Studies on Seed Germination, Seedling Growth, and In Vitro Shoot Induction of *Aloe ferox* Mill., a Commercially Important Species

Michael W. Bairu, Manoj G. Kulkarni, Renée A. Street, Rofhiwa B. Mulaudzi, and Johannes Van Staden

Research Centre for Plant Growth and Development, School of Biological and Conservation Sciences, University of KwaZulu-Natal Pietermaritzburg, Private Bag X01, Scottsville 3209, South Africa

Abstract. A study was done to investigate the effects of some physical and chemical factors on growth and development of *Aloe ferox* ex vitro and in vitro. The effects of light, temperature, and smoke–water on seed germination, ex vitro seedling growth requirements, and effect of germination medium and cytokinins on shoot induction and multiplication in vitro were investigated. The highest germination percentage of *A. ferox* seeds was recorded between 15 and 30 °C. Seed germination was inhibited at 10 and 35 °C. There was a nonsignificant increase in germination percentage (78%) for seeds that were germinated under constant dark conditions. Smoke–water-treated seeds showed significant improvement in percentage germination (80%) over the control (66%) at 25 °C with a 16-h photoperiod. Seedlings of *A. ferox* subjected to alternating temperatures (30/15 °C) or being irrigated three times weekly at 25 °C showed better seedling growth. In an in vitro experiment, seedlings germinated in distilled water and one-tenth strength Murashige and Skoog medium had superior shoot induction competence compared with the other germination media. The cytokinins meta-Tropolin (mT) and meta-Tropolin riboside (mTR) at 5 μM gave significantly higher shoot multiplication rates compared with the control and benzyladenine (BA)-treated plants. A higher abnormality index was recorded for BA-treated plants. The findings of this study will be beneficial for commercial propagation of *Aloe ferox*.

*Aloes* grow in a wide range of habitats and are an important group of medicinal plants in Africa. Traditionally, the harvesting of plant parts was sustainable and limited to household use. Currently, many *Aloe* species are threatened in Africa as a result of commercial use (Maundu et al., 2004). Pfab and Scholes (2004) reported that harvesting only one adult plant annually from a population of 100 *Aloe peglerae* produces an extinction probability of 100%. This means harvesting of only 0.12% or less of the mature plants of *A. peglerae* per year can be considered as sustainable. *Aloe ferox* is another widely harvested South African species (Sachedina and Bodeker, 1999). The bitter yellow juice and gel obtained from the leaves of this species are used as a laxative and for healthcare products, respectively (Van Wyk et al., 1997). Newton and Vaughan (1996) estimated that in 1990, *A. ferox* leaves were officially harvested from ≈10 million plants for 400 t of bitters. They also reported that there was an unofficial trade of another 300 t of bitters, suggesting a need for the safeguarding of this species. The export details of *A. ferox* extract from South Africa between 1994 and 2003 is documented by Knapp (2006). Overall, the value of the *A. ferox* industry in South Africa is estimated to be R150 million (U.S. $15 million) per year. Government and private sectors are expanding this industry for the benefits of rural communities as a result of the increasing demand for *A. ferox* bitters and gel (Shackleton and Gambiza, 2007).

At present, *A. ferox* is not vulnerable in the wild. However, there is concern that overharvesting of leaves, although this does not harm the plant, may affect growth, flowering, and fruit production. The plant less resistant to drought (Donaldson, 1989; Newton and Vaughan, 1996), which can lead to local extinction (Van Wyk and Smith, 1996). Pfab and Scholes (2004) emphasized that sustainability can never be achieved without ex situ cultivation.

