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Posted Date: 25 March 2026

doi: 10.20944/preprints202603.2028.v1

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Article

Karrikinolide Maximises Seed Use Efficiency for Global Ecosystem Restoration and Nature Repair

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+ *In memoriam*

Abstract

Ecological restoration and nature repair combat ecosystem and land degradation, biodiversity loss and climate change. Yet seedling recruitment failure and perilously low plant survival (6-11%, often less) result in mass seed wastage. To understand its ability to improve seed use efficiency for restoration, the powerful germination stimulant karrikinolide or KAR₁ (3-methyl-2*H*-furo [2,3-*c*]pyran-2-one) was evaluated for on-demand seed germination stimulation and plant recruitment in core restoration species. In line with global trends, our research demonstrated that KAR₁ promoted on-demand germination in 82% of species across nine families. Our pioneering work also showed improved outcomes for deteriorating (aged) seeds and higher seedling recruitment, thereby enhancing seed use efficiency. The commercially available stimulant, gibberellic acid (GA₃), provided no assistance beyond seed germination, suggesting KAR₁ cannot be readily substituted. We recommend that KAR₁ has the potential to meaningfully enhance seed use efficiency for nature restoration once challenges like cost and KAR₁ delivery issues are overcome.

Keywords: Karrikinolide; KAR₁; gibberellic acid; GA₃; seed use efficiency; recruitment; germination; restoration; plant growth regulators; PGRs

1. Introduction

The UN Decade on Ecosystem Restoration (2021-2030) aims to restore 1 billion hectares of degraded land into resilient, self-sustaining ecosystems [1]. This global effort is logical for reversing biodiversity loss, improving human well-being and for business acumen: every dollar invested in restoration represents up to US\$30 in economic benefit, and restored environments store carbon and create resilience against climatic extremes and public health threats, such as pandemics [2]. Encouragingly, by restoring just 15% of priority area lands, it is estimated that 63% of extinctions may be avoidable and up to 34% of total atmospheric CO₂ released since the Industrial Revolution could be sequestered (i.e., 335 gigatonnes) [3]. Nonetheless, ecological systems are simultaneously complex and complicated, making them wicked systems in which humans can struggle to accomplish restoration-done-right [4]. For example, native plant reintroductions over large scales (≥ 100 km²) can only be achieved economically by directly sowing seeds into the targeted area for repair [5–7], but at least 89% of these directly sown seeds experience establishment failure, resulting in perilously low plant survival (6-11%, often less) and mass seed wastage [8,9]. This directly impacts the success of large-scale restoration operations, such as those needed to meet UN goals.

One tool to minimise seed wastage is 'on-demand' seed germination stimulation [5]. This enhances seed use efficiency by allowing seeds to germinate upon sowing during the ideal

recruitment window, thereby providing seedlings the best chance to outgrow in-field challenges, such as predation, pathogens, weed competition and seed bank decline [9,10]. Chemical stimulants can directly promote on-demand germination by overcoming the most common dormancy class found in more than 80% of species worldwide, known as physiological dormancy (Figure 1) [9,11]. In this space, the most promising stimulant in literature is 3-methyl-2H-furo[2,3-c]pyran-2-one or karrikinolide (karrikin-1), abbreviated as KAR₁ [12,13]. KAR₁ was first discovered in nature as the most biologically active compound released by burning vegetation (*viz.*, in charred plant materials and smoke), which enhances seed germination upon activation of molecular signalling pathways [12,14]. The subsequent availability of KAR₁ through chemical synthesis [15], combined with the nanomolar quantities required for promoting germination and the effectiveness of KAR₁ across a broad range of species from both fire-prone and non-fire-prone environments, as well as across taxonomic groups and continents, gave the restoration sector hope for cost-effective, on-demand nature repair with substantially enhanced seed use efficiency [13,16]. Yet, although >1200 native plant species across >80 genera are responsive to chemical cues from smoke [17] and more than 20 years have passed since KAR₁ chemical synthesis [15] and patent filing [13], there is still confusion around KAR₁'s true usefulness for restoration.

To provide clarity for end-users in restoration and to guide future research and technology development, we appraise KAR₁ for its capacity to enhance seedling recruitment (germination and establishment), plant survival and seed use efficiency and therefore the success of restoration activities. While we were the first to identify KAR₁ in biochar and pyrolytic extracts and link its presence at biologically active levels to the germination of native seeds and crop abiotic stress tolerance [14], pyrolytic products and smoke water/aerosols are not the focus here. This omission is because such products contain complex organic compounds and toxins (4000-odd in smoke) [16] which are 'hit-and-miss', being permissive or inhibitory depending on the specific chemical cocktail formed and species-specific sensitivity to this chemistry [18].

Instead, here, we screen the effectiveness of pure KAR₁ for promoting on-demand seed germination in 22 species across ten plant families that are critical for rebuilding nature and are therefore categorised as 'core restoration species.' Additionally, many are difficult-to-germinate and/or establish in nursery and/or in field environments (pers. comm. Greening Australia, 2024). For fourteen species, we compare KAR₁ responses to the industry-accessible and registered germination stimulant, gibberellic acid (GA₃). GA₃ costs less than \$600 USD for 10 g (Merck Millipore, USA), and is known to overcome seed dormancy in many native species [19] and sometimes effectively replaces smoke-related cues [20]. By contrast, KAR₁ is presently used only for research, partly because of its prohibitively expensive chemical synthesis, estimated at \$12,500 USD g⁻¹ upon up-scaling [21].

