|                | LT + MAN + GSH            | 75 ± 8        | 61       | 14 ± 8b  | 65 ± 20       | 73       | 69 ± 39a|
| DH19           | LT (control)              | 90 ± 22NS     | 100      | 0NS      | 84 ± 19NS     | 100      | 0        |
|                | LT + GSH                  | 76 ± 19       | 104      | 0.5      | 64 ± 7        | 82       | 1        |
|                | LT + MAN                  | 71 ± 9        | 90       | 1.1      | 76 ± 23       | 86       | 0        |
|                | LT + MAN + GSH            | 98 ± 10       | 126      | 310 ± 99b| 26 ± 4NS      | 100      | 1.4 ± 1.4NS/662 ± 188bc |
| cv. Igri       | LT (control)              | 19 ± 2b       | 100      | 310 ± 99bc| 39 ± 20       | 147      | 0/237 ± 93c|
|                | LT + GSH                  | 10 ± 2b       | 57       | 130 ± 49bc| 36 ± 14       | 133      | 2.1 ± 1.3/852 ± 71ab |
|                | LT + MAN                  | 47 ± 16a      | 238      | 22 ± 13c | 25 ± 5        | 97       | 0/1294 ± 199a |
|                | LT + MAN + GSH            | 27 ± 6ab      | 141      | 413 ± 114a| <0.1NS       | 10 ± 3NS | 0.2NS    |
| cv. Golden Promise | LT (control)               | 14 ± 6NS      | 100      | <0.1NS   | 8 ± 1         | 91       | 0        |
|                | LT + GSH                  | 12 ± 5        | 90       | <0.1     | 14 ± 4        | 138      | 1        |
|                | LT + MAN                  | 10 ± 4        | 72       | <0.1     | 15 ± 2        | 183      | 0.3      |
|                | LT + MAN + GSH            | 12 ± 6        | 155      | <0.1     | 14 ± 4        | 138      | 1        |

Microspore isolation started in: May/July (A) or April (B) in the case of cv. Igri prolonged to June that’s why data for cv. Igri represent the effectiveness of ME induction in March/June. ELS—embryo-like structures; LT—low-temperature tiller pre-treatment (21 days at 4 °C); LT + GSH—low-temperature tiller pre-treatment combined with the application of 0.3 mmol dm⁻³ reduced glutathione during the last 4 days before microspore isolation; LT + MAN—low-temperature tiller pre-treatment combined with the application of 0.3 mol dm⁻³ mannitol during the last 4 days before microspore isolation; LT + MAN + GSH—low-temperature tiller pre-treatment combined with the application of reduced 0.3 mmol dm⁻³ glutathione and 0.3 mol dm⁻³ mannitol during the last 4 days before microspore isolation.
turned out to be highly sensitive to the environmental factors. Mean microspore viability decreased significantly from 52% to 69% received in the experiments when isolation procedures started in May/July (Exp. 1) or November (Exp. 2), through 38–42% detected in April, to 15–22% received in March (Exp. 4). In contrast, mean viability of microspores of spring barley (cv. Golden Promise), which varied from 33% to 69%, was the highest in March (Exp. 4) and the lowest in May/July (Exp. 1). Seasonal variation was not detected in microspore viability of cv. Igri, which changed to a minor extent—from 50% to 68%. In addition, variation in microspore viability was found even within each experiment between subsequent isolation procedures. Depending on the growing season and the donor plant’s genotype, the highest number of viable cells was detected in microspore population isolated from the primary tillers, secondary tillers or from tillers developed from next axillary buds (Fig. 1).

Characteristic for cv. Golden Promise, in the majority of experiments, was a time-sensitive decrease in microspore viability. To eliminate the variation originating from the effect of tiller order, microspore viability and the yield of isolated microspores were also presented as the percentage of the control (Tables 1 and 2; Figs. 2 and 4).

Significant variation was also observed in the response of the studied cultivars/DH lines to the standard ME-inducing treatments (control, Tables 1 and 2). In Exp. 1, where the isolation of microspores started in May/July, the effectiveness of ME induction was highest for cv. Igri (>300 ELS per spike), moderate for DH28 (ca. 30 ELS per spike) and very low for both recalcitrant genotypes: DH19 and cv. Golden Promise (<1 ELS per spike). However, ME effectiveness was significantly influenced by the donor plant’s growing season.
and varied significantly among the conducted experiments (Tables 1 and 2). This effect was the most pronounced in the case of cv. Igri as its huge embryogenic potential observed when the isolation of microspores started in May/July was not revealed in March/April and November (Exps. 2, 3 and 4). It was clearly visible in Exp. 3 where the cultivation of cv. Igri was prolonged to June, and the effectiveness of ME induction varied drastically from about 1 to over 600 ELS per spike in April and June, respectively (Table 1). Similarly in the responsive triticale line DH28, the effectiveness of ME initiation varied in all conducted experiments from 0 to 144 ELS per spike (Tables 1 and 2).

