Possible protective effects of trans-resveratrol against 1,4-dioxane induced toxicity in meristematic cells of plant bioassays

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Abstract

Background In this study, the protective effects of trans-resveratrol against 1,4-dioxane induced toxicity in meristematic cells were investigated. For this purpose, Allium test which is a reliable test was used and the alterations in all experimental groups were examined by using physiological, cytogenetic, biochemical and anatomical parameters.

Results As a result of the analysis, it has been determined that 1,4-dioxane causes serious abnormalities in Allium cepa meristematic cells. It was determined that in 1,4-dioxane treated group germination percentage regressed 1.6-times, root length reduced 12.7-times and weight gain decreased 7.7-times compared to control group. It has been observed that trans-resveratrol administration with 1,4-dioxane causes improvement in physiological parameters and reduces the damage rate from 0.4 to 0.16. Mitotic index, micronucleus and chromosomal abnormality frequency were investigated as cytogenetic parameters. It was determined that mitotic index decreased, chromosomal abnormalities and micronucleus frequency increased in 1,4-dioxane treated group. Trans-resveratrol treatment was found to cause a dose dependent improvement in genotoxic effects induced by 1,4-dioxane. Changes in the antioxidant system in all experimental groups were determined by measuring malondialdehyde, glutathione, superoxide dismutase and catalase enzyme levels. It was determined that 1,4-dioxane administration caused an increase in malondialdehyde level, decreased glutathion rate and induced antioxidant enzyme activity. Trans-resveratrol application was found to cause improvement in these alterations induced by 1,4-dioxane. It was observed that the 200 mg/mL trans-resveratrol+1,4-dioxane treatment caused a 1.9-fold decrease in malondialdehyde level which is indicator of lipid peroxidation compared to only 1,4-dioxane treated group. The abnormalities caused by 1,4-dioxane application in the meristematic cells are also found in the anatomical structure. In 1,4-dioxane treatment group, anatomical changes such as cell deformation and cortex wall thickening were observed. The frequency of these changes decreased with trans-resveratrol administration.

Conclusions As a result, it was determined that 1,4-dioxane caused a versatile toxicity in Allium cepa meristematic cells, while trans-resveratrol was found to have a dose-dependent protective feature
against 1,4-dioxane-induced toxicity.

Background
With the increasing industrialization, the use of chemicals has increased in various sectors and the use of many additives to improve the properties of industrial products has become widespread. 1,4-dioxane, which is used to fix solvents with its hydrophilic and anti-chlorination properties, is frequently used in personal care and cleaning products, varnishes, paints and disinfectants. 1,4-dioxane which is also present in the structure of adhesives and inks, is also used in the manufacture of insecticides, herbicides and softeners [1]. Due to its frequent use in many products, contamination to the environment and people has also become inevitable. Especially in agricultural areas, direct use as an insecticide or as a component of pesticides leads to serious contamination. As a result of its use as an insecticide, it accumulates in agricultural areas, pollutes water environments and shows toxic effects on non-target organisms such as plants. In particular, accumulation in plants causes 1,4-dioxane to be transmitted to other living organisms and humans through the food chain. 1,4-dioxane has been reported to cause damage to the liver and kidneys as exposure time and dose increase and is included in the list of carcinogenic substances [2,3]. However, in the literature, both negative and positive results were obtained in terms of genotoxic effect in 1,4-dioxane-related studies. Roy et al. [4] reported that there was no genotoxic effect of 1,4-dioxane, while the Morita and Hayashi [5] stated that 1,4-dioxane is mutagenic in Ames test. In this study, 1,4-dioxane toxicity was investigated with different parameters and a contribution has provided to the solution of the complexity of 1,4-dioxane toxicity in the literature. It was also tested whether trans-resveratrol (t-resv) could neutralize 1,4-dioxane toxicity. Powerful antioxidant compounds can be used to reduce or neutralize the harmful effects of toxic substances that we are exposed to in daily life through diet or different contamination. However, considering the diversity in plant species and herbal products, such studies are inadequate. In this study, protective properties of t-resv against 1,4-dioxane toxicity were investigated with multiple parameters.

