

# HOKKAIDO UNIVERSITY

Title	Antioxidant and fibroblast-activating activities of the by-product of skate chondroitin extractive production
Author(s)	Li, Wen; Terauchi, Naoya; Meng, Dawei; Miyamoto, Nobuyuki; Tsutsumi, Naonobu; Ura, Kazuhiro; Takagi, Yasuaki
Citation	Sustainable Chemistry and Pharmacy, 23, 100499 https://doi.org/10.1016/j.scp.2021.100499
Issue Date	2021-10
Doc URL	http://hdl.handle.net/2115/90463
Rights	© 2021. This manuscript version is made available under the CC-BY-NC-ND 4.0 license https://creativecommons.org/licenses/by-nc-nd/4.0/
Rights(URL)	https://creativecommons.org/licenses/by-nc-nd/4.0/
Туре	article (author version)
File Information	manuscript received 2021-08-27.pdf



## Highlights

- Skate chondroitin sulfate (sCS) production generated a by-product (BP-sCS)
- BP-sCS is a mixture of CS, type II collagen peptides, and non-collagenous peptides
- BP-sCS displayed antioxidant activity and protected fibroblasts from oxidative stress
- BP-sCS promoted fibroblast proliferation and activated collagen deposition
- BP-sCS could be a superior healing promoter of chronic wounds



1	Antioxidant and fibroblast-activating activities of the by-product of skate chondroitin
2	extractive production
3	
4	Wen Li <sup>a*</sup> , Naoya Terauchi <sup>a</sup> , Dawei Meng <sup>a,b</sup> , Nobuyuki Miyamoto <sup>c</sup> , Naonobu Tsutsumi <sup>d</sup> ,
5	Kazuhiro Ura <sup>e</sup> , Yasuaki Takagi <sup>e</sup>
6	
7	<sup>a</sup> Graduate School of Fisheries Sciences, Hokkaido University, 3-1-1 Minato-Cho, Hakodate,
8	Hokkaido 041-8611, Japan
9	<sup>b</sup> Zhejiang Province Joint Key Laboratory of Aquatic Products Processing, Institute of
10	Seafood, Zhejiang Gongshang University, Hangzhou, China
11	° Marukyo Bio Foods, 4-18-18 Chuo, Wakkanai, Hokkaido 097-0022, Japan
12	<sup>d</sup> Sapporo Fine Chemical Research Center, Marukyo Bio Foods, Nishimiyanosawa 4-2-1-40,
13	Teine, Sapporo, Hokkaido 006-0004, Japan
14	<sup>e</sup> Faculty of Fisheries Sciences, Hokkaido University, 3-1-1 Minato-Cho, Hakodate,
15	Hokkaido 041-8611, Japan
16	
17	E-mail: fenzhongbahe@outlook.com (W. Li), naoya19950728708@eis.hokudai.ac.jp (N.
18	Terauchi), dawei@mail.zjgsu.edu.cn (D. Meng), miyamoto@mbf-net.com (N. Miyamoto),
19	naonobu_tsutsumi@mbf-net.com (N. Tsutsumi), kazu@fish.hokudai.ac.jp (K. Ura), and
20	takagi@fish.hokudai.ac.jp (Y. Takagi)
21	

22 \*Corresponding author:

- 23 Wen Li, Graduate School of Fisheries Sciences, Hokkaido University, 3-1-1 Minato-Cho,
- 24 Hakodate, Hokkaido 041-8611, Japan
- 25 Email: fenzhongbahe@outlook.com
- 26 Tel./fax: +81 138 40 5551
- 27
- 28
- 29

30 Abstract

31	Owing to the increasing popularity of chondroitin sulfate (CS) for joint pain treatment,
32	the CS-production industry has been producing an increasing amount of waste, which
33	includes type II collagen, non-collagenous proteins, and residual CS. To effectively utilize
34	these resources, we intended to develop new products from the by-product of skate
35	chondroitin sulfate production (BP-sCS). In this study, we examined the antioxidant and
36	fibroblast-activating properties of BP-sCS, intending to apply it for a wound-healing
37	promoter. BP-sCS exhibited ABTS and DPPH radical scavenging activities, protected L929
38	fibroblasts from H <sub>2</sub> O <sub>2</sub> - or AAPH-induced oxidative stress, and scavenged intracellular
39	reactive oxygen species. Moreover, BP-sCS promoted L929 fibroblast
40	proliferation/metabolism and stimulated collagen deposition into the extracellular matrix. In
41	addition, BP-sCS counteracted AAPH-induced oxidative stress damage that inhibited
42	fibroblast migration. These effect were attributed to the cooperation among the molecules of
43	BP-sCS, namely, type II collagen peptides, non-collagenous peptides, and CS
44	polysaccharides. Our findings indicate that BP-sCS has the potential as a novel wound-
45	healing promoter. This study is the first step toward the realization of a sustainable CS-
46	production industry by waste utilization in healthcare products.
47	Keywords: by-product, wound-healing promoter, type II collagen peptide, chondroitin sulfate
48	polysaccharide
49	
50	Abbreviations: AAPH, 2,2'-azobis(2-methylpropionamidine) dihydrochloride; ABTS, 2,2'-

51 azino-bis(3-ethylbenzothiazoline-6-sulfonic acid; BP-sCS, the by-product of skate

52	chondroitin sulfate production; CP, collagen peptide; CS, chondroitin sulfate; DCFH,
53	dichlorofluorescein; DPPH, 2,2-diphenyl-1-picrylhydrazyl; Pa-CP, papain-hydrolyzed CP;
54	CSp, CS polysaccharides; MW, molecular weight; ROS, reactive oxygen species

## **1. Introduction**

57	Chondroitin sulfate (CS) is a polysaccharide chain consisting of a repeated disaccharide
58	unit, which comprises glucuronic acid and N-acetylgalactosamine. CS is used as a
59	symptomatic slow-acting drug in Europe and a dietary supplement in the United States
60	(Volpi, 2007) for joint pain treatment. Currently, CS is extracted and purified from animal
61	cartilaginous tissues. Although microbial production, as in the case of hyaluronic acid, is an
62	attractive technology in the future (Restaino et al., 2017), it is still a challenge (Schiraldi et
63	al., 2012). Thus, industrial CS production uses cartilage sources derived from terrestrial or
64	marine animals; however, a large amount of the cartilage residue, including type II collagen,
65	non-collagenous proteins, and residual CS, is wasted during CS extraction and purification.
66	Due to the aging population and the discovery of new CS bioactivities, such as anti-obesity
67	(Li et al., 2019) and antiviral (Vázquez et al., 2013) activities and intestinal microbiota
68	modulation (Shang et al., 2016), the demand for CS production has rapidly increased
69	(Restaino et al., 2019; Vázquez et al., 2013). This, in turn, rapidly increased the waste
70	generated from CS production. To realize sustainable development and maximize profit, the
71	generated waste must be utilized. However, the utilization of the CS production waste has not
72	been demonstrated to date.
73	Generally, cartilage proteins are hydrolyzed during CS purification; therefore, the by-

product of CS production contains a mixture of molecules, such as CS polysaccharide
residues, type II collagen peptides, and non-collagenous peptides, alongside a small number
of minerals that bind to CS. Since CS polysaccharides and peptides with a high molecular
weight (MW) exhibit low absorbability in the skin and digestive system (Li et al., 2015;