The influence of stress on the microspore yield, viability and the effectiveness of ME induction (Exps. 1 and 3)

The effects induced by various stress treatments were genotype-specific. Generally, modifications introduced in Exp. 1 had no effect on the microspore yield (Table 1). The only exception was LT + MAN treatment applied to tillers of cv. Igri, which increased the number of isolated microspores, but at the same time significantly decreased their viability (Figs. 1 and 2) and dramatically diminished the number of produced ELS (numerical data in Table 1; exemplary pictures in Fig. 3). No other variation was observed in both responsive genotypes (cv. Igri, DH28) with respect to microspore viability. In both recalcitrant genotypes, microspore viability was enhanced when LT was combined with the application of exogenous GSH, alone or concomitantly with MAN (Fig. 2). Only in DH19, also LT + MAN significantly increased the frequency of viable microspores.

None of the treatments effectively induced the process of ME in isolated microspore cultures of highly recalcitrant DH19 and cv. Golden Promise (Table 1). At the same time, the only one significant effect in microspore cultures of DH28 was induced by exogenously applied GSH resulting in almost four times higher production of ELS. This effect was not observed in isolated microspore cultures of cv. Igri. However in this case, application of GSH concomitantly with MAN reversed the dramatic decrease in the effectiveness of ELS production observed after MAN treatment (Table 1, Fig. 3).

Interestingly, the effects of the same treatments applied in the second replication of the experiment (Exp. 3) were not completely reproducible (Table 1). The received results confirmed strong genotype control of the microspore yield, not influenced by any of the applied modifications. Again, this parameter was significantly higher in triticale in comparison with both barley cultivars (Table 1). However, positive effects of LT + GSH and LT + GS + MAN on microspore viability of the recalcitrant genotypes (DH19 and cv. Golden Promise) were not observed (Fig. 1). On the contrary, these treatments significantly diminished (by about 20% in comparison with the control) the number of viable cells in DH19 while only LT + MAN + GSH enhanced significantly (by about 70%) cell viability of cv. Golden Promise. Surprisingly, this had no effect on the effectiveness of ME induction. The strong negative effect of LT + MAN on microspore viability of cv. Igri and on the effectiveness of ME initiation observed in Exp. 1 was not confirmed either (Table 1, Fig. 1). This time, the negative effect of LT + GSH and LT + MAN application on ELS production was observed in microspore cultures of DH28 and similarly it was, at least partially, reversed by combined tiller pre-treatment (LT + MAN + GSH; Table 1).

The influence of disturbances in antioxidative defense system on the microspore yield, viability and the effectiveness of ME induction (Exps. 2 and 4)

The effects of two inhibitors of antioxidative enzymes (DDC, 3-AT) and two modulators of endogenous glutathione biosynthesis (BSO, OTC) were examined in Exp. 2, with microspore isolation procedures starting in November. Once again, the applied modifications did not induce significant changes in the microspore yield (Table 2). The only exception was the treatment with BSO, which increased by around 70% the number of isolated microspores per spike of the responsive DH28. Interestingly, although at the same time BSO diminished the number of viable cells estimated after the washing procedure (numerical data in Fig. 1; exemplary pictures in Fig. 4), it did not reduce the effectiveness of ELS formation (Table 2). Generally, the viability of microspores after LT tiller pre-treatment was relatively high, ranging from 55% to 69% of total microspore populations (Fig. 1). All tested chemicals induced genotype-specific and usually negative effects on microspore viability (Figs. 1 and 4). The exception was cv. Igri, for which tiller pre-treatments with BSO, OTC and 3-AT significantly increased the number of viable cells by about 18-33% in comparison with the control (LT).

The effectiveness of ME induction acquired in Exp. 2 is presented in numerical form in Table 2 and exemplary pictures are shown in Fig. 5. For control (LT) tiller pre-treatment, the number of produced ELSs was similar (for cv. Golden Promise) or even higher (for DH28 and DH19) when compared to earlier results in Exp. 1. At the same time the standardly high embryogenic potential of cv. Igri was not expressed. Among various tested chemicals, DDC had no significant effect on triticale ELS formation, although some positive effects seem to be induced in the case of DH19 (Table 2). The same treatment completely inhibited ELS production in both barley cultivars. Conversely, treatment with OTC visibly diminished embryogenic potential of DH28 but increased ELS production in isolated microspore cultures of...
| Genotype       | Treatment | Viability [% ± Se] | Experiment 1 | Viability [% ± Se] | Experiment 3 |
|---------------|-----------|-------------------|--------------|-------------------|--------------|
|               |           |                   | Biological replication |                   | Biological replication |
|               |           |                   | 1            | 2    | 3    | 1            | 2    | 3    |
| DH28          | LT        | 69 ± 3 NS          |              |                   |              |
|               | LT+GSH    | 62 ± 3             |              |                   |              |
|               | LT+MAN    | 64 ± 5             |              |                   |              |
|               | LT+MAN+GSH| 70 ± 3             |              |                   |              |
| DH19          | LT        | 52 ± 3 b           |              |                   |              |
|               | LT+GSH    | 61 ± 3 a           |              |                   |              |
|               | LT+MAN    | 63 ± 2 a           |              |                   |              |
|               | LT+MAN+GSH| 65 ± 3 a           |              |                   |              |
| cv. Isri      | LT        | 55 ± 6 a           |              |                   |              |
|               | LT+GSH    | 63 ± 6 a           |              |                   |              |
|               | LT+MAN    | 43 ± 5 b           |              |                   |              |
|               | LT+MAN+GSH| 48 ± 4 ab          |              |                   |              |
| cv. Golden Promise | LT    | 33 ± 21 ab         |              |                   |              |
|               | LT+GSH    | 67 ± 4 a           |              |                   |              |
|               | LT+MAN    | 13 ± 13 b          |              |                   |              |
|               | LT+MAN+GSH| 65 ± 20 a          |              |                   |              |