Resveratrol (3,4′,5-trihydroxy-trans-stilbene) is a phytoalexin commonly found in consumable foods such as grape, peanuts and red wine [6]. Resveratrol is known to have properties such as lipid
peroxidation inhibitory, copper chelating, radical scavenging, anti-inflammatory activity and anti-carcinogenic activity [7]. Resveratrol has been reported to be protective against many diseases such as arteriosclerosis and cancer in the literature and this feature has been explained by many mechanisms. Activating the antioxidant system, facilitating antioxidant mechanisms, suppressing the formation of superoxide radical and $\text{H}_2\text{O}_2$, inhibiting lipid peroxidation induced by reactive oxygen are some of these mechanisms [8,9]. These mechanisms may be in the form of a specific protection against direct toxic effect or indirect protection by activating other mechanisms. Resveratrol has two geometric isoforms, cis-resveratrol and trans-resveratrol. And the main form, the trans isomer concentration, contributes significantly to its biological activity [7]. Resveratrol is a polyphenol with antioxidant properties, while t-resv is a polyphenolic phytoalexin with anti-inflammatory, antioxidant and anticancer properties. The trans-resveratrol has a planar backbone, while the resveratrol comprises two main planes. Due to its structural properties, t-resv is a form associated with a potential protective property. The protective properties of t-resv have been demonstrated by various studies in the literature. T-resv is reported to inhibit metal-induced lipid peroxidation and to reduce lipid peroxidation caused by ethanol [10,11]. In addition, it has been reported that due to its antioxidant properties, it has a cardioprotective effect against ischemia [12]. In the literature, the protective properties of cis-resveratrol or trans-resveratrol against various agents have been investigated, but no protective properties of t-resv against 1,4-dioxane toxicity have been investigated. In this study, the dose-dependent protective properties of t-resv against biochemical, genetic, physiological and anatomical toxicity induced by 1,4-dioxane were investigated.

Results

Germination Parameters

In this study, the toxic effects of 1,4-dioxane and the ameliorative effects of t-resv against this toxicity have been investigated in Allium cepa which is used as a model organism in toxicity tests. Firstly, effects on germination and germination related parameters were examined and the results were given in Table 1. No abnormalities were detected in the germination parameters of Group I, Group II and Group III, while similar germination rate, root length and weight gain levels were found between
the groups. Root length, germination rate and weight gain decreased by 12.6, 1.6 and 7.7 times, respectively in 100 mg/L 1,4-dioxane treated group compared to control group. T-resv treatment with 1,4-dioxane in Group V and Group VI showed an improvement in germination parameters, but the levels were far behind the control group. In Group VI where 200 mg/L t-resv and 1,4-dioxane were applied together, germination rate was 1.4 times higher than in 1,4-dioxane treated group and 1.19 times lower than in control group. When the relative injury rates were examined, the highest injury was observed in Group IV as 0.40, whereas it was found to decrease with the application of 100 mg/L and 200 mg/L t-resv. In summary, 1,4-dioxane inhibited all parameters related to germination in the model organism and caused significant relative injury rate. However, it was determined that t-resv treatment regressed the negative effects of 1,4-dioxane and caused significant improvements. It was observed that these improvements increased as t-resv dose increased and more dominant at 200 mg/L.

**Antioxidant-oxidant balance**

1,4-dioxane induced changes in Glutathion (GSH) and malondialdehyde (MDA) levels, which are parameters related to oxidative stress, were given in Fig. 1. For both parameters, there was no significant difference between the control group, 100 mg/L and 200 mg/L t-resv treated groups (p>0.05). The fact that MDA level was increased 5.09-fold in 1,4-dioxane treated group compared to control group indicates lipid peroxidation and oxidative stress formation in meristematic cells. It was determined that this oxidative damage induced by 1,4-dioxane regressed with t-resv application. In group V and VI, it was found that t-resv treatment with 1,4-dioxane caused an significantly improvement in MDA and GSH levels (p>0.05). This improvement was found to increase with increasing t-resv dose, but there was no direct proportional increase. The most significant protection against oxidative damage was observed at a dose of 200 mg/mL t-resv treatment. It was determined that MDA level decreased 1.93 fold and GSH level increased 1.85 fold in the 200 mg/mL t-resv+dioxane treated group compared to Group IV and these changes showed the protective role of t-resv against oxidative stress.

