Seed dormancy regulates germination response to smoke and temperature in a rhizomatous evergreen perennial

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Abstract. Seed dormancy status regulates the response of seeds to environmental cues that can trigger germination. Anigozanthos flavidus (Haemodoraceae) produces seeds with morphophysiological dormancy (MPD) that are known to germinate in response to smoke, but embryo growth dynamics and germination traits in response to temperatures and after-ripening have not been well characterized. Seeds of A. flavidus, after-ripened for 28 months at 15 °C/15 % relative humidity, were incubated on water agar, water agar containing 1 μM karrikinolide (KAR1) or 50 μM glyceronitrile at 5, 10, 15, 20, 25, 20/10 and 25/15 °C for 28 days. After incubation at 5, 10 and 25 °C for 28 days, seeds were transferred to 15 °C for another 28 days. Embryo growth dynamics were tested at 5, 10, 15 and 25 °C. Results demonstrated that fresh seeds of A. flavidus had MPD and the physiological dormancy (PD) component could be broken by either glyceronitrile or dry after-ripening. After-ripened seeds germinated to ≥80 % at 15–20 °C while no additional benefit of germination was observed in the presence of the KAR1 or glyceronitrile. Embryo growth significantly increased at 10 °C, and only slightly increased at 5 °C, while growth did not occur at 25 °C. When un-germinated seeds were moved from 5–10 °C to 15 °C for a further 28 days, germination increased from 0 to >80 % in significantly less time indicating that cold stratification may play a key role in the germination process during winter and early spring in A. flavidus. The lower germination (<50 %) of seeds moved from 25 to 15 °C was produced by the induction of secondary dormancy. Induction of secondary dormancy in seeds exposed to warm stratification, a first report for Anigozanthos species, suggests that cycling of PD may be an important mechanism of controlling germination timing in the field.

Keywords: Cold stratification; dormancy cycling; embryo growth; germination stimulant; secondary dormancy; temperature.

Introduction

Seed germination is the initial and most crucial stage in the life cycle of most flowering plants. Seeds of the majority of plant species are dormant at maturity (Baskin and Baskin 2014). Dormancy is an adaptive trait that helps regulate the timing of seed germination ensuring that the chances of seedling survival are greatest. Seeds respond to a host of environmental factors...
Five primary kinds (classes) of seed dormancy are recognized, i.e. physical dormancy (PY), physiological dormancy (PD), morphological dormancy (MD), morphophysiological dormancy (MPD) and combinational dormancy (PY+PD), with PD being the most common (Baskin and Baskin 2004). Seeds with non-deep PD, the least stringent form of PD control, commonly cycle between dormant and non-dormant states in the soil seed bank in response to seasonal changes in environmental conditions (Baskin and Baskin 2004; Chauhan et al. 2006; Long et al. 2011).

Since seed dormancy status regulates the response of seeds to environmental cues that can trigger germination (Flematti et al. 2011; Abdelgadir et al. 2013; Florentine et al. 2016), unravelling the interplay between dormancy status and the efficacy of germination cues is important in resolving the seed germination ecology of species in their natural environments (Tieu et al. 2001a, b; Baker et al. 2005a, b; Merritt et al. 2007; Hidayati et al. 2012) and for managing seeds stored under ex situ conditions (Merritt and Dixon 2011; Erickson et al. 2017).

For seeds of the species-rich flora of southwestern Australia, primary processes involved in dormancy loss include dry after-ripening (i.e. dormancy loss during warm, dry conditions such as those that seeds experience in summer), warm stratification (warm, moist conditions such as those seeds experience in autumn) and wet/dry cycling (conditions common during the sporadic rainfall of early autumn) (Turner et al. 2006, 2009; Merritt et al. 2007; Hidayati et al. 2012). Temperatures of 25–35 °C commonly promote dormancy loss, with seeds then germinating in winter (the period of reliable rainfall) upon the onset of cooler temperatures of 15–20 °C (Bellairs and Bell 1990; Merritt et al. 2007). Dormancy loss through cold stratification is not often reported, although it is known to occur in some species such as Xyris lanata (Merritt et al. 2007).

