Regulation of seed germination by diurnally alternating temperatures in disturbance-adapted banana crop wild relatives (Musa acuminata)

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Abstract
Seed conservation of banana crop wild relatives (Musa L. spp.) is limited because of lack of knowledge about their germination ecology. Musa acuminata Colla, the most important banana crop wild relative, is distributed in tropical and subtropical Asian and Pacific rainforests and colonizes disturbed sites. The role of temperature in stimulating/inhibiting germination to detect disturbance when canopy gaps are formed is not well known. We assessed seed germination thermal requirements of three subspecies of M. acuminata using nine seed accessions which had been stored in the Millennium Seed Bank. Diurnally alternating temperature cycles were almost completely essential for germination compared with constant temperatures. Germination was optimal when the upper temperature of a diurnal cycle was at 35°C; the lower temperature of the cycle was less important. Subspecies occurrence coordinates were used to extract climate temperature data which were then compared against the temperature requirements for germination from our experiment results. Maximum temperatures of the warmest month across subspecies ranges were close to but below optimal germination temperatures, as were diurnal ranges, suggesting soil-warming at the micro-climate level following gap creation is important for M. acuminata seed germination. Additionally, pre-treatment for 3 months at 60% relative humidity at constant 25°C improved germination from 14 ± 10 (mean, standard deviation) to 41 ± 29% suggesting a period in the soil seed bank under the canopy may increase sensitivity to alternating temperature cycles. Overall viability was low (49 ± 28%), and considerable variance was caused by the different accessions. Germination remained somewhat inconsistent.

Introduction
Multiple biotic and abiotic issues threaten the future of banana production (Ramirez et al., 2011; Ploetz and Evans, 2015), the potential consequences of which will likely have a large impact on the nutrition and livelihoods of many millions of people (Ploetz et al., 2015). The limited gene pool of clonal crops like bananas (Perrier et al., 2011) means they have evolved little resistance to many pathogens, making them particularly vulnerable (Strange and Scott, 2005; McKey et al., 2010). Thus, there is urgent need for the genetic resources present in banana crop wild relatives (CWRs) (Musa L. spp.) to be conserved and made available to breeders (Dempewolf et al., 2017).

Storing and accessing genetic resources as seeds is an efficient method for ex situ conservation (Li and Pritchard, 2009; Convention on Biological Diversity, 2012; FAO, 2012; Laliberté, 2016). The management of seed genetic resources requires their routine germination for viability monitoring (FAO, 2014), as does accessing plants for phenotyping or breeding; for bananas, low and inconsistent germination rates are a considerable limitation to these (Laliberté, 2016; Brown et al., 2017). Improvement in the ability to germinate banana seeds will therefore have important applications for food security and plant genetic resource conservation (Panis et al., 2020).

There are around 80 species in the genus Musa (Govaerts and Häkkinen, 2006); cultivated bananas mostly derive from two of these: Musa acuminata Colla and Musa balbisiana Colla...
(Boonruangrod et al., 2008; De Lange et al., 2009; Perrier et al., 2009, 2011). Overall, *M. acuminata* is the most important contributor of genetic material for cultivated bananas (Raboin et al., 2005; Martin et al., 2020); for instance, they are the sole contributor to the Cavendish group which accounts for approximately 47% of global production (FAO, 2019). There are ten accepted subspecies (used hereafter to also include varieties and subspecies) of *M. acuminata* (Govaerts and Håkkinen, 2006), which are distributed in Asia, northern Australasia and Melanesia (Perrier et al., 2009).

In contrast to *M. balbisiana* (McGahan, 1961a, 1961b; Stotzky et al., 1961, 1962; Stotzky and Cox, 1962; Bhat et al., 1994), surprisingly little attention has been paid to *M. acuminata* seeds. In the 1950s, Simmonds (1952, 1959) examined both *M. acuminata* and *M. balbisiana* with various non-invasive treatments (looking at ripeness, maturity and moisture content) and invasive treatments (using chipping, scourching, soaking and acid treatments). The physical permeability of *M. acuminata* seed coats to water has received most attention but still remains unresolved (Darjo and Bakry, 1990; Wattanachaiyingenroen and Turner, 1990; Fortescue and Turner, 2011; Puteh et al., 2011). Significantly, the thermal requirements for germination of *M. acuminata* seeds have not been investigated.

