Effect of copper and silver ions on sequence and DNA methylation changes in triticale regenerants gained via somatic embryogenesis

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Received: 22 April 2022 / Revised: 1 August 2022 / Accepted: 5 August 2022 / Published online: 19 August 2022 © The Author(s) 2022

Abstract
Somatic embryogenesis is a plant regeneration method that can be exploited in tissue culture systems for a variety of tasks, such as genetic modification or the selection of somaclones with advantageous characteristics. Therefore, it is crucial to create efficient regeneration procedures and comprehend how medium components affect regeneration effectiveness or the degree of variation created in plant tissue cultures. The level of tissue culture-induced variation in triticale regenerants was examined in the current study in relation to the concentration of copper and silver ions in the induction media as well as the length of time immature zygotic embryo explants were incubated on these media. The high degree of variation (45%) revealed by the methylation-sensitive amplified fragment length polymorphism approach for estimating variation included 38% DNA sequence alterations, 6% DNA demethylation, and 1% de novo DNA methylation. Different levels of variance were found in relation to various DNA sequence settings. The CHG context had the most alterations, whereas CG experienced the fewest; sequence variation predominated in each sequence context. Lower copper ion concentrations showed the most variance. However, it could not be connected to the duration of in vitro culture or the effect of silver ions. Accordingly, we think that altering the concentration of copper ions in the induction medium may throw off the equilibrium of the metabolic processes in which copper is involved, resulting in tissue culture-induced variation.

Keywords CG · CHG · CHH · Copper · Epigenetic variation · Genetic variation · Immature zygotic embryos · metAFLP · Silver

Introduction
Plant tissue culture (PTC) methods take advantage of the induced plant cell totipotency, which finds its outstanding reflection in the phenomenon of somatic embryogenesis (SE). Somatic embryogenesis is the multi-step developmental process in which somatic cells of plant explants differentiate into bipolar structures known as somatic embryos. Induction of SE can occur by acting on the explants with stress (Zavattieri et al. 2010) or through appropriate growth regulators (Méndez-Hernández et al. 2019). In addition, the success of plant regeneration through SE is also influenced by genotype (Fiuk and Rybczyński 2008), type and age of explant (Atak et al. 2008), condition of donor plants (Dahleen 1999), the composition of culture media (Dahleen and Bregitzer 2002), and especially type and concentration of growth regulators (Przetakiewicz et al. 2003) or carbon source (Ganesan and Jayabalan 2005) as well as the release of organic molecules by the explants into the culture medium (Nic-Can et al. 2015). Furthermore, depending on the mode, somatic embryo formation can occur either directly (without the callus phase) (Eudes et al. 2003) or indirectly (Vega et al. 2009), when somatic embryos are formed on the produced calli.

Over 60 years of studies on the phenomenon of SE has led to developing protocols for plant regeneration for many species, including cereals. In vitro, culture methods allowing obtaining plants via the SE pathway have also been elaborated for triticale, a new, and synthetic crop with increasing importance for agriculture and plant breeding. Successful induction of SE in triticale was first reported for immature inflorescence-derived callus cultures (Eapen and Rao 1985)
and then in callus cultures obtained from immature (Zimny and Rybczyński 1985) and mature zygot embryos (Padmaja et al. 1992). When it comes to the last two types of explants, embryogenic callus from immature embryos is more likely to produce somatic embryos than tissue grown from mature embryos (Vikrant and Rashid 2001). Although, in the case of triticale, more emphasis seems to have been put on systems of androgenic embryogenesis to obtain plants that are double haploids (DH) (Oleszczuk et al. 2004), the induction of SE with the use of zygotic embryos as explant is constantly practiced (Machczyńska et al. 2015).

Regenerants obtained by SE should be morphologically and genetically identical to the donor plant. However, this assumption is not always valid. Some changes (phenotypic, cytological, genetic) are observed in regenerants (de la Puente et al. 2008). The changes are defined by the common term somaclonal variation (Larkin and Scowcroft 1981) or tissue culture-induced variation (TCIV) (Machczyńska et al. 2014). The first term describes changes affecting regenerants (Eeckhaut et al. 2020) and sometimes also their generative progeny (Zehr et al. 1987), while the second one refers only to changes affecting regenerants (Orłowska et al. 2021b). The causes of variation induced by in vitro culture are complex (Bednarek and Orłowska 2019; Bednarek et al. 2021); indeed, the components forming the culture media (Elena and Ginzo 1988; Immonen 1996), time (Etienne and Bertrand 2003), and conditions of PTC can be the source of variation. For example, among the culture media components, micronutrients such as copper or silver seem to influence the course of SE and, as recently shown, genetic and epigenetic changes (Orłowska et al. 2021b).

At a concentration of 0.1 M, copper sulfate (CuSO₄) is one of the main components of the MS medium, and it exhibits positive effects on callus production and regeneration (Malik et al. 2021). Copper ions have been widely used in PTC to promote plant regeneration via SE (Dordević et al. 2019) and androgenesis pathways (Warchol et al. 2021). In the latter case, copper ions reduced the production of albino plants (Graudà et al. 2014). Also, CuSO₄ in a concentration of 0.1–100 μM significantly improved the regeneration of triticale shoots (Purnhauser and Gyulai 1993). On the other hand, the concentration of CuSO₄ from 1 to 80 μM had a positive effect on the induction of SE and plant regeneration and, in the case of androgenesis, it decreased the number of albino regenerants (Cho et al. 1998; Jacquard et al. 2009). However, the concentration of 100 μM had a toxic effect and suppressed the formation of embryogenic calli, adversely influenced regeneration, and promoted the formation of albino wheat plants (Miroshchnichenko et al. 2021).

Silver nitrate (AgNO₃) (Rojas-Lorz et al. 2019), silver thiosulfate (Ag₂S₂O₃) (Diab 2017), and silver nanoparticles (AgNPs) (Malik et al. 2021) are all added to in vitro culture media. Their primary function is to accelerate the callogenesis process and improve plant regeneration. Silver nitrate, usually added to the culture media at a concentration of 6–88 μM, impacts SE, shoot formation, efficient root growth, and organogenesis, prerequisites for successful genetic transformation (Paladi et al. 2017). The use of AgNO₃ improved SE in wheat species Triticum durum (a more than 22-fold increase) (Fernandez et al. 1999) and Triticum aestivum (Wu et al. 2006), barley (Orłowska et al. 2020), maize (Carvalho et al. 1997), and rice (Ghobeishavi et al. 2015). Silver nitrate has been shown to mediate the inhibition of ethylene (Beyer 1976), a plant hormone involved in many developmental processes, including fruit ripening, abscission, senescence, growth, and flowering (Hobson et al. 1984). As an ethylene inhibitor, AgNO₃ has significantly improved shoot regeneration in wheat (Purnhauser et al. 1987) and counteracted the aging process of callus of this species (Wu et al. 2006).

