Inhibition of browning problem during the callogenesis of Spartium junceum L.

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Abstract

During different phases of in vitro culture, plant tissues may be exposed to some stresses that never encounter in their natural habitats. The most significant stresses which interfere with in vitro culture are pathogenic contamination and browning disorder. Since browning sign is occurred during all phases of in vitro culture of Spartium junceum L., the present study was done preventing explants from browning during disinfection and callogenesis phases using exposure time of sterilants (ethanol 0, 30, 60 s and home bleach 0, 10, 15 min), antioxidant compounds (PVP 0.5%, Activated charcoal 0.1%, Curcumin 0.1%), Running water (30 and 60 min) plant growth regulators (2,4-D 0, 0.5, 1 and 2 mg L\(^{-1}\) and BA 0, 0.1 and 0.2 mg L\(^{-1}\)), and by changing light/dark conditions was designed. The results showed that ethanol 70% (30 s) in combination with home bleach 20% (10 min) had the best effect in control contaminations and browning sign in nodal explants of S. junceum. The application of PVP 0.5% in medium was the best treatment to control of browning nodal explants in callus induction phase. The highest callus formation and the lowest explant browning were obtained on the medium supplemented with 0.5 mg L\(^{-1}\) 2,4-D under the darkness condition. According to the results of this study, how disinfection methods, culture medium compositions and light conditions were effective on the browning and callogenesis of Spartium junceum L.

Keywords: callus, disinfection, explant, PVP, woody plant.

Introduction

During different phases of in vitro culture, plant tissues may be exposed to some stresses that never encounter in their natural habitats. The most significant stresses which interfere with in vitro culture include plant pathogens contamination and browning (Taghizadeh et al., 2016). Browning disorder is often occurred in plant species containing high level of tannin and other hydroxid phenols. The wound due to cutting of tissue from stock plants induces polyphenol oxidase enzyme production which is crucial contributor to the oxidation of phenolic compounds, and...
accordingly makes tissues and culture medium brown and inhibits the explants growth or loss them (Meziani et al., 2016). One of the too serious difficulties in micropropagation process is in vitro pathogens; they cause infection and so loss the tissue culture process at early stages, even though sterilization methods are applied (Al Ghasheem et al., 2018). Therefore, the first step to start the tissue culture process involves disinfection and sterilization. There are a number of substances with mentioned property including variety of fungicides and antibiotics, sodium hypochlorite, calcium hypochlorite, ethyl alcohol, silver nitrate, mercury chloride, etc. (Ranjan et al., 2017). Among these, mercury chloride and silver nitrate have more powerful effects than others and have been applied frequently in past. Both mercury chloride and silver nitrate are of heavy metals group and have high reactivity to proteins which gives rise to block cell metabolism. Sodium hypochlorite and calcium hypochlorite make explants less injured because of their lower activity degree. However, they are able to induce necrotic explant and cell death in high concentration, and so cause browning of explants (Mahmoud and Al-Ani, 2016). In order to reduce browning degree, a number of approaches have been suggested (Krishna et al., 2008; Momeni et al., 2017).

*Spartium junceum* L. is a tall shrub-like ornamental plant from family Fabaceae. It has a characteristic specific to xerophilous plants (Gavilán et al., 2016). *In vitro* studies so far carry out on *Spartium junceum* L. involve *in vitro* culture the rooted micropropagated plantlet with the purpose of providing mycorhizal symbiosis via inoculating explant by *Arbuscular mycorhiza* during ex-vitro compatibility. Also, in order to investigate isoflavone compounds, the stem and nodal segments with axillary buds were cultured on medium MS containing 0.5 mg L⁻¹ BA. After one month, shoot proliferation was found (Clematisa et al., 2014). Since browning disorder is occurred during all phases of *in vitro* culture of *S. junceum* and causes plant material to be lose, the present study was done with the purpose of control and preventing explants from browning during disinfection and Callogenesis phases using exposure time of sterilants (ethanol and sodium hypochlorite), antioxidant compounds (PVP, activated charcoal, Curcumin), plant growth regulators (2,4-D and BA), and by changing light/dark conditions was designed.

