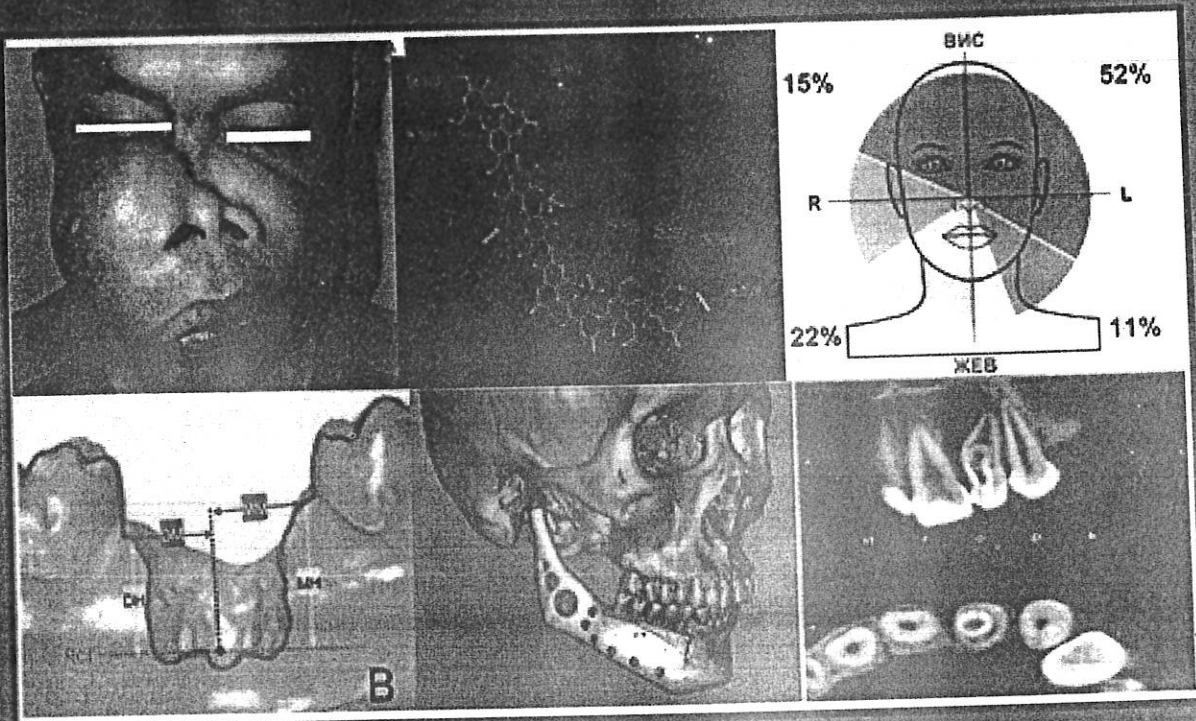


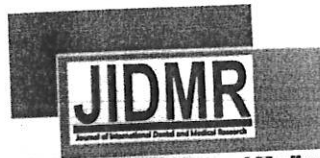
E-ISSN: 1309-100X

Journal of
International
Dental and Medical
Research



2021 - Vol. 14 - No. 1

<http://www.jidmr.com>



Journal of International Dental and Medical Research



Editorial Board of JIDMR

Prof. Dr. Izzet YAVUZ
Editor-in-Chief and General Director

Advisory Board

Prof. Dr. Refik ULKU Editor for Medicine
Prof. Dr. Zulkuf AKDAG Editor for Biomedical Research
Prof. Dr. Ozkan ADIGUZEL Associate Editor

Gajanan Kiran KULKARNI (CANADA)
Betul KARGUL (TURKEY)
Diah Ayu MAHARANI (INDONESIA)
Francisco Cammarata-Scalisi (Venezuela)
Myroslav Goncharuk-Khomyn (UKRAINE)

Ferranti WONG (UK)
Zeki AKKUS (TURKEY)
Michele CALLEA (ROME, ITALY)
Zelal ULKU (TURKEY)

Mosc
Linda
Yase
Yuliy
Nik N

Editorial Board

Abdel Fattah BADAWI (EGYPT)
Abdurrahman ONEN (TURKEY)
Ahmet YALINKAYA (TURKEY)
Ahmet DAG (TURKEY)
Ali Al-ZAAG (IRAQ)

Guvenc BASARAN (TURKEY)
Guvenc ERBIL (TURKEY)
Halimah AWANG (MALAYSIA)
Halit AKBAS (TURKEY)
Heloisa Fonseca MARAO (BRAZIL)

Nezahat A
Nihal HAM
Nik Noriah
Nicola Pra
Nurten AK

Ali BUMIN (TURKEY)	Hilal TURKER (TURKEY)	Nurten ER
Ali FADEL (EGYPT)	Huseyin ASLAN (TURKEY)	Orhan TAC
Ali GUR (TURKEY)	Igor BELYAEV (SWEDEN)	Ozant ONI
Ali Kemal KADIROGLU (TURKEY)	Ilhan INCI (ZURICH)	Ozgur UZI
Ali Riza ALPOZ (TURKEY)	Ilker ETIKAN (TURKEY)	Ozkan AD
Ali Riza Tunçdemir (TURKEY)	Isil TEKMEN (TURKEY)	Pafat Ali S
Allah Bakhsh HAAFIZ (USA)	Isin ULUKAPI (TURKEY)	Refik ULK
Alpaslan TUZCU (TURKEY)	Jalen DEVECIOGLU KAMA (TURKEY)	Sabiha Ze
Alpen ORTUG (TURKEY)	Kemal CIGDEM (TURKEY)	Sabri BATI
Armelia Sari WIDYARMAN (INDONESIA)	Kemal NAS (TURKEY)	Sadullah K
Ashish AGGARWAL (INDIA)	Kewal KRISHAN (INDIA)	Saul Marti
Ayşe GUNAY (TURKEY)	King Nigel MARTYN(HONG KONG, CHINA)	Sedat AKI
Aziz YASAN (TURKEY)	Kursat ER (TURKEY)	Seher GUI
Balasubramanian MADHAN (INDIA)	Levent ERDINC (TURKEY)	Selahattin
Benik HARUTUNYAN (ARMENIA)	Luca TESTARELLI (ROME)	Selahattin
Betul KARGUL (TURKEY)	Lucianne Cople MAIA (BRAZIL)	Serdar ER
Betul URREHMAN (UAE)	Luciane Rezende COSTA (BRAZIL)	Serdar ON
Bugra OZEN (TURKEY)	Marri Sai ARCHANA (INDIA)	Sergio Adr
Carlos Menezes AGUIAR (BRAZIL)	Manoj KUMAR (INDIA)	Serhan Ak
Cemil SERT (TURKEY)	Marcelo Rodrigues AZENHA (BRAZIL)	Sertac PE
Chiramana SANDEEP (INDIA)	Marcia Cancado FIGUEIREDO (BRAZIL)	Seyed Am
Christine Bettina STAUDT (SWITZERLAND)	Marco MONTANARI (ITALY)	Seyit Burh
Cihan AKGUL (TURKEY)	Margaret TZAPHLIDOU (GREECE)	Shailesh L
Claudia DELLAVIA (ITALY)	Maria Elisa Oliveira dos SANTOS (BRAZIL)	Sinerik N.
Diah Ayu MAHARANI (INDONESIA)	Medi GANIBEGOVIĆ (BOSNIA and HERZEGOVINA)	Smaragda
Dinesh Rokaya (NEPAL)	Mehmet DOGRU (TURKEY)	Sossani S
Edoardo BAUNER (ROMA)	Mehmet Emin ERDAL (TURKEY)	Stefano Di
Emmanuel Joao N. Leal da SILVA (BRAZIL)	Mehmet Sinan DOGAN (TURKEY)	Sunit Kr. J
Emin Caner TUMEN (TURKEY)	Mehmet Ünal (TURKEY)	Stephen D
Emrullah BAHSI (TURKEY)		Susumu T

