






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The effectiveness of Virkon® S disinfectant against an invasive insect and implications for Antarctic biosecurity practices

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Abstract: The flightless midge *Eretmoptera murphyi* is thought to be continuing its invasion of Signy Island via the treads of personnel boots. Current boot-wash biosecurity protocols in the Antarctic region rely on microbial biocides, primarily Virkon® S. As pesticides have limited approval for use in the Antarctic Treaty area, we investigated the efficacy of Virkon® S in controlling the spread of *E. murphyi* using boot-wash simulations and maximum threshold exposures. We found that *E. murphyi* tolerates over 8 h of submergence in 1% Virkon® S. Higher concentrations increased effectiveness, but larvae still exhibited > 50% survival after 5 h in 10% Virkon® S. Salt and hot water treatments (without Virkon® S) were explored as possible alternatives. Salt water proved ineffective, with mortality only in first-instar larvae across multi-day exposures. Larvae experienced 100% mortality when exposed for 10 s to 50°C water, but they showed complete survival at 45°C. Given that current boot-wash protocols alone are an ineffective control of this invasive insect, we advocate hot water (> 50°C) to remove soil, followed by Virkon® S as a microbial biocide on 'clean' boots. Implications for the spread of invasive invertebrates as a result of increased human activity in the Antarctic region are discussed.

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Key words: biosecurity, Chironomidae, invertebrate control, Signy Island, species management

Introduction

Throughout history, humans have acted as agents of change in ecological systems through the deliberate or unintentional introduction of species to various areas. Antarctica's geographical isolation and challenging environmental conditions have, to date, acted as barriers to non-native species dispersal and establishment, thereby minimizing non-native species impacts on the continent itself (Frenot *et al.* 2005, Hughes *et al.* 2015). In 1959, the Antarctic Treaty was signed, coming into force in 1961 and establishing the Antarctic Treaty area as all land, ice shelves and surrounding ocean south of 60°S latitude. From its inception, the Antarctic Treaty placed a high priority on the preservation of Antarctic ecosystems, although this has been achieved by different mechanisms over time, currently by the Protocol on Environmental Protection to the Antarctic Treaty (e.g. the Committee for Environmental Protection Non-native Species Manual; CEP 2016). The remote,

lower-latitude sub-Antarctic islands are closely linked to the Antarctic Treaty area in biological terms, and similarly are of high conservation value, but they are instead regulated under national sovereignty. In recent decades, increasing levels of human activity are progressively breaking down the geographical barriers between Antarctica and the sub-Antarctic region, as well as the rest of the world, thereby increasing the risk of species introductions.

To date, most non-native species occurrences in the Antarctic and sub-Antarctic regions have been the result of historical intentional introductions, but with human activity in the region rapidly rising, the risk of unintentional introductions is becoming an increasing threat to Antarctic ecosystems (Frenot *et al.* 2005, Hughes *et al.* 2015). Human activity has already led to > 200 species of non-native animals and plants successfully establishing in the Antarctic and sub-Antarctic regions, the majority of these being in the sub-Antarctic, but with increasing numbers recorded

111 from the maritime Antarctic (Frenot *et al.* 2005, Hughes
112 *et al.* 2015). These include introductions of Acari,
113 Collembola, Diptera, Coleoptera and Araneae (Pugh,
114 1994, 2004, Ernsting *et al.* 1995, Greenslade & Convey
115 2012). Furthermore, the transfer of pathogens may risk
116 disease in local wildlife populations that may be
117 'immunologically naïve' due to evolution in microbial
118 isolation (Grimaldi *et al.* 2014).

119 Through increased liquid water availability and extent
120 of ice-free habitat, reduced numbers of extreme cold
121 events and extending growing seasons, areas of
122 Antarctica previously unsuitable for colonization are
123 becoming available to both native and non-native species
124 alike (Lee *et al.* 2017). Species introductions can have
125 significant impacts within the simple terrestrial
126 ecosystems of the Antarctic regions. For example, the
127 introduction of a single detritivore to the maritime
128 Antarctic, the midge *Eretmoptera murphyi* (Schaeffer,
129 1914), has been found to increase litter turnover within
130 the local environment where it is established by almost
131 an order of magnitude (Hughes *et al.* 2013). In the
132 sub-Antarctic, a new non-native predatory ground beetle
133 has led to significant declines in native terrestrial
134 invertebrate species (Lebouvier *et al.* 2012). All Parties
135 to the Antarctic Treaty are therefore responsible for
136 developing and enacting measures to prevent or
137 minimize the introduction of non-native species, to
138 control and, if feasible, to eradicate any that have
139 established (Hughes & Pertierra 2016). Available
140 practical response measures are limited, however, by the
141 requirement to keep collateral damage to native habitats
142 and species to a minimum and by associated costs and
143 practicability, as well as by sometimes contradictory
144 existing legislation. For instance, Article 7 of Annex III
145 Waste Disposal and Management bans the use of
146 pesticides within Antarctica, unless under certain
147 necessary circumstances (Hughes *et al.* 2015). Thus, the
148 traditional and most widely used insecticides applied
149 elsewhere (pyrethroids, neonicotinoids and insect growth
150 regulators) may not be options for use in Antarctica.
151 Disinfectants, in contrast, are permitted and are
152 routinely deployed to destroy microbial pathogens and
153 to prevent their spread (Curry *et al.* 2002).

