

A Rapid Molecular Assay for Detecting the Mediterranean Fanworm Sabella spallanzanii Trialed by Non-Scientist Users

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Zirngibl M, von Ammon U, Pochon X and Zaiko A (2022) A Rapid Molecular Assay for Detecting the Mediterranean Fanworm Sabella spallanzanii Trialed by Non-Scientist Users. Front. Mar. Sci. 9:861657. doi: 10.3389/fmars.2022.861657 The invasive Mediterranean fanworm Sabella spallanzanii (Gmelin, 1791) is a notifiable organism under New Zealand's Biosecurity Act and is recognized as a marine pest of particular concern, that must be reported to the Ministry for Primary Industries (MPI), New Zealand. Since its first detection in 2008, great effort and financial resources are put into surveillance and removal of individuals to contain population growth and spread. Sensitive molecular detection techniques gain great interest and are being increasingly tested for the fanworm detection in marine high-risk sites (i.e., ports and marinas) around New Zealand. However, conventional molecular detection via PCR assays from environmental DNA (eDNA) samples requires specific laboratory resources and technical expertise. This restricts the wider applicability of this approach by biosecurity practitioners or communities willing to be engaged in biosecurity surveillance. To provide end-users with a fast, easy and highly specific way to detect S. spallanzanii directly at the site of interest, a species-specific recombinase polymerase amplification (RPA) assay was designed to be read-out with lateral flow strips (RPA-LF). The RPA generates amplification within 20 minutes at 37-39°C, with a detection limit of 10 pg of the target DNA and was matching the detection limit of digital droplet PCR (ddPCR) when performed on eDNA samples. A simplified visual protocol for non-scientist users of the assay was developed and improved through independent trials with different end-user groups. The assay applicability was verified in a final validation trial with participants without scientific background resulting in 50 percent of the participants successfully detecting S. spallanzanii. Participants rated the ease of use and performance and read-out mostly as easy-to-very easy with overall positive written feedback on its usability for citizen science applications.

Keywords: recombinase polymerase amplification (RPA), point-of-need (PON), environmental DNA (eDNA), lateral flow detection, Sabella spallanzanii

INTRODUCTION

The Mediterranean fanworm Sabella spallanzanii (Gmelin, 1791) is a tube-forming annelid, native to the Mediterranean Sea and Atlantic coast of Europe (Patti and Gambi, 2001). It has successfully invaded Australia in the 1990s (Clapin and Evans, 1995) and was first recorded in New Zealand in 2008 in Lyttelton Harbor (Read et al., 2011). Despite efforts to eradicate S. spallanzanii at the initial incursion area and mitigate its further spread (Fletcher, 2014), soon it was found in multiple harbors across the North and South Islands (Fletcher, 2014). Being a highly effective suspension filter feeder (Clapin, 1996) and an ecosystem engineer (O'Brien et al., 2006), the fanworm can impact abundance and recruitment patterns of invertebrate larvae (O'Brien et al., 2006; Ross et al., 2007) and alter benthic macrofauna and microbial communities adjacent to S. spallanzanii canopies (Atalah et al., 2019). Because of its high invasive potency and adverse ecological impacts, the uncontrolled spread of S. spallanzanii threatens species native to New Zealand and puts local economies at risk. In New Zealand, S. spallanzanii invasions are particularly problematic for the shellfish aquaculture sector. Dense biofouling by the fanworm, competing with mussels for food and space, may cause reduced mussel growth and compromise their attachment, therefore leading to crop losses (Bannister et al., 2019). It is estimated that S. spallanzanii could cause financial losses of roughly 14 Million NZ Dollar to the green shell mussel industry (Soliman and Inglis, 2018).

Therefore, Sabella spallanzanii is listed as a notifiable and unwanted organism in New Zealand under the Biosecurity Act 1993 (Hewitt et al., 2004; Inglis et al., 2005; Read et al., 2011) and is currently subjected to targeted surveillance as part of the national Marine High Risk Site Surveillance (MHRSS) program (Inglis et al., 2005; Wood et al., 2019b). Although the juvenile/adult stage of S. spallanzanii is morphologically distinct from most New Zealand native marine organisms and could be identified easily, when juveniles/adults are abundant in a marina, eradication or population management at this stage of invasion is near to impossible. In most cases, detecting juvenile/adult individuals at early incursion stages is crucial for effective response but very difficult to accomplish via visual surveys from the surface, i.e. when it is part of biofouling on a boat hull or harbor structures. Therefore, to effectively monitor its spread, detect incursions at the earliest instance and mitigate establishment of new populations, biannual diver surveys are conducted in New Zealand's major ports and marinas (Inglis et al., 2005). Visual surveying and removal by divers can be highly efficient with a 80 - 100% detection rate for juvenile/adult individuals (Inglis et al., 2008; Inglis et al., 2009), however smaller individuals and pelagic larvae are difficult to detect and identify without a microscope and expert taxonomist (Giangrande et al., 2000; Floerl et al., 2010). Additionally, organizing and conducting underwater visual surveys are logistically complex, expensive and subject to restrictions like weather and site access and pose health and safety risks to divers in the busy port or marina environments (Bowers et al., 2021). Therefore, biosecurity authorities are interested in alternative, ideally more sensitive and cost-efficient surveillance technologies.