78	Shang et al., 2016; Shen & Mastsui, 2017), we hypothesized that the by-product could be
79	used for wound healing as they can directly get in contact with the wound site. CS has been
80	reported to activate fibroblasts and their migration during wound healing (Zou et al., 2009),
81	while peptides, especially collagen peptides, exhibit several biological activities, such as
82	antioxidant activity, antimicrobial activity, and extracellular matrix synthesis activation (Pal
83	& Suresh, 2016). Consequently, it is possible that the by-product of CS production, which is
84	the waste now, may promote wound healing.
85	Wound healing is a complex and highly regulated process divided into three phases:
86	hemostasis and inflammation, proliferation, and remodeling (Broughton et al., 2006; Han &
87	Ceilley, 2017). Failure to progress through these normal stages of healing results in chronic
88	wounds, such as diabetic wounds and pressure ulcers (Dhivya et al., 2015; Han & Ceilley,
89	2017). An essential feature of these wounds is oxidative stress, which is caused by
90	inflammatory cells in wound tissues that produce large amounts of reactive oxygen species
91	(ROS), pro-inflammatory cytokines, and oxidases (Kurahashi & Fujii, 2015; Schäfer &
92	Werner, 2008). Although low ROS levels are required for defense against invading pathogens
93	and modulating signaling molecules, excessive ROS levels cause cellular apoptosis in the
94	surrounding tissues and imbalanced redox homeostasis, resulting in chronic wounds
95	(Kurahashi & Fujii, 2015; Schäfer & Werner, 2008). Chronic wounds are an important global
96	health problem that causes significant discomfort and distress to patients. Even in the
97	developed countries, almost 1.5% of the population experiences problematic wounds,
98	accounting for 2%–4% of all healthcare expenses (Ahmed et al., 2019). Thus, the market
99	demands superior wound-healing promoters that can achieve rapid healing at a reasonable

100 cost.

101	In northern Hokkaido, Japan, skate fishery is one of the essential industries. From the
102	newest governmental data of the aquatic product market, the total skate catch of Hokkaido is
103	1798 tons in 2018. As only their fins (about 33% of wet weight, unpublished observation) are
104	used as foods, a massive volume of cartilaginous by-products is generated from skate
105	processing. Our preliminary survey revealed that 1 ton of CS is produced per year from 270
106	tons of by-products obtained from 400 tons of skate. Thus, still, a considerable number of by-
107	products are being discarded even after CS production. Transforming this waste into value-
108	added products can mitigate environmental pollution, improve the economic profit of fishery
109	and aquaculture industries, and contribute to the circular economy in Hokkaido.
110	This study obtained a by-product of the final stage of skate CS production (BP-sCS) and
111	examined its bioactivities as a wound-healing promoter. During the proliferative phase of
112	wound healing, fibroblasts migrate to the wound site from surrounding tissues, become
113	activated, proliferate, synthesize extracellular matrix, and finally differentiate into
114	myofibroblasts to close the wound (Broughton et al., 2006). At the same time, excessive ROS
115	must be detoxified during chronic wound healing. Therefore, we focused on the antioxidant
116	and fibroblast-activating roles of BP-sCS. In addition, we examined the activities of pure CS
117	polysaccharides and type II collagen peptides to determine the active components of BP-sCS.
118	This study is the first step toward a sustainable CS production industry as our results indicate
119	that CS production wastes can be used in various healthcare products.

## 120 2. Materials and methods

122	Skate cartilage, chondroitin sulfate polysaccharide (CSp), and BP-sCS were supplied by
123	Marukyo Bio Foods (Wakkanai, Hokkaido, Japan). CSp was isolated and purified from the
124	cartilage of Raja pulchra (skate), and the remaining material was spray-dried to produce BP-
125	sCS containing CSp (14%), type II collagen peptides (60%), non-collagenous peptides (14%),
126	minerals (8%), and other materials (Supplementary Fig. 1, Japan Food Analysis Center
127	no.17091776001-0101). CSp was obtained as a CS sodium salt, which includes CS (83.1%)
128	and sodium (16.9%). Thus, the content of CSp in BP-sCS as CS sodium salt was 22%. Type
129	II collagen was purified from skate cartilage largely according to the method described by
130	Meng et al. (2019). The papain-hydrolyzed collagen peptides (Pa-CP) were prepared by
131	hydrolyzing the purified type II collagen using papain (2.5% w/w) at 50 $^\circ$ C for 4 h. The
132	formulations and molecular weight of CSp are presented in Supplementary Table 1. No
133	keratan sulfate existed in the cartilage glycosaminoglycan of most skate species (Murado et
134	al., 2010).

#### 135 *2.2. Antioxidant assays*

136 *2.2.1. ABTS radical scavenging assay* 

ABTS (7 mM; Wako Pure Chemical, Osaka, Japan) and potassium persulfate ( $K_2S_2O_8$ ,

138 2.45 mM; Wako Pure Chemical) solutions (2:1 v/v) were reacted in the dark at room

temperature (21 °C–23 °C) for 12–16 h to generate ABTS radical solution. The solution was

diluted with phosphate-buffered saline (PBS, pH 7.4) to an absorbance of  $0.70 \pm 0.02$  at 734

nm, reacted (500 µL) with 500 µL of sample solution for 10 min at room temperature (21 °C-141 23 °C), and the absorbance determined at 734 nm using a microplate reader (Infinite F50R, 142 Tecan Japan, Kanagawa, Japan). Each sample was dissolved in deionized water to 143 concentrations of 0.04–5 mg/mL. L (+)-ascorbic acid (VC, Wako Pure Chemical) was used as 144 the positive control. Each measurement was performed in triplicate, and the percentage of the 145 scavenging effect was calculated as follows: 146 ABTS radical scavenging activity (%) =  $(A_b - A_s)/A_b \times 100$ , 147 where  $A_b$  and  $A_s$  denote the absorbance of deionized water and sample solution, respectively. 148 2.2.2. DPPH radical scavenging assay 149

DPPH (Tokyo Chemical Industry, Tokyo, Japan) was dissolved in methanol at a 150 concentration of 115 µM. Each sample was dissolved in deionized water at concentrations of 151 0.625–5 mg/mL. VC was used as a positive control. Next, the sample solution (500  $\mu$ L) was 152 reacted with DPPH methanol solution (500 µL) in the dark at room temperature (21 °C-23 153 °C) for 30 min, and the absorbance was measured at 517 nm using a microplate reader 154 (Infinite F50R, Tecan Japan, Kawasaki, Japan). Each measurement was performed in 155 triplicate, and the percentage of the scavenging effect was calculated as follows: 156 DPPH radical scavenging activity (%) =  $(A_b - A_s)/A_b \times 100$ , 157 where  $A_b$  and  $A_s$  denote the absorbance of deionized water and sample solution, respectively. 158

159 2.3. Molecular weight distribution of peptides

160 Peptides and CSp residues in the BP-sCS were separated *via* ultrafiltration using the

161	molecular weight cutoff (MWCO) of 3000 and 10000 Da (Merck Millipore, Darmstadt,
162	Germany). A 5-mL aliquot of BP-sCS solution was firstly centrifuged at $4,000 \times g$ for 20 min
163	and then eluted with 5-mL deionized water at 4,000 $\times$ g for 20 min using a filter with an
164	MWCO of 10000 Da. The elution process was repeated several times to obtain <10000 Da
165	fraction. Then, the <10000 Da fraction was separated into 10000–3000 Da and <3000 Da
166	fractions using a filter with an MWCO of 3000 Da. Since the MW of the CSp in BP-sCS is
167	much larger than 3000 Da (Supplementary Fig. 1B), the <3000 Da fraction of BP-sCS did not
168	contain CS (Supplementary Fig. 1C). However, the separation also excluded peptides larger
169	than 3000 Da in BP-sCS. Therefore, the <3000 Da fraction of Pa-CP was obtained to
170	compare its bioactivity with the <3000 Da fraction of BP-sCS. The collected fractions were
171	freeze-dried. The MW distribution of the <3000 Da fractions was determined via size
172	exclusion chromatography using a TSK gel G2500PW column (7.5 mm $\times$ 30 cm, Tosoh,
173	Tokyo, Japan) with UV detection at 214 nm and a mobile phase of $35\%$ acetonitrile in $0.05\%$
174	TFA (pH 2.1) at a flow rate of 0.6 mL/min. An MW calibration curve was obtained using the
175	following standards: insulin (5700 Da), vitamin B <sub>12</sub> (1355 Da), and triglycine (189 Da). A 30-
176	$\mu$ L aliquot of each sample or standard was analyzed using High Performance Liquid
177	Chromatography system (HPLC, SCL-10AVP, Shimadzu, Tokyo, Japan), and the percentage
178	of each MW fraction was calculated as follows:

179 MW fraction (%) =  $(S_x/S_{total}) \times 100$ ,

where  $S_x$  and  $S_{total}$  denote the area of each fraction (>5000 Da, 3000–5000 Da, 1000–3000

181 Da, and <1000 Da) and the total area of the chromatogram, respectively.

## *2.4.1 Cell culture condition*

184	L929 fibroblast cells from the RIKEN Cell Bank (Tsukuba, Japan) was cultured at 37 °C
185	with 5% CO <sub>2</sub> in the minimum essential medium (MEM, Gibco, Grand Island, NY, USA)
186	containing 1% penicillin/streptomycin (Thermo Fisher Scientific, Waltham, MA) with 5%
187	fetal bovine serum (FBS, Lot. No 451456, Gibco). The medium was changed every 2–3 d.
188	The samples dissolved in endotoxin-free water (EFW) were added to the culture medium at
189	various concentrations for different experiments, and EFW was used as a negative control.
190	The endotoxin levels in MEM and the samples (1 mg/mL) were less than 1.0 EU/mL.
191	2.4.2. Antioxidant assay in cell culture
192	Cell viability/activity assay under oxidative stress was performed using the strong
193	oxidant H2O2 (Wako Pure Chemical) and the weak oxidant AAPH (Tokyo Chemical
194	Industry). Briefly, L929 cells were seeded in each well of a 96-well plate ( $2 \times 10^3$ cells/well).
195	When the cells reached >90% confluency, samples were added to the culture medium to
196	produce a final concentration of 1 mg/mL. After 24 h, the cells were exposed to ${\rm H_2O_2}$ (0.5
197	mM) or AAPH (5 mM) and incubated for another 24 h. After incubation, the cells were
198	washed with the Hank's balanced salt solution (HBSS, Sigma-Aldrich, Saint Louis, MO,
199	USA), and cell viability was assessed using Cell Counting Kit-8 (CCK-8, Dojindo,
200	Kumamoto, Japan), which measures the total metabolic activity. For the assay, the culture
201	medium was replaced with that containing 10% (v/v) CCK-8, and the cells were incubated at
202	37 °C for 30 min, and the absorbance at 450 nm was measured using a microplate reader

(Infinite F50R, TECAN, Kawasaki, Japan). This assay was also used to estimate the total cell 203 number in the well to evaluate cell proliferation; this measurement postulates that the 204 metabolism of each cell remains constant during the assay. As we cannot ascertain whether 205 the cell metabolism was affected by the addition of samples, the term 206 "proliferation/metabolism" was used for the measurements of cell proliferation assessed in 207 this experiment. The data were expressed as the relative absorbance normalized to that of the 208 control well (no oxidant). 209 Intracellular ROS formation was assessed according to the method described by Ahn et 210 211 al. (2012) with minor modifications. The oxidation-sensitive dye H<sub>2</sub>DCFDA was used as a probe as it diffuses through the cell membrane and is hydrolyzed into its nonfluorescent form 212 dichlorofluorescein (DCFH) by intracellular esterase. Then, DCFH reacts with the 213 214 intracellular H<sub>2</sub>O<sub>2</sub> to produce an oxidized form of DCFH, which is a green fluorescent dye. L929 cells ( $2 \times 10^3$  cells/well) were grown in a black/transparent 96-well microtiter plate 215 until they reached >90% confluency and were treated with the samples for 24 h. After 216 217 discarding the culture medium, the cells were washed with HBSS and labeled with H<sub>2</sub>DCFDA (20 µM; Sigma-Aldrich) in HBSS for 20 min in the dark at 37 °C and then 218 washed with HBSS and incubated with 0.5 mM H<sub>2</sub>O<sub>2</sub> in HBSS for 30 min in the dark at 37 219 °C. Fluorescence was read at an excitation wavelength of 485 nm and an emission 220 221 wavelength of 528 nm using a fluorescence microplate reader (WallAc ARVO 1420 Multilabel Counter, PerkinElmer, Waltham, MA, USA). VC was used as a positive control. 222 Data have been expressed as the relative fluorescence intensity normalized to that of the 223

224 control well (no oxidant).

226	L929 cells (5 × 10 <sup>3</sup> cells/well) were seeded in a 48-well plate. After 24 h, the culture
227	medium was replaced with a medium containing 1-mg/mL sample (Day 0). Fibroblast
228	proliferation was assessed based on a method described by Li et al. (2019) using the CCK-8
229	assay. Then, type I collagen gene expression was analyzed via real-time quantitative PCR
230	(qPCR). Total RNA was extracted from each well using the ISOGEN II reagent (Nippon
231	Gene, Tokyo, Japan) and quantified using a spectrophotometer (ND-1000, Thermo Fisher
232	Scientific, Waltham, MA, USA). cDNA was synthesized from 500 ng total RNA using a
233	PrimeScript RT Reagent Kit with the gDNA Eraser Perfect Real Time (Takara Holdings,
234	Ohtsu, Japan), according to the manufacturer's instructions. qPCR was performed using a
235	real-time PCR system (LightCycler® Nano System, Basel, Switzerland) with the synthesized
236	cDNA, which was diluted five times, as the template. Amplification was performed at a final
237	volume of 15 $\mu$ L, containing 1 $\mu$ L of the cDNA template, 7.5 $\mu$ L of 2X SYBR (FastStart
238	Universal SYBR Green Master, Roche, Basel, Switzerland), 1.5 $\mu$ L of each primer (5 $\mu$ M
239	each), and 3.5 $\mu L$ of sterilized water. PCR was performed as follows: 95 °C for 10 min,
240	followed by 40 cycles of 95 °C for 10 s, 56 °C for 10 s, and 72 °C for 15 s. The reaction used
241	primers for the type I procollagen al gene Collal (COL1A1-F: 5'-AAC CCG AGG TAT
242	GCT TGA TCT-3' and COL1A1-R: 5'-CCA GTT CTT CAT TGC ATT GC-3') and Gapdh
243	(GAPDH-F: 5'-TCC CAC TCT TCC ACC TTC-3' and GAPDH-R: 5'-CTG TAG CCG TAT
244	TCA TTG TC-3') (Kanazawa et al., 2008; Yoshimoto et al., 2009). Relative gene expression
245	was calculated using the $2^{-\Delta\Delta CT}$ method with <i>Gapdh</i> as the internal control (Schmittgen &
246	Livak, 2008). Data have been expressed as relative values normalized based on the control

247 well (EFW only).