Time is another factor that may impact somatic embryogenesis. The extended duration of in vitro culture increases the number of mutations in regenerated plants (Duncan 1996), which may involve an increased mutation rate (Murashige and Nakano 1965) and sequential accumulation of mutations over time (Kaeppler et al. 2000). Thus, prolonged tissue culture time resulted in increased variation, manifested in different phenotypic variants (Armstrong and Phillips 1988). Also, increased callus induction time during SE decreases plant morphogenic potential and increases the number of albino plants and calli (Wen et al. 2004). It was documented that DNA methylation changes in regenerants may be related to time on the tissue culture run (Bednarek and Orłowska 2020).

Assessment of TCIV variation can be performed at different levels using a range of techniques, from the use of morphological characteristics of the clones (Eeckhaut et al. 2020) through the use of biochemical methods (Thomas et al. 2006) to molecular methods (Bednarek et al. 2007; Orłowska and Bednarek 2020; Patzak et al. 2021). The development of molecular techniques such as methylation-sensitive amplified fragment length polymorphism (metAFLP) has made it possible to characterize the variation induced by in vitro cultures at the level of DNA sequence and its methylation in one analysis. The metAFLP exploits the properties of restriction enzymes which are sensitive to the presence of methylated cytosine in or around the cutting site. The metAFLP method utilizes the Acc65I and KpnI isoschizomers to recognize the 5′-GGT-3′ sequence. Acc65I does not hydrolyze a DNA strand if the cytosine is methylated. On the other hand, the restriction enzyme KpnI cuts the DNA strand regardless of the presence of cytosine methylation. In the metAFLP technique, the differences between DNA profiles result from cutting the DNA of the donor plant and the regenerant with two pairs of restriction enzymes, namely Acc65I/MsEl and KpnI/MsEl. The method allows the estimation of changes both qualitatively
and quantitively. Using the isoschizomers permits quantitative assessment of TCIV and its components (sequence variation-SV, demethylation-DMV, and de novo DNA methylation-DNMV). In addition, the development of the method enabled the estimation of changes in specific DNA nucleotide sequences (hereafter referred to as sequence contexts) reflecting symmetric (CG and CHG) and asymmetric (CHH) cytosine methylation (H stands for A, C, T).

Although the function of copper and silver ions is relatively well studied concerning obtaining regenerants in PTC, including cereal ones, there is a lack of information on how these ions, in combination with various incubation times of immature zygotic embryos on induction medium (IM), can influence TCIV in triticale regenerants obtained by SE. Thus, the aim of this study was to evaluate the impact of copper and silver ions and incubation time of explants on TCIV in triticale regenerants obtained via SE in immature zygotic embryo cultures.

Material and methods

Acquisition of donor plants

The cultivar T28/2 of winter triticale (X Triticosecale spp. Wittmack ex A. Camus 1927) derived from cv. Presto × cv. Mungis cross provided by dr Sylwia Oleszczuk (Plant Breeding and Acclimatization Institute-NRI, Radzików, Poland) served to prepare donor plants via in vitro anther culture. The precise procedure for getting donor plants was previously published (Pachota et al. 2022). Simply put, we first obtained regenerants that were DH in anther culture. The final donor plants in the experiment were then obtained as the generative progeny of DH regenerants. Twenty donor plants were used to conduct additional research on triticale plant regeneration via SE from immature zygotic embryos under various in vitro growth conditions.

Plant regeneration via somatic embryogenesis

Twenty donor seedlings were cultivated in a growth chamber at 16/12 °C (day/night). Photoperiod (16 h/8 h; light/dark) and light intensity (190 μE m⁻² s⁻¹) were maintained using high-pressure sodium lamps. After vernalization at 4 °C for 6 weeks under short-day photoperiod (8 h/16 h) in 20 μE m⁻² s⁻¹ light intensity plants were grown in greenhouse conditions. Spikes with unripe caryopses were harvested after 12–16 days of self-pollination. The caryopses removed from the tillers were surface-sterilized with 70% ethanol for 1 min and then transferred to 10% sodium hypochlorite solution (NaOCl) for 20 min. The caryopses were then thoroughly rinsed four times with distilled water. Immature zygotic embryos were dissected from the disinfected caryopses and plated on an MS induction medium (Murashige and Skoog 1962) with 2 mg l⁻¹ 2,4-dichlorophenoxyacetic acid. Nine variants of tissue culture (trials: T1–T9) included IM with various concentrations of CuSO₄ × 5H₂O (0.1, 5, 10 μM) and AgNO₃ (0, 10, 60 μM), and the incubation time (35, 42, 49 days) of immature embryos on these media (Table 1). The combination of the factors tested (Cu, Ag, time), as well as the concentration levels and the number of days, was determined according to the design of experiment principles based on Taguchi’s orthogonal arrays (Taguchi 1986). The induction step’s length (incubation time) covered the time from plating embryos on IM to calli collection and transferring them on regeneration media. Trial T1 was considered a control. Immature zygotic embryos were incubated on IM at 26 °C under a photoperiod of 16 h/8 h (light/dark). After several days, first calli and subsequent somatic embryos were recorded and transferred on a solid regeneration medium 190–2. Calli and embryos were incubated at 26 °C under 16 h/8 h (light/dark) conditions. Regenerated plants were transferred to a glass flask with rooting medium N6I (Chu 1978) supplemented with 2 mg l⁻¹ indole-3-acetic acid. Seedlings with a well-developed root system were transferred into pots (soil to sand 3:1) and grown to maturity under controlled conditions in the greenhouse on a 16 h/8 h (light/dark) photoperiod. Immature zygotic embryos from each donor plant were plated on all the IM tested (T1–T9); however, regenerants for each trial were not obtained for every donor plant. Between 0 and 12 regenerants were obtained per trial for different donor plants. Finally, regenerants from a single donor plant representing all in vitro culture conditions tested (T1–T9) were selected for analysis. After all, forty-five regenerants originating from one donor plant were prepared for analysis. Five regenerants from each trial (T1–T9) were tested.