**Materials and Methods**

The effect exposure times of Ethanol and home bleach treatments

For this experiment, nodal segments were used as explants of *S. junceum* L. At first, semi hard woody and juvenile shoots were cut from stock plant. These were cleaned with one drop of washing-up liquid and then rinsed in tap water for 20 min to remove any dust or pests. In order to optimize disinfection and sterilization conditions, nodal segment explants were submerged in ethanol 70% for three periods (0, 30 and 60 s), then the explants were placed into commercial sodium hypochlorite 20% (home bleaching solution, containing 5% active agent) for three periods (0, 10 and 15 min). After disinfection process was completed, explants were cut into 0.5 cm segments and were cultured. The callogenesis culture medium MS used in this experiment was based on some plant growth regulators treatments containing 2 mg L⁻¹ BA, 0.5 mg L⁻¹ IBA, 30 g L⁻¹ sucrose and 7 g L⁻¹ agar. The factors studied in this experiment included type and concentration of disinfectants (ethanol and home bleach) and the type of explants. It was a factorial experiment in a completely randomized design implemented with three replications for each treatment. After 7-10 days of culture, bacterial and fungal infections of any kinds and browning degree of explants were graded and recorded (Table 1 and Figure 1).

Table 1. The effect of Ethanol 70% and home bleach 20% treatments at various exposure times on different types of contaminations in *S. junceum* nodal explants. Means followed by the same letters are not significantly different from each other (*p* < 0.01) as determined by Duncan’s Multiple Range Test (DMRT).

| Ethanol 70% (s) | Home Bleach 20% (min) | Fungi infection | Bacterial infection |
|----------------|-----------------------|----------------|-------------------|
| 0              | 0                     | 6⁴              | 6⁴                |
| 0              | 10                    | 3.33⁴           | 1.66⁶ab           |
| 0              | 15                    | 3bc             | 2.33⁵ab           |
| 30             | 0                     | 2⁴ab            | 3⁶d               |
| 30             | 10                    | 1⁴              | 2⁴bc              |
| 30             | 15                    | 1⁴              | 1⁴                |
| 60             | 0                     | 1.66⁴           | 3.66⁴             |
| 60             | 10                    | 1⁴              | 1.66⁶ab           |
| 60             | 15                    | 1⁴              | 1.66⁶ab           |
The antioxidant treatments in initiation culture medium

Since browning phenomenon was occurred in explants after first experiment, this experiment was implemented. Here, the same semi hard woody shoots and nodal segments of *S. junceum* L. in 0.5 cm long were used as explants. At the beginning, explants were sterilized using alcohol 70% for 30 s and commercial sodium hypochlorite 20% concentration (home bleaching solution, containing 5% activated agent) for 15 min. In all treatments, nodal segments were cultured on medium MS containing 0.5 mg L\(^{-1}\) 2,4-D, 30 g L\(^{-1}\) sucrose and 7 g L\(^{-1}\) agar. Antioxidant treatments for control browning of nodal explant in this experiment included submerging explants in solution PVP 0.5% for 0, 20, 30, 60, 120, 240, and 480 min, rinsing explant in tap water for 30 and 60 min, applying PVP 0.5% in media, curcumin (turmeric) 0.1% in media and activated charcoal 0.1% in culture medium. This experiment was implemented in a completely randomized design with three replications. The factors studied here were the type and the method of using antioxidants. Browning and fungal or bacterial infections and the amount of callogenesis were graded and recorded at the end of 1\(^{th}\) week and at the end of month 1, respectively (Table 1, Figure 1 and 2).