Collagen production was quantified by the Sircol<sup>TM</sup> Soluble Collagen Assay (Biocolor 248 Ltd., Carrickfergus, Northern Ireland, UK) according to the manufacturer's instruction. In 249 this experiment, L929 cells  $(1.2 \times 10^4 \text{ cells/well})$  were seeded in 24-well plates. The culture 250 medium was sampled on Days 3 and 6 to measure the collagen content in the medium. On 251 Day 3, a new culture medium was added after the sampling. On Day 6, 0.1 mg/mL pepsin in 252 0.5 M acetic acid was added after the sampling of the medium, and the wells were incubated 253 overnight at 4 °C to extract collagen from the extracellular matrix (ECM) secreted by L929 254 255 cells during the culture. In the ECM, collagen molecules assembled into fibrils and became insoluble. Pepsin can remove the terminal non-helical telopeptides of collagen molecules to 256 release them into the extract. Prior to the assay, the samples were concentrated using the 257 258 isolation and concentration reagent provided in the kit. In the case of the extracted samples, those from three wells were mixed to make one test-sample as the collagen concentration was 259 low. Then, the collagen concentrations were quantified using a Sircol assay kit according to 260 the manufacturer's instructions. The collagen test was conducted in triplicate for each sample. 261 The amount of soluble collagen secreted into the culture medium was obtained as the sum of 262 collagen in the media on Days 3 and 6. 263

264

## 2.4.4. Fibroblast migration

To assess the effects of BP-sCS on the migration of fibroblasts, the scratch assay was conducted using the methods described by Liang et al. (2007) with minor modifications. Briefly, L929 cells ( $1.2 \times 10^4$  cells/well) were seeded in 24-well plates with a reference point

268	in each well, and when the cells reached confluence, a scratch through the reference point in
269	the cell monolayer was generated using a sterile 200- $\mu$ L pipette tip. The wells were washed
270	with HBSS to remove debris, and the culture medium containing each sample was added. For
271	the scratch-oxidative stress tests, AAPH was added to the culture medium at a final
272	concentration of 5 or 2.5 mM (Alvarez-Suarez et al., 2016).
273	The scratch was gradually occupied with the fibroblasts migrating from the non-scratched
274	area during the culture. Thus, time-course changes on the area of each scratch were observed
275	and photographed under a microscope (DMI600B, Leia, Wetzlar, Germany), and the images
276	obtained for each well were quantitatively analyzed using the ImageJ software (1.52a,
277	National Institutes of Health, USA). In brief, the scratch was defined as the region of interest,
278	and its area was measured using the software. The fibroblast-migration activity was assessed
279	as the repair rate (percentage scratch closure) calculated as follows:
280	Repair rate (%) = [area (0 h) – area (x h)]/area (0 h) × 100,
281	where area (0 h) and area (x h) indicate the scratch area at time 0 and time x ( $x = 6, 12, 24,$
282	30, and 36 h), respectively.
283	2.5. Statistical analyses
284	Data have been expressed as the mean $\pm$ standard error. Statistical analyses were
285	conducted using Student's t-test, Dunnett's test, or the Steel-Dwass test after ANOVA with

the Microsoft Excel add-in statistical software (SSRI, Tokyo, Japan).

**3. Results and discussion** 

# *3.1.1 Free-radical scavenging activity*

290	We examined the antioxidant activity of BP-sCS using ABTS and the DPPH scavenging
291	assays and found that BP-sCS exhibited a strong antioxidant activity, particularly in the
292	ABTS radical scavenging assay (Fig. 1A). At concentrations over 1 mg/mL, BP-sCS was able
293	to scavenge almost all the ABTS radicals, which was the same as VC. Although CSp
294	displayed no activity in the DPPH or ABTS assays, Pa-CP was able to scavenge radicals in a
295	dose-dependent manner; therefore, type II CP, not CSp, is one of the critical antioxidant
296	components in BP-sCS.
297	Next, we separated CSp and the peptides in BP-sCS using a 3000 Da MWCO
298	ultrafiltration membrane. Tricine-SDS-PAGE revealed that the <3000 Da fraction of BP-sCS
299	contained only peptides (Supplementary Fig. 1C), which may include type II CP and non-
300	collagenous peptides. Similarly, we separated the <3000 Da fraction of Pa-CP and compared
301	the ABTS radical scavenging activity of the <3000 Da BP-sCS and Pa-CP fractions (1
302	mg/mL). Antioxidant activity was significantly higher in the <3000 Da BP-sCS fraction than
303	that in Pa-CP (Fig. 2A), suggesting that non-collagenous peptides in BP-sCS contribute to its
304	antioxidant activity.
305	Further, we analyzed the MW distribution of the <3000 Da BP-sCS and Pa-CP fractions
306	via size exclusion chromatography (Fig. 2B) and found that BP-sCS contained more
307	<1000 Da peptides and less 3000–5000 Da peptides than Pa-CP. Membrane retention depends
308	not only on the solute's molecular size but also on its shape. The MWCO membrane cannot

309	fully reject the molecules whose MW is above their nominal MWCO (Sun et al., 2011).
310	Therefore, the <3000 Da fractions of BP-sCS included 3000–5000 Da peptides. Many studies
311	have reported that peptides with a lower MW exhibit a stronger antioxidant activity (Agrawal
312	et al., 2019; Nwachukwu & Aluko, 2019; Zou et al., 2016), and it has been suggested that
313	smaller peptides may expose more bioactive fragments via hydrolysis (Zou et al., 2016).
314	Therefore, the peptides with a lower MW in BP-sCS may also be responsible for its high
315	antioxidant activity.

3.1.2 Protection of cells from oxidative stress 316

H<sub>2</sub>O<sub>2</sub> generates free radicals; thus, it is commonly used to induce cell death and study 317 oxidative stress (Canas et al., 2007). In this study, we established an H<sub>2</sub>O<sub>2</sub>-induced oxidative 318 319 stress model (Fig. 3A), in which the addition of 0.5 mM H<sub>2</sub>O<sub>2</sub> significantly reduced cell viability/activity to approximately 70% that of the control cells, as quantified by the CCK-8 320 assay. Treating cells with 1 mg/mL of BP-sCS significantly reduced the H<sub>2</sub>O<sub>2</sub>-induced 321 damage, increasing their viability/activity to approximately 50% that of the control cells. Pa-322 CP demonstrated a smaller protective effect than BP-sCS, and the treatment with CSp did not 323 affect cell viability/activity. 324

We also produced another oxidative stress model using AAPH, which initiates free-325 radical reactions by generating radicals at a constant rate of 37 °C. Thus, AAPH acted more 326 slowly and had gentler effects than H<sub>2</sub>O<sub>2</sub>. AAPH (5 mM) treatment for 24 h reduced the cell 327 328 viability to approximately 50% that of the control group, which was gentler than the H<sub>2</sub>O<sub>2</sub> treatment (70%) (Fig. 3B). Treatment with BP-sCS or Pa-CP significantly increased the cell 329

330	viability/activity compared with that in the EFW group. CSp was again ineffective. Taken
331	together, the results from the H <sub>2</sub> O <sub>2</sub> and AAPH models indicate that peptides, but not CSp,
332	were the major compounds in BP-sCS, which have significant cytoprotective activities
333	against oxidative stress. This is consistent with the previous studies in which peptides derived
334	from the tilapia skin, lantern fish hydrolysate, and microalgae were found to exert
335	cytoprotective effects under AAPH- and H <sub>2</sub> O <sub>2</sub> -induced oxidative stresses (Chai et al., 2016;
336	Zeng et al., 2018; Zheng et al., 2018). Davalos et al. (2004) reported that among the amino
337	acids, tryptophan, tyrosine, and methionine exhibited the highest antioxidant activity,
338	followed by cysteine, histidine, and phenylalanine. Glycine and proline also play significant
339	roles in the antioxidant activity of peptides (Li et al., 2017). Thus, the antioxidant activity
340	must be the common biological activity of peptides, and the activity seems to be based on
341	their amino acid sequence. In addition, BP-sCS contained more <1000 Da peptides and less
342	3000–5000 Da peptides than Pa-CP (Fig. 2B). Therefore, the peptides with a lower MW in
343	BP-sCS, which may expose more bioactive fragments, has a higher potency of antioxidant
344	activity.

## *3.1.3 Reduction of intracellular ROS production*

Next, we examined whether the compounds in BP-sCS could reduce H<sub>2</sub>O<sub>2</sub>-induced intracellular ROS production using a fluorescent probe, whose fluorescence increases as free radicals are generated within a cell (Ahn et al., 2012). Exposing L929 cells to 0.5 mM H<sub>2</sub>O<sub>2</sub> significantly increased the intracellular ROS production (Fig. 4), whereas the ROS production was significantly reduced when the cells were treated with the positive control 351 (VC) or 1 mg/mL of BP-sCS. The same concentration of Pa-CP and CSp could prevent
352 intracellular ROS production, although they exhibited no significant cell-protective effects
353 against H<sub>2</sub>O<sub>2</sub>-induced oxidative stress.