154 The Virkon® S range of disinfectants is currently
155 recommended by the Council of Managers of National
156 Antarctic Programs (COMNAP) and the International
157 Association of Antarctica Tour Operators (IAATO) as
158 an approved biocide (IAATO 2018, COMNAP 2019).
159 These products are also marketed in the UK as
160 Department for Environment, Food and Rural
161 Affairs-approved virucides for farms ([http://disinfectants.
162 defra.gov.uk/DisinfectantsExternal/Default.aspx?Module=
163 ApprovalsList_SI](http://disinfectants.defra.gov.uk/DisinfectantsExternal/Default.aspx?Module=ApprovalsList_SI)), and they claim effectiveness through
164 oxidation against bacteria, viruses and certain strains of
165 fungi at temperatures as low as 4°C (Hernández *et al.*

2000). Virkon® S powder is easy to transport and has
low dermal toxicity, does not give off toxic vapour and,
should it end up in an aqueous environment, will
decompose over time into a harmless mixture of
non-toxic salts (Curry *et al.* 2005, see also manufacturer
declaration [https://syndel.com/wp-content/uploads/
2019/01/Information-Virkon-Aquatic-degradability-in-
the-environment.pdf](https://syndel.com/wp-content/uploads/2019/01/Information-Virkon-Aquatic-degradability-in-the-environment.pdf)). The efficacy and low-toxicity of
Virkon® S has led to its application in Antarctica, where
it has proven effective at preventing the spread of
microbial pathogens under ambient conditions when
used to wash equipment or footwear (Curry *et al.* 2005).

The convenience of Virkon® S products has prompted
toxicity testing against higher-order organisms beyond
its intended use against microbial pathogens, in
particular against invasive marine invertebrate species
within aquatic environments that are more vulnerable to
the off-target effects of harsher chemicals (Stockton-Fiti
& Moffitt 2017). Tests on the New Zealand mud snail,
Potamopyrgus antipodarum, found that 20 min exposure
to 2% Virkon® S solution resulted in 100% mortality at
15°C and 22°C, but that a 1% solution only achieved
total mortality at the lower temperature (Stockton-Fiti &
Moffitt 2017). In the same study, 2% Virkon® S solution
was highly effective against quagga mussels, *Dreissena
rostriformis bugensis*. An invasive tunicate that affects
mussel farming in Canada, *Ciona intestinalis*, has also
been found to be vulnerable to Virkon® S at 1%
concentration (Paetzold & Davidson 2011), whilst the
faucet snail, *Bithynia tentaculata*, proved to be resistant
to dilutions of 1% and 2% at 20–23°C over 1–24 h
(Mitchell & Cole 2008). Its efficacy against insects
remains largely untested, although soaking eggs of the
yellow mealworm, *Tenebrio molitor*, in 1% Virkon® S
for 10 min did not prevent hatching (Li *et al.* 2016), and
mixing it with certain insecticides reduced its efficacy
against the house fly, *Musca domestica* (Watson *et al.*
2008).

The flightless chironomid midge *E. murphyi* is endemic
to the sub-Antarctic island of South Georgia (54°S, 36°W)
(Fig. 1), but it was discovered in 1980 in the maritime
Antarctic on Signy Island (South Orkney Islands, 60°S,
45°W) (Fig. 1) at the site of a previous plant
transplantation experiment (Burn, 1982). Originally
reported to be restricted to a 1 m² introduction site, the
midge has since colonized an area of ~85 000 m² and
can be found along footpaths regularly used by staff and
visitors at the research station. It is now on the verge of
entering into new valley systems (Bartlett *et al.* 2020).

At present, anthropogenic transfer of *E. murphyi* is the
greatest known introduction risk in Antarctica. In 2005,
a British Antarctic Survey (BAS) vessel carried
construction vehicles contaminated with soil containing
various invertebrate species, including *E. murphyi*, from
South Georgia to Rothera Research Station on Adelaide