Figure 1. Browning ranking of nodal explants in *S. junceum*: (a) degree 0: no brown signs, (b) degree 1: partial browning at the cut end, (c) level 2: browning at the cut end and bottom surface of the explant, (d) level 5: Browning the whole surface of the explant

Figure 2. Callogenesis ranking of nodal explants in *S. junceum*: (a) No callogenesis, (b) Low, (c) Low to medium, (d) High

The effects of PGR treatments and light condition

In this experiment, the effect of light (16 hours’ daylight and 8 hours’ dark cycle) and continues darkness, and different concentrations of BA in combination with different concentrations of 2,4-D on callogenesis of nodal explants of *S. junceum* were studied. To this experiment, three concentrations of BA (0, 0.1 and 0.2 mg L\(^{-1}\)) in combination with four concentrations of 2,4-D (0, 0.5, 1.0 and 2.0 mg L\(^{-1}\)) were applied. The 300 mg L\(^{-1}\) PVP was added to medium to prevent explants from browning. Disinfected nodal explants (by ethanol 70% for 30 s and sodium hypochlorite 20% for 15 min) were cut into 0.5 cm segments and cultured on callus induction medium. The cultures were kept at temperature 25 ± 1 °C at light/dark and darkness conditions.

This experiment was carried out in factorial plan in completely randomized design with four replications. The results were evaluated every week and the time of callus initiation was recorded. At the end of 4\(^{th}\) week, callus induction percentage, the amount of callogenesis, texture
and color of callus (callus morphology) and browning of explants were recorded. Browning degree and the amount of callogenesis in explants were evaluated according to grading represented in Figure 1 and 2.

**Results**

The effect exposure times of Ethanol and home bleach treatments

This experiment was scheduled to determine the most effective procedure for surface disinfection of explants by the way of identifying the best disinfectant and the best duration for explants to be exposed to them to control infections and browning disorder in *S. junceum* L.

The results from comparison of the mean interaction effects of ethanol duration and home bleach duration on the evaluated qualities in nodal explants showed that home bleach alone reduced the fungal infection to half in comparison with control explant, whereas ethanol duration alone (30 and 60 s) showed better results in controlling fungal infection in nodal explant in comparison with control explants. The least fungal infection corresponded to simultaneous ethanol and home bleach treatment. However, there wasn’t significant difference between each of treatments alone. Also, the results from comparison of means showed that the most bacterial infection was in control explants and then the ethanol treatment alone came after that. Once home bleach was applied alone, bacterial infection level reduced significantly compared with control treatment and didn’t differ from combined simultaneous ethanol and home bleach treatment significantly, whereas it showed significant effect in comparison with ethanol alone. The least bacterial infection corresponded to applying combination of ethanol for 30s and home bleach for 15 min (Table 1).

Studying the simple effects of different durations of alcohol treatment on browning degree of nodal explants showed that the highest degree of browning corresponded to applying ethanol for 60 s which didn’t differ from control significantly, while the lowest degree was observed in ethanol treatment for 30 s (Figure 3a). The results showed that applying home bleach for 10 min and 15 min increased browning degree in nodal explants. The browning increased significantly up to two times in 15 min treatment compared with control (Figure 3b).

![Figure 3. Influence of various exposure times sterilizing agents on *S. junceum* explant browning: (a): Home bleach 20%, (b): Ethanol 70%. Means followed by the same letters are not significantly different from each other (p<0.01) as determined by Duncan’s Multiple Range Test (DMRT).](image)

**Antioxidant treatments in initiation culture medium**

In all treatments, calluses initiated after 14 days of culture, in this study. The results showed that in pre-treating explants with PVP 0.5% and most of treatments no bacterial infections were found, compared with control. Bacterial contamination just increased in some treatments such as pre-treating with PVP (240 and 480 min). Furthermore, the effect of culture medium supplemented with PVP on explant browning was so significant that any tissue browning wasn’t occurred in explants.