Tree and shrub species of *Aloes* are propagated through stem cuttings, which are dried and planted. A more or less similar method is adopted for suckering species in which small plantlets are detached from the mother plant that are dried and planted. However, seed propagation is more feasible and recommended for survival of rare species (Van Wyk and Smith, 1996). If this species has to be propagated on a large scale by means of seed or tissue culture methods, then currently there is no basic information available on these aspects. *Aloes* are succulent and warm-climate plants, where both temperature and water play an important role in establishing them. This study was therefore conducted to examine 1) the effects of different temperatures, growth-promoting substances, and watering frequencies on seed germination and seedling growth of *A. ferox*; and 2) to assess the applicability of an in vitro propagation protocol developed for other *Aloe* spp.

Materials and Methods

Seed collection. Dried seeds of *A. ferox* were collected between the middle to the end of August from the Botanical Garden, University of KwaZulu-Natal Pietermaritzburg, South Africa. Seeds of *A. ferox* are black in color, triangular in shape with a size ranging between 0.3 and 0.5 mm. The average weight of 100 seeds determined was 174 ± 4 mg with 8% moisture content at harvest. Collected seeds were used immediately.

Seed germination experiments. To determine the optimum requirements for seed germination of *A. ferox*, the seeds were subjected to different temperature regimes, light conditions, plant growth regulators (PGRs), potassium nitrate, and smoke solutions. Seeds were decontaminated with 0.1% mercuric chloride for 1 min and then rinsed with distilled water before germination tests. All treatments consisted of four replicates with 25 seeds in each. The seeds were placed on two layers of Whatman No. 1 filter paper in disposable plastic petri dishes (9 cm). The filter paper was wetted with 4 mL distilled water, PGRs, potassium nitrate, or smoke solutions and kept moist by adding aliquots of the respective solutions when required until the end of the experiment. Plant growth regulators used in this experiment were gibberellic acid (GA3) and kinetin (Sigma Chemical Co., St. Louis, MO) at concentrations of 10⁻³, 10⁻⁴, and 10⁻⁵ M (GA3 346.4, 34.64, and 3.464 mg L⁻¹, respectively; kinetin 215.2, 21.52, and 2.152 mg L⁻¹ respectively). Potassium nitrate (KNO₃) (Merck, Darmstadt, Germany) was tested at 10⁻¹, 10⁻², and 10⁻³ M (101.10, 10.10, and 1.010 mg L⁻¹, respectively). Seeds were incubated with smoke–water (smoke concentration and water dilution of 1.500 m/v) or an active compound butenolide [3-methyl-2H-furo[2,3-c]pyran-2-one] (10⁻⁶ M) isolated from smoke. The smoke–water was prepared by the methods outlined in Baxter et al. (1994), and the butenolide was isolated from plant-derived smoke–water according to the methods described by Van Staden et al. (2004). To determine the effects of different temperatures, the seeds were incubated at 10, 15, 20, 25, 30, 35, and 30/15 °C. The experiments were...
Conducted under a 16-h photoperiod with cool-white fluorescent lamps, which provided a photosynthetic photon flux density (PPFD) of 75.6 ± 3.8 μmol·m⁻²·s⁻¹. For continuous dark conditions, the petri dishes were placed in lightproof boxes at 25 ± 0.5 °C, and the seeds were inspected daily under green “safe light” conditions with a PPFD of 0.3 μmol·m⁻²·s⁻¹. In continuous light conditions, the PPFD was 76.4 ± 3.5 μmol·m⁻²·s⁻¹ at 25 °C.

Germination was recorded daily and was considered complete once the radicle protruded ≥2 mm in length. The experiments were continued for 14 d. Mean germination time (MGT) was calculated by the equation: MGT = Σ(n x d)/N, where n = number of seeds germinated on each day, d = number of days from the beginning of the test, and N = total number of seeds germinated at the termination of the experiment (Ellis and Roberts, 1981). The optimum temperature for germination was calculated on the basis of constant temperature as: To = Σtp/Σ p, where p is the percentage germination at temperature t (Ollf et al., 1994).