*Musa* spp. are tall forest herbs of tropical to subtropical forests. Their inflorescences are pollinated by bats, birds and probably insects (Itino et al., 1991; Liu et al., 2002), and seeds are dispersed by bats, birds and mammals (Ge et al., 2005; Tang et al., 2005, 2007; Marod et al., 2010). They are observed in disturbed sites such as farm or plantation edges, craggy cliffs and steep mountain gullies, or roadside and track edges. Specifically, *M. acuminata* is known to colonize and dominate disturbed sites (Zhang et al., 2000; Shi et al., 2002; Meng et al., 2012). *Musa* are therefore widely recognized as disturbance-adapted ‘jungle weeds’ (Simmonds, 1959).

Several adaptations may be expected in *M. acuminata* seeds to facilitate germination following disturbance. Firstly, an environmentally regulated gap detection mechanism may stimulate germination immediately following disturbance and, conversely, prevent germination prior to disturbance (Raich and Khoon, 1990; Vázquez-Yanes and Orozco-Segovia, 1993). For larger-seeded disturbance-adapted species, germination may be stimulated by changes in micro-climate temperature (Pearson et al., 2002; Aud and Ferraz, 2012). This may involve changes in diurnal temperature fluctuations, as soil is exposed to sunlight when a canopy gap forms and therefore warming and cooling fluctuations are altered (Vázquez-Yanes and Orozco-Segovia, 1982; Aud and Ferraz, 2012; Poschlod et al., 2013; Jaganathan, 2018).

Sensitivity to differences in alternating temperature range plays an important role in limiting seedling growth under unfavourable conditions (Kos and Poschlod, 2007; Saatkamp et al., 2011). Secondly, the hard seed coat of *Musa* spp. may limit imbibition and delay germination by physical dormancy. This may be relieved according to environmental factors in relation to disturbance, including temperature and moisture regimes, as well as changes in the ecological community such as predators and dispersers. Alternatively, after-ripening or stratification (a pretreatment period prior to dispersal and germination) may delay germination immediately after dispersal as observed for some seeds of tropical rainforests (Vázquez-Yanes and Orozco-Segovia, 1993, and references therein).

The aim of this study is to test hypotheses that *M. acuminata* seed germination is regulated in accordance with expectations of disturbance-adapted tropical species. In particular, we focus on germination responses to temperature, including diurnally alternating temperature regimes as a mechanism for gap detection. Therefore, we (1) describe the morphology and mass of the seeds used in this study, (2) assess whether seed coats are permeable to water, (3) investigate optimal temperature regimes for seed germination, (4) investigate whether stratification relieves dormancy, and finally (5) evaluate the optimal temperature regimes from our germination tests against the macro-climate temperatures of wider distributions of the subspecies under investigation. However, because the seeds used in this study have undergone processes which may influence behaviour, namely drying and sub-zero temperature storage (Baskin et al., 2006), a true definition of germination and dormancy class is not possible, but rather the aim is application to seed conservation.

**Materials and methods**

**Plant material**

Nineteen seed accessions of four *M. acuminata* subspecies were used for the initial viability assessment (Fig. 1 and Supplementary Table S1). Following the viability assessment described below, nine of these accessions, which consist of three subspecies, were selected for a series of experiments (*M. acuminata* subsp. *acuminata*, *M. acuminata* subsp. *malaccensis* and *M. acuminata* subsp. *microcarpa*; Fig. 1 and Table 1). Seeds were collected from ‘Malaysian peninsular rainforest’ and ‘montane rainforest’ ecoregions in the ‘tropical and subtropical moist broadleaf forest’ biome (Olson et al. 2001). Mean annual temperature and mean annual precipitation at the collecting locations are 26.4 ± 0.7°C and 2332 ± 342 mm (mean and standard deviation), respectively (Fick and Hijmans, 2017). Seeds were collected in 2015 and 2016 and provided to the Millennium Seed Bank (MSB) by the Malaysian Agricultural Research and Development Institute.