Oxidative stress in cells is removed by enzymatic antioxidant system. Superoxide dismutase (SOD)
and catalase (CAT) are two important enzymes for the removal of radical products, and no change in the activity of these enzymes was found in the control group and the groups treated with only t-resv (Fig. 2). It was found that 1,4-dioxane treatment caused changes in SOD and CAT activity in meristematic cells and enzyme activities increased 2.1 and 2.6 times, respectively, compared to control group. This increase is a response to oxidative damage evidenced by increased MDA and decreased GSH content by 1,4-dioxane effect. The enhancing effect of 1,4-dioxane on SOD and CAT enzymes began to normalize with t-resv application. As a result of t-resv application with 1,4-dioxane, SOD and CAT activities were observed to approach the control group levels, but the levels remained above the control group. Increased SOD activity in the group treated with 1,4-dioxane decreased 34.7% in the group receiving 200 mg/mL t-resv with 1,4-dioxane. CAT activity in the same group decreased by 47.4% compared to Group IV.

**Cytogenetic parameters**

The frequency of chromosomal abnormalities (CAs) induced by 1,4-dioxane in the meristematic cells were given in Table 2 and Fig. 3. Only a few unequal distribution of chromatine and vagrant chromosome were observed in Group I, II, and III, but these abnormalities were not statistically significant (p>0.05). In 1,4-dioxane treated group, chromosomal abnormalities such as fragment, sticky chromosome, unequal distribution of chromatine, bridge and vagrant chromosome were observed with a high level. While the highest fragment frequency was observed among CAs, vagrant chromosome formation was observed to be low compared to other CAs. It was determined that the t-resv application with 1,4-dioxane decreased the frequency of CAs formations. Briefly, t-resv exhibited dose-dependent protective properties, but this protective property was not directly proportional to the dose. In 200 mg/L t-resv+dioxane treated group, fragment frequency (the highest CAs type) observed in this study was found to be reduced by 40.0% compared to the 1,4-dioxane-treated group. Abnormalities observed in Group V and Group VI were still higher than in the control group, so t-resv reduced the toxic effect of 1,4-dioxane but could not completely neutralize it.

Micronucleus (MN) frequency and mitotic index (MI) ratios of all treatment groups were given in Fig 4. Very low MN formation was detected in the control group and the groups receiving only t-resv, and
there was no significant difference between these groups (p>0.05). This result shows that t-resv does not cause the formation of MN at the doses tested. Similarly, a parallel level was obtained in MI ratios for all three groups and no difference was observed between Groups I, II and III. It was determined that MN ratio increased significantly and reached 49.00 ± 3.39 in 1,4-dioxane treated group. In Group IV, the MI rate was significantly redounded and the MI rate was reduced by 50.1% compared to the control group. The application of t-resv resulted in improvement in 1,4-dioxane induced MN formation and abnormalities in MI. It was determined that 100mg/L and 200mg/L t-resv treatment in Group V and VI reduced MN formation by 1.58 and 2.13 times, respectively compared to only 1,4-dioxane treated group. T-resv showed similar amerolative effect in MI rates and the most significant improvement was achieved in Group IV treated with 200 mg/L t-resv. It was determined that 100 mg/L and 200 mg/L t-resv application with 1,4-dioxane increased MI ratio by 1.50 and 1.79 times, respectively compared to only 1,4-dioxane treated group.

**Anatomical changes**

The anatomical changes caused by the application of 1,4-dioxane in *A.capa* meristematic cells were shown in Fig 5. While no abnormality was observed in the anatomical structures of Group I, Group II and Group III, various anatomical changes were observed in the 1,4-dioxane treated group. These changes are flattened cell nuclei, cell wall thickening and cell deformation. In the root tissue where the nutrient uptake of the plant is provided, the epidermis cells are the place where the first contact with exogenous substances is provided. The fact that 1,4-dioxane application leads to deformation in epidermis cells confirms this hypothesis. It was found that the frequency of 1,4-dioxane induced anatomic abnormalities decreased with t-resv administration (Table 3). Anatomical improvements were more pronounced in Group VI treated with 200 mg/mL t-resv.

