Online ISSN: 2249-4618 Print ISSN: 0975-5888 DOI: 10.17406/GJMRA

# GLOBAL JOURNAL

OF MEDICAL RESEARCH: K

# Interdisciplinary



**VOLUME 20** 

ISSUE 13

**VERSION 1.0** 



# Global Journal of Medical Research: K Interdisciplinary

# Global Journal of Medical Research: K Interdisciplinary

VOLUME 20 ISSUE 13 (VER. 1.0)

OPEN ASSOCIATION OF RESEARCH SOCIETY

# © Global Journal of Medical Research. 2020.

All rights reserved.

This is a special issue published in version 1.0 of "Global Journal of Medical Research." By Global Journals Inc.

All articles are open access articles distributed under "Global Journal of Medical Research"

Reading License, which permits restricted use.

Entire contents are copyright by of "Global
Journal of Medical Research" unless
otherwise noted on specific articles.

No part of this publication may be reproduced or transmitted in any form or by any means, electronic or mechanical, including photocopy, recording, or any information storage and retrieval system, without written permission.

The opinions and statements made in this book are those of the authors concerned.

Ultraculture has not verified and neither confirms nor denies any of the foregoing and no warranty or fitness is implied.

Engage with the contents herein at your own risk

The use of this journal, and the terms and conditions for our providing information, is governed by our Disclaimer, Terms and Conditions and Privacy Policy given on our website <a href="http://globaljournals.us/terms-and-condition/">http://globaljournals.us/terms-and-condition/</a>

menu-ju-1465/

By referring / using / reading / any type of association / referencing this journal, this signifies and you acknowledge that you have read them and that you accept and will be bound by the terms thereof.

All information, journals, this journal, activities undertaken, materials, services and our website, terms and conditions, privacy policy, and this journal is subject to change anytime without any prior notice.

Incorporation No.: 0423089 License No.: 42125/022010/1186 Registration No.: 430374 Import-Export Code: 1109007027 Employer Identification Number (EIN): USA Tax ID: 98-0673427

# Global Journals Inc.

(A Delaware USA Incorporation with "Good Standing"; Reg. Number: 0423089)

Sponsors: Open Association of Research Society

Open Scientific Standards

# Publisher's Headquarters office

Global Journals® Headquarters 945th Concord Streets, Framingham Massachusetts Pin: 01701, United States of America

USA Toll Free: +001-888-839-7392 USA Toll Free Fax: +001-888-839-7392

# Offset Typesetting

Global Journals Incorporated 2nd, Lansdowne, Lansdowne Rd., Croydon-Surrey, Pin: CR9 2ER, United Kingdom

# Packaging & Continental Dispatching

Global Journals Pvt Ltd E-3130 Sudama Nagar, Near Gopur Square, Indore, M.P., Pin:452009, India

# Find a correspondence nodal officer near you

To find nodal officer of your country, please email us at *local@globaljournals.org* 

# *eContacts*

Press Inquiries: press@globaljournals.org
Investor Inquiries: investors@globaljournals.org
Technical Support: technology@globaljournals.org
Media & Releases: media@globaljournals.org

# Pricing (Excluding Air Parcel Charges):

Yearly Subscription (Personal & Institutional) 250 USD (B/W) & 350 USD (Color)

# EDITORIAL BOARD

# GLOBAL JOURNAL OF MEDICAL RESEARCH

# Dr. Apostolos Ch. Zarros

DM, Degree (Ptychio) holder in Medicine, National and Kapodistrian University of Athens MRes, Master of Research in Molecular Functions in Disease, University of Glasgow FRNS, Fellow, Royal Numismatic Society Member, European Society for Neurochemistry Member, Royal Institute of Philosophy Scotland, United Kingdom

# Dr. Alfio Ferlito

Professor Department of Surgical Sciences University of Udine School of Medicine, Italy

# Dr. Jixin Zhong

Department of Medicine, Affiliated Hospital of Guangdong Medical College, Zhanjiang, China, Davis Heart and Lung Research Institute, The Ohio State University, Columbus, OH 43210, US

# Rama Rao Ganga

**MBBS** 

MS (Universty of Health Sciences, Vijayawada, India) MRCS (Royal Coillege of Surgeons of Edinburgh, UK) United States

# Dr. Izzet Yavuz

MSc, Ph.D., D Ped Dent.

Associate Professor, Pediatric Dentistry Faculty of Dentistry, University of Dicle Diyarbakir, Turkey

# Sanguansak Rerksuppaphol

Department of Pediatrics Faculty of Medicine Srinakharinwirot University NakornNayok, Thailand

# Dr. William Chi-shing Cho

Ph.D.,

Department of Clinical Oncology Queen Elizabeth Hospital Hong Kong

# Dr. Michael Wink

Ph.D., Technical University Braunschweig, Germany Head of Department Institute of Pharmacy and Molecular Biotechnology, Heidelberg University, Germany

# Dr. Pejcic Ana

Assistant Medical Faculty Department of Periodontology and Oral Medicine University of Nis, Serbia

# Dr. Ivandro Soares Monteiro

M.Sc., Ph.D. in Psychology Clinic, Professor University of Minho, Portugal

# Dr. Sanjay Dixit, M.D.

Director, EP Laboratories, Philadelphia VA Medical Center Cardiovascular Medicine - Cardiac Arrhythmia Univ of Penn School of Medicine Web: pennmedicine.org/wagform/MainPage.aspx?

# Antonio Simone Laganà

M.D. Unit of Gynecology and Obstetrics Department of Human Pathology in Adulthood and Childhood "G. Barresi" University of Messina, Italy

# Dr. Han-Xiang Deng

MD., Ph.D

Associate Professor and Research Department

Division of Neuromuscular Medicine

Davee Department of Neurology and Clinical

Neurosciences

Northwestern University Feinberg School of Medicine

Web: neurology.northwestern.edu/faculty/deng.html

# Dr. Roberto Sanchez

Associate Professor

Department of Structural and Chemical Biology

Mount Sinai School of Medicine

Ph.D., The Rockefeller University

Web: mountsinai.org/

# Dr. Feng Feng

Boston University

Microbiology

72 East Concord Street R702

Duke University

United States of America

# Dr. Hrushikesh Aphale

MDS- Orthodontics and Dentofacial Orthopedics.

Fellow- World Federation of Orthodontist, USA.

# Gaurav Singhal

Master of Tropical Veterinary Sciences, currently pursuing Ph.D in Medicine

# Dr. Pina C. Sanelli

Associate Professor of Radiology

Associate Professor of Public Health

Weill Cornell Medical College

Associate Attending Radiologist

NewYork-Presbyterian Hospital

MRI, MRA, CT, and CTA

Neuroradiology and Diagnostic Radiology

M.D., State University of New York at Buffalo,

School of Medicine and Biomedical Sciences

Web: weillcornell.org/pinasanelli/

# Dr. Michael R. Rudnick

M.D., FACP

Associate Professor of Medicine

Chief, Renal Electrolyte and Hypertension Division (PMC)

Penn Medicine, University of Pennsylvania

Presbyterian Medical Center, Philadelphia

Nephrology and Internal Medicine

Certified by the American Board of Internal Medicine

Web: uphs.upenn.edu/

# Dr. Seung-Yup Ku

M.D., Ph.D., Seoul National University Medical College, Seoul, Korea Department of Obstetrics and Gynecology

Seoul National University Hospital, Seoul, Korea

# Santhosh Kumar

Reader, Department of Periodontology,

Manipal University, Manipal

# Dr. Aarti Garg

Bachelor of Dental Surgery (B.D.S.) M.D.S. in Pedodontics and Preventive Dentistr Pursuing Phd in Dentistry

# Sabreena Safuan

Ph.D (Pathology) MSc (Molecular Pathology and Toxicology) BSc (Biomedicine)

# Getahun Asebe

Veterinary medicine, Infectious diseases, Veterinary Public health, Animal Science

# Dr. Suraj Agarwal

Bachelor of dental Surgery Master of dental Surgery in Oromaxillofacial Radiology.

Diploma in Forensic Science & Oodntology

# Osama Alali

PhD in Orthodontics, Department of Orthodontics, School of Dentistry, University of Damascus. Damascus, Syria. 2013 Masters Degree in Orthodontics.

# Prabudh Goel

MCh (Pediatric Surgery, Gold Medalist), FISPU, FICS-IS

# Raouf Hajji

MD, Specialty Assistant Professor in Internal Medicine

# Surekha Damineni

Ph.D with Post Doctoral in Cancer Genetics

# Arundhati Biswas

MBBS, MS (General Surgery), FCPS, MCh, DNB (Neurosurgery)

# Rui Pedro Pereira de Almeida

Ph.D Student in Health Sciences program, MSc in Quality Management in Healthcare Facilities

# Dr. Sunanda Sharma

B.V.Sc.& AH, M.V.Sc (Animal Reproduction,
Obstetrics & gynaecology),
Ph.D.(Animal Reproduction, Obstetrics & gynaecology)

# Shahanawaz SD

Master of Physiotherapy in Neurology PhD- Pursuing in Neuro Physiotherapy Master of Physiotherapy in Hospital Management

# Dr. Shabana Naz Shah

PhD. in Pharmaceutical Chemistry

# Vaishnavi V.K Vedam

Master of dental surgery oral pathology

# Tariq Aziz

PhD Biotechnology in Progress

# CONTENTS OF THE ISSUE

- i. Copyright Notice
- ii. Editorial Board Members
- iii. Chief Author and Dean
- iv. Contents of the Issue
- Comparison of ATP Values on Meat and Fish Cutting Boards before and after Alcohol Disinfection. 1-6
- 2. Study of Biochemical and Anthropometric Variables among Pancagavya and Non-Pancagavya Diet Population: A Cross-Sectional Comparative Study. *7-11*
- 3. Comparison of ATP Values on Vegetables Cutting Boards before and after Alcohol Disinfection. *13-18*
- 4. A New Medical Device for Platelet Rich Plasma Filler. 19-29
- 5. Deep Learning for Classification of Sleep EEG Data during the Epidemic of Coronavirus Disease. 31-34
- 6. Hidradenitis Suppurativa- The Imaging Spectrum. 35-41
- v. Fellows
- vi. Auxiliary Memberships
- vii. Preferred Author Guidelines
- viii. Index



# Global Journal of Medical Research: k Interdisciplinary

Volume 20 Issue 13 Version 1.0 Year 2020

Type: Double Blind Peer Reviewed International Research Journal

Publisher: Global Journals

Online ISSN: 2249-4618 & Print ISSN: 0975-5888

# Comparison of ATP Values on Meat and Fish Cutting Boards before and after Alcohol Disinfection

By Akemi Ito, Naomi Katayama, Mayumi Hirabayashi, Natuki Sasaki & Moe Inuzuka

Nagoya Women's University

Abstract- Sanitary control of cutting boards in the kitchen is important to prevent food poisoning. Using ATP and microbiological tests, we investigated the cleaning and 70% alcohol spraying effects of cutting boards for meat and fish. As a result, the ATP value and the number of microbial bacteria decreased after washing the cutting board but decreased more after spraying with 70% alcohol. The ATP value was 100 or less after spraying with 70% alcohol. The number of microbial bacteria decreased after spraying with 70% alcohol. However, not all bacterialeliminated even after spraying with 70% alcohol. If the cutting boardleft in a moist state at room temperature, microorganisms could grow again.

Keywords: Gender: ATP wiping test, Microbial stamp test, Cutting board, alcohol disinfection.

GJMR-K Classification: NLMC Code: QV 250



Strictly as per the compliance and regulations of:



© 2020. Akemi Ito, Naomi Katayama, Mayumi Hirabayashi, Natuki Sasaki & Moe Inuzuka. This is a research/review paper, distributed under the terms of the Creative Commons Attribution-Noncommercial 3.0 Unported License http://creativecommons.org/licenses/by-nc/3.0/), permitting all non-commercial use, distribution, and reproduction in any medium, provided the original work is properly cited.

# Global Journal of Medical Research (K) Volume XX Issue XIII Version I - Year 2020

# Comparison of ATP Values on Meat and Fish Cutting Boards before and after Alcohol Disinfection

Akemi Ito a, Naomi Katayama a, Mayumi Hirabayashi b, Natuki Sasaki a & Moe Inuzuka a

Abstract- Sanitary control of cutting boards in the kitchen is important to prevent food poisoning. Using ATP and microbiological tests, we investigated the cleaning and 70% alcohol spraying effects of cutting boards for meat and fish. As a result, the ATP value and the number of microbial bacteria decreased after washing the cutting board but decreased more after spraying with 70% alcohol. The ATP value was 100 or less after spraving with 70% alcohol. The number of microbial bacteria decreased after spraying with 70% alcohol. However, not all bacterialeliminated even after spraying with 70% alcohol. If the cutting boardleft in a moist state at room temperature, microorganisms could grow again.

Keywords: Gender: ATP wiping test, Microbial stamp test, Cutting board, alcohol disinfection.

#### Introduction I.

anitary control of cutting boards in the kitchen is important to prevent food poisoning. In the past, we reported the results of hygiene management by repeatedly cleaning the cutting board with detergent and running water for 30 seconds or more<sup>1)</sup>. Currently, the COVID-19 epidemic requires stricter hygiene control. To control invisible microorganisms, it is necessary to take measures to avoid the risk of food poisoning accidents due to familiarity with cooking work; as the O-JT education, it is necessary to create a hygiene management manual and protect it with all the cooks<sup>2,3,4,5)</sup>. However, if the procedure is complicated and difficult, it will not last long. We need easy and reliable procedures and methods that anyone can do. The ATP tests<sup>6,7,8)</sup> and HACCP-based microbiological tests<sup>9)</sup> are useful in hygiene management to transform invisible bacteria into visible forms and educate them. Therefore, in this study, and the cutting board cleaning method we performed last time, a step of spraying 70% alcohol added. The effects of this alcohol disinfection compared by adding a stamping test (General bacteria. Staphylococcus aureus, Escherichia coli, Salmonella, Vibrio parahaemolyticus) in addition to the same ATP test as in the previous report.

Author α σ ρ: Graduate School of Nagoya Women's University, Nagoya City, Japan.

Author σ ω ¥: Nagoya Women's University, Nagoya City, Japan.

Author o: Department of Otorhinolaryngology, Nagoya University Graduate School of Medicine, Nagoya, Japan.

Corresponding Author o: Nagoya Women's University, Nagoya City, Japan. e-mail: naomik@nagoya-wu.ac.jp

#### Materials and Methods II.

# a) Kitchen cutting board

The six kitchen meat or fish thick cutting board (cutting board 1) and the six kitchen meat for the fish thin cutting board (cutting board 2) prepared in the kitchen were stored in the sterilization storage the day before cooking.

# b) ATP inspection procedure

Each of the 12 cooks carried a kitchen cutting board for meat or fish at the start of their work and brought it to the cooking table. The work start time depends on the working conditions of the cooks. Still, the inspector always performed an ATP inspection before using meat or fish with a kitchen cutting board. Then, each cook finished the work, washes the cutting board firmly with detergent and sponge, rinse with running water for 30 seconds or more. Then, each cook repeated this process twice (as same as the last report<sup>1)</sup>). The inspector performed an ATP inspection after using meat or fish with a kitchen cutting board, again. Then, each cook sprayed 70% alcohol on the cutting board after washing. At last, the inspector performed an ATP inspection after using meat or fish with a kitchen cutting board. The ATP test kit used manufactured by KIKKOMAN.

## c) Stamp test inspection procedure

Five types of stamp test (General bacteria, Staphylococcus aureus, Escherichia coli, Salmonella, Vibrio parahaemolyticus) used. The stamp test conducted by the inspector at the same time as the ATP. The stamp test was colony-counted after culturing in an incubator at 38 degrees for three days. The stamp test made by NISSUI.

# d) Statistical processing

The results obtained compared using statistical methods. The data were statistically processed, was subjected to an F test to determine whether to use a parametric test or nonparametric test. When there is no difference in the F test, the presence or absence of a significant difference was confirmed using the student ttest with or without a correspondence. If there was a difference in the F test, the presence or absence of a significant difference was confirmed using the Wilcoxon

test with a pair or the Mann-Whitney test without correlation.

#### RESULTS III.

ATPvalue results before and after alcohol disinfection

The table 1 and 2 shows the results of ATP wiping tests on cutting board before and after alcohol disinfection. It can see that the average value of the ATP values measured after washing before and after cleaning, after cleaning, the ATP value is low. The ATP value after 70%alcohol spraying was statistically significantly lower than that before alcohol spraying. The ATP value dropped below 100 for both cutting boards.

Table 1. ATP test value and statistical processing result of cutting board 1.

	No alcohol	treatment	Alcohol t	reatment	
For meat	Before washing	After washing	After washing	After alcohol	
1	8414	50121	50121	13	
2	210	56	56	31	
3	132205	103	103	20	
4	59141	62	62	31	
5	30814	272	272	18	
6	76010	70	70	72	
Average value	51132.3	8447.3	8447.3	30.8	
SD	49166.2	20416.0	20416.0	21.4	
F test	P-0.0	025*	P=0.0001**		
Student-t*					
Wilcoxon	P=0	.116	P=0.046*		
F test	P=0.0001**				
Student-t*					
Wilcoxon		P=0	0.028*		

\*Paired Student-t test \* P<0.05, \*\* P<0.01

Table 2. ATP test value and statistical processing result of cutting board 2.