The highest browning sign (browning of the whole explant tissues) was occurred in control medium and in explants pre-treated with PVP (60, 240 and 480 min) and in medium supplemented with curcumin. Also, to apply tap water (30 and 60 min) increased the explant browning and there wasn’t significant difference among these tap water treatments and curcumin treatment and PVP-soaked explant treatment (60, 120, 240 and 480 min). Adding activated charcoal to culture medium and soaking explants in PVP (20 and 30 min) decreased browning in comparison with control which didn’t differ significantly from application of PVP to medium, but they had significant difference with other treatments (Table 2). At the end of week 4, at first, the treatments injured due to infections were discarded then the extent of callus induction of remainder was evaluated.
The results from mean comparisons demonstrated that the greatest extent of callogenesis associated with application PVP to culture medium and soaking explants in PVP (20 and 60 min), and it wasn’t found significant difference between these three treatments, but application of PVP to medium differed significantly from other treatments. The lowest amount of callus was found in medium containing curcumin 1%, compared with other treatments; however, there wasn’t significant difference between curcumin treatment, tap water treatments and activated charcoal 1% treatment (Table 2).

The effects of PGR treatments and light conditions
In this experiment, all formed calluses had compact and hard textures, and were in light green (in presence of light) and in milky white color (under darkness). It wasn’t found any callus formation in absence of plant growth regulators (control) or in culture medium supplemented with BA to alone either in presence or in absence of light. Callus initiation was occurred in all treatments containing 2,4-D. In culture medium supplemented with 0.5 mg L\(^{-1}\) 2,4-D in absence of BA, callus formation was initiated in 3 weeks in presence of light, and in 2 weeks under darkness conditions.

Calluses started to initiate after two weeks in culture medium supplemented with 0.5 mg L\(^{-1}\) 2,4-D and 0.1 and 0.2 mg L\(^{-1}\) BA, under light and darkness conditions, while callus initiation time declined as 2,4-D concentration increased. So that in all treatments supplemented with 1 and 2 mg L\(^{-1}\) 2,4-D alone or in combination with different concentration of BA, callus formation was observed just in one week of culture. It showed that there is a contrary link between concentration of 2,4-D and callus initiation time. It can be concluded based on comparison of means that 0.5 mg L\(^{-1}\) 2,4-D in combination with BA (0, 0.1 and 0.2 mg L\(^{-1}\)) under darkness is the best PGR composition to get the maximum mass of callus.

Considering the results, the lowest level of 2,4-D (0.5 mg L\(^{-1}\)) and BA (0 mg L\(^{-1}\)) and darkness condition had appropriated the high callus mass, and used more concentrations of 2,4-D has reduced the mass of callus. The highest browning was obtained in medium supplemented with 2 mg L\(^{-1}\) 2,4-D and 0.2 mg L\(^{-1}\) BA under darkness, while the least browning degree was obtained by used 0.5 mg L\(^{-1}\) 2,4-D in medium and under darkness, so in most cases, low or zero concentration of 2,4-D had brought about least degree of browning in explants. It demonstrates that applying 2,4-D in low concentration alone or in combination with BA under both light and darkness conditions didn’t have any influence on browning of explants, whereas browning degree increased either in presence or in absence of BA under both light and darkness conditions, as concentration of 2,4-D increased (Table 3).

### Table 2. Effect of antioxidant treatments on contamination, browning and callogenesis of survival of *S. junceum* nodal explants. Means followed by the same letters are not significantly different from each other (*p* < 0.01) as determined by Duncan’s Multiple Range Test (DMRT).