| Genotype       | Treatment | Viability [% ± Se] | Experiment 2 | Viability [% ± Se] | Experiment 4 |
|---------------|-----------|-------------------|--------------|-------------------|--------------|
|               |           |                   | Biological replication |                   | Biological replication |
|               |           |                   | 1            | 2    | 3    | 1            | 2    | 3    |
| DH28          | LT        | 69 ± 4 a           |              |                   |              |
|               | LT+GSH    | 55 ± 2 b           |              |                   |              |
|               | LT+2AT    | 53 ± 3 b           |              |                   |              |
|               | LT+BSO    | 41 ± 3 c           |              |                   |              |
|               | LT+OTC    | 55 ± 3 b           |              |                   |              |
| DH19          | LT        | 55 ± 2 a           |              |                   |              |
|               | LT+GSH    | 48 ± 3 a c         |              |                   |              |
|               | LT+2AT    | 41 ± 3 c           |              |                   |              |
|               | LT+BSO    | 52 ± 3 ab          |              |                   |              |
|               | LT+OTC    | 46 ± 4 b           |              |                   |              |
| cv. Isri      | LT        | 63 ± 4 c           |              |                   |              |
|               | LT+GSH    | 64 ± 3 bc          |              |                   |              |
|               | LT+2AT    | 78 ± 3 a           |              |                   |              |
|               | LT+BSO    | 71 ± 2 ab          |              |                   |              |
|               | LT+OTC    | 74 ± 2 a           |              |                   |              |
| cv. Golden Promise | LT    | 59 ± 5 a           |              |                   |              |
|               | LT+GSH    | 41 ± 5 b           |              |                   |              |
|               | LT+2AT    | 42 ± 3 b           |              |                   |              |
|               | LT+BSO    | 56 ± 4 a           |              |                   |              |
|               | LT+OTC    | 52 ± 3 ab          |              |                   |              |

| Genotype       | Treatment | Viability [% ± Se] | Experiment 3 | Viability [% ± Se] | Experiment 4 |
|---------------|-----------|-------------------|--------------|-------------------|--------------|
|               |           |                   | Biological replication |                   | Biological replication |
|               |           |                   | 1            | 2    | 3    | 1            | 2    | 3    |
| DH28          | LT        | 38 ± 2 a           |              |                   |              |
|               | LT+GSH    | 37 ± 2 a           |              |                   |              |
|               | LT+MAN    | 33 ± 3 a           |              |                   |              |
|               | LT+MAN+GSH| 27 ± 3 b           |              |                   |              |
| DH19          | LT        | 42 ± 2 a           |              |                   |              |
|               | LT+GSH    | 33 ± 2 b           |              |                   |              |
|               | LT+MAN    | 38 ± 2 a           |              |                   |              |
|               | LT+MAN+GSH| 31 ± 2 b           |              |                   |              |
| cv. Isri      | LT        | 50 ± 4 NS          |              |                   |              |
|               | LT+GSH    | 47 ± 4             |              |                   |              |
|               | LT+MAN    | 44 ± 4             |              |                   |              |
|               | LT+MAN+GSH| 53 ± 4             |              |                   |              |
| cv. Golden Promise | LT    | 41 ± 4 b           |              |                   |              |
|               | LT+GSH    | 52 ± 3 ab          |              |                   |              |
|               | LT+MAN    | 52 ± 4 ab          |              |                   |              |
|               | LT+MAN+GSH| 55 ± 3 a           |              |                   |              |
cv. Golden Promise. The treatments with 3-AT and BSO had no significant effect on ELS production.

Two treatments (LT + DDC, LT + OTC) inducing diverse effects on triticale/barley ME initiation were used again in Exp. 4. Here, the isolation of microspores started in March, viability assay revealed only 15–22% of viable triticale microspores after LT tiller pre-treatment (Fig. 1). At the same time, viability of barley microspores ranged from 54% to 69%. Mean microspore viability for both species varied from 42% observed in control tillers to 31% in DDC- and OTC-treated tillers. For both studied DH lines of triticale, OTC significantly increased the number of viable tillers (1), secondary tillers (2) and tillers developed from next axillary buds (3). Mean viability of the responsive genotypes (DH28, cv. Igri) demonstrated a significantly lower level of H2O2 generation in DH28 microspores was significantly reduced, especially in microspores isolated from LT + GSH pre-treated tillers (Fig. 6c).

No significant difference in CAT activity was observed in comparison with the control (LT) in any studied DH line/cultivar (Fig. 6b).

Regardless of the treatment, consistently low SOD activity was characteristic for isolated microspores of DH19 (18–23 U mg⁻¹ protein; Fig. 6a). It was associated with relatively low level of H2O2 generation, especially in microspores isolated from control tillers (0.1 nmol H2O2 g⁻¹ FW; Fig. 6c), whereas the amount of H2O2 generated in microspores isolated from LT + MAN pre-treated tillers was significantly higher (ca. 10 nmol H2O2 g⁻¹ FW).

