Effectiveness of Colchicine and Oryzalin at Inducing Polyploidy in Watsonia lepida N.E. Brown

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Abstract. Genetic modification and manipulation offers the possibility of introducing novel traits into existing plants, thereby increasing marketability. Polyploid induction has in the past produced plants that are more compact and have larger flowers, leaves, and fruit, making them more desirable to consumers. The effect of pulse treatments (0, 24, 48, and 72 h) of colchicine (25, 50, 125, and 250 μM) or oryzalin (30, 60, 90, and 120 μM) on in vitro-grown Watsonia lepida N.E. Brown shoots was investigated. Explant survival was higher and more consistent with oryzalin treatment compared with treatment with colchicine. More mixoploids than tetraploids were produced with both compounds. The optimum treatment for producing tetraploids was 120 μM oryzalin for 24 h. Of the 30% explants that survived this treatment, 33% were found to be stable tetraploids.

Sexual reproduction results in the fusion of two gametophytes (n) to form a new sporophyte (2n) zygote containing genetic material from both parents. It is possible that over extended periods of time, mutations, discrepancies in reduction division during gamete formation, spontaneous somatic polyploidization, and wide hybridization cause plants to accumulate additional sets of chromosomes from closely or distantly related species. Polyploidy has been an important factor in plant evolution and is often associated with speciation and development of novel adaptations (Levin, 2002). Approximately 70% of all angiosperms are considered polyploids (Goldblatt, 1980; Masterson, 1994).

It is possible to artificially create polyploids by interfering with cell division. A number of natural and synthetic compounds can be used and are either applied to plants ex vitro or in vitro. Colchicine is naturally occurring, and it was initially thought that it promoted polyploidy induction by disrupting spindles formation and preventing nuclear and cell division (Ganga and Chezhiyan, 2002, and references therein).

Recently, however, Caperta et al. (2006) showed that spindle disruption alone is insufficient for the production of polyploid cells. Low concentrations (0.5 mM) of colchicine inhibited microtubule formation in all phases of the cell cycle but resulted in abnormalities, including reduced viability from irregular-shaped nuclei and micronuclei. In contrast, treatment with high concentrations (5 mM) of colchicine induced microtubule polymerization to form new structures in c-metaphase cells. These new structures are thought to aid reconstitution of polyploid nuclei and subsequent re-entry into the cell cycle (Caperta et al., 2006). Oryzalin and trifluralin are synthetic herbicides used for weed control and disrupt microtubule assembly during cell division. They are often more effective than colchicine because they have a higher affinity for plant tubulins (Dolezel et al., 1994).

Effectiveness of these compounds in vitro depends highly on the concentration applied, duration of treatment, type of explant, and the penetration of the compound (Allum et al., 2007). Colchicine has been effectively used in the concentration range 0.25 μM (Chen et al., 2006) to 38,000 μM (Stanys et al., 2006); oryzalin from 2.5 μM (Allum et al., 2007) to 150 μM (Contreras et al., 2007); and trifluralin from 250 μM to 2500 μM (field application ex vitro; Zlesak et al., 2005). In many instances, oryzalin and trifluralin are more effective at stable ploidy induction, have an increased survival of explants, and are used at lower concentrations than colchicine (Ganga and Chezhiyan, 2002; Zlesak et al., 2005).

Ploidy manipulation offers some benefits for horticultural, pharmaceutical, and agricultural improvement of plants. Rhododendron tetraploids induced by oryzalin had larger leaves, flowers, and pollen compared with diploids (Conterras et al., 2007). Similarly, tetraploid watermelons had greater leaf area, larger flowers and ovaries, larger seeds, and a thicker rind (Jaskani et al., 2005). From a horticultural perspective, polyploid plants often have a shorter, more squat stature, and the larger, thicker leaves and bigger flowers are more attractive to consumers (Chen et al., 2006). Production of secondary metabolites from polyploid plants was increased in Panax ginseng (Kim et al., 2004), Artemisia annua (De Jesus-Gonzalez and Weathers, 2003), Datura stramonium (Berkov and Philipov, 2002), and Scutellaria biacalensis (Gao et al., 2002).

A number of explants can be used as starting material to induce polyploidy. Young, actively growing explants containing a meristem usually give best results. For example, success has been found using germinating seeds (Pringle and Murray, 1992), ex vitro shoots (Contreras et al., 2007), roots (Kim et al., 2004), embryogenic callus (Gmitter et al., 1991), nonembryogenic callus (Gao et al., 2002), nodal segments (Chen et al., 2006), cotyledons (Stanys et al., 2006), and hypocotyls (De Carvalho et al., 2005). During micropropagation of Watsonia species, we observed that only the hypocotyl region of seedlings (immediately above the root) was capable of regenerating adventitious shoots, probably because they contain the shoot apical meristem and thus would make suitable explants for ploidy induction (Ascough et al., 2007).

