#### BUKTI KORESPONDENSI JURNAL INTERNASIONAL BEREPUTASI

- Judul Artikel : (C15) Biocompatibility of crab shell- derived graft sterilized by ultraviolet light towards human gingival fifibroblast cell culture
- Jurnal : World Journal of Advanced Research and Reviews
- Penulis : *Michael Josef Kridanto Kamadjaja*\*, Bambang Agustono dan Innocencio Kresna Pratama

No	Perihal	Tanggal	Halaman
1	Bukti submit dan artikel yang disubmit	12 Maret 2023	1-2
2	Bukti Revisi 1	14 Maret 2023	3-5
3	Bukti Revisi 2	17 Maret 2023	6-8
4	Bukti Revisi 3	18 Maret 2023	9-10
5	Decision 1	18 Maret 2023	11-20
6	Decision 2		
7	Decision 3		
8	Decision 4		
9	Bukti accepted	20 Maret 2023	21-24
10	Bukti published	22 Maret 2023	25



#### Call for Paper: World Journal of Advanced Research and Reviews, Volume 17, Issue 3

Editor WJARR <editor@wjarr.com> Reply-To: Editor WJARR <editor@wjarr.com> To: Michael-j-k-k@fkg.unair.ac.id Sun, Mar 12, 2023 at 11:33 AM

View web version

## World Journal of Advanced Research and Review

#### (WJARR)

#### e-ISSN: 2581-9615 CODEN (USA): WJARAI

#### **CALL FOR PAPER**

Dear Colleague,

We cordially invite you to submit your precious research manuscripts (Original research, review articles, Short communication and letter to editor etc.) under various disciplines of Biological, Pharmaceutical and Health Sciences that falling within the scope of **World Journal of Advanced Research and Reviews (WJARR)** for publication in upcoming issue of this journal.

#### Submit manuscript

Click here for Online Submission

or E-mail editor@wjarr.com

Go through Authors guide before submission

**IMPACT FACTOR: 7.8** 

Google Scholar Indexing

valid Crossref DOI and Cross ref-mark

Publication certificate to each author

#### **Benefits to author**

Rapid, easy and high quality of Publication

Low article processing charges

**Online tracking** of articles submitted

Publish articles within 3 days from acceptance

Looking forward for your kind response and quality submission for possible publication in World Journal of Advanced Research and Reviews (WJARR).

Please feel free to contact us, if you have any query or concerns.

With best regards, Managing Editor World Journal of Advanced Research and Reviews (WJARR) editor@wjarr.com We hope you enjoy receiving update news and offer emails from https://wjarr.com/.

If you would prefer not receiving our emails, please click here to unsubscribe



#### Call for Paper: World Journal of Advanced Research and Reviews, Volume 17, Issue 3

editor@wjarr.com <editor@wjarr.com> To: michael josef kridanto kamadjaja <michael-j-k-k@fkg.unair.ac.id>

Tue, Mar 14, 2023 at 10:15 PM

Manuscript No.: WJARR-2023-0438 Submitted by: Michael Josef E-mail of Corresponding Author: michael-j-k-k@fkg.unair.ac.id Article Title: Biocompatibility of Crab Shell-Derived Graft Sterilized by UV Light towards Human Gingival Fibroblast Cell Culture Status of Article: Your article is Submitted to the Journal [Quoted text hidden]



# Your Manuscript (WJARR-2023-0438) is submitted to World Journal of Advanced Research and Reviews (WJARR)

**World Journal of advanced Research and Reviews** <editor@wjarr.com> To: michael-j-k-k@fkg.unair.ac.id Tue, Mar 14, 2023 at 10:15 PM

Dear Author, Greetings for the day...

Your manuscript entitled "Biocompatibility of Crab Shell-Derived Graft Sterilized by UV Light towards Human Gingival Fibroblast Cell Culture" has been received to "World Journal of Advanced Research and Reviews (WJARR)". Manuscript number assigned to your manuscript is WJARR-2023-0438. Use this reference number in the future communications. Also, you will be able to check the status of your submitted manuscript by logging manuscript number and corresponding author's email id in Track manuscript form of World Journal of Advanced Research and Reviews (WJARR) website. http://wjarr.com/content/trackmanuscript-status

If you need more information, feel free to contact us.

With Best Regards, Managing Editor World Journal of Advanced Research and Reviews (WJARR) editor@wjarr.com



#### Call for Paper: World Journal of Advanced Research and Reviews, Volume 17, Issue 3

**michael josef kridanto kamadjaja** <michael-j-k-k@fkg.unair.ac.id> To: Editor WJARR <editor@wjarr.com> Tue, Mar 14, 2023 at 2:04 PM

Dear Editor WJARR,

I would like to submit a research manuscript to your esteemed journal. The manuscript is entitled "Biocompatibility of Crab Shell-Derived Graft Sterilized by UV Light towards Human Gingival Fibroblast Cell Culture ", authored by Michael Josef Kridanto Kamadjaja, Bambang Agustono, and Innocencio Kresna Pratama.

Hereby I also attached the copyright form, cover letter, and manuscript file.

Should you require any further queries regarding the submission of the manuscript, please feel free to contact me.

