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Assessment of the effects of sulfated polysaccharides extracted from the red seaweed Irish moss *Chondrus crispus* on the immune-stimulant activity in mussels *Mytilus spp*.

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Abstract

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2 Seaweeds contain a number of health enhancing and antimicrobial bioactive compounds 3 including sulfated polysaccharides (SP). In the present study, SP extracted from a European 4 red seaweed Irish moss Chondrus crispus was chemically analyzed, SP content extracted and 5 the immune-response effect on wild Irish mussels *Mytilus* spp. investigated for the first time. A high percent yield of SP was extracted from C. crispus and the immune-stimulant activity 6 7 of SP was assessed in a laboratory trial with mussels exposed to three different treatments of low (10 µg mL⁻¹), medium (20 µg mL⁻¹) and high (50 µg mL⁻¹) SP dose concentrations and a 8 9 control mussel group with no exposure to SP. An initial mussel sample was processed prior to the trial commencing and mussels were subsequently sampled on Days 1, 2, 3, 4, 7, and 10 10 11 post SP exposure. Both cell, humoral and immune related gene responses including 12 haemocyte cell viability, haemocyte counts, lysozyme activity and expression of immune 13 related genes (defensin, mytimycin and lysozyme mRNA) were assessed. No mussel mortalities were observed in either the treated or non-treated groups. Mussels exposed with 14 15 SP showed an increase in haemocyte cell viability and the total number of haemocytes compared to control mussels. Lysozyme activity was also higher in treated mussels. 16 17 Additionally, up-regulated expression of defensin, mytimycin and lysozyme mRNA was 18 observed in SP treated mussels shortly after exposure (on Days 1, 2, and 3) to SP. These 19 results indicate that a high quality yield of SP can be readily extracted from C. crispus and 20 more importantly based on the animal model used in this study, SP extracted from C. crispus 21 can rapidly induce health enhancing activities in Mytilus spp. at a cellular, humoral and 22 molecular level and with a prolonged effect up to ten days post treatment.

Keyword: Chondrus crispus; immune responses; Mytilus spp; sulfated polysaccharides

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1. Introduction

Marine natural products have received much attention in recent years due to their health benefits and bioactivity. More than 16,000 natural products have been isolated from marine organisms including proteins, amino acids, lipid, fibers, pigments and polysaccharides, all of which have the potential to modulate the immune system [1,2]. In particular, seaweeds are recognized as a rich source of diverse bioactive compounds such as sulfated polysaccharides (SP). Marine seaweeds contain many different polysaccharides related to their taxonomic classifications [3]. It is acknowledged that SP from seaweed groups

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include carrageenan and agaran from red seaweeds, ulvan from green seaweeds and fucoidan from brown seaweeds [4,5]. SP present in the extracellular matrix of seaweeds [6] have been increasingly reported to induce many health enhancing biological activities and are considered be antioxidant, anticancer, anti-inflammatory, to antiviral, and immunomodulatory [7]. The immunostimulatory effect induced by SP derived from marine seaweeds has been widely studied in vertebrate and other invertebrate animal models [4,8]. In other invertebrate animals, laminaran and \(\beta\)-glucan from the brown macroalgae Laminaria hyperborea showed immunomodulatory effects on Atlantic salmon Salmo salar macrophages [9]. Alginate extracted from seaweed increased the proportion of neutrophils, phagocytic activity, respiratory burst and expression of interleukins in rainbow trout Oncorhynchus mykiss post injection [10]. Administration of fucoidans from the brown macroalgae Sargassum wightii by immersion in Pacific white shrimp Litopenaeus vannamei culture was demonstrated to increase haemocyte proliferation and the mitotic index of haematopoietic tissue [11]. Oral administration of carrageenan, a red seaweed polysaccharide from Gigartina sp., supplemented diets has been reported to increase the immune-related expression in L. vannamei [12]. Additionally, treatment of alginate and fucoidans from the brown seaweed, Macrocystis pyrifera and Fucus vesiculosus enhanced the immune response activity in head kidney leukocytes of cod, Gadus morhua [13].