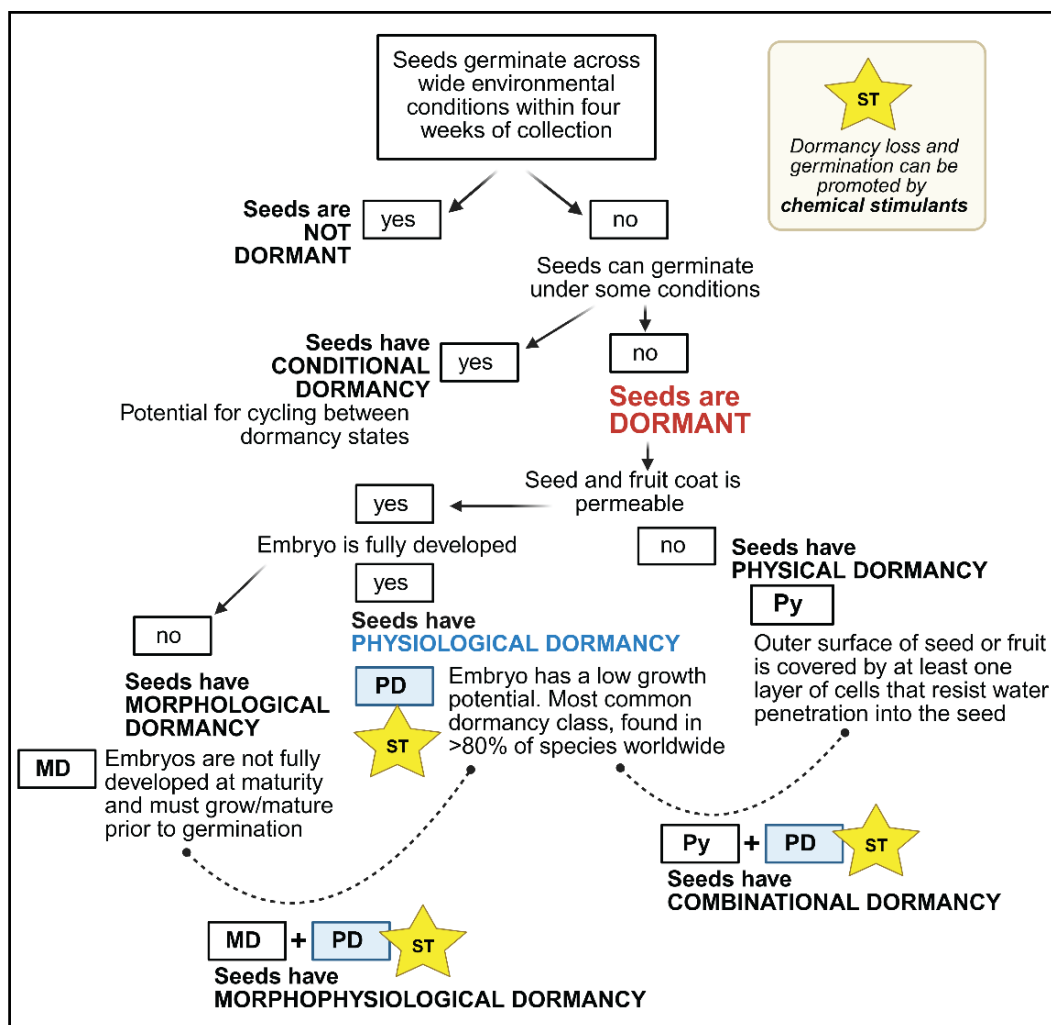


Figure 1. Chemical stimulants, including KAR_1 and GA_3 , are recommended to overcome the most common dormancy class found in >80% of species worldwide, which is physiological dormancy (PD). Seeds with PD have a fully developed embryo with a low growth potential and stimulants can initiate internal molecular signalling, which promotes dormancy loss and germination [9,22]. Stimulants can also assist in germination where seeds have a physical barrier to water penetration plus PD (combinational dormancy, also requires physical dormancy alleviation) or embryos that are not fully developed plus have PD (morpho-physiological dormancy, also requires seeds to mature). The full complexity of dormancy and broader seed treatments is detailed by [9,11] but is beyond our scope here.

Thus, to determine if KAR_1 can be replaced by the vastly cheaper GA_3 , we compare the effects of both stimulants on germination of healthy seeds and, as pioneering research, germination of deteriorating seeds and plant recruitment across critical life phases - specifically, establishment and survival in soil. Indeed, the use of KAR_1 for extending the life of deteriorating seeds is a critical value proposition for restoration practitioners since it is common to store seeds for years before use (pers. comm. Greening Australia, 2025), yet it has never been studied in native species. Similarly, native plant survival after KAR_1 application to seeds has only been monitored in one identified study without control treatments [23], while most studies solely monitor germination in a Petri dish [21]. Yet in crops and *Arabidopsis*, KAR_1 is known to stimulate post-germination seedling growth by accelerating transition to photoautotrophy and enhancing root growth and development, phloem formation, meristem growth and even tolerance to abiotic stress [21,24]. Hence, here we compare the trajectory of seedling establishment and survival in soil after KAR_1 versus GA_3 seed treatments to understand implications for seed use efficiency. Finally, to provide a realistic pathway forward for

restoration practitioners, researchers, and policy makers, we conclude by providing future perspectives for KAR₁, discussing the barriers to KAR₁ commercial applications and upscaling, as well as suggesting future research and technology development to overcome such obstacles.

2. Materials and Methods

2.1. Plant Material and Storage Conditions

Seeds of native species were sourced from Nindethana Seed Services (Albany and Richmond, Australia) in sealed plastic or foil packets and stored at 15 °C until use (and re-dried at 15-17% relative humidity after use). Twenty-two species across ten plant families were screened and Table S1 in Supplementary Material provides collection date, location and previously available information on dormancy and seed pre-treatments for each species.

2.2. Seed Germination and Recruitment

Seed quality and fill for all collections were determined under an Olympus SZ60 Stereomicroscope (Olympus, Australia) fitted with a 1x eyepiece and variable zoom (Figure 2a). Seeds that were damaged or empty were removed from the analysis [19]. As documented in literature, Poaceae seeds were cleaned using a 500 µm-sized mesh to enhance their germination, except for *Austrostipa scabra* and *Chloris truncata*, because manual cleaning does not enhance their germination [25].

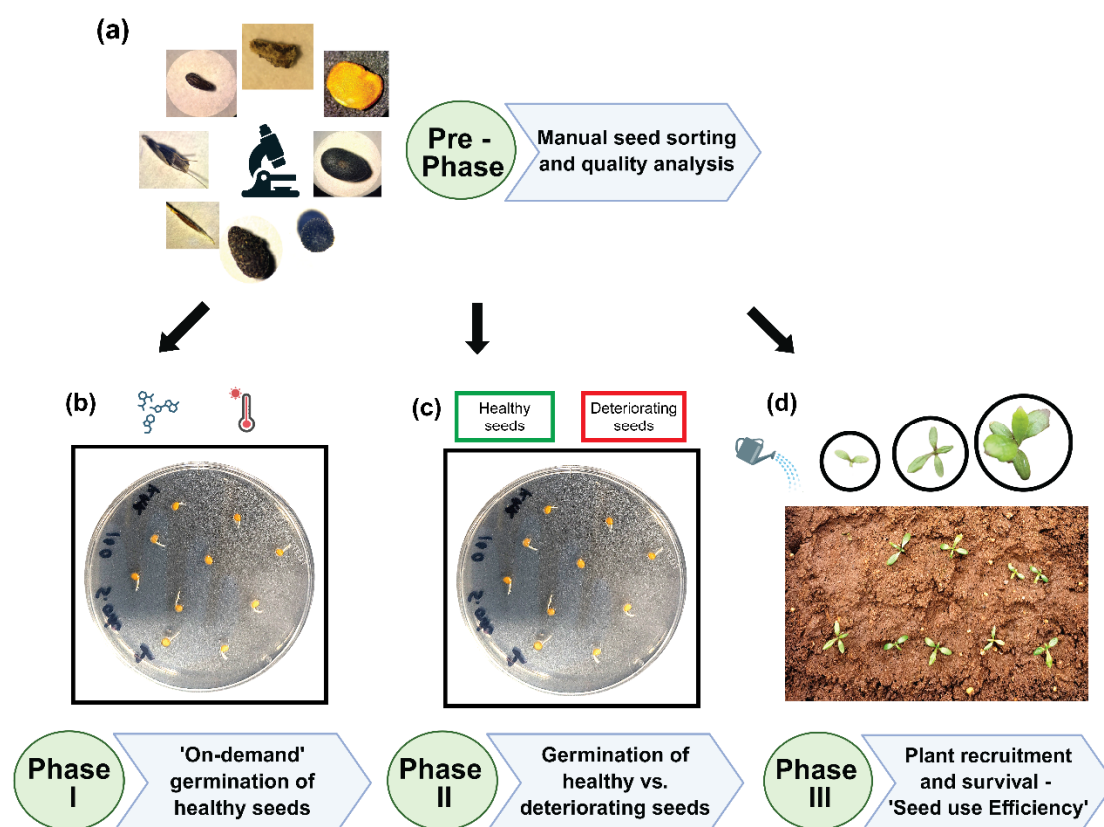


Figure 2. Study phases included: (a) A pre-phase whereby damaged or empty seeds were removed from the analysis to ensure high seed quality; (b) Phase 1 which determined 'on-demand' germination of healthy seeds using bioassays on agar with and without KAR₁ and GA₃; (c) Phase 2 which determined germination of healthy versus deteriorating seeds for four species from two plant families and; (d) Phase 3 which determined plant recruitment (germination and establishment) and long-term survival in soil, thereby allowing calculation of seed use efficiency.