DNA extraction and metAFLP procedure

The leaves of young seedlings of 20 donor plants and 45 regenerants were ground in liquid nitrogen. The extraction of DNA was performed using the DNeasy MiniPrep Kit (Qiagen, Hilding, Germany) according to the procedure recommended by the manufacturer. The amount of DNA was determined spectrophotometrically (Nanodrop, Thermo Fisher

| Trial | T1 | T2 | T3 | T4 | T5 | T6 | T7 | T8 | T9 |
|-------|----|----|----|----|----|----|----|----|----|
| CuSO₄ × 5H₂O (μM) | 0.1 | 0.1 | 0.1 | 5  | 5  | 5  | 5  | 10 | 10 |
| AgNO₃ (μM)         | 0   | 10 | 60 | 60 | 0  | 10 | 10 | 60 | 0  |
| Incubation time (days) | 35 | 42 | 49 | 42 | 49 | 35 | 49 | 35 | 42 |

Table 1 Conditions of triticale plant regeneration on induction media via somatic embryogenesis
Scientific, Wilmington, USA) at A$_{260}$/A$_{280}$ ∼ 1.8 and A$_{260}$/A$_{230}$ ∼ 2.0. DNA quality and integrity were tested in a 1.2% agarose gel by staining DNA fragments with ethidium bromide. For the metAFLP technique, two samples of genomic DNA (2 × 500 ng) were prepared from each plant.

Obtaining DNA band profiles by metAFLP was performed for donor plants and regenerants according to the methodology described previously (Bednarek et al. 2007) and then modified (Machczyńska et al. 2014). The prepared DNA samples, two from each plant, were subjected to restriction enzyme digestion; the 6-bp cutting enzymes Acc65I and KpnI and the 4-bp cutting enzyme MseI were used (NEB Ipswich, Massachusetts, USA). Adapters were ligated onto the digested DNA fragments, and these fragments were then amplified using a polymerase chain reaction (PCR). After pre-selective PCR amplification, products were diluted with sterile distilled water (1 : 19 dilutions), of which 1.5 μL was used as the template for selective PCR amplification with radio-labeled primer (labeling with γ$^{32}$P) and unlabeled MseI + YYY (Table 2). Electrophoresis was carried out on 7% polyacrylamide in 1× TBE buffer. Gels were exposed to X-ray film, then developed by a conventional procedure.

**Analysis of data obtained by metAFLP**

The metAFLP profiles were scored and generated binary presence-absence data matrices (1 for presence, 0 for absence of a band). Two matrices were prepared: one reflected fragments obtained with Acc65I/MseI (A) enzymes and the second with KpnI/MseI (K). The binary data from the two matrices were collated and compared. The Acc65I and MseI enzyme pair for cutting genomic DNA provided information regarding changes in DNA sequence and methylation. On the other hand, cutting genomic DNA with KpnI and MseI enzymes provided information on DNA sequence variations (genetic variation). Comparing the two matrices allows DNA fragments to be extracted and describe changes in DNA methylation (epigenetic variation) and creates a third virtual matrix (M). Based on matrices K and M, we calculated the total number of loci generated in the analysis and expected heterozygosity (He) under the assumption of Hardy–Weinberg equilibrium in GenAlex (Excel add-in software). XLSTAT software was applied to agglomeration (UPGMA and dissimilarity Jaccard coefficients).

The quantitative characterization of TCIV was previously described in detail (Bednarek et al. 2007; Machczyńska et al. 2014), with some modifications accompanied by an Excel file capable of carrying out the relevant calculations (Ołowska and Bednarek 2020). This description included formulas for estimating SV, DMV, DNVM, and TCIV also in specific sequence contexts (CG, CHG—symmetrical, and CHH—asymmetrical) in which in vitro culture-induced changes may occur.

The values obtained during the estimation of in vitro culture-induced changes were analyzed using analysis of variance (one-way ANOVA) and ANOVA with Brown-Forsythe $F$ when the data did not meet the assumptions of the Levene test (lack of homogeneity of variances in the study groups). The means for different metAFLP characteristics (TCIV, SV, DMV, DNVM) were compared with each other, also in different sequence contexts (CHH, CG, CHG) for regenerants obtained under different in vitro conditions (trials T1–T9). Multiple comparisons were checked with the Games-Howell post hoc test when the homogeneity of variance was violated. ANOVA was performed using XLSTAT software.

| Table 2 | Arrangement of primers used for selective DNA amplification of triticale donors and regenerants by the metAFLP technique |
|---------|--------------------------------------------------------------------------------------------------|
| **metAFLP oligomer** | **Sequence 5’ → 3’** |
| Labeled γ$^{32}$P selective oligonucleotides |  |
| CG-GAC | CA TGC GTA CAG TAC CGA C  |
| CG-GCA | CA TGC GTA CAG TAC CGC A  |
| CG-GGC | CA TGC GTA CAG TAC CGG C  |
| CG-TCG | CA TGC GTA CAG TAC CTC G  |
| CXG-AGA | CA TGC GTA CAG TAC CAG A  |
| CXG-AGC | CA TGC GTA CAG TAC CAG C  |
| CXG-AGG | CA TGC GTA CAG TAC CAG G  |
| CXG-ATG | CA TGC GTA CAG TAC CAT G  |
| CXG-TGC | CA TGC GTA CAG TAC CTA C  |
| CXG-TTG | CA TGC GTA CAG TAC TGG T  |
| CXX-ATT | CA TGC GTA CAG TAC CAT T  |
| CXX-TAA | CA TGC GTA CAG TAC CTA A  |
| Selective oligonucleotides MseI + YYY |  |
| M-CAC | GAT GAG TCC TGA GTA ACA C  |
| M-CGT | GAT GAG TCC TGA GTA ACG T  |
Results

Twenty donor plants that were the generative progeny of a regenerant obtained through androgenesis showed no morphological differences from each other. Only one donor plant out of twenty from which immature zygotic embryos were obtained for the somatic embryogenesis process yielded 45 regenerants in all in vitro culture variants tested (T1–T9). The regenerants were morphologically identical to the donor plant independently of in vitro plant regeneration conditions (trials T1–T9).