In recent years, molecular techniques have gained increasing popularity for bio-surveillance applications, e.g., community screening using metabarcoding or species-specific detections using targeted assays (Pochon et al., 2017; Wood et al., 2017; Zaiko et al., 2018; Hernandez et al., 2020; Gold et al., 2021). For example, a quantitative real-time polymerase chain reaction (qPCR) (Wood et al., 2017) and digital droplet polymerase chain reaction (ddPCR) assays (Wood et al., 2019a) have been developed and validated for environmental DNA (eDNA) based detection of S. spallanzanii in New Zealand's coastal environments. These assays, however, require specific laboratory gear and technical expertise, which restricts their wide use and point-of-need applicability by non-scientist users, such as biosecurity practitioners or communities willing to be engaged in biosecurity surveillance. Engaging non-scientists to perform assays for invasive species detections could support biosecurity surveillance efforts in mitigating the establishment and spread of invasive organisms as early as possible.

To address the current need in molecular surveillance techniques that are easy, fast, specific, and usable with minimal equipment and expertise, we have designed a recombinase polymerase amplification (RPA) assay for species-specific S. spallanzanii detection. Recombinase polymerase amplification is an isothermal amplification method, that amplifies target DNA within 20 minutes at a temperature range of 25°C to 45°C (Piepenburg et al., 2006; James and Macdonald, 2015; Daher et al., 2016), which makes it possible to run assays at body temperature (~37°C), i.e., hand-held (Crannell et al., 2014). With the introduction of labelled probes and primer, RPA can be used for fluorometry-based quantitative real time amplification. Detection is also possible with lateral flow (LF) strips, that provide easy visual read-outs (Piepenburg et al., 2006). The combination of RPA and LF (RPA-LF) strip is especially valuable in resource-limited environments. RPA-LF was developed successfully for multiple disease carrying organisms/ viruses (Li et al., 2019a), SARS-CoV-2 (Lau et al., 2021) or for invasive species like the Northern Pacific Seastar, Asterias amurensis (Unpublished).

Here we present the results of the successful *in silico* and *in vitro* validation of a newly designed RPA-LF assay for the Mediterranean fanworm *S. spallanzanii*, as well as the trialing of a simplified RPA protocol with different end-user groups.

MATERIALS AND METHODS

Samples for Recombinase Polymerase Amplification (RPA) Assay Development and Validation

New Zealand native Sabellidae species were obtained from the NIWA Invertebrate Collection, (NIC, Wellington, New Zealand) and the Marine Invasives Taxonomic Service (MITS, Wellington, New Zealand) (**Table 1**), and *S. spallanzanii* specimens were collected by commercial divers in Tasman Bay, New Zealand. Genomic DNA was extracted from roughly 25 mg tissue of one sample of each Sabellidae species (**Table 1**) using the QIAamp[®]

TABLE 1	New Zealand Sabellidae species used for cross-reactivity testing.
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Species	Type locality	Coordinates	Sample Date	Catalogue #	Archived Specimen Collection
Acromegalomma suspiciens	Port Auckland, NZ	41.2844400°S	07.10.2009	75089	MITS
		174.4136110°E			
Bispira bispira	Frenchman Island, NZ	35.8608341°S	17.04.2010	60432	NIC
		174.5275116°E			
Branchiomma curtum	Port Lyttelton, NZ	N/A	28.09.2009	41167	MITS
Euchone pallida	Kahuwhera Bay, NZ	35.2592400°S	21.09.2010	155832	NIC
		174.1846000°E			
Parasabella aberrans	Port Nelson, NZ	41.26365278°S	28.02.2014	70868	MITS
		173.2803806°E			
Pseudobranchiomma grandis	Port Lyttelton, NZ	N/A	22.07.2009	41021	MITS
Pseudopotamilla laciniosa	West Coast, NZ	N/A	29.03.2007	57786	MITS
Sabellastarte spectabilis	Port Whangarei, NZ	35.7577000°S 174.3490000°E	19.10.2012	70319	MITS

NIC, National Institute of Water and Atmospheric Research Invertebrate Collection; MITS, Marine Invasives Taxonomic Service; N/A, not available.

DNA Mini Kit (Qiagen, Germany) with 12 h Proteinase K digest and following the manufacturer's instructions. Additionally, five eDNA samples collected in Waitematā Harbor (New Zealand) (**Supplementary Table 2**) which is known to harbor *S. spallanzanii* populations were also included to further validate the RPA assay (von Ammon, in review). The sampling sites were a minimum of 1 km apart. We used a previously developed digital droplet PCR assay specific to *S. spallanzanii* (Wood et al., 2019a) to confirm that *S. spallanzanii* DNA was either present or absent in the environmental samples in order to verify the results obtained from our RPA assay on those same samples.