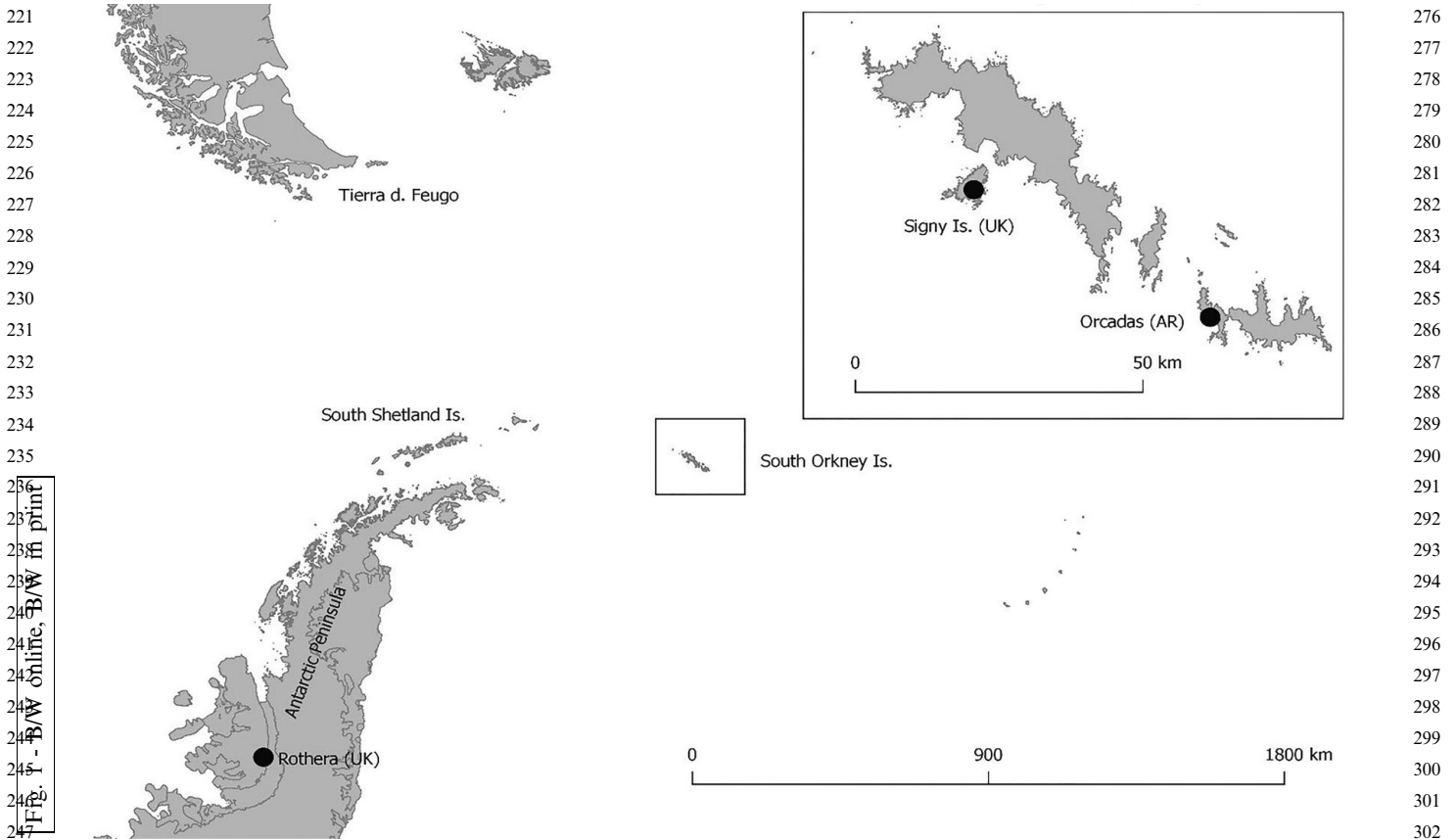


Fig. 1. Location of the South Orkney Islands and Signy Island in the Southern Ocean. Created using *Arc-Map*® 10.4.1 software by Esri. Copyright © Esri.

Island, off the Antarctic Peninsula (68°S), where they were alive when discovered after arrival (Hughes *et al.* 2010). In this instance, no establishment has been detected, probably due to a lack of suitable habitat immediately adjacent to the offloading site, but many suitable locations across the maritime Antarctic are at risk, with the South Shetland Islands being a particularly suitable candidate region and a major logistical hub for the northern Antarctic Peninsula (Perterra *et al.* 2019). Current biosecurity measures employed by BAS encompass the whole supply chain and include cleaning of containers and cargo, where pyrethrum-based insecticides may be used to fumigate shipping containers prior to transportation to Antarctica. Relevant to Signy Island and *E. murphyi*, BAS biosecurity regulations require the cleaning of soil from equipment, boots and clothing, and the use of Virkon® S products at a 1% dilution in boot-wash baths prior to entry and exit from the island (BAS 2019). However, Virkon® S is primarily an antimicrobial agent, and even then its effectiveness is limited without physical removal of any soils/organic loads from contaminated surfaces (Guan *et al.* 2013). The efficacy of Virkon® S to potentially control the spread of any Antarctic invertebrate remains untested.