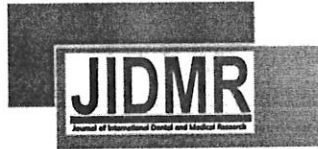
Ertunc Dayı (TURKEY)	Mehmet Zulkuf AKDAG (TURKEY)	Suha TUR
Fadel M. ALI (EGYPT)	Meral ERDINC (TURKEY)	Suleyman
Fahinur ERTUGRUL (TURKEY)	Michele CALLEA (ITALY)	Taskin GU
Feral OZTURK (TURKEY)	Mohamed TREBAK (USA)	Ufuk ALUC
Feridun BASAK (TURKEY)	Mohammad Khursheed Alam (KSA)	Ugur KEKI
Ferranti WONG (UNITED KINGDOM)	Mohammed Mustahsen UKREHMAN (UAE)	Xiong-Li Y.
Feyzi Çelik (TURKEY)	Moschos A. PAPADOPOULOS (GREECE)	Vatan KAV
Feyzullah Uçmak (TURKEY)	Mostaphazadeh AMROLLAH (IRAN)	Yasar YILI
Figen SEYMEN (TURKEY)	M.S. Rami REDDY (INDIA)	Yasemin Y
Filippo BATTELLI (ITALY)	Muhammad FAHIM (INDIA)	Yavuz SAI
Filiz Acun KAYA (TURKEY)	Mukadder ATMACA (TURKEY)	Yu LEI (US)
Flavio Domingues Das NEVES (BRAZIL)	Murat AKKUS (TURKEY)	Yuri LIMAI
Folakemi OREDUGBA (NIGERIA)	Murat SOKER (TURKEY)	Zafer C. C
Francesca De Angelis (ITALY)	Mustafa KELLE (TURKEY)	Zeki AKKL
Gajanan Kiran KULKARNI (CANADA)	Mustafa ZORTUK (TURKEY)	Zeynep Aı
Gamze AREN (TURKEY)	Muzeyyen YILDIRIM (TURKEY)	Zuhal KIRI
Gauri LELE (INDIA)	Neval Berrin ARSERIM (TURKEY)	Zurab KOI
Gonul OLMEZ (TURKEY)		
Gulsen YILMAZ (TURKEY)		
Gulten UNLU (TURKEY)		

Design: TwTDizayn

www.jidmr.com

www.journalofinternationaldentalandmedicalresearch.com





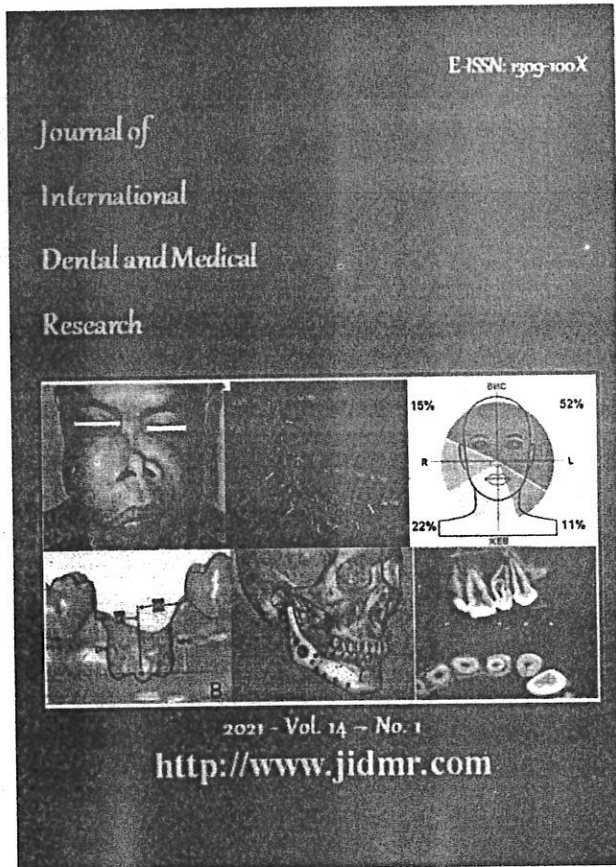
Journal of International Dental and Medical Research

Contents of JIDMR-2021-Vol.14- No.1

31/03/2021 /

Journal of International Dental and Medical Research

ISSN: 1309-100X




Download Cover page

 [http://www.jidmr.com/journal/wp-content/uploads/2021/03/Cover-2021.14.1-1.jpg]

Current Issue of JIDMR

Table of Contents 2021 Vol.14–No.1


 [http://www.jidmr.com/journal/wp-content/uploads/2021/03/1_1_14_21_Table_of_Contents.pdf]

DENTISTRY

EXPERIMENTAL ARTICLE

1. Efficiency of the Algorithm of Examination, Treatment and Rehabilitation of Dental Patients with Comorbid Pathology


Viktoriya N. Naumova, Yuliya A. Makedonova, Dmitriy V. Mikhailchenko, Tatyana V. Kolesova, Olga Yu. Afanaseva, Elena E. Maslak, Hosityain Yu. Salyamov

Pages 1-4 **Full Text**  [http://www.jidmr.com/journal/wp-content/uploads/2021/03/1-E-D21_1407_Yuliya_A_Makedonova_Russia.pdf]

EXPERIMENTAL ARTICLE

2. Effects of an Obesogenic Diet on Craniofacial Morphology in Rats


Daniela Botero-González, María Carolina Pustovrh, Mario Ortiz, Adriana María Herrera-Rubio

Pages 5-11 **Full Text**  [http://www.jidmr.com/journal/wp-content/uploads/2021/03/2-E-D20_1293_Daniela_Botero_Gonzalez_Colombia.pdf]

EXPERIMENTAL ARTICLE

3. Chewing gum with added chitosan reduces the number of cariogenic bacteria colonies in human saliva

Magdalena Nowosielska, Joanna Baginska, Anna Kierklo

Pages 12-16 **Full Text**  [http://www.jidmr.com/journal/wp-content/uploads/2021/03/3-E-D20_1277_Magdalena_Nowosielska_Poland.pdf]