Chondrus crispus, a red seaweed from the family Gigartinaceae, is found abundantly on the Atlantic coasts of Europe and North America [14]. It has been recognized that C. crispus is a source of SP since the 1960s [15] and besides having a relatively high SP content, C. crispus is also rich in proteins, amino acids, lipids and pigments [16]. The biochemical composition of SP from seaweeds is dependent on the species, anatomical regions, growing conditions, extraction procedures and analytical methods [4]. Protein content in seaweeds has not been studied as well as polysaccharide content but it is acknowledged that seaweed protein content can be species dependent with protein content in green and red seaweeds being higher (up to 35-44% of dry mass) compared to brown seaweeds (less than 5%) [16,17]. In addition to protein and polysaccharide content, polyphenols such as phenolic compounds (2.5%) and flavonoids (0.1%), are also produced by seaweeds [18]. In a recent study, water-soluble polysaccharides from C. crispus were observed to enhance the immune response in the free-living nematode Caenorhabditis elegans and to suppress the expression of quorum sensing and the virulence factors of the gram-negative bacteria Pseudomonas aeruginosa [19]. In that study, it was suggested that the water-soluble polysaccharides derived from C. crispus may play a health-promoting role in animals and humans. A growing

interest in the biomedical prospects of seaweed-derived SP in human health care has emphasized the need for strategies to maximize SP extraction, bioavailability and investigate precisely the therapeutic mechanisms of SP.

Mussels *Mytilus spp.* belonging to the Mytillidae family are important aquatic animals that are harvested for human consumption worldwide [20] and also play an important ecological role in the marine environment, forming biogenic reefs [21]. The blue mussel Mytilus edulis are boreo-temperate in their distribution on both coasts of the Atlantic Ocean in Europe and North America and are found in abundance, intertidally and subtidally, in both sheltered and exposed sites, attached to hard substrates [22,23]. The Mediterranean mussel Mytilus galloprovincialis is endemic to the Mediterranean, Black Sea and Adriatic Sea and has expanded its range to the British Isles [24]. Evidence of hybridisation and hybrid zones of M. edulis and M. galloprovincialis in the south west of England and Ireland were first recorded in the 1970s and subsequent studies have further documented this phenomenon [25,26,27]. The immune response of Mytilus spp. like other invertebrates includes a cellular response and soluble haemolymph factors including lysozyme, an antibacterial enzyme, and other antimicrobial peptides (AMPs) including defensins and mytimycins [28,29]. Haemocytes play an important role in invertebrate cellular immune response carrying out phagocytosis, melanization, encapsulation and cell-to-cell communication and indirect humoral immune response [30]. Several studies have shown that lysozymes are able to kill Gram-negative bacteria, which has been demonstrated in bivalves [31,32].

The objectives of the present study were (a) to extract, quantify and chemically analyze the SP derived from *C. crispus* and (b) assess the immune effects of exposure to the extracted SP using mussels *Mytilus spp*. as the animal model for the first time. Findings from this study will contribute to a better understanding of naturally derived biotherapeutics and their contribution not only to animal health and well-being, in particular in species involved in the aquaculture industry, but possibly to vertebrate health including humans.

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2. Materials and methods

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2.1. Cold water extract of sulfated polysaccharides from Chondrus crispus

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Red seaweed Irish moss *C. crispus* (50 g) was collected in November 2015 from Inch Strand, Midleton Co. Cork, Ireland (51.79508, -8.180008). Seaweed samples were freshly

harvested, returned to the laboratory and sand and epiphytes were removed. The *C. crispus* was dried in an oven (35-40 °C) for 20-24 h. SP was extracted as previously described by Wongprasert et al. (2014) [33]. Briefly, the dry seaweed was ground and de-pigmented with benzene (24 h) and acetone (24 h). Five grams of de-pigmented seaweed powder was stirred at 35-40 °C in 500 mL distilled water (DW) for 4 h. The extract was diluted with 500 mL of hot water (100 °C) and centrifuged at $6000 \ x \ g$ for 5 min. The seaweed residue was reextracted again by adding 800 mL of DW and stirring for 4 h and its supernatant was filtered through a white cloth (35-48 mesh). The filtrate was allowed to cool to room temperature (RT) and was subsequently kept at -20 °C overnight. The frozen supernatant was thawed and centrifuged at $6000 \ x \ g$ for 5 min to separate gel and non-gel fractions. The gel fraction was discarded and the non-gel fraction SP was collected, freeze-dried and stored at -20 °C for further study.

2.2. Chemical analysis

2.2.1. Sulfate content analysis

The sulfate content of SP was measured using a K_2SO_4 solution as a standard. Briefly, 20 mg of SP was hydrolyzed for 2 h at 100 °C in 0.5 mL of HCl (2N). The SP solution was then transferred and made to volume in a 10 mL volumetric flask. Humic substance was removed from the SP solution by centrifugation ($3000 \ x \ g$, 10 min). The 2 mL of the supernatant was diluted with 18 mL of DW and 2 mL of HCl (0.5 N). Then 1 mL of BaCl₂-gelatin reagent was added, swirled and retained for 30 min at RT. The mixture solution was measured at 550 nm using a spectrophotometer and the percentage of sulfate was calculated with reference to the standard curve given by a K_2SO_4 solution of known concentrations in the ranges of 10-50 μg sulfate mL⁻¹.