Seedling growth experiments. Two-week-old seedlings grown in petri dishes were transplanted into 20-cm pots filled with sterile quartz sand (sulfuric acid-washed, thoroughly rinsing with water cycles, and dried in an oven at 80 °C for 7 d) and moistened (1.3 kg) with half-strength Hoagland’s nutrient solution (HS) (75 mL) (Hoagland and Snyder, 1935). Each pot consisted of five seedlings with eight replications per treatment. The pots were arranged randomly in the plant growth chambers under a 16-h photoperiod with continuous cool fluorescent tubes (Osram L75 W/20X) light intensity of 45 μmol·m⁻²·s⁻¹ and a temperature of 24 ± 1 °C for 9 weeks. The data on shoot multiplication, shoot length, and rooting were collected and analyzed after 9 weeks of growth in culture.

Shoot multiplication. After bulking sufficient explants using the previously described procedure, experiments to investigate the effects of type and concentration of cytokinin on shoot multiplication and incidence of abnormality were designed. Benzyladenine and the meta-topolins, mT and mTR, were investigated at 2.5 and 5.0 μM concentrations. Five shoot-tip (2 cm each) explants (derived from seedlings germinated in one-tenth strength MS and agar) with two to three young leaves attached were cultured in screw-cap jars (300 mL) containing 50 mL of media. A total of 15 explants per treatment was used. Cultures were then incubated under the same growth conditions mentioned previously. Growth parameters such as multiplication rate (total number of shoots per jar divided by initial number of explants) and abnormality index (the ratio of abnormal to normal shoots) were calculated and analyzed after 9 weeks. Plantlets exhibiting undifferentiated growth, hyperhydricity, and deformity were considered abnormal.

Statistical analysis. Ex vitro results were analyzed using MINITAB release 14 statistical package (Minitab Inc., State College, PA). One-way analysis of variance was conducted to test Fisher’s significance level at 5%. Percentage germination data were arc-sine transformed before analysis. Data from the tissue culture experiments were analyzed using SPSS release 10 statistical package (SPSS Inc., Chicago, IL) at 5% probability levels. Mean separation for significance test was made using the Duncan’s multiple range test. Kruskal-Wallis test was conducted using the MINITAB package to analyze nonparametric data.

Results

Effect of temperatures, light conditions, plant growth regulators, KNO3, and smoke solutions on seed germination. Seeds of A. ferox incubated at 30 °C showed the highest percentage germination (76%) in comparison with the other temperatures (Fig. 1A). On the other hand, 35 °C had significantly lower germination (10%). Alternating temperatures (30/15 °C) showed an intermediate response to percentage germination (Fig. 1A). Different light conditions had no significant effect on the percentage germination. However, greater percentage germination (78%) was recorded under constant dark than with the seeds that were exposed to constant and alternating light conditions (Fig. 1B). Smoke-water (1:500 v/v) significantly increased percentage germination over the control seeds (Fig. 2). The different PGRs and KNO3 solutions tested in this study did not significantly affect germination.

Effect of temperatures and watering frequencies on seedling growth. The longest

![](image.png)

Fig. 1. (A) Effect of different temperature (16-h photoperiod) and (B) light conditions (AL = alternating light; CL = constant light; CD = constant dark) on percentage seed germination of Aloe ferox. All bars with different letter(s) are significantly different (P < 0.05) by Fisher’s test. Numbers above the letter(s) indicate mean germination time in days. To = calculated optimum temperature.
shoots were recorded at alternating temperatures (30/15 °C), which was significantly different from other temperatures with the exception of 30 °C (Table 1). The root length also significantly increased at 30/15 °C with the exception of 25 °C. Maximum number of leaves was recorded at 30/15 °C, whereas a greater number of roots were developed at 30 °C. The seedlings that were grown at 30 °C showed significantly greater fresh weight than at the other temperatures with the exception of 30/15 °C (Table 1). Watering frequencies of twice and three times weekly (W2 and W3) showed significantly longer root lengths (69 and 72 mm, respectively) than once weekly watering (W1) (51 mm) (Fig. 3). Although, in these watering frequencies, there was an increase in the length of shoots, these results were not significantly different from the seedlings that were watered once weekly. Watering of seedlings three times weekly had significantly more roots than watering once weekly. Seedling fresh weight was significantly greater for W2 and W3 treatments than W1 (Fig. 3).