The metAFLP resulted in fully stable and reproducible banding patterns amplified for donors and regenerants. Four hundred forty-four bands amplified with 13 primer pairs based on the KpnI/MseI (K) metAFLP platform were evaluated for donor plants. The Acc65I/MseI – KpnI/MseI (M) data reflecting DNA methylation changes resulted in 34 bands. There were 426 unique bands for donors derived under specific trial conditions and related to sequence variation. In contrast, a unique banding pattern related to DNA methylation was exhibited by 16 signals (Table 3). The metAFLP analysis (with 11 primer pairs) concerning regenerants revealed the presence of 213 related to sequence variation and 39 to DNA methylation change; 186 and 12 were unique for regenerants representing selected trials. The He was lower for donors (0.007 for K and M) than for the regenerants set (0.014 for K and M markers) (Table 3). The polymorphic loci (%P) percentage was slightly lower for the K matrix than for donors. Similar differences in the %P level were observed for K and M matrices for regenerants (Table 1). The %P for regenerants also indicated a lower value for sequence data (K) and a higher value for methylation data (M) (Table 3).

Agglomeration analysis of donor plants based on (K) and (M) data (Fig. 1a, b) shows donor plants exhibited higher variation based on M than on K. In both analyses, a single donor plant was somewhat apart from the other samples, which were more uniform but still formed three additional clusters.

Factor analyses carried on K and M markers are congruent with the UPGMA analyses (Fig. 2a, b), where a single donor plant is apart from the other samples, possibly forming clusters encompassing individuals from different trials.

The metAFLP quantitative characteristics (Table 4) showed that the highest level of TCIV was observed for trial T2, with the lowest concentration of copper ions, intermediate levels of silver ions, and the intermediate incubation time of immature zygotic embryos on the IM (Table 1). In contrast, the highest level of copper ions applied, the intermediate level of silver ions, and the highest incubation time resulted in the slightest changes induced by the in vitro culture (T7). Among the components of TCIV, SV was the highest, and maximum and minimum levels of DNA sequence changes were recorded for regenerants from T2 and T7 trials, similarly to TCIV. However, DNA methylation changes were lower than SV and amounted to about 6% for DMV and 1% for DNMV. Furthermore, DMV was favored by the highest levels of copper and silver ions and the intermediate incubation time of the explants on the inducing media (T8), whereas the slightest demethylation events were recorded for the T2–T4 trials at the lowest copper and intermediate and highest silver ion concentrations; the incubation time of immature embryos on the IM for these trials was intermediate and the longest. Similar to DMV, the fewest de novo DNA methylation events were observed for T2–T4 and the highest for T5–T9 (Table 4).

According to Levene’s test, the assumption of homogeneity of variance for trials in each metAFLP characteristic was violated (Table 4). ANOVA indicated differences between TCIV, SV, and DMV trials, while no differences were observed for DNMV (Table 4). Games-Howell’s post hoc analysis for TCIV trials revealed that trials T1–T4 and T7–T9 compose separate groups; each group contains a trial with the highest or lowest in vitro induced variation level. Similar to TCIV, the Games-Howell test noted intertrial significant differences for SV. For changes concerning DNA methylation, significant differences were noted only for DMV. The post hoc Games-Howell test indicated a homogeneous group that included regenerants from T1–T7 trials and distinct regenerants representing trial T8. Trial 8 included the regenerants with the highest DMV.

ANOVA and grouping according to the Games-Howell post hoc test indicated significant differences between individual metAFLP characteristics concerning symmetric and asymmetric sequence contexts. For SV and DMV, the most significant changes were observed in the symmetric CHG context and the least in the symmetric CG context. On the other hand, in the case of changes concerning DNMV, no such changes were observed in the symmetric CG context, and the highest DNMV was observed in the asymmetric CHH context (Table 5).
A detailed examination of the differences between the regenerants belonging to the different trials in the different characteristics of the metAFLP considering the CG, CHG, and CHH contexts showed that a higher variation was observed in the DNA sequence in the CHG context (21.55%) and the least in the CG (6.21%) (Fig. 3). Subsequently, between 4 and 0.45% of the changes were related to DMV in the CHG and CG contexts, and the least number of changes was related to DNMV (0–0.76%).

According to Levene’s test, the assumption of homogeneity of variance for almost all tested metAFLP characteristics and sequence contexts violated the assumptions \(p < 0.0001\), apart from SV_CHG. Thus, ANOVA with a Tukey HSD post hoc test for SV_CHG indicated differences between trials. T1–T3, T4–T6, and T7–T9 trials are divided into separate groups. In the cases of SV_CG and DMV_CG, ANOVA with a Games-Howell post hoc test was used. This analysis for SV_CG indicated differences between regenerants belonging to trials T1-T4, T8, and T9. The same analysis was conducted for DMV_CG grouped regenerants from trials T1–T7, showing their distinctiveness from regenerants belonging to trials T8 and T9 (Table 6). Differences between regenerants obtained in separate in vitro culture conditions for the other metAFLP characteristics and sequence contexts were insignificant.

**Discussion**

Regenerating plants via in vitro cultures is often affected by TCIV (Orłowska et al. 2021a), possibly due to in vitro culture conditions (Bednarek and Orłowska 2019). Such variation was also observed in triticale regenerants obtained by SE, where somaclones/regenerants differ in resistance to the *Fusarium* head blight (Góral and Arseniuk 1997), showing the variation of the mitochondrial atp6 gene region (Schmidt et al. 1996), or showing confirmed rearrangements in the mtDNA (Weigel et al. 1995). The use of the metAFLP method to study variation induced by tissue culture allowed the estimation of TCIV concerning changes in the sequence and methylation of genomic DNA (Machczyńska et al. 2014). Running experiments with the metAFLP technique require the appropriate research material, consisting of selected donor plants and their regenerants (Bednarek et al. 2007). Such material can attribute the observed changes to the action of in vitro culture. Hence, in the
present study, evaluating the level and characteristics of variation in triticale regenerants obtained under different in vitro conditions (T1–T9), donor plants were used, which are the generative progeny of the regenerant obtained by androgenesis. All obtained donor plants showed no phenotypic changes and were identical to the initial plant, indicating their genetic uniformity. This result was also confirmed at the level of molecular patterns obtained
Table 4 Mean values for in vitro induced variation and its components for triticale regenerants, including grouping trials by the post hoc test