·	No alcohol	treatment	Alcohol t	reatment
For meat	Before washing	After washing	After washing	After alcohol
1	4817	3828	3828	38
2	1302	12	12	58
3	99080	456	456	16
4	61864	33	33	56
5	161792	293	293	17
6	243	85	85	50
Average value	54849.7	784.5	784.5	39.2
SD	66022.3	1500.9	1500.9	18.9
F test	P=0.00	01**	P=0.0	0001**
Student-t*				
Wilcoxon	P=0.0	28*	P=0	.173
F test		P=0.0	0001**	
Student-t*				
Wilcoxon	P=0.028*			

\*Paired Student-t test \* P<0.05, \*\* P<0.01

Stamp test results before and after alcohol disinfection

Tables 3,4,5,6,7,8.9.10.11 and 12 show the results of ATP wiping tests on cutting board before and after 70% alcohol disinfection. Results of general bacteria show in Tables 3 and 4. Results of E. coli show in Tables 5 and 6. Results of Staphylococcus aureus shown in Tables 7 and 8. Result of Salmonella show in Tables 9 and 10. Result of Vibrio parahaemolyticus show in Tables 11 and 12. The number of all microbial bacteria was lower after washing than after cooking and after spraying 70% alcohol. However, there was no statistically significant difference in the number of microbial bacteria.

Table 3 Number of general bacteria on cutting board 1. and statistical processing result

	No alcohol	treatment	Alcohol	treatment
For meat	Before washing	After washing	After washing	After alcohol
1	82	40	40	0
2	4	0	0	0
3	200	46	46	0
4	13	0	0	0
5	200	9	9	0
6	60	0	0	14
Average value	93.2	15.8	15.8	2.3
SD	87.7	21.4	21.4	5.7
F test	P=0.00	02**	P=0.	.003**
Student-t*				
Wilcoxon	P=0.0	28*	P=0	0.273
Ftest	•	P=0.	0001**	-
Student-t*				
Wilcoxon		P=0	0.028*	
	*Doing J Carr	James & Band * Dan	05 ** D -0 01	

\*Paired Student-t test \* P<0.05, \*\* P<0.01

Table 4 Number of general bacteria on cutting board 2. and statistical processing result

	No alcohol treatment				
F					
For meat	Before washing	After wasning	After washing	After alcohol	
1	270	7	7	0	
2	1	18	18	8	
3	200	8	8	0	
4	2	7	7	0	
5	200	23	23	0	
6	61	20	20	0	
Average value	122.3	13.8	13.8	1.3	
SD	115.6	7.3	7.3	3.3	
F test	P=0.00	001**	P=0.	.035*	
Student-t*					
Wilcoxon	P=0.	.116	P=0.	.028*	
F test	P=0.0001**				
Student-t*					
Wilcoxon		P=	0.075		

<sup>\*</sup>Paired Student-t test \* P<0.05, \*\* P<0.01

Table 5 Number of E. coli on cutting board 1. and statistical processing result

	No alcohol	treatment	Alcohol t	treatment
For meat	Before washing	After washing	After washing	After alcohol
1	0	0	0	0
2	2	1	1	0
3	200	8	8	0
4	5	0	0	7
5	200	0	0	0
6	6	0	0	16
Average value	68.8	1.5	1.5	3.8
SD	101.6	3.2	3.2	6.6
F test	P=0.00	001**	P=0	.052
Student-t*			P=0	.518
Wilcoxon	P=0.0	)43*		
F test	·	P=0.0	001**	·
Student-t*				
Wilcoxon		P=0.	.418	
	*Paired Stud	ent-t test * P<0.05	5, ** P<0.01	

Table 6 Number of E. coli on cutting board 2. and statistical processing result

	No alcohol	treatment	Alcohol t	treatment
For meat	Before washing	After washing	After washing	After alcohol
1	3	2	2	0
2	0	0	0	1
3	200	23	23	0
4	0	0	0	0
5	15	0	0	0
6	35	0	0	0
Average value	42.2	4.2	4.2	0.2
SD	78.5	9.3	9.3	0.4
F test	P=0.00	001**	P=0.0	001**
Student-t*				
Wilcoxon	P=0.	068	P=0.	.285
F test		P=0.00	001**	
Student-t*				
Wilcoxon		P=0.	080	
	*Paired Stude	ent-t test * P<0.05	, ** P<0.01	•

Table 7 Number of Staphylococcus aureus on cutting board 1. and statistical processing

		result			
	No alcohol	l treatment	Alcohol	treatment	
For meat	Before washing	After washing	After washing	After alcohol	
1	0	0	0	0	
2	5	1	1	0	
3	0	0	0	7	
4	0	0	0	3	
5	152	2	2	0	
6	212	0	0	0	
Average value	61.5	0.5	0.5	1.7	
SD	95.3	0.8	0.8	2.9	
F test	P=0.00	001**	P=0.	004**	
Student-t*					
Wilcoxon	P=0.	109	P=0	0.465	
F test	P=0.0001**				
Student-t*					
Wilcoxon		P=0	).345		
	*Doined Ctu	dont t toot * D o	05 ** D <0.01		

Table 8 Number of Staphylococcus aureus on cutting board 2. and statistical processing

	No alcohol	treatment	Alcohol	treatment
For meat	Before washing	After washing	After washing	After alcohol
1	0	0	0	0
2	0	1	1	16
3	24	0	0	0
4	0	1	1	0
5	200	0	0	0
6	432	0	0	0
Average value	109.3	0.3	0.3	2.7
SD	176.3	0.5	0.5	6.5
F test	P=0.00	001**	P=0.0001**	
Student-t*				
Wilcoxon	P=0.	225	P=0	.655
F test		P=0.0	0001**	
Student-t*				
Wilcoxon		P=0	0.144	

<sup>\*</sup>Paired Student-t test \* P<0.05, \*\* P<0.01

Table 9 Number of Salmonella on cutting board 1. and statistical processing result

	No alcohol	treatment	Alcohol t	reatment	
For meat	Before washing	After washing	After washing	After alcohol	
1	508	0	0	0	
2	80	0	0	0	
3	1	0	0	2	
4	168	0	0	5	
5	1	9	9	0	
6	26	15	15	5	
Average value	130.7	4.0	4.0	2.0	
SD	195.4	6.5	6.5	2.4	
F test	P=0.00	001**	P=0.0	016*	
Student-t*					
Wilcoxon	P=0.	075	P=0.	465	
F test	P=.0001**				
Student-t*					
Wilcoxon		P=0.059			
*Paired_Student-t test_* P<0.05, ** P<0.01					

Table 10 Number of Salmonella on cutting board 2. and statistical processing result

	No alcoho	l treatment	Alcohol	treatment
For meat	Before washing	After washing	After washing	After alcohol
1	0	1	1	0
2	0	5	5	0
3	34	1	1	2
4	0	23	23	0
5	21	1	1	0
6	55	0	0	0
Average value	18.3	5.2	5.2	0.3
SD	22.8	8.9	8.9	0.8
F test	P=0	.019	P=0.0	001**
Student-t*				
Wilcoxon	P=0	.463	P=0	.138
F test	-	P=0.0	0001**	-
Student-t*				
Wilcoxon		P=0	0.109	
	*Paired St	udent-t test * P<0.0	05, ** P<0.01	-

Table 11 Number of Vibrio parahaemolyticus on cutting board 1. and statistical processing result

_	No alcohol	treatment	Alcohol	treatment	
For meat	Before washing	After washing	After washing	After alcohol	
1	21	0	0	0	
2	6	0	0	0	
3	1	7	7	0	
4	119	0	0	0	
5	0	38	38	2	
6	18	0	0	0	
Average value	27.5	7.5	7.5	0.3	
SD	45.7	15.2	15.2	0.8	
F test	P=0.0	09**	P=0.0001**		
Student-t*					
Vilcoxon	P=0.	402	P=0.180		
F test		P=0.0	001**		
Student-t*					
Wilcoxon		P=0.	.075		

<sup>\*</sup>Paired Student-t test \* P<0.05, \*\* P<0.01

Table 12 Number of Vibrio parahaemolyticus on cutting board 2. and statistical processing result

	No alcohol	treatment	Alcohol ti	reatment
For meat	Before washing	After washing	After washing	After alcohol
1	0	0	0	0
2	0	4	4	2
3	5	0	0	0
4	0	0	0	0
5	200	0	0	0
6	256	0	0	0
Average value	76.8	0.7	0.7	0.3
SD	118.4	1.6	1.6	0.8
F test	P=0.00	01**	P=0.	.68
Student-t*			P=0.3	363
Wilcoxon	P=0.	144		
F test		P=0.00	001**	
Student-t*				
Wilcoxon		P=0.	144	
	*D : 1 G: 1	* B 0.0	5 ** D 0.01	

\*Paired Student-t test \* P<0.05, \*\* P<0.01

#### IV. DISCUSSION

To manage the hygiene of meat and fish cutting board that has a high risk of causing secondary contamination in cooking. We tried to verify using the ATP test and microbial stamp test by spraying 70% alcohol after cleaning instead of controlling only by the cleaning method<sup>1)</sup>. The ATP value decreased after washing then after cooking and after spraying 70% alcohol than after washing. The ATP value was a statistically significant decrease, which was less than 100 after 70% alcohol spraying. However, the microbial stamp test results were not statistically significant reductions in bacterial counts. The cutting board inspected by spraying 70% alcohol after cleaning. But if 70% of alcohol not sprayed after sufficiently wiping off the water, the alcohol may be dilute, and the bactericidal effect may weakened. In the future, we would like to verify the sterilization of microorganisms by spraying 70% alcohol on the cutting board by thoroughly wiping off the water after cleaning and then spraying 70% alcohol. Not all microorganisms are killed even after spraying 70% alcohol, so when using a cutting board left at room temperature (with moist), it is better to wash repeatedly and cook after spraying 70% alcohol.

#### V. Conclusions

The effects of 70% alcohol spraying investigated using cutting boards for meat and fish. Both cutting boards had high ATP and microbiological test values after cooking. However, although the value of the cutting board decreased after cleaning, the ATP value did not fall below 100. Microbial test values were also high in many cases. After spraying with 70% alcohol, the ATP value was 100 or less, and the value decreased statistically significantly. Microbial test values were decreasing with or without statistically significant reductions. Providing safe and secure meals by further spraying 70% alcohol after cleaning the cooking utensils helps prevent food poisoning. However, since the microorganisms are present even after spraying with 70% alcohol, the bacteria may grow again if the cooking utensils left for a long time. It is advisable to clean and spray 70% alcohol before using the equipment.

# ACKNOWLEDGMENTS

We would like to thank all the cooks who participated in this experiment. Also, we would like to thank the inspectors who also performed the ATP inspection.

# References Références Referencias

- Katayama N, Hirabayashi M, Ito A, Kondo S, Nakayama Y, Naka A, Sasaki N, Inuzuka M, Tamura T. Results of Hygiene Education of Kitchen Cutting Board by using ATP Inspection - Comparison of vegetable Cutting Board and Meat Cutting Board. (2020). Global Journal of Medical Research. 20(5): 13-16.
- Lee JH (2018) An investigation of Factors that influence Hygiene Practices at a small Day Care Center. (2018). J Food Prot. 81(1): 158-164.
- Stanley PE. A review of bioluminescent ATP techniques in papid microbiology. (1989) J Biolumin Chemilumin 4(1): 375-380.
- Stannard CJ, Gibbs PA. Rapid microbiology: application s of bioluminescence in the food industry—a review. (1986) J Biolumin Chemilumin 1(1): 3-10.5.
- Griffith CJ, Coooper RA, Gilmore J, Davies C, Lweis M. An evaluation of hospital cleaning refimes and standards. (2000) J Hosp Infect. 45(1): 19-28.
- Nante N, Ceriale E, Messina G, Lenzi D, Manzi P. Effectiveness of ATP bioluminescence to assess hospital cleaning: a review. (2017) J Prev. Med. Hyg. 58(2): E177-E183.
- 7. Amodio E, Dubi C. Use of ATP bioluminescence for assessing h eclealiness of hospital surfaces: a review of the published literature (1990-2012).(2014) J infect Public Health 7(2): 92-98.
- Aycieck H, Oquz U. Karci K. Comparison of results of ATP bioluminescence and traditional ygiene swabbing methods fro the determination of surface

- cleanliness at a hospital kitchen. (2006). Int J Hyg Environ Heatth. 209(2): 203-206.
- 9. Osimani A, Garofalo C, Clementi F, Tavoletti S, Aquilanti L.Bioluminescence ATP monitoring for the routine assessment fo food contact surface cleanliness in a university canteen. (2014). Int J Environ Res Public Health 17; 11(10): 10824-10837.



# Global Journal of Medical Research: K Interdisciplinary

Volume 20 Issue 13 Version 1.0 Year 2020

Type: Double Blind Peer Reviewed International Research Journal

Publisher: Global Journals

Online ISSN: 2249-4618 & Print ISSN: 0975-5888

# Study of Biochemical and Anthropometric Variables among Pancagavya and Non-Pancagavya Diet Population: A Cross-Sectional Comparative Study

By Neeraj, Itagi Ravi Kumar, Dwivedi Krishna & Pandey Mangesh

Deemed-to-be University

Abstract- Background: Diseases negatively affect people's work performance, joy in life, emotional and physical health, quality of life, and spiritual well-being. The pancagavya (five cow derivatives) diet based on the consumption of five cow derivatives like milk, curd, ghee, urine and dung from Bos indicus cow not only helps to provide physical health but also useful in other aspects of life.

Objective: To study the biochemical and anthropometric variables among the pancagavya diet population.

*Materials and method:* Both male and female sample size 80 with an age range between 20 to 80 years were recruited from different states of India. The current study is between the pancagavya diet and non-pancagavya diet groups and had more than two years in their diet.

Keywords: pancagavya diet, non-pancagavya diet, bhramari time, health.

GJMR-K Classification: NLMC Code: QT 235



Strictly as per the compliance and regulations of:



© 2020. Neeraj, Itagi Ravi Kumar, Dwivedi Krishna & Pandey Mangesh. This is a research/review paper, distributed under the terms of the Creative Commons Attribution-Noncommercial 3.0 Unported License http://creativecommons.org/licenses/by-nc/3.0/), permitting all non-commercial use, distribution, and reproduction in any medium, provided the original work is properly cited.

# Study of Biochemical and Anthropometric Variables among Pancagavya and Non-Pancagavya Diet Population: A Cross-Sectional Comparative Study

# Impact of Panchagavya Diet on Health

Neeraj a, Itagi Ravi Kumar , Dwivedi Krishna & Pandey Mangesh a

Abstract- Background: Diseases negatively affect people's work performance, joy in life, emotional and physical health, quality of life, and spiritual well-being. The pancagavya (five cow derivatives) diet based on the consumption of five cow derivatives like milk, curd, ghee, urine and dung from Bos indicus cow not only helps to provide physical health but also useful in other aspects of life.

Objective: To study the biochemical and anthropometric variables among the pancagavya diet population.

Materials and method: Both male and female sample size 80 with an age range between 20 to 80 years were recruited from different states of India. The current study is between the pancagavya diet and non-pancagavya diet groups and had more than two years in their diet.

Result: Pancagavya diet group had significantly less random blood glucose levels and had no change in hemoglobin level compared to the non-pancagavya diet group. Pancagavya diet group have more bhramari (humming honey bee sound produced at the back of the throat during the practice time) and respiratory rate with exponential significance, less pulse rate with highly significant, and was having lesser blood pressure compared to the non-pancagavya diet group.

Conclusion: In the biochemical and anthropometric variables studied pancagavya diet group had a better healthy lifestyle compared to the non-pacagavya diet group.

Keywords: pancagavya diet, non-pancagavya diet, bhramari time, health.

Author α: M.Sc. (Yoga Therapy), Division of Yoga and Life Sciences, Swami Vivekananda Yoga Anusandhana Samsthana (S-VYASA), Deemed-to-be University, #19, Eknath Bhavan, Gavipuram Circle, K.G. Nagar, Bangalore, India.

Corresponding Author o: Associate Professor, Division of Yoga and Physical Sciences, Swami Vivekananda Yoga Anusandhana Samsthana (S-VYASA), Deemed-to-be University, #19, Eknath Bhavan, Gavipuram Circle, K.G. Nagar, Bangalore, India. e-mail: itagi.ravi@gmail.com

Author p: Junior Researcher, Division of Yoga and Life Sciences, Swami Vivekananda Yoga Anusandhana Samsthana (S-VYASA), Deemed-to-be University, #19, Eknath Bhavan, Gavipuram Circle, K.G. Nagar, Bangalore, India.

Author ω: Assistant professor, Division of Yoga and Humanities, Swami Vivekananda Yoga Anusandhana Samsthana (S-VYASA), Deemed-to-be University, #19, Eknath Bhavan, Gavipuram Circle, K.G. Nagar, Bangalore, India.

# Introduction

ealth is a positive concept accentuating social and personal assets as well as physical and psychological aptitudes. In a healthy condition, an individual can satisfy needs and deal with interpersonal, social, biological, and physical environments. Therefore, it is a resource for every day [1]. In the perspective of understanding health, it is required to focus on the concept of global health. It is an area for study, investigation, and practice that places precedence on refining health and attaining justice of health for all people internationally. Global health emphasizes worldwide health issues, causes, and solutions, includes numerous corrections within and beyond the health sciences and indorses interdisciplinary association [2]. In addition to health, quality of life involves the standard of living, the quality of housing, and the neighborhood in which one lives, job satisfaction, and many other factors. According to the World Health Organisation (WHO), health is defined as a "state of complete mental, physical, and social well-being and not only the absenteeism of disease or disability. Health brings "broadness"- nothing is missing from the person; it brings "proper functions"-everything proficiently [3]. Biochemical anthropometric variables are directly connected to health. For Cardiovascular Disease (CVD), diabetes is one of the cause. Recent decades have seen a striking rise in diabetes dominance across the globe [4].