Best regards, Michael Josef

[Quoted text hidden]

#### 3 attachments WJARR Authors Declaration INNO.docx 31K [DJMKG] Original article template.docx

2835K

B WJARR Authors Declaration INNO.pdf



# Editorial decision on your Manuscript (WJARR-2023-0438): accepted for further publication process after initial editorial review

**World Journal of advanced Research and Reviews** <editor@wjarr.com> To: michael-j-k-k@fkg.unair.ac.id Fri, Mar 17, 2023 at 7:46 PM

Dear Author, Greetings for the day...

We are pleased to inform you that your manuscript (Manuscript ref. no. WJARR-2023-0438) entitled "Biocompatibility of Crab Shell-Derived Graft Sterilized by UV Light towards Human Gingival Fibroblast Cell Culture" is approved for further publication process after initial editorial review.

In order to proceed for publication of your manuscript as fast as possible, follow the below steps

1) Provide Author's Declaration form (If not submitted earlier) and Author's name details. Your article cannot be published until the publisher has received the appropriate signed Author's Declaration form and filled Author's name details form (Download both forms from website. https://wjarr.com/content/downloads)

2) Make the payment of Article processing charges: within seven working days after receipt of this e-mail using as details given below

A] Online payment through RazorPay: Using debit cards, or credit cards or Razorpay log in credential. Amount payable through RazorPay is Total USD 32.00 (USD Thirty two only) [Article processing fee (USD 30.00) + Transaction charges (USD 2.00)] Click on link and follow instructions: https://rzp.io/I/NLH0xdL

**B] Online payment through PayPal:** Using PayPal log in credential, debit cards, or credit cards. Amount payable through PayPal is Total USD 32.00 (USD Thirty two only) [Article processing fee (USD 30.00) + Transaction charges (USD 2.00)] Click on below link and follow instructions: https://wjarr.com/content/paypalrjp

Author must e-mail copy of Payment details (Transaction ID) along with details involving Name of corresponding author, e-mail of corresponding author, manuscript number and Title of the manuscript to the editor of World Journal of Advanced Research and Reviews (WJARR) at editor@wjarr.com.

Please be aware that your manuscript can not be published until all above steps are followed.

If you need more information, feel free to contact us.

With Best Regards, Managing Editor World Journal of Advanced Research and Reviews (WJARR) editor@wjarr.com



#### World Journal of Advanced Research and Reviews (WJARR) <u>Author's Declaration Form</u>

**Title of article:** Biocompatibility of Crab Shell-Derived Graft Sterilized by UV Light towards Human Gingival Fibroblast Cell Culture

Corresponding Author's name: Michael Josef Kridanto Kamadjaja

Names of co-authors: Michael Josef Kridanto Kamadjaja, Bambang Agustono, Innocencio Kresna Pratama

E- mail of corresponding author: Michael-j-k-k@fkg.unair.ac.id

The author(s) acknowledged that:

- 1. All authors have read, understand and agreed to the submission guidelines, policies and submission declaration of the journal.
- 2. The manuscript is the authors' original work and has not been published or is not being submitted or considered for publication elsewhere.
- 3. All authors have checked the manuscript for plagiarism and declare that manuscript is plagiarism free. However, in future if any one report/complain about plagiarized part or text in the manuscript, authors will be responsible for the action (retraction) taken by journal editors/publisher..
- 4. All authors have seen and approved the manuscript as submitted.
- 5. All authors participated in the work in a substantive way and are prepared to take public responsibility for the work.
- 6. All authors of the manuscript have no conflict of interests to declare.
- 7. The manuscript submitted to the journal is not copied or plagiarized version of some other published work. All the data taken from other sources is written in authors own language and properly cited.
- 8. The text, illustrations, and any other materials included in the manuscript do not infringe upon any existing copyright or other rights of anyone.
- 9. Submission of false or incorrect information/undertaking would invite appropriate penal actions as per norms/rules of the journal.
- 10. All authors shall not submit the paper for publication in any other Journal or Magazine till the decision is made by journal editors.
- 11. All authors confirm that upon acceptance of the paper, they will publish the paper by paying Article Processing Charges (APC), or withdraw it by paying Withdrawal Fee.

Date: 14<sup>th</sup> March 2023 Place: Surabaya Signature (Corresponding Author)



#### Your Manuscript (WJARR-2023-0438): accepted for further publication

World Journal of advanced Research and Reviews <editor@wjarr.com> To: michael-j-k-k@fkg.unair.ac.id Fri, Mar 17, 2023 at 7:48 PM

Dear Author, Greetings for the day...

We are pleased to inform you that your manuscript (Manuscript ref. no. WJARR-2023-0438) entitled "Biocompatibility of Crab Shell-Derived Graft Sterilized by UV Light towards Human Gingival Fibroblast Cell Culture" is accepted for further publication process.

We already have sent a separate e-mail regarding this with details for submission of required documents as well as Payment of article processing charges. Please check it in your inbox or Spam folder and do the needful.

If you need more information, feel free to contact us.

With Best Regards, Managing Editor World Journal of Advanced Research and Reviews (WJARR)



#### **Payment of article**

michael-j-k-k <michael-j-k-k@fkg.unair.ac.id> To: editor@wjarr.com

Sat, Mar 18, 2023 at 10:13 AM

WJARR-2023-0438

"Biocompatibility of crab shell derived graft sterilized by UV light towards Human Gingival Fibroblast Cell Culture" Corresponding author : Michael Josef Kridanto Kamadjaja

Sent from my Galaxy



Screenshot\_20230318\_081032\_Samsung Internet.jpg



#### **Payment of article**

editor@wjarr.com <editor@wjarr.com> To: michael-j-k-k <michael-j-k-k@fkg.unair.ac.id> Sat, Mar 18, 2023 at 8:41 PM

Dear author, Greetings...