EXPERIMENTAL ARTICLE

Pages 384-393 **Full Text PDF** [http://www.jidmr.com/journal/wp-content/uploads/2021/03/60-R-D20_1265_Sharul_Nisha_Ali_Malaysia.pdf]

REVIEW

61. COVID-19: Unraveling 10 Most Significant Answers about The Current Pandemic

Asmaa Tahseen Uthman, Noor Natheer Al-Rawi, Bidaa Othman, Natheer Hashim Al-Rawi

Pages 394-403 **Full Text PDF** [http://www.jidmr.com/journal/wp-content/uploads/2021/03/61-R-D20_1218_Noor_Al_Rawi_UAE.pdf]

MEDICINE

EXPERIMENTAL ARTICLE

62. Biocompatibility Testing of Hydroxyapatite-Chitosan-Chondroitin Sulfate Composite Scaffold as Bone Graft

Dolfi Varton, Prihartini Widiyanti, Sri Puji Astuti Wahyuningsih, Aminatun

Pages 404-411 **Full Text PDF** [http://www.jidmr.com/journal/wp-content/uploads/2021/03/62-E-M20_1222_Aminatun_Indonesia.pdf]

EXPERIMENTAL ARTICLE

63. Effects of Addition of Gonadotropin in the Media of Oocyte Maturation on the Embryo Cleavage

63. A. A. M. N. Kasman, B. Santoso, E. M. Luqman, Widjiati

Pages 412-418 **Full Text PDF** [http://www.jidmr.com/journal/wp-content/uploads/2021/03/63-E-M20_1231_Widjiat_Indonesia.pdf]

EXPERIMENTAL ARTICLE

64. Nanocurcumin Protective Effect on Gpx Scavenger Enzyme Expression and Apoptosis of Lead Acetate-Induced Rats Ovarian Granulosa Cells

Anis Satus Syarifah, Sri Agus Sudjarwo, Hendy Hendarto, Reny I'tishom, Supriyanto

Nanocurcumin Protective Effect on Gpx Scavenger Enzyme Expression and Apoptosis of Lead Acetate-Induced Rats Ovarian Granulosa Cells

Anis Satus Syarifah^{1*}, Sri Agus Sudjarwo², Hendy Hendarto³, Reny I'tishom⁴, Supriyanto⁵

1. Assistant Professor, Nursing Professional Education Program, STIKES Pemkab Jombang, Indonesia.
2. Professor, Department of Pharmacology, Faculty of Veterinary Medicine, Universitas Airlangga, Surabaya, Indonesia.
3. Professor, Department of Obstetrics and Gynecology, Faculty of Medicine, Universitas Airlangga, Surabaya, Indonesia.
4. Associate Professor, Department of Medical Biology, Faculty of Medicine, Universitas Airlangga, Surabaya, Indonesia.
5. Assistant Professor, Department of Nursing, Politeknik Kesehatan Kemenkes Surabaya, Indonesia.

Abstract

This study aims to prove the protective effect of nanocurcumin against the reduction of glutathione peroxidase (GPx) scavenger enzyme and the increase of apoptosis in lead acetate-induced rat ovarian granulosa cells.

A total of 45 female white rats were divided into 5 groups, the negative control group (receiving corn oil and one hour later receiving distilled water), positive control group (receiving corn oil and one hour later receiving lead acetate (Pb) of 40 mg/kg bw), and experimental groups 1, 2 and 3 (receiving nanocurcumin 50, 100 and 200 mg/kg, and one hour later receiving lead acetate (Pb) 40 mg/kg bw). All groups received treatments per oral once a day for 26 days. On day 27 the rats were sacrificed and then the expression of the GPx scavenger enzyme and apoptosis was performed by immunohistochemistry.

Lead acetate (Pb) decreased the expression of the GPx scavenger enzyme and increased apoptosis. The administration of nanocurcumin increased the expression of GPx scavenger enzyme and decreased the expression of apoptosis in lead acetate-induced rat ovarian granulosa cells.

Nanocurcumin functions as an antioxidant by providing protective effect against GPx scavenger enzyme and apoptosis in lead acetate (Pb)-induced rat granulosa cells.

Experimental article (J Int Dent Med Res 2021; 14(1): 419-425)

Keywords: Antioxidants; lead acetate; scavenger enzymes and apoptosis; nanocurcumin.

Received date: 30 August 2020

Accept date: 07 November 2020

Introduction

Lead (Pb) is an easily found toxic heavy metal, one of environmental pollutants which can affect the function of organs such as testes, liver, brain, hematopoietic and kidneys¹. Accumulation of lead in the body may cause toxicity and potentially affect the reproductive system². Lead exposure can also cause abortion, preterm birth and fetal death³ and has been shown to cause rat's nephron toxicity⁴ and testicular toxicity⁵.

As a part of heavy metals, lead acetate has a tendency to catalyze oxidation reactions and lead to the formation of reactive oxygen species (ROS)^{6,7}. If exposure to ROS is not able

to be balanced by scavenger enzyme, where ROS ratio is higher than that of antioxidant enzymes, cells experience oxidative stress which causes lipid peroxidation and DNA damage which will end in apoptosis and decreased endogenous antioxidant defense systems in cells⁸. Endogenous antioxidants known as scavenger enzymes include the GPx enzyme⁹. GPx is the most important enzyme as an endogenous antioxidant that can scavenge free radicals. GPx is a first-line defense antioxidant and requires selenium to perform its function, so it is also called a selenoprotein antioxidant¹⁰. Glutathione peroxidase (GPx) is an important intracellular enzyme that breaks down hydrogen peroxide (H₂O₂) into water (H₂O) and lipid peroxides into alcohol¹¹.

Pb has effect on antioxidant enzyme GPx because Pb has a high affinity and reactivity for selenium, which is a component of the GPx enzyme, so that it can act as an antioxidant. The binding of selenium to Pb will cause glutathione

*Corresponding author:

Anis Satus Syarifah, S.Kep., Ns., M.Kes
Nursing Professional Education Program
STIKES Pemkab Jombang, Indonesia
E-mail: syarifah_anissatus@yahoo.co.id

peroxidase to lose its ability as an antioxidant¹².

Lead acetate exposure causes increased apoptosis of ovarian granulosa cells through oxidative stress mechanisms. Hydroxyl radicals (OH*) formed due to lead exposure can translocate into ovarian granulosa nuclei and stimulate the release of P53. P53 reacts with mitochondrial membrane and activates pro-apoptosis (Bax) and causes decreased anti-apoptosis (Bcl-2 and Bcl-x) which makes cytochrome c release to granulosa cell cytosol. In the cytosol, cytochrome c binds to Apaf-1 (apoptosis-activating factor 1) to form a caspase recruitment domain (CARD) which stimulates caspase 9 granulosa cells and caspase 9 will stimulate caspase-3, an effector that carries out granulosa cell apoptosis¹⁰.