2.2.2. Total polysaccharide analysis

Total polysaccharide content of SP was estimated by phenol-sulfuric acid method using a galactose solution as a standard. One mL of SP solution (1 mg mL⁻¹) was mixed with 1 mL of 5% phenol in DW and 5 mL of sulfuric acid. The mixture was vortexed and allowed to stand for 10 min at RT. The mixture solution was then cooled in an ice bath (15-20 °C) for 15 min and its absorbance was measured at 490 nm using a spectrophotometer. The

percentage of polysaccharide in SP was calculated with reference to the standard curve given by a galactose solution of known concentrations in the ranges of 10-2000 µg mL⁻¹.

2.2.3. Total protein analysis

Total protein content of SP was measured by BCA protein assay [34]. An aliquot (25 μ L) of SP or standard BSA solution (final concentrations at 0-2000 μ g mL⁻¹) was added to 200 μ L by BCA solution (BCATM protein assay kit, Thermo scientific, Inc., USA), mixed and incubated at 37 °C for 30 min. After incubation, the solution was measured the absorbance at 562 nm using a spectrophotometer. The total protein content of SP was calculated with reference to the standard curve given by a BSA solution of known concentrations.

2.2.4. Total phenolic analysis

The total phenolic content of SP was determined by using the Folin-Ciocalteu assay [35]. An aliquot (25 μ L) of SP or standard solution of gallic acid (final concentrations at 0-100 mg L⁻¹) was added to 250 μ L of DW. A reagent blank was DW. Then, 25 μ L of Folin-Ciocalteu's phenol reagent was added to the mixture and shaken. After 5 min, 250 μ L of 7 % Na₂CO₃ solution was added to the mixture. The solution was diluted to 625 μ L with DW and mixed by vortex. After incubation for 90 min at RT, the absorbance was measured at 750 nm using a spectrophotometer. Total phenolic content of SP was expressed as mg gallic acid equivalents (GAE) per 100 g of SP.

2.2.5. Total flavonoid analysis

Total flavonoid content of SP was measured by the aluminum chloride colorimetric assay [36] and modified as described by Marinova et al. (2005) [37]. An aliquot (65 μ L) of SP or standard solution of catechin (final concentrations at 0-100 mg L⁻¹) was added to 325 μ L of DW and mixed. Then, 19.5 μ L of 5% NaNO₂ was added. After 5 min, 19.5 μ L of 10% AlCl₃ was added. At 6 min, 130 μ L of 1 M NaOH was added and the total volume was made up to 650 μ L with DW. The solution was mixed and shaken and the absorbance measured at 510 nm using a spectrophotometer. Total flavonoid content of SP was expressed as mg catechin equivalents (CE) per 100 g of SP.

168 2.3. Immune-stimulant activity of SP derived from C. crispus in the mussel Mytilus spp.

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170 *2.3.1. Experimental design*

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Wild mussels Mytilus spp. (n=148) were randomly sampled from rocks at the intertidal at Ringaskiddy, Cork Harbour, Co. Cork (51°50'N, 8°19'W), a known hybrid zone for M. edulis, M. galloprovincialis and hybrids of both parent species, and a Class C water quality site influenced greatly by anthropogenic effects such as agricultural and industrial run-off, leachate from landfills and untreated waste water and sewage [27]. A total of 136 mussels were arbitrarily divided into 4 groups, each with duplicates of 17 mussels in standalone rectangular plastic aquaria containing 8 L of artificial seawater (ASW) at a salinity of 34 and at 12 °C (8 tanks in total) and aerated using an air stone. The photoperiod was 12/12 h of day/night cycles. The four groups consisted of Group 1, control mussels, not exposed to SP and Groups 2, 3 and 4 consisting of mussels exposed to SP at final concentrations of 10, 20 and 50 µg mL⁻¹, respectively. Water changes were not carried out for the trial duration to ensure that the SP dose amount added to the water at the beginning of the trial was not altered and/or removed from each system. The trial was carried out in a constant temperature room and with a constant photoperiod. Holding conditions were optimal for the mussels as no mortalities were observed in either the control or treated mussels. Mussels were fed with 2 mL of Shellfish Diet 1800 (Reed Mariculture) consisting of a heteromorphic mix of phytoplankton at a total concentration of 2 x 10⁹ cells mL⁻¹ containing *Isochrysis sp*, *Pavlova* sp, Thalossiosira weissflogii, and Tetraselmis sp at day 4 of the experiment. At days 1, 2, 3, 4, 7 and 10 of the trial, 4 mussels were arbitrarily selected from each treatment (n = 2) replicate/ treatment) for immunological analyses including haemocyte viability, total haemocyte counts, lysozyme activity assay and immune related mRNA expression. All assays were performed in triplicate.