In vitro seed germination and shoot induction. In vitro seed germination of *A. ferox* was very erratic. Seeds started germinating on Day 10 and continued to germinate for more than 1 month. Seeds germinated on all the different germination media tested. There was, however, a difference in the quality of seedlings produced and their subsequent response to in vitro culture treatments. Seeds germinated in distilled water and agar and one-tenth strength MS solid media showed better in vitro multiplication (Fig. 4). The different germination media also had a significant effect on shoot growth as indicated by shoot length (Fig. 5). These different germination media, however, did not have significant effects on rooting when transferred to shoot induction medium as revealed by the Kruskal-Wallis test on rooting scores (data not presented).

Seedlings germinated in one-tenth strength MS media were used for the multiplication experiment as a result of their superior response to in vitro treatment. Subculturing plants to fresh multiplication medium significantly improved shoot multiplication. This was evident when we compare the 12.0 ± 1.4 shoots per explant obtained from the 5 μM mT treatment (Table 2) with the 4.57 ± 0.9 (Fig. 4) on the same treatment during the shoot induction phase. The mTR at 5 μM gave the highest multiplication rate, which was significantly higher than benzyladenine at the same concentration but not significantly different when compared with mT. Benzyladenine also appears to be more detrimental than the toponis based on the abnormality index.

Plantlets failed to root in a multiplication medium, but rooting was achieved easily by transferring elongated shoots to cytokinin-free

### Table 1. Effect of different temperatures on seedling growth of *Aloe ferox* under a 16-h photoperiod.

| Temperature (°C) | Shoot length (mm) | Root length (mm) | Leaf (no.) | Root (no.) | Seedling fresh wt (g) |
|-----------------|------------------|-----------------|------------|------------|---------------------|
| 10              | 17 ± 0.46 e       | 7 ± 0.19 e      | 1.61 ± 0.04 d | 0 d        | 0.259 ± 0.020 cd    |
| 15              | 25 ± 0.59 d       | 14 ± 1.54 de    | 3.5 ± 0.33 c | 0.445 ± 0.021 d |
| 20              | 22 ± 0.64 d       | 25 ± 1.99 c     | 5.0 ± 0.41 ab | 0.663 ± 0.038 cd |
| 25              | 50 ± 3.17 c       | 61 ± 2.47 ab    | 5.8 ± 0.39 a | 1.550 ± 0.159 b   |
| 30              | 61 ± 3.41 ab      | 53 ± 3.30 b     | 6.4 ± 0.49 a | 2.341 ± 0.231 a   |
| 35              | 57 ± 4.13 bc      | 19 ± 2.47 cd    | 5.4 ± 0.29 ab | 0.858 ± 0.145 c   |
| 30/15           | 67 ± 2.02 a       | 64 ± 3.90 a     | 4.2 ± 0.22 bc | 1.973 ± 0.153 ab  |

*Mean value and SE (±) in the column with different letter(s) are significantly different (P<0.05) by Fisher’s test.*

---

**Fig. 2.** Effect of different concentrations of plant growth regulators [gibberellic acid (GA₃) and kinetin], potassium nitrate (KNO₃), and smoke solutions (SW = smoke–water; B = butenolide) on percentage seed germination of *Aloe ferox* at 25 °C with a 16-h photoperiod. Asterisk indicates significant difference from the control at 5% level (**P**<0.05) by Fisher’s test. Mean germination time was 8 d for all treatments.