In protecting against the reduction of scavenger enzymes, including GPx, and the occurrence of apoptosis due to ROS, body's cells require exogenous antioxidants, such as curcumin rhizome (*Curcuma longa*). Curcumin has benefits as anti-bacterial, anti-inflammatory, chemopreventive, wound healing, anti-parasitic, and has a phenolic group that has great potential as an antioxidant substance¹³.

Analysis of the antioxidant activity of curcumin also showed that curcumin had considerable antioxidant activity¹⁴. Several studies have shown that curcumin can reduce oxidative stress due to lead toxicity by inhibiting oxidative stress very effectively¹⁵. but its clinical application is still limited, both in vascular and oral administration.

The limitation of clinical application of curcumin is due to its poor solubility and absorption, leading to its low bioavailability¹³. Orally administered curcumin absorption undergoes presystemic elimination. Once absorbed, curcumin is conjugated by sulfates and glucuronates at various tissue sites. The poor absorption pattern makes it difficult to find high levels of curcumin in blood some time after administration, so the effect is less effective¹⁶. With innovations to improve bioavailability, longer circulation, and better permeability, curcumin is formulated in the form of nanoparticles. This study aims to investigate the protective effect of nanocurcumin against the reduction of the GPx scavenger enzyme and the increase in lead acetate-induced apoptosis of white rats' granulosa cells.

Materials and methods

Chemical materials

Lead acetate (Product No: CAS 6080-56-4, molecular weight (MW): 379.33 g/mol, Linear formula: $Pb(CH_3CO_2)_2 \cdot 3H_2O$), purchased from Sigma-Aldrich.co. USA and Curcumin (*Curcuma longa* (turmeric) powder, $\geq 65\%$, Product No: CAS 458-37-7 molecular weight (MW): 368.38 g/mol, product of China).

2.1. Making process of nanocurcumin

The method used for the making of nanocurcumin is by milling using a High Energy Milling (HEM) machine. Nanocurcumin was made at the Physics Laboratory, Universitas Airlangga, Surabaya, Indonesia. The milling time was 20 minutes with a setting of 5 minutes milling, 5 minutes resting until total effective processing time of 20 minutes outside the resting time. Curcumin powder from the rhizome of *Curcuma longa* (turmeric) was milled by pulverizing cubic zirconia balls, with the ratio of curcumin : cubic zirconia balls = 1 : 10. Curcumin and cubic zirconia balls were fed into the tubes and milled in a High Energy Milling (HEM) machine.

2.2. Analysis of nanocurcumin characteristics

Analysis of nanocurcumin size characteristics used the Scanning Electron Microscopy (SEM) method and was carried out at the Robotics Laboratory, ITS, Surabaya, Indonesia. The morphology of curcumin after milling was compared with that of before. The morphology of curcumin after milling showed a more regular crystal shape with an average diameter of less than 200 nm. The morphology of curcumin before milling appeared as irregular plates with a mean diameter of more than 1000 nm as shown in Figure 1.

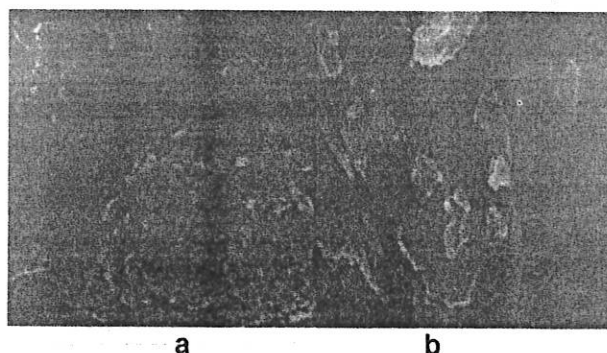


Figure 1. The morphology of nanocurcumin by scanning electron microscopy (SEM). (a) Curcumin after milling with a more regular crystal form with a diameter of < 200 nm. (b) Curcumin before milling with irregular slab shape, diameter > 1000 nm.

2.3. Preparation of nanocurcumin solution

A solution was made by dissolving 2 grams of nanocurcumin with corn oil into 200 ml, so that 1 ml of the solution contained 10 mg of nanocurcumin. Nanocurcumin was diluted with corn oil, because corn oil was the best carrier compared to butter, milk and water¹⁷.

Experimental Animals

Female wistar rats weighing about 180-200 g, aged 2.5 - 3 months were obtained from Institut Teknologi Bandung (ITB), Bandung, Indonesia, for experimental purposes. The rats were placed in cages in an air-conditioned room with temperatures maintained at 26°C - 2°C and 12 hours in a light and dark cycle. Rats were given with feed and mineral water ad libitum. This study has passed the ethical test by the Ethics Committee, Faculty of Veterinary Medicine, Universitas Airlangga, Indonesia, and obtained ethical eligibility No: 2.KE.170.08.2019 on 29 August 2019.

Experimental design

A total of 45 female rats were divided into 5 groups, the negative control group (rats receiving corn oil, one hour later receiving distilled water), positive control group (rats receiving corn oil, one hour later receiving lead acetate of 40 mg/kg bw), and experimental groups 1, 2 and 3 (rats receiving 50 mg, 100 mg and 200 mg/kg bw of nanocurcumin orally, one hour later receiving lead acetate of 40 mg/kg). All groups received treatment orally once a day for 26 days. On day 27 the rats were sacrificed with neck dislocation, then the peritoneum was excised, the ovaries were taken to examine the expression of GPx scavenger enzyme and the apoptosis of granulosa cells by immunohistochemical examination.

Immunohistochemical examination

The removed ovaries were implanted in a paraffin block, then sliced, and a representative slice was selected from the tissue sample for immunostaining procedure. Each slice was stained with streptavidin method using immunoperoxidase. Serial cutting of paraffin blocks was carried out in a thickness of 4 - 6 µm.

The best slices were selected to examine the expression of GPx scavenger enzyme and apoptosis at 400X magnification. The presence of the GPx scavenger enzyme and apoptosis was characterized by dark brown color intensity. Observations were made quantitatively by counting the number of positive cells per visual

field, counting up to 10 fields.

The numbers of positive cells for each visual field were summed up and divided by 10, and the final result was the mean number of positive cells per visual field.

Statistical analysis

Data were presented with mean ± standard deviation. The comparative test used Kruskal-Wallis Test to determine the differences between groups, followed by Mann-Whitney test to determine the differences between the groups.

Results

Nanocurcumin protection against lead acetate-induced reduction of GPx scavenger enzyme expression in rats ovarian granulosa cells

The results of Kruskal-Wallis test showed differences in the expression of GPx scavenger enzyme (Kruskal-Wallis H = 21,787; df = 4; p = .000). Then, to identify differences between groups, we used Mann-Whitney Test, as in Table 1.

Groups	n	GPx expression (%/micro)	Minimum	Maximum
Negative control	9	3.5 ± 0.5 ^a	2.9	4.5
Positive control	8	2.4 ± 0.3 ^b	1.7	2.8
Experimental 1	9	2.5 ± 0.2 ^b	2.3	2.9
Experimental 2	9	3.2 ± 1.0 ^a	2.1	5.4
Experimental 3	8	3.5 ± 0.6 ^a	2.6	4.3

Table 1. Effect of nanocurcumin on GPx scavenger enzyme expression of lead acetate-induced rat ovarian granulosa cells (mean ± standard deviation).