According to the World Health Organization (WHO), the global prevalence of anemia is 24.8%, which means about 1.62 billion people worldwide [5]. A decrease in the level of hemoglobin is associated with reduced health-related quality of life, congestive heart failure, and increased mortality in chronic kidney disease [6]. The previous studies has described that bhramari (humming bee breath) is a yoga practice, in which subjects should sit in any meditative posture, inhale through both nostrils, and while exhaling, produce the sound of a humming bee. More the bhramari timing, the more will be the lungs capacity [7]. The effect of pulse rate is also measured at the time of diagnosis. High pulse rate, at the time of diagnosis, is strongly associated with cardio related risks [8]. For the person's health condition and physiological stability, respiratory rate also provides information. An abnormal respiratory rate is a strong pointer that a health crisis is about to happen [9]. Blood pressure is the pressure exerted by blood on the walls of blood vessels while flowing. Globally, the cause of death is high blood pressure, and also it is the second foremost cause of debility next to childhood malnutrition. More than 80% of the adults are at threat from their blood pressure [10].

According to modern science, the gross (physical) body is made up of packets of energy. The ancient indian scripture taittiriya Upanishad has mentioned that the physical body is made of Annam, and the Annam is called food, which consists of five elements (earth, water, space, air, and fire). The yogic diet mentioned in Katha Upanishad and Hatha yoga scriptures consists of cow milk, cow ghee, sprouts, fruits, which is easy to digest and helps to maintain the physical and mental health. The Bhagavadgita highlights three categories of food tamasika (which is stale, tasteless, stinking, cooked overnight and impure), rajasika (that are bitter, sour, saline, over-hot, pungent, dry and burning), and satvika (that increase vitality, energy, vigour, health, joy and cheerfulness) based on the characteristics of food and its influence on human personality. The quantity of food, place, time, the mental state also contributes equally to maintain positive health [11]. Medical research centers emphasize lifestyle modification consisting of diet, normalization of body weight, and aerobic exercise as factors in treating noninsulin-dependent diabetes mellitus (NIDDM). Diet and lifestyle modification can be in controlling non-insulindependent diabetes mellitus (NIDDM) and reducing risk factors linked with macrovascular complications [12].

The other diet known as the pistachio diet also improved endothelial function, blood glucose level, some indices of inflammation, and oxidative status in healthy young men. Studies have also shown that frequent nut consumption decreases the risk of coronary artery disease [13]. A low-carbohydrate ketogenic diet (LCKD) has also shown beneficial effects in patients with type 2 diabetes, including reducing anti-diabetic medication dosage [14]. Hemoglobin determination is considered as a screening index valuable in describing various degrees of iron deficiency anemia. Dietary factors play a role in the growth of iron deficiency [15]. A Diet of calorie consisted of moderate carbohydrate, high protein, and rich in vitamins with a high amount of vegetables and fruits can increase the hemoglobin level [16]. The study also shows; changes in anthropometric variables like body weight, hip circumference, and waist circumference due to specific dietary intake [17]. There is an intensive investigation of the relationship between diet and blood pressure in recent years. A vegetarian diet shows lesser BP values in hypertensive subjects [18]. The diet approach is recommended to lower blood pressure. The diet improves cardiovascular risk factors

and beneficial in increased cardiometabolic risk [19]. The dietary approach to stop hypertension shows a high reduction in blood pressure and improvement in autonomic and vascular functions [20].

Pacagavya, as given in Ayurveda, consist of five substances obtained from cow namely, urine, dung, milk, ghee, and curd [21], and this diet is called as a pancagavya diet (PD). The Bos indicus (Indian) cow is known as kamadhenu (divine bovine-goddess/cow of plenty), signifying its nourishing nature, similar to a mother. According to the Indian scripture, The Indian sage maharshi vashistha served the divine kamadhenu cow, and Indian sage maharshi dhanavantari offered a wonderful medicine pancagavya to humanity [22]. Many formulations mentioned in Ayurveda describe the use of pancagavya components either as a single ingredient or in combination with drugs of herbal, animal, or mineral origin [21]. The cow milk consists of essential nutrients that are good for health, such as vitamins A, B, C, carotenes, and proteins. It contains the low calorific value and less cholesterol. It is a good animator for human health, easily digestible, and it also plays a bioprotective role [22].

Cow curd is the removal of three humors of the body and a blood purifier. It is beneficial for gastrointestinal disorders, piles, and blood-related problems. It is one of the most health-giving among all food items. In a non-drug manner, it helps to manage infections as it is an efficient anti-infection. Buttermilk and cow curd helps to control the growth of harmful microorganisms [23]. Cow's ghee enhances the body's resistance to infections, intelligence, eyesight, voice quality, and memory. It is for cholesterol and a heart patient as well as it is an anti-aging agent. It purifies the blood to an extent, and it also improves physical and mental health [23]. Ayurveda mentioned the formulation of pancagavya ghee, which is useful against anemia, fever, inflammations, and liver disorder [21]. Cow urine is used to remove the blockage in arteries, used for arthritis, psoriasis, eczema, diabetes, heart attack, piles, prostrate, fits, migraine, ulcer, acidity, constipation, avnecological problems, nose and ear problems [24]. Recently cow urine has been granted U.S. Patents (No. 6896907 and 6410059) for its use along with antibiotics for the fight against cancer and to control bacterial infections [25]. Cow urine helps to enhance immune responses in the body. Several elements in the body can be balanced by cow urine. Total salts present in cow urine are 24 in numbers [23]. In treating diseases like respiratory diseases, chronic renal failure, hepatitis A, B, and C, urological disorders, asthma, and cancer, cow urine plays an important role. It also acts as a disinfectant against many diseases like various kinds of allergies, acne vulgaris, scabies, eczema, and psoriasis [26]. In ancient times cow dung was widely used as fertilizer. Goumayarash is used as a skin tonic and useful in many skin related disorders like gangrene, psoriasis, eczema. The properties which cow dung includes are antibacterial, antifungal, and antiseptic [23].

#### Materials and Methods II.

Pancagavya diet (PD) group and nonpancagavya diet (NPD) group were recruited from Delhi, Haryana, and Rajasthan states of India, and its demographic details are given in table 1. The data for both the groups were collected between the period of January 2020 and February 2020. The sample size was calculated by using G-power software; based on; the previous study, the sample size was 76 with alpha 0.05, power 0.95, effect size 0.84 [27]. The assessments of the two groups were based on people adhering to PD and NPD for more than two years were considered. People with psychiatric ailments underwent any recent surgery, infectious disease, and female under menstruation and pregnancy were excluded from the study. Group of PD was directly or indirectly consumers of Bos indicus cow's products mainly of milk, curd, and clarified butter (ghee), cow urine, and cow dung. Nonpancagavya diet group was consumers of NPD diet, including buffalo, jersey cow, or any other animal's milk, ghee, curd, and grains produced by UREA/DAP and other pesticides more than two years are considered. In the present study biochemical variables, blood glucose, hemoglobin, and anthropometric variables bhramari time, pulse rate, respiratory rate, and blood pressures are measured. Bhramari (breath-holding time is a yoga practice, in which subjects sits in any meditative posture, inhale through both nostrils, and exhale, produce the sound of a humming bee) [7]. Data analysis was done by using JASP software with Shapiro-Wilk test for normality, and independent sample t-test was performed.

#### III. RESULT

The result showed that there was a significant difference between the pancagavya diet group compared to the non-pancagavya diet group for blood glucose (p<0.05), but there is no significant difference between both groups (p < 0.975) in levels of hemoglobin as shown in table 2. For anthropometric variables, there is a significant difference in bhramari time (p < 0.001), pulse rate (p=0.02), respiratory rate (p<0.001), systolic blood pressure (p < 0.01), and for diastolic blood pressure (p < 0.05).

#### IV. DISCUSSION

The random blood glucose level used as a biomarker showed a significantly lower in the pancagavya diet subjects than the subjects consuming nonpancagavya diet. The bhramari time with exponential significance, pulse rate with the highly significant and respiratory rate with exponential significance in PD group compared to the NPD group. The Subjects following PD had less measurement in both systolic and diastolic blood pressure compared to NPD. These indicate that the PD group have an opportunity for a better life-style, which comprises physical, mental, and spiritual well-being. The pancagavya have much application like in treating many diseases and to increase the body resistance to fight diseases medicines prepared from panchagavya are effective [22], therapeutic benefits of cow urine in managing cancer [27], practical application of pancagavyha products in the field of agriculture, to rejuvenate the soil health [24], based on the synergistic and systematic harnessing of energies from cows, plants, and earth [22], pancagavya ghrita, is also one of the formulations mentioned in Ayurveda which is prepared with all five components of panchagavya viz cow milk, ghee, urine, dung and curd in equal proportions useful for rejuvenation [21]. As the side effects of antibiotic medicine have harmful, one need to look into new therapeutic approach like panchagavya to remove diseases and to control infections.

#### V. Conclusions

Pancagavya diet group had less random blood glucose level, and have more bhramari time, less pulse rate, respiratroy rate, and blood pressure. Pancagavya diet had shown a more positive impact on health compared to the non-pancagavya diet.

Sources of funding

None.

Conflict of interest

None.

Acknowledament:

Author would like to thank Mr. Vikas Yadav, Mr. Pawan, and Mr. Kamal Phogat for their help in data collection.

Table 1: Demographic details

Partio	culars	PD	NPD
Number o	f subjects	40	40
Age(year) me	an <u>+</u> SD value	42.12 <u>+</u> 13.66	42.22 <u>+</u> 16.17
Gender	Male	29	29
	female	11	11
Occupation	Agriculture	29	24
	Job	1	4
	Business	5	3
	Student	1	5
	Housewife	2	3
	Others	2	1
Diet (from	2 years	5	0
years)	3 years	6	1
	More than 5 years	29	39

Legend: PD- Pancagavya diet.

NPD-Non-pancagavya diet.

Table 2: Result of statistical analysis

Domain	PD Mean±SD	NPD Mean±SD	T-value	<i>p</i> -value	Cohen's d
Biochemical variables					
Blood glucose (mg/dl)	110.63±20.08	135.34±63.08	2.361	0.021*	0.528
Hemoglobin (gm/dl)	13.55±1.90	13.52±2.08	0.062	0.975	0.014
Anthropometric variables					
Bhramari time (sec)	20.17±6.49	12.92±5.67	5.315	<0.001***	1.188
Pulse rate (bpm)	76.92±4.99	83.37±11.65	3.218	0.002**	0.720
Respiratory rate (cpm)	13.97±3.39	18.22±3.97	5.141	<0.001***	0.720
Sys. BP(mmhg)	118.65±7.67	127.35±15.67	3.152	0.002**	0.705
Dia. BP(mmhg)	78.40±6.93	82.45±10.20	2.077	0.041*	0.464

Legend: PD- Pancagavya diet.

NPD-Non-pancagavya diet

Grant support or other sources of funding None

Conflict of interest None

# References Références Referencias

- 1. McCartney G, Popham F, McMaster R, Cumbers A. Defining health and health inequalities. Public Health 2019;172;22-0.
- Koplan JP, Bond TC, Merson MH, Reddy KS, Rodriguez MH, Sewankambo NK, et al. Towards a common definition of global health. The Lancet 2009;373;1993-5.
- 3. John EWJ. Standards for validating health measures: Definition and content. Journal of Chronic Diseases 1987;40;473-0.
- Bragg FLiL, Bennett D, Guo Y, Lewington S, Bian Z, Yang L, et al. Association of random plasma glucose levels with the risk for cardiovascular disease among Chinese adults without known diabetes. JAMA Cardiol 2016;1;813-3.

- 5. Vibha BKM. Comparative study of prevalence of anaemia in vegetarian and nonvegetarian women of udaipur city, Rajasthan. J Nutr Food Sci 2015; s3; 1-6.
- Cote C, Zilberberg MD, Mody SH, Dordelly LJ, Celli B. Haemoglobin level and its clinical impact in a cohort of patients with COPD. ERJ Open Res 2007;29;923-9.
- Mooventhan AVK. Effect of Bhramari pranayama and OM chanting on pulmonary function in healthy individuals: A prospective randomized control trial. Int J Yoga 2014;7;104-0.
- Kurgansky KE, Schubert P, Parker R, Djousse L, Riebman JB. Gagnon DR. et al. Association of pulse rate with outcomes in heart failure with reduced ejection fraction: A retrospective cohort study. BMC Cardiovasc Disord 2020;20;1-1.
- Lim WS, Carty SM, Macfarlane JT, Anthony RE, Christian J, Dakin KS, et al. Respiratory rate measurement in adults - how reliable is it? Respir Med 2002;96;30-3.

- 10. He FJ, MacGregor GA. Blood pressure is the most important cause of death and disability in the world. Eur Heart J Suppl 2007;9;23-8.
- 11. Kwon DY, Tamang JP. Religious ethnic foods. Journal of Ethnic Foods 2015;2;45-6.
- 12. James RB, Tiffany J, Inkeles SB. Diet and exercise in the treatment of NIDDM. The need for early emphasis. Diabetes Care 1994;17;1469-2.
- 13. Sari I, Baltaci Y, Bagci C, Davutoglu V, Erel O, Celik H, et al. Effect of pistachio diet on lipid parameters, endothelial function, inflammation, and oxidative status: A prospective study. Nutrition 2010; 26; 399-4.
- 14. Hussain TA, Mathew TC, Dashti AA, Asfar S, Al-Zaid N, Dashti HM. Effect of low-calorie versus lowcarbohydrate ketogenic diet in type 2 diabetes. Nutrition 2012; 28;1016-1.
- 15. Vibha BKM. Comparative study of prevalence of anaemia in vegetarian and non vegetarian women of Udaipur City, Rajasthan. J Nutr Food Sci 2015;S3;1-6.
- 16. Bajpai S, Choudhary V, Sahu GK. Changes in body mass index, blood pressure and haemoglobin level of carcinoma patients: A study on the effect of designed diet. International Journal of Health Science and Research 2016; 6; 389-4.
- 17. Kasim SE, Martino S, Kim PN, Khilnani S, Boomer A, Depper J, et al. Dietary and anthropometric determinants of plasma lipoproteins during a longterm low-fat diet in healthy women. Am J Clin Nutr 1993:57:146-3.
- 18. Jenner DA, English DR, Vandongen R, Beilin LJ, Armstrong BK, Miller MR, et al. Diet and blood pressure in 9-year-old Australian children. Am J Clin Nutr 1988;47;1052-9.
- 19. Siervo M, Lara J, Chowdhury S, Ashor A, Oggioni C, Mathers JC. Effects of the dietary approach to stop hypertension (DASH) diet on cardiovascular risk factors: A systematic review and meta-analysis. Br J Nutr 2015;113;1-5.
- 20. Blumenthal JA, Babyak MA, Hinderliter A, Watkins LL, Craighead L, Lin PH, et al. Effects of the DASH diet alone and in combination with exercise and weight loss on blood pressure and cardiovascular biomarkers in men and women with high blood pressure: The ENCORE study. Arch Intern 2010;170; 126-5.
- 21. Achliya GS, Kot NR, Wadodka, SG, Dorle AK. Hepatoprotective activity of panchagavya ghrita hepatoprotective activity of panchagavya ghrita against carbontetrachloride induced hepatotoxicity in rats. Indian J Pharmacol 2003;35;308-1.
- 22. Dhama K, Chakraborty S, Tiwari R. Panchgavya therapy (Cowpathy) in safeguarding health of animals and humans - A review. Res. Opin. Anim. Vet. Sci 2013;3;170-8.

- 23. Dhama K, Khurana S, Karthik K, Tiwar R, Malik Y, Chauhan R. Panchgavya: Immune-enhancing and therapeutic perspectives. Journal of Immunology and Immunopathology 2016;16;1-1.
- 24. Mohanty I, Senapati MR, Jena D, Palai S. Innovare academic sciences diversified uses of cow urine. Int J Pharm Pharm Sci 2014;6;6-8.
- 25. Ritu P, Sahni YP, Swatantra KS and Sulochana S. Effect of panchgavya on central actions in albino rats. Pharma science monitor 2013;4;3940-6.
- 26. Vats S and Miglani K. Synergistic antimicrobial effect of cow urine and azadirachta indica on infectious microbes. Int J Pharm Sci 2011;2;1781-5.
- 27. Jain NK, Gupta VB, Garg R, Silawat N. Efficacy of cow urine therapy on various cancer patients in Mandsaur District, India-A survey. International Journal of Green Pharmacy 2010;4;29-5.

# This page is intentionally left blank



# Global Journal of Medical Research: k Interdisciplinary

Volume 20 Issue 13 Version 1.0 Year 2020

Type: Double Blind Peer Reviewed International Research Journal

Publisher: Global Journals

Online ISSN: 2249-4618 & Print ISSN: 0975-5888

# Comparison of ATP Values on Vegetables Cutting Boards before and after Alcohol Disinfection

By Naomi Katayama, Akemi Ito, Mayumi Hirabayashi, Natuki Sasaki & Moe Inuzuka

Nagoya Women's University

Abstract- Sanitary control of cutting boards in the kitchen is important to prevent food poisoning. To preventing secondary and tertiary contamination of food poisoning bacteria, it is necessary to know the hygiene status of cooking utensils. Therefore, in this study, we compared the values after cooking, washing, and spraying 70% alcohol on cutting boards for vegetables using the ATP test and microbiological test. As a result, the ATP value after spraying with alcohol was 100 or less, which was better than that after washing. Microbial test results showed that microorganisms were present on the vegetable cutting board even after spraying with 70% alcohol. Since microorganisms are present even after spraying with alcohol, it is possible that the growth of microorganisms will occur again if the vegetable cutting boardleft in a moist state at room temperature. When using a vegetable cutting board left unattended, it is necessary to wash repeatedly and spray it with alcohol.