| Antioxidant treatments                      | Bacterial infection (ranking) | Explant browning (ranking) | Callogenesis (ranking) |
|--------------------------------------------|-------------------------------|-----------------------------|------------------------|
| Control                                    | 1.00\(^a\)                   | 5\(^f\)                     | Not survival           |
| Medium supplemented with 0.5% PVP          | 1.00\(^a\)                   | 0\(^a\)                     | 5\(^a\)                |
| Explant immersed in 0.5% PVP (20min)       | 1.00\(^a\)                   | 2.26\(^b\)                  | 4.26\(^abc\)           |
| Explant immersed in 0.5% PVP (30min)       | 1.00\(^a\)                   | 2.13\(^b\)                  | 3.37\(^bc\)           |
| Explant immersed in 0.5% PVP (60min)       | 1.00\(^a\)                   | 4.13\(^def\)               | 4.76\(^ab\)           |
| Explant immersed in 0.5% PVP (120min)      | 1.00\(^a\)                   | 3.90\(^abc\)               | 3.50\(^c\)           |
| Explant immersed in 0.5% PVP (240min)      | 2.66\(^b\)                   | 4.33\(^def\)               | Not survival           |
| Explant immersed in 0.5% PVP (480min)      | 3.00\(^b\)                   | 4.73\(^ef\)                | Not survival           |
| Medium supplemented with turmeric 0.1%     | 1.00\(^a\)                   | 4.20\(^abc\)               | 2.20\(^d\)           |
| Medium supplemented with 0.1% AC           | 1.00\(^a\)                   | 2.46\(^bc\)                | 3.33\(^de\)           |
| Running water (30 min)                     | 1.00\(^a\)                   | 3.40\(^d\)                 | 3.20\(^e\)           |
| Running water (60 min)                     | 1.00\(^a\)                   | 3.60\(^a\)                 | 3.26\(^de\)           |
Table 3. Effect of PGR and treatments light conditions on Callogenesis and browning of nodal explants in *S. junceum*. Means followed by the same letters are not significantly different from each other (*p* < 0.01) as determined by Duncan’s Multiple Range Test (DMRT).

| 2,4-D (mg L⁻¹) | BA (mg L⁻¹) | Treatment of light regime | Callus induction time (week) | Callogenesis (ranking) | Explant browning (ranking) |
|----------------|-------------|--------------------------|-----------------------------|------------------------|---------------------------|
| 0              | 0           | Light                    | 0                            | 0                      | 2.12*                     |
| 0              | 0           | Dark                     | 0                            | 0                      | 0.99*                     |
| 0              | 0.1         | Light                    | 0                            | 0                      | 0.45*                     |
| 0              | 0.2         | Dark                     | 0                            | 0                      | 0.29*                     |
| 0.5            | 0           | Light                    | 3                            | 4.24 abc               | 0.17 ab                  |
| 0.5            | 0           | Dark                     | 2                            | 4.57 a                 | 0.17 ab                  |
| 0.5            | 0.1         | Light                    | 2                            | 3.79 cd                | 0.31 ab                  |
| 0.5            | 0.2         | Dark                     | 2                            | 4.33 ab                | 0.37 ab                  |
| 0.5            | 0.2         | Light                    | 2                            | 2.74 f                 | 0.45 ac                  |
| 1              | 0           | Light                    | 1                            | 3.16 ef                | 2.34 gh                  |
| 1              | 0           | Dark                     | 1                            | 3.99 g                 | 0.99 fc                 |
| 1              | 0.1         | Light                    | 1                            | 2.99 i                 | 1.66 gb                  |
| 1              | 0.1         | Dark                     | 1                            | 3.45 ik                | 0.66 bc                  |
| 1              | 0.2         | Light                    | 1                            | 2.78 f                 | 1.52 d                  |
| 1              | 0.2         | Dark                     | 1                            | 3.78 cd                | 0.66 bc                  |
| 2              | 0           | Light                    | 1                            | 1.62 ef                | 1.91 gi                  |
| 2              | 0           | Dark                     | 1                            | 1.83 e                 | 1.77 gh                  |
| 2              | 0.1         | Light                    | 1                            | 1.70 gb                | 2.49 gh                  |
| 2              | 0.1         | Dark                     | 1                            | 1.53 gh                | 2.16 ey                  |
| 2              | 0.2         | Light                    | 1                            | 1.28 h                 | 2.74 f                  |
| 2              | 0.2         | Dark                     | 1                            | 1.24 i                 | 3.48 l                  |

Discussion

One of the main restricting factors in micropropagation process, and especially in woody plants is a fungal and bacterial infection (Nazary Moghaddam Aghayeh et al., 2019). Death explants associated with infections was one of the limitations in initiation phase on nodal segment culture of *S. junceum* L. in present study, as in other woody plants. At disinfection phase, various chemical compounds are applied to control *in vitro* infections. However, some of them are less effective and some are too toxic for plants to apply in disinfection. So, it is difficult to adopt a standard disinfection method for all plants (Jan et al., 2013; Taghizadeh et al., 2016). The results from this experiment demonstrated that ethanol in combination with sodium hypochlorite had the best effect in control fungal infections for nodal explants of *S. junceum* L.