^{a,b}Different superscript within each column indicates significant difference between the means (p < .05).

Table 1 shows that the mean expression of GPx scavenger enzyme of granulosa cells in rats' ovaries was the highest in the experimental group 3 (3.5 ± 0.6%/micro), the same (p = .481) with that of negative control group (3.5 ± 0.5%/micro) and also the same (p = .139) with that of experimental group 2 (3.2 ± 1.0%/micro). The lowest value in control group was positive (2.4 ± 0.3%/micro) and the same (p = .263) with that of the experimental group 1 (2.5 ± 0.2%/micro).

These results indicated that the nanocurcumin at a dose of 100 mg/kg bw and 200 mg/kg bw increased the expression of GPx scavenger enzyme in lead acetate-induced rat

ovarian granulosa cells. This difference is also shown in Figure 2.

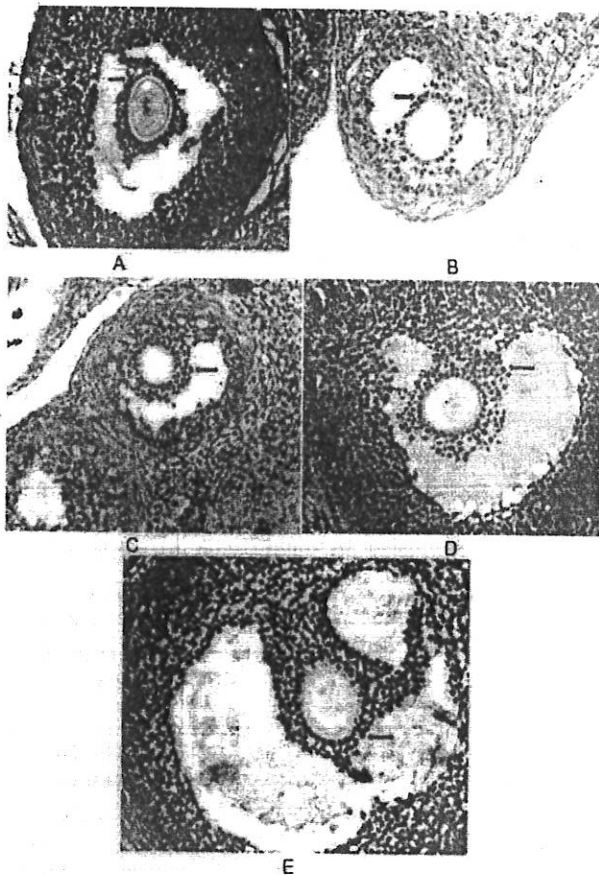


Figure 2. Comparison of GPx scavenger enzyme expression in rats' ovarian granulosa cells. (A) K- group; (B) K+ group; (C) P1 group; (D) P2 group; (E) P3 group. Observation used light microscope in a magnification of 400x. Arrow (➡) shows an example of GPx scavenger enzyme expression.

Groups	n	Apoptosis expression (%/micro)	Minimum	Maximum
Negative control	9	2.2 ± 0.8 ^a	1.0	3.5
Positive control	8	4.3 ± 0.8 ^b	2.7	5.6
Experimental 1	9	3.4 ± 1.0 ^c	2.1	4.8
Experimental 2	9	2.4 ± 0.8 ^a	1.6	3.8
Experimental 3	8	2.0 ± 1.2 ^a	1.2	4.9

Table 2. Effect of nanocurcumin on the expression of apoptotic granulosa in ovaries of lead acetate-induced rats (mean ± standard deviation).

^{a,b,c}Different superscript within each column indicates significant difference between the means ($p < .05$).

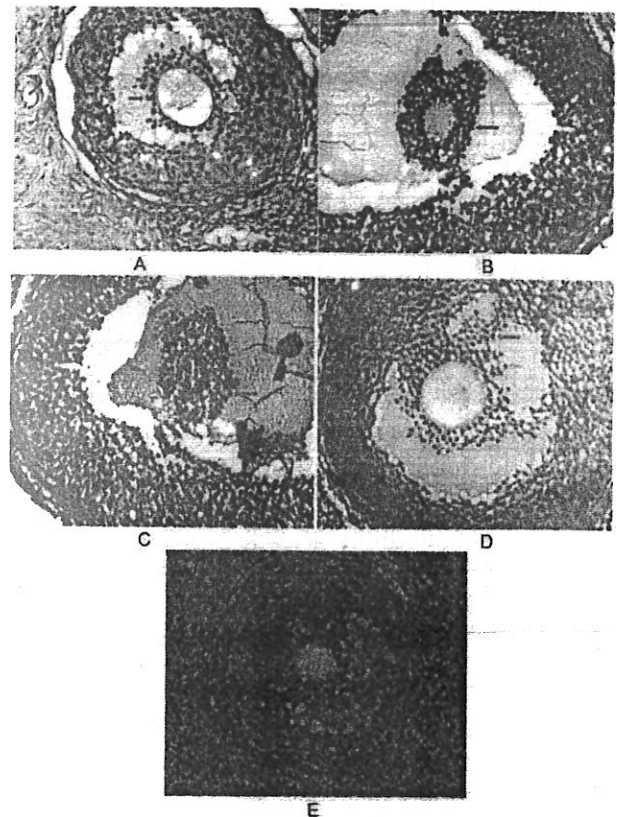


Figure 3. Comparison of the expression apoptotic of rats' ovarian granulosa cells with 400 x magnification. (A) K- group; (B) K+ group; (C) P1 group; (D) P2 group; (E) P3 group. Observation using a light microscope with a magnification of 400x. Arrow (➡) shows one example of apoptotic expression.

Discussion

The results showed that exposure to lead acetate as much as 40 mg/kg bw reduced the expression of GPx scavenger enzyme in rats' ovarian granulosa cells. The decrease in the expression of GPx scavenger enzyme due to lead acetate induction was in line with previous studies. Rats receiving lead acetate injection of 20 mg/kg bw/day intraperitoneally for 5 days experienced a decrease in renal GPx compared to control group receiving distilled water injection¹⁸. Rats injected with lead acetate of 20 mg/kg bw for 11 days had a lower mean testicular GPx than control group injected with distilled water alone⁵. Rats that were given with normal saline (0.9% NaCl) of 3 ml/kg bw/day orally and one hour later injected intraperitoneally with lead acetate of 20 mg/kg bw/day for 7 days experienced a significant reduction in hepatic GPx compared to

that of control group with the same treatment and received a normal saline injection of 1 ml/kg bw/day¹⁹

As the first line defense besides SOD and caspase-3, GPx is the most important enzyme as an endogenous antioxidant capable of scavenging free radicals. As an antioxidant, GPx requires selenium to carry out its functions, so it is also called a selenoprotein antioxidant. GPx breaks down hydrogen peroxide (H₂O₂) as a result of ROS into water (H₂O); and lipid peroxides into alcohol¹⁰.