In fire-prone ecosystems, fire plays a major role in the recruitment of plant species via seed (Dixon et al. 1995; Merritt et al. 2006; Turner et al. 2009). Seedlings emerge en masse from the soil seed bank in the winter growing season immediately following a fire, with seed germination being promoted by chemicals in smoke that are deposited into the soil during the fire (Flematti et al. 2013). Chemicals present in smoke, including karrikins (KAR) and glyceronitrile, along with crude smoke products (aerosol smoke and smoke water) promote the germination of at least 1200 Australian species (Dixon et al. 2009; Flematti et al. 2013, 2015). While likely to produce the same ecological response in the field, the effect of the compounds is species specific, with seeds of some species responding only to KAR, but not glyceronitrile and vice versa (Flematti et al. 2011; Erickson et al. 2017; Lewandrowski et al. 2018).

Recent studies have highlighted the complexities of the germination response to smoke and that seed receptivity to smoke-derived chemicals is strongly regulated by seed dormancy status (Merritt et al. 2007), prior-hydration history (Long et al. 2010) and even seed age (Liyanage and Ooi 2017). For example, in seeds with PD, seed sensitivity to smoke can cycle seasonally as the depth of dormancy varies in response to changing soil temperature and moisture conditions (Baker et al. 2005b; Long et al. 2011). Other seeds appear receptive to smoke regardless of the previous conditions experienced (Merritt et al. 2007), and it has been proposed that seeds can be classified according to whether their response to smoke is inherent or inducible (Long et al. 2011). Thus, to understand the ecology of seed germination in fire-prone floras, studies are required on the role of dormancy-breaking factors in regulating the response to smoke and also on the germination requirements and characteristics of non-dormant seeds and how these determine the expression of the smoke response.

The genus Anigozanthos is a member of the ancient Gondwanan family Haemodoraceae which consists of highly divergent and relictual taxa in southwest Western Australia (Hopper et al. 2009; Ayre et al. 2017). Anigozanthos species produce seeds with MPD. Seeds with MPD have an underdeveloped embryo that must grow inside the seed prior to germination and physiological blocks to germination (i.e. PD) that must also be overcome (Baskin and Baskin 2005; Baskin et al. 2015). Dormancy release and germination of a number of Anigozanthos species have long been known to be associated with fire-related cues including heat and smoke (Dixon et al. 1995; Tieu et al. 2001a, b). More recently, it has been shown that seed germination of Anigozanthos manglesii, A. humilis and A. viridis is promoted by glyceronitrile (Flematti et al. 2011; Downes et al. 2014), a smoke-derived chemical that, in the presence of water, hydrolyzes to release cyanide (Flematti et al. 2011).

Seeds of a few Anigozanthos species after-ripen in laboratory storage, with the germination percentage of a seed population exposed to smoke or glyceronitrile becoming progressively higher during after-ripening and germination in water also increasing, albeit to a lesser extent (Tieu et al. 2001c; Downes et al. 2014). This response indicates that as dormancy is released there is a widening of the conditions suitable for germination.
However, in seeds of several Anigozanthos species, the germination potential varies substantially within and between species in response to dormancy alleviation pretreatments, exposure to a suite of germination stimulants and the post-harvest handling and storage conditions (Tieu et al. 2001c; Merritt et al. 2007; Downes et al. 2014). Seeds of Anigozanthos flavidus provide one such example (Flematti et al. 2011; Downes et al. 2013, 2014; Phillips et al. 2014).