It has been proven that two-phase disinfection method is beneficial for certain plant species. Ethanol usually exerts the influence in combination of sodium hypochlorite (Khanam and Chandra, 2017). In this study, it was found that sodium hypochlorite alone brought about reduction of bacterial infection, compared with ethanol alone. To apply ethanol for 30s simultaneously with sodium hypochlorite for 10 min was the best treatment for control bacterial infections in nodal explants. Ethanol may increase the efficiency of disinfection with sodium hypochlorite and other disinfectant by eliminating air bubble from surface of explants. Ethanol also improves penetration of disinfectants through the plant tissues by removing vaxes surface plant organs. During disinfection procedure, live material shouldn’t lose their biological activity but just microorganisms should be eliminated. So, to sterilize explants, just disinfectant solutions in proper concentrations and with proper durations should be used (Putri et al., 2019).

Sodium hypochlorite as a bleaching substance is usually the main choice for surface disinfection. Because of toxic effects on explants, balance between concentrations and exposure durations should be determined experimentally for any explants (Putri et al., 2019). According to results of this study, ethanol and sodium hypochlorite were both effective for disinfecting nodal explants of *S. junceum* L. In *Fragaria × ananassa* hort., among treatments ethanol 70% for 30s in combination with sodium hypochlorite for (0, 15, 20 min), the highest percentage of sterile cultures was achieved when nodal segments were treated with ethanol 70% for 30s and sodium hypochlorite 1.5% for 20 min, but viable explants percentage was less because of browning and injured status of tissues (Jan et al., 2013).
Increase in the concentration of NaOCl and duration of immersion decreased the percentage of survival of explants taken from *Hevea brasilensis* (Moradpour et al., 2016). In this experiment, the highest nodal explant browning was achieved when sodium hypochlorite was applied alone for 15 min and ethanol was applied for 60s. This fact demonstrated that increased submerging duration in disinfectant solution gave rise to increase stress in tissues, and subsequently phenolic compounds would be produced. In Sour cherry (*Prunus cerasus* L.), by applying sodium hypochlorite 3% for 10 min and 1% for 20 min, high viable explants were achieved (Mihaljevic et al., 2013).

In plant tissue culture, browning phenomenon and subsequently explant death is one of the main problems for initiation in woody plant culture, which is generally attributed to phenolic compounds (Hesami et al., 2020). Phenolic compounds brought about browning of nodal segments of *S. junceum* L. as they cultured in medium MS, which prevented *S. junceum* L. from callus initiation. Phenolic compounds could activate polyphenol oxidase enzyme and could alter cellular metabolism and then make brown precursor Quinine through oxidation. Quinine gradually penetrated explant tissues in culture medium, and therefore suppressed the activity of other enzymes which eventually caused compounds in medium to be toxic.

Antioxidants are able to suppress oxidation of phenolic compounds by changing oxidation potential of phenolic compounds explants, so they can decrease browning symptoms (Raj et al., 2020). Antioxidants are able to prevent phenolic compounds from oxidation, while the absorbents are able to absorb quinine (Nishchal et al., 2018). Many authors have reported some techniques to overcome phenolic compounds, including pre-soaking explants in tap water in order for phenolic compounds to get eliminated, adding antioxidants to culture medium, such as polivynil pyrrolidone (PVP) or activated charcoal which will cause elimination of phenols (Meziani et al., 2016). In present study, we succeeded to get free-infection cultures in all treatments, and bacterial infection was occurred just when the duration of pre-treatment with PVP was increased (240 and 480 min).