Against this background, this study investigates whether current Virkon® S boot-wash protocols are effective biosecurity measures against the midge. We also examine *E. murphyi*'s tolerance to seawater and hot water immersion as possible alternatives to chemical control.

Materials and methods

Sample collection

Eretmoptera murphyi larvae were collected in soil on Signy Island (Fig. 1) close to the BAS's Signy Research Station during the 2016–17 summer. Samples were maintained on soil substrate from the site of collection, which is both the species' natural habitat on the island and source of food. Samples were returned to the UK by ship (4°C, constant darkness for 10 weeks) and then maintained under the same control conditions at the University of Birmingham. Soil containing larvae was kept moist and larvae hydrated using field water (water from a 3:1 mix of deionized water and Signy soil). Individual larvae were extracted by breaking apart soil substrate with a fine brush and tweezers or by washing through stacked

331 **Table I.** Summary of all treatments and methods explored in this study. Concentration refers to either salinity dilutions with a soil control or Virkon® 386
 332 dilutions. See 'Materials and methods' section for full details. 387

333 Treatment type	Life stage	Condition/concentration (%)	Temperature (°C)	Exposure duration	Survival assessment	N
335 Virkon® boot-wash simulation	Larvae	0.1	~20	10 s	72 h post-exposure	24
336		1.0	~20	10 s	72 h post-exposure	24
337		10	~20	10 s	72 h post-exposure	24
338 Virkon® thresholds	Larvae	0	4	18 h	Hourly	30
339		1.0	4	18 h	Hourly	30
340		4.0	4	18 h	Hourly	30
341		10	4	18 h	Hourly	30
342		0	20	8 h	Hourly	30
343		1.0	20	8 h	Hourly	30
344		4.0	20	8 h	Hourly	30
345		10	20	8 h	Hourly	30
346 Hot water boot wash	Larvae	0	40	10 s	72 h post-exposure	15
347		0	45	10 s	72 h post-exposure	15
348		0	50	10 s	72 h post-exposure	15
349 Salinity thresholds	Larvae	Soil	4	7 days	72 h post-exposure	30
350		0	4	7 days	72 h post-exposure	30
351		25	4	7 days	72 h post-exposure	30
352		50	4	7 days	72 h post-exposure	30
353		75	4	7 days	72 h post-exposure	30
354		100	4	7 days	72 h post-exposure	30
355	Eggs	Soil	4	35 days	35 days	30
356		0	4	35 days	35 days	30
357		25	4	35 days	35 days	30
358		50	4	35 days	35 days	30
359		75	4	35 days	35 days	30
360		100	4	35 days	35 days	30

359 2.0 mm and 0.5 mm mesh sieves. In the latter instance, all 414
 360 larvae were rested in control conditions for 48 h to ensure 415
 361 that the extraction process was not an additional stressor 416
 362 prior to treatment. All larvae were subsequently assigned 417
 363 to instars based on size (Bartlett *et al.* 2018a). 418
 364 Experiments using eggs were conducted in laboratories 419
 365 at Signy Research Station during January 2017, using 420
 366 recently laid egg sacs collected from moss banks 421
 367 surrounding the research station. Egg sacs were removed 422
 368 from the substrates as described in Bartlett *et al.* 423
 369 (2018b). As egg sacs are only available in quantity from 424
 370 the field, these were not included in the later Virkon® S 425
 371 experiments conducted in the UK. It has previously 426
 372 been shown that *E. murphyi* larvae can respire 427
 373 underwater (freshwater) for up to 28 days, so the effect 428
 374 of submersion itself is not considered a stressor within 429
 375 the timeframe of these experiments (Everatt *et al.* 430
 376 2014b). A summary of all treatments and associated 431
 377 methods is presented in Table I. 432

379 Preparation of Virkon® S solutions

380
 381 Correspondence with the manufacturers of Virkon® S 436
 382 (Lanxess, Germany, sourced from Fisher Scientific UK 437
 383 Ltd) indicated that Virkon® S begins to degrade at 438
 384 temperatures > 40°C and that, while a 10% Virkon® S 439
 385 solution can be prepared under laboratory conditions, 440

414 the maximum recommended concentration for practical 415
 416 use is 5% at room temperature (~20°C). Therefore, all 417
 418 Virkon® S treatments took place at room temperature or 419
 420 below. Dilutions were measured using a colorimeter, and 421
 422 it was found that we were able to mix a 10% dilution 423
 424 that showed no re-granulation during the course of any 425
 426 treatments. Virkon® S solutions were thus made up in 427
 428 concentrations of 0% (control), 0.1%, 1.0% and 10% 429
 430 with deionized water and stored at 4°C. 431
 432