**Discussion**

In this study, it has been determined that 1,4-dioxane causes a versatile toxic effect in *A.capa* meristematic cells, which is a eukaryotic model organism, and t-resv reduces this toxicity. In 1,4-dioxane treated group significant reductions in radicle lenght, weight increase and germination rates were determined and high realtive injury rate was observed. Abnormalities observed in germination
parameters may be related to toxic effects of 1,4-dioxane. Dioxane has direct or indirect toxic effects on living organisms. The direct effect occurs as a result of the reaction of 1,4-dioxane and molecular oxygen, which causes the production of free radicals and the formation of oxidative stress \([13,14]\). Oxidative stress in plants causes autocatalytic peroxidation of membrane lipids and pigments, modification of membrane permeability and consequently damage to cell structure \([15]\). In particular, the rapidly dividing meristem cells are more affected by oxidative stress, and root growth is inhibited and this inhibition is reflected in germination and weight gain. Similarly, it has been reported in the literature that application of 1,4-dioxane causes inhibition in weight gain, germination rate and root growth of test organisms \([1,16]\). T-resv showed an ameliorative effect against 1,4-dioxane induced damage in germination parameters. Resveratrol is present in cis and trans isomeric form, and the concentration of the trans isomer, the main form, significantly contributes to biological activity \([7]\). In the literature, resveratrol is reported to be a potent antioxidant component that has been proven in various in vitro and in vivo studies. Chanbitayapongs et al. \([11]\) reported that resveratrol inhibited metal-derived lipid peroxidation and showed antioxidant properties in in-vitro studies. Sun et al. \([10]\) reported that resveratrol reduced lipid peroxidation induced by iron and ethanol. Since there is no data on the protective properties of t-resv against 1,4-dioxane toxicity in plants, this study is the first data on this subject.

MDA, GSH, SOD and CAT analyzes were performed to determine the effects of dioxane and resve applications on antioxidant and oxidan balance. It was determined that dioxane application increased MDA level, decreased GSH level and induced antioxidant enzyme activities. It was found that these changes observed in antioxidant-oxidant balance started to normalize with resveratrol application. It is known that 1,4-dioxane causes free radical production and oxidative stress in living systems. Free radicals in the cells attack the unsaturated lipids containing carbon-carbon double bonds, causing lipid peroxidation. Low levels of MDA have been reported to act as regulators of gene expression in cells. However, high levels of MDA can easily interact with functional groups of molecules such as proteins, lipoproteins, DNA and RNA in the cell, causing adduct formation and different pathological conditions \([17]\). MDA is highly toxic and its toxicity is associated with Michael's ability to form adducts
with thiol groups, facilitate protein cross-linking and induce mutagenesis [18]. Along with the increase in MDA, changes in glutathione levels were observed in meristematic cells treated with 1,4-dioxane. It was determined that glutathione level which is an antioxidant with tripeptide structure decreased 2.38 times in 1,4-dioxane treated group compared to control group. Briefly, the increase in MDA level and the decrease in GSH level in the same cells treated with 1,4-dioxane are indicative of obvious oxidative damage. Glutathions have been reported to be found in almost all cell parts of plant tissues such as cytosol, endoplasmic reticulum, vacuole, mitochondria, chloroplast, peroxisome. Reduced glutathione reacts with lipid peroxides and oxidizes during detoxification of these molecules and the level of reduced glutathione decreases [19]. In cells, the decreased glutathione level leads to reduced antioxidant capacity and the increased MDA level causes an enhanced oxidation. The decrease in GSH level and the increase in MDA level indicate the deterioration of antioxidant-oxidant balance. It was observed that this disrupted balance induced by 1,4-dioxane, started to improve with t-resv application. Resveratrol has scavenging activity on reactive oxygen species and shows significant effects on the radical induced cellular response [20]. Similarly, Al-Hussaini and Kilarkaje [21] reported that lipid peroxidation and macromolecule oxidation in the cell decreased with t-resv administration. In another study, Mikstacka et al. [22] reported that t-resv administration was effective in recovering significantly depleted GSH content in tested cells. Gupta et al. [23] reported that in oxidative stress-induced subjects the administration of 20 and 40 mg/kg t-resv administration resulted in a reduction in MDA levels, but no effect on GSH levels.