Cellular antioxidant enzymes can deactivate intracellular ROS before they attack cellular 354 components and maintain cellular redox homeostasis (Tao et al., 2018). Non-collagenous 355 peptides and CS-A from cartilage have been reported to increase the intracellular levels of 356 antioxidant enzymes under oxidative stress (Canas et al., 2007; Tao et al., 2018). CS-A is the 357 second major component and occupies 26.6% of the skate CS (Supplementary Table 1). 358 359 Therefore, BP-sCS could regulate the antioxidant enzymes, at least in part, through its CS-A and non-collagenous peptides and reduce intracellular ROS. Although the relationships 360 between type II CP and antioxidant enzymes remain unclear, the collagen hydrolysate from 361 362 Nile tilapia skin (type I CP) enhanced the activities of antioxidant enzymes to alleviate oxidative stress (Wang et al., 2018). Thus, we believe that type II CP also influences 363 antioxidant enzymes. 364

365 In these experiments, the sample-containing culture medium had already been changed to that without samples before H<sub>2</sub>O<sub>2</sub> was added; thus, peptides and CS may have been 366 transported into cells via cell surface transporters or membrane channels to exert their 367 antioxidant activities. Alternatively, they may have bound cell surface receptors to produce 368 intracellular effects. Several peptide transporters (PEPTs) and CS receptors have been 369 reported. For instance, PEPT-1 was found to be involved in the cellular uptake of peptides in 370 skin keratinocytes (Kudo et al., 2016). Moreover, Zheng et al. (2018) reported that fish skin 371 peptides failed to inhibit ROS production in H<sub>2</sub>O<sub>2</sub>-exposed porcine enterocytes with PEPT1 372

373	knockdown. In addition, CS chain receptors, such as Toll-like receptor 2 and annexin 6, have
374	been identified on the surface of fibroblasts (Takagi et al., 2002; Wu et al., 2018). These
375	studies strongly suggest that the antioxidative effects of BP-sCS depend on transporter- and
376	receptor-related signaling pathways.
377	3.2. Effect of BP-sCS on L929 fibroblast activity
378	3.2.1 Fibroblast proliferation/metabolism
379	To evaluate their effect on fibroblast proliferation/metabolism, L929 cells were treated
380	with 0.01 mg/mL, 0.1 mg/mL, or 1 mg/mL of BP-sCS, Pa-CP, or CSp for up to 7 d (Fig. 5).
381	No significant effects were observed after 3 d (Fig. 5A); however, Pa-CP and CSp
382	significantly enhanced L929 fibroblast proliferation/metabolism in a dose-dependent manner
383	after 7 d (Fig. 5B). The lowest effective concentrations of Pa-CP and CSp were 0.01 and 0.1
384	mg/mL, respectively, whereas BP-sCS significantly activated fibroblast
385	proliferation/metabolism at 1 mg/mL. Thus, high BP-sCS concentrations promote fibroblast
386	proliferation/metabolism, and its activity is mainly due to type II CP and CSp.
387	Previously, we found that skate CS polysaccharides accelerated 3T3-L1 fibroblast
388	proliferation/metabolism (Li et al., 2019), whereas several studies have reported the fibroblast
389	proliferation activity of type I collagen-derived peptides. For example, collagen peptides
390	from the Asian sea bass were found to promote L929 fibroblast proliferation (Benjakul et al.,
391	2018), whereas those from tilapia scales were shown to stimulate human skin fibroblast
392	proliferation (Chai et al., 2010). However, this study is the first to report the effects of type II
393	peptides on fibroblast proliferation/metabolism.

395	Type I collagen is secreted by skin fibroblasts during wound healing to construct a new
396	extracellular matrix (Broughton et al., 2006). In this study, we first analyzed the type I
397	collagen al chain (Collal) mRNA expression. Cell proliferation assays determined the
398	effective BP-sCS dose (1000 $\mu$ g/mL) for subsequent experiments. As can be seen from Figure
399	6A, BP-sCS slightly and transiently activated the Collal gene expression compared with that
400	in control. Neither Pa-CP nor CSp increased the Collal expression. These results indicate
401	that BP-sCS stimulates type I collagen synthesis at the mRNA transcriptional level, whereas
402	the minimal effect of Pa-CP and CSp suggests that the BP-sCS activity is mainly due to non-
403	collagenous peptides.
404	Next, we monitored the effects of BP-sCS on collagen production by fibroblasts in two
405	phases: the soluble collagen secreted into the culture medium and the insoluble collagen
406	incorporated into the ECM. The results indicated that BP-sCS significantly lowered the
407	soluble collagen level of the culture medium but significantly increased the collagen level in
408	the ECM (Fig. 6B). These data indicated that BP-sCS changed the distribution ratio of
409	collagen produced by fibroblasts, i.e., BP-sCS induced more collagen to deposit into the
410	ECM and less collagen to secrete into the culture medium. The cell number increased by
411	1.94-fold, and the ECM collagen levels increased by 2.83-fold in the BP-sCS group
412	compared with that in the EFW group during a week of culture (Figs. 5, 6B). This suggests
413	that the increase in ECM collagen is due to not only the cell proliferation-promoting activity

414 but also the collagen deposition-promoting activity of BP-sCS. Several previous reports have

415 demonstrated that type I CP and non-collagenous peptides increased the intracellular collagen

416	contents of fibroblasts (Zeng et al., 2018) or accelerated the secretion of collagen into the
417	culture medium by fibroblasts (Benjakul et a., 2018; Chotphruethipong et al., 2019; Pozzolini
418	et al., 2018; Zague et al., 2018). However, to the best of our knowledge, this is the first study
419	that reports the promotion of collagen deposition into the ECM. Although the precise
420	mechanism of BP-sCS that stimulates collagen deposition into the ECM is not revealed by
421	this study, non-collagenous ECM proteins, such as small leucine-rich proteoglycans, secreted
422	by fibroblasts may regulate it as they play critical roles in collagen fibrillogenesis (Taye et al.,
423	2020). The effects of BP-sCS on non-collagenous ECM proteins should be studied in the
424	future.

### 425 *3.2.3 Fibroblast migration*

426 The scratch assay is particularly suitable for studying cell migration during wound healing (Liang et al., 2007). As presented in Figure 7A, treatment with BP-sCS, Pa-CP, or 427 CSp (1 mg/mL) tended to transiently and slightly increase L929 fibroblast migration during 428 429 the first 12 h, whereas the scratch healed in all groups after 24 h. Hu et al. (2017) prepared peptides via hydrolyzation of tilapia skin type I collagen with neutral protease and papain and 430 demonstrated that the peptides significantly enhanced the migration of HaCaT keratinocytes. 431 Therefore, the different amino acid compositions and MW distributions of the peptides may 432 vary their effects on cell migration. Moreover, peptides may differently affect the migration 433 of keratinocytes and fibroblasts. In addition, Zou et al. (2009) showed pure CS-C and CS-A 434 435 had a migration-promoting activity of human dermal fibroblasts. CS polysaccharides obtained from animal sources are a mixture of non-, mono-, and disulfated disaccharides. The 436