Seeds derive from one or two plants per accession. Populations were wild in origin and occurred either in secondary rainforest, or oil palm/orchard plantation edges. Bunches were only collected from light green to yellow fruits. Fruits from unhealthy or injured plants were avoided. Cut tests were carried out at the time of collection to assess seed maturity, and only bunches with mature seeds were collected. Seeds were considered mature if the endosperm was dry and powdery, as opposed to wet or milky, and embryos were well developed with a mushroom-like capitellate shape. Seeds were extracted by hand from ripe fruit on return to the laboratory, and the pulp containing seeds was washed thoroughly using running tap water and sieves, until all the flesh was removed. Unripe fruits were left in the laboratory at room temperature (~25°C) to ripen until they began to yellow and soften and then seeds were removed in the manner described above. After extraction, seeds were air-dried in the laboratory at room temperature and then further dried in a large sealed plastic drum containing silica gel to a maximum of 25% equilibrium relative humidity (eRH). Seeds were then packed in sealed aluminium envelopes and stored for an average of 6 months at 4°C until air shipped to the MSB. On arrival at the MSB seeds were further dried at 15% eRH and 18°C in a dry room to approximately 7% moisture content. Seed mass of dried seeds was measured by weighing five replicates of 50 seeds. Seeds were then sealed in airtight glass containers and stored at −20°C for 12–18 months prior to use in this study. Before use, selected accessions were removed from cold storage for at least 24 h to

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equilibrate to 18°C in a dry room at 15% relative humidity (RH). Morphological observations were recorded by dissecting ten seeds per accession and inspecting using a binocular microscope. Seed mass was measured using five samples of 50 seeds.

Viability

Post-storage seed viability was assessed using two methods to improve estimation methods for this and future studies. The first method was the tetrazolium chloride test (TTC), following the approach of Leist and Krämer (2011). Seeds were imbibed on agar for 3 d at 20°C. Then, a proportion of the testa was removed using a scalpel on two lateral sides to expose the endosperm. Seeds were then soaked in 1% buffered 2,3,5-triphenyl tetrazolium chloride (pH6–8) for 2 d at 30°C in the dark. Staining patterns were then recorded. Embryos that completely stained dark red, or that showed dark red staining at the embryonic axis were considered viable; light pink staining or white embryos were considered unviable. Fifty seeds per accession were tested.

Secondly, embryo rescue (ER) was carried out as a viability test. In a laminar flow, embryos were extracted from seeds. This was done using an incision in the seed coat next to the micropyle and manipulating the seed in order to split the testa, the embryo was then gently removed. Embryos were subsequently transferred onto autoclaved half MS medium (Murashige and Skoog, 1962) in tubes using long forceps with the haustorium in contact with medium and the embryonic axis upwards. Tubes containing embryos were incubated in the dark at 27°C for 14 d after which they were put in a growth chamber in the light at 27°C for an additional 14 d. Six possible observations were recorded: shoot, callus, blackened colouration, contamination, no change or no embryo as observed during extraction. Ten seeds per accession were tested.

Imbibition

Seed coat permeability to moisture was tested by assessing mass change of dry seeds during imbibition. Seeds were either left intact and incubated at 25°C, or scarified by removing a sliver of lateral seed coat with a scalpel in order to expose the white seed endosperm (3.36 ± 1.35 mg) and incubated at 25°C, both with a 12 h light/dark regime. To imbibse seeds, they were placed on agar with the micropyyle in contact with 1% agar in Petri dishes (100 × 150 mm). Seeds were weighed individually on days 0, 1, 2, 5, 7 and 21. Prior to measurement, surface moisture and agar were blotted off with tissue paper. Twenty-five seeds from accession M. acuminata subsp. malaccensis (882899) were used for each treatment. This accession was selected because it had the most seeds. The increase in mass percentage was calculated individually for each seed and plotted against time in order to estimate equilibrium.