It has also been determined that 1,4-dioxane application induces SOD and CAT activities in A. cepa meristematic cells. SOD and CAT are involved in the removal of radical products in the cell. SOD catalyzes the dismutation of the highly reactive superoxide anion to O$_2$ and the less reactive product H$_2$O$_2$ and the peroxide which is formed as a result of this reaction is destroyed by CAT enzyme [24]. Although there are studies showing that 1,4-dioxane causes oxidative damage and increases MDA level [25], there is no direct study investigating its effect on SOD and CAT activity. However, there are many studies in the literature that report changes in SOD and CAT activity in the presence of
oxidative stress. Malar et al. [26] observed that in the presence of induced lipid peroxidation and oxidative stress SOD activity increased by 251% and CAT activity increased by 60% in *Eichhornia crassipes* leaf tissues compared to control. Macczak et al. [27] reported a reduction in GSH levels and alterations in SOD and CAT activities in the presence of oxidative stress caused by various chemicals inducing the formation of reactive oxygen species. In this study, t-resv treatment with dioxane ameliorated the enhanced SOD and CAT activity induced by 1,4-dioxane. These improvements can be explained by the role of t-resv in reducing oxidative stress and regulating enzyme induction. Similarly, Sadi et al. [28] specified that SOD activity increased in the presence of induced oxidative stress and decreased again after t-resv treatment, briefly they reported that t-resv treatment caused an improvement in the level of oxidative biomarkers. Pintea et al. [29] reported that t-resv directly contributes to antioxidant defense by scavenging reactive oxygen species and causing changes in superoxide dismutase, catalase and glutathione peroxidase activities.

1,4-dioxane, which causes oxidative stress in *A. cepa* meristematic cells, has also been found to cause chromosomal abnormalities such as fragment, sticky chromosome, unequal distribution of chromatine, bridge and vagrant chromosome. These genotoxic effects indicate that 1,4-dioxane disrupts genome stability, and these effects may be associated with 1,4-dioxane-induced oxidative damage. 1,4-dioxane leads to free radical formation, oxidative stress and lipid peroxidation in living systems [13,14]. The effects of oxidative stress on DNA have been investigated in detail and have been demonstrated by many studies in literature. Oxidative stress and free radicals cause DNA adducts, phosphodiester bond cleavages, chain breaks and mutations in bases [30]. Abnormalities in bases caused by 1,4-dioxane exposure cause A:T-T:A transversions and this mutation results from adenosine adducts [31,32]. All these changes in the DNA structure induced by 1,4-dioxane cause genome instability and the formation CAs. Similarly, Sağır et al. [16] reported that the application of 1,4-dioxane resulted in high CAs formations such as fragment, ascentric and dicentric chromosomes. In this study, it was also determined that genotoxic effects induced by 1,4-dioxane were decreased with t-resv treatment. Oxidative stress caused by 1,4-dioxane in organisms has been proved by the increased MDA levels in *A. cepa* root cells. Oxidative stress in cells also affects DNA and causes the
formation of DNA base oxidation products. Agents that increase oxidative DNA damage increase cancer development. Inhibiting the formation of oxidative stress, which is the starting point of these abnormalities, will also inhibit mutation and cancer development. It has been shown in many studies that consumption of antioxidant-containing plants in daily diet reduces oxidative DNA damage levels and the incidence of human cancers [33]. Resveratrol increases the expression of glutathione peroxidase and catalase enzymes and induces scavenging of $\text{H}_2\text{O}_2$, thereby providing resistance to oxidative damage [34]. Agents such as resveratrol that reduce oxidative stress and consequently DNA damage have antimutagenic and anti-cancer effect. Jang et al. [35] reported that resveratrol has an anticarcinogenic effect and this effect is associated with inhibition of the initiation and promotion of carcinogenesis. Al-Hussaini and Kilarkaje [21] reported that DNA oxidation in the cell decreased in subjects treated with t-resv. Ungvari et al. [34] showed that DNA damage induced by oxidative stressor was reduced by $10^{-6}$–$10^{-4}$ mol/L resveratrol application in a cell culture.

The cytogenetic effects of 1,4-dioxane and t-resv applications in meristematic cells were also supported by MN and MI analysis. It has been proven by the increase in the frequency of MN and the decrease in MI rate that 1,4-dioxane causes genomic instability. The decrease in the MI ratio of 1,4-dioxane treated group can be explained by the oxidative stress created in the meristematic cell.