| Trial | metAFLP characteristics†† |
|-------|---------------------------|
|       | TCIV | SV | DMV | DNMV |
| T1    | 45.55<sup>a</sup> 38.45<sup>a</sup> 5.97<sup>b</sup> 1.12<sup>a</sup> |
| T2    | 45.62<sup>a</sup> 38.57<sup>a</sup> 5.93<sup>b</sup> 1.11<sup>a</sup> |
| T3    | 45.47<sup>a</sup> 38.43<sup>a</sup> 5.93<sup>b</sup> 1.11<sup>a</sup> |
| T4    | 45.40<sup>b</sup> 38.35<sup>a</sup> 5.93<sup>b</sup> 1.11<sup>a</sup> |
| T5    | 45.10<sup>b</sup> 38.01<sup>b</sup> 5.97<sup>b</sup> 1.12<sup>a</sup> |
| T6    | 45.10<sup>b</sup> 38.01<sup>b</sup> 5.97<sup>b</sup> 1.12<sup>a</sup> |
| T7    | 44.36<sup>b</sup> 37.29<sup>b</sup> 5.95<sup>b</sup> 1.12<sup>a</sup> |
| T8    | 44.84<sup>b</sup> 37.39<sup>b</sup> 6.33<sup>a</sup> 1.12<sup>a</sup> |
| T9    | 44.69<sup>b</sup> 37.32<sup>b</sup> 6.26<sup>a</sup> 1.12<sup>a</sup> |
| Levene’s test, p | 0.037 0.027 0.0001 0.0001 |
| Browne-Forsythe F-ratio | 8.750 15.534 27.199 1.160 |
| p | 0.0001 0.0001 0.0001 0.429 |

††The TCIV, SV, DMV, and DNMV characteristics reflect tissue culture-induced variation, sequence variation, demethylation, de novo methylation, respectively.

The results of ANOVA statistics and the Games-Howell post hoc tests demonstrating differences between DNA sequence contexts for the metAFLP characteristics

Table 5

| Methylation context | metAFLP characteristics |
|---------------------|--------------------------|
|                     | SV | DMV | DNMV |
| CHG                 | 21.55<sup>a</sup> 4.44<sup>a</sup> 0.37<sup>b</sup> |
| CG                  | 6.21<sup>c</sup> 0.45<sup>c</sup> 0.00<sup>c</sup> |
| CHH                 | 10.21<sup>b</sup> 1.13<sup>b</sup> 0.76<sup>a</sup> |
| Levene’s test, p    | 0.0001 0.0001 0.0001 |
| Browne-Forsythe F-ratio | 26.015,670 22,620,932 439,028,173 |
| p                   | 0.0001 0.0001 0.0001 |

The a, b and c superscript letters indicate Games-Howell grouping

with the metAFLP technique. We obtained donor plants that revealed as little as 3.04% and 3.26% of polymorphic metAFLP loci concerning sequence and methylation changes, respectively. Also, the expected heterozygosity confirmed a low level of polymorphism of donor plants in the case of both matrices K and M. Cluster analysis showed that donors’ genetic and epigenetic dissimilarities varied depending on whether genetic or methylation data were analyzed, indicating higher variation regarding changes in DNA methylation (0.7% K vs. 14% M). One plant was randomly selected from the pool of donor plants, and regenerants derived from this plant were included in the analysis to avoid the impact of pre-existing variation (Flinn et al. 2020) on TCIV. This procedure avoided disturbances in TCIV that could result from potential variations introduced by the donors.

The evaluation of molecular data for regenerants obtained under different in vitro culture conditions showed that He and %P for regenerants were higher than for donors, which was expected due to tissue culture-induced stress. Furthermore, in regenerants also, more polymorphic fragments were observed for methylation data. Visualization of the factor analysis results for regenerants illustrated the clustering of regenerants forming individual trials and therefore obtained under different in vitro culture conditions. For sequence and DNA methylation data, the donor plant was located at the periphery of the regenerants. In contrast, the regenerants formed two groups, each reflecting Cu and Ag ion supplementation and modifications regarding the incubation time of immature zygotic embryos on IM. Thus, pointing to the applied modifications of the substrates and the conditions for obtaining regenerants as those that can influence the resulting changes detected at the molecular level.

The metAFLP analysis performed for triticale regenerants obtained in immature zygotic embryo cultures indicated a high level of TCIV ranging from 44.36 to 45.55%. The result obtained is higher than the TCIV observed in barley regenerants (8.74%) acquired in the same type of tissue culture (Orłowska 2021). However, the results received for TCIV do not generally differ from the level of changes induced in plant tissue culture in triticale regenerants derived by androgenesis (Orłowska et al. 2022), although they are slightly lower (45.13 somatic embryogenesis vs. 51.84 androgenesis). Such significant differences in TCIV levels for regenerants for two different kinds of grain cereal may be due to the genome specificity of each species. Barley is a self-pollinating, diploid species with 2n < 100 enzymes involved in crucial life processes of plants (Burkhead et al. 2009); obtaining plants through in vitro cultures may indirectly lead to sequence variation in the genome of regenerants (Orłowska et al. 2021b). The highest

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The level of TCIV observed here was for the T2 trial, where the lowest level of CuSO₄×5H₂O (0.1 μM) was used. In contrast, the highest concentration of CuSO₄×5H₂O (10 μM) resulted in the lowest observed TCIV level. These results are consistent with data for triticale regenerants obtained by androgenesis (Orłowska et al. 2022).

The variation induced in plant tissue cultures is genetic and epigenetic. Genetic variation, i.e., changes in DNA sequence, had the highest contribution to TCIV. Similarly, the highest contribution of SV to TCIV was observed in regenerants obtained by androgenesis (Machczyńska et al. 2015; Orłowska et al. 2022) and somatic embryogenesis (Machczyńska et al. 2015), although in the study by Machczyńska and co-authors, the level of SV was lower and oscillated around 19%. The in vitro culture conditions with the lowest CuSO₄×5H₂O concentration showed the most remarkable genetic changes. It cannot be excluded that the minimum amount of copper does not provide an adequate amount of copper cofactor for the superoxide dismutase Cu/Zn-SOD.

Consequently, the dysfunction of the enzyme and oxidative stress, causing mutations in DNA (Poetsch 2020), may be elevated. Apart from SV, epigenetic changes were also observed in the regenerants, which concerned a decrease in DNA methylation (demethylation) and an increase in genomic DNA methylation compared to the donor plants; these changes were described as de novo
methylation. Changes in up- (DNMV-3.34%) or down- (5.15%) methylation were a lot smaller than changes in the sequence variation. Furthermore, differences between regenerants obtained under different in vitro culture conditions were noted only for DMV. ANOVA did not reveal differences between trials concerning DNMV. Regenerants from the T3–T4 trials had the lowest DMV in the presence of a minimal concentration of CuSO₄ × 5H₂O.