'On-demand' germination of healthy seeds after KAR₁ and GA₃ pre-treatments was determined for 14 species across 10 families in Phase 1 studies (Figure 2b) using bioassays on agar, as this technique has been shown to outperform priming in solution [26]. Bioassays were prepared according to the protocol from [19], using 1% w/v agar (Sigma-Aldrich, USA) with and without stimulants in 90-mm plastic Petri dishes, which were wrapped in aluminium foil for darkness and sealed in plastic zip-lock bags. KAR₁ was synthesised as described in [15], and GA₃ was obtained from Sigma-Aldrich (USA). KAR₁ concentrations tested were 10 and 100 µg L⁻¹ (67 and 670 nM, respectively), but also at up to 1000 µg L⁻¹ (6.7 µM) for selected species, and GA₃ concentrations were 200 and 500 mg L⁻¹ (0.577 mM and 1.444 mM, respectively) [19,27]. Bioassays were incubated in a Conviron growth cabinet (Winnipeg, Canada) at 20/15 °C for winter species and 25/20 °C for summer species (Table S1 in Supplementary Material). The experimental unit was a Petri dish containing 10 seeds (Figure S1a in Supplementary Material) and Petri dishes were maintained in a randomised complete block design using three blocks per treatment (true biological replications). For two *Eucalyptus* species where seed numbers were low, four seeds were placed per Petri dish and five blocks were used. The criterion for seed germination was radicle emergence to >2 mm, followed by normal seedling growth [19], recorded every *c.* 3-4 days.

To determine germination of healthy versus deteriorating seeds for Phase 2 studies (Figure 2c), we used the same protocol as in Phase 1 bioassays. Seeds of four species from the Solanaceae and Poaceae families were tested for their responses to KAR₁ and/or GA₃ in 2021, when seeds were healthy and in 2024, when seeds were deteriorating. Deteriorating seed health was evidenced by fungal contamination on ungerminated seeds [28].

To determine plant recruitment (germination and establishment) and survival in soil for Phase 3 studies (Figure 2d), we first calculated seed germination outcomes, using the same protocol as from Phase 1 with agar that contained optimised stimulant dosages. To determine seedling establishment, germinates were transferred into trays (380 × 140 mm, Bunnings Group Ltd., Victoria, Australia) containing an optimised soil blend with a pH of *c.* 5 (90% sandy loam, 10% AS-4454 certified compost, Centenary Landscaping Supplies, Darra, QLD, Australia; mixed with propagation sand and peat moss, Brunnings Garden Products Pty Ltd., Victoria, Australia; ratio of 60:30:10; Figure S1b in Supplementary Material). Plants were maintained for 8-12 weeks in growth cabinets under a 16/8 h light/dark cycle using 20/15 °C for winter species and 25/20 °C for summer species (Table S1 in Supplementary Material). To monitor long-term survival, plants at the two true leaf stage were transferred into 40 mm-sized forestry tubes (Garden City Plastics, Australia) with one plant per tube (Supplementary Figure 1c). A low-phosphorus native plant liquid fertiliser was used once per week (Australian Native Focus, Growth Technology, Perth, Australia) and irrigation was undertaken every second day until saturation (Figure S1d in Supplementary Material). Once roots had filled the tubes, plants were transferred to 145 mm pots and moved to a temperature-controlled glasshouse with moderate conditions (25 ± 10 °C) with daily automated watering and fertiliser application once per week (Figure S1e in Supplementary Material), where they were maintained for 3-6 months. Growth was monitored using a protocol modified from [19].

2.3. Statistical Analysis and Software

Generalised linear models (GLMs) with binomial error structure and the log link function were used to evaluate the effects of different treatments on seed germination, seedling establishment and plant survival [29]. Tukey's test at the 5% level of significance ($\alpha = 0.05$) was used for multiple mean comparisons unless stated otherwise. All statistical analyses and graphs were created using GraphPad Prism, version 10.2.2 (GraphPad Software, Boston, MA, USA). Figures were designed and assembled using BioRender (Toronto, Canada) and/or Adobe Illustrator 2025. All software licenses were provided by The University of Queensland. Raw data is deposited within UQRDM [2025-RD001812]. Minor adjustments to plant images were made using ImageJ. No generative artificial intelligence has been used in this paper.

3. Results

3.1. Phase 1: KAR₁ as an 'On-Demand' Germination Stimulant Outperforms GA₃

KAR₁ outperformed GA₃ with more germinates and/or faster germination for seven out of fourteen species across six out of nine families tested (Goodeniaceae, Figure 3a; Dilleniaceae, Figure 3b; Asparagaceae, Figure 3d; Asteraceae, Figure 3f; Poaceae, Figure 3h; and two species in Myrtaceae, Figure 3j, k). By contrast, GA₃ outperformed KAR₁ for only one Campanulaceae species (Figure 3e) and one species in the Myrtaceae (Figure 3l) family. Both stimulants were equally effective for Fabaceae (Figure 3c) and another species of Myrtaceae (Figure 3i). GA₃ more commonly reduced germination below the control than KAR₁, specifically at 500 mg L⁻¹ for plant species in the Fabaceae (Figure 3c) and Myrtaceae (Figure 3k) families and at both GA₃ dosages for one Asteraceae species (Figure 3f). Interestingly, both stimulants at one or both dosages suppressed germination for c. 30% of species tested (Asparagaceae, Figure 3d; Poaceae, Figure 3g; Fabaceae, Sapindaceae, Figure S2a, b in Supplementary Material, respectively). For the former two species, physical dormancy is reported (Table S1 in Supplementary Materials), and although beyond our scope here, stimulants should be re-tested after physical dormancy is removed [30].