424 Short-term exposures

425 Different life stages of insects can have various levels of 426
 427 pesticide tolerance (Athanasassiou *et al.* 2012). It was 428
 429 therefore important to measure any difference in the 430
 431 boot-wash effects between the different larval instars of 432
 433 *E. murphyi*. Volumes of 20 ml of 0.1%, 1.0% and 10% 434
 435 Virkon® S were measured out using a graduated syringe 436
 437 and deposited into separate 100 ml beakers. Three 438
 439 replicates ($n = 8$) of either L4, L3 or L2 larvae were 440
 441 placed on a 250 µm nylon net, which was folded and 442
 443 gathered together so that the larvae were together at the 444
 445 base. Larvae were then completely submerged in the 446
 447 different Virkon® S dilutions for 10 s (to simulate a 448
 449 typical boot-wash period). Upon removal, the net was 450
 451 blotted on tissue paper to remove excess Virkon® S, and 452
 453 the larvae were quickly returned to control conditions. 454

Survival was assessed after 72 h by visual monitoring of larvae movement, either spontaneously or with gentle stimulation with a brush. Independent peristalsis of the gut and/or movement of the mandibles were registered as live movement.

Long-term exposures

To assess the efficacy of warming Virkon® S and/or increasing exposure times, Virkon® S solutions of 0% (control), 1%, 4% and 10% were prepared and stored at either 4°C or 20°C. Three groups of $n = 10$ mixed L3/L4 larvae were placed in a Petri dish with 2 ml of each dilution at each temperature (no soil). The Petri dishes were kept at either 4°C or 20°C and survival was assessed every hour for 8 h. The time taken to reach 50% (lethal time, LT_{50}) or 100% (lethal time, LT_{100}) mortality was noted. Based on the results from the 8 h experiments, hourly assessments were repeated at only 4°C for all dilutions for a duration of 18 h, then left overnight and assessed again at 27 h, in order to assess the LT_{100} for each dilution.

High-temperature exposures

In order to establish the potential for hot water boot washes to act as an alternative biosecurity measure against *E. murphyi*, the above 'net and dip' method was used on three groups of $n = 5$ L4 larvae. A 28 ml test tube containing ~15 ml of field water was placed in an alcohol bath (Haake Phoenix II C50P) and heated to 40°C, 45°C or 50°C. A minimum of 40°C was chosen as *E. murphyi* larvae are known to survive short exposures to temperatures up to 39°C (Everatt *et al.* 2014a). Larvae were submerged in the heated water for 10 s, removed to control conditions and survival assessed immediately after exposure and then again at 24 and 72 h.

Salinity exposures

To assess the ability of *E. murphyi* to withstand immersion in seawater, we exposed both larvae and egg sacs to a range of salinities. For experiments on eggs, conducted on Signy Island, seawater was collected locally. All eggs within the egg sacs were confirmed to be at the first (opal) developmental stage prior to the start of experiments and were then used for the entire gestation period of 35 days (Bartlett *et al.* 2018a). If any eggs showed signs of yellowing or embryonic development, the whole egg sac was discarded and not used in this study. Experiments on larvae, conducted at the University of Birmingham, used Antarctic seawater obtained from stocks at the BAS. In all instances and for all dilutions, pH and salinity (μS) were measured using a Hanna HI-98129 Combimeter.

Three groups of $n = 10$ egg sacs were submerged for 35 days at 4°C in either a soil control, 0% (field/fresh water), 25%, 50%, 75% or 100% seawater. Development was noted weekly and, at the end of the gestation period (35 days), the egg sacs were carefully dissected and the percentage of eggs that had hatched recorded. For comparison with larvae, the same dilution experiment was conducted on three groups of $n = 10$ L4 larvae that were kept submerged for 7 days. After treatment, the larvae were returned to soil control conditions and survival assessed after 72 h, as described previously.

Results

Efficacy of the disinfectant Virkon® S and use of boot-wash protocols

Short (10 s) exposure to all concentrations of Virkon® S resulted in 0% mortality in both L4 and L2 larvae. Only one death was observed among L3 larvae. In the long-term experiments, immersion of larvae in water (control) over 18 h resulted in 0% mortality at both 4°C and 20°C (Fig. 2a). Exposure to the 1% Virkon® S resulted in some mortality after 4 h, but with no significant difference between 4°C and 20°C after 8 h (Mann-Whitney $U = 3$, $P = 0.7$), and with survival remaining > 50% even after 18 h at 4°C (Fig. 2b). There was a marked decline in survival in 4% Virkon® S, with LT_{50} observed after ~5 h at 20°C and after ~9 h at 4°C (Fig. 2c). LT_{100} was reached after 8 h at 20°C and after 14 h at 4°C. In 10% Virkon® S, mortality occurred after 3 h at 20°C, reaching LT_{50} at 5 h. Survival at 4°C also declined more rapidly at this concentration, with the LT_{50} being reached after ~7 h and LT_{100} being reached after 13 h (Fig. 2d). Overall, mortality after 8 h of exposure for all dilutions was significantly higher than controls at both 20°C (Kruskal-Wallis $H = 9.9$, $P < 0.0001$) and 4°C (Kruskal-Wallis $H = 8.2$, $P = 0.01$).