437	functionality of CS polysaccharides depends on their composition, which varies depending on
438	their species of origin (Li et al., 2019). Therefore, the difference in fibroblast migration
439	activity between the report of Zou et al. (2019) and the present study may be due to the
440	compositional differences of CS.
441	Next, we examined fibroblast migration under oxidative stress to simulate chronic
442	wound conditions. Treatment with 5-mM AAPH significantly reduced fibroblast migration
443	but was significantly counteracted by Pa-CP (Fig. 7B). A lower concentration of AAPH (2.5
444	mM) had a weaker inhibitory effect on fibroblast migration (Fig. 7C). Under this condition,
445	Pa-CP and BP-sCS stimulated fibroblast migration, although the differences were
446	insignificant. After 24 h, fibroblasts in the EFW and CSp groups started to die, and the
447	scratch site enlarged (Fig. 7C); however, the cells did not die, and the scratch did not enlarge
448	in the BP-sCS and Pa-CP groups (Fig. 7C). After 48 h, the fibroblasts in the control group (no
449	AAPH) contacted each other, and the scratch completely closed (Fig. 7D), whereas the
450	AAPH-stressed fibroblasts (EFW (AAPH) group in Fig. 7D) shrank and separated from each
451	other, which could be the result of cell contraction and reduced cell number due to oxidative
452	damage. Although the scratch in the BP-sCS and Pa-CP groups did not close, the AAPH-
453	stressed fibroblasts in these groups (BP-sCS + AAPH and Pa-CP + AAPH groups in Fig. 7D)
454	retained their normal morphology, and no further enlargement of the scratch was observed.
455	Therefore, BP-sCS may counteract the inhibitory effects of oxidative stress on fibroblast
456	migration, likely due to type II CP (Pa-CP), which significantly enhances fibroblast migration
457	under oxidative stress (Fig. 7B), consistent with the results of the antioxidant assay.
458	Excessive ROS levels are an essential feature of chronic, non-healing wounds (Schäfer &

Werner, 2008); therefore, the counteractivity of BP-sCS against ROS-induced loss of
fibroblast migration may be beneficial for chronic wound healing. A similar scratch-based
oxidative stress model previously demonstrated the antioxidant activity of honey, which
protected fibroblasts against oxidative damage and promoted fibroblast migration (AlvarezSuarez et al., 2016). This study is the first to report the activity of fish by-products on
fibroblast migration under oxidative stress.

465 **4. Conclusion** 

BP-sCS, a presently wasted by-product of skate CS extraction, is a combined 466 preparation of skate CSp, type II CP, and non-collagenous peptides. It exhibits antioxidant 467 activities, protects fibroblasts from oxidative stresses, promotes fibroblast 468 proliferation/metabolism, and counteracts oxidative stress damage that inhibits fibroblast 469 migration. We also found that the type II CP and non-collagenous peptides exhibited 470 antioxidant activities, whereas CSp and type II CP showed fibroblast activating properties; 471 however, none of these purified compounds exhibited all the bioactivities of BP-sCS 472 independently. Although many studies have reported the activities and applications of CS or 473 peptides as bioactive compounds, the industrial applications of BP-sCS, involving low-cost, 474 environmentally friendly products that are uncontaminated with zoonosis pathogens and are 475 subject to no religious objections, have not been considered. These advantages make BP-sCS 476 a competitive potential bioactive compound, such as a healing promoter of chronic wounds. 477 478 This study is the first step toward the realization of a sustainable CS production industry based on the utilization of wastes with bioactivity capabilities. However, our preliminary 479

480	estimation revealed that CS and BP-sCS occupy approximately 6% of by-products after skate
481	processing. Although skin (10% of by-products) is used for collagen production, a large part
482	of by-products is still not efficiently used. Further studies on the under-utilized sections of
483	by-products will be conducted in the future.

#### 484 Acknowledgments

This study was supported in part by a grant-in-aid for the Research and Development for the Recycle Technology from the Hokkaido Local Government Office. We would like to thank Editage group (www.editage.jp) for English language editing.

### 488 Declaration of interest: None

#### 489 **References**

- 490 Agrawal, H., Joshi, R., Gupta, M., 2019. Purification, identification and characterization of
- 491 two novel antioxidant peptides from finger millet (*Eleusine coracana*) protein
- 492 hydrolysate. Food Res. Int. 120, 697–707.
- Ahmed, T. A., Suso, H. P., Maqbool, A., Hincke, M. T., 2019. Processed eggshell membrane

494 powder: Bioinspiration for an innovative wound healing product. Mat. Sci. Eng. C. 95,

- 495 192–203.
- 496 Ahn, C. B., Je, J. Y., Cho, Y. S., 2012. Antioxidant and anti-inflammatory peptide fraction
- 497 from salmon by-product protein hydrolysates by peptic hydrolysis. Food Res. Int. 49,
  498 92–98.
- 499 Alvarez-Suarez, J. M., Giampieri, F., Cordero, M., Gasparrini, M., Forbes-Hernández, T. Y.,

500	Mazzoni, L., Afrin, S., Beltran-Ayala, P., Gonzalez-Paramas, A. M., Santos-Buelga, C.,
501	Varela-Lopez, A., Quiles, J. L., Battion, M., 2016. Activation of AMPK/Nrf2 signalling
502	by Manuka honey protects human dermal fibroblasts against oxidative damage by
503	improving antioxidant response and mitochondrial function promoting wound healing. J.
504	Funct. Foods. 25, 38–49.
505	Benjakul, S., Karnjanapratum, S., Visessanguan, W., 2018. Hydrolysed collagen from Lates
506	calcarifer skin: its acute toxicity and impact on cell proliferation and collagen
507	production of fibroblasts. Int. J. Food Sci. Tech. 53, 1871–1879.
508	Broughton, G. 2nd, Janis, J. E., Attinger, C. E., 2006. The basic science of wound healing.
509	Plast. Reconstr. Surg. 117, 12S–34S.
510	Canas, N., Valero, T., Villarroya, M., Montell, E., Verges, J., García, A. G., Lopez, M. G.,
511	2007. Chondroitin sulfate protects SH-SY5Y cells from oxidative stress by inducing
512	heme oxygenase-1 via phosphatidylinositol 3-kinase/Akt. J. Pharmacol. Exp. Ther. 323,
513	946–953.
514	Chai, H. J., Li, J. H., Huang, H. N., Li, T. L., Chan, Y. L., Shiau, C. Y., Wu, C. J., 2010.
515	Effects of sizes and conformations of fish-scale collagen peptides on facial skin qualities
516	and transdermal penetration efficiency. J. BioMed Res. 2010, 757301.
517	Chai, H. J., Wu, C. J., Yang, S. H., Li, T. L., Pan, B. S., 2016. Peptides from hydrolysate of
518	lantern fish (Benthosema Pterotum) proved neuroprotective in vitro and in vivo. J. Funct.
519	Foods. 24, 438–449.
520	Chotphruethipong, L., Aluko, R. E., Benjakul, S., 2019. Hydrolyzed collagen from porcine
521	lipase-defatted seabass skin: antioxidant, fibroblast cell proliferation, and collagen

- 522 production activities. J. Food Biochem. 43, e12825.
- 523 Davalos, A., Miguel, M., Bartolome, B., Lopez-Fandino, R., 2004. Antioxidant activity of
- peptides derived from egg white proteins by enzymatic hydrolysis. J. Food Prot. 67,
  1939–1944.
- 526 Dhivya, S., Padma, V. V., Santhini, E., 2015. Wound dressings a review. BioMed. 5, 24–28
- Han, G., Ceilley, R., 2017. Chronic wound healing: a review of current management and
  treatments. Adv. Ther. 34, 599–610.
- Hu, Z., Yang, P., Zhou, C., Li, S., Hong, P., 2017. Marine collagen peptides from the skin of
- 530Nile Tilapia (*Oreochromis Niloticus*): Characterization and wound healing evaluation.
- 531 Mar. Drugs, 15, 102–113.
- 532 Kanazawa, I., Yamaguchi, T., Yano, S., Yamauchi, M., Sugimoto, T., 2008. Metformin
- enhances the differentiation and mineralization of osteoblastic MC3T3-E1 cells via AMP
- kinase activation as well as eNOS and BMP-2 expression. Biochemi. Biophysi. Res.
- 535 Commun. 375, 414–419.
- 536 Kudo, M., Katayoshi, T., Kobayashi-Nakamura, K., Akagawa, M., Tsuji-Naito, K., 2016.
- H<sup>+</sup>/peptide transporter (PEPT2) is expressed in human epidermal keratinocytes and is
  involved in skin oligopeptide transport. Biochem. Bioph. Res. Co. 475, 335–341.
- Kurahashi, T., Fujii, J., 2015. Roles of antioxidative enzymes in wound healing. J. Dev. Biol.
  3, 57–70.
- 541 Li, H., Low, Y. S. J., Chong, H. P., Zin, M. T., Lee, C. Y., Li, B., Leolukman, M., Kang, L.,
- 542 2015. Microneedle-mediated delivery of copper peptide through skin. Pharm. Res. 32,
  543 2678–2689.