Chemicals that trigger the formation of oxidative damage and cause glutathione reduction are reported to delay cell cycle by causing delayed progression through G1 and S phases and even stopping the cell cycle at the G$_2$ point [36]. The fact that the 1,4-dioxane application causes an increase in MDA level, a decrease in gsh level and changes in antioxidant enzyme activities in meristematic cells is evidence of oxidative damage and this damage causes an inevitable delay in cell cycle. Similarly, Sağır et al. [16] reported that 1,4-dioxane administration caused a decrease in the MI rate and reduced the number of dividing cells from $835\pm45.38$ to $438.5\pm23.31$. The decrease in MI ratio also explains the abnormalities caused by 1,4-dioxane in physiological parameters. Root growth, germination and weight gain are directly related to cell division, and the decrease in division rate negatively affects these parameters. DNA damage and MN formation also cause disruptions and
abnormalities in the cell cycle. The formation of MN in a cell is an indicator of toxic effect and MN is caused by all chromosomes or chromosome fragments that do not belong to the main nucleus. MN is usually caused by abnormalities in the mitotic spindle, kinetocor or mitotic apparatus and chromosomal damage [37,38]. The fact that MN induction in meristematic cells indicates that 1,4-dioxane causes mitotic abnormalities and has genotoxic effect. Similarly, Teker et al. [1] reported that 1,4-dioxane treatment caused an increase in MN frequency in root tip cells in a dose dependent manner and they achieved a MN level of 46.70±11.91 in the group treated with 100 ppm 1,4-dioxane. In this study, the protective properties of t-resv proven in previous parameters were also observed in MN and MI analysis. T-resv application with 1,4-dioxane was found to cause improvement in MN and MI rates. This protective property of t-resv can be explained by its reducing effect against oxidative damage induced by 1,4-dioxane. In the previous analysis of this study, t-resv was found to reduce the rate of increased MDA and regulate the GSH level, thereby reducing oxidative damage. These results are the evidence that t-resv administration is protective against oxidative stress and same amerolative effect was also observed in MN and MI results. Similarly, Ranjini and Manonmani [39] reported that 100 µM resveratrol treatment has protective effects against induced toxicity and has a reducing effect on MN formation and an an enhancing effect on cell viability.

The application of 1,4-dioxane has been found to cause changes such as flattened cell nuclei, cell wall thickening and cell deformation in the anatomical structure of A. cepa root. Root tissue can develop various adaptation mechanisms against toxic agents. Thickening of the cortex cell wall is one of the adaptation mechanisms, thereby reducing the access of the toxic agent to the internal tissues and the central cylinder [40,41]. In addition to the general structure of the cell, anatomical changes were observed in the cell nucleus. Flattening of the cell nucleus was observed with 1,4-dioxane application and this was associated with the cumulative effect of genotoxic and biochemical changes caused by 1,4-dioxane. A possible change in intracellular pressure and cell skeleton as a result of toxicity that disrupts the overall integrity of the cell may lead to shape changes of the organelles. The cell nucleus is generally spherical or ellipsoidal, but may change its shape in response to intracellular changes. Alterations in nuclear volume and protein concentration, degradation in DNA integrity and density
may result in nuclear shape changes [42,43]. 1,4-dioxane induced peroxidation in membrane component lipids, disruption of DNA integrity by MN and CAs formations are the possible explanation for nuclear shape changes in Group IV. Although there is no study on the effect of 1,4-dioxane on anatomical damage, many studies have shown that many toxic agents cause changes and necrosis in epidermis and cortex cells [44,45]. With previous analyzes of this study it was found that 1,4-dioxane induced genotoxic effects and abnormalities in antioxidant system were found to be reduced by t-resv. Ameliorative effects of t-resv in other parameters have also been shown against anatomical damages. treatment with t-resv caused a significant decrease in anatomical changes especially in the frequency of flattened cell nucleus. Similar, Macar et al. [45] reported that 400 mg/L and 800 mg/L resveratrol application alleviate the anatomic damages in meristematic cells induced by CuCl₂.

Conclusion
In this study, 1,4-dioxane, which has been used for various purposes in many sectors, has been shown to cause toxic effects on frequently dividing meristematic cells and it has been shown that t-resv has a ameliorative effect against these toxicity. It has been determined that 100 mg/L 1,4-dioxane induces genotoxic effects, causes a regression in germination and growth and serious changes in anatomical structure of meristematic cells. However, it has caused a change in antioxidant enzyme levels, a decrease in GSH level and an increase in MDA level, which is an indicator of lipid peroxidation. By altering the oxidant-antioxidant balance that is in equilibrium in living things, it decreased the cell's strength against oxidative damage. T-resv was found to have a ameliorative effect against all these 1,4-dioxane-induced abnormalities and this effect was dose-dependent. 200 mg/L t-resv, one of the two tested doses, has a higher protective activity. However, it was also found that all the parameters tested were correlated and that the abnormalities observed in one parameter triggered other abnormalities. The decrease in MI rate naturally affect the root growth, weight gain and germination percentage. High concentrations of MN and CAs were observed in 1,4-dioxane-treated group, where abnormalities were observed in the antioxidant system. The improvement in antioxidant parameters with t-resv administration decreased the genotoxic effects. In short, all the data obtained in the study support each other and it is possible to elucidate the mechanism of toxicity.
with a multi-parameter study. Considering the amerolative effects of t-resv against genotoxic, physiological and biochemical toxicity, the importance of natural antioxidant foods emerges. Although there are many studies investigating the multiple biological properties of antioxidant plant foods, these studies are insufficient considering the high plant diversity. For this reason, studies on this subject will contribute to the literature and will guide other studies.