We received details of payment of article processing charges. We already have sent you a galley proof of your manuscript. That e-mail may have been landed in your spam folder (You may check there). Further action is pending from your side. Do the needful.

Thanks and regards, Editor

Quoting michael-j-k-k <michael-j-k-k@fkg.unair.ac.id>:

[Quoted text hidden]



#### WJARR-2023-0438 Article proof for final check and approval

Editor WJARR <editor@wjarr.com> Reply-To: editor@wjarr.com To: michael-j-k-k@fkg.unair.ac.id Sat, Mar 18, 2023 at 6:57 PM

Dear Author, Greetings for the day...

We are grateful for your association with us. Please find herewith attached final edited version (as a word file) of your manuscript (ref no. WJARR-2023-0438) entitled "Biocompatibility of Crab Shell-Derived Graft Sterilized by UV Light towards Human Gingival Fibroblast Cell Culture" which is accepted for publication in World Journal of Advanced Research and Reviews (WJARR).

### Download Author proof copy of your manuscript: https://wjarr.com/sites/default/files/Attached%20Files/WJARR-2023-0438%2...

Download Author query form: https://wjarr.com/content/downloads

#### Special Note:

- 1. Provide the name(s) of authors as First Name (Given name)-Middle Name- Last name (Family name/Surname) for each author and check that all names are accurately spelled. It is important from Indexing purpose.
- 2. Authors should write complete name of their department.
- 3. Scientific names of plant/animal/microbes should be written in Italics and follow nomenclature rules like *Genus species*. Check this throughout the manuscript.
- 4. Author should put °C sign with one space after numerical as 50 °C. Do this throughout the manuscript.
- 5. While writing numerical results in table author should use decimal point instead of comma. Ex Table 2.
- 6. Provide X and Y axis lable for figure 5.
- 7. Author should provide an appropriate conclusion to the article. Write conclusion as single para. Conclusion should be concise, informative and can be started with summarizing outcome of the study in 1-2 sentence and ended with one line stating: how this study will benefit to the society and way forward.
- 8. Provide appropriate disclosure of conflict of interest in the manuscript.
- 9. While writing reference citation in the main text at the end of sentence, author has put full stop before square bracket ".[1]". It should be corrected by writing full stop after square bracket as "[1].". Check and correct throughout the manuscript.
- 10. Ensure that, all the references appearing in the text should be enlisted at the end in reference section and vice versa.

Without this we will not able to process your manuscript further.

#### We hereby request you that,

1. Read the proof at least thrice, note the errors and make necessary corrections (in blue colored text) to make the manuscript error free.

- 2. Do not change the format of the word file, just check the spelling and other errors in the attached text part.
- 3. Use this proof for checking the typesetting, editing, completeness and correctness of the text, tables and figures.
- 4. Significant changes to the article will only be considered at this stage with permission from the Editor.

5. Make sure that you send final approved copy of article proof along with duly filled Author proof form within 3 days after receipt of this e-mail.

6. Your manuscript will not be further processed for publication until and unless we receive your final approved article proof and Author query form.

7. It is responsibility to author to check each and every word and spelling including italic and normal formatting of specific names. No changes will be accepted after this. If any changes are expected after publication of manuscript, author will have to pay charges accordingly.

We would appreciate to receive your corrected proof as a word file by e-mail within stipulated time period so that we can process your article quickly and efficiently.

Thank you for submitting your work to this journal.

If you need more information, please do not hesitate to contact us.

With Best Regards, Managing Editor World Journal of Advanced Research and Reviews (WJARR) editor@wjarr.com



eISSN: 2581-9615 CODEN (USA): WJARAI Cross Ref DOI: 10.30574/wjarr Journal homepage: https://wjarr.com/

WJARR	USSN 2581-961 CODEN (USA): WJARU
W	JARR
World Journal of	
Advanced	
Research and	
Reviews	
	World Journal Series INDIA
	World Journal Ser INDIA

(RESEARCH ARTICLE)

Check for updates

# Biocompatibility of crab shell-derived graft sterilized by ultraviolet light towards human gingival fibroblast cell culture

Michael Josef Kridanto Kamadjaja<sup>1,\*</sup>, Bambang Agustono<sup>1</sup> and Innocencio Kresna Pratama<sup>2</sup>

<sup>1</sup> Department of Prosthodontics, Faculty of Dental Medicine, Universitas Airlangga, Surabaya, Indonesia. <sup>2</sup> Faculty of Dental Medicine, Universitas Airlangga, Surabaya, Indonesia.

World Journal of Advanced Research and Reviews, 2023, XX(XX), XXX-XXX

Publication history: Received on 07 February 2023; revised on 15 March 2023; accepted on 18 March 2023

#### Abstract

**Background:** *Portunus pelagicus* crab species meat is generally taken, and their shells that is thrown away becoming untapped waste. Their shells contain so many useful minerals such as calcium carbonate which can be processed into hydroxyapatite (HA). HA has been extensively used in the medical field for bone grafting, to repair damaged in bone structures.