**Fig. 3.** Effect of watering frequencies (W1, once; W2, twice; and W3, three times weekly) on seedling growth of *Aloe ferox* at 25 °C under a 16-h photoperiod. SE bars with different letter(s) are significantly different (**P**<0.05) by Fisher’s test.
medium as described previously. Furthermore, some browning was observed (Fig. 6A). This problem, however, had little effect on the formation of shoot clusters (Fig. 6B) when cultures were transferred to fresh medium 2 weeks after the initial culture. The severity of the problem was reduced with subsequent subcultures.

**Discussion**

**Ex vitro seed germination and seedling growth requirements.** Temperature and light are two important factors that influence seed germination of a number of species. In this study, the seeds of *A. ferox* showed better percentage germination (greater than 70%) between 15 and 30 °C with a lower MGT than at 10 and 35 °C. A broad range of temperature for germination can be attributed to the wide distribution of this species in South Africa (Shackleton and Gambiza, 2007; Van der Bank et al., 1995). Lower (10 °C) and higher (35 °C) temperatures were detrimental for seed germination of *A. ferox* (Fig. 1). Alternating temperatures (30/15 °C) were not effective in optimizing germination. Seeds of *A. ferox* incubated at all temperatures did not achieve optimum germination, although the seeds of *A. ferox* showed a viability of 95% or greater that was determined by the TTC method (International Seed Testing Association, 1999). This can be attributed to the need of an after-ripening period for *A. ferox* seeds to reach full maturity for maximum germination. However, this aspect needs further study. In regard to the effects of temperatures, the maximum percentage germination (76%) was obtained at 30 °C with a 16-h photoperiod. The best germination (78%) was achieved at 25 °C under constant dark conditions (Fig. 1). Smoke–water treatment significantly improved the germination of *A. ferox* seeds at 25 °C with a 16-h photoperiod. It is apparent that smoke–water shows similar effects to PGRs (Kulkarni et al., 2006) and therefore smoke solutions are now widely used to enhance seed germination of many plant species (Light and Van Staden, 2004). The PGRs examined were not as effective as smoke–water.

Temperature has a vital influence on early developmental stages of the seedlings. *Aloe ferox* seedlings subjected to different temperatures under controlled conditions showed variations in growth. Alternating temperatures (30/15 °C) were the best for shoot and root growth and increasing the number of leaves. Subsequently, most of the growth parameters were better at 30 °C than the other constant temperatures examined (Table 1). Low (10, 15, and 20 °C) and high temperatures (35 °C) suppressed the growth of seedlings, indicating that these temperatures are not suitable for propagation of *A. ferox*. However, in the case of *A. vera*, it is suggested that this species can be considered as a potential crop in regions with low temperatures with negligible losses in gel production (Saks and Ish-shalom-Gordon, 1995). A study by Rodriguez-Garcia et al. (2007), on the same species, showed that low soil water potential reduced leaf weight, plant growth rate, and leaf number, indicating sensitivity of newly developed leaves to water stress. This indicates that watering level in the soils is important for the growth of *Aloes*. Irrigating seedlings of *A. ferox* once weekly showed a similar response with decline in shoot and root length, number of roots, and fresh weight of seedling. On the other hand, watering of *A. ferox* seedlings three times weekly enhanced most of the growth parameters studied. This result suggests that regular watering frequencies are crucial for growing *A. ferox* seedlings.

**In vitro seed germination and shoot multiplication.** The unsynchronized in vitro germination of *A. ferox* seeds warrants the need for some pretreatments. Because optimizing the in vitro seed germination of this species was not the main objective of this study, it was set aside for future investigation. Of considerable importance is the role the germination media played in vitro shoot multiplication potential of the seedlings. Generally, media high in salt and sugar content reduced germination efficacy and subsequent shoot multiplication potential. This
Table 2. Effect of cytokinin type and concentration on in vitro shoot multiplication and incidence of abnormality in Aloe ferox.3