Anigozanthos flavidus is distributed along the most southwestern parts of Western Australia and forms large, densely clumped plants to 3 m tall. This species produces numerous yellow-green flowers with seeds maturing in early autumn (March), and these seeds will drop off from the mother plant after maturity. To date, research on seeds of A. flavidus has focussed on the progression of dormancy loss during soil burial, the effects of after-ripening in the laboratory on dormancy break and the germination response of dormant seeds to smoke-derived products. For example, two recent studies on the germination traits of A. flavidus seeds found that freshly collected seeds were dormant at dispersal (0 % germination), and following a period of dry after-ripening in the laboratory (e.g. 10 weeks at 35 °C, or 0 % germination), and following a period of dry after-ripening in the laboratory on dormancy break and the germination response of dormant seeds to smoke-derived products. For example, two recent studies on the germination traits of A. flavidus seeds found that freshly collected seeds were dormant (average germination 32 %) but highly responsive to glyceronitrile (average germination 86 %) (Phillips et al. 2014).

Although dry after-ripening is thought to be the predominant means of dormancy alleviation in seeds of Anigozanthos spp., the response of the seeds to cold or warm stratification has not been tested. In a study by Downes et al. (2014), after-ripening resulted in >50 % germination of smoke/glyceronitrile-treated A. flavidus seeds, but seeds buried for 1 year in the soil and retrieved in autumn germinated to only 17 % following smoke or glyceronitrile treatment. A similar discrepancy between after-ripened and soil-buried seeds was reported by Tieu et al. (2001c) for seeds of A. manglesii. These results suggest that more complex interactions beyond after-ripening may be influencing dormancy status and germination in the soil environment. In particular, suppression of germination of seeds buried in soil compared to laboratory after-ripened seeds and the apparent variation in dormancy depth across different populations suggest that induction of secondary dormancy is possible under unfavourable conditions.

In this study, we examined the germination traits of A. flavidus seeds aiming to determine whether the (i) incubation temperatures and warm and/or cold stratification will influence the seed germination potential by affecting the elongation of embryos; (ii) seed sensitivity to the smoke-derived products glyceronitrile and/or KAR, is dictated by dormancy status.

**Methods**

**Seeds**

Seeds of A. flavidus were collected in late March 2011 from 50 plants in a natural population near Northcliffe in southwest Western Australia (34°38′32″S, 116°07′25″E), cleaned in the laboratory and then pooled. The annual average temperature and annual precipitation of the Northcliffe area is 15 °C and 1158 mm, respectively. From November to April, the monthly maximum and minimum temperatures are 25-30 °C and 14–18 °C, respectively (http://www.bom.gov.au/climate/averages/tables/cw_009034_All.shtml). Precipitation predominantly falls during winter from June to August (ca. 50 % of total precipitation), with ca. 6 % occurring in summer (from December to February). Cleaned seeds were allowed to after-ripen in a controlled environment room at 15 °C and 15 % RH for 28 months, until germination experiments commenced in July 2013. Mean (±SE) length and mass of individual seeds were 2.39 ± 0.01 mm and 0.65 ± 0.02 mg, respectively. The mass of seeds was determined for seeds stored at 15 °C and 15 % RH. The viability of seeds (four replicates of 100 seeds) was tested via x-ray analysis (Faxitron MX-20 Digital X-ray Cabinet, Tucson, AZ, USA). Final viability was deemed to be 99–100 % due to the presence of uniformly, white/grey x-ray images of seeds and well-developed endosperm.

**Germination of freshly collected seeds**

To confirm the presence and the initial effect of dormancy on seeds of A. flavidus, freshly collected seeds were incubated in Petri dishes containing 0.7 % (w/v) water agar, water agar containing 50 μM glyceronitrile, and assessed for germination at constant 15 °C on a 12/12 h light/dark regime in July 2011. Prior to germination testing, seeds were sterilized in a 2 % (w/v) calcium hypochlorite (Ca(OCl)₂) solution for 30 min and rinsed twice in sterilized distilled water.

**Effects of temperature, KAR, and glyceronitrile on germination of after-ripened seeds**

Seeds from the same accession described above, after-ripened for 28 months, were incubated in Petri dishes containing either: 0.7 % (w/v) water agar, water agar containing 1 μM karrikinolide (KAR), or water agar containing 50 μM glyceronitrile. Glyceronitrile and KAR were used at concentrations that enhanced germination and were within the range of smoking products. Glyceronitrile and KAR were selected as, at this stage, the smoke or glyceronitrile treatment. A similar discrepancy between after-ripened and soil-buried seeds was reported by Tieu et al. (2001c) for seeds of A. manglesii. These results suggest that more complex interactions beyond after-ripening may be influencing dormancy status and germination in the soil environment. In particular, suppression of germination of seeds buried in soil compared to laboratory after-ripened seeds and the apparent variation in dormancy depth across different populations suggest that induction of secondary dormancy is possible under unfavourable conditions.