Pb has high affinity and reactivity to selenium, a component of GPx, so that it can act as an antioxidant. The binding of selenium to Pb causes GPx to lose its ability as an antioxidant. The increase in Pb in the body will cause the binding of Pb with selenium. As a result, the availability of selenium in the body decreases. The reduced availability of selenium causes GPx not to act as an antioxidant²⁰. This was proved by the decrease in the expression of the enzyme GPx in rats ovarian granulosa cells induced with lead acetate of 40 mg/kg bw/day by sonde.

This study also proved that the administration of nanocurcumin increased GPx expression in rats granulosa cells induced with lead acetate of 40 mg/kg bw. This study found that mean GPx expression in P3 group was higher than that in P2 group and that mean GPx in P2 group was higher than in P1 group. This finding is in line with previous studies which reported that curcumin as an antioxidant can increase catalase, superoxide dismutase (SOD), and glutathione peroxidase (GPx)²¹.

Curcumin has a protective effect on reproductive organs, such as anti-inflammatory, anti-apoptotic and antioxidant in normal cells and acts as a pro apoptosis in malignant cells. Curcumin's effects depend on the dose and type of cells used for the trial²².

The protective mechanism of nanocurcumin against lead acetate-induced decreased GPx expression in rats' ovarian granulosa cells is through the inhibition of superoxide radical formation (O₂⁻) by suppressing the activity of cytochrome P450. P450 is an important isoenzyme for initial bioactivation of reactive oxygen species²³. Inhibition of cytochrome P450 activity will inhibit oxidative phosphorylation, which is an important part of metabolism that produces reactive oxygen species, the superoxide (O₂⁻). The reduction of

superoxide radicals (O₂⁻) will increase GPx expression²⁴.

This study proved that lead acetate exposure of 40 mg/kg bw/day for 26 days in rats increased ovarian granulosa cell apoptosis compared to that of the control group. The results of this study were in line with previous studies, that the administration of intra-peritoneal injection of lead acetate of 20 mg kg bw for 7 days to rats increased pro-apoptosis markers (Bax and caspase-3²⁵). Researchers also reported that administering intraperitoneal injection of lead acetate of 20 mg/kg bw/day for 7 days to rats increased nephron apoptosis, as evidenced by the increase of pro-apoptotic nephron markers, the Bax and caspase-3 proteins, and, conversely, the decrease of anti-apoptotic marker, the Bcl-2 protein, as compared to control group receiving injection of 1 ml/kg bw/day normal saline²⁶.

Lead acetate exposure causes increased H₂O₂ causes oxidative stress which can trigger apoptosis²⁷. Hydroxyl radicals (OH^{*}) formed as a result of lead exposure translocate to ovarian granulosa cell nucleus and stimulate the release of P53. P53 reacts with mitochondrial membrane and activates pro-apoptosis (Bax) and causes decreased anti-apoptosis (Bcl-2 and Bcl-x) leading to the release of cytochrome c into granulosa cell cytosol. In the cytosol, cytochrome c binds to Apaf-1 (apoptosis-activating factor 1), forming the caspase recruitment domain (CARD) which stimulates caspase-9 granulosa cells, and caspase-9 stimulates caspase-3 which is an effector that carries out apoptosis of granulosa cells¹⁰.

This study also proved that administering nanocurcumin reduced lead acetate-induced apoptosis of granulosa cells in rats ovaries. at a dose of 40 mg/kg bw. This was evident in this study that the mean apoptosis expression in P3 rats was lower than in P2 group, the mean apoptosis in P2 group was lower than in P1 group, and that in P3, P2, and P1 groups were lower than in K+ group. This suggests that nanocurcumin provides a protective effect against ovarian granulosa cell apoptosis.

Curcumin isolated from *Curcuma longa* (turmeric) is very potential as an antioxidant which is thought to be caused by phenolic and 1,3-diketone groups¹⁴. This natural polyphenolic antioxidant compound is multifunctional and can function as: (1) antidote to free radicals such as superoxide (O₂^{*}) and hydroxyl radicals (*OH), (2)

chelating metals such as iron (Fe), (3) inhibiting oxidative enzyme activity such as cytochrome P-450, and (4) reducing the formation of oxygen radicals²¹. The antioxidant activity of curcumin compound can occur because the formation of free radicals is inhibited by this compound by suppressing the activity of cytochrome p450²⁸.

As an antioxidant, nanocurcumin plays an important role in preventing apoptosis caused by oxidative stress due to lead toxicity by inhibiting the formation of hydroxyl radicals (OH^{*}). The inhibition of hydroxyl radicals (OH^{*}) formation is by preventing Haber Weiss reaction and Fenton reaction. The prevention is carried out by chelating the transition metals F ++ and C + which act as catalysts for OH^{*} formation²⁹. The prevention of OH^{*} formation suppresses Bax, and Bax increases the release of Bcl-2 and Bcl-xl expression and suppresses cytochrome c expression out of mitochondrial membrane, resulting in the absence of binding between cytochrome c and Apaf-1, which is called apoptosome³⁰. Thus, there is a decrease in stimulation to caspase-9 which results in the decrease of caspase-3 and ends with the decrease in apoptosis³¹.

Conclusions

Nanocurcumin acts as protection against the reduction of GPx scavenger enzyme and the increase in apoptosis of lead acetate-induced rats ovarian granulosa cells.

Acknowledgements

The authors are grateful to the Physics Laboratory Unit, the Faculty of Science, and the Electron Microscope Laboratory Unit, Faculty of Medicine, Airlangga University, Surabaya, Indonesia, who have helped manufacture nanocurcumin and carry out immunohistochemical examinations. We also thank the Robotics Laboratory, ITS, Surabaya, Indonesia, for helping to analyze nanocurcumin.

Declaration of Interest

Tidak ada konflik kepentingan dalam artikel ini.