Germination and temperature

Nine accessions were selected for germination tests based on adequate viability and availability (Table 1 and Supplementary Table S1). Incubator temperature regimes were based on the climate at the location of collection. Temperatures were extracted from WorldClim version 2.0 (Fick and Hijmans, 2017) using seed accession collection coordinates to extract climate data at 30 arc seconds resolution. The collecting locations’ mean monthly temperature to the nearest 5°C was selected for a constant incubation regime: 25°C; whilst alternating temperatures were based on the monthly maximum and minimum temperatures where the upper temperature was rounded both up and down: 35/20 and 30/20°C. Minimal temperature seasonality occurred in the climate data for the collecting sites (mean coefficient of variability 1.81 ± 0.16), so no seasonal differences were included in the design.

Seeds were placed on moist sand (100 g fine silica sand and 14 ml of deionized water) in sealed square plastic boxes (100 × 100 × 20 mm) which were then sealed in clean plastic sealable bags to minimize moisture loss and contamination. These were then placed in the corresponding incubators with 12 h at each temperature. Photoperiod was also on 12 h cycles, light was during the warmer period and darkness during the cooler; light quantity was approximately 7 μmol m⁻² s⁻¹. Twenty-three to fifty-three seeds were used in one to three replicates per accession. Sample sizes were constrained by limited seed availability. Germination tests were scored every 7 d, and tests were continued for 6 months to account for potential long sporadic germination times previously observed and to ensure reasonable maximum germination. Germination was defined by radicle emergence. Germinated seeds were removed at scoring.

Thermogradient

For one accession of M. acuminata subsp. malaccensis (accession 882899) (the accession with the most seeds), 25 temperature regimes were tested using a thermogradient plate (GRD1, Grant Instruments Ltd, Cambridge, UK). All combinations of constant
and alternating temperatures from 15 to 35°C with an interval of 5°C were used. An additional six temperature regimes were tested in incubators to 40°C, with all combinations of 15–40°C under 5°C temperature intervals. Temperature was cycled on a 6/18 hourly basis to allow for a comparison of short and long periods at the warmer/cooler part of the cycles estimated from local temperature data (Meteoblue, 2020). Accordingly, for each temperature regime, there was a sample that had the hotter temperature at 6 h and one at 18 h, except for the samples at 40°C which only had the hotter temperature for 6 h because of a limitation of incubators. Incubation was carried out in the dark because light could not be controlled differently in the thermogradient plate for the short and long cycles, and seeds were previously observed to germinate in the dark. For each condition, 30 seeds were placed on 10 g of silica sand with 8 ml of deionized water in a circular Petri dish (100 × 15 mm), and dishes were sealed with film to avoid moisture loss. An additional 1–2 ml of water was added if the colour of the sand lightened due to water loss. Tests were scored in daylight every 7 d. These germination tests were continued for 70 d.

### Germination and pre-treatment

To simulate the transition of seeds in the soil seed bank before and after gap formation, seeds were incubated (‘stratified’) at constant 25°C and then moved to alternating 30/20°C after 3 months. There were two levels of stratification at 25°C, seeds were either placed on moist sand as described above (at 100% RH), or suspended in sealed jars over lithium chloride solutions controlled to be at 60% RH (60 g LiCl in 200 ml deionized water), for these a humidity meter was placed into the jar with the seeds to be at 60% RH (60 g LiCl in 200 ml deionized water), for 7 d to ensure the correct RH. After 3 months, seeds were then incubated for three further months at 30/20°C light/dark as previously described. Scoring of germination tests was carried out as previously described. All germination tests were carried out at the MSB.