Two of the applied treatments—LT + MAN and LT + MAN + GSH—significantly increased SOD activity in microspores of cv. Igri in comparison with the control (LT, Fig. 6a). The effect of LT + MAN was most pronounced, resulting in the highest activity of both antioxidative enzymes and the highest level of H2O2 generation (Fig. 6a–c). Interestingly, although combined LT + MAN + GSH treatment also activated SOD, the amount of generated H2O2 diminished significantly.

All modifications reduced SOD activity in microspores of cv. Golden Promise (Fig. 6a). However, this effect did not correlate with H2O2 generation as both higher (by 48% under LT + MAN + GSH treatment) and lower (by 87% under LT + GSH treatment) amount of H2O2 was detected in comparison with the control (Fig. 6a, c).

Analyses conducted in Exp. 4 revealed that microspores of the responsive genotypes (DH28, cv. Igri) demonstrated significantly lower activity of antioxidative enzymes in comparison with the previous experiment (Fig. 6a, b, d, e). Interestingly, this time, a much higher activity of SOD was detected in microspores of the recalcitrant cv. Golden Promise, which, was not associated with a higher activity of CAT (Fig. 6c, d). In both experiments, microspores of DH19 were characterized by a low SOD activity (Fig. 6a, c), whereas a relatively high CAT activity detected in Exp. 3 was not observed this time (Fig. 6b, d). Comparing both...
Fig. 2 Microspore viability of two DH lines of winter triticale (responsive DH28 and recalcitrant DH19) and two cultivars of barley (responsive winter cv. Igri and recalcitrant spring cv. Golden Promise) after low temperature (LT) tillers pre-treatment (control) and LT treatment combined with the application reduced glutathione (GSH) and/or mannitol (MAN). Microspores stained with FDA (fluorescence of vital cells intensely green). Bar = 50 μm. Included are normalized values of microspore viability. Results received in Experiment 1 with the isolation of microspores started in May/July. LT—low-temperature tiller pre-treatment (21 days at 4 °C); LT+GSH—low-temperature tiller pre-treatment combined with the application of 0.3 mmol dm⁻³ reduced glutathione; LT+MAN—low-temperature tiller pre-treatment combined with the application of 0.3 mol dm⁻³ mannitol; LT+MAN+GSH—low-temperature tiller pre-treatment combined with the application of 0.3 mol dm⁻³ mannitol and 0.3 mmol dm⁻³ reduced glutathione. All chemicals were applied during the last 4 days before microspore isolation.
experiments it could be noticed that LT tiller pre-treatment induced lower (by 60%) accumulation of H$_2$O$_2$ in microspores of DH28, but drastically increased its generation in microspores of DH19 (from 0.1 to 53 nmol g$^{-1}$ FW) and cv. Golden Promise (from 15 to 134 nmol g$^{-1}$ FW). Significant variation between the studied genotypes was observed only in barley, again with higher H$_2$O$_2$ accumulation detected in microspores of cv. Golden Promise.

The effect of tiller pre-treatment modifications applied in Exp. 4 was significant only in antioxidative defense of cv. Golden Promise (Fig. 6d, e). Interestingly, both the decrease in SOD activity induced by LT + OTC treatment and the decrease in CAT activity resulting from LT + DDC treatment had no effect on H$_2$O$_2$ generation (Fig. 6f).

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|                | LT (control) | LT+GSH | LT+MAN | LT+MAN+GSH |
|----------------|--------------|--------|--------|------------|
| **DH28**       | ![Image](image1.png) | ![Image](image2.png) | ![Image](image3.png) | ![Image](image4.png) |
| **DH19**       | ![Image](image5.png) | ![Image](image6.png) | ![Image](image7.png) | ![Image](image8.png) |
| **cv. Igri**   | ![Image](image9.png) | ![Image](image10.png) | ![Image](image11.png) | ![Image](image12.png) |
| **cv. Golden Promise** | ![Image](image13.png) | ![Image](image14.png) | ![Image](image15.png) | ![Image](image16.png) |

**Fig. 3** The effect of low temperature (LT, control) tillers pre-treatment and LT treatment combined with the application of reduced glutathione (GSH) and/or mannitol (MAN) on the effectiveness of ME in in vitro cultures of two DH lines of winter triticale (responsive DH28 and recalcitrant DH19) and two cultivars of barley (responsive winter cv. Igri and recalcitrant spring cv. Golden Promise). Results received in Experiment 1 with the isolation of microspores started in May/July. LT—low-temperature tiller pre-treatment (21 days at 4 °C); LT + GSH—low-temperature tiller pre-treatment combined with the application of 0.3 mmol dm$^{-3}$ reduced glutathione; LT + MAN—low-temperature tiller pre-treatment combined with the application of 0.3 mol dm$^{-3}$ mannitol; LT + MAN + GSH—low-temperature tiller pre-treatment combined with the application of 0.3 mol dm$^{-3}$ mannitol and 0.3 mmol dm$^{-3}$ reduced glutathione. All chemicals were applied during the last 4 days before microspore isolation.
Correlation analysis

Data received in Exp. 3 revealed a significant positive correlation between H$_2$O$_2$ generation and ME effectiveness ($r=0.85$, $p \leq 0.05$; Table 3). Moreover, a negative correlation between the microspore yield and cell viability was also detected ($r=-0.75$).