High-temperature treatments

At 40°C, there was no effect on survival after a 10 s exposure, whilst at 45°C, immediately post-exposure, all larvae were in a heat coma, but fully recovered to 100% survival within 24 h. Exposure to 50°C water resulted in 100% mortality of L4 larvae with no recovery over the post-exposure period of up to 72 h (Fig. 3).

Salinity exposure

The pH of field-collected vs laboratory-stored seawater (means of 6.7 ± 0.4 SEM and 6.2 ± 0.4 , respectively) were not significantly different (Mann-Whitney $U = 7$, $P = 0.3$). Salinity values (means of $25\,400 \mu\text{S} \pm 9030$ SEM and $27\,385 \mu\text{S} \pm 8312$ SEM, respectively) were also not significantly different (Mann-Whitney $U = 12$, $P > 0.99$).

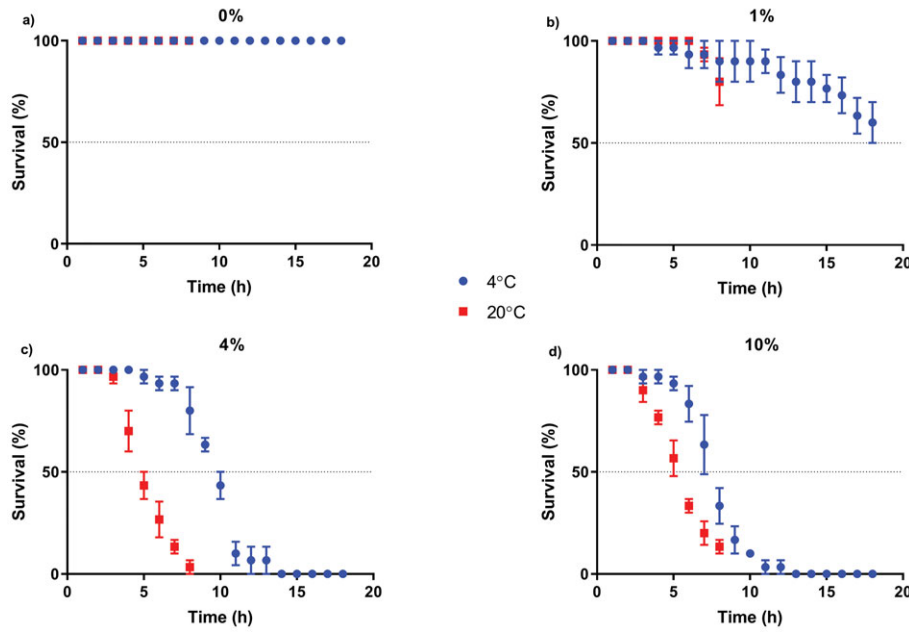


Fig. 2. Virkon® S exposures of **a.** control (0%), **b.** 1%, **c.** 4% and **d.** 10% dilutions at either 4°C (blue) or 20°C (red), with three groups of $n = 10$ larvae exposed for 1–18 h (4°C) or 1–8 h (20°C). Shown as mean survival at each time point \pm SEM.

Survival of L4 larvae in 0%, 25%, 50%, 75% or 100% seawater over a period of 7 days was not significantly different from that in the non-submerged soil control (Kruskal-Wallis $H = 7$, $P = 0.17$) (Fig. 4a), although there was a slight trend of declining survival with increasing salinity. In contrast, in egg sacs exposed to the same salinity range for their gestation of 35 days, the proportion of eggs hatching was greatly reduced even at low salinity (amongst all treatments: Kruskal-Wallis $H = 50.5$, $P < 0.001$; multiple comparisons between treatments: 100%, 75% and 50% vs soil, $P < 0.0001$; 25% and 0% vs soil, $P < 0.001$) (Fig. 4b). No eggs hatched under 50%, 75% and 100% seawater treatments. Exposure to 25% seawater or to field/fresh water led to hatching success rates of $4.5\% \pm 2.1\%$ SEM and $9.1\% \pm 5.8\%$

SEM, respectively, while hatching success rate in the soil control was $59\% \pm 7.7\%$ SEM. Observations made throughout the 35 day exposure period confirmed that the eggs developed within the egg sacs as described in previous studies, but that in all submergence exposures development slowed at maturation and, of the few eggs that did hatch under saline treatments, the L1 hatchlings did not survive and often did not fully escape from the egg casings within the egg sac, although they did survive freshwater treatments.