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were synthesized in pure form following Flematti et al. (2011) and Flematti et al. (2004), respectively. Seeds were incubated at 5, 10, 15, 20, 25, 20/10 or 25/15 °C with a daily 12 h photoperiod of 30 μmol m\(^{-2}\) s\(^{-1}\) 400–700 nm, cool white fluorescent light in growth chambers (Contherm BIOSYN 6000CP, Hutt City, New Zealand). For the alternating temperatures, the high temperature was implemented during the 12-h light period. Seeds were considered to be germinated upon radicle emergence, and germination was scored every 2–3 days for 28 days. Four dishes of 25 seeds were used for each treatment.

**Movement of seeds to 15 °C**

After incubating seeds at 5, 10 and 25 °C for 28 days, no germination was observed. Therefore, all the Petri dishes containing non-germinated seeds were moved to the 15 °C incubator. Subsequent germination was then scored at 2- to 3-day intervals for another 28 days.

**Effects of temperature on embryo growth**

To observe the rate of embryo growth, four replicates of 25 seeds were incubated on water agar, or 50 μM glyceronitrile at 5, 10, 15 and 25 °C and embryo length was determined by dissecting seeds and measuring the embryo and seed lengths under a binocular microscope equipped with an ocular micrometre. For seeds at 15 °C, the embryo length was determined after 0, 3, 6, 9, 12 and 15 days of incubation, and for seeds at 5, 10 and 25 °C embryo length was determined after 28 days of incubation.

**Data analysis**

Generalized linear models (GLMs) with a binomial error structure and logit link function were used to compare the proportional data for the final germination of *A. flavidus* for the temperatures and smoke-related chemical treatments. The time to obtain 50 % germination (\(T_{50}\)) was calculated using the equation (Farooq et al. 2005):

\[ T_{50} = t_j + \left( \frac{N_j - N_i}{N_j - N_i} \right) \left( t_j - t_i \right) \]

where \(N\) is the final germinated seed number, \(N_i\) and \(N_j\) are the cumulative number of seeds germinated by adjacent counts at times \(t_i\) and \(t_j\) respectively, when \(N_i < N/2 < N_j\). We used a one-way or two-way analysis of variance (ANOVA; \(P < 0.05\)) to compare \(T_{50}\) and embryo length (mm) data with temperatures and smoke chemicals as factors. Tukey’s test was used for multiple comparisons when the final germination among treatments was significant. All analyses were carried out using the R statistical platform (R Core Development Team 2014).

**Results**

**Germination of fresh seeds**

Most freshly collected seeds were dormant with a germination percentage of 41 ± 3.4, but dormancy could be broken by the application of glyceronitrile with a germination percentage of 84 ± 2.8 (Fig. 1). Significant differences were obtained between germination of the control and the glyceronitrile-treated seeds (\(P < 0.0001\)).

**Effects of temperature, KAR\(_1\), and glyceronitrile on germination of after-ripened seeds**

For the after-ripened seeds, high germination was obtained under the intermediate temperature regimes of 15, 20 and 20/10 °C. A significant reduction in germination was evident at 25/15 °C while no germination was observed at 5, 10 and 25 °C (Fig. 2). Seed germination of *A. flavidus* was significantly affected by the incubation temperature (df = 3, \(\chi^2 = 100.902\), \(P < 0.0001\)) and smoke stimulants (df = 2, \(\chi^2 = 7.242\), \(P = 0.0268\)), but the interaction of germination stimulants × incubation temperature did not significantly affect seed germination (df = 6, \(\chi^2 = 11.914\), \(P < 0.064\)).