### Macro-climate assessment

We composed a dataset of occurrence records of wild M. acuminata subspecies from 16 sources including recent field missions (Supplementary Fig. S1 and Supplementary Table S2). Accurate locality descriptions without coordinates were georeferenced using Google Earth pinpoints (Google Earth, 2018). Duplicate records, outliers and zero coordinates were detected and removed with the online tool CoordinateCleaner (Ziarka et al., 2019). This resulted in a dataset consisting of 222 occurrence records of nine (of the ten) M. acuminata subspecies. Coordinates from these were used to extract temperature-related bioclimatic data from WorldClim version 2.0 (Fick and Hijmans, 2017) to compare to the optimal germination temperatures from the previous germination tests, and to look for variation between subspecies. Additionally, precipitation across subspecies distributions was extracted to assess further climate variation by taxa.

#### Statistical analysis

Seed mass increase during imbibition was plotted against time, and a point was selected where equilibrium with the environment was reached across treatments. Percentage mass increase at this point for the two treatments was compared using a two-sample t-test. Descriptive indices were calculated for the germination data in the R package GerminaR (Isla et al., 2019). These included germination percentage (Labouriau and Valadares, 1983), mean germination rate defined as the reciprocal of the average time to germination (Ranal and Santana, 2006), mean time to germination (Czabator, 1962) and synchrony index (Primack, 1980; Ranal and Santana, 2006). Generalized linear models (GLMs) were made to analyze final germination with binary responses. Models had quasibinomial error structure (to account for overdispersion) and logit link functions. Overdispersion of models were assessed by simulating residuals from the fitted models and comparing the standard deviation of the simulated and actual data using the DHARMa package in R (Hartig, 2019). Maximum models, with fixed factors of accession, subspecies, viability and a random factor of temperature treatment, were fitted. Minimum adequate models were achieved by removing factors according to ANOVA and Chi-squared tests. Post hoc contrast analysis was carried out between treatments and controls using multiple comparisons of means and Tukey contrasts. In all cases, effective sample sizes were used which corrected sample sizes according to the viability estimates from the TTC test result. This viability measure was used because it was not possible to correlate the two viability measures (ER and TTC) and the sample size used in the TTC was greater than for ER (50 rather than

### Table 1. M. acuminata seed accessions selected for use in germination tests

<table>
<thead>
<tr>
<th>Accession</th>
<th>Subspecies</th>
<th>State</th>
<th>Mother plants</th>
<th>Date collected</th>
<th>Date donated</th>
</tr>
</thead>
<tbody>
<tr>
<td>882741</td>
<td>malaccensis</td>
<td>Pahang</td>
<td>1</td>
<td>20/10/2015</td>
<td>23/06/2016</td>
</tr>
<tr>
<td>882763</td>
<td>acuminata</td>
<td>Pahang</td>
<td>1</td>
<td>21/10/2015</td>
<td>23/06/2016</td>
</tr>
<tr>
<td>882800</td>
<td>acuminata</td>
<td>Negeri Sembilan</td>
<td>2</td>
<td>11/05/2015</td>
<td>23/06/2016</td>
</tr>
<tr>
<td>882811</td>
<td>acuminata</td>
<td>Negeri Sembilan</td>
<td>2</td>
<td>11/05/2015</td>
<td>23/06/2016</td>
</tr>
<tr>
<td>882833</td>
<td>acuminata</td>
<td>Selangor</td>
<td>1</td>
<td>19/11/2015</td>
<td>23/06/2016</td>
</tr>
<tr>
<td>882844</td>
<td>acuminata</td>
<td>Selangor</td>
<td>1</td>
<td>19/11/2015</td>
<td>23/06/2016</td>
</tr>
<tr>
<td>882888</td>
<td>malaccensis</td>
<td>Johor</td>
<td>1</td>
<td>30/12/2015</td>
<td>23/06/2016</td>
</tr>
<tr>
<td>882899</td>
<td>malaccensis</td>
<td>Johor</td>
<td>1</td>
<td>30/12/2015</td>
<td>23/06/2016</td>
</tr>
<tr>
<td>928500</td>
<td>microcarpa</td>
<td>Pahang</td>
<td>1</td>
<td>20/07/2016</td>
<td>31/03/2017</td>
</tr>
</tbody>
</table>