In Exp. 4, the microspore yield was not only strongly negatively correlated with microspore viability ($r=-0.92$) but also with the effectiveness of ME induction ($r=-0.79$). Negative correlation was also found between SOD activity and the microspore yield ($r=-0.74$). At the same time, the activity of this enzyme was positively correlated with the viability of microspores ($r=0.60$) and H$_2$O$_2$ generation ($r=0.81$). Conversely, CAT activity was positively correlated with the microspore yield ($r=0.78$) but negatively with its viability ($r=-0.75$) and ME effectiveness ($r=-0.79$).

|               | LT (control) | LT + DDC | LT + 3-AT | LT + BSO | LT + OTC |
|---------------|--------------|----------|-----------|----------|----------|
| DH28          | 1.00         | 0.81     | 0.78      | 0.60     | 0.81     |
| DH19          | 1.00         | 0.84     | 0.73      | 0.93     | 0.76     |
| cv. Igri      | 1.00         | 1.10     | 1.33      | 1.18     | 1.23     |
| cv. Golden Promise | 1.00 | 0.89 | 0.75 | 1.13 | 0.93 |

Fig. 4 Microspore viability of two DH lines of winter triticale (responsive DH28 and recalcitrant DH19) and two cultivars of barley (responsive winter cv. Igri and recalcitrant spring cv. Golden Promise) after LT tillers pre-treatment (control) and LT treatment combined with the application of chemicals inducing disturbances in the activity of the antioxidative system. Results received in Experiment 2 with the isolation of microspores started in November. Microspores stained with FDA (fluorescence of vital cells intensely green). Bar=50 μm. Included are normalized values of microspore viability.
Discussion

Despite years of research, many aspects of ME regulation on molecular, cytological and physiological levels have yet to be fully explained. It remains unclear why not only various plant species or cultivars, but even closely related plant genotypes differ significantly with respect to their embryogenic potential. Such variation was observed, for example in the triticale mapping population composed of 90 DH lines derived from the cross Saka 3006 × Modus, being the standard model of our studies (Krzewska et al. 2012, 2015). Moreover, this kind of variation was also revealed among individual plants of the same genotype and among subsequent spikes produced by an individual plant. Several authors also reported strong seasonal variation in ME effectiveness (Wang et al. 2000; Ritala et al. 2001; Datta 2005; Jacquard et al. 2006; Ercan et al. 2006; Silva 2010). The answers to these questions would be very interesting but they are not easy to obtain due to the multifactorial nature of the process and the complex network of interactions between genetic and environmental factors controlling and determining its effectiveness. The data presented in this work can help understand why even subtle changes in the physiological condition of the donor plant associated with a disturbed redox balance can have a huge impact on the effectiveness of microspore reprogramming towards embryogenic development.

The two sibling DH lines of winter triticale (DH19, DH28) used in the study were selected as significantly different in ME induction effectiveness based on the results of our previous research (Krzewska et al. 2012, 2015; Żur et al. 2014, 2015). The two cultivars of barley (spring cv. Golden Promise and winter cv. Igri) were selected according to the same criterion based on literature data (Coronado et al. 2005; Lippmann et al. 2015). Our study confirmed significant differences between the studied plant species as well as between individual DH lines/cultivars with respect to several parameters determining ME effectiveness. We also...
Fig. 6  Activity of SOD (a, d) and CAT (b, e), and H$_2$O$_2$ generation (c, f) in microspores of two DH lines of winter triticale (responsive DH28 and recalcitrant DH19) and two cultivars of barley (responsive winter cv. Igri and recalcitrant spring cv. Golden Promise) isolated after various tillers pre-treatments. Data represent means from three biological replications (isolations) ± SE. Data marked with the same letter do not differ according to Duncan's multiple range test ($p \leq 0.05$). Figures a–c represent data received in Experiment 3 with the isolation of microspores started in April/June, figures d–f—data received in Experiment 4 with the isolation of microspores started in March. LT—low temperature tillers pre-treatment (21 days at 4 °C); LT+GSH—low temperature tillers pre-treatment combined with the application of 0.3 mmol dm$^{-3}$ reduced glutathione; LT+MAN—low temperature tillers pre-treatment combined with the application of 0.3 mol dm$^{-3}$ mannitol; LT+MAN+GSH—low temperature tillers pre-treatment combined with the application of 0.3 mol dm$^{-3}$ mannitol and 0.3 mmol dm$^{-3}$ reduced glutathione; LT+DDC—low temperature tillers pre-treatment combined with the application of 0.25 mmol dm$^{-3}$ diethyldithiocarbamate trihydrate; LT+OTC—low temperature tillers pre-treatment combined with the application of 0.25 mmol dm$^{-3}$ L-2-oxothiazolidine-4-carboxylic acid. All chemicals were applied during the last 4 days before microspore isolation.
Table 3 Pearson correlation (r) matrix between parameters describing the effectiveness of ME initiation: microspore yield (mY), microspore viability (mV), the effectiveness of microspore embryogenesis (ME), hydrogen peroxide (H$_2$O$_2$) generation and the activity of antioxidative enzymes: superoxide anion (SOD) and catalase (CAT)

| Parameters | mY | mV | H$_2$O$_2$ | SOD | CAT | ME |
|------------|----|----|-----------|-----|-----|----|
| mY         | –  | $-0.922$ | $-0.319$ | $-0.742$ | $0.780$ | $-0.612$ |
| mV         | $-0.751$ | –  | $0.206$  | $0.603$  | $-0.753$ | 0.514  |
| H$_2$O$_2$ | 0.204 | –  | –        | $0.806$  | 0.047  | 0.152  |
| SOD        | 0.094 | –  | $0.286$  | –        | $-0.289$ | 0.279  |
| CAT        | 0.212 | 0.064 | $0.091$  | $-0.563$ | –      | $-0.787$ |
| ME         | 0.391 | $-0.330$ | **0.852** | 0.103    | $-0.068$ | –      |