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2.3.2. Cell viability in haemolymph of mussels Mytilus spp.

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The neutral red retention time (NRRT) assay used to determine the haemocyte viability in mussels as previously described [38] was carried out. Briefly, haemolymph (750 μ L) was collected from individual mussels using a 20-gauge sterile needle (Microlance) fitted on a 1 mL syringe containing 250 μ L of tris-buffered saline solution (TBS). Haemolymph samples were constantly vortexed to resuspend haemolymph and prevent aggregation. One-

hundred microliters of haemolymph sample was then transferred into microplate well, $20 \,\mu\text{L}$ of 2% neutral red solution (Sigma Aldrich, USA) was added and then sample was held in a dark humid chamber for 30 min. The supernatant was subsequently drained by tilting the plates at an angle, facilitating drainage, followed by a gentle rinse with TBS, to remove any excess neutral red dye. The neutral red was extracted by displacing the haemocytes using 100 μ L of extraction solution (1% acetic acid in 50% ethanol) for 30 min. The plates were measured at 450 nm and 570 nm using a UV max spectrophotometer (ELx808IU, Mason Technology, USA).

2.3.3. Total haemocyte counts

Haemolymph sample (200 μ L) was individually collected and fixed with 6% formalin in Alsever's solution (200 μ L) to prevent cell aggregation. After 10 min, 20 μ L portions of the fixed haemocyte suspension were mixed with 20 μ L Rose Bengal solution (1.2% Rose Bengal in 50% ethanol) and incubated at RT for 20 min before being used to determine the total haemocyte counts using a hemocytometer under light microscopy at 20 x magnification (Eclipse 80i microscope, Nikon Instruments Inc., USA). Haemocytometer (improved Newbauer bright line) counts were made for 5/25 squares (volume of 1 square = 0.2 x 0.2 x 0.1 mm³) to calculate total haemocyte count mL⁻¹ of hemolymph (5 x count x 10⁴ x dilution factor) [39].

2.3.4. Lysozyme activity

Lysozyme activity was measured in serum as previously described by Prado-Alvarez et al. (2015) [40]. Haemolymph sample (200 μ L) was individually collected from mussels and mixed with EDTA (200 μ L) to prevent coagulation. Samples were centrifuged at *1200 x g* for 10 min to separate cells from the serum. Serum samples and serial dilution of standard hen egg white lysozyme suspensions (Sigma Aldrich, USA) in 0.1 M phosphate buffer (final concentrations at 0-5 μ g mL⁻¹) (30 μ L) were dispensed in triplicate in 96 well plates before adding a *Micrococcus luteus* (*M. lysodeikticus*) suspension (170 μ L) at 0.2 mg mL⁻¹ in 0.2 M monobasic sodium phosphate and 0.2 M dibasic sodium phosphate (Sigma Aldrich, USA). A set of five measurements of the optical density at 620 nm was recorded every minute using a UV max spectrophotometer (ELx808IU, Mason Technology, USA).

2.3.5. Expression of immune-related mRNA in mussels

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The haemocytes from mussels (4 mussels/ group) was RNA extracted in 200 µL TRI reagents according to the manufacturer's protocol (Sigma Aldrich, USA). The concentration and quality of RNA was determined by measuring the absorbance at 260/280 nm using a NanoDrop 2000 spectrophotometer (Thermo Scientific, USA). cDNA was transcribed from RNA (1 µg) using the Thermo Scientific RevertAid First Strand cDNA Synthesis Kit (Thermo Scientific, USA) containing RevertAid reverse transcriptase (200 U µL⁻¹), RiboLock RNase inhibitor (20 U μL⁻¹), Oligo (dT)18 primer (100 μM), dNTP mix (10 mM), and 5X reaction buffer (250 mM tris-HCl (pH 8.3), 250 mM KCl, 20 mM MgCl₂, 50 mM DTT) at 42 °C for 1 h. The immune related mRNA was amplified by PCR using 1 µL (100 ng) of cDNA with specific primer sets and conditions (Table 1). 18S specific primer was also amplified as an internal control. Twenty-five microliters of PCR reactions contained Phusion DNA polymerase (2 U µL⁻¹), 10 mM dNTP mix, 10 µM of each forward and reverse primers and 5x Phusion HF buffer containing 7.5 mM MgCl₂ (Thermo Scientific, USA). The hot start PCR program used for immune related mRNA was performed with 98 °C for 30 s, followed by 40 cycles of 98 °C for 10 s, annealing temperature for each particular mRNA as shown in table 1, extension at 72 °C for 30 s followed by final extension at 72 °C for 5 min. The RT-PCR product was analyzed by 1.5% agarose gel electrophoresis, stained with 3% ethidium bromide, and visualized under ultraviolet light and documented using the EpiChemi3 darkroom (UVP, Inc., Upland, CA). Expression was semi-quantitatively determined from the ratio of band intensity to the internal control (18S) using ImageJ analysis program (from NIH website by Scion Corporation, Frederick, MD). Each assay was carried out in triplicate.