The full set of accessions used for viability testing is in Supplementary Table S1. Date donated is accessioning date on arrival at the MSB.
Results

Morphology and seed mass

Observations showed that *M. acuminata* seeds have two chambers within a two-layer integument (Fig. 2A). The larger first chamber contains the embryo and endosperm. Embryos are undifferentiated and capitate, 1–2 mm in size and the embryonic axis extends into the micropylar collar (Fig. 2B). The cotyledonary haustorium of the embryo extends below the embryonic axis and is surrounded by a powdery white endosperm. Above the embryo is a micropyle, which is filled by a micropyle plug and forms an operculum. The second chamber consists of a chalazal mass. Seed mass was 40.45 ± 6.80 mg (mean and standard deviation, used from hereon).

Viability

Viability was 49 ± 28% according to the TTC test and 28 ± 29% for the embryo recue (i.e. the ‘shoot’ category; Supplementary Table S1). No linear relationship could be found between the two parameters of viability ($r^2 = 0.055, P = 0.811$). Embryos that did not germinate mainly remained white (Supplementary Fig. S2). This means that viability was low because embryos were most likely dead, rather than infested by insects, fungal/bacterial contaminated or lacking embryos. The combined results of the viability tests allowed us to select the nine accessions used for germination tests.

Imbibition

During imbibition, both scarified and non-scarified seeds rapidly increased mass in 1 d and reached equilibrium after 5 d or earlier (Fig. 3). The difference in percentage mass increase was not statistically significant between scarified (49.6 ± 11.8%) and non-scarified seeds (53.3 ± 8.7%, two-sample *t*-test, $t = 1.1189, df = 33.145, $P = 0.27$). The moisture content of non-scarified seeds increased from 7% on day 0 to 36% after 21 d imbibition.
Germination and temperature

Seeds germinated to higher levels at alternating temperatures compared with constant temperature (Table 2 and Fig. 4). Optimal germination was at 35/20°C ($P = 0.006$) followed by 30/20°C ($P = 0.031$) compared with constant temperature at 25°C; however, there was no significant difference between 35/20 and 30/20 ($P = 0.164$) following multiple comparison of means using Tukey contrasts. The minimum adequate GLM included only treatment temperatures, as germination was highly variable between accessions in all tests. Mean time to germination under optimal temperature (35/20°C) was 56 d, and this was highly variable ($±39$ d).

**Table 2.** Indices from germination results from selected *M. acuminata* accessions

<table>
<thead>
<tr>
<th>Treatment (°C)</th>
<th>Final germination (%)</th>
<th>Time to germination (d)</th>
<th>Germination rate</th>
<th>Synchrony index</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean</td>
<td>SD</td>
<td>Mean</td>
<td>SD</td>
</tr>
<tr>
<td>25</td>
<td>1.19</td>
<td>3.40</td>
<td>85</td>
<td>67.88</td>
</tr>
<tr>
<td>30/20</td>
<td>14.38</td>
<td>10.20</td>
<td>41.17</td>
<td>43.87</td>
</tr>
<tr>
<td>35/20</td>
<td>25.63</td>
<td>27.47</td>
<td>55.69</td>
<td>39.18</td>
</tr>
<tr>
<td>100% RH$^a$</td>
<td>8.22</td>
<td>12.79</td>
<td>41.44</td>
<td>46.90</td>
</tr>
<tr>
<td>60% RH$^a$</td>
<td>41.54</td>
<td>29.26</td>
<td>25.62</td>
<td>12.06</td>
</tr>
<tr>
<td>60% RH control$^b$</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
</tr>
</tbody>
</table>

Germination time is mean time to germination in days (Czabator, 1962), germination rate defined as the reciprocal of the average time to germination (Ranal and Santana, 2006), and synchrony index define by Primack (1980) and Ranal and Santana (2006). Alternating temperatures were for 12 h thermo and photo cycles. Germination tests were continued for 6 months. Results were corrected to take into account viability assessment with previous TTC test ($n = 23–53$ seeds in 1–3 replicates).