In contrast, the highest copper ion concentration led to the most significant changes in DMV. This data is only partially consistent with the results for regenerants obtained by anther cultures. While concordance was observed for in vitro culture conditions leading to minimal DMV levels, the highest DMV levels in regenerants obtained by androgenesis were associated with minimal CuSO₄ × 5H₂O concentrations. It appears that the high DMV associated with maximum CuSO₄ × 5H₂O levels may reflect the action of copper as a factor responsible for ROS production. Elevated levels of ROS may lead to oxidative changes in methylated cytosine (Kurek et al. 2019) and, through repair mechanisms, to sequence variation (Pachota et al. 2022). Finally, comparing the presented data with the results for barley, we observed that in the genome of barley regenerants received both by somatic embryogenesis and androgenesis, the changes between regenerants concerned all the sequence contexts (Orłowska 2021).

Analyzing different in vitro culture conditions for the changes assigned to SV in the CHG and CG contexts and DMV in the CG context, it can be seen that the highest level of SV in the CHG context is combined with minimal supplementation with copper ions and with the full range of silver concentrations and the full range of days of incubation of immature zygotic embryos on IM. This arrangement of data also applies to SV in the CG context and generally replicates the variation analysis performed for TCIV and SV without splitting the data into methylation contexts. In contrast, DMV in the CG context was highest in maximum CuSO₄ × 5H₂O concentration (T8, T9) and the presence (60 μM) (T8) and absence (T9) of Ag ion supplementation. Furthermore, the incubation time of immature zygotic embryos on IM where the highest DMV was detected in the CG context was the lowest (37 days, T8) and moderate (42 days, T9). In contrast, differences between trials for the CG context concerning DMV were not observed for triticale regenerants obtained by androgenesis (Pachota et al. 2022). This relationship between in vitro culture conditions and DMV was similar to an analysis where demethylation changes were studied without considering methylation contexts. Therefore, an analysis that considers the level of variation for regenerants from different trials concerning sequence contexts potentially typifies copper ions as a factor that may influence the observed changes. However, other analyses would be required to determine how copper may affect SV and DMV in symmetric CHG and CG contexts.

**Conclusions**

The emergence of TCIV has an impact on the production of triticale regenerants from immature zygotic embryo cultures; furthermore, the magnitude of this variation seems to be influenced by the tissue culture conditions. The modifications that were noticed were impacted by the addition of copper salts to IM. However, the analyses carried out do not suggest the relevance of supplementation with silver salts or the influence of time as factors that can determine genetic and epigenetic alterations in triticale regenerants, at least within the scope of elementary statistical tests. Additionally, an in-depth investigation of the metAFLP technique’s molecular data revealed certain DNA sequence contexts (CHG and CG) in which sequence-related alterations (SV_CG, SV_CHG), as well as DNA demethylation (DMV_CG), were noted. Therefore, we have a tendency to believe that
altering copper ion supplementation in the IM affects the balance of metabolic pathways and causes TCIV.

**Author contribution**  R. O. conceptualization; K. A. P. and R. O. methodology and formal analysis and investigation; R. O. and K. A. P. writing—original draft preparation and writing—review and editing.

**Funding** This study was funded by the Ministry of Agriculture and Rural Development (Poland), project no. HORN-801-PB-22/15–18.

**Data availability** All data generated during this study are included in this published article.

**Declarations**

**Conflict of interest** The authors declare no competing interests.

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**References**

Anantasaran J, Kanchanapoom K (2008) Influence of medium formula and silver nitrate on in vitro plant regeneration of Zinnia cultivars. Songklanakarin J Sci Technol 30:1–6

Armstrong CL, Phillips RL (1988) Genetic and cytotgenetic variation in plants regenerated from organogenic and friable, embryogenic tissue-cultures of maize. Crop Sci 28:363–369. https://doi.org/10.2135/cropsci1988.0011183X002800020038x

Atak M, Kaya M, Khawar KM, Saglam S, Ozcan S, Ciftci CY (2008) Influence of medium formula and silver nitrate on in vitro plant regeneration of Zinnia cultivars. Crop Sci 42:934–938. https://doi.org/10.2135/cropsci2002.0934

de la Puente R, González AI, Ruiz ML, Polanco C (2008) Somaclonal variation in rye (Secale cereale L.) analyzed using polymorphic and sequenced AFLP markers. In: Vitro Cell Dev Biol-Plant 44:419–426. https://doi.org/10.1007/s11627-008-9152-z

Diab MI (2017) In vitro propagation of the endangered date palm (Phoenix dactylifera L.) cv. Ghazal: 2- Improvement of germination and development of embryos by silver nitrate and silver thiosulfate. IOSR-JBB 3:66–74. https://doi.org/10.9790/264X-03046674

Dordević B, Nedela V, Thilaříková E, Trojan V, Havel L (2019) Effects of copper and arsenic stress on the development of Norway spruce somatic embryos and their visualization with the environmental scanning electron microscope. New Biotechnol 48:35–43. https://doi.org/10.1016/j.nbt.2018.05.005

Duncan RR (1996) Tissue culture-induced variation and crop improvement. Adv Agron 58:201–240. https://doi.org/10.1016/S0065-2113(08)60256-4

Eaep S, Rao PS (1985) Plant regeneration from immature inflorescence callus cultures of wheat, rye and triticale. Euphytica 34:153–159. https://doi.org/10.1007/BF0022875

Eckhart T, Van Houtven W, Bruznican S, Leus L, Van Huylenbroeck N (2017) Effects of age on somatic embryogenesis from immature zygotic embryos of 5 Turkish triticale genotypes. Afr J Biotechnol 16:4483–4490. https://doi.org/10.5897/AJB2016.08608

Eeckhaut T, Van Houtven W, Bruznican S, Leus L, Van Huylenbroeck N (2017) Effects of age on somatic embryogenesis from immature zygotic embryos of 5 Turkish triticale genotypes. Afr J Biotechnol 16:4483–4490. https://doi.org/10.5897/AJB2016.08608