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2.4. Statistical analysis

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All assays were performed in triplicate. The data were presented as mean \pm SD and analyzed by one-way ANOVA followed by Tukey's multiple comparison and statistically significant difference was required at p-value less than 0.05.

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3. Results

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3.1. Extraction and chemical analyses of SP from C. crispus

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270	SP extracted from the red seaweed C. crispus by cold water extraction was 15.36%
271	yield of dry weight. The chemical analysis showed that SP contained a sulfate content of 9.78
272	\pm 0.42% w/w, a polysaccharide content of 73.94 \pm 20.61% w/w, a total protein content of
273	$7.08 \pm 0.45\%$ w/w, phenolic content of $2.55 \pm 0.24\%$ w/w and flavonoid content of 0.1002 ± 0.000
274	0.003% w/w (Table 2).
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276	3.2. C. crispus SP extract effect on immune parameters in mussel Mytilus spp.
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278	3.2.1. Effect of SP on haemocyte cell viability in Mytilus spp. using the NRRT assay
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280	The results of haemocyte cell viability, analysed using the NRRT assay, are shown in
281	Fig. 1. Haemocyte cell viability of SP treated groups (final concentrations of 10, 20 and 50
282	μg mL ⁻¹) were higher than the control group on Days 1, 2, 3, 4 and 10. While on day 7, the
283	haemocyte cell viability was not different.
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285	3.2.2. Effect of SP on total haemocyte counts (THCs) in Mytilus spp.
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287	The results of THCs from SP treated groups (final concentration of 10, 20 and 50 μg
288	mL ⁻¹) and control group are shown in Fig. 2. THCs were higher in the SP exposed groups
289	than control mussels on Days 1, 2, 3, 4 and 10, with the exception of day 7. Moreover, the
290	results also show a positive correlation between THCs and SP dose concentration, with an
291	increase in THCs with increasing dose concentration. For example, on Day 3, mean THCs of
292	control and SP treated groups at the concentrations of 10, 20 and 50 μg mL ⁻¹ were 0.993 \pm
293	0.24×10^6 , $1.262 \pm 0.36 \times 10^6$, $1.915 \pm 0.15 \times 10^6$ and $1.724 \pm 0.31 \times 10^6$ cells mL ⁻¹ ,
294	respectively. On Day 10, THCs of control and SP treated groups at the concentrations of 10,
295	20 and 50 μg mL ⁻¹ were 1.593 \pm 0.47 x 10 ⁶ , 1.599 \pm 0.90 x 10 ⁶ , 1.864 \pm 0.32 x 10 ⁶ and 2.263
296	$\pm 0.67 \times 10^6$ cells mL ⁻¹ , respectively.
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298	3.2.3. Effect of SP on lysozyme activity in Mytilus spp.
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300	Lysozyme activity of mussels treated with SP from C. crispus was evaluated by
301	measuring the lysozyme activity in serum. Compared to control, lysozyme activity in the
302	serum of mussels treated with SP significantly increased on Days 1 and 2 (Fig. 3). On Day 3,
303	lysozyme activity in the serum of mussels treated with SP declined to control levels. On Days

4 and 7, lysozyme activity increased in treated and control groups from Day 3 but this increase was not significant in both the control and treated mussels. On Day 10, lysozyme activity in each groups had decreased from Days 4 and 7.

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3.3. The effects of SP stimulation on immune related genes expression in Mytilus spp.