Data from Experiment 3 is presented in plain font, data from Experiment 4—in italic font. Significant correlations with $p \leq 0.05$ are indicated in bold.

observed a substantial effect of the donor plant’s growing season, tiller order and stress tiller pre-treatment on microspore viability, effectiveness of microspore reprogramming and the activity of the antioxidative system.

The first tested parameter—the microspore yield, dependent on microspore productivity and their condition—was determined mainly genetically and remained almost unchanged across different growing seasons and tiller pre-treatments. Although both studied plant species are predominantly self-pollinating, higher microspore productivity in triticale is probably inherited from rye, its cross-pollinated parent (Nguyen et al. 2015). Rye anthers are relatively long and produce around 19,000 pollen grains per anther, which is roughly ten times higher in comparison with wheat pollen production. Surprisingly, this parameter was not only uncorrelated with ME effectiveness but also negatively correlated with microspore viability measured through microspore enzymatic activity and cell membrane integrity (FDA assay). This result, obtained also in our earlier study (Zur et al. 2019), suggests that the washing procedure based on cell density gradient centrifugation is not so effective in the separation of fully viable microspores. On the other hand, a very low microspore yield revealed for cv. Golden Promise associated with middling or low cell viability makes it difficult to establish isolated microspore cultures. It is also important to note that only for this cultivar the quality of the produced microspores declined visibly with the tillers developed from the next axillary buds. A similar observation on barley was described by Wang et al. (2000) and Jacquard et al. (2006), who recommended the first three-five spikes as the source of donor plant material. Observations made on winter wheat showed that the timing of tiller initiation affected leaf development and influenced tiller size, the number of kernels per spike and their contribution to the final yield (Tilley et al. 2019).

For the majority of the studied DH lines/cultivars (with the exception of cv. Igri), the viability of microspores was strongly influenced by the donor plant’s growing season. Equally high viability of winter triticale microspores was received when microspore isolation started in May/July and November, with strong reduction in cell viability observed in March. In contrast, microspore viability of spring cv. Golden Promise declined from March through November to July. It should be underlined that decreased microspore viability did not result from weakened physiological condition of donor plants as no morphological symptoms of any (a)biotic stress were observed.

This seasonal effect was observed not only with respect to microspore viability but also their embryogenic potential. The most effective ME induction took place in November for both triticale DH lines and spring barley or in June in the case of winter barley. The observed phenomenon is probably determined by a kind of biological clock regulating the natural rhythm of plant development. This type of dependency has often been reported by authors using in vitro cultures as models for their study (e.g. Ritala et al. 2001; Jacquard et al. 2006; Ercan et al. 2006; Silva 2010; Ari et al. 2016). In this case, with plants growing in glass chambers illuminated additionally by HPS lamps SON-T + AGRO (Philips), the signalling system may be at least partially, based on light as its integration with other environmental stimuli determines plant adaptation to the changing environment. Using several photoreceptors (phytochromes, cryptochromes, phototropins, UV-A and UV-B detecting receptors), plants can not only measure the quantity of light but also react to subtle changes in its quality, direction and periodicity (Franklin et al. 2014; Fernando and Schroeder 2016). All these parameters changed significantly between different growth seasons, as did the ratio between the quantity of natural and artificial light. Published data confirmed that light intensity and spectrum regulated photosynthetic activity, accumulation of amino acids and antioxidants (phenolic compounds, ascorbic acid and glutathione) what significantly influenced on plant growth and development (Lee et al. 2010; Toldi et al. 2019).