Keywords: ATP wiping test, cutting board, hygiene education, double wash.

GJMR-K Classification: NLMC Code: QV 250



Strictly as per the compliance and regulations of:



© 2020. Naomi Katayama, Akemi Ito, Mayumi Hirabayashi, Natuki Sasaki & Moe Inuzuka. This is a research/review paper, distributed under the terms of the Creative Commons Attribution-Noncommercial 3.0 Unported License http://creativecommons.org/licenses/by-nc/3.0/), permitting all non-commercial use, distribution, and reproduction in any medium, provided the original work is properly cited.

# Comparison of ATP Values on Vegetables Cutting Boards before and after Alcohol Disinfection

Naomi Katayama <sup>a</sup>, Akemi Ito <sup>a</sup>, Mayumi Hirabayashi <sup>a</sup>, Natuki Sasaki <sup>a</sup> & Moe Inuzuka <sup>a</sup>

Abstract- Sanitary control of cutting boards in the kitchen is important to prevent food poisoning. To preventing secondary and tertiary contamination of food poisoning bacteria, it is necessary to know the hygiene status of cooking utensils. Therefore, in this study, we compared the values after cooking, washing, and spraying 70% alcohol on cutting boards for vegetables using the ATP test and microbiological test. As a result, the ATP value after spraying with alcohol was 100 or less, which was better than that after washing. Microbial test results showed that microorganisms were present on the vegetable cutting board even after spraying with 70% alcohol. Since microorganisms are present even after spraying with alcohol, it is possible that the growth of microorganisms will occur again if the vegetable cutting boardleft in a moist state at room temperature. When using a vegetable cutting board left unattended, it is necessary to wash repeatedly and spray it with alcohol.

Keywords: ATP wiping test, cutting board, hygiene education, double wash.

# I. Introduction

are should taken when cleaning vegetable cutting boards, as compared to meat and fish cutting boards, it may not be possible to wash them carefully due to the lack of sliminess<sup>1)</sup>. In this study, we used the ATP test and microbial test to compare the ATP value and the number of microbial bacteria immediately after cooking, washing, and spraying 70% alcohol on cutting boards for vegetables. The ATP value is preferably 100 or less, and the number of microbial bacteria is preferably free. We reported the results of the actual ATP test and the microbiological test.

# II. Materials and Methods

## a) Kitchen vegetable cutting board

The twelve vegetables cutting board prepared in the kitchen were stored in the sterilization storage the day before cooking. Cooking done in two places, and six cutting boards used for each.

Author  $\alpha$   $\sigma$   $\rho$ : Graduate School of Nagoya Women's University, Nagoya City, Japan.

Author α ω ¥: Nagoya Women's University, Nagoya City, Japan.

Author a: Department of Otorhinolaryngology, Nagoya University Graduate School of Medicine, Nagoya, Japan.

Corresponding Author α: Nagoya Women's University, Nagoya City, Japan. e-mail: naomik@nagoya-wu.ac.jp

# b) ATP inspection procedure

Each of the twelve cooks carried a vegetable cutting board for the vegetable of their work and brought it to the cooking table. Still, the inspector always performed an ATP inspection before using vegetables with a cutting board. Then, each cook finished the work, washes the cutting board firmly with detergent and sponge, rinse with running water for 30 seconds or more. Then, each cook repeated this process twice (as same as the last report<sup>1)</sup>). The inspector performed an ATP inspection after using vegetable with a cutting board, again. Then, each cook sprayed 70% alcohol on the cutting board after washing. At last, the inspector performed an ATP inspection after using vegetables with a kitchen cutting board. The ATP test kit used manufactured by KIKKOMAN.

# c) Stamp test inspection procedure

Five types of stamp test (General bacteria, Staphylococcus aureus, Escherichia coli, Salmonella, Vibrio parahaemolyticus) used. The stamp test conducted by the inspector at the same time as the ATP. The stamp test was colony-counted after culturing in an incubator at 38 degrees for three days. The stamp test by MISSUI.

# d) Statistical processing

The results obtained compared using statistical methods. The data statistically processed, was subjected to an F test to determine whether to use a parametric test or nonparametric test. When there is no difference in the F test, the presence or absence of a significant difference confirmed using the student t-test with or without a correspondence. If there was a difference in the F test, the presence or absence of a significant difference was confirmed using the Wilcoxon test with a pair or the Mann-Whitney test without correlation.

# III. RESULTS

a) ATP value results before and after alcohol disinfection

Tables 1 and 2 show the results of ATP wiping tests on vegetables cutting board before and after alcohol disinfection. The ATP value was statistically significantly lower after washing than after cooking. However, the ATP value did not fall below 100. The ATP

value after 70% alcohol spraying was 100 or less. The ATP value was statistically significantly lower after 70% alcohol spraying than after cooking.

Table 1. ATP test value and statistical processing result of cutting board 1.

	No alcohol	treatment	Alcohol treatment	
For vegetables	Before washing After washing		After washing	After alcohol
1	176205	863	863	10
2	909793	68	68	10
3	6543	39	39	44
4	15	42	42	26
5	38244	283	283	11
6	14200	5790	5790	17
Average value	190833.3	1180.8	1180.8	19.7
SD	358322.3	2279.9	2279.9	13.4
F test	P=0.00	001**	P=0.0001**	
Student-t*				
Wilcoxon	P0.0-	46*	P-0.0	)46:
F test	P=0.0001**			
Student-t*				
Wilcoxon		P=0.0	)46*	

\*Paired Student-t test \* P<0.05, \*\* P<0.01

Table2. ATP test value and statistical processing result of cutting board 2.

	No alcohol	treatment	Alcohol treatment	
For vegetables	Before washing	After washing	After washing	After alcohol
1	9194	2630	2630	18
2	1103	56	56	35
3	48126	449	449	45
4	3168	52	52	22
5	136610	259	259	3
6	3983	616	616	23
Average value	33697.3	677.0	677.0	24.3
SD	53435.9	981.9	981.9	14.4
F test	P=0.00	001**	P=0.00	01**
Student-t*				
Wilcoxon	P=0.	28*	P-0.0	28*
F test	P=0.0001**			
Student-t*				
Wilcoxon	P=0.028*			
	*D: 1 G: 1 * D 0 05 ** D 0 01			

\*Paired Student-t test \* P<0.05, \*\* P<0.01

b) Stamp test results before and after alcohol disinfection

Tables 3,4,5,6,7,8.9.10.11 and 12 show the results of ATP wiping tests on vegetables cutting board before and after alcohol disinfection. The result of common bacteria, Staphylococcus aureus and Vibrio parahaemolyticus was that microorganisms could be present on the cutting board even after 70% alcohol spraying. However, the number of microorganisms reduced compared to after cooking. In the case of E. Coli and Salmonella, the number of microorganisms decreased statistically significantly after spraying with 70% alcohol.

Table 3 Number of general bacteria on cutting board 1. and statistical processing

		result			
	No alcohol treatment		Alcohol to	reatment	
For vegetables	Before washing	After washing	After washing	After alcohol	
1	298	110	110	0	
2	8	10	10	13	
3	22	3	3	0	
4	50	0	0	3	
5	7	42	42	0	
6	8	1	1	0	
Average value	65.5	27.7	27.7	2.7	
SD	115.1	43.3	43.3	5.2	
F test	P=0.0	)16*	0.000	1**	
Student-t*					
Wilcoxon	P=0.	249	P-0.3	345	
F test	P=0.0001**				
Student-t*					
Wilcoxon		P=0.046*			
	*Paired_Student-t test_* P<0.05, ** P<0.01				

Table 4 Number of general bacteria on cutting board 2. and statistical processing

	result				
	No alcohol treatment		Alcohol t	reatment	
For vegetables	Before washing	After washing	After washing	After alcohol	
1	87	120	120	0	
2	1	2	2	18	
3	9	1	1	0	
4	90	14	14	0	
5	200	47	47	0	
6	3	0	0	3	
Average value	65.0	30.7	30.7	3.5	
SD	78.0	47.3	47.3	7.2	
F test	P=0	.124	P=0.0001**		
Student-t*	P=0	.273			
Wilcoxon			P=0.	345	
F test	P=0.0001**				
Student-t*					
Wilcoxon		P=0.	.138		
·					

\*Paired Student-t test \* P<0.05, \*\* P<0.01

Table 5 Number of E. coli on cutting board 1. and statistical processing result

	No alcohol treatment		Alcohol treatment		
For vegetables	Before washing	After washing	After washing	After alcohol	
1	14	3	3	0	
2	3	3	3	2	
3	6	1	1	0	
4	23	0	0	0	
5	30	30	30	0	
6	3	0	0	0	
Average value	13.2	6.2	6.2	0.3	
SD	11.3	11.8	11.8	0.8	
F test	P=0	.463	P=0.00	01**	
Student-t*	P=0	.110			
Wilcoxon		P=0.043*			
F test	P=0.0001**				
Student-t*					
Wilcoxon		P=0.028*			
	*Paired Stud	ant t tact * D/O	15 ** P/0.01		

\*Paired Student-t test \* P<0.05, \*\* P<0.01

 $Table\ 6\quad Number\ of\ E.\ coli\ \ on\ cutting\ board\ 2.\ and\ statistical\ processing\ result$ 

	No alcohol treatment		Alcohol treatment		
For vegetables	Before washing	After washing	After washing	After alcohol	
1	7	1	1	0	
2	0	0	0	0	
3	21	11	11	0	
4	16	4	4	4	
5	200	0	0	0	
6	60	0	0	0	
Average value	50.7	2.7	2.7	0.7	
SD	76.1	4.4	4.4	1.6	
F test	P=0.00	001**	P=0.0001**		
Student-t*					
Wilcoxon	P=0.0	043*	P=0.	075	
F test	P=0.0001**				
Student-t*					
Wilcoxon		P=0.0	)43*		

\*Paired Student-t test \* P<0.05, \*\* P<0.01

Table 7 Number of Staphylococcus aureus on cutting board 1. and statistical processing result

	No alcohol	treatment	Alcohol t	Alcohol treatment		
For vegetables	Before washing	After washing	After washing	After alcohol		
1	28	18	18	4		
2	1	2	2	3		
3	100	0	0	1		
4	7	0	0	0		
5	4	200	200	0		
6	15	1	1	0		
Average value	25.8	36.8	36.8	1.3		
SD	37.6	80.2	80.2	1.8		
F test	P=0.0	)44*	P=0.0001**			
Student-t*						
Wilcoxon	P=0.	463	P=0.	345		
F test	P=0.0001**					
Student-t*						
Wilcoxon		P=0.046*				
	*Doined Ctude	mt t toot * D <0.04	*Pointed Student t toot * D <0.05 ** D <0.01			

<sup>\*</sup>Paired Student-t test \* P<0.05, \*\* P<0.01

### 8 Number of Staphylococcus aureus on cutting board 2. and statistical processing

	No alcohol treatment		Alcohol tr	eatment
For vegetables	Before washing	After washing	After washing	After alcohol
1	2	2	2	0
2	0	0	0	2
3	0	0	0	0
4	7	0	0	0
5	250	56	56	0
6	33	0	0	0
Average value	48.7	9.7	9.7	0.3
SD	99.4	22.7	22.7	0.8
F test	P=0.0	01**	P=0.00	01**
Student-t*				
Wilcoxon	P=0.109 P=0.423			23
F test	P=0.0001**			
Student-t*				
Wilcoxon		P=0.1	06	

<sup>\*</sup>Paired Student-t test \* P<0.05, \*\* P<0.01

Table 9 Number of Salmonella on cutting board 1. and statistical processing result

	No alcohol treatment		Alcohol tro	eatment
For vegetables	Before washing	After washing	After washing	After alcohol
1	210	3	3	0
2	48	0	0	27
3	3	100	100	2
4	240	35	35	8
5	1	9	9	0
6	175	0	0	0
Average value	112.8	24.5	24.5	6.2
SD	107.9	39.3	39.3	10.7
F test	P=0.0	)13*	P=0.00	3**
Student-t*				
Wilcoxon	P=	173	P=0.2	81
F test		P=0.00	001**	
Student-t*				
Wilcoxon		P=0.0	)28*	
*D-11 C1				

Table 10 Number of Salmonella on cutting board 2. and statistical processing result

	No alcohol treatment		Alcohol tı	reatment
For vegetables	Before washing	After washing	After washing	After alcohol
1	378	0	0	0
2	3	2	2	0
3	8	0	0	6
4	25	0	0	0
5	1	5	5	0
6	43	0	0	0
Average value	76.3	1.2	1.2	1.0
SD	148.6	2.0	2.0	2.4
F test	P=0.00	001**	P=0.3	335
Student-t*			P=0.9	914
Wilcoxon	P=0.	075		
F test	P=0.0001**			
Student-t*				
Wilcoxon	P=0.028*			
*D-:1 Ct-:1+				

\*Paired Student-t test \* P<0.05. \*\* P<0.01

Table 11 Number of Vibrio parahaemolyticus on cutting board 1. and statistical processing result

	No alcohol treatment		Alcohol treatment		
For vegetables	Before washing	After washing	After washing	After alcohol	
1	11	15	15	0	
2	0	1	1	1	
3	20	0	0	0	
4	0	15	15	0	
5	61	0	0	0	
6	0	92	92	0	
Average value	15.3	20.5	20.5	0.2	
SD	23.8	35.8	35.8	0.4	
F test	P=0.	172	P=0.00	001**	
Student-t*	P=0.811				
Wilcoxon	P=0.109				
F test	P=0.0001**				
Student-t*					
Wilcoxon	P=0.144				
*D-:1 Ct-1+ * D-0.05 ** D-0.01					

\*Paired Student-t test \* P<0.05, \*\* P<0.01

Table 12 Number of Vibrio parahaemolyticus on cutting board 2. and statistical processing result

No alcohol treatment		Alcohol treatment		
Before washing	After washing	After washing	After alcohol	
192	31	31	0	
40	0	0	0	
0	0	0	0	
0	40	40	2	
0	100	100	1	
28	3	3	0	
43.3	29.0	29.0	0.5	
74.8	38.8	38.8	0.8	
P=-0.067		P=0.0001**		
P=0.	706			
P=0.068				
P=0.0001**				
P=0.225				
	Before washing 192 40 0 0 28 43.3 74.8 P=-0	Before washing	Name	

\*Paired Student-t test \* P<0.05, \*\* P<0.01

#### IV. Discussion

On cutting boards for vegetables, hygiene tests performed on the ATP value and the number of microorganisms. For the microbiological test, a selective medium of general bacteria, Escherichia coli, Staphylococcus aureus, Salmonella, and Vibrio parahaemolyticus used. The ATP level and the number of microorganisms decreased after washing as compared with after cooking. Furthermore, after alcohol spraying, the ATP level, the number of E. Coli, and the number of Salmonella bacteria decreased statistically significantly. However, the bacteria did not disappear. Microorganisms are more likely to grow if they are moist, at the right temperature, and hove nutrients. If the cutting board is left unattended after cooking, it may be necessary to wash repeatedly and spray it with alcohol before use. The ATP test can show invisible microorganisms on the spot with visible numbers 1,2,3,4). Therefore, it is used in many places and is useful for hygiene education and food poisoning prevention<sup>5,6,7,8)</sup>. Although it takes time, it is useful for hygiene education to know the condition of food poisoning bacteria by conduction microbiological tests.

# Conclusions

As a result of the ATP test and microbiological test performed on the cutting board for vegetables, there are surviving bacteria that even after spraying 70% alcohol, so spray 70% alcohol firmly, and the cutting board left for a while is washed repeatedly and sprayed with alcohol before cooking. We think it's better to use it.

# Acknowledgments

We would like to thank all the cooks who participated in this experiment. Also, we would like to thank the inspectors who also performed the ATP inspection.

# References Références Referencias

- 1. Nante N, Ceriale E, Messina G, Lenzi D, Manzi P. Effectiveness of ATP bioluminescence to assess hospital cleaning: a review. (2017) J Prev. Med. Hyg. 58(2): E177-E183.
- 2. Amodio E, Dubi C. Use of ATP bioluminescence for assessing h eclealiness of hospital surfaces: a review of the published literature (1990-2012).(2014) J infect Public Health 7(2): 92-98.
- Aycieck H, Oquz U. Karci K. Comparison of results of ATP bioluminescence and traditional ygiene swabbing methods fro the determination of surface cleanliness at a hospital kitchen. (2006). Int J Hyg Environ Heatth. 209(2): 203-206.
- Osimani A, Garofalo C, Clementi F, Tavoletti S, Aquilanti L.Bioluminescence ATP monitoring for the routine assessment fo food contact surface cleanliness in a university canteen. (2014). Int J Environ Res Public Health 17; 11(10): 10824-10837.
- 5. Lee JH (2018) An investigation of Factors that influence Hygiene Practices at a small Day Care Center. (2018). J Food Prot. 81(1): 158-164.
- Stanley PE. A review of bioluminescent STP techniques in papid microbiology. (1989) J Biolumin Chemilumin 4(1): 375-380.
- Stannard CJ, Gibbs PA. Rapid microbiology: application s of bioluminescence in the food industry—a review. (1986) J Biolumin Chemilumin 1(1): 3-10.
- Griffith CJ, Coooper RA, Gilmore J, Davies C, Lweis M. An evaluation of hospital cleaning refimes and standards. (2000) J Hosp Infect. 45(1): 19-28.