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Post treatment with SP (final concentrations of 10, 20 and 50 µg mL⁻¹), the relative expression of defensin, mytimycin and lysozyme with 18S mRNA was evaluated in treated and control mussels on Days 1, 2, 3, 4, 7 and 10. The levels of defensin mRNA expression in mussels from the SP treated groups was significantly higher than that of mussels in the control group as shown in Fig. 4A and B. On Days 7 and 10, the levels of defensin mRNA expression showed higher than that of the control group but it was not significant. For mytimycin mRNA expression, on Days 1 and 2, the levels of mytimycin mRNA expression in mussels in the SP treated groups were significantly higher than that of mussels in the control group. On Day 3, the levels of mytimycin mRNA expression of SP treated at 50 µg mL⁻¹ remained significantly high from the control group. Whereas, on Days 4, 7 and 10, the levels of mytimycin mRNA in mussels treated with the three SP concentrations showed no difference to the mussels from the control group (Fig. 4A and C). For lysozyme mRNA expression, on Days 1 and 2, the levels of lysozyme mRNA expression in mussels, from the treated groups, was significantly higher than that of mussels in the control group. On Day 3, the levels of lysozyme mRNA in mussels treated with the three SP concentrations showed no difference to the mussels from the control group. An increase of lysozyme mRNA expression in SP treated groups was observed again on Day 4. A decrease in lysozyme mRNA expression was observed from Day 7 onwards and this was relative to SP dose concentration. For example, on Day 7, the levels of lysozyme/18S mRNA expression in SP treated mussels at concentrations of 10, 20 and 50 µg mL⁻¹ were 0.67, 1.1 and 2.0 folds of control, respectively and on Day 10, it was 1.6, 3.1 and 3.1 folds of control, respectively (Fig. 4A and D).

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4. Discussion and Conclusion

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In the present study, a high quality yield of SP from the red seaweed *C. crispus* was extracted successfully by cold-water extraction following the methodology previously reported for the red seaweed *G. fisheri* in Wongprasert et al. (2014) [33]. The yield of SP

from *C. crispus* (15.36%) was high and similar to other seaweeds, which have yielded a range of 14.2-19.7%, using cold water extraction of the red seaweed *G. corticata* (2.8-19.7%) [41] and the green seaweed *U. pertusa* (14.2%) [42]. Chemical analysis in this study indicated that the SP in *C. crispus* contained high levels of carbohydrate including sulfate ester and low levels of protein, which is comparable with other red seaweeds (*Porphyra*, *Palmaria*, *Gracilaria*) [16,41]. It has been demonstrated that different thermal extractions obtain different levels of polysaccharide and sulfate contents [41], with the cold-water extraction method yielding higher levels of sulfate ester compared to the hot water extraction method, as was also observed in this study.

The immune response in Mytilus spp. was observed in the present study to be enhanced at a cellular, humoral and molecular level shortly after exposure to C. crispus SP (on day 1). Under optimal conditions, bivalves such as Mytilus spp. will filter seawater at a maximum rate [43] to facilitate feeding and respiration. Valve opening and continuous filtering activity was observed in the mussels in this study thus readily exposing the treated mussels to the SP dissolved in the tank seawater. Haemocyte cell viability and total haemocyte counts were increased with SP treated mussel groups compared to the nontreated mussel group. The immune system of bivalves lacks immune specificity and memory, however, innate defense mechanisms that can identify and protect against non-self-material have evolved, with invertebrate immune response centered largely on the multifunctional haemocytes [44]. Early effects of physiological alterations are often seen as changes in haemocyte counts with elevated cell counts a common response to environmental stress [45] or as observed in this study exposure to an immunostimulant biocompound. The rapid SP induced increase in haemocyte cell count and cell viability, observed in the treated mussels in this study, highlights the effective delivery mechanism of SP to the mussels via their filter feeding activity.

Similar findings including haemocyte proliferation and an increase in the mitotic index of haematopoietic tissue in response to SP exposure have been observed in crustaceans and fish species [10,11]. However, SP showed a differentially distinctive increase in percentage of total haemocyte counts, for instance *Mytilus spp.* exposed to SP from *C. crispus* and rainbow trout injected with alginate from *L. digitata* showed a similar increase in number of haemocyte. While shrimp immersion with fucodains from *S. wightii* showed increased percentage of total haemocyte counts in a lesser degree than that of mussel and fish at the same concentration and exposure time.

In the present study, lysozyme activity was measured in the serum of SP-treated compared with SP-nontreated mussels, and it was observed that lysozyme activity increased shortly (on Days 1 and 2) after exposure to SP and the level of lysozyme mRNA expression was also up-regulated on Days 1 and 2 of the trial. Previous studies reported that marine microalgae, *Chaetoceros calcitrans* (*C. calcitrans*) and *Tetraselmis suecica* (*T. suecica*), presented in Shellfish Diet 1800, have been reported to evoke an immune response in bivalves [46]. It was reported that *C. calcitrans* had a positive effect on total haemocyte count, granulocyte percentage, phagocytic rate and oxidative activity of clam haemocytes. Moreover, *T. suecica* had a positive effect on the phagocytic rate of oyster haemocytes. Interestingly, our study demonstrated that on Day 4 when the SP exposed mussels were fed the heteromorphic commercial Shellfish Diet 1800 (containing microalgae), lysozyme activity and lysozyme mRNA expression increased more once again than the nontreated control mussels, suggesting the SP possibly enhanced increased lysozyme activity after Shellfish Diet 1800 feeding.