Regardless of the source and perception of the environmental stimulus, its transduction regulating plant cell fate and the direction of its development is based on ROS.
Fig. 7 Summary. (a) The effect of low temperature (LT) tillers pre-treatment on reactive oxygen species (ROS) generation, the activity of the antioxidative system and the effectiveness of ME and its modulation induced by season × genotype interaction. (I) Depending on the physiological condition of donor plants, LT tillers pre-treatment induces ROS accumulation, which activates (responsive DH/cv.) or deactivates (recalcitrant DH/cv.) the antioxidative defence system. (II) Depending on the physiological condition of donor plants, isolation of microspores associated with excessive ROS generation results in: (A) high activity of superoxide dismutase (SOD) and catalase (CAT), intensive or moderate generation of hydrogen peroxide (H$_2$O$_2$) and effective induction of microspore embryogenesis (ME); (B) low activity of SOD, low generation of H$_2$O$_2$ and high frequency of viable microspores which continue gametophytic development; (C) low activity of SOD and CAT, moderate generation of H$_2$O$_2$, and high frequency of dead or damaged microspores. However, high activity of SOD and intensive generation of H$_2$O$_2$ are not the only prerequisites for effective ME induction, which could be seen on the example of cv. Golden Promise (D). V—microspore viability [%] determined by FDA assay; ME—effectiveness of ME expressed as the number of ELS per spike. A—embryogenic cultures of isolated microspores characterized by high frequency of star-like structures (triticale) and symmetrically divided microspores (barley). B—non-embryogenic cultures of isolated microspores characterized by high frequency of asymmetrically divided microspores. C—non-embryogenic cultures of isolated microspores characterized by high frequency of plasmolyzed, shrunk microspores. Data are presented in colour according to an arbitrary adopted scale, with colour intensity corresponding to the number of ELS developed per spike of the donor plant. Asterisk marks values statistically different from control (LT). GP—Golden Promise; LT—low-temperature tiller pre-treatment (21 days at 4 °C); LT+GSH—low-temperature tiller pre-treatment combined with the application of 0.3 mmol dm$^{-3}$ reduced glutathione; LT+MAN—low-temperature tiller pre-treatment combined with the application of 0.25 mmol dm$^{-3}$ mannitol; LT+MAN+GSH—low-temperature tiller pre-treatment combined with the application of 0.25 mmol dm$^{-3}$ diethyldithiocarbamate trihydrate; LT+OTC—low temperature tillers pre-treatment combined with the application of 0.25 mmol dm$^{-3}$ 1,2-oxothiazolidine-4-carboxylic acid. All chemicals were applied during the last 4 days before microspore isolation.

Fig. 7. Summary. 

The effect of low temperature (LT) tillers pre-treatment on reactive oxygen species (ROS) generation, the activity of the antioxidative system and the effectiveness of ME and its modulation induced by season × genotype interaction. (I) Depending on the physiological condition of donor plants, LT tillers pre-treatment induces ROS accumulation, which activates (responsive DH/cv.) or deactivates (recalcitrant DH/cv.) the antioxidative defence system. (II) Depending on the physiological condition of donor plants, isolation of microspores associated with excessive ROS generation results in: (A) high activity of superoxide dismutase (SOD) and catalase (CAT), intensive or moderate generation of hydrogen peroxide (H$_2$O$_2$) and effective induction of microspore embryogenesis (ME); (B) low activity of SOD, low generation of H$_2$O$_2$ and high frequency of viable microspores which continue gametophytic development; (C) low activity of SOD and CAT, moderate generation of H$_2$O$_2$, and high frequency of dead or damaged microspores. However, high activity of SOD and intensive generation of H$_2$O$_2$ are not the only prerequisites for effective ME induction, which could be seen on the example of cv. Golden Promise (D). V—microspore viability [%] determined by FDA assay; ME—effectiveness of ME expressed as the number of ELS per spike. A—embryogenic cultures of isolated microspores characterized by high frequency of star-like structures (triticale) and symmetrically divided microspores (barley). B—non-embryogenic cultures of isolated microspores characterized by high frequency of asymmetrically divided microspores. C—non-embryogenic cultures of isolated microspores characterized by high frequency of plasmolyzed, shrunk microspores. Data are presented in colour according to an arbitrary adopted scale, with colour intensity corresponding to the number of ELS developed per spike of the donor plant. Asterisk marks values statistically different from control (LT). GP—Golden Promise; LT—low-temperature tiller pre-treatment (21 days at 4 °C); LT+GSH—low-temperature tiller pre-treatment combined with the application of 0.3 mmol dm$^{-3}$ reduced glutathione; LT+MAN—low-temperature tiller pre-treatment combined with the application of 0.3 mmol dm$^{-3}$ mannitol; LT+MAN+GSH—low-temperature tiller pre-treatment combined with the application of 0.3 mmol dm$^{-3}$ mannitol and 0.3 mmol dm$^{-3}$ reduced glutathione; LT+DDC—low temperature tillers pre-treatment combined with the application of 0.25 mmol dm$^{-3}$ diethyldithiocarbamate trihydrate; LT+OTC—low temperature tillers pre-treatment combined with the application of 0.25 mmol dm$^{-3}$ 1,2-oxothiazolidine-4-carboxylic acid. All chemicals were applied during the last 4 days before microspore isolation.

The effect of different tillers pre-treatments on the effectiveness of ME and its modulation induced by season × genotype interaction. Data are presented in colour according to an arbitrary adopted scale (presented below), with colour intensity corresponding to the number of ELS developed per spike of the donor plant. Asterisk marks values statistically different from control (LT). GP—Golden Promise; LT—low-temperature tiller pre-treatment (21 days at 4 °C); LT+GSH—low-temperature tiller pre-treatment combined with the application of 0.3 mmol dm$^{-3}$ reduced glutathione; LT+MAN—low-temperature tiller pre-treatment combined with the application of 0.3 mmol dm$^{-3}$ mannitol; LT+MAN+GSH—low-temperature tiller pre-treatment combined with the application of 0.3 mmol dm$^{-3}$ mannitol and 0.3 mmol dm$^{-3}$ reduced glutathione; LT+DDC—low temperature tillers pre-treatment combined with the application of 0.25 mmol dm$^{-3}$ diethyldithiocarbamate trihydrate; LT+OTC—low temperature tillers pre-treatment combined with the application of 0.25 mmol dm$^{-3}$ 1,2-oxothiazolidine-4-carboxylic acid. All chemicals were applied during the last 4 days before microspore isolation.