Discussion

Given the increasing number of non-native species found in Antarctica, improvements in biosecurity practice will be essential to ensure the ongoing protection of marine

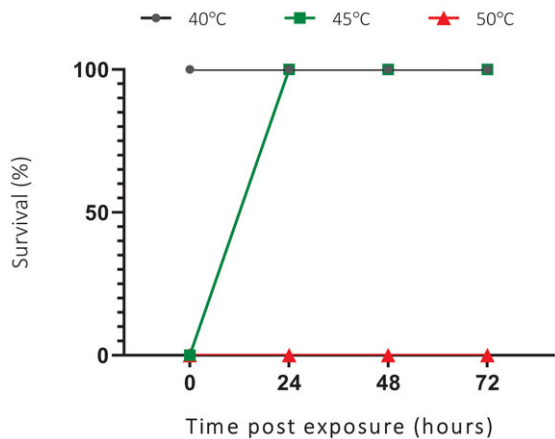


Fig. 3. Mean \pm SEM tolerance of larvae to 10 s exposures to hot water temperatures of 40°C, 45°C and 50°C and recovery over 72 h.

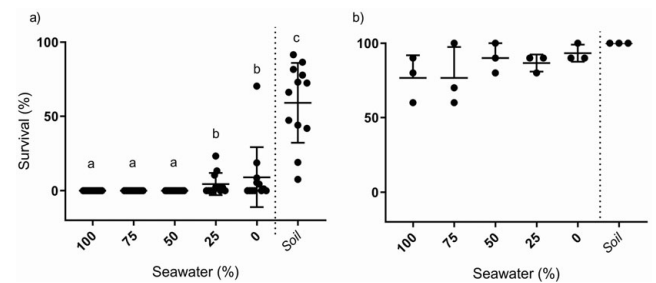


Fig. 4. **a.** Mean \pm SEM tolerance of eggs to seawater dilutions and a soil control after exposure for the whole gestation period (35 days). Three groups of $n = 10$ egg sacs, with ~ 70 eggs in each sac. Dilutions with the same letter are not significantly different. **b.** Mean \pm SEM tolerance of L4 larvae to seawater dilutions after 7 days of continuous exposure with a soil control.

661 and terrestrial ecosystems from biological invasion.
662 Current biosecurity protocols concerning the cleaning of
663 footwear primarily focus on reducing the risk of
664 microbial transfer, with the standard practice consisting
665 of dipping footwear into baths containing a 1% Virkon®
666 S solution for a few seconds and boot scrubbing to
667 directly eliminate any visible soil and macro-biology
668 (IAATO 2018, COMNAP 2019). While the consistency
669 of implementation of this procedure varies across
670 different operators in the region, both scrubbing and
671 boot-wash dips are mandatory on arrival at research
672 stations and deployment to field sites under the current
673 BAS biosecurity regulations (BAS 2019). BAS operates
674 a research station located on Signy Island (South
675 Orkney Islands, maritime Antarctic). Here, a principal
676 biosecurity threat is the transfer of two known
677 non-native invertebrate species to locations beyond their
678 current distribution on the island or to various islands
679 and the Antarctic Peninsula: the flightless midge
680 *E. murphyi* and the enchytraeid worm *Christensenidrilus*
681 *blocki*. Both are thought to have been introduced to
682 Signy Island in the 1960s during plant transfer
683 experiments involving material from South Georgia and
684 the Falkland Islands (Burn 1982). The physiological
685 capacity of *E. murphyi* to survive conditions further
686 south (Everatt *et al.* 2012), as well as to alter soil
687 processes (Hughes *et al.* 2013), makes further transfer of
688 this species to other sites in the region a particular
689 concern. Currently, BAS regulations specify boot
690 washing and scrubbing as a method to restrict the
691 transfer of these species from Signy Island itself to other
692 locations. However, recent evidence indicates that human
693 footfall is also a primary mechanism extending the
694 range of *E. murphyi* on Signy Island (Bartlett *et al.*
695 2020), and thus assessing the efficacy of boot-wash
696 protocols in limiting the spread of this (and potentially
697 other) invertebrate species on Signy Island is very timely.
698 There is clear evidence that Virkon® S can be lethal to
699 aquatic invertebrates and mud snails (Stockton-Fiti &
700 Moffitt 2017), but it has been ineffective in the only
701 studies in which it has been applied to terrestrial insects
702 to date: eggs of the yellow mealworm *T. molitor*
703 (Li *et al.* 2016) and the house fly *M. domestica* (Watson
704 *et al.* 2008). We present evidence that larvae of this
705 invasive midge experienced 0% mortality in
706 concentrations of up to 10% Virkon® S over periods of
707 well over 1 h. Indeed, LT₅₀ values at this highest
708 concentration were only reached after 8 h at field
709 temperatures (4°C), or after ~5 h at elevated
710 temperatures (20°C). Importantly, these experiments
711 were conducted with zero soil load (i.e. assuming 100%
712 removal of soil from footwear, but with a chance that
713 some larvae remained attached). This means Virkon® S
714 boot-wash protocols alone are totally ineffective
715 biosecurity measures for controlling the spread of

E. murphyi, and only meticulous boot scrubbing under
current protocols could prevent transfer of this species
from Signy Island to other locations or limit its spread
on the island.