Primers and Probes Design

All primers and probes used in this study were designed in Geneious Prime version 2020.2.2 (https://www.geneious.com) following the TwistDxTM (Cambridge, England) guidelines. Recombinase polymerase amplification primers and probe were designed to amplify the mitochondrial Cytochrome C Oxidase subunit 1 (COI) gene of *S. spallanzanii* based on a multiple sequence alignment previously generated (Wood et al., 2017). For detection on the lateral flow strip, the reverse primer was labelled with biotin at the 5' end, the LF probe was labelled with 6-FAM (5') and the polymerase extension blocker C3-spacer (3') contained an internal abasic tetrahydrofuran (idSp) (**Table 2**). All primers and probes were checked *in silico* for cross-reactivity with other species using the Primer-BLAST tool (https://www.ncbi.nlm.nih.gov/tools/primer-blast/) and checked with the OligoAnalyzerTM (IDT) for *in situ* parameters. All

primers and probes were synthesized by Integrated DNA Technologies, Inc. (IDT, Singapore).

Recombinase Polymerase Amplification (RPA) and Droplet Digital PCR (ddPCR) Analyses

Testing of RPA primers was carried out using the TwistDxTM TwistAmp[®] Liquid Basic kit, with reaction volumes adapted to a total reaction volume of 20 μ L. Reactions contained 10 μ L 2 x reaction buffer, 2.1 μ L H₂O, 2 μ L 10 x basic E-mix, 1 μ L 20 x Core Mix, 0.9 μ L dNTPs (InvitrogenTM, USA), 0.96 μ L primer each (10 μ M), 1 μ L genomic DNA (2 ng DNA total unless stated otherwise) and 1 μ L magnesium acetate to start the reaction. The reaction mixture was incubated at 37°C for 20 minutes. To ensure even distribution of reaction components, the reaction tube was flicked after 5 minutes. Amplified products were cleaned up using QIAquick[®] PCR Purification Kit (Qiagen, Germany) to remove any interference of RPA proteins in the electrophoresis process. The products were visualized on 1.5% agarose gel, using RedSafeTM Nucleic Acid Staining Solution (iNtRON Biotechnology, Korea).

Recombinase Polymerase Amplification Lateral Flow (RPA-LF) Reaction Mixture

Due to unavailability of the TwistDxTM Nfo kit (implications of the restricted supplies caused by SARS-CoV-2 pandemic), Endonuclease IV (EN0591, Thermofisher, USA) was supplemented into the Basic mixture to replicate the functionality of the TwistDxTM Nfo kit. We tested different

TABLE 2 | Primer and probe sequences targeting Sabella spallanzanii, designed and tested for Recombinase Polymerase Amplification (RPA) and Recombinase Polymerase Amplification with lateral flow read-out (RPA-LF) analyses used in this study.

Assay	Name	Sequence (5'-3')		
RPA	Sab_RPA_F	GGATTTGGAAATTGATTATTGCCCCTGATG		
	Sab_RPA_R	AATAAAGTTTACTGCCCCTATAATAGAAGAAGC		
RPA-LF	Sab_LF_F	GGATTTGGAAATTGATTATTGCCCCTGATG		
	Sab_LF_R	/5Biosg/AATAAAGTTTACTG CCCTATAATAGAAGAAGC		
	Sab_LF_Probe	/56-FAM/ATTTATCCTCCTTTGTCAAGAGGAGTTGGA/idSp/ATAGAGGGTCTTCCA/3SpC3/		

Modifications as per IDT nomenclature: /5Biosg/ = 5' Biotin; /56-FAM/ = 5' 6- Carboxyfluorescein (FAM): /idSp/ = abasic tetrahydrofuran (THF); /3SpC3/ = 3'extension block. Bold values in sequence represent primer and probe modifications as per Integrated DNA Technologies (IDT) nomenclature. concentrations of Endonuclease IV (0.2 and 0.5 U) and incubation times (5 – 40 min) at 37°C to determine the optimal reaction setup (2 ng total *S. spallanzanii* DNA per reaction). The resulting optimized RPA-LF reaction was: 10 μ L 2x Reaction buffer, 2.1 μ L H₂O, 2 μ L 10x basic E-mix, 1 μ L 20x Core Mix, 0.9 μ L dNTPs (InvitrogenTM, USA), 0.84 μ L primer each (10 μ M final), 0.24 μ L Probe (10 μ M), 0.5 U Endonuclease IV (0.25 μ L of Endonuclease IV), 1 μ L genomic DNA and 1 μ L magnesium acetate (280mM stock concentration) to start the reaction. The reaction mixture was incubated at 37°C for 20 minutes and subsequently heat treated (5 minutes 90°C) to stop the reaction. To further increase sensitivity, we used 3 μ L template (excluded H₂O in the mixture) and incubated the reaction for 40 minutes (for amplification of low target DNA concentration e.g., environmental samples).