Fernandez S, Michaux-Ferrière N, Coumans M (1999) The embryogenic response of immature embryo cultures of durum wheat (Triticum durum Desf.). histology and improvement by AgNO3. Plant Growth Regul 28:147–155. https://doi.org/10.1023/A:1006142504577
Fliuk A, Rybczyński JJ (2008) Genotype and plant growth regulator-dependent response of somatic embryogenesis from Gentiana spp. leaf explants. In Vitro Cell Dev Biol-Plant 44:90–99. https://doi.org/10.1007/s11627-008-9124-3

Flinn B, Dale S, Disharoon A, Kresovich S (2020) Comparative analysis of in vitro responses and regeneration between diverse bioenergy sorghum genotypes. Plants-Basel 9:248. https://doi.org/10.3390/plants9020248

Ganesan M, Jayabalanan N (2005) Carbon source dependent somatic embryogenesis and plant regeneration in cotton, Gossypium hirsutum L. cv. SVPR2 through suspension cultures. Indian J Exp Biol 43:921–925

Ghobeishavi H, Dorani Uliaie A, Alavikia SS, Valizadeh M (2015) Study of factors influencing somatic embryogenesis in rice (Oryza sativa L.). Int J Adv Biol Biomed Res 3:43–50

Góral T, Arseniuk E (1997) Somaclonal variation in winter triticale for breeders to enhance anther culture response in wheat (Triticum aestivum L.). Heliyon 7:e07075–e07075. https://doi.org/10.1016/j.heliyon.2021.e07075

Grauda D et al (2014) Anther culture effectiveness in producing doubled haploids of cereals. Proc Latv Acad Sci Sect B 68:142–147. https://doi.org/10.2478/prolas-2014-0016

Hassan MF, Islam SMS (2021) Effect of silver nitrate and growth regulators to enhance anther culture response in wheat (Triticum aestivum L.). Front Plant Sci 10:77–77. https://doi.org/10.3389/fpls.2019.01328

Hobson GE, Nichols R, Davies JN, Atkey PT (1984) The inhibition of tomato fruit ripening by silver. J Plant Physiol 116:21–29. https://doi.org/10.1016/S0176-1617(84)80080-2

Immonen AST (1996) Influence of media and growth regulators on somatic embryogenesis and plant regeneration for production of primary triticales. Plant Cell Tissue Organ Cult 44:45–52. https://doi.org/10.1007/BF00045912

Jacquard C et al (2009) Microspore embryogenesis and programmed cell death in barley: effects of copper on albimism in recalcitrant cultivars. Plant Cell Rep 28:1329–1339. https://doi.org/10.1007/s00299-009-0773-z

Kaeppler SM, Kaeppler HF, Rhee Y (2000) Epigenetic aspects of somaclonal variation in plants. Plant Mol Biol 43:179–188. https://doi.org/10.1023/A:1006423101343

Kurek K, Plitta-Michalak B, Ratajczak E (2019) Reactive oxygen species as potential drivers of the seed aging process. Plants-Base 8:174. https://doi.org/10.3390/plants8060174

Larkin PJ, Scowcroft WR (1981) Somaclonal variation - a novel source of variability from callus cultures for plant improvement. Theor Appl Genet 60:197–214

Ma X-F, Gustafson JP (2008) Allopolyoidization-accommodated genomic sequence changes in triticate. Ann Bot (Lond) 101:825–832. https://doi.org/10.1093/aob/mcm331

Machczyńska J, Orlowska R, Zimny J, Bednarek PT (2014) Extended metaAFLP approach in studies of the tissue culture-induced variation (TCIV) in case of triticale. Mol Breeding 34:845–854. https://doi.org/10.1007/s11032-014-0079-2

Machczyńska J, Zimny J, Bednarek P (2015) Tissue culture-induced genetic and epigenetic variation in triticale (× Triticosecale Wittmack) cv. Bogo Plant Cell Rep 22:885–893. https://doi.org/10.1007/s11103-015-0079-2

Malik WA et al. (2021) Exploring potential of copper and silver nano particles to establish efficient callogenesis and regeneration system for wheat (Triticum aestivum L.). GM Crop Food. 1–22. https://doi.org/10.1080/21646598.2021.1917975

Méndez-Hernández HA et al (2019) Signaling overview of plant somatic embryogenesis. Front Plant Sci 10:77–77. https://doi.org/10.3389/fpls.2019.00077

Miroshnichenko D, Klementyeva A, Dolgov S (2021) The effect of daminozide, dark/light schedule and copper sulphate in tissue culture of Triticum timopheevii. Plants-Basel 10:2620

Murashige T, Nakano R (1965) Morphogenetic behavior of tobacco tissue cultures and implication of plant senescence. Am J Bot 52:819–827. https://doi.org/10.1002/0001-1377(1965)52:7<819::AID-AMJ790>3.0.CO;2-6

Murashige T, Skoog F (1962) A revised medium for rapid growth and bio assays with tobacco tissue culture. Physiol Plant 15:473–497. https://doi.org/10.1111/j.1399-3054.1962.tb00802.x

Oleszczuk S, Grzechnik N, Mazon AS, Zimny J (2019) Heritability of meiotic restitution and fertility restoration in haploid triticale. Plant Cell Rep 38:1515–1525. https://doi.org/10.1007/s00299-019-02462-6

Oleszczuk S, Sowa S, Zimny J (2004) Direct embryogenesis and green plant regeneration from isolated microspores of hexaploid triticale (Triticosecale Wittmack) cv. Bogo. Plant Cell Rep 22:885–893. https://doi.org/10.1007/s00299-004-0796-9

Orłowska R (2021) Barley somatic embryogenesis—an attempt to modify variation induced in tissue culture. J Biol Res 28:9. https://doi.org/10.1186/s40709-021-00138-5

Orłowska R, Bednarek PT (2020) Precise evaluation of tissue culture-induced variation during optimisation of in vitro regeneration regime in barley. Plant Mol Biol 103:33–50. https://doi.org/10.1007/s11103-020-00973-5

Orłowska R, Pachota KA, Andrasiuk P, Bednarek PT (2022) Triticale green plant regeneration is due to DNA methylation and sequence changes affecting distinct sequence contexts in the presence of copper ions in induction medium. Cells 11:84

Orłowska R, Pachota KA, Dynkowska WM, Niedziela A, Bednarek PT (2021a) Androgenic-induced transposable elements dependent sequence variation in barley. Int J Mol Sci 22:6783. https://doi.org/10.3390/ijms22136783