# Global Journal of Medical Research: K Interdisciplinary

Volume 20 Issue 13 Version 1.0 Year 2020

Type: Double Blind Peer Reviewed International Research Journal

Publisher: Global Journals

Online ISSN: 2249-4618 & Print ISSN: 0975-5888

# A New Medical Device for Platelet Rich Plasma Filler

By Araco Antonino, M.D.

Univerity Tor Vergata of Roma

Introduction- Injectable fillers are used in aesthetic medicine to reduce visible signs of facial aging.

It involves loss of volume in the skin, muscle and superficial and deep fat compartments, superficial wrinkles and deep folds (1-2).

There are different products that vary on filling capacity, longevity, potentiality for causing allergic reaction, safety, indication.

Physicians select the most suitable agent for each patient by considering the advantage and disadvantage.

In fact, hyaluronic acid (HA) has the property to increase the skin's water-binding capacity (3-5).

Calcium hydroxylapatite (6) and poly-L-lactic acid (7) have been proved to stimulate autologous collagen (8-9).

Keywords: Growth factors, PRP, regenerative medicine, face wrinkles, perioral wrinkles, dermal matrix.

GJMR-K Classification: NLMC Code: WH 300



Strictly as per the compliance and regulations of:



© 2020. Araco Antonino, M.D. This is a research/review paper, distributed under the terms of the Creative Commons Attribution-Noncommercial 3.0 Unported License http://creativecommons.org/licenses/by-nc/3.0/), permitting all non-commercial use, distribution, and reproduction in any medium, provided the original work is properly cited.

# A New Medical Device for Platelet Rich Plasma Filler

Araco Antonino, M.D.

Keywords: Growth factors, PRP, regenerative medicine, face wrinkles, perioral wrinkles, dermal matrix.

#### Ī. Introduction

njectable fillers are used in aesthetic medicine to reduce visible signs of facial aging.

It involves loss of volume in the skin, muscle and superficial and deep fat compartments, superficial wrinkles and deep folds (1-2).

There are different products that vary on filling capacity, longevity, potentiality for causing allergic reaction, safety, indication.

Physicians select the most suitable agent for each patient by considering the advantage and disadvantage.

In fact, hyaluronic acid (HA) has the property to increase the skin's water-binding capacity (3-5).

Calcium hydroxylapatite (6) and poly-L-lactic acid (7) have been proved to stimulate autologous collagen (8-9).

Although fillers have lower immunogenicity and a good safety profile, several scientific articles have showed side effects (10).

Those can vary from allergic reaction (11), inflammatory nodule formation (12), infection (13), vision and the limited use in human loss (14)immunodeficiency (15).

Platelet rich plasma (Prp) has proved to achieve regenerative capacity on human tissues (16) and positive effect on facial dermal fibroblasts (17-18).

Autologous plasma filler was first introduced by Krajcik et al. in 1999 (19) and used for the treatment of different types and grades of facial wrinkles (20-22).

The main issues of the clinical use of plasma filler is its poor filling effect and duration.

In fact, the volume achieved just after the injection lasts only few hours because the plasma is reabsorbed.

The goal of our study was the preparation of a new medical filler device able to fill soft tissues for a reasonable period of time and also containing the regenerative Prp property.

Our first step was the formulation of a thermosensitive gelableto embed Prp, the second to evaluate the behavior of the formulation alone, in particular, its dispersibility and the homogeneity and the third the study of platelets and the growth factors behavior inside the formulation.

After a detailed research on the scientific literature (23,24) poloxamer 407 was chosen for our purpose.

Multiple reasons have corroborated this choice: first of all, the family of "poloxamer" polymers is listed in the US and European Pharmacopoeia (25) and is approved by FDA for parenteral use in humans and this ensure safety, biocompatibility and tolerability required for parenchymal use.

Furthermore, poloxamer 407, being a non-ionic block copolymer, does not negatively interact with the biomolecules embed in the gel, such as platelets and growth factors.

Again, it allows to develop thermosensitive formulations without the need of excipients in preparations.

#### Materials and Methods II.

# 1. Preparation of poloxamer formulation

Aqueous solutions of poloxamer 407, purchased from Sigma-Aldrich (Milano, Italy), prepared by the so-called "cold method".

Poloxamer powder was dissolved in bidistillate water at 277.15 K (4°C) under gentle stirring to facilitate the copolymer dissolution (26).

Solutions at poloxamer concentration of 15% (w/v) were prepared with this method and stored at 277.15 K (4 °C).

Differential scanning calorimetry (DSC) was used to determine the micellization and the gelation point.

Measurements were performed calorimeter Mettler 821<sup>e</sup> (Mettler-Toledo, Greifensee, Switzerland) equipped with a cooling system with liquid nitroaen.

Scans were recorded from 273.15 (0 °C) to 313.15 K (40 °C) at a heating rate of 5 K/min (27).

Measurements were performed in duplicate and the results were expressed as the mean of the two measurements.

Author: Contracted professor at Univerity Tor Vergata of Roma, Italy. e-mail: aracoantonino@gmail.com

## 2. Analysis of the Platelet

The analysis of the platelet contained inside the formulation could not be address by normal hemochrome automatic analysis systems.

For this reason, platelet numbers were firstly estimated by using a Burker's chamber and the Breker -Cronkite method.

Observations were performed with a Nikon Eclipse 80i microscope provided of a digital camera Nikon Coolpix 8400.

Breker - Cronkite solution was prepared adding 0,01 g of brilliant cresyl blue to 100 mL of 1% (w/v) ammonium oxalate aqueous solution.

Medical Device with Prp sample was diluted with Breker - Cronkite solution (dilution 1/100) in a Thoma pipette.

After five minutes of pipette manual shaking, a drop of diluted solution was charged in the Burker's chamber and observed at microscope magnification 40X.

We counted platelets presents in a big square of 1 mm<sup>2</sup> (Figure 1).

To obtain the number of platelets within 1  $\mu$ L of PRP, the average of platelets counts in three squares was corrected for the initial dilution (100) and for the height of chamber (10) (28).

For this reason, the final formula is: Platelet number counted X 1000.

Prp platelet size and number were also determined using an AccusizerTM 770 Optical Particle Sizer (PSS Inc., Santa Barbara, CA, USA), using the technology "single particle optical counting", coupled with an auto-dilution system AccusizerTM 770A Autodiluter PAT (PSS Inc., Santa Barbara, CA, USA).

100  $\mu$ L of solution with Prp were injected in the system and measurements were performed for 30 minutes to allow the analysis of all the particles present in the sample.

Saline (0.9% NaCl) was used for the analysis and results were expressed as total particles size and mean volume diameter (Figure 2).

# 3. HPLC analysis of growth factors

We analyzed the Epidermal Growth Factor (EGF) which is centrally involved in the regulation of key processes of the epithelia.

Calibration curve for RP-UHPLC analysis was obtained using an HPLC Jasco LC-2000 plus, equipped with the quaternary gradient pump PU 2089 Plus, the UV-vis detector diode array MD 2010 plus and injector Rheodyne.

Controlled EGF was purchased from Sigma-Aldrich (Milano, Italy).

The column used was a C18 Ascentis Express Peptide 150  $\times$  4.6 mm (L  $\times$  I.D.), 2.7  $\mu m$  and analysis were made combining two different mobile phases:  $H_2O/CH_2CN$  (95/5 %) + 0.1 % TFA (solvent A) and  $CH_3CN/H_2O$  (95/5 %) + 0.1 % TFA (solvent B), with a flow of 2.0 mL/min (29).

The mobile phases were degassed prior to use: cell and column temperature at 313.15 K (40 °C).

The calibration curve was performed in the concentration range of 0.086-0.4 mg/mL, with an injection volume of 5 µL.

# 4. Rheological analysis

Poloxamer rheological characterization was performed with the aim of better understanding the thermal behavior of poloxamer formulation.

A Stresstech HR Rheometer (Rheologia Instruments AB, Milano, Italy), equipped with a Peltier device for temperature control, was used for the rheological characterization.

Viscosity was measured with a cone-plate geometry (cone angle 1°, cone-plate diameter 40 mm).

Poloxamer formulation was analyzed in duplicate at different temperature (278, 283, 288, 293, 298, 303 and 308 K corresponding to 5, 10, 15, 20, 25, 30 and 35 °C).

For formulation in sol state, measurements were performed applying a stress range from 3.044 10<sup>-3</sup> to 1 Pa, while for temperatures at which the formulations were already at the gel state, measurements were carried out applying a stress range from 3.044·10<sup>-3</sup> to 10 Pa. 12.

## Spreadability

The gel was tested by two independent persons for the spreadability.

The gels stored at 277.15 K (4 °C) were withdraw from the recipients and spread on the skin surface.

## Preparation of Prp

For the preparation of the Prp, Plasma Active system (CE0373 - Medical Device s.r.l. via artigianato n.6, 52022 Meleto, Craviglia (AR) – Italy) was used.

It is composed of two vacutainer tubes (BD Vacutainer® Brand SST II) of 9 mm each with separator gel and anti-coagulant for harvesting 18 ml of peripheral blood.

Tubes were centrifugated for 5 minutes at 1800 rpm (Sorvall Legend XTR Centrifuge - Thermo Fisher Robert-Bosch-Straße 1 D -Langenselbold Germany) and 8 ml of Prp were collected from the upper part of the tube in a sterile matter and placed inside the Medical Device.

# 7. Preparation of poloxamer containing PRP

Five mL of Prp were added to 10 mL of thermosensitive gel.

The ratio of Prp/gel was chosen by considering the future application of the thermosensitive gel and the size of the final device (30).

Results have shown that 5 mL of Prp could be easily dispersed within 10 mL of the hydrogel by stirring the cold gel (4 °C) or by agitation of the closed container.

#### RESULT III.

# 1. Differential scanning calorimetry (DSC) results

DSC data show broad enthalpic transitions in the formulation due to poloxamer micellization.

In fact, in aqueous solution, with increasing temperature, poloxamer aggregates in micelles to minimize the free energy of solution.

Micellization temperatures decrease with increasing poloxamer concentration (Figure 3).

Also, micellization energy transition (ΔH<sub>mic</sub>) per amount of poloxamer increases with increasing concentration.

Data of micellization temperature and  $\Delta H_{mic}$  of poloxamer preparation is reported (Table 1).

Very interesting, in DSC thermograms is possible to see also a little enthalpic transition at temperature slightly higher than micellization point.

This peak is more evident in solutions with higher poloxamer concentrations and it has been previously attributed to the gelation transition (Figure 4).

In agreement with this little peak, gelation energy ( $\Delta H_{Qel}$ ) is very small (Table 2).

Gel temperatures found from this approach agree with results reported in the literature (27).

This DSC analysis give a first information about poloxamer solution thermal behavior.

In fact, the decrease of micellization and gelation temperature with increasing poloxamer concentration clearly shows how formulations with high poloxamer concentration (e.g. 25-30% w/v) pass to gel state at lower temperatures compared to formulations with a low poloxamer concentration.

Micellization and gelation temperatures plotted in Figure 5.

These results demonstrate that gelation point is strongly temperature and concentration dependent.

# 2. Rheological analysis result

In Figure 6 is shown the behavior of poloxamer formulation viscosities as a function of temperature.

In the first part of curves, solution viscosity decreases slightly on warming but, reached a certain temperature, is possible to see a steep increase in viscosity.

This viscosity increase can be attributed to the sol-to-gel transition.

Verv interesting, results obtained rheological characterization agree with gel temperatures obtained from the DSC data (Figure 3).

## 3. Spreadability results

The best spreading was individuated for the formulation containing 15% of poloxamer that did not gelify immediately after contact with the skin.

However, once the gel was spread it remained on surface as a thick gel.

## 4. Result of poloxamer containing PRP

Prp dispersed homogeneously into the matrix without forming lipid globules but changing the appearance of the gel from completely transparent to a light-yellow color (Figure 7A).

The formulation pH did not change with respect to the control formulation (blank gel).

The smell of the formulation containing PRP was slightly different from the blank gel, but it was still pleasant.

Prp loaded thermosensitive gel was stored at 4 °C for 1 month and its characteristics were evaluated 7 and 30 days after preparation.

After 7 days of storage, the gel color and smell were unchanged and, more important, Prp was still homogeneously dispersed (Figure 7B).

After 1 month of storage, the smell was unchanged while a slight phase separation was observed (Figure 7C).

This phase separation was reversible by simpler agitation.

In fact, by gently shaking the closed container, the gel assumed the same aspect and homogeneity of a fresh prepared formulation (Figure 7D).

This finding is of particular interest for the final use and we store the formulation at: 2-8 °C and shake if before use.

## Platelet behaviour

We analyzed 10 different samples of solution of poloxamer containing PRP after 12, 23, 73, 96, 120, 144, 168 hours from the preparation and expressed the results as mean.

Platelet remained intact for 72 hours and then numbers dropped rapidly as shown on Figure 8.

## 6. HPLC results of growth factors

Analysis made on the 10 different samples of solution of poloxamer containing Prp after 12, 23, 73, 96, 120, 144, 168 hours from the preparation showed that the EGF concentration increased progressively and reached his peak after 4 days and then decreased progressively.

The Egf concentration remained above 2,5  $\mu$ g/mL after 7 days (Figure 9).

#### IV. Discussion

Fillers are among the most performed cosmetic medicine procedures worldwide (31-32).

They are used mostly on the face for the reduction of superficial and deep wrinkles (32-34), for increasing the volumes of lips (35) and cheekbones (36) and for the redefinition of the mandibular profile (37).

There are different materials used for fillers including hyaluronic acid (38), collagen (39), calcium hydroxyapatite (40-41), and polycaprolactone (42), polymethylmethacrylate (43).

They vary depending on the filling capacity, due to their ability of recalling water (44), or even for the ability to stimulate collagen (45).

The advantages of commercial fillers are the simplicity of use, as they are supplied in single-use vials and the very low incidence of major complications as embolisms (46-48).

The disadvantages are the high cost, since to obtain appreciable results, several vials are required, and because of the limited duration, the treatment must be repeated periodically (49-50).

Again, possible side effects, although minor, create discomfort to the patients and stress to the doctors (51).

And definitely, fillers do not have regenerative activities.

Prp is used in dermatological clinic for its regeneration effects on dermal cells (17).

Different studies have already shown the positive effects on the treatment of facial wrinkles.

"Plasma filler" has always meant the simple injection of Prp into subcutaneous tissues (18, 20, 21).

But the main limitation of plasma fillers is the fact that it does not generate filling effects, if not for a few hours.

For this reason, we set up a new filler device able to combine volumetric and regenerative effects.

We faced different problems: first of all, the Prp is a fluid and to be well mixed inside the new filler device it is necessary that this is also in the liquid state.

Moreover, once injected into the tissues, this liquid would have to become a gel to guarantee the volumetric effect.

Furthermore, since both the platelets and the Prp have a low half-life outside the plasma, we had to guarantee the new filler device a chemical and physical composition that allows the survival of platelet and growth factors for several days.

So, we have chosen a polymer (poloxamer 407) that contains all these characteristics (23-24).

In fact, at low temperatures (2-8 C°) it is found in the liquid phase and this allows the simple mixing of the Prp inside it.

When the temperature reaches 32-37 ° C, the liquid passes to the gel state.

This ensures either the volumizing effect and the progressive release of growth factors within the tissues.

We performed precise measurements on both the number of platelets and EGF present in the new filler device at regular time intervals for 7 days.

EGF is known to be a potent stimulator of cell proliferation of various cells including keratinocytes, fibroblasts and vascular endothelial cells.

EGF stimulates the migration of keratinocytes and also stimulates fibroblasts and endothelial cells to promote the formation of granulation tissue (52-54).

These measurements were also difficult because platelets and the EGF could not be measured with the common automatic measurement systems.

The results found were encouraging for a clinical point.

In fact, the platelets were maintained vitality until 72 hours and then their number felled by freeing the granules containing the growth factors.

EGF, progressively increased its concentration with a peak after 72-96 hours and then slowly decreased until 7 days.

This new filler device could open new scenarios in facial rejuvenation and revolutionize the treatment of face wrinkles.

In fact, the advantages are evident: it would have volumizing and regenerative capacities.

This would allow wrinkles and grooves to be filled and regenerated at each application.

Furthermore, being autologous growth factors, there would be no side effects related to allergic reactions or inflammatory nodules.

Again, since the immune system is also present inside the plasma, this would help prevent the infections.

Finally, the cost of treatment would be cheaper than normal commercial filler, since for each session of treatment 8 ml of Prp are added to 16 ml of hydrogel to obtain a 24 ml filler.

The disadvantage lies on the Prp preparation.

## Conclusions

Our study has shown that it is possible to obtain a new type of filler able to have both filling capacity, due to the gelling effect of the material used, and a regenerative effect due to the presence of Prp.

Therefore, new clinical trials will be necessary to assess the duration of the volumizing effect in the different areas of the face.