For detection of RPA products on the Milenia HybriDetect 2T lateral flow strips (Milenia Biotec GmbH, Germany), the RPA reactions were diluted 1:5 with HybriDetect 2T Assay Buffer and 10 μ L of the diluted mixture was applied to the sample area. The LF strip was placed vertically into a test tube filled with 80 μ L running buffer. Results were read after 5 - 10 minutes (longer incubation increases strength and clarity of bands), but within a maximum of 24h. The amplification was regarded as successful when a purple band was visible in the test-line. The run was only regarded as valid, when a purple band in the control lane (CTRL-lane) was visible, as this ensured proper functionality of the test strip itself. Assays were run repeatedly (min. 3 times) to test for consistency.

Assay Specificity

The specificity of the RPA-LF assay was tested with DNA from Sabellidae species native to New Zealand and closely related to S. spallanzanii (Table 1), using the above-described RPA-LF reaction mixture. Additionally, the environmental samples A -D containing S. spallanzanii (Supplementary Table 2) were amplified with the RPA primers Sab_RPA_F and Sab_RPA_R (Table 2). Successful amplification was confirmed using agarose gel electrophoresis (1.5% agarose gel with RedSafeTM Nucleic Acid Staining Solution [iNtRON Biotechnology, Korea]). Reactions were purified using QIAquick[®] PCR Purification Kit (Qiagen, Germany). Purified amplification products were bidirectionally sequenced using an ABI 3730xl DNA Analyzer at the University of Otago Genetic Analysis Services. The sequencing reads were aligned in Geneious Prime version 2020.2.2 (https://www.geneious.com). The consensus sequence of the alignment was checked against the GenBank reference database (Nucleotide collection) using the Nucleotide BLAST tool (https://blast.ncbi.nlm.nih.gov/Blast.cgi).

Limits of Detection

To assess the limit of detection, the RPA-LF assay was run on a 10-fold dilution series of *S. spallanzanii* DNA ranging from 1 ng to 100 fg of total DNA per reaction (0.05 ng/µl and 5 fg/µl respectively). The limit of detection was determined as the lowest concentration of *S. spallanzanii* DNA that produced a visible band on the lateral flow strip.

Comparative Sensitivity Assessment With the Droplet Digital PCR (ddPCR)

A ddPCR analysis specifically targeting *S. spallanzanii* was performed as described in (Wood et al., 2019a) for comparative assay sensitivity assessment. Briefly, the reaction mixture contained $1 \times BioRad$ ddPCR Supermix for probes (no UDP), 450 nM of each primer and probe (Sab3F, Sab3R, Taqman probe Sab3: Wood et al., 2017), 1 µl environmental DNA and sterile water at a total volume of 21 µL. After droplet generation in a BioRad QX200 droplet generator, the reactions were amplified at 95°C for 10 min (initial denaturation), followed by 40 cycles of 94°C for 30 s, 60°C for 1 min, and a final enzyme deactivation step at 98°C for 10 min. Positive droplets were quantified using a QX200 droplet reader using the "Direct quantification" setting.

Assay Applicability Assessment by Different User Groups

To assess the wider usability of our RPA assay, we designed a simplified step-by-step protocol focused on minimizing steps that might pose difficulties to people with minimal or no background in molecular biology and laboratory procedures. This protocol was tested in two preliminary trials by groups of 5 (Preliminary Trial 1) and 4 (Preliminary Trial 2) volunteer participants, representing different educational backgrounds and occupations to improve the protocol. A final validation trial with 10 participants was performed using the improved protocol and its usability was assessed based on the final trial.

Before each trial, all participants were briefed on the protocol with a quick demonstration of how to use micropipettes. During the experiment, participants were asked to individually follow the protocol, running blinded RPA tests on 4 standardized sample types spiked with S. spallanzanii DNA to a final concentration of: 0 ng/ μ L [negative sample (neg)], 2 ng/ μ L [low concentration (low)], 20 ng/µL [medium concentration (mid)], and 200 ng/µL (high concentration [high]. Due to limited availability of necessary material at the time, participants did not perform replicate runs of each sample. Upon completion of the tests, participants were asked to report detection/non-detection of the target, based on their LF strip read-outs, and return their results in a post-experiment survey including their opinion on the usability of the assay and other suggestions on how to improve the protocol (Supplementary File "Survey and Result Sheet"). The involvement of voluntary participants for the validation trials and the survey were approved by the Cawthron Institute Ethics board (Approval number: CAW-ETH-210427). During preliminary trials (i.e., Trial 1 and 2), at least two molecular scientists were present to observe if participants struggled and supported the participants to proceed. The observations of the scientists and feedback from the participants from Preliminary Trial 1 were used to adjust the protocol before Preliminary Trial 2. Our observations and the participants feedback from Preliminary Trial 2 were in turn used to adjust the protocol before the Validation trial. During the final validation trial, no help was provided to the participants. Due to the fact that we assisted some of participants in preliminary trials

1 and 2, we decided to only use the survey and results from the final validation trial to draw conclusions on the usability of this assay as a point-of-need tool.