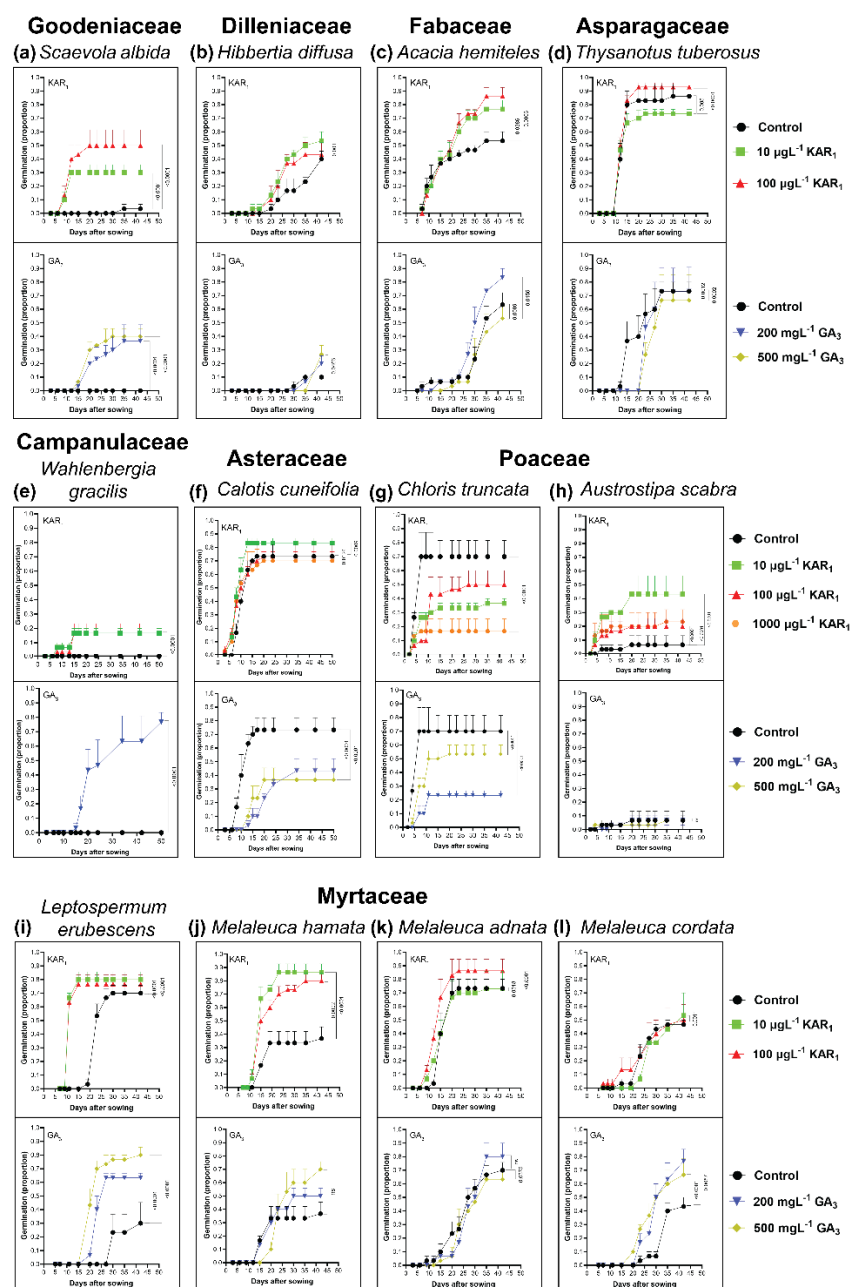


Figure 3. 'On-demand' germination of healthy seeds for core restoration species across eight native plant families in response to KAR₁ and GA₃. Species are from families including (a) Goodeniaceae: *Scaevola albida*; (b) Dilleniaceae: *Hibbertia diffusa*; (c) Fabaceae: *Acacia hemiteles*; (d) Asparagaceae: *Thysanotus tuberosus*; (e) Campanulaceae: *Wahlenbergia gracilis*; (f) Asteraceae: *Calotis cuneifolia*; Poaceae: (g) *Chloris truncata*, (h) *Austrostipa scabra*; and Myrtaceae: (i) *Leptospermum erubescens*, (j) *Melaleuca hamata*, (k) *M. adnata*, (l) *M. cordata*. Plotted values are the mean + SEM of seed germination (proportion, n = 3) recorded on agar containing no stimulants (control) or 10, 100 and/or 1000 µg L⁻¹ KAR₁ and 200 or 500 mg L⁻¹ GA₃. The numbers represent P values after a post-hoc Tukey test (α = 0.05).

3.2. Phase 2: Studies with Healthy Versus Deteriorating Seeds

3.2.1. KAR₁ Outperforms GA₃ for Germination of Deteriorating Seeds

Healthy seed germination was significantly stimulated by both GA₃ and KAR₁ for two Solanaceae species (Figure 4). Specifically for healthy seeds in 2021, GA₃ (200 mg L⁻¹) and KAR₁ (100 µg L⁻¹) germinated 92-94% of *S. orbiculatum* seeds relative to 57% in the control (Figure 4a, 2021) and up to 70% of seeds of *S. prinophyllum* relative to 23% in the control (Figure 4b, 2021).

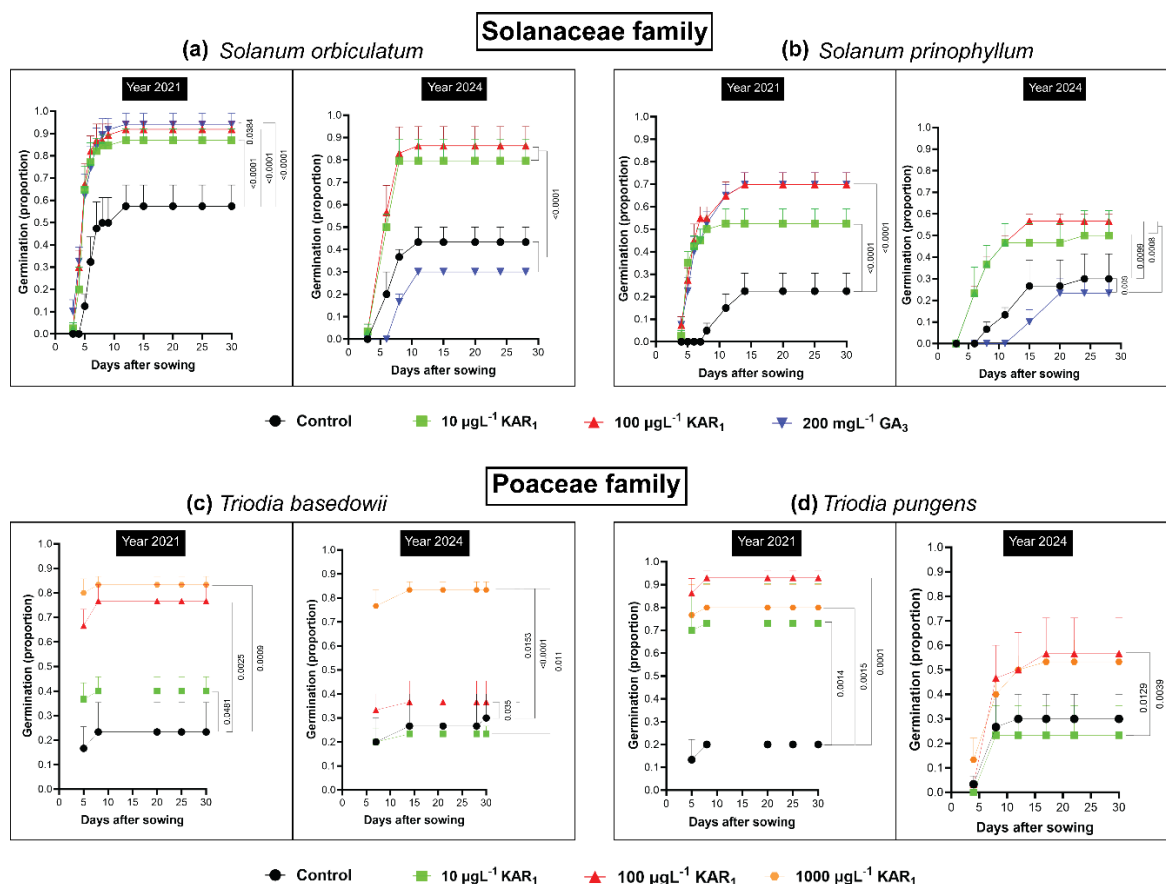


Figure 4. Germination of healthy versus deteriorating seeds sown in 2021 versus 2024, respectively, for core restoration species across two native plant families. Species are from families including, Solanaceae: (a) *Solanum orbiculatum*; (b) *S. prinophyllum*; and Poaceae: (c) *Triodia basedowii*; (d) *T. pungens*. Plotted values are the mean + SEM of seed germination (proportion, n = 3) recorded on agar containing no stimulants (control) or 10, 100 or 1000 µg L⁻¹ KAR₁ and/or 200 mg L⁻¹ GA₃. The numbers represent P values after a post-hoc Tukey test, α = 0.05.

By contrast, when seed quality had deteriorated in 2024, only KAR₁ (at 100 µg L⁻¹) resulted in higher germination for both species, doubling germinates to 86% for *S. orbiculatum* relative to 43% in the control (Figure 4a, year 2024) and increasing germinates to 57% for *S. prinophyllum* relative to 30% in the control (Figure 4b, 2024). Meanwhile, GA₃ reduced germination for aged seeds of both species

in 2024 below the control to 30% and 23%, respectively (Figure 4a, b, 2024). Thus, GA₃ performed poorly as seeds aged, while KAR₁ continued to enhance germination across years, hinting towards its broader application for seeds that have been stored for extended periods. Indeed, seeds are often stored for years before use in restoration (pers. comm. Greening Australia, 2024), hence this is an important value proposition for the sector.

3.2.2. Active KAR₁ Dosage Increases as Seeds Deteriorate

To investigate further, we tested the germination effects of KAR₁ for healthy versus deteriorating seeds for two native grass species (Poaceae) from arid Australia. For healthy seeds of *Triodia basedowii*, KAR₁ improved germination to >75% relative to 23% in the control (Figure 4c, year 2021) at medium and high dosages (100 and 1000 µg L⁻¹). By contrast, for aged seeds in 2024, only a high dosage of KAR₁ improved germination to >75% relative to 30% in the control (Figure 4c, year 2024, KAR₁ at 1000 µg L⁻¹). Similarly, for healthy *T. pungens* seeds in 2021, low, medium and high KAR₁ concentrations improved germination to ≥73% relative to 20% in the control (Figure 4d, year 2021, KAR₁ at 10-1000 µg L⁻¹). Meanwhile, for deteriorating seeds in 2024, only medium and high KAR₁ dosages showed a trend towards improved germination relative to the control (Figure 4d, year 2024, KAR₁ at 100 and 1000 µg L⁻¹) while the low KAR₁ dosage had no effect. These studies thus confirmed that KAR₁ continued to elicit a positive effect on germination for healthy and deteriorating seeds, although in the case of the Poaceae family, the active dosage became higher as the seeds aged.