References

1. Singh N, Kumar A, Gupta VK, Sharma B. Biochemical and Molecular Bases of Lead-Induced Toxicity in Mammalian Systems and Possible Mitigations. *Chem Res Toxicol*. 2018;31(10):1009–10021.
2. Thangarajan S, Vedagiri A, Somasundaram S, Sakthimananogaran R, Murugesan M. Neuroprotective effect of morin on lead acetate- induced apoptosis by preventing cytochrome c translocation via regulation of Bax/Bcl-2 ratio. *Neurotoxicol Teratol*. 2018;66(1):35–45.
3. Dumitrescu E, Chiurciu V, Muselin F, Popescu R, Brezovan D, Cristina RT. Effects of long-term exposure of female rats to low levels of lead: Ovary and uterus histological architecture changes. *Turkish J Biol*. 2015;39(2):284–289.
4. Sudjarwo SA, Eraiko K, Sudjarwo GW, Koerniasari. Protective effects of piperine on lead acetate induced-nephrotoxicity in rats. *Iran J Basic Med Sci*. 2017;20(11):1227–1231.
5. Sudjarwo SA, Anwar C, Wardani G, Eraiko K, Koerniasari. Antioxidant and anti-caspase 3 effect of chitosan-Pinus merkusii extract nanoparticle against lead acetate-induced testicular toxicity in rat. *Asian Pacific J Reprod*. 2019;8(1):13–19.
6. Abdelhamid FM, Mahgoub HA, Ateya AI. Ameliorative effect of curcumin against lead acetate – induced hemato-biochemical alterations, hepatotoxicity, and testicular oxidative damage in rats. *Environ Sci Pollut Res*. 2020;3(1):1–16.
7. Gurer H, Ercal N. Can antioxidants be beneficial in the treatment of lead poisoning? *Free Radic Biol Med*. 2000;29(10):927–945.
8. Zhai Q, Narbad A, Chen W. Dietary strategies for the treatment of cadmium and lead toxicity. *Nutrients*. 2015;7(1):552–571.
9. Assi MA, Noor M, Hezme M, Haron AW, Yusof M, Sabri M. The detrimental effects of lead on human and animal health. *Vet World*. 2016;9(6):660–671.
10. Ighodaro OM, Akinloye OA. First line defence antioxidants-superoxide dismutase (SOD), catalase (CAT) and glutathione peroxidase (GPX): Their fundamental role in the entire antioxidant defence grid. *Alexandria J Med*. 2018;54(4):287–293.
11. Parwata MOA. Teaching Material: Antioxidant [Internet]. Applied Chemistry Postgraduate Program Universitas Udayana. 2016;1(3) 1–54.
12. Kamilatussaniah, Yuniastuti A, Iswari R. Effect of *madu kelengkeng* supplementation on lead-induced white rats TSA and MD level. *J MIPA*. 2015;38(2):108–114.
13. Bodhankar MM, Chikhle S. Various Approaches Towards Enhancement of Bioavailability of Curcumin-a Potent Phytochemical. *World J Pharm Res*. 2019;8(1):606–626.
14. Kabeer A, Muhammad Mailafiya M, Danmaigoro A, Abdul Rahim E, Bu Bakar MZA. Therapeutic potential of curcumin against lead-induced toxicity: A review. *Biomed Res Ther*. 2019;6(3):3053–3066.
15. Abu-Taweel GM. Curcumin attenuates lead (Pb)-induced neurobehavioral and neurobiochemical dysfunction: A Review. *Int J Pharm Pharm Sci*. 2018;10(8):23–28.
16. Le MH, Do HD, Thi HHT, Dung LV, Nguyen HN, Thi HNT, et al. The dual effect of curcumin nanoparticles encapsulated by 1-3/1-6 β-glucan from medicinal mushrooms *Hericium erinaceus* and *Ganoderma lucidum*. *Adv Nat Sci Nanosci Nanotechnol*. 2016;7(4):1–9.
17. Shishu, Maheshwari M. Comparative bioavailability of curcumin, turmeric and Biocurcumax™ in traditional vehicles using non-everted rat intestinal sac model. *J Funct Foods*. 2010;2(1):60–65.
18. Dkhill MA, Al-Khalifa MS, Al-Quraishy S, Zrieq R, Moneim AEA. Indigofera oblongifolia mitigates lead-acetate-induced kidney damage and apoptosis in a rat model. *Drug Des Devel Ther*. 2016;10(7):1847–1856.

19. Al-Megrin WA, Alkhuriji AF, Yousef AOS, Metwally DM, Habotta OA, Kassab RB, et al. Antagonistic efficacy of luteolin against lead acetate exposure-associated with hepatotoxicity is mediated via antioxidant, anti-inflammatory, and anti-apoptotic activities. *Antioxidants*. 2020;9(10):1-18
20. Wang S, He G, Chen M, Zuo T, Xu W, Liu X. The Role of Antioxidant Enzymes in the Ovaries. *Oxid Med Cell Longev*. 2017;2017(9):1-14.
21. Alhusaini A, Fadda L, Hasan IH, Zakaria E, Alenazi AM, Mahmoud AM. Curcumin ameliorates lead-induced hepatotoxicity by suppressing oxidative stress and inflammation, and modulating akt/gsk-3 β signaling pathway. *Biomolecules*. 2019;9(11):1-17
22. Mohebbati R, Anaegoudari A, Khazdair MR. The effects of *Curcuma longa* and curcumin on reproductive systems. *Endocr Regul*. 2017;51(4):220-228.
23. Flora G, Gupta D, Tiwari A. Toxicity of lead: A review with recent updates. *Interdiscip Toxicol*. 2012;5(2):47-58.
24. Grivennikova VG, Vinogradov AD. Generation of superoxide by the mitochondrial Complex I. *Biochim Biophys Acta - Bioenerg*. 2006;1757(4):553-561.
25. AL-Megrin WA, Soliman D, Kassab RB, Metwally DM, Dina DM, El-Khadragy MF. Coenzyme Q10 Activates the Antioxidant Machinery and Inhibits the Inflammatory and Apoptotic Cascades Against Lead Acetate-Induced Renal Injury in Rats. *Front Physiol*. 2020;11(2):1-13.
26. Albarakati AJA, Baly RS, Aljoudi AM, Habotta OA, Elmahallawy EK, Kassab RB, et al. Luteolin protects against lead acetate-induced nephrotoxicity through antioxidant, anti-inflammatory, anti-apoptotic, and Nrf2/HO-1 signaling pathways. *Mol Biol Rep*. 2020; 1(3):1-13
27. Susanti E, Sudiana I., K, Hendarto H. Smoke effect of disturbances folliculogenesis (Mda, Gnrh, Hsp70, Apoptosis, and follicles) in ovarian on mice balb/C. *Journal of International Dental and Medical Research*. 2020;13(2):774-777
28. Majeed M, Badmaev V, Shivakumar U, Rajendranm R. Curcuminoids - Antioxidant Phytonutrients. 1995;1-24.
29. Sudiana IK. Hantaran sinyal pada proses inflamasi. 1st ed. Surabaya: Airlangga University Press; 2017; 1(1): 5-28.
30. Banu MAF, Abhilasha R. Patterns of Apoptosis – A Review. *Int J Pharm Sci Rev Res* 2018;50(2):1-6.
31. Redza-Dutordoir M, Averill-Bates DA. Activation of apoptosis signalling pathways by reactive oxygen species. *Biochim Biophys Acta - Mol Cell Res* .2016;1863(12):2977-2992.