RESULTS

Design and Validation of the Recombinase Polymerase Amplification With Lateral Flow Read-Out

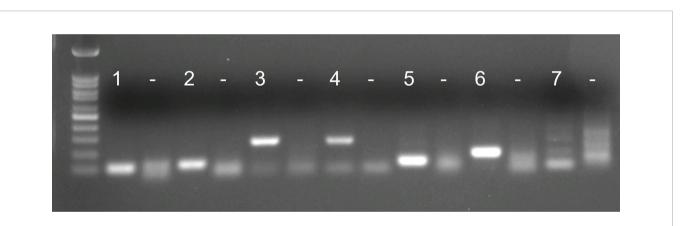
Of the initially designed eight forward and eight reverse primers targeting the COI gene region, only six primer combinations (1 - 6) – with acceptable *in silico* primer parameters, checked with the IDT OligoAnalyzerTM tool – and the existing *S. spallanzanii* qPCR/ddPCR primers (7) (Wood et al., 2017) were chosen for further testing (**Supplementary Table 1**). All combinations of newly designed RPA primers produced a singular band (lane 1 – 6) corresponding to the expected length (**Figure 1**). The diffuse bands in the negative control are attributed to primer dimers. The existing qPCR/ddPCR primers (lane 7) produced multiple weak bands and were not considered further. Primer

combination 3 (lane 3) (Figure 1, Supplementary Table 1) performed best and was chosen for the development of the RPA-LF and modified to be compatible with lateral flow readout and an appropriate probe was designed (Table 2)

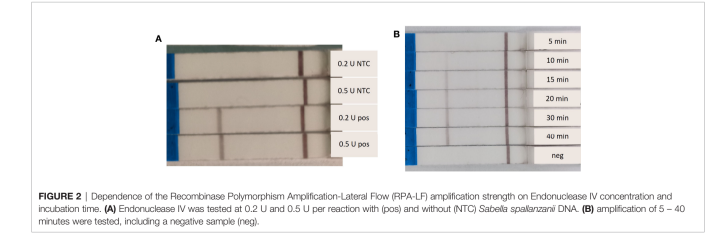
Different Endonuclease IV concentrations and amplification times were tested. The RPA-LF reactions were affected by the concentration of Endonuclease IV. Both tested concentrations (0.2 and 0.5 U) resulted in amplification after 20 minutes at 37°C, however the reaction containing 0.5 U produced a slightly stronger band (**Figure 2A**). We also observed a clear dependence of the band strength on the incubation time. A band was visible on the flow stick after 10 minutes of amplification time. The band strength increased with increasing amplification time, with maximum band strength after 40 minutes (**Figure 2B**).

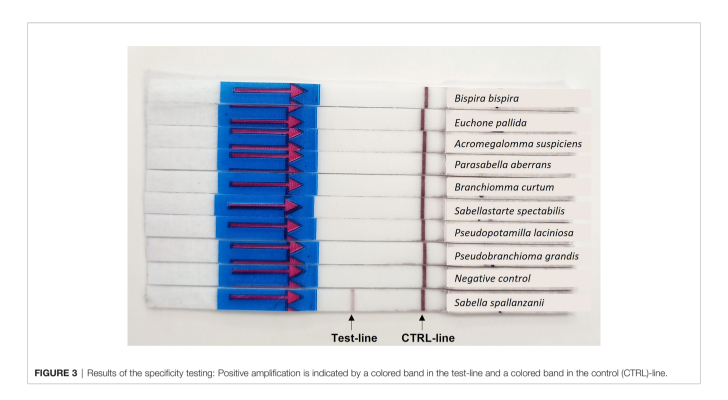
Species Specificity and Limit of Detection

Out of all tested Sabellidae species only *S. spallanzanii* was amplified by RPA-LF, suggesting high specificity of the designed assay (**Figure 3**). No false positive result in the negative control sample was detected either.







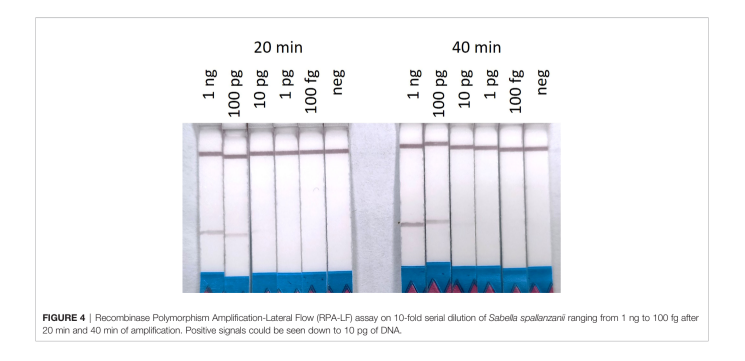


The RPA-LF limit of detection was established after the standard 20 minutes and extended 40 minutes amplifications. The limit of detection for *S. spallanzanii* DNA, that resulted in a visible band after 20 minutes, was 10 pg (0.5 pg/ μ L). After 40 minutes amplification time, the lowest concentration that produced a visible band was still 10 pg, however the band was slightly brighter (**Figure 4**). The ddPCR assay performed on the same dilution series, consistently produced positive signals at the

100 fg concentration but also returned a positive result in the 1 fg sample for one out of the two runs (**Supplementary Figure 1**).