**Purpose:** To determine the biocompatibility of hydroxyapatite from *Portunus pelagicus* crab's shells extract after sterilized using UV on human gingival fibroblast cell.

**Methods:** This research is an experimental laboratory research using post-test only control group design. The shells were first heated using furnace. Once hydroxyapatite was obtained, it was sterilized using ultraviolet (UV) with certain concentration (25 ppm, 50 ppm, and 100 ppm) to kill the microorganisms. MTT assay was done to test the biocompatibility of the sterilized hydroxyapatite on human gingival fibroblast cell.

**Results**: The highest viability of fibroblast cells was the group with 50 ppm UV (98,10%), followed by 100 ppm UV (97,93%), and 25 ppm (93,28%).

**Conclusion:** hydroxyapatite from *Portunus pelagicus* crab's shells extract sterilized using UV is biocompatible towards human gingival fibroblast cell.

Keywords: Biocompatibility; UV light; Hydroxyapatite; Portunus pelagicus; Human gingival fibroblast cell; Medicine

#### 1. Introduction

Scylla serrata and *Portunus pelagicus* crab species are one of Indonesia's seafoods commodities that are often exported to the United States. In 2012, the export value of the two marine animals reached 9% and occupied the third place in the export of Indonesian seafood products after shrimp and tuna. About 60% of the volume of crabs exported is in processed form (canned crab).[1] The consumption of crab meat from the domestic community also contributes to shell waste. The by-product of crab meat processing in the form of shell waste reaches about 40-60% of the total weight of crabs. This waste has not been put to good use and is not efficient, in fact most of it is a waste that pollutes the environment.[2,3]

Crab shell waste contains organic and inorganic compounds, including protein (15.60% - 23.90%), CaCO<sub>3</sub> (53.70 - 78.40%), and chitin (18.70% - 32.20%). [4] Minerals contained in crab shells can be utilized and processed into

Copyright © 2023 Author(s) retain the copyright of this article. This article is published under the terms of the Creative Commons Attribution Liscense 4.0.

<sup>\*</sup> Corresponding author: Michael Josef Kridanto Kamadjaja

hydroxyapatite (HA) and can be synthesized in many ways. HA is widely used in orthopedic and biomedical field applications. The structure of HA has a similar structure with human bones which makes HA have the potential and innovation as a manufacture of synthetic bones (bone graft). In dentistry, bone graft is used to increase the alveolar bone volume, remodeling the jawbone, and reshaping the alveolar crest.[5,6]

To be used as a graft material, hydroxyapatite graft from crab shells (*Portunus pelagicus*) must meet biocompatibility requirements. Biocompatibility relates to the nature of the material that demonstrates biologic compatibility or is acceptable to the body without providing a harmful local or systemic response. This property is very important for synthetic bones (bone graft) because it is related to the success of osteoconductivity, osteoinduction and osteogenesis. Osteoconductive and osteoinductive are paramount to resorbable biomaterials to direct and encourage tissue growth formations.[5,7]

In the manufacture of hydroxyapatite, a heating / calcination process has been carried out which can kill various microorganisms but at the time of heating there is a possibility of exposure to the external environment so that additional sterilization process is needed. [8] Sterilization is a process that destroys all forms of life. From a microbiological point of view, a sterile object is free from living microorganisms. In the sterilization process, bacterial spores are the most resistant among all living organisms. Currently, there are various sterilization equipment with different working mechanisms, each of which has its own limitations in its practical application. The method chosen usually depends on the nature of the material to be sterilized. [9] Sterilization of ultraviolet (UV) light is one method that can emit rays with wavelengths between 200-400 nm. A wavelength of 200-300 nm is usually referred to as germicidal because UV light of this wavelength is absorbed by DNA and RNA from microorganisms. Photons absorbed from these UV rays will cause changes in DNA and RNA structure, resulting in microorganisms not being replicable.[10] Based on research by Nishikawa (2003) another effect that can be produced from UV light sterilization on hydroxyapatite is the formation of free radicals (OH and  $O_2$ ) due to the breakdown of  $H_2O$  and  $O_2$  molecules which can cause decomposition of organic pollutants.[11]

To find out the biocompatibility of a material towards human body, a toxicity test is conducted so that it is known how far the adverse effects can be caused, so as not to harm the body. In this study, human gingival fibroblast (HGF) cell culture was used to get fast results and to expose cell cultures directly with experimental materials. Cell cultures are also more sensitive to toxic materials. Thus, not only toxicity can be measured quantitatively, but the response to living cells can be seen. Fibroblasts are often used by researchers to test dental materials, because fibroblasts are one of the constituent components of the tissue around the tooth.[12]

Based on the description above, the purpose of this study is to determine the biocompatibility of hydroxyapatite of *Portunus pelagicus* crab shells that have been sterilized with UV light against HGF cell cultures.

#### 2. Material and methods

#### 2.1. Portunus pelagicus crab shell preparation

This study used a post-test only control group design. The crab population used comes from Semedu Sari, Grati, Pasuruan District. Cleaned crab shells were soaked with 5.25% sodium hypochlorite (Bayclin, SC Johnson) and fed into a furnace engine with an initial temperature of  $\pm$ 50°C. The temperature was raised by 5°C/min until it reaches a temperature of 1000°C for 2 hours. The next process was to sift the powder using a sifting machine to obtain hydroxyapatite powder measuring less than 150 µm particle. The transparent plastic tubes with an additional layer of polyethylene plastic were used as a container to keep hydroxyapatite so that it can be sterilized with UV rays and not contaminated. Hydroxyapatite was sterilized with a UV sterilizer for 2 x 30 minutes by turning it over.