| Treatment (µM) | Shoot/explant (no.) | Normal shoot/explant (no.) | Abnormal shoot/explant (no.) | Abnormality index |
|---------------|---------------------|-----------------------------|------------------------------|-------------------|
| Control       | 2.1 ± 0.4           | 2.1 ± 0.6                   | 0                            | 0 c               |
| 2.5 BA        | 7.0 ± 1.5           | 4.1 ± 1.2                   | 2.9 ± 0.8                    | 0.71 ± 0.2 a      |
| 2.5 mT        | 8.2 ± 1.7           | 6.8 ± 1.5 bc                | 1.4 ± 0.4                    | 0.21 ± 0.06 b     |
| 2.5 mTR       | 8.5 ± 1.9           | 7.0 ± 1.6                   | 1.5 ± 0.4                    | 0.21 ± 0.07 b     |
| 5 BA          | 10.0 ± 1.8 ab       | 5.1 ± 1.4 c                 | 4.9 ± 1.1 a                  | 0.90 ± 0.029 a    |
| 5 mT          | 12.0 ± 1.4 a        | 10.2 ± 2.4 a                | 1.8 ± 0.6 bc                 | 0.17 ± 0.05 b     |
| 5 mTR         | 13.0 ± 2.0 a        | 10.8 ± 2.5 a                | 2.2 ± 0.9 bc                 | 0.20 ± 0.08 b     |

3Abnormality index = number of abnormal shoots/number of normal shoots. n = 15.

Effect on shoot multiplication of Aloe ferox: (A) shoot clusters in a jar showing some browning problem; (B) shoot clusters derived from five shoot-tip explants.

Conclusions

This study indicates that seeds of A. ferox require a temperature of 21.5 °C for optimum germination. Treating A. ferox seeds with smoke–water (1:500 v/v) may help in improving percentage germination. Alternating temperatures (30/15 °C) and high irrigation frequencies (three times weekly) would be useful to raise seedlings of A. ferox.

In vitro propagation of A. ferox was achieved using a previously developed protocol. It should, however, be noted that the germination medium plays a significant role in the shoot induction competence of the seedlings. Seeds germinated in either distilled water or one-tenth strength MS medium gave a superior shoot induction response.

Literature Cited

Bairu, M.W., W.A. Stirk, K. Doležal, and J. Van Staden. 2007. Optimizing the micropropagation protocol for the endangered Aloe polyphylla. Can meta-topolin and its derivatives serve as replacement for benzyladenine and zeatin? Plant Cell Tissue Organ Cult. 90:15–23.

Bairu, M.W., W.A. Stirk, K. Doležal, and J. Van Staden. 2008. The role of topolins in micropropagation and somaclonal variation of banana cultivars ‘Williams’ and ‘Grand Naine’ (Musa spp. AAA). Plant Cell Tissue Organ Cult. 95:373–379.

Baxter, B.J.M., J. Van Staden, J.E. Granger, and N.A.C. Brown. 1994. Plant-derived smoke and smoke extracts stimulate seed germination of the fire-climax grass Themeda triandra. Environ. Exp. Bot. 34:217–223.

Chukwujekwu, J.C. Fennell, and J. Van Staden. 2002. Optimisation of the tissue culture protocol for the endangered Aloe polyphylla. S. Afr. J. Bot. 68:424–429.

Donaldson, J. 1989. Criteria for listing on Appendix I and Appendix II. Test of the applicability of the criteria, p. 88–89. In: Munoz, M.C. (comp.). Aloe ferox—Proposed revision of Resolution Conf. 9.24 (CoP12 Com.1.3).

Ellis, R.H. and E.H. Roberts. 1981. The quantification of ageing and survival in orthodox seeds. Seed Sci. Technol. 9:373–409.

Hardegere, S.P. and W.E. Emmerich. 1990. Partitioning water potential and specific salt effects on seed germination of four grasses. Ann. Bot. (Lond.) 66:587–595.