# References Références Referencias

- Nkengne A, Bertin C. Aging and facial changes-documenting clinical signs, part 1: clinical changes of the aging face. Skinmed. 2013 Sep-Oct; 11(5): 281-6.
- In the shadow of the wrinkle: theories. Humbert P, Viennet C, Legagneux K, Grandmottet F, Robin S, Oddos T, Muret P. J Cosmet Dermatol. 2012 Mar; 11(1):72-8.
- Lim HK, Suh DH, Lee SJ, Shin MK. Rejuvenation effects of hyaluronic acid injection on nasojugal groove: Prospective randomized split face clinical

- controlled study. J Cosmet Laser Ther. 2014; 16: 32-36.
- 4. Farhi D, Trevidic P, Kestemont P, et al.; Emervel French Survey Group. The Emervel French survey: A prospective real-practice descriptive study of 1,822 patients treated for facial rejuvenation with a new hyaluronic acid filler. J Drugs Dermatol. 2013; 12:e88-e93.
- 5. Huang X, Liang Y, Li Q. Safety and efficacy of hyaluronic acid for the correction of nasolabial folds: A meta-analysis. Eur J Dermatol. 2013;23:592–599.
- 6. PavicicT.Calcium hydroxylapatite filler: an overview of safety and tolerability. J Drugs Dermatol. 2013 Sep;12(9):996-1002. Review.
- 7. Alam M, Havey J, Pace N, Pongprutthipan M, YooS. Large particle calcium hydroxylapatite injection for correction of facial wrinkles and depressions. J Am Acad Dermatol. 2011 Jul;65(1):92-6.
- 8. Breithaupt Fitzgerald Collagen Α, R. Stimulators: Poly-L-Lactic Acid and Calcium Hydroxyl Apatite. FacialPlast Surg Clin North Am. 2015 Nov: 23(4):459-69. doi: 10.1016/j.fsc. 2015.07.007. Review.
- 9. Stein P, Vitavska O, Kind P, Hoppe W, Wieczorek H, SchürerNY.The biological basis for poly-L-lactic acid-induced augmentation. J Dermatol Sci. 2015 Apr: 78(1):26-33.
- 10. Gilbert E, Hui A, Meehan S, Waldorf HA. The basic science of dermal fillers: past and present Part II: adverse effects. J Drugs Dermatol. 2012 Sep; 11(9):1069-77. Review.
- 11. Lucey P, Goldberg DJ. Complications of collagen fillers. Facial Plast Surg. 2014 Dec; 30(6):615-22. doi: 10.1055/s-0034-1396904. Epub 2014 Dec 23. Review.
- 12. Ledon JA, Savas JA, Yang S, Franca K, Camacho I. Nouri K. Inflammatory nodules following soft tissue filler use: a review of causative agents, pathology and treatment options.
- 13. Am J Clin Dermatol. 2013 Oct; 14(5):401-11.
- 14. Schlesinger TE, Cohen JL, Ellison S. Purpura and fillers: a review of pre-procedural, intraprocedural, and post-procedural considerations. J Drugs Dermatol. 2013 Oct; 12(10):1138-42. Review.
- 15. Loh KT, Chua JJ, Lee HM, Lim JT, Chuah G, Yim B, Puah BK. Prevention and management of vision loss relating to facial filler injections. Singapore Med J. 2016 Aug; 57(8):438-43.
- 16. Shuck J, Iorio ML, Hung R, Davison SP. Autologous fat grafting and injectable dermal fillers for human immunodeficiency virus-associated lipodystrophy: a comparison of safety, efficacy, and long-term treatment outcomes. PlastReconstr Surg. 2013 Mar; 131(3):499-506.
- 17. Yuksel EP, Sahin G, Aydin F, Senturk N, Turanli AYEvaluation of effects of platelet-rich plasma on

- human facial skin.. J Cosmet Laser Ther. 2014 Oct;16(5):206-8.
- 18. Kim DH, Je YJ, Kim CD, Lee YH, Seo YJ, Lee JH, Lee Y. Can Platelet-rich Plasma Be Used for Skin Rejuvenation? Evaluation of Effects of Platelet-rich Plasma on Human Dermal Fibroblast.AnnDermatol. 2011 Nov;23(4):424-31.
- 19. Cameli N. Mariano M. Cordone I. Abril E. Masi S. Foddai ML. Autologous Pure Platelet-Rich Plasma Dermal Injections for Facial Skin Rejuvenation: Flow Instrumental, Clinical, and Cytometry Assessment. Dermatol Surg. 2017 Jun; 43(6): 826-835.
- 20. Krajcik R, Orentreich DS, Orentreich N. A novel injectable autologous material for soft tissue augmentation. J Aesthetic Dermatol and Cosmet Surg 1999; 1:109-115.
- 21. Choi YJ1. Kim HS1. Min JH1. Nam JH1. Lee GY1. Kim WS1. A clinical study on the usefulness of autologous plasma filler in the treatment of nasolabial fold wrinkles. J Cosmet Laser Ther. 2017 Jun; 19(3):174-180. doi:10.1080/14764172.2016. 1248443. Epub 2017 Feb 21.
- 22. Naema Y Elnehrawy, MD, <sup>1</sup> Zeinab A Ibrahim, MD, PhD, <sup>1</sup> Azza M Eltoukhy, MD, PhD, <sup>1</sup> & Hala M Nagy, MD, PhD<sup>2</sup>. Assessment of the efficacy and safety of single platelet-rich plasma injection on different types and grades of facial wrinkles. Journal of Cosmetic Dermatology, 1-9, 2017.
- 23. Lynch MD, Bashir S. Applications of platelet-rich plasma in dermatology: A critical appraisal of the literature. J Dermatolog Treat. 2016;27(3):285-9.
- 24. E. Ruel-Gariepy et al., In situ-forming hydrogels review of temperature-sensitive systems. Eur. J. Pharm. Biopharm. 58 (2004)409-426. G. Dumortier et al., A review of Poloxamer 407 pharmaceutical and pharmacological characteristics, Pharm. Res. 23 (2006) 2709-2728.
- 25. R. Rowe et al., Pharmaceutical handbook of pharmaceutical excipients, 5<sup>th</sup>edn., Pharmaceutical, London UK and American Pharmaceutical Association, Washington, USA, 2005.
- 26. I. R. Schmolka et al., Artificial skin. I. Preparation and properties of pluronic F-127 gels for treatment of burns, J. Biomed. Mater. Res. 6 (1972) 571-582.
- 27. A. Cabana et al., Study of the gelation process of polyethylene oxide – polypropylene oxide polyethylene oxide copolymer (Poloxamer 407) aqueous solutions, J. Colloid Interface Sci. 190 (1997) 307-312.
- 28. F. Pasquanelli, Diagnostica e tecniche laboratorio, Rosini Editrice (1981).
- 29. A. W. Burgess et al., Two forms of murine epidermal growth factor: rapid separation by using reverse-

- phase HPLC, Proc. Natl. Acad. Sci. USA 79 (1982) 5753-5757.
- 30. Y. Liu et al., Controlled delivery of recombinant hirudin based on thermo-sensitive Pluronic® F127 hydrogel for subcutaneous administration: in vitro and in vivo characterization, J. Control. Release 117 (2007) 387-395. 8. I. R. Schmolka et al., Artificial skin. I. Preparation and properties of pluronic F-127 gels for treatment of burns, J. Biomed. Mater. Res. 6 (1972) 571-582.
- 31. Pierre S, Liew S, Bernardin A. Basics of dermal filler rheology. Dermatol Surg. 2015 Apr;41 Suppl 1:S120-6.
- 32. Moradi Watson J.Current Α, Concepts in Filler Injection. Facial Plast Surg Clin North Am. 2015 Nov;23(4):489-94.
- 33. Bass LS.Injectable Filler Techniques for Facial Rejuvenation, Volumization, and Augmentation. Facial Plast Surg Clin North Am. 2015 Nov; 23(4):479-88.
- 34. Rhee do Y, Won CH, Chang SE, Noh TK, Kim MS, Kim BJ, Park GH, An JS, Lee MW, Choi JH, Moon KC, Lim SH.Efficacy and safety of a new monophasic hyaluronic acid filler in the correction of nasolabial folds: a randomized, evaluator-blinded, split-face study. J Dermatolog Treat. 2014 Oct; 25(5):448-52.
- 35. San Miguel Moragas J, Reddy RR, Hernández Alfaro F, Mommaerts MY.Systematic review of "filling" procedures for lip augmentation regarding types of material, outcomes and complications. J Craniomaxillofac Surg. 2015 Jul;43(6):883-906.
- 36. Few J, Cox SE, Paradkar-Mitragotri D, Murphy DK.A Multicenter, Single-Blind Randomized, Controlled Study of a Volumizing Hyaluronic Acid Filler for Midface Volume Deficit: Patient-Reported Outcomes at 2 Years. Aesthet Surg J. 2015 Jul; 35(5):589-99.
- 37. Belmontesi M, Grover R, Verpaele A.Transdermal injection of Restylane SubQ for aesthetic contouring of the cheeks, chin, and mandible. Aesthet Surg J. 2006 Jan-Feb;26(1S):S28-34.
- 38. Ho D. Jagdeo J.Biological properties of a new volumizing hyaluronic acid filler: a systematic review.J Drugs Dermatol. 2015 Jan;14(1):50-4. Review.
- 39. Lee JH, Choi YS, Kim SM, Kim YJ, Rhie JW, Jun YJ.Efficacy and safety of porcine collagen filler for nasolabial fold correction in Asians: a prospective multicenter, 12 months follow-up study. J Korean Med Sci. 2014 Nov; 29 Suppl 3:S217-21.
- 40. Yutskovskaya Y, Kogan E, Leshunov randomized, split-face, histomorphologic study comparing a volumetric calcium hydroxylapatite and a hyaluronic acid-based dermal filler. J Drugs Dermatol. 2014 Sep; 13(9):1047-52.
- 41. Emer J, Sundaram H.Aesthetic applications of hydroxylapatite volumizing filler: calcium

- evidence-based review and discussion of current concepts: (part 1 of 2).J DrugsDermatol. 2013 Dec:12(12):1345-54. Review.
- 42. Galadari H, van Abel D, Al Nuami K, Al Faresi F, Galadari I.A randomized, prospective, blinded, splitsingle-center study comparing polycaprolactone to hyaluronic acid for treatment of nasolabial folds.J Cosmet Dermatol. Mar; 14(1):27-32.
- 43. Lee YB, Song EJ, Kim SS, Kim JW, Yu DS.Safety and efficacy of a novel injectable filler in the treatment nasolabial of folds: polymethylmethacrylate and cross-linked dextran in hydroxypropyl methylcellulose. J Cosmet Laser Ther. 2014 Aug; 16(4):185-90.
- 44. Paliwal S, Fagien S, Sun X, Holt T, Kim T, Hee CK, Van Epps D, Messina DJ.Skin extracellular matrix stimulation following injection of a hyaluronic acid-based dermal filler in a rat model. PlastReconstr Surg. 2014 Dec; 134(6):1224-33.
- 45. Lorenc ZP, Bass LM, Fitzgerald R, Goldberg DJ, Graivier MH. Physiochemical Characteristics of Calcium Hydroxylapatite (CaHA). Aesthet Surg J. 2018 Apr 6; 38(suppl 1):S8-S12.
- 46. Casabona G.Blood Aspiration Test for Cosmetic Fillers to Prevent Accidental Intravascular Injection in the Face. Dermatol Surg. 2015 Jul; 41(7):841-7.
- 47. Tansatit T, Moon HJ, Apinuntrum P, Phetudom T. Verification of Embolic Channel Causing Blindness Following Filler Injection. Aesthetic Plast Surg. 2015 Feb: 39(1):154-61.
- 48. Park KH, Kim YK, Woo SJ, Kang SW, Lee WK, Choi KS, Kwak HW, Yoon IH, Huh K, Kim JW; latrogenic occlusion of the ophthalmic artery after cosmetic facial filler injections: a national survey by the Korean Retina Society. Korean Retina Society. JAMA Ophthalmol. 2014 Jun; 132(6):714-23.
- 49. Callan P, Goodman GJ, Carlisle I, et al. Efficacy and safety of a hyaluronic acid filler in subjects treated for correction of midface volume deficiency: a 24 month study. Clin CosmetInvestig Dermatol. 2013; 6:81-89.
- 50. DeLorenzi C, Weinberg M, Solish N, Swift A. The long-term efficacy and safety of a subcutaneously injected large-particle stabilized hyaluronic acidbased gel of nonanimal origin in esthetic facial contouring. Dermatol Surg. 2009; 35(Suppl 1): 313-321.
- 51. Kim JH, Ahn DK, Jeong HS, Suh IS. Treatment algorithm of complications after filler injection: based on wound healing process. J Korean Med Sci. 2014 Nov; 29 Suppl 3:S176-82.
- 52. Carpenter G. Cohen S. Human epidermal growth factor and the proliferation of human fibro- blasts. J. Cell. Physiol. 1976; 88:227-237.
- 53. Carpenter G, Cohen S. Epidermal growth factor. Annu. Rev. Biochem. 1979; 48:193-216.

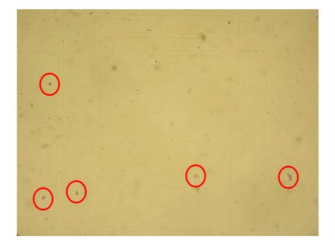
54. Cooper DM, Yu EZ, Hennessey P, Ko F, Robson MC. Determination of endogenous cytokines in chronic wounds. Ann. Surg. 1994; 219:688-691.

Table 1: DSC data of micellization transition of poloxamer preparation

Poloxamer concentration (% w/v)	Micellization onset (K)			Micellization peak (K)			ΔH mic (-J/g)		
	Sample 1	Sample 2	Mean ± D.S.	Sample 1	Sample 2	Mean ± D.S.	Sample 1	Sample 2	Mean ± D.S.
15	287.02	286.22	286.62 ± 0.57	289.68	288.65	289.17 ± 0.73	3.58	3.79	3.68 ± 0.15

Table 2: DSC data of gelation transition of various poloxamer preparations

Poloxamer concentration (% w/v)	Gelation onset (K)		Gelation peak (K)			ΔH gel (-J/g)			
	Sample 1	Sample 2	Mean ± D.S.	Sample 1		Mean ± D.S.	Sample 1	120	Mean ± D.S.
15	295.95	-	295,95	296.36	-	296,36	0.00015998	-	0.00016



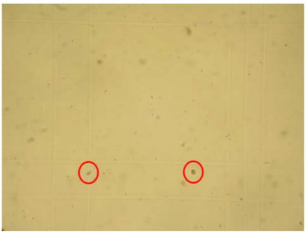
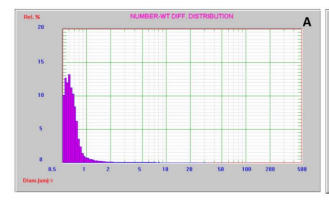


Figure 1: A schematic representation of Burker's chamber



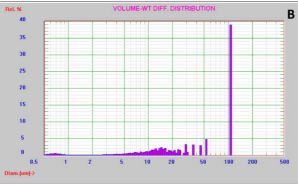


Figure 2: Optical Particle sizer.