#### 2.2. Human gingival fibroblast cell culture preparation

Human gingival fibroblast cell cultures were given Phosphate-Buffered Saline (PBS) of 2.5 mL to separate or remove the media in the culture, then given the enzyme trypsin as much as 2 mL to release cells in the medium. After that, cell cultures were incubated at a temperature of  $37^{\circ}$ C, CO<sub>2</sub> levels of 5% for 5 minutes. Observations were made to make sure all the cells were detached. Cells were taken using a pipette and incubated again for 5 minutes. Cells that have been detached were given alpha MEM and trypsin medium in a ratio of 1:1 (2 mL each). Then, cells were inserted into the conical tube so that the cell would stick to the tube wall. Several washes were carried out using trypsin and alpha MEM so that the monolayer cells became single cells in the conical tube. Centrifugation of 1800 rpm was carried out for 5 - 6 minutes. The medium and enzymes were removed using a pipette, then alpha MEM was given up to 50 mL. The vibrating process was carried out using a tube mixer and cells were poured into a microplate of 96 well as much as 100  $\mu$ L with a density of 3 – 5x10<sup>3</sup> and incubated for 24 hours with a temperature of 37°C.

#### 2.3. Experimental procedure

This study consisted of 4 groups, control group (without hydroxyapatite, consist of control of media and control of HGF cell), group with 25 ppm of *Portunus pelagicus* crab shell, 50 ppm, and 100 ppm. Each group consisted of 5 samples, was poured into microplate with HGF cells. This microplate underwent UV sterilization for 60 minutes. After UV sterilization, that was underwent incubation for next 24 hour inside the incubator with a temperature of 37°C.

#### 2.4. MTT assay

Tetrazolium salt (MTT) was dissolved in 5 mg/mL PBS. MTT was added directly to the plate containing 10  $\mu$ l of culture medium, then re-incubated for approximately 4 hours at 37°C. The whole medium inside the well was taken out. Then, each well was added DMSO (Dimethylsufoxide) of 50  $\mu$ l. The plate was stirred mechanically with a plate shaker until the formazan crystals dissolved for 5 minutes. Living fibroblast cells will be stained with formazan (turning into blue colour), while the dead cells do not form blue colour. Formazan was read by spectrophotometry with an ELISA reader at a wavelength of 620 nm. The amount of absorbance indicated the number of living cells in the media culture. The calculation of the viability of fibroblast cells in each group was carried out with the formula below:

$$cell \, viability = \frac{OD \, of \, treatment \, group \, + \, OD \, of \, control \, media \, group}{OD \, of \, control \, cell \, group \, + \, OD \, of \, control \, media \, group} \, \times 100\%$$

Notes:

- Cell viability: percentage of living cells after experimental procedure
- OD refers to optical density.
- All variables were taken from mean value of each group

Data obtained was analyzed with Kolmogorov-Smirnov, Levene test, and one-way ANOVA test with confidence level of 95%.

#### 3. Results

Precursor compound found in the *Portunus pelagicus* crab shell that had been used in this study has been observed and the results were shown below:

Table 1 The result of assessment of calcium carbonate in Portunus pelagicus crab shell

Main molecule	Percentage	
Calcium carbonate	61,82%	
Chitin	28,60%	
Phosphate	13,86%	

On the SEM micrograph observation, the fine structure of hydroxyapatite crystal could be seen along with its low porosity, clustered, and the similar size (Figure 1).



Figure 1 SEM micrograph samples of hydroxyapatite *Portunus pelagicus* shell with an enlargement of 8650x with two different field of view

It could be seen the smoothness and homogeneity of the hydroxyapatite structure because the sifting process has been carried out using a sifting machine so that a size of less than 150  $\mu$ m was obtained (Figure 2)



Figure 2 SEM micrograph samples of hydroxyapatite *Portunus pelagicus* shell with an enlargement of 500x with two different field of view

The composition of hydroxyapatite was dominated by oxygen (O)  $67.59\pm6.9\%$ , followed by calcium (Ca)  $29.16\pm0.7\%$ , and phosphorus (P)  $3.25\pm0.1\%$  for normalized weight and oxygen (O):  $83.54\pm6.9\%$ , calcium (Ca):  $14.39\pm0.7\%$ , and phosphorus (P):  $2.08\pm0.1\%$  for the calculation of the number of atoms (Table 2, Figure 3).



Figure 3 The EDX spectrum samples of hydroxyapatite from *Portunus pelagicus* shell with three main constituent elements dominated by Ca, O, and P

Element	Atom number	Weight measurement (wt. %)		Atom measurement (at. %)	Error (%)
		Not normalized	normalized		
0	8	<mark>56,23</mark>	<mark>67,59</mark>	<mark>83,54</mark>	<mark>6,9</mark>
Са	20	<mark>24,25</mark>	<mark>29,16</mark>	<mark>14,39</mark>	<mark>0,7</mark>
Р	15	<mark>2,71</mark>	<mark>3,25</mark>	<mark>2,08</mark>	<mark>0,1</mark>
Total		<mark>83,19</mark>	<mark>100,00</mark>	100,00	

**Table 2** Weight measurement of not normalized, normalized, and atom measurement from three main constituentelements dominated by Ca, O, and P



Figure 4 Left: mapping hydroxyapatite elements using EDX. Right: Combining the three mappings of hydroxyapatite elements using EDX

Figure 4 showed the mapping of Ca, O, and P atomic elements and combination of three of them using EDX on hydroxyapatite. The Ca atom was shown in blue, the O atom was shown in red, and the P atom was shown in green.