**Methods**

**Test materials and grouping principles**

1,4-dioxane was purchased from Sigma-Aldrich and the other chemicals were supplied from Sigma-Aldrich. *Allium cepa* bulbs were used as target organism for determining the 1,4-dioxane toxicity. The identification of plant materials used in this study was applied by Dr. Zafer Türkmen from GRU Botanic Department. An example of plant material was deposited in a herbarium located in Botanical laboratory (GRU-FEF-23/2020). *A. cepa* bulbs were divided into six groups, one control and five application groups. Each group contained 6 bulbs and experimental stages given in Table 4. In each group bulbs were sterilized and germinated with related solution in individually controlled tubes at 24 °C for 72 hours.

**Germination parameters**

The effects of 1,4-dioxane and t-resv on physiological parameters were investigated by germination percentage, root length, weight gain and injury rate analysis. Germination percentage (GP) and Relative injury rate were calculated by using the Equation 1 and 2 [46]. Root length analysis was based on 10 root lengths and a mean value was calculated by random measurement from the ampoules of each group. For the weight gain analysis, the initial weight of each bulb and the weight at the end of the application were measured and the weight gain was determined using the difference between the first and last weight.

\[
\text{GP (\%)} = \left[ \frac{\text{Number of germinated bulb}}{\text{Total number of bulb}} \right] \times 100
\]  
(1)

\[
\text{Relative injury rate} = \frac{\text{\%GP in control} - \text{\%GP in each group}}{\text{\%GP in control}}
\]  
(2)

**Cytogenetic parameters**

The effects of 1,4-dioxane and t-resv on MI, MN and CAs frequency and MI levels were determined by using root tip preparations. After the germination period, healthy root tips from each ampoule were
subjected to fixation procedures in serial ethanol solutions. After fixation, the root ends which were hydrolyzed with 1N HCl were stained with acetocarmine and crushing preparations were prepared. Entellan-fixed preparations were examined using a research microscope [47]. 1000 for MN and CA frequency analysis and 10000 for MI analysis were counted. MI was calculated using Equation 3.

\[
\text{MI (\%)} = \frac{\text{Cell number in mitosis}}{\text{Total cell number}} \times 100
\]

(3)

**Antioxidant-oxidant balance**

The effects of 1,4-dioxane and t-resv on the antioxidant system were evaluated by examining MDA level and GSH, SOD and CAT levels, which are antioxidant parameters. Samples (0.5 g) from the meristematic tissues were homogenized in sodium phosphate buffer before all analyzes. GSH levels in homogenates were measured by acid soluble sulfhydryl level determination as described by Vecchia et al. [48]. The level of MDA, an indicator of oxidative stress was measured by a protocol reported by Unyayar et al. [49]. SOD activity was determined according to the procedure developed by Beauchamp and Fridovich [50] and expressed as U / mg FW. CAT activity was calculated according to the method proposed by Beers and Sizer [51] and the activity was expressed as OD240 nm / min.

**Anatomical changes**

All anatomical changes were examined by taking cross-sections from root tip. Cross-sections taken from the root tip of each group were stained with methylene blue by routine staining procedure and the preparations were fixed with entellan. The anatomical structures of each group were examined and a total of 1000 cells were examined for the frequency of anatomical changes [52].

**Statistical analysis**

Statistical analyzes were performed using the “IBM SPSS Statistics 22 SP” package program. Data were shown as mean ±SD (standard deviation). The statistical significance between the means was determined by One-way ANOVA and Duncan's test and p value <0.05 was considered statistically significant.