**Effects of temperature, KAR\(_1\), and glyceronitrile on germination speed**

Germination rate was significantly faster at 20 °C and slowest at 25/15 °C (Table 1) and temperature strongly affected \(T_{50}\) (\(F_{3, 12} = 57.187\), \(P < 0.001\)). However, neither smoke chemicals (\(F_{2, 9} = 1.098\), \(P = 0.344\)) nor the interaction of chemicals × temperature (\(F_{6, 36} = 0.607\), \(P = 0.723\)) had a significant effect on germination rate.
Seeds incubated at 5 and 10 °C exhibited significant embryo growth, but there were no growth observed in seeds incubated at 25 °C (Fig. 6). The embryo length of seeds incubated in glyceronitrile was significantly longer than that of seeds in water at 10 °C, but no differences were recorded at the other temperatures. The ANOVA analysis showed that temperature had the greatest effect on embryo growth ($F_{2,9} = 112.368, P < 0.01$), and there were no significant effects of the presence/absence of glyceronitrile ($F_{1,6} = 1.380, P = 0.06$). There was also a significant interaction between temperature and the presence/absence of glyceronitrile on embryo growth ($F_{2,18} = 6.722, P < 0.01$).

**Discussion**

It is well recognized that temperature regulates both seed dormancy break and germination worldwide (Baskin and Baskin 2004; Finch-Savage and Leubner-Metzger 2006; Liyanage and Ooi 2017). Seeds may be dormant at maturity, and then come out of, or go back into, dormancy in response to the moisture and temperature of their environment (Kebreab and Murdoch 1999; Baskin and Baskin 2006; Vandelook et al. 2008; Scholten et al. 2009; Baskin and Baskin 2014; Liyanage et al. 2017). In this study, we showed that the freshly collected seeds of *A. flavidus* have MPD and the PD component could be overcome by either exposure to glyceronitrile, a smoke-derived germination stimulant previously shown to promote germination more consistently in seeds of the Haemodoraceae (Flematti et al. 2011), or by dry after-ripening (e.g. in this study storage at 15 °C/15% RH for up to 28 months was utilized).

Following the dispersal of the *A. flavidus* seeds at the end of March or April, when autumn temperatures are cooling and early season rains commence, PD is most likely alleviated under natural conditions due to a combination of dry after-ripening and periods of stratification when soils are periodically wet (Merritt et al. 2007). The loss of PD in seeds of *Anigozanthos* species, through mechanisms such as dry after-ripening and exposure to fire-related cues, has long been recognized (Tieu et al. 2001c; Flematti et al. 2011; Downes et al. 2013, 2014; Phillips et al. 2014).

Unique to this study, however, is the previously unreported benefit of cold stratification of *A. flavidus* seeds in the promotion of embryo growth and germination at temperatures aligned to the winter season. For some Mediterranean climate species, seeds germinate over a wide range of temperatures (e.g. 5–30 °C), once non-dormant (Zaidi et al. 2010; Martínez-García et al. 2012; Copete et al. 2014; Veiga-Barbosa and Perez-Garcia 2014; Martinez-Baniela et al. 2016; Cross et al. 2017). However, temperatures at which maximum germination...
occurs usually correspond with seasonal conditions that are most favourable for the growth and survival of seedlings (Bellairs and Bell 1990; Mackenzie et al. 2016), and seeds of *Anigozanthos* spp. have consistently been observed to germinate at a narrow temperature range of 15–20 °C, including in this study (Tieu et al. 2001c; Flematti et al. 2011; Downes et al. 2013, 2014). Unsurprisingly, embryo growth also was centred around these optimum germination temperatures. For instance, we found that once PD was alleviated through dry storage, growth of the underdeveloped embryo occurred on moist substrates at ca. 15 °C (Figs 4 and 5). Embryo length significantly increased at 10 °C, but did not protrude the seed coat, and only slightly increased at 5 °C. Embryo growth did not occur at 25 °C, meaning there is strict temperature regulation, aligned to cooler winter season conditions, with respect to initiation of embryo growth. Interestingly, when non-germinated seeds were moved from 5–10 °C to 15 °C for a further 28 days, germination increased from 0 to >80 % in significantly less time (i.e. time to 50 % germination reduced from ca. 13–16 days down to ≤3 days). This indicates that cold stratification may play a key role in the germination process during winter and early spring in *A. flavidus*, whereby embryo growth commences at temperatures <15 °C but is not completed until a shift into the optimum temperature range of 15–20 °C is achieved. This response to simulated seasonal temperatures of southwest Western Australia suggests that germination would be aligned to late winter and early spring when temperature increases are in unison with high soil moisture conditions (sensu Fig. 5 of Merritt et al. 2007).