3.2.3. Active KAR₁ Dosage for Healthy Seeds

As summarised in Table 1, 100 µg L⁻¹ of KAR₁ (670 nM, 100 ppb) resulted in the most germinates and/or fastest germination for species from Goodeniaceae (*S. albida*), Fabaceae (*A. hemiteles*), Myrtaceae (*M. adnata*), Solanaceae (*S. orbiculatum*, *S. prinophyllum*) and Poaceae (*T. pungens*, *T. triandra*, *S. leiocladum*) families and showed a trend for improved germination for three species from Asparagaceae (*T. tuberosus*) and Myrtaceae (*E. melliodora*, *E. tereticornis*) families. Two species responded equally to 10 and 100 µg L⁻¹ of KAR₁ from Campanulaceae (*W. gracilis*) and Myrtaceae (*L. erubescens*), while four species germinated best at 10 µg L⁻¹ of KAR₁ from Dilleniaceae (*H. diffusa*), Asteraceae (*C. cunefolia*), Poaceae (*A. scabra*) and Myrtaceae (*M. hamata*). Germination was reduced at 10 µg L⁻¹ of KAR₁ for one Asparagaceae species (*T. tuberosus*). Although 1000 µg L⁻¹ of KAR₁ was most beneficial for one Poaceae species (*T. basedowii*), this dose reduced germination below the control for two other Poaceae species (*C. truncata*, *T. triandra*).

Table 1. The effects of KAR₁ (K) and GA₃ (G) for on-demand seed germination of twenty-two core restoration species across ten plant families. Data is combined from Figure 2-3 and Figure S3-5 in the Supplementary Material. Taxon-specific information for Figure S2-3 is detailed in the Supplementary Material.

Figure	Family	Species	Stimulant summary	Dosage summary (µg L ⁻¹)	Mean germination (%) and difference from control ^P value (Tukey test, α = 0.05)						
					KAR ₁ (µg L ⁻¹)			GA ₃ (mg L ⁻¹)			
Phase 1. On-demand seed germination, KAR ₁ versus GA ₃ (Figure 3, Figure S2 in Supplementary Materials)					Co				Co		
					ntr	10	100	1000	ntr	200	500
					ol				ol		
3a	Goodeniaceae	<i>Scaevola albida</i>	↑K > ↑G	↑K (100>10), ↑G (500>200)	3.3	30 ^{<0}	50 ^{<0}		0	37 ^{<0}	40 ^{<0.001}
						.0001	.0001			.0001	.001

3b	Dilleniaceae	<i>Hibbertia diffusa</i>	↑K > ↑G	↑K (10), ↑G (500)	40	53 ^{0.001}	43 ^{ns}		10	20 ^{ns}	27 ^{0.04} ₆
3c	Fabaceae	<i>Acacia hemiteles</i>	↑K = ↑G	↑K (100>10) = ↑G (200), ↓GA (500)	53	77 ^{0.04}	86 ^{0.006}		63	83 ^{0.156}	↓ _{53^{0.009}}
3d	Asparagaceae	<i>Thysanotus tuberosus</i>	↑K trend > ↓K, ↓G	↑K trend (100), ↓K (10), ↓G (200, 500)	86	↓ _{73^{0.005}}	93 ^{ns}		73	↓ _{73^{0.012}}	↓ _{67^{0.002}}
3e	Campanulaceae	<i>Wahlenbergia gracilis</i>	↑G >> ↑K	↑G (200), ↑K (100=10)	0	17 ^{<0.0001}	17 ^{<0.0001}	0	-	77 ^{<0.0001}	0
3f	Asteraceae	<i>Calotis cunefolia</i>	↑K >> ↓G	↑K (10), ↓G (200, 500)	73	83 ^{0.0009}	73 ^{<0.0124}	70 ^{ns}	-	↓ _{43^{<0.0001}}	↓ _{37^{<0.001}}
3g	Poaceae	<i>Chloris truncata</i>	↓K, ↓G	↓K (1000), ↓G (200, 500)	70	37 ^{ns}	50 ^{ns}	↓ _{17^{<0.0001}}	70	↓ _{23^{<0.0001}}	↓ _{53^{<0.0001}}
3h	Poaceae	<i>Austrostipa scabra</i>	↑K >> G (ns)	↑K (10>100=1000), G (ns)	7	43 ^{<0.0001}	20 ^{<0.0001}	23 ^{<0.0001}	7	7 ^{ns}	7 ^{ns}
3i	Myrtaceae	<i>Leptospermum erubescens</i>	↑K = ↑G	↑K (10=100) = ↑G (500>200)	70	80 ^{<0.0001}	77 ^{<0.0001}		30	63 ^{<0.0001}	80 ^{<0.001}
3j	Myrtaceae	<i>Melaleuca hamata</i>	↑K > G (ns)	↑K (10>100) = G (ns)	37	86 ^{<0.0001}	80 ^{0.0002}		37	50 ^{ns}	70 ^{ns}
3k	Myrtaceae	<i>Melaleuca adnata</i>	↑K > ↓G	↑K (100>10), ↓G (500)	73	73 ^{0.0318}	86 ^{<0.0001}		70	80 ^{ns}	↓ _{63^{0.008}}
3l	Myrtaceae	<i>Melaleuca cordata</i>	↑G >> K	↑G (200>500) >> K (ns)	47	53 ^{ns}	50 ^{ns}		43	77 ^{0.067}	67 ^{<0.001}
S2a	Fabaceae	<i>Acacia haviandiorum</i>	K(ns), ↓G	↓G (200, 500)	23	23 ^{ns}	20 ^{ns}		23	↓ _{17^{0.008}}	↓ _{7^{<0.001}}
S2b	Sapindaceae	<i>Dodonaea viscosa</i> var. <i>cuneata</i>	↓K, ↓G	↓K (1000) = ↓G (200, 500)	13	3 ^{ns}	10 ^{ns}	↓ _{3^{0.025}}	-	↓ _{3^{0.007}}	↓ _{7^{0.020}}

Phase 2. On-demand seed germination of healthy versus deteriorating seeds (Figure 4)					Co ntr ol	KAR ₁ (µg L ⁻¹)			GA ₃ (mg L ⁻¹)	
						1	10	100	100 0	200
4a	Solanaceae	<i>S. orbiculatum</i> (2021 - healthy)	↑K = ↑G	↑K (100>10), ↑G (200)	57		87 ^{<0} .0001	92 ^{<0} .0001		94 ^{<0} .0001
4a	Solanaceae	<i>S. orbiculatum</i> (2024 - aged)	↑K >> ↓G	↑K (100>10), ↓G (trend)	43		80 ^{<0} .0001	86 ^{<0} .0001		↓ 30 ns
4b	Solanaceae	<i>S. prinophyllum</i> (2021 - healthy)	↑K = ↑G	↑K (100>10), ↑G (200)	23		53 ^{<0} .0001	70 ^{<0} .0001		70 ^{<0} .0001
4b	Solanaceae	<i>S. prinophyllum</i> (2024 - aged)	↑K >> ↓G	↑K (100>10), ↓G (200)	30		50 ⁰ .0099	57 ^{0.0} .008		↓ 23 ^{0.0} .091
4c	Poaceae	<i>Triodia basedowii</i> (2021 - healthy)	↑K	↑K (1000>100>10)	23	23 ns	40 ⁰ .0481	77 ^{0.0} .025	83 ^{0.0} .009	
4c	Poaceae	<i>T. basedowii</i> (2024 - aged)	↑K	↑K (1000>>100)	30		23 ns	37 ^{0.0} .350	83 ^{<0} .0001	
4d	Poaceae	<i>Triodia pungens</i> (2021 - healthy)	↑K	↑K (100>1000>1>10)	20	77 ⁰ .0031	73 ⁰ .0014	93 ^{0.0} .001	80 ^{0.0} .015	
4d	Poaceae	<i>T. pungens</i>	↑K trend	Trend: ↑K (100=1000), ↓K (10)	30		23 ns	57 ns	53 ns	