Validation on Environmental Samples

When tested on five environmental samples (**Supplementary Table 2**), the RPA-LF assay was able to detect *S. spallanzanii* at ddPCR detection signals as low as 0.49 copies/ μ L, producing overall consistent results with the ddPCR assay. Only the lowest



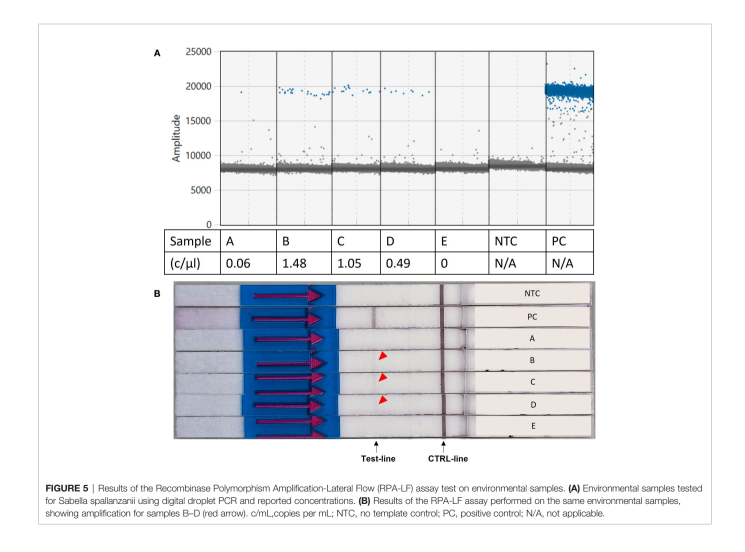
reported concentration (sample "A", 0.06 copies/ μ L) failed to produce a visible band on the flow strip. Sample "E" was negative for both the ddPCR assay and the RPA-LF (**Figure 5**).

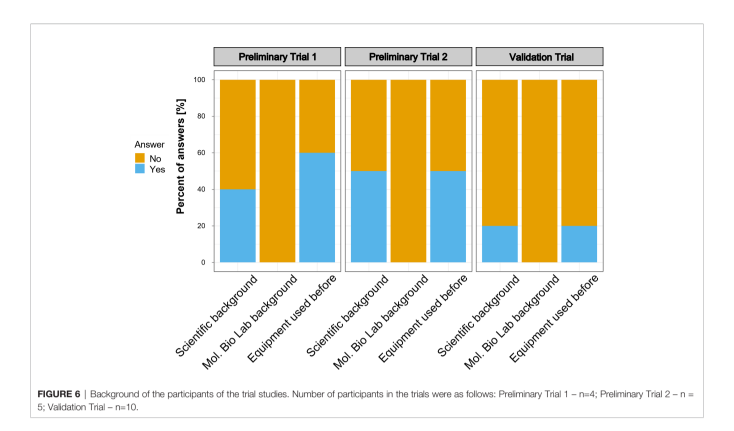
The Nucleotide BLAST of the amplified environmental samples returned the *S. spallanzanii* COI region as 100% sequence identity and query cover with an e-value of 2e-100 (highest hit: MN184708.1, **Supplementary Table 3**). Two other species matched the submitted consensus sequences with similar value were *Sabella pavonina* (Sequence identity: 100%; Query cover: 100.00%; e-value: 2e-100) and *Hermodice carunulata* (Sequence identity: 99.02%; Query cover: 100.00%; e-value: 4e-97, GenBank Accession number: AY495947.1).

Assay Trials With Different User Groups

The participants of the preliminary assay trials and the validation trial represented different groups of potential users: biosecurity practitioners, scientists, high-school students, indigenous communities, and public representatives. None of the nine preliminary trial participants had molecular laboratory background. Five participants reported some experience operating a micropipette at least once in the past. Similarly, none of the participants in the validation trial had a background in molecular biology and only 2 out of 10 participants reported having a scientific background and used pipettes before (**Figure 6**). In the post-experiment surveys of the preliminary trials, participants mainly suggested to include a section about materials and an induction on micropipette usage. The issue regarding micropipette usage was consistent with our observations during the preliminary trials; therefore, we incorporated all observations and feedback from the participants into the final version of the simplified protocol (**Supplementary Figure 2**) and included a pipetting guide (**Supplementary Figure 3**) and provided these updated guides to the participants of the validation trial.

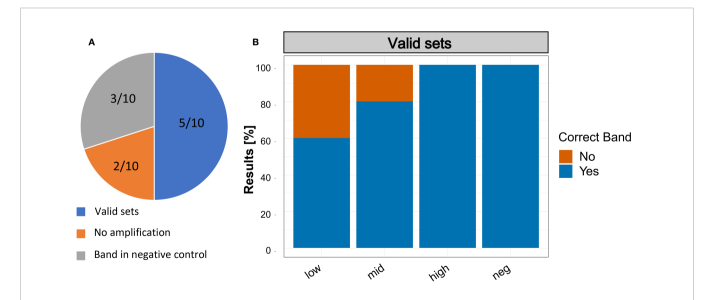
The assay results reported by the participants of the validation trial showed that three participants detected a positive band in the negative control. Also, two participants reported no positive band in any sample (**Figure 7A**). Looking only at valid sets – by removing the five participants, that had either no band at all or a band in the negative control – all participants detected *S. spallanzanii* in the 'high' concentration range. Two out of 5 did not detect *S. spallanzanii* in the 'hiw' sample and only one out of 5 did not detect it in the 'mid' concentration range (**Figure 7B**).