Although warm stratification breaks dormancy in many species with MPD (Baskin and Baskin 2004), we found the opposite in seeds of *A. flavidus*, whereby seeds exposed to extended periods of warm (25 °C), moist conditions lost the ability to germinate and appeared to enter into secondary dormancy; a first report for the *Anigozanthos* genus. The presence of secondary dormancy is supported by the lack of germination occurring at 25 °C, and

Table 1. Time to obtain 50 % germination (T50) of after-ripened *Anigozanthos flavidus* seeds incubated at 5, 10, 15, 20, 25, 20/10 and 25/15 °C and movement from 5, 10, 25 to 15 °C in a 12/12 h light/dark regime on water (control), KAR, or glyceronitrile. Data in the table represent mean ± SE. No germination was observed at 5, 10 and 25 °C and their T50 are expressed as /. The 5→15, 10→15 and 25→15 represent seeds were moved from 5, 10 and 25 °C to 15 °C, respectively.

| Temperature (°C) | 5   | 10  | 15  | 20  | 25  | 10/20 | 15/25 | 5→15 | 10→15 | 25→15 |
|------------------|-----|-----|-----|-----|-----|-------|-------|------|-------|-------|
| H2O              | /   | /   | 16.4 ± 0.2 | /   | 16.8 ± 0.3 | 17.7 ± 0.8 | 10.2 ± 0.4 | 2.6 ± 0.2 | 18.4 ± 0.7 |
| KAR              | /   | /   | 16.4 ± 0.2 | 13.7 ± 0.1 | /   | 17.5 ± 0.3 | 17.9 ± 0.7 | 10.4 ± 0.4 | 2.8 ± 0.1 | 18.0 ± 0.7 |
| Glyceronitrile   | /   | /   | 16.2 ± 0.2 | 13.9 ± 0.1 | /   | 17.3 ± 0.3 | 18.8 ± 0.7 | 10.3 ± 0.3 | 3.2 ± 0.3 | 17.7 ± 0.7 |

Figure 4. Embryo growth (mean ± SE) dynamics of after-ripened *Anigozanthos flavidus* seeds incubated at 15 °C in Petri dishes containing 0.7 % (w/v) water agar (control, white circle) or water agar containing 50 μM glyceronitrile (black circle). The embryo length was determined after 0, 3, 6, 9, 12 and 15 days of incubation. Four dishes of 100 seeds were used for each sampling time point.

Figure 3. Mean germination (% ± SE) of after-ripened *Anigozanthos flavidus* seeds cold stratified (5 °C, 10 °C) or warm stratified (25 °C) for 4 weeks in water, KAR, or glyceronitrile and then moved to 15 °C for an additional 4 weeks with seed germination at 15 °C as control. Before movement, no germination occurred at 5, 10 and 25 °C. Different letters indicate significant differences (P < 0.05) between mean germination time across all treatments.
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40–50 % germination occurring after shifting seeds from 25 to 15 °C. Further, the embryos of non-germinated A. flavidus seeds under these conditions were assessed for viability post-germination testing and were confirmed to be firm, white, and therefore, presumed viable. It should be noted that our assessment of dormant viable embryos from non-germinated A. flavidus seeds was based on a cut test, which has been previously identified as a robust method for dormant species in Australia (Ooi et al. 2004). Induction of non-dormant seeds into secondary dormancy (i.e. MD → MPD) by warm stratification is not uncommon. For instance, it has been reported in seeds of Delphinium fissum subsp. sordidum from the Iberian Peninsula, southwestern Europe (Herranz et al. 2010). Further, secondary dormancy in Narcissus alcaracensis seeds from southern Spain, induced by high temperature, could be broken by certain cold stratification treatments (Herranz et al. 2017). These examples of cycling between a non-dormant state and secondary dormancy, induced by warmer temperatures, indicate that seed germination of A. flavidus is synchronized with the beginning of winter.