Source details

Journal of International Dental and Medical Research

Open Access ⓘ

Scopus coverage years: from 2009 to Present

Publisher: Ektodermal Displazi Grubu

SSN: 1309-100X

Subject area: Dentistry: General Dentistry

Source type: Journal

CiteScore 2020 **1.3** ⓘ

SJR 2020 **0.259** ⓘ

SNIP 2020 **0.759** ⓘ

[View all documents >](#) [Set document alert](#) [Save to source list](#) [Source Homepage](#)

[CiteScore](#) [CiteScore rank & trend](#) [Scopus content coverage](#)

Improved CiteScore methodology

CiteScore 2020 counts the citations received in 2017-2020 to articles, reviews, conference papers, book chapters and data papers published in 2017-2020, and divides this by the number of publications published in 2017-2020. [Learn more >](#)

CiteScore 2020

1.3 = $\frac{1,250 \text{ Citations } 2017 - 2020}{953 \text{ Documents } 2017 - 2020}$

Calculated on 05 May, 2021

CiteScoreTracker 2021 ⓘ

0.9 = $\frac{774 \text{ Citations to date}}{828 \text{ Documents to date}}$

Last updated on 04 June, 2021 • Updated monthly

CiteScore rank 2020 ⓘ

Category	Rank	Percentile
Dentistry		
— General Dentistry	#75/111	32nd

[View CiteScore methodology >](#) [CiteScore FAQ >](#) [Add CiteScore to your site ↗](#)

About Scopus

- What is Scopus
- Content coverage
- Scopus blog
- Scopus API
- Privacy matters

Language

- 日本語に切り替える
- 切换到简体中文
- 切换到繁體中文
- Русский язык

Customer Service

- Help
- Contact us



Get peer reviews certified

ReviewerCredits - Certification and rewarding of peer review and conference talks

reviewercredits.com

Journal of International Dental and Medical Research

COUNTRY	SUBJECT AREA AND CATEGORY	PUBLISHER	H-INDEX
Turkey Universities and research institutions in Turkey	Dentistry Dentistry (miscellaneous)	Ektodermal Displazi Grubu	13

PUBLICATION TYPE	ISSN	COVERAGE
Journals	1309100X	2009-2020

Scopus Indexed Jo

Call for Papers August Iss
Fast Track Peer Reviewed Put
tojqi.net

OPEN

SCOPE

Information not localized

Join the conversation about this journal

Scopus Indexed Journal

Fast Track Peer Reviewed Publication.

tojqi.net

OPEN

Quartiles

Reviewer Certifications

Why Join Peer Reviewers - Free Registration - ReviewerCredits

reviewercredits.com

OPEN

FIND SIMILAR JOURNALS ?

1
Journal of International Oral Health
IND

40%
similarity

2
Pesquisa Brasileira em Odontopediatria e Clinica
BRA

38%
similarity

3
Journal of International Society of Preventive and
IND

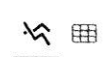
37%
similarity

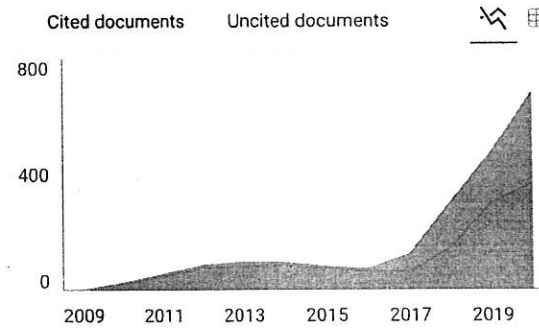
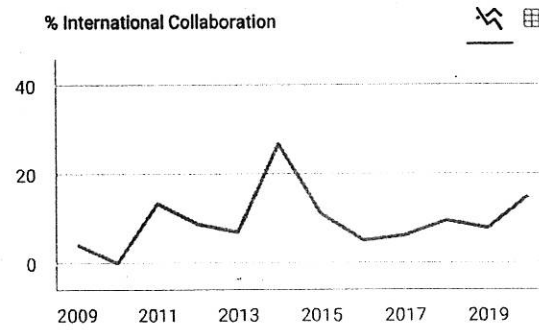
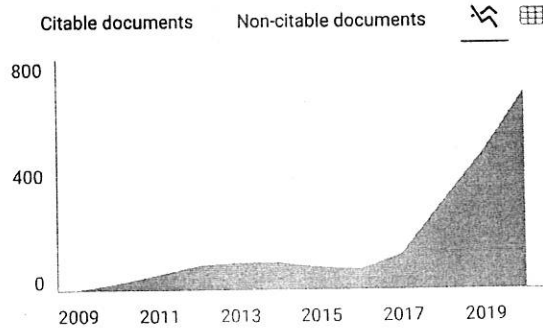
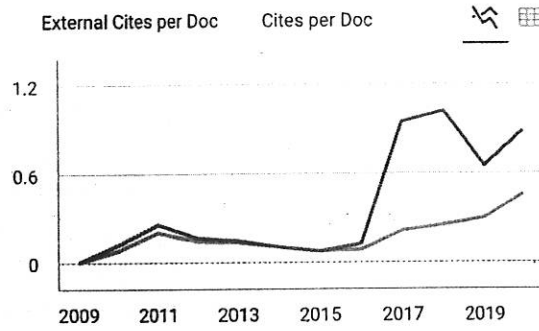
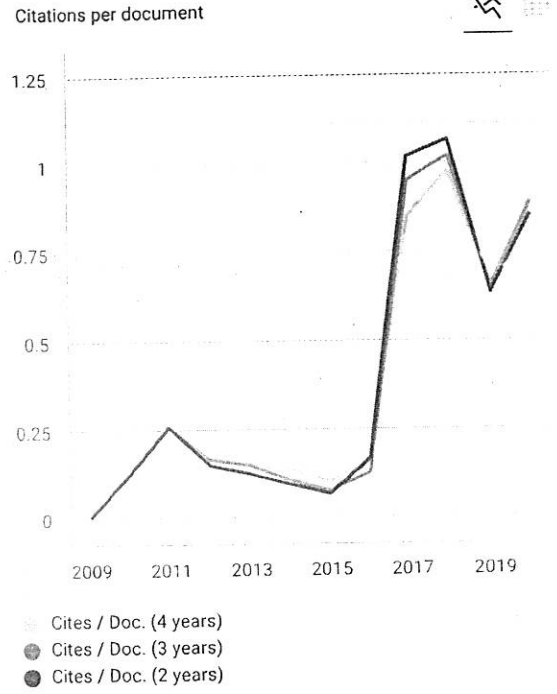
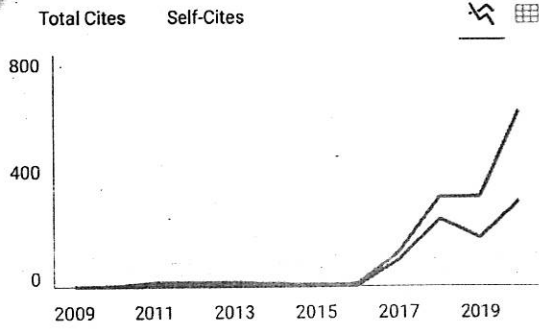
4
Indian Jourr Research
IND

SJR



Total Documents





Journal of International Dental and Medical...

Q3 Dentistry (miscellaneous)

best quartile

SJR 2020

0.26

powered by scimagojr.com

← Show this widget in your own website

Just copy the code below and paste within your html code:

```
<a href="https://www.scimagojr.com"
```