Bacterial infection associated with increased duration of pre-treatment with PVP solution (240 and 480min) might have caused by increased moisture to the greatest extent, which provided suitable conditions for growth of pathogens. No bacterial infection in other PVP-pretreated explants was observed because PVP was capable of reducing phenol leakage, and hence it supported preservation of high levels of phenolic compounds in explants to keep pathogens back from growing. In mango, significant reduction in infection extent was observed on explants treated with PVP (Krishna et al., 2008). In present study, it was found that applying PVP 0.5% to callogenesis culture medium of *S. junceum* L. prevented explants from browning. Similarly, in *Curculigo latifolia*, a medicinal plant, applying PVP 0.5% to culture medium reduced browning and infection in shoot-tip explants (Babaei et al., 2012).

Browning intensity increased as duration of soaking explants in PVP solution was increased. It might have been occurred as a result of undesirable effect of reduction of solution pH by antioxidants (Karantaca, 2008). In fact, limited leakage of phenolic compounds from tissue treated implies that viable explants rate is increased in culture duration. In *Strelitzia reginae*, soaking explants in citric acid (200 ppm) for 24 hours preceding culture in medium MS containing activated charcoal (1%) decreased tissue browning effectively (Karantaca, 2008). In *Cinnamomum tamala*, phenolic compounds and other secondary compounds entered into the antioxidant solution when nodal explants and immature seeds were soaked in various antioxidant solutions immediately after harvest, but the explants got brown and damaged when they cultured without soaking in antioxidant solutions (Chitta et al., 2013).

The least browning was found in PVP treatments containing activated charcoal for 30 and 60 min in explant *S. junceum* L. It can be assumed that these treatments are capable of leakage phenols away from explant tissues effectively. The culture medium to supplemented with PVP to the better controlled browning disorder than to soak explants in PVP for 20 and 30 min, and PVP in culture medium could make tissue browning disappeared, completely.

Activated charcoal absorbs both toxic phenolic compounds and growth regulators, so the amounts of activated charcoal should be reduced critically (Nayanakantha et al., 2010). In *Saccharum officinarum* L., medium MS supplemented with 0.2% and 0.3% PVP, 100% and 80% of explants had viable tissues respectively, while in medium containing 0.3% and 0.4% activated charcoal 46% and 40% of explants survived respectively (Shimelis et al., 2015). Activated charcoal made culture medium dark, and hence it didn’t allow free radicals to form through oxidation of phenolic compounds (Nayanakantha et al., 2010). In this study, the most callogenesis was observed in medium containing 0.5% PVP, while the least callogenesis belonged to use of curcumin to the medium.

Antioxidant compounds can affect explants growing by the way of stimulating leaf expansion process; stimulating growing of callus, and promoting root development in cuttings (Huh et al., 2017). Therefore, the positive effect of PVP on callogenesis was related to absorb phenolic compounds out of explants. So, applying PVP to culture medium in concentration 0.5% can be reported as the treatment with lowest level of infections and with enough callus induction and as the best treatment for control browning in callogenesis medium of *S. junceum* L.

According to these results, it wasn’t found any callus in free PGR medium and in medium supplemented with BA as well. This non-induction of callus might have caused by insufficient phyto-hormone in nodal explants of *S. junceum* L. needed for inducing. So, it is necessary to apply synthetic auxins, such as 2,4-D for stimulating callus induction (Budisantoso et al., 2017). The results from respective study showed that the existence of 2,4-D in low concentration was necessary for inducing callus formation in this shrub, even if there isn’t any cytokinin in medium. The effective contribution of 2,4-D to inducing callus
formation was attributed to stimulates cell division in plant tissues and prevents tissues from organogenesis extremely (Shinta and Minarno, 2020). Maximum percentage of callus induction and the highest mass of callus was occurred in medium supplemented with 0.5 mg L\(^{-1}\) 2,4-D, with induction percentage and mass of callus were decreased as concentration of 2,4-D was increased. It showed that the concentration of auxin affected callogenesis in nodal explants of \textit{S. junceum} L. differently. 2,4-D is widely applied either alone or in combination with cytokinin, specially BA, in order to stimulate in vitro callus induction. However, the combination and concentration of these plant growth regulators should be defined for species (Dawa et al., 2017) and efficiency of these reactions depends on species and tissue of the plant (Castro et al., 2016).