Figure 5 Diagram of optical density mean value from each group along with standard deviation

From the data represented by Figure 5 and Table 3, it could be seen that other than control group, the 100 ppm group showed higher living cells, followed by 50 ppm and 25 ppm group. ANOVA test also showed that there was no significant difference between each group.

**Table 3** Percentage of living cells from each group. The numbers below were obtained from the formula mentioned onMaterial and methods section

Group	N	Viability (%)
Control of HGF cell	5	100%
25 ppm	5	93.28%
50 ppm	5	98.10%
100 ppm	5	97.93%

#### 4. Discussion

Hydroxyapatite (HA) is the main mineral component of human hard tissues (especially bones and teeth). It is a storage area to control the absorption and release of calcium from the human body. HA belongs to the class of calcium phosphate-based biochemistry with the chemical formula  $Ca_{10}(PO_4)_6(OH)_2$ .[13] HA can be used as a bone substitute in orthopedics and dental care due to its good biocompatibility and osteoconductive properties.[14]

Crab shells are rich in inorganic and organic materials that can be processed into useful materials. The results of component on *Portunus pelagicus* crab shells originating from the population of Semedu Sari, Grati, Pasuruan Regency: 61.82% CaCO3, 28.60 chitin, and 13.86% PO<sub>4</sub>. CaCO<sub>3</sub> and PO<sub>4</sub> are precursor materials for hydroxyapatite. [13] Based on this study, it showed that the shell of *Portunus pelagicus* has the potential to be extracted into hydroxyapatite.

Characterization of hydroxyapatite samples using SEM-EDX, where the SEM micrograph images (Figure 4) show the formation of small-sized crystals (<150  $\mu$ m), smooth surfaces with low porosity.[15] The small particle size is good for filling bone because it is quickly absorbed, has a larger surface area, and increases osteogenesis.[16] On examination of the EDX spectrum (Table 2) the hydroxyapatite sample was dominated by oxygen (0): 67.59 ± 6.9%, calcium (Ca): 29.16 ± 0.7 %, and phosphorus (P): 3.25 ± 0.1 % for normalized weight and oxygen (0): 83.54 ± 6.9 %, calcium (Ca) : 14.39 ± 0.7 %, and phosphorus (P): 2.08 ± 0.1 % for calculation of the number of atoms. The distribution map (Figure 4) of atoms also shows the evenness of 0, Ca, and P atoms in the hydroxyapatite sample. The composition of these three elements confirms the hydroxyapatite compound.

Hydroxyapatite extract from *Portunus pelagicus* shell was sterilized with UV light to kill microorganisms that can harm surrounding tissues by releasing toxic substances that can damage them. UV light with a wavelength of 200-300 nm is usually referred to as germicidal because UV light with this wavelength is absorbed by the DNA and RNA of microorganisms. Photons absorbed from UV rays will cause changes in the structure of DNA and RNA. [10] Photon energy causes the breaking of hydrogen bonds in nitrogenous bases in DNA or RNA, resulting in chemical modifications of nucleoproteins and causing cross-links between adjacent thymine molecules by covalent bonds. This relationship can cause misreading of the genetic code in the process of protein synthesis, which will produce mutations that will further damage or weaken the vital functions of the organism. The existing electrons will be transferred to the surrounding oxygen and followed by the formation of  $O_2^-$  or superoxide radicals. This reaction occurs due to the mechanism of photoactivation. These superoxide radicals will oxidize organic molecules and react with water molecules and OH<sup>-</sup> ions, produce hydrogen peroxide ( $H_2O_2$ ) The  $H_2O_2$  will further separate and produce OH radicals which oxidize organic molecules (microorganisms) during sterilization. UV sterilization also has the following advantages, namely relatively inexpensive, easy to operate, no chemicals needed, and fast sterilization.[10]

According to Akhila *et al.*, 2007, the determination of a material as toxic or non-toxic was based on Lethal Dose 50 (LD50) which is the first step for assessment and evaluation. [17] It was said that a material is non-toxic or biocompatible if it has cell viability greater than 50%. Of the three results, the percentage of cell viability was above 50% which indicated that the hydroxyapatite graft extract from *Portunus pelagicus* shell which was sterilized by UV light had good biocompatibility towards human gingival fibroblast cells. The treatment with a concentration of 50 ppm had the highest cell viability at 98.10%. The highest cell viability in certain groups can be influenced by differences in the ability of hydroxyapatite receptors on fibroblast cells. In a study conducted by Kasaj et al., it is proven that hydroxyapatite plays a role in stimulating the proliferation of human periodontal fibroblast cells. This is related to the activation of the epidermal growth factor receptor (EFGR) and its subsequent targets ERK1/2 and Akt. Activation of pYFAK 397 also occurs, this can be seen from the increased cellular attachment. The reason for the increased attachment of human periodontal fibroblast cells to hydroxyapatite is an increase in the activation of  $\alpha 5\beta 1$  integrin will cause an increase in fibroblast cell proliferation. Integrins act as

attachment receptors for extracellular matrix proteins transducing signaling pathways through FAK phosphorylation and lead to ERK1/2 activation. This is related to the activation of the epidermal growth factor receptor. Another mechanism that enhances proliferation is activation of the Akt pathway. After the epidermal growth factor receptor is activated, this receptor is phosphorylated at tyrosine 1173, so that Akt increased. The high ability of the epidermal growth factor receptor and the expression of a5ß1 integrin on fibroblast cells in group 2 (50 ppm) caused an increase in fibroblast cell proliferation so that this group had a high number of living fibroblast cells. This is supported by the statement of Yuan et al. that the active epidermal growth factor receptor signaling is responsible for the proliferation of fibroblast cells.[18]