Antimicrobial peptides such as defensins, mytilins, myticins and mytimycin, have been identified in mussel species (*M. galloprovincialis* and *Bathymodiolus azoricus*) and are associated with a humoral immune response [28]. Consistent with lysozyme mRNA, defensin and mytimycin mRNA expressions were up-regulated shortly (on Days 1, 2 and 3) after SP treatment in this study. It was reported that polysaccharides, sulfated galactans from the red seaweed *Gracilaria fisheri* stimulated immune AMPs-mRNA expressions in *L. vannamei* shrimp haemocyte [47], however the response in *Mytilus spp.* is much more rapid following exposure to SP.

In conclusion, the present study demonstrated that SP was efficiently extracted from *C. crispus* using the cold-water extraction methodology and this extracted SP has a rapid immune-stimulant effect in mussel *Mytilus spp*. This prompt immune response on a cellular, humoral and molecular level, observed in *Mytilis spp*. and instigated by SP exposure, may be beneficial to animals in assisting them in overcoming the challenges of parasites and disease. More importantly the short-term energy output required for such a rapid immune response may not be too costly to individuals whose health maybe already be compromised due to stress and poor health.

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407	the study.	
408		
409	Legends	
410		
411	Table 1 Specific primers and conditions used for the determination of immune-related	
412	mRNA expression in mussels Mytilus spp.	
413		
414	Table 2 Contents of sulfated polysaccharides (SP) from the red seaweed C. crispus. SP	
415	showed sulfate, polysaccharide, protein, phenolic and flavonoid contents. Data are presented	
416	as a mean of triplicate independent experiments.	
417		
418	Fig. 1. Haemocyte cell viability of mussels Mytilus spp. exposed with SP from the red	
419	seaweed C. crispus at the concentrations of 10, 20 and 50 µg mL ⁻¹ determined by neutral red	
420	retention time (NRPT) assay. The bar graphs show the haemocyte cell viability presented as	
421	the percent of control (mean \pm standard deviation of triplicate independent experiments). *	
422	indicates value significantly different from the control (P $<$ 0.05), n = 4.	
423		
424	Fig. 2. Total haemocyte counts of mussels Mytilus spp. exposed with SP from the red	
425	seaweed C. crispus at the concentrations of 10, 20 and 50 µg mL ⁻¹ . Data is presented as a	
426	mean ± standard deviation of triplicate independent experiments. * indicates value	
427	significantly different from the control (P $<$ 0.05), n = 4.	
428		
429		
430	Fig. 3. Lysozyme activity of mussels <i>Mytilus spp.</i> exposed with SP from the red seaweed <i>C</i> .	
431	crispus at the concentrations of 10, 20 and 50 μg mL ⁻¹ . Data is presented as a mean \pm	
432	standard deviation of triplicate independent experiments. * indicates value significantly	
433	different from the control (P $<$ 0.05), n = 4.	
434		
435	Fig. 4. Expression levels of immune related mRNA of mussels Mytilus spp. in the initial	
436	samples, control and SP from the red seaweed C. crispus at the concentrations of 10, 20 and	
437	50 μg mL ⁻¹ . (A) RT-PCR analysis of defensin, mytimycin and lysozyme mRNA expression	
438	by 1.5% agarose gels. (B) Densitometry value of defensin mRNA relative to 18S in different	

- groups, (C) densitometry value of mytimycin mRNA relative to 18S in different groups and
- 440 (D) densitometry value of lysozyme mRNA relative to 18S in different groups. Bars indicate
- 441 mean ± standard deviation. * indicates value significantly different from control group
- 442 (P<0.05), n = 4.

443

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Primers Primer sequences (5' to 3')		References	Annealing temp
Lysozyme-F Lysozyme-R	ATGTGGAATCTGAAGGACTTGT CCAGTATCCAATGGTGTTAGGG	[48]	60 °C for 30 s
Defensin-F Defensin-R	GTGGCGTCTGCTGGGTTT GAATGGACTTACAATGTCGATGACA	[49]	58 °C for 30 s
Mytimycin-F Mytimycin-R			58 °C for 30 s
18S-F 18S-R	TTACGTCGGCGCAACTTCT CTGTTCCAAGGACTTTAATG	[50]	57.6 °C for 30 s

Table 1

Contents	% of SP extract (w/w)	
Yield	15.36	
Sulfate	9.78 ± 0.424	
Polysaccharide	73.94 ± 20.61	
Protein	7.08 ± 0.455	
Phenolic	2.55 ± 0.236	
Flavonoid	0.10 ± 0.003	