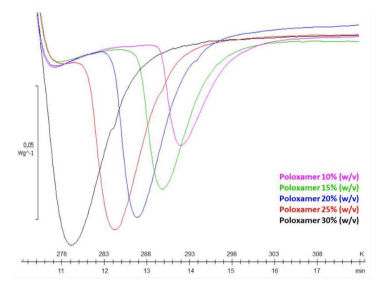


Figure 3: DSC thermograms of different poloxamer concentration samples. Micellization temperature (represented by the enthalpicpeak) decreases with increasing poloxamer concentration

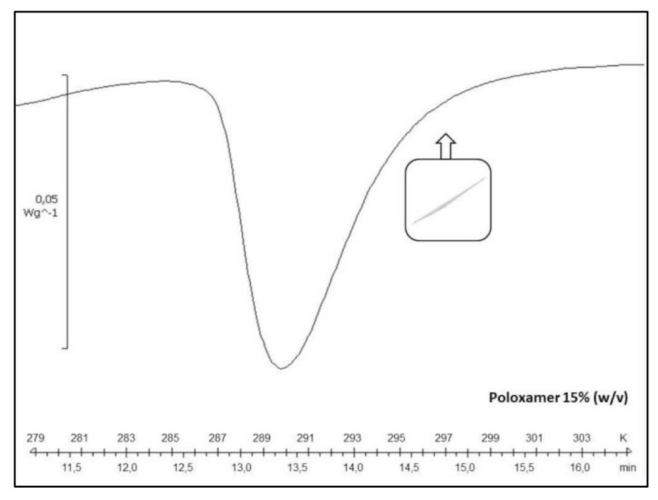


Figure 4: DSC thermogram of poloxamer concentration sample. In the pane there is an enlargement of gelation transition

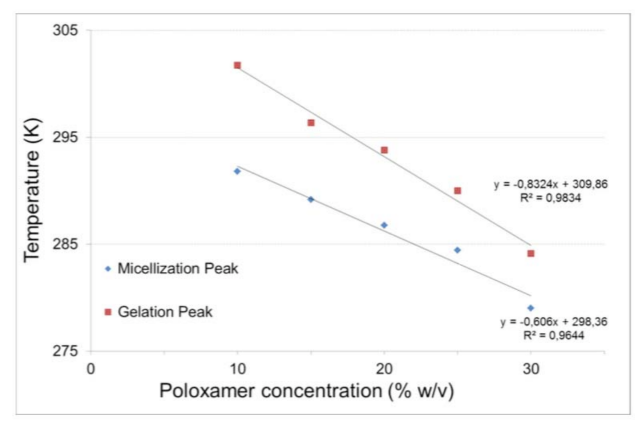


Figure 5: Micellization and gelation temperatures at different poloxamer concentrations

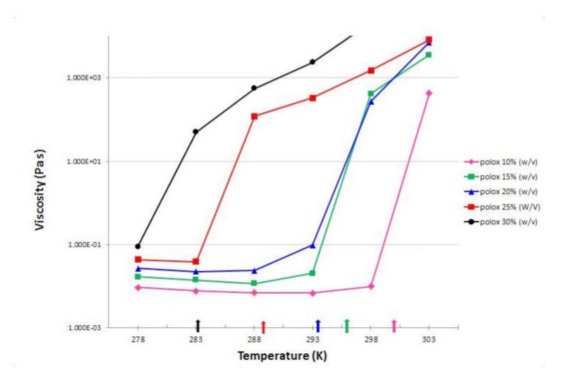


Figure 6: Trend of poloxamer formulation viscosity as a function of temperature. The arrows in the x-axis indicate the gel temperatures obtained from DSC measurements

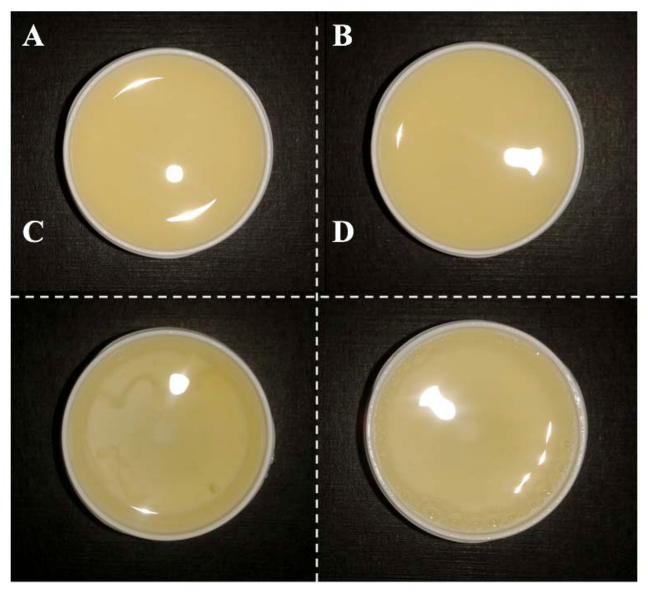


Figure 7: Thermosensitive gel containing PRP fresh prepared (A) and after 7 days from preparation (B). Formulation after 1 month of storage at ~ 4 °C before (C) and after (D) manual shaking

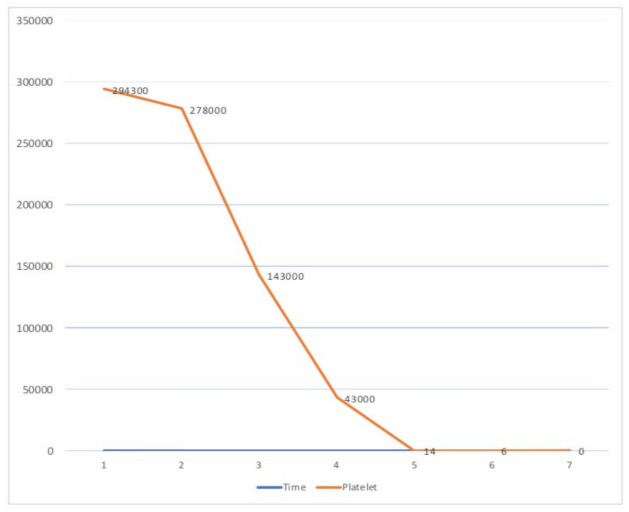


Figure 8: Platelet behaviour on preparation of poloxamer containing PRP

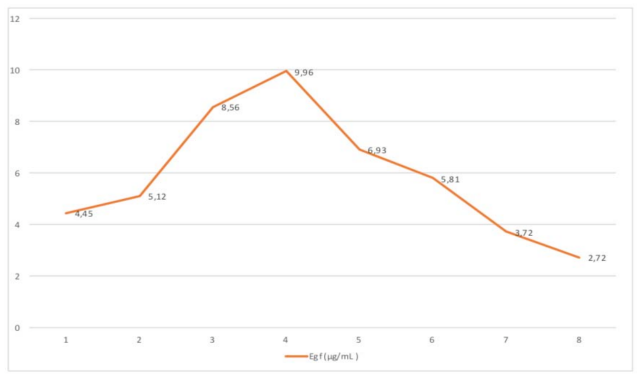


Figure 9: Egf behaviour on preparation of poloxamer containing PRP

# This page is intentionally left blank



# Global Journal of Medical Research: K Interdisciplinary

Volume 20 Issue 13 Version 1.0 Year 2020

Type: Double Blind Peer Reviewed International Research Journal

Publisher: Global Journals

Online ISSN: 2249-4618 & Print ISSN: 0975-5888

# Deep Learning for Classification of Sleep EEG Data during the Epidemic of Coronavirus Disease

By Mingzhe E, Jinming Cao & Bin Zhao

Hubei University of Technology

Abstract- Sleep is an important part of the body's recuperation and energy accumulation, and the quality of sleep also has a significant impact on people's physical and mental state during the epidemic of Coronavirus Disease. It has attracted increasing attention on how to improve the quality of sleep and reduce the impact of sleep-related diseases on health during the Epidemic of Coronavirus Disease.

The electroencephalogram (EEG) signals collected during sleep belong to spontaneous EEG signals. Spontaneous sleep EEG signals can reflect the body's changes, which is also an basis for diagnosis and treatment of related diseases.

Therefore, the establishment of an effective model for classifying sleep EEG signals is an important auxiliary tool for evaluating sleep quality, diagnosing and treating sleep-related diseases.

Keywords: Sleep EEG; deep learning; softmax function; adam algorithm; multiple classifications problem.

GJMR-K Classification: NLMC Code: WC 532



Strictly as per the compliance and regulations of:



© 2020. Mingzhe E, Jinming Cao & Bin Zhao. This is a research/review paper, distributed under the terms of the Creative Commons Attribution-Noncommercial 3.0 Unported License http://creativecommons.org/licenses/by-nc/3.0/), permitting all noncommercial use, distribution, and reproduction in any medium, provided the original work is properly cited.

# Global Journal of Medical Research (K) Volume XX Issue XIII Version I 😢 Year 2020

# Deep Learning for Classification of Sleep EEG Data during the Epidemic of Coronavirus Disease

Mingzhe E α, Jinming Cao α & Bin Zhao ρ

sleep and REM.

Abstract- Sleep is an important part of the body's recuperation and energy accumulation, and the quality of sleep also has a significant impact on people's physical and mental state during the epidemic of Coronavirus Disease. It has attracted increasing attention on how to improve the quality of sleep and reduce the impact of sleep-related diseases on health during the Epidemic of Coronavirus Disease.

The electroencephalogram (EEG) signals collected sleep belong to spontaneous EEG signals. Spontaneous sleep EEG signals can reflect the body's changes, which is also an basis for diagnosis and treatment of related diseases.

Therefore, the establishment of an effective model for classifying sleep EEG signals is an important auxiliary tool for evaluating sleep quality, diagnosing and treating sleep-related diseases.

In this paper, outliers of each kind of original data were detected and deleted by using the principle of 3 Sigma and k-means clustering + Euclidean distance detection method. Then, using the Adam algorithm with adaptive learning rate constructs the Softmax multi-classification BP neural network the model, and relatively high accuracy and AUC values were finally obtained during the Epidemic of Coronavirus Disease.

Keywords: Sleep EEG; deep learning; softmax function; adam algorithm; multiple classifications problem.

### I. Introduction

he sleep process is a complex process of dynamic changes. According to R&K, the international standard for the interpretation of sleep stages, there are different states during sleep.

In addition to the awake period, the sleep cycle consists of two alternate sleep states, namely rapid eye movement(REM), and non-REM.

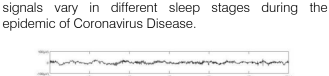
In non-REM, according to the gradual change of sleep state from shallow to deep, it is further divided into sleep, I, II, III, and sleep IV. Sleep stage III and sleep stage IV can be combined with a deep sleep stage.

Figure 1: The sequence of sleep EEG signals in sleep stages. Automatic staging based on EEG signals can

reduce the artificial burden on expert physicians and it is a useful auxiliary tool for assessing sleep quality, diagnosing and treating sleep-related diseases. In this paper, Python is used to build a neural network, and design a sleep staging prediction model. Based on as few training samples as possible, it can obtain relatively high prediction accuracy.

# Overview of BP Neural Network

An artificial neural network gets widely used in some aspects, including pattern recognition, function approximation, data compression, data classification, data prediction, etc. [1-6] BP neural the network is an

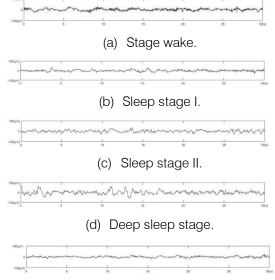


corresponding to different sleep stages, from top to

bottom, namely, wakefulness, sleep I, sleep II, deep

Figure 1 shows the time series of EEG signals

Figure 1 shows the characteristics of EEG



(e) Rapid eye movement.

Author α ρ: School of Science, Hubei University of Technology, Wuhan, Hubei. China.

Author σ: School of Information and Mathematics, Yangtze University, Jingzhou, Hubei, China.

Corresponding Author p: School of Science, Hubei University of Technology, Wuhan, Hubei, China. e-mail: zhaobin@hbut.edu.cn

algorithm in ANN. Figure 2 shows the basic structure of the BP neural network.

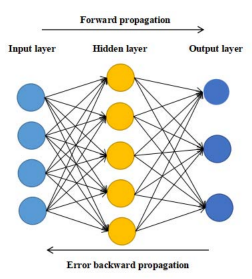


Figure 2: Basic structure diagram of BP neural network. Introduction to activation function and algorithm:

ReLU = max(0, x)

Softmax function:

ReLU function:

$$S_i = \frac{e^i}{\sum_j e^j}$$

Figure 3 shows the operating principle.

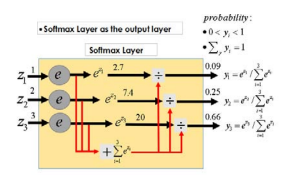


Figure 3: Principle of Softmax function.

Adam algorithm:

 $m_0 v_0 t \leftarrow 0$  Initialize 1st, 2st moment vedor and time step do while:

$$t \leftarrow t + 1$$
$$g_t \leftarrow \nabla_{\theta} f(\theta_{t-1})$$

Computing the gradient.

$$m_t \leftarrow \beta_1 \cdot m_{t-1} + (1 - \beta_1) \cdot g_t$$

Update biased first moment estimate.

$$v_t \leftarrow \beta_2 \cdot v_{t-1} + (1 - \beta_2) \cdot g_t^2$$

Upgrade biased second moment estimate.

$$\hat{m}_{t} \leftarrow \frac{m_{t}}{\left(1 - \left(\beta_{1}\right)^{t}\right)}$$

Compute bias-corrected first moment estimate.

$$\hat{v}_t \leftarrow \frac{v_t}{\left(1 - (\beta_2)^t\right)}$$

Compute bias-corrected second draw moment estimate.

$$\theta_t \leftarrow \theta_{t-1} - \alpha \cdot \frac{\hat{m}_t}{\sqrt{\hat{v}_t} + \varepsilon}$$

Upgrade parameters.

Where  $\alpha$  is the step length,  $\beta_1$ ,  $\beta_2 \in [0,1)$  is the moment estimation of the exponential decay rate, and  $f(\theta)$  is the random objective function of the parameter  $\theta$ .

In this paper, the whole training process of the improved BP neural network model is:

Step 1: Parameter initialization. Determine the node number of the network input layer, hidden layer and output layer, and initialize the weight, bias between each layer, then initialize learning rate.

Step 2: Calculate the output of the hidden layer. The hidden layer output is calculated by the weight and bias between the input vector and the connection layer and the ReLU activation function.

Step 3: Calculate the output of the output layer. through the hidden layer output and connection weights and bias and the Softmax activation function calculate the predicted output.

Step 4: Calculate Softmax cross-entropy as cost function according to predicted output and real label.

Step 5: Back propagation, and this paper use the adaptive learning rate Adam algorithm [7] to update the weight and bias.

Step 6: Determine whether the cost reaches the error range or the number of iterations. If not, return step 2.

# III. Data Description and Preprocessing

Data were collected from 3000 sleep EEG samples and their labels are taken from different healthy adults during overnight sleep. The first is a "known label," which represents the different sleep stages in digital form: stage wake (6), rapid eye movement (5), sleep I (4), sleep II (3), and deep sleep (2); The second to fifth columns are the characteristic parameters calculated from the original time sequence, successively including "Alpha", "Beta", "Theta" and "Delta", which correspond to the energy proportion of EEG signals in the frequency range of "8-13Hz", "14-25Hz", "4-7Hz" and "0.5-4Hz" respectively. The unit of characteristic parameters is the percentage.

This paper gives raw data stage wake (6), and REM. (5), sleep I (4), sleep II (3), deep sleep (2), four brain electrical signal energy proportion of five sleep stages of brain electrical signal energy proportion, but the original data are generally given there are some abnormal data outliers or missing value, therefore we to each index of the five sets of data make a boxplot graph, the result is as follows in figure 4.

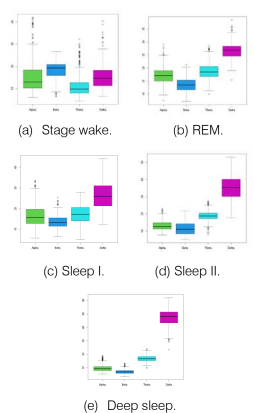


Figure 4: Boxplot of each sleeping period.

Five sleep period by Figure 4 shows, there are some outliers, namely, these all belong to the original data of abnormal points, this paper uses the principle of 3 sigmas [8] will each table of data deletion, then after the processing of five tables to merge, and then using the K-means clustering + Euclidean distance outlier test [9], to find and remove outliers, as shown in figure 5, a total of 2883 samples after pretreatment.

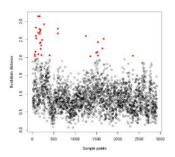


Figure 5: Diagram of outliers.

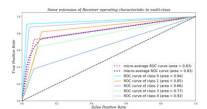
# Model Training and Prediction

We divided the data into a training set and test set in a ratio of 2:8. We trained and tested the data using the traditional decision tree model [10] (DT) and support vector machine model (SVM), and compared the classification effect with the accuracy rate and AUC value as evaluation indexes. The results are as follows:

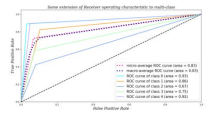
Table 1: Comparison table of several classifications accuracy rates.

Classifier	Accuracy rate
DT	0.59
SVM	0.68
Adam-BPNNet	0.73

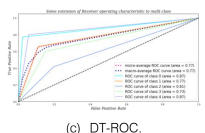
As can be seen from Table 1, the accuracy of Adam-BPNNet in several traditional methods is relatively high. Figure 6 shows the ROC curve of each classification method.

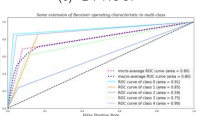


(a) Two-layer BPNNet-ROC.



(b) Three-layer BPNNet-ROC.





(d) SVM-ROC.

Figure 6: ROC curve of each classification method.

Table 2: Comparison table of AUC values of different classification models.

Classifier	AUC
DT	0.77
SVM	0.80
Adam-Bennett	0.83

Table 2 shows that in the Adam-BPNNet model, fewer training sets will still have a better classification effect.

The prediction result is the best classification effect obtained after many experiments. In the early stage of experiment, the classification accuracy is low. After repeated debugging of the number of hidden layers and nodes, the best AUC value of this experiment is 0.83.

# Conclusion

This study is mainly based on theoretical research and combines theory with practice. This paper uses BP neural network based on an adaptive learning rate Adam algorithm for data classification. Also, this paper selects Softmax as the activation function in the output layer, enabling the model to have good selflearning and self-adaptive ability. The most important thing is that the network has good generalization ability. When designing the classifier, it should consider whether the network can correctly classify the objects it needs to classify, and whether the network can correctly classify the unseen or noise-polluted patterns after training. The classification AUC value of this study is 0.83, which is scientific to a certain extent and can be used as auxiliary tool for the evaluation of sleep quality, diagnosis and treatment of sleep-related diseases.

# Conflict of Interest

We have no conflict of interests to disclose and the manuscript has been read and approved by all named authors.

# ACKNOWLEDGEMENT

This work was supported by the Philosophical and Social Sciences Research Project of Hubei Education Department (19Y049), and the Staring Research Foundation for the Ph.D. of Hubei University of Technology (BSQD2019054), Hubei Province, China.