In unison with dormancy cycling, fire-related germination cues have also been shown to assist in field recruitment of many Mediterranean species (Dixon et al. 1995). The most commonly applied fire-related chemicals stimulating seed germination are karrikinolide (KAR,) (Flematti et al. 2004, 2005; Daws et al. 2007; Stevens et al. 2007) and glyceronitrile (Flematti et al. 2011; Downes et al. 2013; Phillips et al. 2014). These germination stimulants are widely used to study the influence of fire on seed dormancy and plant recruitment patterns in both conservation and restoration programs (Phillips et al. 2014; Commander et al. 2017; Cross et al. 2017; Erickson et al. 2017; Lewandrowski et al. 2018). Changes of seeds in sensitivity to smoke with alleviation of seed dormancy have been reported in several species (Brown and van Staden 1997; Baker et al. 2005b; Merritt et al. 2007). Seeds of A. manglesii become more smoke responsive with increasing shelf storage time (Tieu et al. 2001c). Seed germination of Actinotus leucocephalus (Apiaceae) gradually increases with the combination of increasing laboratory storage time and smoke water treatment (Baker et al. 2005b). Significant promotion of seed germination by glyceronitrile in A. flavidus (Fig. 1) has been previously reported in freshly collected (Phillips et al. 2014) and after-ripened seeds (Downes et al. 2013, 2014). In our current study, glyceronitrile slightly promoted germination of after-ripened A. flavidus seeds but not significantly. As shown in Figs 4 and 6, significant interaction between temperature and glyceronitrile on embryo growth was observed. This demonstrated that glyceronitrile promoted embryo growth only at certain temperatures. Therefore, it is likely that the dormancy status of the seeds, and hence their response to some of the germination stimulating treatments, changed as a result of dry storage at 15 °C (after-ripening) for 28 months.

Figure 5. Embryo growth for after-ripened seeds of Anigozanthos flavidus in this study was demonstrated by continually measuring the increase in length (mm) of embryos during incubation at suitable temperatures (see Fig. 4). Examples of an intact seed (A), embryo before growth (B) and embryo before germination (C) are shown visually here. Scale bar = 1 mm.

Figure 6. Embryo length (mean ± SE) of after-ripened Anigozanthos flavidus seeds incubated at 5, 10 and 15 °C for 28 days on filter paper moistened with water or glyceronitrile. Control represents mean embryo length of seeds prior incubation. Bars with different letters are significantly different at P < 0.05.
Conclusions
Fresh seeds of *A. flavidus* possess a type of non-deep MPD that could be broken by smoke-derived chemical glyceronitrile. However, after-ripened seeds of *A. flavidus* germinated readily at temperatures 15–20 °C and neither glyceronitrile nor KAR1 improved germination percentage or speed following loss of PD. Embryo growth could occur at 5 and 10 °C but seeds could not germinate until the temperature is about 15 °C in winter when precipitation is also suitable for germination and establishment of *A. flavidus*. Temperatures above 25 °C induced secondary PD in seeds and no embryo elongation occurred. The dormancy cycling of MPD→MD→ND→PD in *A. flavidus* seeds may be an important mechanism of controlling germination timing in the field.

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Contributions by the Authors
H.M., T.E.E. and D.J.M. designed the experiments. H.M. conducted the experiments and analysed the data. H.M., T.E.E. and D.J.M. interpreted the data. H.M., T.E.E. and D.J.M. wrote the manuscript.

Conflict of Interest
None declared.

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