**Abbreviations**

CAs Chromosomal abnormalities

CAT Catalase
GP    Germination Percentage
GSH   Glutathion
MN    Micronucleus
MI    Mitotic index
MDA   Malondialdehyde
SOD   Superoxide dismutase
T-RESV Trans-resveratrol

Declaration

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Not applicable.

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Authors’ contributions
Initiated and designed the research: EY, KÇ
Performed the experiments: DK, EY, KÇ
Analyzed the data: DK, EY, KÇ
Wrote the paper: EY, KÇ
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Not applicable
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Tables
Table 1 The effects of 1,4-dioxane and the ameliorative effects of t-resv on germination parameters

| Groups | GP (%) | Relative Injury Rate | Root length (cm) | Final weight (g) | Weight Gain (g) |
|--------|--------|----------------------|-------------------|------------------|-----------------|
| Group I | 100    | 0.00                 | 9.50±1.64         | 11.00±1.43       | 6.00            |
| Group II | 100   | 0.00                 | 9.63±1.47         | 11.13±1.63       | 6.13            |
| Group III | 100   | 0.00                 | 9.70±1.68         | 11.36±1.61       | 6.32            |
| Group IV | 60    | 0.40                 | 0.75±0.31         | 5.83±0.42        | 0.78            |
| Group V  | 70    | 0.30                 | 2.00±0.63         | 6.57±0.51        | 1.50            |
| Group VI | 84    | 0.16                 | 4.50±0.80         | 8.47±0.88        | 3.45            |

Group I: Tap water, Group II: 100 mg/L t-resv, Group III: 200 mg/L t-resv, Group IV: 100 mg/L 1,4-dioxane, Group V: 100 mg/L 1,4-dioxane+100 mg/L t-resv, Group VI: 100 mg/L 1,4-dioxane+200 mg/L t-resv. Different letters in the same column indicate statistical significance.

Table 2 The effects of 1,4-dioxane and t-resv treatment on CAs frequency
Group I: Tap water, Group II: 100 mg/L t-resv, Group III: 200 mg/L t-resv, Group IV: 100 mg/L 1,4-dioxane, Group V: 100 mg/L 1,4-dioxane+100 mg/L t-resv, Group VI: 100 mg/L 1,4-dioxane+200 mg/L t-resv. Different letters in the same column indicate statistical significance. FRG: fragment, SC: sticky chromosome, UDC: unequal distribution of chromatine, B: bridge, VC: vagrant chromosome.

### Table 3 Frequency of anatomical changes in all experimental groups

|                          | Group I | Group II | Group III | Group IV | Group V | Group VI |
|--------------------------|---------|----------|-----------|----------|---------|----------|
| Thickening in cell wall of cortex | -       | -        | -         | +++      | +++     | ++       |
| Flattened cell nucleus   | -       | -        | -         | +++      | ++      | +        |
| Epidermis cell deformation| -       | -        | -         | ++++     | +++     | ++       |
| (-): No change, (+): mild change, (++): moderate change, (+++): severe change, (+++++): serious change |

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Table 4 Experimental groups
| Group     | Treatment                                         |
|-----------|--------------------------------------------------|
| Group I   | Tap water                                        |
| Group II  | 100 mg/L t-resv                                 |
| Group III | 200 mg/L t-resv                                 |
| Group IV  | 100 mg/L 1,4-dioxane                            |
| Group V   | 100 mg/L 1,4-dioxane + 100 mg/L t-resv          |
| Group VI  | 100 mg/L 1,4-dioxane + 200 mg/L t-resv          |

Figures
Figure 1

The effects of 1,4-dioxane and t-resv treatment on MDA and GSH levels

Figure 2

The effects of 1,4-dioxane and t-resv treatment on SOD and CAT activities
Figure 3

CAs and MN formations induced by 1,4-dioxane a MN. b fragment. c sticky chromosome. d unequal distribution of chromatine. e bridge. f vagrant chromosome

Figure 4

The effects of 1,4-dioxane and t-resv treatment on MN frequency and MI
Figure 5

Anatomical changes in A. cepa meristematic cell induced by 1,4-dioxane a epidermis cells in control. b cortex cells in control. c normal view of cell nucleus. d epidermis cell deformation in 1,4-dioxane treated group. e cortex cell wall thickening in 1,4-dioxane treated group. f flattened cell nucleus in 1,4-dioxane treated group