(2024 - aged)					KAR ₁ (µg L ⁻¹)			GA ₃ (mg L ⁻¹)	
On-demand seed germination – KAR ₁ only (Figure S3 and S4 in Supplementary Materials)					Concentration				Concentration
					10	100	1000	200	500
					ol				ol
S3c	Poaceae	<i>Themeda triandra</i>	↑K	↑ K (100)	27	27 ns	30 ^{0.0189} ns	10 ns	
S3d	Poaceae	<i>Sorghum leiocladum</i>	↑K*	↑ K (100) (100 ^{Tukey test, α=0.1})	80	83 ns	86 ^{0.0614}		
S3a	Myrtaceae	<i>Eucalyptus melliodora</i>	↑K trend	↑K trend (100)	55	60 ns	70 ns		
S3b	Myrtaceae	<i>Eucalyptus tereticornis</i>	↑K trend	↑K trend (100)	69	67 ns	75 ns		
S4a	Myrtaceae	<i>Leptospermum erubescens</i> #2	↑K	2 ND PROVENA NCE, ↑K (10=100)	60	63 ^{<0.0001}	70 ^{<0.0001}		
S4b	Myrtaceae	<i>Melaleuca hamata</i> #2	↑K	2 ND PROVENA NCE, ↑K (100)	60	67 ns	87 ^{0.0491}		

Abbreviations and symbols: K, KAR₁; G, GA₃; ↑, increased germination; ↓, decreased germination; trend, a distinctive but nonsignificant trend; *, significant at $\alpha = 0.1$; #2, second provenance for a given species. Note: Certain species showed high germination variation in the control without stimulants, possibly because these species simultaneously contained seeds that were dormant, non-dormant and conditionally dormant, as is common for some native species (Baskin & Baskin, 2004).

It is noteworthy that KAR₁ effectiveness may show similar outcomes across provenances within species, after similarly positive KAR₁ outcomes were observed when two Myrtaceae species were retested from a second provenance (relative to Phase 1 species *L. erubescens* and *M. hamata*, respectively, Figure S4a, b in Supplementary Materials versus Figures 3h, i).

3.3. Phase 3: KAR₁ Improves Plant Recruitment in Soil in Comparison to GA₃

For native species, large seedling losses during recruitment have been observed, even with KAR₁ pre-treatments [22,23]. Here, KAR₁ drastically outperformed GA₃ for seedling establishment and plant survival (Figure 5). Investigations used four restoration species that had significant germination responses to stimulants in Phase 1 studies and stimulant dosages used were those that elicited the

best germination outcomes (Figure 3, Table 1). Seed use efficiency (SUE) is here defined as the number of plants that emerged and survived out of the total number of seeds sown.

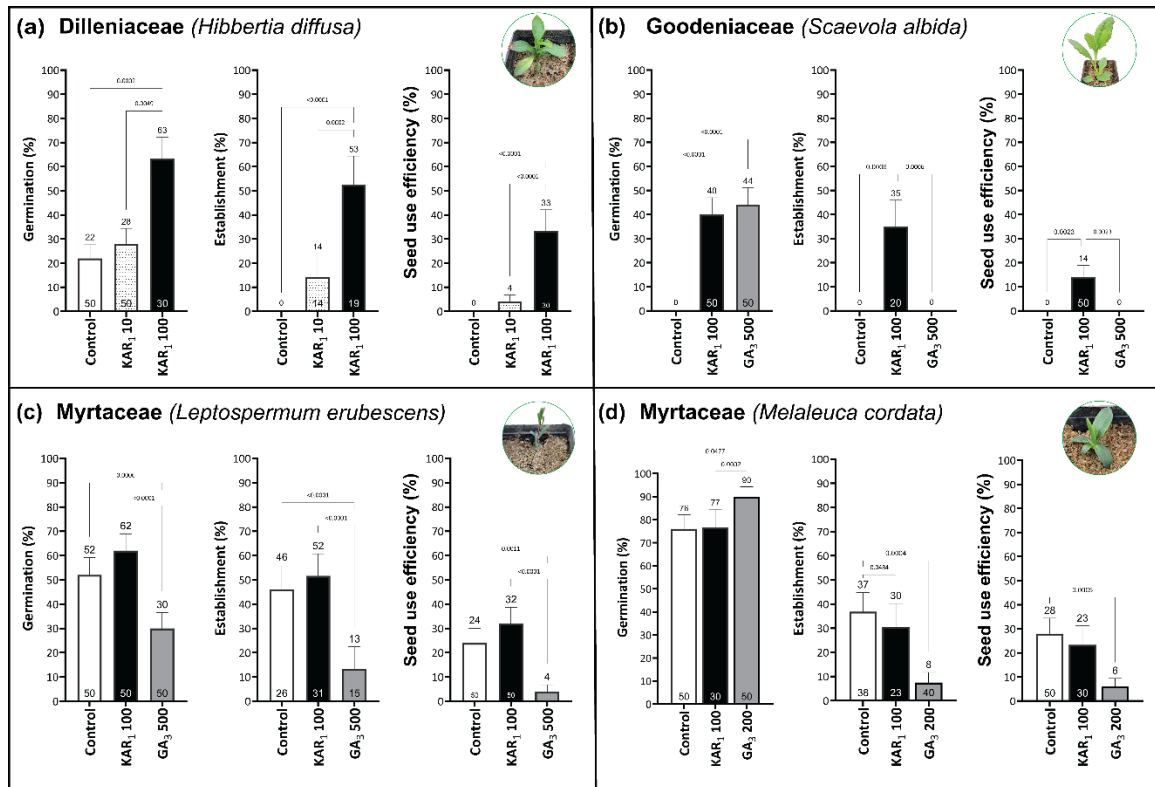


Figure 5. Plant recruitment and seed use efficiency for four core restoration species from three native plant families. Species were selected from three families, including (a) Dilleniaceae: *H. diffusa*; (b) Goodeniaceae: *S. albida*; (c) Myrtaceae: *L. erubescens*; (d) *M. cordata*, and treatments included no stimulants (control) or 10 or 100 $\mu\text{g L}^{-1}$ KAR₁ or 200 or 500 mg L^{-1} GA₃. Bars and numbers above each bar are the mean + SEM of percentage seed germination, seedling establishment and seed use efficiency. The numbers above the lines represent mean separation, whereby *P* values indicate significant differences after a post-hoc Tukey test, $\alpha = 0.05$. Germination is defined as the number of germinates per agar plate out of the total number of seeds sown; establishment is the number of seedlings that grew in soil out of the total number of germinates transferred from agar; seed use efficiency is the number of plants that survived for 3-6 months out of the total number of seeds sown on agar.

Especially the medium KAR₁ dose (100 $\mu\text{g L}^{-1}$) significantly enhanced germination (40-76%), establishment (30-53%) and SUE (14-33%) for three out of four core restoration species, relative to control treatments (germination, 0-76%; establishment, 0-46%; seed use efficiency, 0-28%). By contrast, GA₃ pre-treatments resulted in high germination (30-89%) but drastically hindered establishment (0-30%) and SUE (0-6%) for all species tested, with outcomes even below the untreated control (Figure 5).