544	Li, W., Kobayashi, T., Moroi, S., Kotake, H., Ikoma, T., Saeki, H., Ura, K., Takagi, Y., 2019.
545	Anti-obesity effects of chondroitin sulfate oligosaccharides from the skate Raja
546	phulchra. Carbohyd. Polym. 214, 303–310.
547	Li, X., Chi, C., Li, L., Wang, B., 2017. Purification and identification of antioxidant peptides
548	from protein hydrolysate of scalloped hammerhead (Sphyrna lewini) cartilage. Mar
549	Drugs. 15, 1–16.
550	Liang, C. C., Park, A. Y., Guan, J. L., 2007. In vitro scratch assay: a convenient and
551	inexpensive method for analysis of cell migration in vitro. Nat. Protoc. 2, 329–333.
552	Meng, D., Tanaka, H., Kobayashi, T., Hatayama, H., Zhang, X., Ura, K., Yunoki, S., Takagi,
553	Y., 2019. The effect of alkaline pretreatment on the biochemical characteristics and
554	fibril-forming abilities of types I and II collagen extracted from bester sturgeon by-
555	products. Int. J. Biol. Macromol. 131, 572-580.
556	Murado, M. A., Fraguas, J., Montemayor, M. I., Vázquez, J. A., González, P., 2010.
557	Preparation of highly purified chondroitin sulphate from skate (Raja clavata) cartilage
558	by-products. Biochem. Eng. J., 49, 126–132.
559	Nwachukwu, I. D., Aluko, R. E., 2019. Structural and functional properties of food protein -
560	derived antioxidant peptides. J. Food Biochem. 43, e12761.
561	Pal, G. K., Suresh, P. V., 2016. Sustainable valorisation of seafood by-products: recovery of
562	collagen and development of collagen-based novel functional food ingredients. Innov.
563	Food Sci. Emerg. 37, 201–215.
564	Pozzolini, M., Millo, E., Oliveri, C., Mirata, S., Salis, A., Damonte, G., Arkel, M., Scarfi, S.,
565	2018. Elicited ROS scavenging activity, photoprotective, and wound-healing properties
	28

- of collagen-derived peptides from the marine sponge *Chondrosia reniformis*. Mar. Drugs
  16, 465–491.
- 568 Restaino, O. F., di Lauro, I., Di Nuzzo, R., De Rosa, M., Schiraldi, C., 2017. New insight into
- chondroitin and heparosan-like capsular polysaccharide synthesis by profiling of the
- 570 nucleotide sugar precursors. Biosci. Rep., 37, 1–11.
- 571 Restaino, O. F., Finamore, R., Stellavato, A., Diana, P., Bedini, E., Trifuoggi, M., Rosa, M.,
- 572 Schiraldi, C., 2019. European chondroitin sulfate and glucosamine food supplements: a
- 573 systematic quality and quantity assessment compared to pharmaceuticals. Carbohyd.
- 574 Polym. 222, 114984.
- 575 Schäfer, M., Werner, S., 2008. Oxidative stress in normal and impaired wound repair.
- 576 Pharmacol. Res. 58, 165–171.
- 577 Schiraldi, C., Alfano, A., Cimini, D., Rosa, M. D., Panariello, A., Restaino, O. F., Rosa, M.
- 578 D., 2012. Application of a 22L scale membrane bioreactor and cross flow
- ultrafiltration to obtain purified chondroitin. Biotechnol. Progr., 28, 1012–1018.
- 580 Schmittgen, T. D., Livak, K. J., 2008. Analyzing real-time PCR data by the comparative CT
- 581 method. Nat. Protoc. 3, 1101–1103.
- 582 Shang, Q., Shi, J., Song, G., Zhang, M., Cai, C., Hao, J., Li, G., Yu, G., 2016. Structural
- 583 modulation of gut microbiota by chondroitin sulfate and its oligosaccharide. Int. J. of
- 584 Boil. Macromol. 89, 489–498.
- 585 Shen, W., Matsui, T., 2017. Current knowledge of intestinal absorption of bioactive peptides.
- 586 Food Funct. 8, 4306–4314.
- 587 Sun, H., Qi, D., Xu, Juan, J., S., Zhe, C., 2011. Fractionation of polysaccharides from

588	rapeseed by ultrafiltration: Effect of molecular pore size and operation conditions on the
589	membrane performance. Sep. Purify. Technol. 80, 670-676.
590	Takagi, H., Asano, Y., Yamakawa, N., Matsumoto, I., Kimata, K., 2002. Annexin 6 is a
591	putative cell surface receptor for chondroitin sulfate chains. J. Cell Sci. 115, 3309-3318.
592	Tao, J., Zhao, Y. Q., Chi, C. Wang, F., B., 2018. Bioactive peptides from cartilage protein
593	hydrolysate of spotless smoothhound and their antioxidant activity in vitro. Mar. Drugs.
594	16, 100–118.
595	Taye, N., Karoulias, S. Z., Hubmacher, D., 2020. The "other" 15-40%: the role of non-
596	collagenous extracellular matrix proteins and minor collagens in tendon. J. Orthop. Res.
597	38, 23–35.
598	Vázquez, J. A., Rodríguez-Amado, I., Montemayor, M. I., Fraguas, J., González, M. D. P.,
599	Murado, M. A., 2013. Chondroitin sulfate, hyaluronic acid and chitin/chitosan
600	production using marine waste sources: Characteristics, applications and eco-friendly
601	processes: A review. Mar. Drugs. 11, 747–774.
602	Volpi, N., 2007. Analytical aspects of pharmaceutical grade chondroitin sulfates. J. Pharm.
603	Sci. 96, 3168–3180.
604	Wang, L., Jiang, Y., Wang, X., Zhou, J., Cui, H., Xu, W., He, Y., Ma, H., Gao, R., 2018. Effect
605	of oral administration of collagen hydrolysates from Nile tilapia on the chronologically
606	aged skin. J. Funct. Foods, 44, 112–117.
607	Wu, F., Zhou, C., Zhou, D., Ou, S., Liu, Z., Huang, H., 2018. Immune-enhancing activities of
608	chondroitin sulfate in murine macrophage RAW 264.7 cells. Carbohyd. Polym. 168,
609	611–619.
	30