#### 5. Conclusion

Author should provide an appropriate conclusion to the article. Write conclusion as single para. Conclusion should be concise, informative and can be started with summarizing outcome of the study in 1-2 sentence and ended with one line stating: how this study will benefit to the society and way forward.

#### **Compliance with ethical standards**

#### Acknowledgments

The authors would like to say thank you to Jennifer Widjaja for her assistance to review and editing this manuscript thoroughly.

#### Disclosure of conflict of interest

If two or more authors have contributed in the manuscript, the conflict of interest statement must be inserted here.

#### References

- [1] Supartono W, Rakhmadhani NR P. Analysis of Rejection of Indonesian Export Products Blue Swimming Crab (*Portunus pelagicus*) and Crab (Scylla serrata) in the United States for the Period of 2002 -2013. Prosiding Seminar Agroindustri Dan Lokakarya Nasional FKPT-TPI 2015:27–32.
- [2] Rochima E. Characterization of Chitin and Chitosan from Crab Waste in Cirebon, West Java. Indonesian Fisheries Processing Journal 2007;10. https://doi.org/10.17844/JPHPI.V10I1.965.
- [3] Rahayu L, Purnavita S. Optimization of Chitosan Manufacture from Chitin Crab Shell Waste (*Portunus pelagicus*) for Mercury Metal Ion Adsorbents. Reaktor 2017;11:45. https://doi.org/10.14710/REAKTOR.11.1.45-49.
- [4] Asni N, Saadilah MA, Saleh D. Optimization of Chitosan Synthesis from Crab Shell as Adsorbent of Heavy Metal Pb(II). Spektra: Jurnal Fisika Dan Aplikasinya 2014;15:18–25.
- [5] Khiri MZA, Matori KA, Zainuddin N, Abdullah CAC, Alassan ZN, Baharuddin NF, et al. The usability of ark clam shell (Anadara granosa) as calcium precursor to produce hydroxyapatite nanoparticle via wet chemical precipitate method in various sintering temperature. Springerplus 2016;5. https://doi.org/10.1186/S40064-016-2824-Y.
- [6] Drismayanti I, T S, B AU, Ruslin Muh, Jubhari EH. Ratio of successful rate of dental implant between distraction osteogenesis with autogenous bone graft. Journal of Dentomaxillofacial Science 2012;11:180. https://doi.org/10.15562/JDMFS.V11I3.335.
- [7] Ichsan MZ, Siswanto S, Hikmawati D. Synthesis of Macropores Collagen Hydroxyapatite Composite as a Candidate for Bone Graft. Jurnal Fisika Dan Terapannya 2013.
- [8] Troy D, Beringer P. Remington: The Science and Practice of Pharmacy. 21st ed. Philadelphia: Lippincott Williams & Wilkins; 2006.
- [9] Adji D, Zulianti Z, Larashanty H. Comparison of the Effectiveness of 70% Alcohol, Infrared, Autoclave and Ozone Sterilization against the Growth of Bacillus subtilis Bacteria. Jurnal Sain Veteriner 2007;25. https://doi.org/10.22146/JSV.275.
- [10] Bolton JR, Cotton CA. The Ultraviolet Disinfection Handbook. 1st ed. American Water Works Association; 2008.

- [11] Nishikawa H. Radical generation on hydroxyapatite by UV irradiation. Mater Lett 2004;58:14–6. https://doi.org/10.1016/S0167-577X(03)00396-3.
- [12] Ariani MD, Yuliati A, Adiarto T. Toxicity testing of chitosan from tiger prawn shell waste on cell culture. Dental Journal (Majalah Kedokteran Gigi) 2009;42:15–20. https://doi.org/10.20473/j.djmkg.v42.i1.p15-20.
- [13] Zhang S. Biological and Biomedical Coatings Handbook: Applications Google Books. Taylor & Francis; 2011.
- [14] Mucalo MR. Hydroxyapatite (HAp) for Biomedical Applications. Hydroxyapatite (HAp) for Biomedical Applications, Elsevier Ltd; 2015, p. 1–381. https://doi.org/10.1016/C2013-0-16440-9.
- [15] Srivastava A, Jain VK, Srivastava A. SEM-EDX analysis of various sizes aerosols in Delhi India. Environmental Monitoring and Assessment 2008 150:1 2008;150:405–16. https://doi.org/10.1007/S10661-008-0239-0.
- [16] Higashi T, Okamoto H. Influence of particle size of hydroxyapatite as a capping agent on cell proliferation of cultured fibroblasts. J Endod 1996;22:236–9. https://doi.org/10.1016/S0099-2399(06)80139-1.
- [17] Akhila JS, Shyamjith D, Alwar M. Acute Toxicity Studies and Determination of Median Lethal Dose. Curr Sci 2007;93:917–20.
- [18] Kartono GS, Widyastuti W, Setiawan HW. Biocompatibility of Hydroxyapatite Graft from Blood Mussel Shells (Anadara granosa) Against Fibroblast Cell Culture. Denta Jurnal Kedokteran Gigi 2014;8:1–8. https://doi.org/10.30649/DENTA.V9I2.8.