The effect of different tillers pre-treatments on the effectiveness of ME and its modulation induced by season × genotype interaction. Data are presented in colour according to an arbitrary adopted scale (presented below), with colour intensity corresponding to the number of ELS developed per spike of the donor plant. Asterisk marks values statistically different from control (LT). GP—Golden Promise; LT—low-temperature tiller pre-treatment (21 days at 4 °C); LT+GSH—low-temperature tiller pre-treatment combined with the application of 0.3 mmol dm$^{-3}$ reduced glutathione; LT+MAN—low-temperature tiller pre-treatment combined with the application of 0.3 mmol dm$^{-3}$ mannitol; LT+MAN+GSH—low-temperature tiller pre-treatment combined with the application of 0.3 mmol dm$^{-3}$ mannitol and 0.3 mmol dm$^{-3}$ reduced glutathione; LT+DDC—low temperature tillers pre-treatment combined with the application of 0.25 mmol dm$^{-3}$ diethyldithiocarbamate trihydrate; LT+OTC—low temperature tillers pre-treatment combined with the application of 0.25 mmol dm$^{-3}$ 1,2-oxothiazolidine-4-carboxylic acid. All chemicals were applied during the last 4 days before microspore isolation.
tiller pre-treatments observed in some cases were the result of excessive decomposition of H₂O₂, whose amount dropped below the threshold initiating microspore reprogramming.

A variation in the response to some tiller pre-treatments suggests that the threshold level of ROS required for microspore reprogramming is genotype-specific and significantly lower for barley in comparison with triticale microspores.

Comparing data received from the analyses of antioxidative enzymes activities in the two experiments, it was confirmed that high activity of SOD, which catalyzes the dismutation of superoxide anion to H₂O₂, was the first feature differentiating embryogenic microspore suspensions from the non-embryogenic ones (Fig. 7a). This conclusion was based on the comparison between responsive and recalcitrant DH lines/cultivars as well as on the comparison between results received for microspores isolated under optimal (Exp. 3) and non-optimal (Exp. 4) conditions for in vitro culture establishment. However, a high SOD activity is not the only prerequisite necessary for effective ME initiation, which could be seen in cv. Golden Promise in Exp. 4. It seems that a high generation of H₂O₂ resulting from high SOD activity also requires tools for its efficient decomposition. The negative correlation between CAT activity, microspore viability and embryogenic development shows only apparent discrepancy with this assumption, as a generally very low CAT activity was probably not sufficient to provide effective defense.

The conducted experiments also revealed genetic specificity of antioxidative defense (Fig. 7b). A relatively high H₂O₂ generation detected in isolated microspores of triticale in association with low activity of antioxidative enzymes (Exp. 4) resulted in extremely low cell viability, which was probably the primary cause of ME induction failure (Fig. 7a). In the case of barley, genotype specific low/high activity of SOD and H₂O₂ level (cv. Igri/cv. Golden Promise) associated with low activity of CAT did not diminish microspore viability and embryogenic development shows only apparent discrepancy with this assumption, as a generally very low CAT activity was probably not sufficient to provide effective defense.

The conducted experiments also revealed genetic specificity of antioxidative defense (Fig. 7b). A relatively high H₂O₂ generation detected in isolated microspores of triticale in association with low activity of antioxidative enzymes (Exp. 4) resulted in extremely low cell viability, which was probably the primary cause of ME induction failure (Fig. 7a). In the case of barley, genotype specific low/high activity of SOD and H₂O₂ level (cv. Igri/cv. Golden Promise) associated with low activity of CAT did not diminish microspore viability and almost completely blocked embryogenic development in typically responsive isolated microspore cultures of cv. Igri. Different relationships between SOD and CAT in the tested lines may suggest that O₂^-/H₂O₂ ratio also plays a role in ME induction, as both of these ROS have been indicated as signalling molecules (Dietz et al. 2016; Noctor et al. 2018; Mhamdi and van Breusegem 2018). The observed variation is in accordance with a well-known genomic specificity of plant cell response to ME-inducing treatment and suggests an important role of other enzymatic or non-enzymatic elements of antioxidative defense. It is known that H₂O₂ can be produced from O₂^- not only through the reaction of disproportionation catalysed by SOD, but also through O₂^- reduction by other reductants, like GSH or ascorbate. Similarly, CAT deficiency can be compensated by the activation of another enzymes— ascorbate or glutathione peroxidases, alternative oxidase or monodehydroascorbate reductase (Mittler 2002). Another almost non-recognized problem is subcellular and organelar specificity of ROS localization and scavenging and the question of how intracellular ROS distribution influences the interaction with other cellular messengers like PGRs, calcium ions or reactive nitrogen species (Rodriguez-Serrano et al. 2012; Zechmann 2014; Mignolet-Spruyt et al. 2016).

Thus, further more detailed analyses of intercellular physiological background and its interaction with environmental stimuli are necessary for better understanding of the mechanism controlling successful ME initiation.

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Authors’ contributions IŻ conceived and designed the research, conducted experiments and wrote the manuscript. ED, MK, AN, KJ and SM performed experiments. PK and ES estimated H₂O₂ generation. KG and GG measured the activity of antioxidative enzymes. ED prepared photographic images. All authors read, reviewed and approved the manuscript.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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