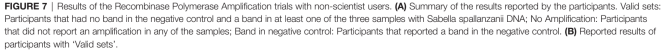


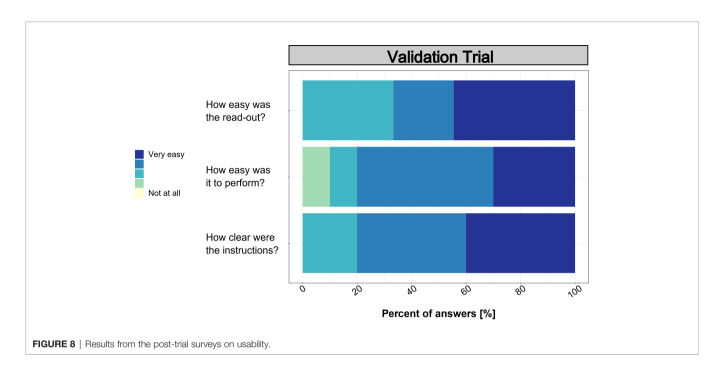


Based on the useability assessment performed after the validation trial (**Figure 8**), the clarity and ease of use values ranged mainly between 3 and 5 (i.e., medium to very easy/clear). Only one participant evaluated the ease of use with a 2 (not easy).

However, written feedback was mainly positive and suggested that the procedure was easy and clear to follow. Still, pipetting was mentioned by multiple participants as difficult as well as the feedback to include a video description.







DISCUSSION

In this study we developed and validated an RPA-LF assay for detecting the Mediterranean fanworm Sabella spallanzanii. Our assay showed no cross-reactivity with other Sabellidae species native to New Zealand. When we sequenced the amplification product obtained from the RPA reaction run on environmental samples and checked the sequence against GenBank using BLAST our top match was indeed S. spallanzanii. However, one closely matching polychaete species, Sabella pavonina could be amplified with our assay but is not reported to occur in New Zealand. The other close match - AY495947.1 - which was published as H. carunculata on GenBank, was later found to be a sample mix-up and indeed corresponded to S. spallanzanii and not H. carunculata (Ahyong et al., 2017). Therefore, we consider that the current assay can be robustly applied for targeted S. spallanzanii detection in New Zealand marine ecosystems but might require additional cross-reactivity validation when intended for other regions where S. pavonina might be present.

The detection limit established for our RPA-LF was 10 pg per reaction, which is higher than those reported for other published RPA-LF assays (Li et al., 2019b; Ravindran et al., 2020). We attribute this variation to the replicated reaction mixture used in this study due to unavailability of the standard RPA (TwistDx) kit. The detection limit of the newly developed RPA (10 pg/ reaction) is also higher than the existing ddPCR assay for *S. spallanzanii* with a detection limit of 100 fg per reaction. However, when tested on environmental samples, RPA-LF consistently detected *S. spallanzanii* even at concentrations close to the ddPCR detection limit and failed to amplify only one ambiguously positive sample, which returned a single droplet above the amplitude threshold in the ddPCR assay. The RPA-LF assay has a higher limit of detection compared to

the ddPCR assay when run on the dilution series of pure *S. spallanzanii* DNA but has comparable sensitivity when run on environmental samples. Considering, that detection of *S. spallanzanii* from eDNA is the main intended application, we conclude that the detection limit of the RPA-assay is acceptable.

In order to use the RPA-LF assay for 'point-of-need' applications by a wide range of potential end-users, the execution of the assay needs to be easy enough for people with no or limited laboratory background, while ensuring robust and reliable results (Larson et al., 2020; Bowers et al., 2021). We found that our visually designed step-by-step protocol was generally easy to follow by participants. One of the biggest hurdles was the use of micropipettes by inexperienced participants, which we tried to overcome by adding a visual explanation on pipette usage in the final version of the protocol. However, to make it accessible to citizens of any background, one needs to find ways around micropipettes e.g. by using so called Exact Volume transfer pipettes - disposable Pasteur pipettes that can be used to transfer a specific amount of liquid. Overall, based on the trial results and feedback provided by the participants, we can conclude that our protocol and RPA-LF assay has the potential to be used as a 'point-of-need' surveillance tool, e.g. for regional councils or similar institutions that already have pipettes and some scientific background.

Half of the ten participants were able to detect *S. spallanzanii* in at least one of the samples, while having no band in the negative control. Among those five, three participants reported all three samples containing *S. spallanzanii* as positive. However, three participants reported a band in the negative, and two did not report a band in any of the samples. Although the results that participants reported were overall promising, we do acknowledge that a 50% failure rate is rather high and further adjustments to the protocol or a more extensive training of volunteers is

warranted. A bigger test group and replication to assess the consistency of the results might be also recommended. Providing a video instruction could give additional visual aid for participants and could be implemented as a standardized training resource for future users. Also, very faint bands, such as seen in the test on environmental samples (**Figure 5**) might pose difficulties for untrained users and could lead to false negative reports. This user-bias might need to be offset by higher sampling effort and replication.