Stomata of induced polyploids are usually larger (Griesbach and Bhat, 1990; Kadota and Niimi, 2002), a simple diagnostic feature that is often used to identify potential polyploids. This technique is rapid, inexpensive, nondestructive, does not require sophisticated equipment, and has a fairly high accuracy rate (up to 90% in some cases; Cohen and Yao, 1996); however, it is an indirect method for ploidy assessment. If mixoploid plants are produced, then stomatal size can be an unreliable method and should be combined with another technique (Chen et al., 2006). Chromosome counts are perhaps the most accurate, and aneuploidy may be detected, although this method can be time-consuming if many plants are being tested. Analysis of DNA content by flow cytometry is often preferred. Because many thousands of cells can be analyzed rapidly, mixoploids can be readily detected. In some cases, for example, in genome doubling from wide hybridization, genomic rearrangement can lead to altered genome size and complex interpretation of ploidy based on DNA content (Levin, 2002). In addition, flow cytometry requires the purchase of expensive equipment, and chromosome behavior cannot be observed with this method.

The genus Watsonia Miller has several members that show potential for horticultural
use. For instance, many of the species grow taller than 1 m, and although this may be good for use in cut flowers, it is too tall for use as a container plant. In *Watsonia aletroides* (Burman fil.) Ker and *W. meriana* (Linnaeus) Miller, populations of triploid (2n = 3x = 27) plants are known, but polyploidy has not otherwise been documented in the genus (Goldblatt, 1989). These triploids do not produce seeds, but produce axillary cormlets at the aerial nodes of the spike after flowering (Goldblatt, 1989). Although chromosome counts have not been performed on *W. lepida*, plants are assumed to be diploid (2n = 2x = 18) because they produce seeds. The 1C value of *W. laccata* (Jacquin) Ker was determined as 0.7 pg (Goldblatt et al., 1984).

Four species (*W. gladioloides* Schlechter, *W. laccata*, *W. lepida*, and *W. vandersypuei* L. Bolus) have been propagated in tissue culture (Ascough et al., 2007, 2008) and chemical dwarfing techniques have been successful on *W. tubularis* Mathews & L. Bolus and *Watsonia* ‘Shrimp pink’ (Thompson et al., 2005). However, to maintain a dwarf stature, corms need to be retreated each year. Thus, the effectiveness of colchicine and oryzalin at inducing polyploidy in *W. lepida* was investigated to establish a working protocol for polyploidization in this genus. The long-term goal is to use polyploidy as a permanent means of dwarfing. Additional morphological change like altered flower phenotypes may also improve marketability of plants to consumers.

**Materials and Methods**

**Plant material.** Seeds from a single seed lot of *W. lepida* were obtained from Silverhill Nurseries, Kenilworth, South Africa, and the experiment initiated within 1 month of receiving the seeds.

**Culture conditions.** Except for seed germination in which one-tenth strength Murashige and Skoog (MS) basal salts (Murashige and Skoog, 1962) were used without sucrose or hormones, all media contained full-strength MS components and 3% sucrose (w/v). Medium pH was adjusted to 5.8 using NaOH before addition of 0.9% agar (w/v) (Agar-Agar no. 1; Marine Chemicals, Kerala, India) and sterilized by autoclaving at 121 °C and 103.4 kPa for 20 min. All cultures were incubated at 25 ± 1 °C under Osram** (Johannesburg, South Africa) 75-W cool white fluorescent tubes providing a 16-h photoperiod and a light intensity of 12.6 μmol·m⁻²·s⁻¹ at culture level.

Seeds were decontaminated for 15 min in a 50% (by volume) commercial bleach [Jik** (Reckitt Colman, Elandsfontein, South Africa), 3.5% sodium hypochlorite] solution with three drops of Tween 20 as a surfactant. Seeds were rinsed three times with sterile distilled water, placed on a one-tenth-strength MS medium, and transferred to the culture room for germination.