For the first species tested from Dilleniaceae (Figure 5a, *H. diffusa*), KAR₁ resulted in the highest germination (63%) relative to the control (22% without stimulants), which translated to vastly improved establishment (53% KAR₁, 0% control) and SUE outcomes (33% KAR₁, 0% control). *H. diffusa* was not tested with GA₃ due to limited seed availability and a vastly stronger KAR₁ response in Phase 1 studies (Figure 3b).

S. albida from the Goodeniaceae family (Figure 5b) had high germination with both stimulants (40-44%) relative to the control (0%). KAR₁ pre-treatments then resulted in 35% of seedlings establishing in the soil, and SUE was 14%. By contrast, no seedlings established after GA₃ pre-treatment, representing 0% SUE (Figure 5b).

For *L. erubescens* from the Myrtaceae family (Figure 5c), recruitment after KAR₁ pre-treatment was high (62% germination, 52% establishment), which was reflected in a high SUE of 32%. This was greater than the control (24%). Meanwhile, recruitment with GA₃ was low (30% germination, 13% establishment), which was reflected in a very low SUE (4%, Figure 5c). For *M. cordata*, also a Myrtaceae (Figure 5d), KAR₁ did not enhance or harm germination, with c. 76% germination with KAR₁ pre-treatment and in the control. This concurs with Phase 1 studies where no significant response to KAR₁ was observed (Figure 3l). KAR₁ then slightly reduced establishment relative to the control (non-significant, 30% KAR₁, 37% control, Figure 5d), and this was reflected as a slightly lower SUE (non-significant, 23% KAR₁, 28% control). Meanwhile, *M. cordata* had the highest germination with GA₃ treatment (90%), but this still translated to the lowest establishment (8%) and a poor SUE (6%), well below the control (28%).

4. Discussion

4.1. KAR₁ as a Stimulant to Enhance Seed Germination and Seedling Establishment

Twenty years after the promise of ‘on-demand’ native plant restoration with KAR₁ [13,17], we show that KAR₁ has the capacity to form a thread within a tapestry of solutions to assist restoration practitioners in achieving successful plant establishment. Across our studies, KAR₁ stimulated germination for 82% of core restoration species tested, including trees, shrubs, herbs and grasses, whereby germination of eighteen out of twenty-two species across nine out of ten families was enhanced by KAR₁. Thus, we concur that as a stimulant that supports ‘on-demand’ germination, KAR₁ has real potential benefits for restoration by enabling a new seedling to emerge in the optimal recruitment window and therefore outgrow challenges that hinder in-field recruitment. This is particularly significant because many in-field challenges will be exacerbated by climate change, such as lower and more variable rainfall, pathogens, seed bank deterioration and competition from weeds [9,19,31,32]. And species from across continents are responsive to KAR₁, such as from North and South America, South Africa, Australia and Europe/Asia [12,33–35], hence this stimulant promises global benefits. We also show that only 10-100 ppb of KAR₁ (10-100 µg L⁻¹, 67-670 nM) was most commonly beneficial, which concurs with the global trend of 10-150 ppb of KAR₁ (10-150 µg L⁻¹, 67 nM - 1 µM) being most commonly bioactive for native seed germination [12,14,36].

4.2. KAR₁ Cannot Be Readily Substituted by Stimulants Such as Gibberellic Acid (GA₃)

We also demonstrate here that KAR₁ cannot be readily substituted, outperforming the relatively cheap commercially available germination stimulant ‘alternative’, gibberellic acid (GA₃). Firstly, in germination bioassays with healthy seeds, KAR₁ outperformed GA₃ with more germinates and/or faster germination for seven out of fourteen species across six out of nine families tested, while GA₃ outperformed KAR₁ for only two species. GA₃ also more commonly reduced germination below the control than KAR₁.

Secondly, ours is the first study to show that KAR₁ outperforms GA₃ for deteriorating native seeds. Here, KAR₁ enhanced deteriorating seed germination for four species, while GA₃ drastically reduced germination, even below the control, once seeds had aged. Oxidative damage increases as seeds age [28] and in crops KAR₁ has suppressed oxidative stress through reduced lipid peroxidation and increased antioxidant activity [21,37,38], improving aged and immature crop seed quality and resulting in higher seedling vigour [39,40]. Thus, we surmise that KAR₁ may assist aged native seed germination via reduced oxidative damage. Given that aged seed may be the only seed available for restoration (pers. comm. Greening Australia, 2024), our result has important implications for seed use and treatments by restoration practitioners to optimise on-ground outcomes.

Thirdly, plant survival studies here determined that KAR₁ outperformed GA₃ for plant recruitment, whereby KAR₁ improved all life phases (germination, establishment, survival) above the control for three out of four core restoration species, while plant survival after GA₃ treatment was far below the control. These results concur with Arabidopsis and crop studies, where karrikins

concomitantly promoted seed germination and healthy seedling growth [41] while gibberellins caused seedling abnormalities [42] and poor crop growth [43]. From previous literature, KAR₁ and GA₃ are both recommended as chemical stimulants to overcome physiological dormancy across species. Our work here suggests that GA₃ is not a viable alternative to KAR₁ for many native species.

It is notable that agar screening with KAR₁ gave a good indication about long-term seed use efficiency, whereby species that germinated well in response to KAR₁ on agar also showed improved recruitment and seed use efficiency outcomes than untreated seeds. Thus, we suggest the use of agar screening to obtain optimal KAR₁ dosages prior to large-scale application for restoration. By contrast, positive results in agar screening with GA₃ did not translate to improved soil establishment or seed use efficiency outcomes.

We note that our study was for Australian species and GA₃ may be more commonly potent for species from other regions, such as North America, Europe or Asia [44–46]. Also, broader stimulants should be tested in future studies, such as broader cues from fire that have assisted germination when combined with KAR₁. Examples are cyanohydrin analogue, mandelonitrile [33,47], and syringaldehyde produced from burning lignin [47,48].

To date, there is very little understanding of long-term recruitment outcomes after KAR₁ treatment of native seeds, and our results here are only the second study to monitor native plant survival after KAR₁ treatment. Indeed, only one study has monitored plant emergence and survival after KAR₁ treatment for green roof applications, but without controls [23]. Three different studies have monitored emergence but not survival [12,22,49] and another monitored germinant density from a seed bank [50]. Most of the remaining studies have monitored germination only, usually in a Petri dish [21]. Clearly, more research is warranted in this space.

4.3. Progressing KAR₁ to a Commercial Product for Large-Scale Restoration

Before KAR₁ can progress to a product for large-scale restoration, there are several obstacles to overcome. Firstly, KAR₁ is expensive, estimated to cost US\$12,500 g⁻¹ upon upscale (currently >US\$600 for 5 mg) [21]. Secondly, there is no known up-scalable technology in the literature to reliably deliver KAR₁ to native seeds.

Optimised priming methods with KAR₁ show promise to improve recruitment for species with simple pre-treatment needs, including species that are ‘inherently’ KAR₁ responsive, whereby KAR₁ germinates freshly collected seeds, or have an ‘inducible’ response that can be overcome with simple pre-treatments, such as extended storage or outer appendage removal [51]. Currently, priming seeds on agar with KAR₁, as used here, is the best delivery method in the literature, with outcomes surpassing KAR₁ priming in solution, but more work is needed to improve such techniques. Although seed coating promises easier seed handling, at present coating pre-primed seeds with standard seed coating materials tends to reduce germination and emergence [22,26,49], highlighting the need for further research.

KAR₁ delivery becomes substantially more complicated for species that have an ‘inducible’ KAR₁ response [51] that requires unrealistic pre-treatments, such as burial in soil [52] and/or specific environmental cues such as warm/humid conditions to mimic a dry season (termed dry after-ripening) [53] or cool/hot conditions to mimic a winter or summer season (termed cold and warm stratification, respectively) [9]. For these species, we propose that the next frontier is to progress commercial development of KAR₁ for restoration involving technologies that maximise recruitment by delivering KAR₁ to seeds in-field. Recruitment outcomes would be especially magnified if the technology allowed species to experience seasonal cues needed to induce KAR₁ responsiveness in-field (e.g., dry, summer, winter) followed by KAR₁ release in the optimal recruitment window. The co-delivery of broader actives, such as other stimulants, microbes, pesticides and/or nutrients, could further magnify recruitment success. Such technologies would ideally boost the number of surviving plants per seed input, thus also enhancing seed use efficiency and enabling ecosystem restoration and nature repair on a large scale.