Hoagland, D.R. and W.C. Snyder. 1933. Nutrition of strawberry plants under controlled conditions. Proc. Amer. Soc. Hort. Sci. 30:288–296.

International Seed Testing Association. 1999. Biochemical test for viability. Seed Sci. Technol. Supplement 27, 333 p.

Knapp, A. 2006. A review of the trade in Aloe ferox with a focus on the role of the European Union. A TRAFFIC Europe report for the European Commission, Brussels, Belgium.

Kulkarni, M.G., S.G. Sparg, M.E. Light, and J. Van Staden. 2006. Stimulation of rice (Oryza sativa L.) seedling vigour by smoke–water and buteanolide. J. Agron. Crop Sci. 192:395–398.

Light, M.E. and J. Van Staden. 2004. The potential of smoke in seed technology. S. Afr. J. Bot. 70:97–101.

Maunud, P., P. Kariuki, and O. Eyog-Matig. 2004. Threats to medicinal plant species—An African perspective. Proc. of a global synthesis workshop on Biodiversity and Species Extinctions: Managing Risk in a Changing World. IUCN, World Conservation Congress, Bangkok, 17–25 Nov.

Murashige, T. and F. Skoog. 1962. A revised medium for rapid growth and bioassays with tobacco tissue cultures. Physiol. Plant. 15:473–497.

Newton, D.J. and H. Vaughan. 1996. South Africa’s Aloe ferox plant, parts and derivatives industry. TRAFFIC East Southern Africa.

Oliff, H., D.M. Pegtel, J.M. Van Groenendael, and J.P. Bakker. 1994. Germination strategies during grassland successions. J. Ecol. 82:69–77.

Pfab, M.F. and M.A. Scholes. 2004. Is the collection of Aloe peglerae from the wild sustainable? An evaluation using stochastic population modelling. Biol. Conserv. 118:695–701.

Rodriguez-Garcia, R., D. Jasso de Rodriguez, J.A. Gil-Marín, J.L. Angulo-Sánchez, and R.H. Lira-Saldivar. 2007. Growth, stomatal resistance and transpiration of Aloe vera under different soil water potentials. Ind. Crops Prod. 25:123–128.

Sachedina, H. and G. Bodeker. 1999. Wild Aloe harvesting in South Africa. J. Altern. Complement. Med. 5:121–123.

Saks, Y. and N. Ish-shalom-Gordon. 1995. Aloe vera L., a potential crop for cultivation under conditions of low-temperature winter and basalt soils. Ind. Crops Prod. 4:85–90.

Shackleton, C.M. and J. Gambiza. 2007. Growth of Aloe ferox Mill. at selected sites in the Makana region of the Eastern Cape. S. Afr. J. Bot. 73:266–269.
Sheng-zuo, F., S. Li-yi, and F. Xiang-xiang. 2006. Effects of NaCl stress on seed germination, leaf gas exchange and seedling growth of *Pteroceltis tatarinowii*. J. For. Res. 17:185–188.

Sławomir, B. and R. Wiktoria. 2002. Sugars as a metabolic regulator of storage protein mobilization in germinating seeds of yellow lupine (*Lupinus luteus* L.). Acta Physiol. Plant. 24:425–434.

Van der Bank, H., B-E. Van Wyk, and M. Van der Bank. 1995. Genetic variation in two economically important *Aloe* species (Aloaceae). Biochem. Syst. Ecol. 23:251–256.

Van Staden, J., A.K. Jäger, M.E. Light, and B.V. Burger. 2004. Isolation of the major germination cue from plant-derived smoke. S. Afr. J. Bot. 70:654–659.

Van Wyk, B.-E. and G. Smith. 1996. Guide to the *Aloe* of South Africa. Briza Publications, Pretoria, South Africa.

Van Wyk, B.-E., B. Van Oudtshoorn, and N. Gericke. 1997. Medicinal plants of South Africa. Briza Publications, Pretoria, South Africa.