After 3 weeks, uniform seedlings 4 cm long were selected for experimental studies. The root and upper portion of the leaf were excised using sterile technique in a laminar flow bench, and sections 1 cm long, containing the hypocotyl and lower leaf, were used as explants. These explants were implanted vertically in media with their basal end in 65 × 10-mm petri dishes. Ten explants were placed in each petri dish. Media consisted of MS basal salts and either colchicine at a concentration of 0, 25, 50, 125, or 250 μM or oryzalin [4-(dipropylamino)-3, 5-dinitrobenzenesulfonyamide] at a concentration of 0, 30, 60, 90, or 120 μM. Both colchicine and oryzalin were purchased from Sigma-Aldrich (St. Louis, MO). Explants were removed after 24, 48, and 72 h and placed singly in cylindrical 33-mL clear glass culture tubes containing 10 mL MS media with 2.2 μM BA to promote shoot production (Ascough et al., 2007). Subculturing was not necessary because plants did not "outgrow" their culture tubes. Ten explants were used for each treatment, and the experiment was repeated twice (for oryzalin treatments) and four times (for colchicine treatments) in time.

After 3 months, explants were examined to determine percentage survival. The few explants that had become infected with microbial contaminants (>5%) or were necrotic as a result of the treatment were discarded. Those that had survived and showed signs of regeneration (elongation, rooting, or multiplication) were analyzed for ploidy level by flow cytometric analysis (Ascough et al., 2006). Leaf tissue samples (0.3 g) from plantlets were cut into very small pieces with a razor blade in 1 mL extraction buffer containing 100 mM MgCl₂, 40 mM trisodium citrate, 22 mM MOPS, and 0.1% (v/v) Triton-X-100 with pH adjusted to 7.1. The suspension was filtered through a 50-μm mesh filter and stained with 500 μL propidium iodide for a final concentration of 0.5 mM. Fluorescence was measured using a Beckman Coulter (Fullerton, CA) Epics XL-MCL flow cytometer and total DNA content was compared with control data from untreated in vitro diploid control plants.

**Data analysis.** Data on survival rate, proportion diploid, mixoploid, and tetraploid plants were arcsine-transformed before statistical analysis by analysis of variance using Minitab® (State College, PA) release 14.

Means were separated using Fisher’s individual error rate at the 5% level of significance.

**Results and Discussion**

Lower concentrations and shorter durations of exposure to colchicine tended to have greater survival (Table 1). Best survival (84%) was observed when explants were treated for the shortest time (24 h) at the lowest concentration (25 μM). However, no polyploid plants were induced, and only diploids were observed, as indicated by the peak in the same position as the untreated control (Fig. 1A). A 48-h treatment at the same concentration resulted in moderate survival and no polyploids were produced. Only one explant treated for 72 h was mixoploid containing both diploid and tetraploid cells (Fig. 1B). A 24-h treatment at 50 μM produced more mixoploids with moderate survival rates, but when the concentration or duration was increased, explant survival declined sharply (Table 1). No tetraploids were observed in any of the surviving colchicine-treated plantlets despite high frequencies of mixoploid induction occurring at 250 μM. It is possible that even the shortest duration was too long, and a shorter duration of 6 or 12 h may be more effective.

Most of the explants produced healthy shoots in vitro, but there was very little adventitious shoot production and hence a low multiplication rate. Explants producing multiple shoots were all found to be diploid (data not shown). It is uncertain if this low multiplication was the result of the treatment regime or simply a suboptimal multiplication procedure. In this study, the concentrations of colchicine used were much lower than those used by Caperta et al. (2006). However, these concentrations are consistent with most other published reports in which tetraploids were successfully created, for example, in *Gladiolus* (Suzuki et al., 2005), *Scutellaria* (Gao et al., 2002), and in *Rhododendron* (Vainolä and Repo, 2001).

A number of recent reports indicate oryzalin as an effective alternative to colchicine. It can be used as a potential replacement because it is safer to work with and, at least in

| Treatment | Survival/treated (%) | Diploid no. (%) | Mixoploid no. (%) |
|-----------|----------------------|----------------|------------------|
| 0 μM      | 38/40 (95) a         | 38 (100) a     | 0 (0) a          |
| 25 μM 24 h| 26/38 (78) b         | 26 (100) a     | 0 (0) a          |
| 48 h      | 14/36 (39) cdef      | 14 (100) a     | 0 (0) a          |
| 72 h      | 20/38 (53) bcd       | 19 (95) ab     | 1 (5) a          |
| 50 μM 24 h| 22/36 (61) bcd       | 19 (96) abc    | 3 (14) a         |
| 48 h      | 10/36 (29) cef       | 9 (92) abcd    | 1 (6) a          |
| 72 h      | 12/36 (33) def       | 11 (92) abede  | 1 (8) a          |
| 125 μM 24 h| 10/36 (29) f         | 7 (70) bcde    | 3 (30) a         |
| 48 h      | 9/36 (25) f          | 7 (78) abcde   | 2 (22) a         |
| 72 h      | 6/36 (17) f          | 4 (67) bcde    | 2 (33) a         |
| 250 μM 24 h| 8/36 (22) f          | 5 (62) e       | 3 (38) a         |
| 48 h      | 5/36 (14) f          | 4 (50) cde     | 1 (20) a         |