610	Yoshimoto, M., Heike, T., Chang, H., Kanatsu-Shinohara, M., Baba, S., Varnau, J. T.,
611	Shinoharac, T., Yoder, M. C., Nakahata, 2009. T., Bone marrow engraftment but limited
612	expansion of hematopoietic cells from multipotent germline stem cells derived from
613	neonatal mouse testis. Exp. Hematol. 37, 1400–1410.
614	Zague, V., do Amaral, J. B., Rezende Teixeira, P., de Oliveira Niero, E. L., Lauand, C.,
615	Machado-Santelli, G. M., 2018. Collagen peptides modulate the metabolism of
616	extracellular matrix by human dermal fibroblasts derived from sun-protected and sun-
617	exposed body sites. Cell Biol. Int. 42, 95–104.
618	Zeng, Q., Fan, X., Zheng, Q., Wang, J., Zhang, X., 2018. Anti-oxidant, hemolysis inhibition,
619	and collagen-stimulating activities of a new hexapeptide derived from Arthrospira
620	(Spirulina) platensis. J. Appl. Phycol. 30, 1655–1665.
621	Zheng, L., Wei, H., Yu, H., Xing, Q., Zou, Y., Zhou, Y., Peng, J., 2018. Fish skin gelatin
622	hydrolysate production by ginger powder induces glutathione synthesis to prevent
623	hydrogen peroxide induced intestinal oxidative stress via the Pept1-p62-Nrf2 cascade. J.
624	Agr. Food. Chem. 66, 11601–11611.
625	Zou, T. B., He, T. P., Li, H. B., Tang, H. W., Xia, E. Q., 2016. The structure-activity
626	relationship of the antioxidant peptides from natural proteins. Molecules. 21, 72-86.
627	Zou, X. H., Jiang, Y. Z., Zhang, G. R., Jin, H. M., Hieu, N. T. M., Ouyang, H. W., 2009.
628	Specific interactions between human fibroblasts and particular chondroitin sulfate

629 molecules for wound healing. Acta Biomater. 5, 1588–1595.

**Fig. 1.** ABTS (A) and DPPH (B) radical scavenging activities of the by-product of skate chondroitin sulfate production (BP-sCS), the papain-hydrolyzed type II collagen peptides (Pa-CP), and the chondroitin sulfate polysaccharides (CSp). Values have been expressed as the mean  $\pm$  standard error (n = 3).

**Fig. 2.** ABTS radical scavenging activities (A) and molecular weight distributions (B) of <br/>
<br/> **Solution**<br/>
<br/> **Solution**<br/>
<br/> **Solution**<br/>
<br/>
<br/

643	<b>Fig. 3.</b> Effects of the by-product of skate chondroitin sulfate production (BP-sCS), the
644	papain-hydrolyzed type II collagen peptides (Pa-CP), and the chondroitin sulfate
645	polysaccharides (CSp) against (A) H <sub>2</sub> O <sub>2</sub> - and (B) AAPH-induced oxidative stress on the
646	L929 fibroblast proliferation/metabolism. Control groups were not treated with oxidants.
647	Oxidative stress was induced in all other groups using $H_2O_2(A)$ or AAPH (B). Sample
648	concentration, 1 mg/mL. EFW, endotoxin-free water. Columns and bars indicate the mean
649	value ± standard error ( $n = 12-16$ ). # $p < 0.05$ , ## $p < 0.01$ compared to the control group (the
650	Steel-Dwass test). Different letters denote significant differences between groups ( $p < 0.05$ ,
651	the Steel-Dwass test).

653	Fig. 4. Effects of the by-product of skate chondroitin sulfate production (BP-sCS), the
654	papain-hydrolyzed type II collagen peptides (Pa-CP), and the chondroitin sulfate
655	polysaccharides (CSp) on the H <sub>2</sub> O <sub>2</sub> -induced intracellular ROS production. Control groups
656	were not treated with $H_2O_2$ . Oxidative stress was induced in all other groups using 0.5 mM
657	H <sub>2</sub> O <sub>2</sub> . EFW, endotoxin-free water; VC, Vitamin C 0.2 mg/mL; BP-sCS 0.01, BP-sCS 0.01
658	mg/mL; BP-sCS 0.1, BP-sCS 0.1 mg/mL; BP-sCS 1, BP-sCS 1 mg/mL; Pa-CP 0.01, Pa-CP
659	0.01 mg/mL; Pa-CP 0.1, Pa-CP 0.1 mg/mL; Pa-CP 1, Pa-CP 1 mg/mL; CSp 0.01, CSp 0.01
660	mg/mL; CSp 0.1, CSp 0.1 mg/mL; CSp 1, CSp 1 mg/mL. Columns and bars indicate the
661	mean value $\pm$ standard error ( $n = 9-10$ ). Different letters denote significant differences
662	between groups ( $p < 0.05$ , the Steel-Dwass test).



674

hydrolyzed type II collagen peptides (Pa-CP), and the chondroitin sulfate polysaccharides 675 (CSp) on collagen mRNA expression (A) and the amount of collagen secreted into the culture 676 medium or the extracellular matrix (ECM) (B) by the L929 fibroblasts. The collagen mRNA 677 expression levels have been normalized to the internal control (Gapdh) and expressed relative 678 679 to the corresponding endotoxin-free water (EFW) control. Sample concentration, 1 mg/mL. Columns and bars indicate the mean value  $\pm$  standard error (n = 11-12 in A and n = 6 in B). \* 680 p < 0.05, compared to the EFW control group (the Dunnett's test). \*\* p < 0.01, Student's *t*-test. 681 682 Fig. 7. Effect of the by-product of skate chondroitin sulfate production (BP-sCS), the papain-683 hydrolyzed type II collagen peptides (Pa-CP), and the chondroitin sulfate polysaccharides 684 (CSp) on the L929 fibroblast migration. (A) The scratch assay without oxidative stress. 685 Samples concentration, 1 mg/mL. Values have been expressed as the mean  $\pm$  standard error (*n* 686 = 5–6). # p < 0.05 compared to the control group (the Dunnett's test). (B, C) The scratch assay 687 under the oxidative stress. The control group was not treated with AAPH. Other groups were 688 treated with 5.0 mM (B) or 2.5 mM (C) AAPH. Samples concentration, 1 mg/mL. Values 689 have been expressed as the mean  $\pm$  standard error (n = 5-6). # p < 0.05 compared to the 690 control group (the Dunnett's test). p < 0.05 compared to the endotoxin-free water (EFW) 691

Fig. 6. Effect of the by-product of skate chondroitin sulfate production (BP-sCS), the papain-

group (the Dunnett's test). (D) Photomicrographs of the L929 fibroblasts at 48 h in the scratch

693 assay under the oxidative stress. Samples concentration, 1 mg/mL. Bars, 250  $\mu$ m in photos

and enlarged photos.



#### Supplementary Fig.1. Composition of the by-product of skate chondroitin sulfate (BP-sCS)

(A) BP-sCS contains CS polysaccharides, peptides, minerals and others (lipid, water, and so

on). The lower graph showed the molecular weight distribution of peptides.

(B) Tricine-SDS-PAGE of BP-sCS with Coomassie Brilliant Blue (CBB) (left) and Alcian blue (right) stains. CBB and Alcian blue stains showed peptides and CS polysaccharides in BP-sCS, respectively. M, protein markers.

(C) Tricine-SDS-PAGE of BP-sCS separated by ultrafiltration membrane. Samples were stained with CBB and Alcian blue. CBB and Alcian blue stains showed peptides and CS polysaccharides in different fractions of BP-sCS, respectively. < 3000 Da fraction was negative

to Alcian blue stain, suggesting it contains no CS. M, protein markers; >10, BP-sCS fraction of more than 10000 Da; 3–10, BP-sCS fraction of between 3000–10000 Da; < 3, BP-sCS fraction of smaller than 3000 Da.

## Supplementary Table 1

	Formulations (%)						
	∆Di-0s	ΔDi-4s (CS-A)	ΔDi-6s (CS-C)	ΔDi-2,6s (CS-D)	ΔDi-4,6s (CS-E)	(kDa)	
CSp	6.5	26.6	60.9	6.5	0.1	37–250	

The formulations and molecular weights of CSp

Data of formulations for CSp were obtained by Marukyo Bio Foods Co. Ltd. Data of molecular

weights were obtained by 16.5% tricine-SDS-PAGE stained with Alcian blue.



Fig. 1



Fig. 2



Fig. 3



Fig. 4



Fig. 5



Fig. 6



(D) 2.5 mM AAPH, 48 h