$a$Pre-treated by incubation at constant 25°C at two levels of RH for 3 months prior to 30/20°C incubation for 3 months at 100% RH.

$b$Incubated only at constant 25°C.

**Fig. 4.** Final germination percentages after 6 months incubation at different conditions for nine *M. acuminata* accessions (subsp. *acuminata*, *malaccensis* and *microcarpa*). (A) Alternating temperature regimes compared with control of constant temperature (30/20, $P = 0.031$; 35/20, $P = 0.006$). (B) Pre-treatment for 3 months at 25°C (60% RH, $P = 0.007$) prior to incubation for 3 months at 30/20°C. Incubation was on moist sand (100% RH) unless otherwise stated. Alternating temperatures (30/20 or 35/20°C) were on 12 hourly cycles. Stars indicate $P$-values (* = <0.05, ** = <0.01) from a GLM with quasibinomial error structure and logit link using the number of seeds germinated and the number of seeds that did not germinate, against the control. Final germination percentages are corrected to take into account viability assessment with previous TTC test ($n = 23–53$ seeds in 1–3 replicates).

**Germination and temperature**

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**Thermogradient**

Maximum germination occurred equally at 35/25°C (6/18 h) and 35/30°C, followed by 30/20°C (90 and 40%, respectively; Fig. 5).
No germination occurred at any constant temperature, or at temperatures below 20°C. There was no germination when the higher temperature was for the longer period (18 h), apart from at 30/35°C. In the GLM, only the short hotter temperature had a significant positive effect on the number of seeds germinated ($P < 0.001$), and all other parameters (long temperature, the average temperature or the temperature differential) could be removed from the model without reducing the explanatory power.

Pre-treatment

Pre-treating the seeds for 3 months at 60% RH increased germination to 41 ± 29% from 14 ± 10% ($P = 0.007$; Table 2 and Fig. 4), but the same treatment at 100% RH did not have an effect ($P = 0.423$). The minimal adequate model only included the treatment, and there was therefore no discernible effect of subspecies.

Macro-climate

Based on our combined germination test results compared with temperature extracted from climate data, we see that the germination temperature requirements are warmer and have a wider diurnal range than estimated from subspecies distributions (Fig. 6). For instance, the temperature of the hotter part of the diurnal cycle from the germination experiments was 35°C; this is close to but warmer than the maximum temperature of the warmest month for all taxa. There were no records at all as high as 35°C, and a few records were close. For diurnal range, no records were as high as the 15°C we observed from the germination test results. Mean diurnal range across taxa distributions was 8 ± 1°C (Fig. 6), and the widest range was 11°C.

Discussion

Stored *M. acuminata* seeds had an almost absolute requirement for alternating temperatures. Within a seed accession, there was non-uniform sensitivity to alternating temperatures, such that by pre-treating seeds with a period of constant temperature, higher final germination percentages were achieved than without such a pre-treatment. Seeds had hard coats, but we found that these were permeable to water.

Our results were comparable to many tropical disturbance-adapted species (reviewed by Vázquez-Yanes and Orozco-Segovia, 1993; Baskin and Baskin, 2014). Within the Musaceae, *M. acuminata* thermal requirements were similar to *M. balbisiana* (Stotzky et al., 1961; Stotzky and Cox, 1962). In these studies, there was also an almost absolute requirement for alternating temperatures, with maximum germination at 35°C for 5 h (mean 59%), and low temperature was 15°C for 19 h (mean 70%). Additionally, low levels of germination of *M. balbisiana* occurred following short and even singular exposure to alternating temperature, but continuous cycles were required for maximum germination (Stotzky and Cox, 1962). As a combination, germination was optimal (80%) at 5 h of 35°C and 19 h of 12°C. The cooler lower temperature requirement for *M. balbisiana* reflects
the subtropical and at times higher elevation distribution of *M. balbisiana* – East India to Yunnan, China.