*pValues containing the same letter within a column are not significantly different from each other at the 5% level of significance.*
some species, plantlet survival and tetraploid induction are higher at a much lower concentration than with colchicine (De Carvalho et al., 2005). Survival of *W. lepida* hypocotyl explants treated with oryzalin for 24 h at the two lowest concentrations (30 and 60 μM; Table 2) was lower than that of the two lowest concentrations (25 and 50 μM) of colchicine used (Table 1). At all other concentrations and treatment durations, survival was higher in oryzalin-treated explants than in those treated with colchicine.

More polyploidization events (mixoploids or tetraploids) were observed when explants were treated with oryzalin compared with the colchicine treatments, indicating possible enhanced action compared with colchicine. Supporting this supposition, a number of tetraploid plants were found in three of the oryzalin treatments (Table 2; Fig. 1C). The treatment producing the most tetraploids was a 24-h duration in 120 μM oryzalin with 33% of surviving explants being tetraploid. Despite explant survival being only 30% in this treatment, it is not so poor as to abandon it completely. Väinölä and Repo (2001) suggest that a concentration and duration of treatment that is highly lethal (low survival) may be an advantage because it reduces the number of diploid and mixoploid plants that need to be screened and tested when the target is tetraploid. However, care must be taken even with this approach because the tetraploids induced by high concentrations may have aberrations like a reduced growth rate. In addition to doubling meristematic cells, high concentrations could adversely affect surrounding non-dividing tissues, thereby reducing the overall vigor of the explant.

In some explants exposed to colchicine or oryzalin for 48 or 72 h, doubling of DNA content happened twice resulting in a plant containing diploid, tetraploid, and octoploid nuclei. In another individual, an intermediate peak between the diploid and tetraploid peak was observed (Fig. 1D). This peak, however, is unlikely to be triploid nuclei, because the peak is not equidistant from the diploid and tetraploid peak but appears to be of intermediate ploidy. Possibly, the oryzalin treatment caused a disruption in cell division in some of the cells such that DNA replication was not fully completed, leading to an incomplete doubling of the chromosomes and a plant that contains some sections/leaves of intermediate ploidy. Although uncommon, incomplete doubling and aneuploid and triploid production can occur when attempting to induce tetraploidy (Zlesak et al., 2005). In this study, more mixoploids were induced, and very few tetraploids were observed in most of the treatments. Similarly, in *Rhododendron*, both colchicine and oryzalin produced more plants of mixed ploidy than stable tetraploids (Väinölä and Repo, 2001). In closely related *Gladiolus*, a high percent (42.9) of plantlets treated with 125 μM colchicine were tetraploid with no mixploids induced (Suzuki et al., 2005). One explanation could be that multicellular tissue was used as the starting material in *Watsonia* compared with callus cultures (containing high numbers of individual cells undergoing rapid division) in *Gladiolus*. Thus, successful polyploid induction in *Watsonia* occurred in some cells and not in others. One possibility of improving this system is to use individual cells as starting material with the aim of producing a plant of only one ploidy level. Alternatively, additional concentrations and treatment times or sizes of explants could be evaluated because these can affect the percent of meristem cells successfully doubled and penetration of the chromosome-doubling agent, respectively (Allum et al., 2007). Most likely, penetration by the agent was through

![Image](image_url)
the cut end, and perhaps a method in which the entire explant is submerged (like in a liquid-shake culture) would improve penetration. It is possible to isolate complete polyploid sectors from mixoploid individuals eventually, but it may be simpler to discard these and produce stable tetraploids from scratch.

_Watsonia_ plants take a number of years to produce a corm large enough to flower; thus, these tetraploid plants will have to be monitored until they flower to determine if altered ploidy level has any significant effect on plant height and flower morphology. Once flowering, fertility will need to be assessed to determine if barriers preventing seed production are present. Tetraploid plants in essence represent a new population with possibilities for epistatic interactions, and time will be needed to ensure genome stability. Superior genotypes with improved characteristics can then be selected for breeding purposes or micropropagated for commercial production and distributed to local nurseries.

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