Orłowska R, Pachota KA, Machczyńska J, Niedziela A, Makowska K, Zimny J, Bednarek PT (2020) Improvement of anther cultures conditions using the Taguchi method in three cereal crops. Electron J Biotechnol 43:8–15. https://doi.org/10.1016/j.ejbt.2019.11.001

Orłowska R, Zimny J, Bednarek PT (2021b) Copper ions induce DNA sequence variation in zygotic embryo culture-derived barley regenerants. Front Plant Sci 11:64837–64837. https://doi.org/10.3389/fpls.2020.614837

Pachota KA, Orłowska R, Bednarek PT (2022) Medium composition affects the tissue culture-induced variation in triticale regenerants. Plant Cell Tissue Organ Cult. https://doi.org/10.1007/s11103-022-02327-z

Padmaja G, Reddy V, Reddy G (1992) Somatic embryogenesis and plant regeneration from mature embryo callus. Indian J Exp Biol 30:181–184

Paladi RK, Rai AN, Penna S (2017) Silver nitrate modulates organogenesis in Brassica juncea (L.) through differential antioxidant defense and hormonal gene expression. Sci Hortic 226:261–267. https://doi.org/10.1016/j.scienta.2017.08.038

Patzak J, Henychová A, Svoboda P, Malířová I (2021) Evaluation of methylation epigenetics changes in micropropagated hop plants. Acta Hortic 1328:17–22. https://doi.org/10.17660/ActaHortic.2021.13282

Pecztsch AR (2020) The genomics of oxidative DNA damage, repair, and resulting mutagenesis. Comp Struct Biotechnol J 18:207–219. https://doi.org/10.1016/j.csbj.2019.12.013

Przetakiewicz A, Orczyk W, Nadowska-Orczyk A (2003) The effect of auxin on plant regeneration of wheat, barley and triticale. Plant Cell Tissue Organ Cult 73:245–256. https://doi.org/10.1023/A:1023030518100

Purnhauser L, Gyulai G (1993) Effect of copper on shoot and root regeneration in wheat, triticale, rape and tobacco tissue cultures. Plant Cell Tissue Organ Cult 35:131–139. https://doi.org/10.1007/bf00032962
Purnhauser L, Medgyesy P, Czakó M, Dix PJ, Márton L (1987) Stimulation of shoot regeneration in Triticum aestivum and Nicotiana plumaginifolia Viv. tissue cultures using the ethylene inhibitor AgNO₃. Plant Cell Rep 6:1–4. https://doi.org/10.1007/bf00269725

Rojas-Lorz L, Arrieta-Espinoza G, Valdez-Melara M, Pereira LFP, Gatica-Arias A (2019) Influence of silver nitrate on somatic embryogenesis induction in Arabica coffee (Coffea arabica L.). Braz Arch Biol Technol 62

Sato K (2020) History and future perspectives of barley genomics. DNA Res 27:dsaa023. https://doi.org/10.1093/dnares/dsaa023

Schmidt M, Walz C, Hesemann CU (1996) Somaclonal variation of the mitochondrial ATPase subunit 6 gene region in regenerated triticale shoots and full-grown plants. Theor Appl Genet 93:355–360. https://doi.org/10.1007/BF0023176

Taguchi G (1986) Introduction to quality engineering; designing quality into products and processes; White Plains, UNIPUB/quality resources; Dearborn, Mich., American Supplier Institute: NY p. 181

Thomas J, Raj Kumar R, Mandal AK (2006) Metabolite profiling and characterization of somaclonal variants in tea (Camellia spp.) for identifying productive and quality accession. Phytochemistry 67:1136–1142. https://doi.org/10.1016/j.phytochem.2006.03.020

Vega R, Vásquez N, Espinoza AM, Gatica AM, Valdez-Melara M (2009) Histology of somatic embryogenesis in rice (Oryza sativa cv. 5272). Rev Biol Trop 57:141–150

Vikrant RA (2001) Comparative study of somatic embryogenesis from immature and mature embryos and organogenesis from leaf-base of Triticale. Plant Cell Tissue Organ Cult 64:33–38. https://doi.org/10.1023/A:1010627630651

Warchol M, Juzoi K, Dziurka K, Czyczylo-Mysza I, Kaploniak K, Marcinińska I, Skrzypek E (2021) The effect of zinc, copper, and silver ions on oat (Avena sativa L.) androgenesis. Plants-Basel 10:248

Weigel R, Wolf M, Hesemann CU (1995) Mitochondrial DNA variation in plants regenerated from embryogenic callus cultures of CMS triticale. Theor Appl Genet 91:1237–1241. https://doi.org/10.1007/bf00220934

Wen F, Sorensen EL, Barnett FL, Liang GHL (2004) Callus induction and plant regeneration from anther and inflorescence culture of Sorghum. Euphytica 52:177–181

Wu LM, Wei YM, Zheng YL (2006) Effects of silver nitrate on the tissue culture of immature wheat embryos. Russ J Plant Physiol 53:530–534. https://doi.org/10.1134/S1021443706040157

Yruela I (2009) Copper in plants: acquisition, transport and interactions. Funct Plant Biol 36:409–430. https://doi.org/10.1071/FP08288

Zavattieri M, Frederico A, Lima M, Sabino R, Arnholdt-Schmitt B (2010) Induction of somatic embryogenesis as an example of stress-related plant reactions. Electronic Journal of Biotechnology. Electron J Biotechnol 13. https://doi.org/10.2225/vol13-issue1-fulltext-4

Zehr BE, Williams ME, Duncan DR, Widholm JM (1987) Somaclonal variation in the progeny of plants regenerated from callus cultures of seven inbred lines of maize. Can J Bot-Rev Can Bot 65:491–499. https://doi.org/10.1139/b87-061

Zhu JK (2009) Active DNA demethylation mediated by DNA glycosylases. Annu Rev Genet 43:143–166. https://doi.org/10.1146/annurev.genet-102108-134205

Zimny J, Rybczyński JJ (1985) Somatic embryogenesis of triticale. In: Horn W, Jensen CJ, Odenbach W, Schieder O (eds) Genetic manipulation in plant breeding: Proceedings International Symposium Organized by EUCARPIA, September 8–13, 1985, Berlin (West), Germany. De Gruyter, pp 507–510. https://doi.org/10.1515/9783110871944-091

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