Everatt *et al.* (2014a) showed that *E. murphyi* larvae
enter heat coma at 31°C, and a few individuals can
survive air temperatures up to 39°C for 1 h.
Consequently, we assessed temperatures > 40°C in the
absence of Virkon® S (which degrades at this
temperature). We found that very short exposures (10 s)
to 40°C or 45°C water, whilst inducing heat coma, were
not lethal. Only 50°C water proved to be effective at
killing *E. murphyi* larvae during typical/short boot-wash
exposure times. Saltwater exposures also proved
ineffective as a biosecurity measure for mature (L4)
larvae, which experienced very little mortality even after
7 days of submersion in 100% seawater (Fig. 3b).
First-instar larvae were highly susceptible to even dilute
saltwater exposure, with very low survival from egg
batches hatching under these conditions (Fig. 3a).

Based on the data obtained in this study, we suggest that
the use of hot water (> 50°C) to scrub soil containing
invertebrates off contaminated items, followed by a
Virkon® S wash on the clean boots, would provide the
most effective control measures currently available
against *E. murphyi* whilst not sacrificing the benefits of
Virkon® S as a microbicide/virucide. This could be
implemented at existing boot-wash stations both prior to
arrival and on departure from islands. To mitigate the
further spread of *E. murphyi* around Signy Island, ideal
scenarios would also include new scrub stations adjacent
to trails at the edge of the known *E. murphyi*
distribution (see Bartlett *et al.* 2020), although this raises
issues of practicality related to sourcing/heating water
and possible health and safety issues.

Whilst the focus of this study has been on the invasive
midge *E. murphyi* on Signy Island, the findings and
suggested additions to the existing protocols may be
relevant to all areas of Antarctica that are vulnerable
to invasive invertebrates or that have already been
colonized. *Eretmoptera murphyi* is not a unique example
in the Antarctic region, but as a flightless species, it is
reliant on mechanical, or potentially oceanic, methods
of dispersal to increase its range. Within the maritime
Antarctic, another dipteran species, *Trichocera maculipennis*,
was recently introduced to King George Island (South
Shetland Islands) (Volonterio *et al.* 2013, Potocka &
Krzemińska 2018). Although most attention has been
given to observations of this species having colonized
research station sewage systems, it is thought that it may
be established in the local natural environment
(Volonterio *et al.* 2013, Potocka & Krzemińska 2018).
As adults of this species can fly, it is capable of greater
natural dispersal than *E. murphyi*, but soil- or
substrate-dwelling life stages could be dispersed through

771 similar mechanisms to those of *E. murphyi* (Volonterio
772 *et al.* 2013). It is probable that all invertebrates will
773 succumb to temperatures > 50°C (Heinrich 1981); what
774 remains to be seen is the minimum exposure time
775 necessary to test this as a viable biosecurity method. We
776 therefore suggest that future work explore simple hot
777 water treatments such as that presented in this study
778 against other non-native invertebrates in the Antarctic
779 region in an attempt to develop a method that could
780 be universally applied throughout the region with
781 comparatively little logistical effort.

783 Conclusions

784
785 The combination of increasing human activity and
786 ongoing regional climate change will probably facilitate
787 further establishment and colonization events of
788 non-native species in continental, maritime and sub-
789 Antarctic regions. Regular review and revision of
790 established biosecurity protocols and the development
791 of new procedures will be necessary if the risk of
792 introductions is to be minimized. Preventing the transfer
793 of soil, and the micro- and macro-organisms contained
794 therein, needs to be a priority action for all stakeholders
795 involved in the protection of Antarctica. Here, using
796 *E. murphyi* as a model species, we have demonstrated
797 important limitations in probably the most widely
798 implemented biosecurity measures, and we suggest
799 alternative actions that could potentially be used to
800 reduce the spread of non-native invertebrate species
801 that, if left unchecked, have the potential to disrupt
802 Antarctica's fragile ecosystems.

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811 Author contributions

812
813 JCB, SALH and PC conceived the study. JCB, SALH and
814 PC designed the methodological approach. JCB and RJR
815 conducted the laboratory experiments. KAH and PC
816 provided policy input. JCB and RJR drafted the
817 manuscript. All authors edited and revised the
818 manuscript.

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831 Details of data deposit

832
833 The experimental data from this study are available online
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835 Richard James; Convey, Pete; Hughes, Kevin; Hayward,
836 Scott (2020), 'The effectiveness of Virkon® S disinfectant
837 against the invasive chironomid *Eretmoptera murphyi*
838 and implications for Antarctic biosecurity practices',
839 Mendeley Data, V4, doi: 10.17632/3686s39g9j.4
840 (available at <http://dx.doi.org/10.17632/3686s39g9j.4>).

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