#### WJARR-2023-0438 Article proof for final check and approval

**michael josef kridanto kamadjaja** <michael-j-k-k@fkg.unair.ac.id> To: editor@wjarr.com

Dear Managing Editor World Journal of Advaned Research and Reviews (WJARR)

I would like to re-send the revised file of the manuscript along with the author's query form. Should you have any other queries regarding this manuscript, please let me know

Best regards, Michael Josef Kridanto Kamadjaja [Quoted text hidden]

#### 2 attachments

WJARR-Authors queries for galley proof form.docx 91K

WJARR-2023-0438 Article Proof.docx 1411K Mon, Mar 20, 2023 at 1:09 PM



#### **Author Queries**

Title of the Manuscript: Biocompatibility of crab shell-derived graft sterilized by ultraviolet light towards human gingival fibroblast cell culture

Queries	Details Required	Author's Response
AQ1	Please check and confirm the type of the article.	Checked &
AQ2	Please check and confirm the title of the article.	Checked & confirmed
AQ3	Please check and confirm the spellings of all the author names, their sequence, affiliations, corresponding author is correctly indicated and details (e-mail, phone)	Corrected & confirmed
AQ4	Please check and confirm that all the keywords given by author are mentioned and correctly spelled.	Checked & confirmed
AQ5	Please check whether all heading and subheading levels are okay.	Checked & confirmed
AQ6	Please check and confirm that all spellings and grammar in the entire text are correctly used.	Checked & confirmed
AQ7	Please check and confirm that all the punctuation marks are appropriately given.	Corrected & confirmed
AQ8	Please check and confirm that all the scientific nomenclature, units, symbols and abbreviations are correctly presented	Corrected & confirmed
AQ9	Please check and confirm that all Tables and Figures are included in the manuscript.	Checked & confirmed
AQ10	Please check and confirm that all the caption of Tables and Figures are correctly presented.	Checked & confirmed
AQ11	Please check and confirm that all footnotes of tables are okay.	Checked & confirmed
AQ12	Please check and confirm that all the formulae (if present) are correctly presented.	Checked & confirmed
AQ13	Please check and confirm that statements about 'Compliance with Ethical Standards' and correctly presented before reference section. These include (i) Acknowledgement (compulsory), (ii) Disclosure of conflict of interest (compulsory), (iii) Statement of ethical approval (if any) and (iv) Informed consent (if any).	Checked & confirmed
AQ14	Please check and confirm that all the References cited in the text are given in the reference list and vice versa. Please provide details in the list or delete the citation from the text if applicable	Checked & confirmed
AQ15	Please confirm whether author want to publish short biography (4-5 lines) or not? If yes, confirm author has provided short biography along with Passport size photograph at the end of the manuscript.	No

Author should check all above queries. If okay then mention "Checked and Confirmed" in the Author's Response column; otherwise, do the necessary changes in the galley proof (in



blue color text) and mention "Checked and Corrected" in the Author's Response column.



#### WJARR-2023-0438 Article proof for final check and approval

editor@wjarr.com <editor@wjarr.com> To: michael josef kridanto kamadjaja <michael-j-k-k@fkg.unair.ac.id> Tue, Mar 21, 2023 at 3:34 PM

#### Dear Author, Greetings for the day.

We received authors proof approved by you. Very soon we will send you article publishing details.

Thanks and Regards Managing Editor

[Quoted text hidden]

[Quoted text hidden]



# Your Manuscript has been published in Volume 17 - Issue 3 (March 2023) of World Journal of Advanced Research and Reviews (WJARR).

**World Journal of advanced Research and Reviews** <editor@wjarr.com> To: michael-j-k-k@fkg.unair.ac.id Wed, Mar 22, 2023 at 12:06 PM

Dear Author, Greetings for the day...

We are pleased to inform you that your manuscript entitled "Biocompatibility of Crab Shell-Derived Graft Sterilized by UV Light towards Human Gingival Fibroblast Cell Culture" has been published in Volume 17 - Issue 3 (March 2023) of World Journal of Advanced Research and Reviews (WJARR).

#### You can view or download your article from the below links

https://wjarr.com/content/biocompatibility-crab-shell-derived-graft-ster...

https://wjarr.com/sites/default/files/WJARR-2023-0438.pdf

Crossref DOI and CrossMark metadata information will be updated very soon. We will inform you regarding this.

I hope you are happy with our services and have good experience while publishing with us. We will be happy to review and publish you next manuscript in this journal. You may also consider our other allied journals...

World Series Journals GSC Online Press Journals Magna Scientia Journals International Research Journals Open Access Research Journals

Visit these journals at: https://wjarr.com/content/our-associated-journals

We will look forward to the submission of your next manuscripts.

With Best Regards, Managing Editor World Journal of Advanced Research and Reviews (WJARR) editor@wjarr.com