In case of \textit{Santalum album} L. similar results were obtained, the most induction and growth of callus was occurred in medium containing low concentration of 2,4-D (1.5 and 2.0 mg L\(^{-1}\)) (Singh et al., 2016). The external application of BA either alone or in combination with 2,4-D didn’t have any influence on induction and growth of callus. It implied that cytokinin didn’t have any positive or negative effect on growth of callus. Abdelmageed and collegues (2012) studied \textit{Michelia champaca} L., and achieved similar results. It was found that darkness had positive effect on percentage of induction and growth of callus. Formation and growth of callus depended on culture conditions, such as light regimen, temperature, and composition of medium in terms of kind and concentration of PGR (Khan et al., 2018).

Abdi and Hedayati (2011) noted that in \textit{Delonix regia} the most percentage of embryogenic callus formed from seeds and immature embryos was achieved by composition 2 mg L\(^{-1}\) 2,4-D and 0.25 mg L\(^{-1}\) BA under darkness conditions. Ye et al. (2012) found that callus induction in leaf explants of \textit{Ziziphus jujube} L. reached to 100% under darkness conditions, whereas under light conditions just 10% of leaves produced low-quality callus. This observation confirmed that callus induction is stimulated by darkness treatment. Similar results were obtained for \textit{Byrsonima verbascifolia} L. The obtained results implied that decreased growth in presence of light might have caused by photodegradation of compounds in culture medium, including auxins (Erst et al., 2018).

Applying high concentration of 2,4-D led to increased browning of explant tissues, whereas nodal explants of \textit{S. junceum} L. could produce callus without any browning in low 2,4-D concentration. 2,4-D is a strong auxin among others which is commonly used (Budisantoso et al., 2017). So, the negative effect of 2,4-D in high concentration on browning of nodal explants of \textit{S. junceum} L. could be attributed to its high stability in media. Moreover, 2,4-D have a relatively high mobility compared with other auxins, so it quickly accumulates in tissues and causes oxidation and degradation of them (Naz et al., 2018). These observations are consistent with those Kayani et al. (2008) had reported that callus got brown and damaged under high concentrations of auxine. According to these results, culture medium supplemented with 0.5 mg L\(^{-1}\) 2,4-D under darkness conditions can be proposed as the best callogenesis treatment in \textit{S. junceum} L.

**Conclusions**

In general, the results from this study showed that to apply ethanol 70% for 30s and sodium hypochlorite 20% for 10 min is the best treatment for disinfection of nodal explants, because the least fungal and bacterial infections, the least explant browning degree. In order to lessen browning in nodal explants due to phenol discharge in conditions of in vitro culture, it is recommended that PVP 0.5% is applied internally to the culture medium because the least browning and the most callus mass were found in this treatment, compared with applying curcumin and activated charcoal to the medium and pre-treating by soaking explants in PVP and in running water. The best hormonal composition to induce callus from explants and get the highest amount of callus was 0.5 mg L\(^{-1}\) 2,4-D under darkness conditions. Although, induction of callus was occurred by applying 1.0 and 2.0 mg L\(^{-1}\) 2,4-D, calluses weren’t able to proceed further because of getting brown. The existence of BA in applied concentration in this experiment didn’t have any significantly effect on callogenesis. Therefore, the best concentration of 2,4-D (0.5 mg L\(^{-1}\)) under darkness conditions is recommended to get callus from nodal explants of \textit{Spartium junceum} L.

**Author Contribution**

MT: participated in all of experiments, coordinated the data-analysis and contributed to the writing of the manuscript. MGD: Coordinated the laboratory work.

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