At the macro-climate scale, temperature and diurnal ranges were rather similar across subspecies distributions (Fig. 6 and Supplementary Fig. S3). In general, temperatures estimated from the climate model (WorldClim) were cooler than in our germination tests; diurnal range was also smaller than may be optimal for germination. This suggests that micro-climate, rather than solely macro-climate, is important in germination of *M. acuminata*. This is because soil temperature, when exposed to direct sunlight, may heat considerably more than the air temperature usually used in climate data and modelling. Furthermore, this observation supports our expectations that seeds are adapted to specifically detect gaps in the forest canopy following disturbance.

Specific optimum temperature fluctuations were different between our two experiments: 5.7 ± 3.9°C for the thermogradient plate experiment and 15°C in the other germination tests, suggesting a degree of plasticity in fluctuation requirement. Pearson et al. (2002) proposed four categories of response to alternating temperature dynamics according to seed mass. Using these same categories, our results place *M. acuminata* into category three, where there is a positive response to increasing temperature fluctuation in the range of 0–16.7°C, but there is no dramatic optimum or cut-off point. For this group, seed mass was the heaviest category in their sample (20.9 ± 14.2 mg, mean and standard error), *M. acuminata* seeds were even heavier than this (40.45 ± 6.80 mg).

At the micro-climate scale, soil mean temperature and diurnal fluctuation are dependent on several factors, including topography, canopy, litter, air temperature and solar irradiance (Kang et al., 2000; Saatkamp et al., 2011; Bilgili et al., 2013). Additionally, mean temperature is correlated to distance from the edge as well as the size of the gap (Pearson et al., 2002; Saner et al., 2009; Takada et al., 2015). Temperature fluctuations are also dependent on the burial depth of the seed in the soil (Pearson et al., 2002). Finally, the composition of the forest effects both the mean temperature and range of the diurnal fluctuation, for instance, *M. acuminata* seeds would be inhibited from germinating in the temperatures of old growth forests, but may occur in oil palm plantations (Hardwick et al., 2015).

Our results show that, after storage, most seeds from the same inflorescence are non-dormant but some seeds are; germination is increased following a period of stratification or after-ripening. Heterogeneity in dormancy could be part of a bet-hedging strategy whereby seeds have variable levels of dormancy to aid dispersal and maximize seedling establishment (Ng, 1980; Tielborger et al., 2012; Gremer et al., 2016).

We found *M. acuminata* seed coats did not limit imbibition. However, for seeds that have not been dried and frozen, physical dormancy cannot be completely ruled out; especially as drying increases imbibition rates in *Musa* seeds (Puteh et al., 2011). Furthermore, *Musa* seeds clearly invest in physical defences, which may correlate with physical dormancy. In other species, alternating temperatures have also demonstrated removal of physical dormancy (De Souza et al., 2012; Jaganathan et al., 2019), and this in relation to disturbance (Jaganathan, 2018).

Much of the variation in our results was the result of the variation between accessions. This is despite seeds being treated and collected in the same way. Additionally, the overall viability of our accessions was low, and *Musa* seeds have been shown to have variable levels of desiccation sensitivity depending on species and maturity at collection (Kallow et al., 2020). The loss of viability observed here may also be the result of immature seeds not having fully developed desiccation tolerance (Ellis et al., 1991;
Hay and Probert, 1995; Leprince et al., 2017). Seed maturity within and between bunches can be important for traits such as desiccation tolerance and germination potential. Despite our efforts, it is possible that immature seeds were collected during field expeditions and this may help explain why it is then difficult to obtain consistent results for seed germination. When collecting bananas in the wild, it is difficult to access mature seeds as it is rare to find ripe or mature bunches in the forest, presumably because of predation and frugivory.

*M. acuminata* seeds demonstrate sensitivity to alternating temperatures for seed germination suitable to detect gaps in forest canopies following disturbance. These results can directly be applied to the management of banana seeds for conservation, and to more easily access material for phenotyping and breeding.

**Supplementary material.** To view supplementary material for this article, please visit: https://doi.org/10.1017/S0960258520000471.

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**Conflict of interest.** None.

**References**


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