4.4. Integrating KAR₁ into Holistic and Multi-Disciplinary Restoration Programs

We acknowledge that many factors must align for effective ecological restoration and nature repair, as illustrated in Figure 6. This includes, for example, optimised seed collection timing, correct seed handling and storage, well-timed in-field seed sowing during optimal recruitment windows and prioritising areas with low/declining ecological stability [5,54,55]. In addition, it is today recognised that the contribution of Indigenous Peoples who occupy lands to be restored forms the cornerstone of international conservation policy and is critical in delivering on international goals [56]. Overall, to maximise the number of surviving plants per seed input we suggest that a combination of best-practice seed handling and timing of sowing, development of new on-demand recruitment technologies and a cross-disciplinary approach, such as involving Traditional Owner knowledge, restoration practitioner know-how, scientific rigour and innovation, will be needed to solve the ‘wicked’ problem of achieving high-quality ecological restoration and nature repair at scale.

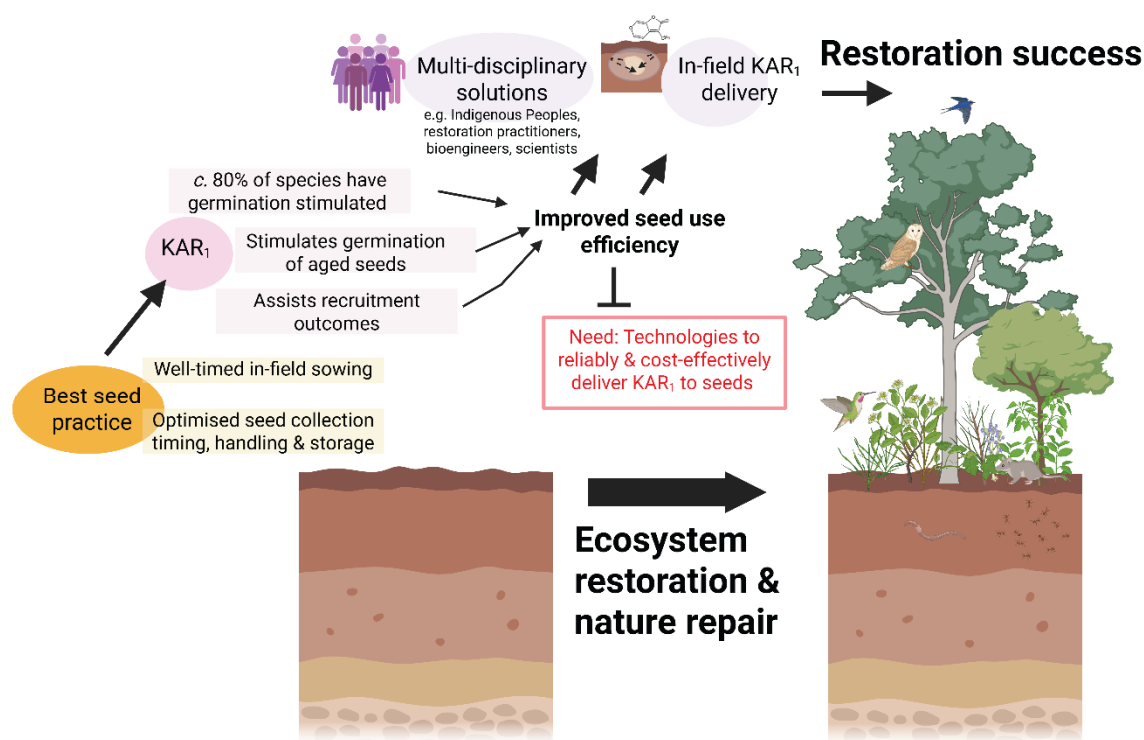


Figure 6. The role of KAR₁ in enhancing seed use efficiency for ecosystem restoration and nature repair.

5. Conclusions

Today we live in an unprecedented ecological overshoot that has manifested as rampant ecosystem and land degradation. Optimistically, by restoring just 15% of priority area lands we may reduce species extinctions, sequester substantial carbon and improve global well-being, however, this requires restoration over large scales that is complex and complicated. Here, we demonstrate that the germination stimulant KAR₁ has the capacity to be a part of the solution through its ability to overcome physiological dormancy commonly found in native species across continents, promoting seed germination in both healthy and deteriorating seeds. Additionally, KAR₁ improved plant recruitment and survival, therefore directly enhancing seed use efficiency. By contrast, the commercially available stimulant GA₃ did not assist seed germination or recruitment, resulting in poor seed use efficiency and failure in comparison to KAR₁. Nonetheless, an up-scalable technology that can reliably and cost-effectively deliver KAR₁ to maximise the number of surviving plants per seed input is needed. The next frontier to progress commercial development of KAR₁ for large-scale

restoration is an innovative technology that delivers KAR₁ in-field, especially if there is a potential to maximise recruitment outcomes. Such innovation, combined with optimised seed handling and cross-disciplinary partnerships, such as with Indigenous Peoples, restoration experts, scientists and bioengineers, sets the landscape for the restoration and repair needed to reinstate the health of our planet and for human prosperity.

Supplementary Materials: The following supporting information can be downloaded at the website of this paper posted on Preprints.org, Figure S1: Plant recruitment and seed use efficiency protocol. Figure S2: ‘On-demand’ germination of healthy seeds for core restoration species across two native plant families in response to KAR₁ and GA₃. Figure S3: ‘On-demand’ germination of healthy seeds for core restoration species across two native plant families in response to KAR₁. Figure S4: ‘On-demand’ germination of healthy seeds for core restoration species from a second provenance from the Myrtaceae family. Table S1: Information on plant species used in the research.

Author Contributions: Conceptualisation, A.B. and J.K.; methodology, A.B., N.B.G., G.M. and K.T.; validation and data curation, A.B., N.B.G., and J.K.; visualisation, A.B., N.B.G. and J.K.; investigation, A.B. and J.K.; formal analysis, A.B., N.B.G. and J.K.; writing – original draft preparation, A.B. and J.K.; writing–review and editing, M.P. and J.K.; resources, J.K., M.P. and C.S.; supervision, J.K. and M.P.; funding acquisition, J.K. and M.P.. All authors have read and agreed to the published version of the manuscript.

Funding: This research was funded by The Ian Potter Foundation as part of a grant to Greening Australia entitled ‘Research and development to build best practice into native seed production in a changing world.’.

Data Availability Statement: The data presented in this study are available on request from the corresponding author. The data is stored within the University of Queensland Research Data Manager (UQRDM) with record identifier number 2025-RD001812 titled ‘Manuscript 2025: Karrikinolide maximises seed use efficiency for global ecosystem restoration and nature repair.’.

Acknowledgments: The authors acknowledge that this research utilises Traditional Indigenous Knowledge, which is wholly owned and shared by Bulugudu Limited (formerly known as Dugalunji Aboriginal Corporation) on behalf of the Indjalandji-Dhidhanu people. Bulugudu Limited has also provided direct financial, equipment, and in-kind support to this project. We acknowledge Associate Professor Gavin Flematti for manufacturing the KAR₁ utilised in this study.

Conflicts of Interest: The authors declare no conflicts of interest.

Abbreviations

The following abbreviations are used in this manuscript:

GA ₃	Gibberellic acid
KAR ₁	Karrikin-1, karrikinolide
PD	Physiological dormancy
SEM	Standard error of the mean
SUE	Seed use efficiency
UN	United Nations

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