Table 2

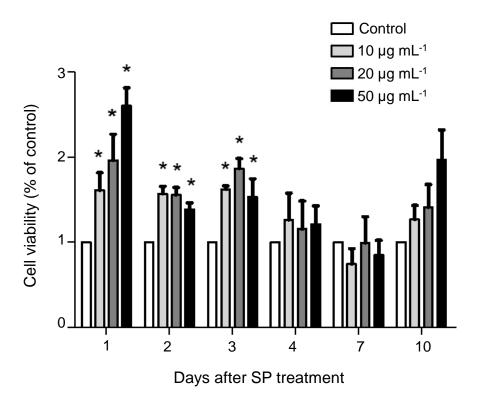


Figure 1

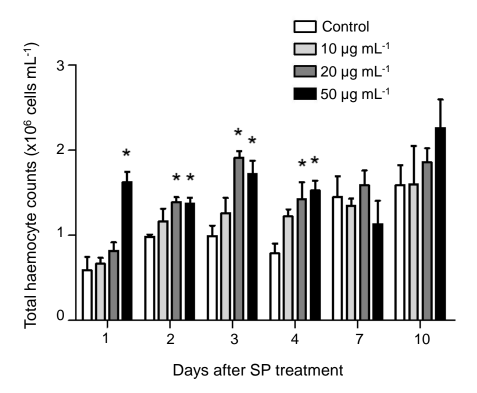


Figure 2

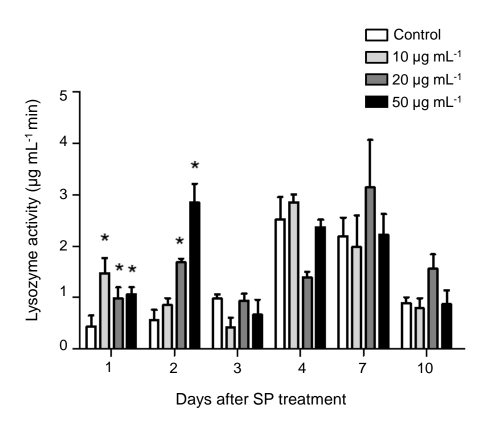


Figure 3

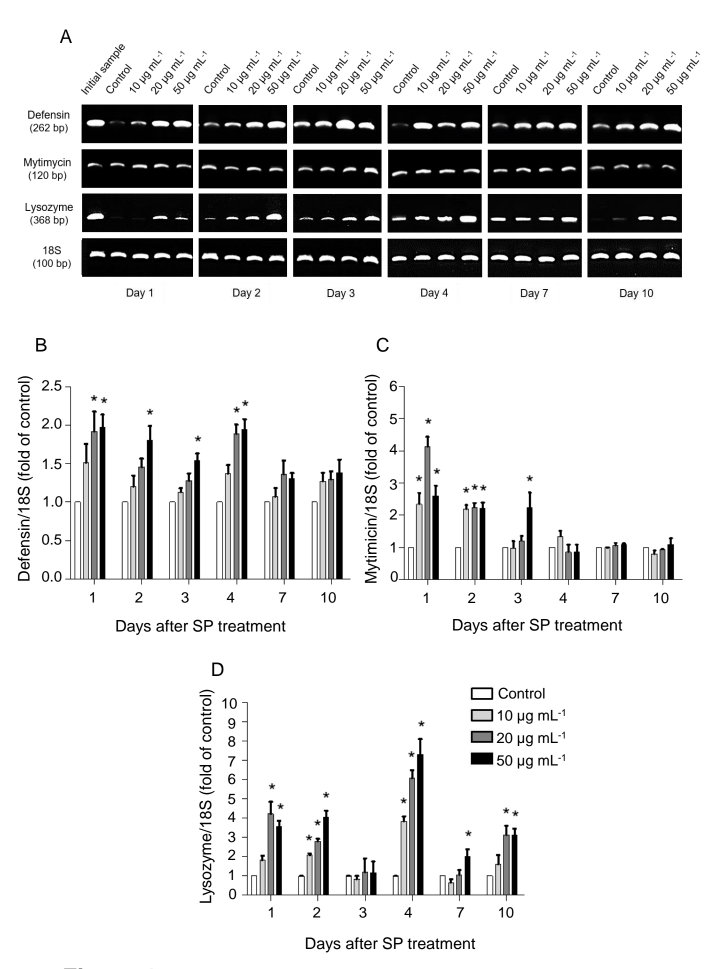


Figure 4

Highlights

- Sulfated polysaccharides (SP) were extracted from Irish moss Chondrus crispus
- SP enhanced immune parameters in mussels *Mytilus spp*.
- SP up-regulated expression of immune genes in mussels shortly after exposure