False positive results in eDNA-based assays represent a major issue for biosecurity surveillance, as false detections may trigger unnecessary response measures and compromise the credibility of molecular detection techniques for routine monitoring (Sepulveda et al., 2020; Jerde, 2021). False positive results can occur due to a lack of specificity or cross-contamination between samples (Ficetola et al., 2016). It is quite common that undesired amplification of negative controls happens in eDNA studies conducted by established molecular laboratories (Maruyama et al., 2014; Sepulveda et al., 2019), despite implementation of widely accepted guidelines to minimize/prevent crosscontaminations (Goldberg et al., 2016). To rule out false positives in the RPA-LF in a citizen science setting, a positive detection of S. spallanzanii should be further confirmed through qPCR/ddPCR. Although we acknowledge, that further improvements and a higher number of participants would need to be involved to make such assays reliably usable by non-scientists, we believe that our protocol and validation with non-scientists serve as a proof-of-concept that such assays can be run by non-scientists.

The workflow for eDNA-based detection of aquatic pests comprises several critical steps, including sample collection and DNA extraction. Traditionally, both steps require specialized equipment and trained personnel, and are therefore restricted to a very small group of dedicated specialists. To fully harness the utility of the RPA-LF assay as a 'point-of-need' tool and for application by non-scientist users, inexpensive, easy-to-use, yet robust and efficient sampling and DNA extraction methods are crucial. Some promising technologies are emerging in that space, that would greatly decrease the complexity of the workflow and make eDNA-based molecular biosecurity surveillance more accessible for a wide range of end-users. For example, the sampling routine can be greatly simplified by using passive sampling devices, that capture eDNA onto a sampling matrix while being deployed in the water (Kirtane et al., 2020; Bessey et al., 2021). As an alternative to time-consuming bulk/planktonnet water sampling and subsequent filtration (von Ammon et al., 2020; Bowers et al., 2021), in situ filtration systems with selfpreserving filter packs are now available on the market (Thomas et al., 2019). For the DNA extraction step some promising tools are reported in the literature to be fast, produce high-quality DNA extracts and utilize equipment that can be battery-operated such as the Pretty Damn Quick Extractor (PDQeX, MicroGem), which has been successfully used in low-resource settings for pathogen detection (Boykin et al., 2019). Another commercially available solution, the SpeedXtract kit (Qiagen) has already been coupled with RPA assays (Mondal et al., 2016). Still, their

efficiency on different sample types as well as compatibility with different molecular detection assays should be validated, and cost/time efficiency formally assessed through scientific and end-users' trials.

We anticipate, that easy-to-follow molecular detection tools can be also utilized for educational purposes. Provided they are affordable and easily accessible, those tools could be used in classrooms as a hands-on experience accompanying theoretical knowledge on marine ecology, molecular biology, biodiversity and biosecurity concepts. Showcasing molecular detection assays in classrooms or public seminars as educational components would contribute to raising the public awareness about nonindigenous marine species, and ultimately help establish citizenled biosecurity surveillance programs that effectively enable the early detection of unwanted invasive species at unprecedented spatial scales.

CONCLUSION

This study designed, tested and validated a novel RPA-LF assay for detection of the invasive Mediterranean fanworm S. spallanzanii from environmental samples. Levels of detection using the lateral flow stick approach were similar to those afforded by ddPCR diagnostics, but sensitivity could presumably be increased when using our primers with the fitfor-purpose TwistDx Nfo kit. Our simplified protocol is a promising means to make molecular tools more accessible to non-scientist users, although we acknowledge that some basic laboratory training would be beneficial for this purpose. Based on this experience, we recommend an iterative process whereby molecular protocols are first trialed with dedicated end-users to optimize useability and uptake, especially when application by citizen-scientists is anticipated. We believe that the combination of RPA-LF assay and the simplified protocol developed in this study can help minimizing resources and time effort for surveying the invasive Mediterranean fanworm in coastal ecosystems. More importantly, it allows bringing eDNA-based diagnostics tools closer to end-users (environmental managers, biosecurity practitioners or citizen scientists), fast-tracking the implementation of point-of-need tools in biosecurity surveillance programs.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/**Supplementary Material**. Further inquiries can be directed to the corresponding author.

ETHICS STATEMENT

The studies involving human participants were reviewed and approved by Cawthron Institute Ethics board (Approval number: CAW-ETH-210427). The patients/ participants provided their written informed consent toparticipate in this study.

AUTHOR CONTRIBUTIONS

MZ designed and carried out experiments, prepared figures and wrote the manuscript. UA, XP, and AZ provided critical input into experimental design and the manuscript. All authors approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmars.2022.861657/full#supplementary-material

Supplementary Data Sheet 1 | Supplementary tables and figures.

Supplementary Data Sheet 2 | Survey and result sheet.

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