# References Références Referencias

- 1. Yun Qi. Experimental Study on NDVI Inversion using GPS-R Remote Sensing Based on BP Neural Network[D]. Xuzhou: China University of Mining and Technology, 2018.
- Jiao Wang. Study of Data Acquistion System for Electric Stair-Climbing Wheelchair Seat Position Regulating Mechanism[D]. Tianjin: Hebei University of Technology, 2016.
- Yongfeng Cao, Yanjun Zhao. Research on Computer Intelligent Image Recognition echnology based on GA-BP Neural Network[J]. Applied Laser, 2017, 37(1): 139-143.
- Faezeh Rasi Marzabadi. Mehran Masdari. Mohammad Reza Soltani. Application of Artificial Neural Network in Aerodynamic Coefficient Prediction of Subducted Airfoil[J]. Journal of Research in Science and Engineering, 2020, 2(1):
- Xiaomin Wang, Rong Chen, Bin Qiao. Application of BP Neural Network in Tea Disease Classification and Recognition[J]. Guizhou Science 2020, 38(4): 93-96.
- Reza Behmanesh, Iman Rahimi. The Optimized 6. Regression Neural Network Combined with Experimental Design and Regression for Control Chart Prediction[J]. 2020, 2(1): 8-12.
- Kingma DP, Ba J. Adam: A Method for Stochastic Optimization[C]. 3rd International Conference for Learning Representations. San Diego, 2015.
- Lin Lou. Design and Implementation of Anomaly Detection System of Web User Behaviors[D]. Zhejiang University, 2018.
- Hua Jiang, Feng Ji, Huijiao Wang, Xin Wang, Yidi Luo. Improved K-means Algorithm for Ocean Data Anomaly Detection[J]. Computer Engineering and Design, 2018, 39(10): 3132-3136.
- 10. Hang Li. Statistical Learning Methods (in Chinese). Beijing: Tsinghua University Press, 2012.



# Global Journal of Medical Research: K Interdisciplinary

Volume 20 Issue 13 Version 1.0 Year 2020

Type: Double Blind Peer Reviewed International Research Journal

Publisher: Global Journals

Online ISSN: 2249-4618 & Print ISSN: 0975-5888

# Hidradenitis Suppurativa- The Imaging Spectrum

By Vasundhara Singh, Chitrangada Singh & Sharmila Patil

Norfolk and Norwich University

Abstract- This was a retrospective study including 10 cases of histologically proven Hidradenitis suppurativa to compare the Histological features with MRI findings and eventually derive a clinical classification including both parameters.

Keywords: hidradenitis, MRI, follicular hyperkeratosis, sinus tract.

GJMR-K Classification: NLMC Code: WR 430



Strictly as per the compliance and regulations of:



© 2020. Vasundhara Singh, Chitrangada Singh & Sharmila Patil. This is a research/review paper, distributed under the terms of the Creative Commons Attribution-Noncommercial 3.0 Unported License http://creativecommons.org/licenses/by-nc/3.0/), permitting all non-commercial use, distribution, and reproduction in any medium, provided the original work is properly cited.

# Hidradenitis Suppurativa- The Imaging Spectrum

Vasundhara Singh °, Chitrangada Singh ° & Sharmila Patil P

Abstract- This was a retrospective study including 10 cases of histologically proven Hidradenitis suppurativa to compare the Histological features with MRI findings and eventually derive a clinical classification including both parameters.

hidradenitis, MRI, follicular hyperkeratosis, Keywords: sinus tract.

### I. Introduction

Iso known as Acne inversa or the Verneuil's disease, Hidradenitis suppurtiva is a chronic disease with recurrent abscess formation progressing to sinus tracts and resultant scarring. It was first described by Velpeau in 1839during his study involving the origin of abscess involving the sebaceous follicles in axillae [1]

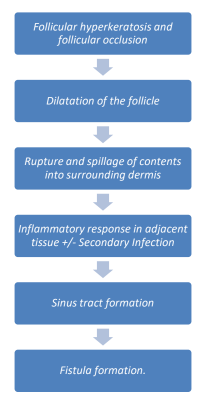
It is commonly seen in females and also incidence involving the axilla in both sexes is nearly equal.

Commonly seen in adolescents, it initially presents like a comedo and progresses with mucopurulent discharge eventually leading to scarring.

### EPIDEMIOLOGY H.

- Prevalence: 1 in 300 adults
- Sex predilection: Females > Males
- Onset: Adolescent to middle age
- Family history: Positive family history with an autosomal-dominant mode of inheritance
- Associations: Crohn's disease. Dowling Dego's. Arthropathy (SAPHO), Smoking, Obesity, Hormonal influence [2]

### III. PATHOPHYSIOLOGY



Author α ρ: MBBS, MD Dermatology and venerology, Department of Dermatology and Venereal Medicine, Dr.D.Y.Patil Hospital and Research Centre, Navi Mumbai (India).

Corresponding Author o: MBBS, MD Radiology, Senior clinical Fellow, Department of Breast Imaging, Norfolk and Norwich University Hospital, UK. e-mail: chitrangada.singh@gmail.com

Clinical Classification: Hurley stages-3 well delineated stages have been described by Hurley emphasizing the clinical diagnosis.

Stage 1: Solitary or multiple isolated abscesses. No scarring or sinus tracts are seen-resembles an acne.

Stage 2: Recurrent abscess may be single or multiple widely separated lesions. Sinus tract may be present. Patient has significant movement restriction.

Stage 3: Diffuse or broad involvement across a regional area with multiple inter-connected sinus tracts and abscesses. Significant scarring is seen and fistula formation is present.

# IV. IMAGING

Imaging evaluation is initially indicated to evaluate the extent and the feasibility to obtain image guided aspirate for culture and sensitivity. A simple ultrasound guided aspiration of the involved part can be used to obtain sample for culture and sensitivity if secondary infections are suspected when there is no sinus tract to directly obtain swabs. [3]

MRI is the preferred modality for evaluating the extent of disease as well as follow up.

# V. MRI SPECTRUM

The protocol

The MRI of the involved area is best suited to evaluate the extent of disease and also may help in monitoring treatment response. The MRI findings parallel with the clinical features therefore avoiding any clinical confusion. STIR (Short tau inversion recovery) and T2W sequences are recommended protocol.

It may initially just show thickening of the skin and subcutaneous tissue which soon progresses to induration best seen on routine T2W and STIR images as subdermal hyperintense signal extending upto skin. In a few days there is formation of subcutaneous abscesses however the disease is confined to the skin and subcutaneous tissue. MRI will show loculated T2W and STIR hyperintense pockets of collection with mild post contrast rim enhancement. This correlates to Stage 1 of clinical classification. (Refer figure A)

This may either heal by mild scarring or progress towards chronic skin involvement in the form of multiple raised subdermal pockets of pus which ultimately rupture to form sinus tract. (Refer figure B) This correlates to Stage 2 of clinical classification.

The stage 3 of clinical classification is includes extensive local involvement in terms of area as well as severity and often includes refractory cases with multiple interconnecting sinus tracts. The chronicity can be identified by thick walls of the sinus tract appearing hypointense on STIR due to scarring.(Refer figure C)

Rarely there is fistulous communication with bladder urethra or rectum etc. in patients with highly

virulent infection or compromised immune status like diabetes.

Reactive inguinal lymph nodes are also seen in conjunction.

Post treatment cases demonstrate residual scarring as STIR hypointense tracts. (Refer figure D)

# VI. DIFFERENTIAL DIAGNOSIS

Carbuncles, Lymphadenitis, Infected Bartholin's cyst, sebaceous cysts, Cellulitis/ erysipelas, Lumbosacral epidural abscess are few of the common differential diagnosis which can be easily ruled out combining the imaging and the clinical picture. [4]

# VII. SEQUELAE

Various sequel can be strictures, disfiguring edema, arthropathy and chronic cases may undergo metaplasia and even lead to squamous cell carcinoma.

# VIII. Treatment

Treatment of HS is directed according to disease severity. Aim is to alleviate symptoms and improve quality of life. Many a times combination therapy is resorted.

- a) Anti-inflammatory Agent [5]
- Intralesional steroids: Triamcinolone acetonide 2-5mg/ml can be used for few lesions
- Anakinra: An IL-1 inhibitor (100mg SC/day for 12 weeks) showed reduction in severity of disease
- Antibiotics: Many topical and oral antibiotics like clindamycin(1%;300mg BD), tetracycline, rifampicin(300mg BD); have been used alone or in combination for their anti-inflammatory and immunomodulatory properties.[6] A study of hyperbaric oxygen therapy with antibiotic combination showed good improvement in sartorius and DLQI score.[7]

Antiandrogens: Although anecdotal in females, a double blinded study in women with Cyproterone acetate(100mg) and Ethinyl estradiol(50 micrograms) as per reversed sequential therapy laid down by Hammerstein and Cupceancu, showed reduced discharge and swelling. [8]

Finasteride (5-10 mg/day) used primarily for prostate cancer showed good results in pediatric patients. [9]

# b) Retinoids

Isotretinoin worked in patients with mild disease when given in low doses 0.5-1.2 mg/kg/day over 4 to 12 months. [10]

A study of Acitretin (0.6mg/kg/day) over 6 to 12 in 12 patients with moderate to severe disease showed improvement. [11]

The mechanism of action is through the keratolytic action thereby reducing ductal occlusions.

# c) Immunosuppresive therapy

Cyclosporine (4.5mg/kg/day) showed raid relief in resistant cases to antibiotics and UVB therapy. [12]

 $TNF\alpha$  inhibitors have been proven to be quite effective in Hurley's II and III stages of the disease.

Infliximab 5mg/kg IV at week 0, 2 and 6 were given to 33 patients; drug was well tolerated and showed good improvement in symptoms and severity of the disease. [13] Etanercept showed varied results when administered twice weekly 50mg for 12 weeks.

Adalimumab when given weekly instead of fortnightly showed superior results when 4 randomized control trials were analysed. [14,15]

Apremilast a selective phosphodiesterase 4 inhibitor (30 mg BD); used primarily for psoriasis; showed moderate results. [16]

# d) Miscellaneous

Botulinum toxin reduces acetylcholine release and in turn reduces the sympathetic activation of apocrine glands. A dose of 40 to 50 Units per session for 3 to 4 times over 3 years reported remission in 4 cases. [17,18]

Metformin helps in decreasing androgen sensitivity by lowering circulating insulin and helps in managing the metabolic syndrome associated with disease. [19]

Others: Zinc, Cryotherapy and Photodynamic therapy

# e) Surgical Intervention

This is the last resort to unresponsive cases. Deroofing is most effective in combination with antibiotics and anti-inflammatory. [20]

# Laser therapy

Nd: YAG laser in 22 patients showed significant improvement in all (65%), axilla (62%), Inguinal (53%), Inframammary (51%). [21]

Carbon dioxide laser for lesions to heal by secondary intention has also been tried. [22]

HURLEY STAGE	TREATMENT
I	Topical antibiotics (clindamycin 1%, Benzoyl Peroxide gel 2.5/5%); Intralesional steroids; Oral Antibiotics: Tetracyclines; Retinoids; Nd YAG; cryotherapy; Botox
II	TNFα inhibitors Antibiotics Nd YAG Surgical/ CO2 deroofing
III	Surgical with Oral antibiotics

# References Références Referencias

- suppurativa: Wiseman MC. Hidradenitis review. Dermatol Ther 2004; 17:50 -54.
- Slade DE, Powell BW, Mortimer PS. Hidradenitis suppurativa: pathogenesis and management. Br J Plast Surg 2003; 56:451 -461.
- Kelly AM, Cronin P. MRI features of hidradenitis suppurativa and review of the literature. AJR Am J Roentgenol. 2005; 185 (5): 1201-4.
- Church JM, Fazio VW, Lavery IC, Oakley JR, Milsom JW. The differential diagnosis and comorbidity of hidradenitis suppurativa and perianal Crohn's disease.
- Tchero H, Herlin C, Bekara F, Fluieraru S et al. Hidradenitis suppurativa: A systematic review and meta-analysis of therapeautic interventions. Indian J Dermatol Venerol Leprol 2019; 85:248-57.
- Clemmensen OJ. Topical treatment of hidradenitis suppurativa with clindamycin. Int J Dermatol 1983; 22: 325-8.
- 7. Yildiz H, Senol L, Ercan E, Bilgili ME, Karabudak Abuaf O. A prospective randomized controlled trial assessing the efficacy of adjunctive hyperbaric

- oxygen therapy in treatment of hidradenitis suppurativa. Int J Dermatol 2016; 55:232-7.
- 8. Sawers RS, Randall VA, Ebling FJ. Control of Hidradenitis suppurativa in women with combined antiandrogen and oestrogen therapy. Br J Dermatol 1986; 115: 269-74.
- Randhawa HK, Hamilton J, Pope E. Finasteride for the treatment of hidradenitis suppurativa in children and adolescents. JAMA Dermatology 149(6):732-5.
- 10. Boer J, van Gemert MJ. Long-term results of isotretinoin in the treatment of 68 patients with hidradenitis suppurativa. J Am Acad Dermatol 1999; 40:73-6.
- 11. Boer J, Nazari M. Long term results of acitretin therapy for Hidradenitis suppurativa. Is Acne-inversa a misnomer? Br J Dermatol 2011; 164(1):170-5.
- 12. Buckley DA, Rogers S. Cyclosporin-responsive hidradenitis suppurativa. J R Soc Med 1995; 88:289P-90P.
- 13. Adams DR, Yankura JA, Fogelberg AC, Anderson BE. Treatment of hidradenitis suppurativa with etarnacept injection. Arch Dermatol 2010;146:501-4.
- 14. Gottlieb A, Menter A, Armstrong A, Ocampo C, Gu Y, Teixeira HD. Adalimumab treatment in women

- with moderate to severe hidradenitis suupurativa from the placebo-controlled portion of a phase 2, randomized, doble-blind study. J Drugs Dermatol 2016;15:1192-6.
- 15. Kimball AB, Okun MM, Williams DA, Gottlieb AB, Papp KA, Zouboulis CC, et al. Two phase 3 trials of adalimumab for hidradenitis ssupurativa. N Eng J Med 2016; 375:422-34.
- 16. Weber P, Seyed Jafari SM, Yawalkar N, Hunger RE. Apremilast in the treatment of moderate to severe hidradenitis suppurativa: A case series of 9 patients. J Am Acad Dermatol 2017; 76:76:1189-91.
- 17. Fieto-Rodriguez M, Sendagorta-Cudos E, Herranz-Pinto P, de Lucas-Laguna R. Prepubertal hidradenitis suppurativa successfully treated with botulinum toxin A. Dermatol Surg 2009;35: 1300-2.
- 18. Khoo AB, Burova EP. Hidradenitis suppurativa treated with clostridium botulinum toxin A. Clin Exp Dermatol 2014;39:749-50.

- 19. Arun B, Loffeld A. Longstanding Hidradenitis suppurativa treated effectively with metformin. Clinical and Exp Dermatology 2009; 34(8):920-1.
- 20. van der Zee HH, Prens EP, Boer J. Deroofing: A tissue saving surgical technique for the treatment of mild to moderate hidradenitis suppurativa lesions. J Am Acad Dermatol 2010;63:475-80.
- 21. Tierney E, Mahmoud BH, Hexsel C, Ozog D, Hamzavi I. Radomized control trial for the treatment of hidradenitis suppurativa with Nd YAG laser. Dermatol Surg 2009; 35:1188-98.
- 22. Finley EM, Ratz JL. Treatment of hidradenitis suppurativa withcarbon dioxide laser excision and second intention healing. J Am Acad Dermatol. 1996; 34:465-9.





Figure A: Hurley Stage 1 MRI Axial STIR image of the perineum shows multiple isolated abscesses within the left natal cleft.

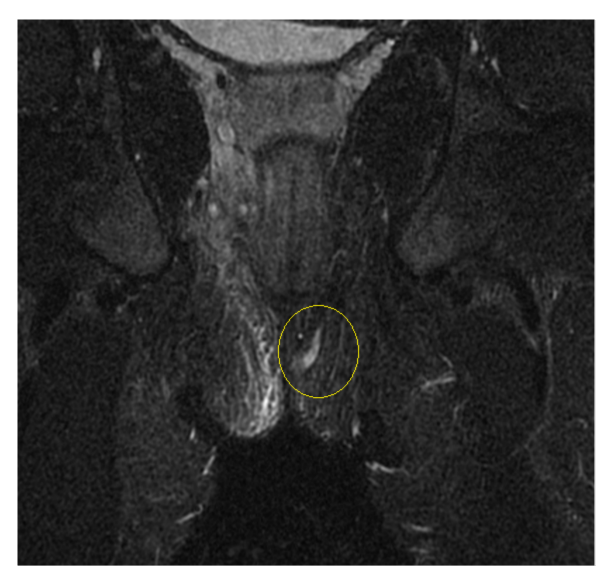


Figure B: Hurley Stage 2 MRI of the perineum: Coronal STIR image shows evolution of the abscess into a linear hyperintense tract/ sinus formation.

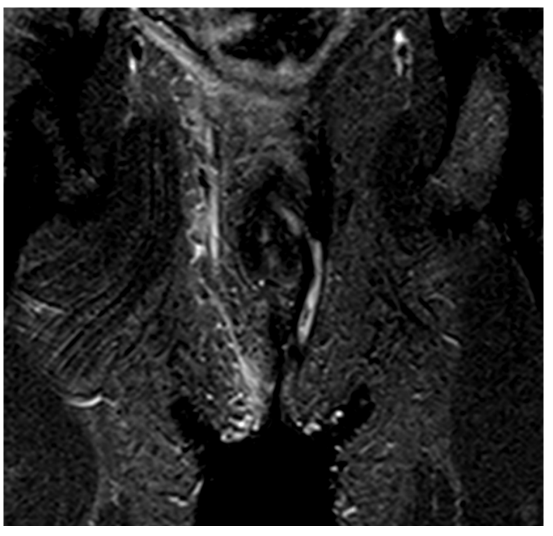


Figure C: Hurley Stage 3.

MRI of the perineum: Coronal STIR image from the selected slices shows evolution of the abscess into a linear hyperintense tract/ sinus formation. Due to chronicity of the sinus, the tract appears more fibrosed and hypointense.

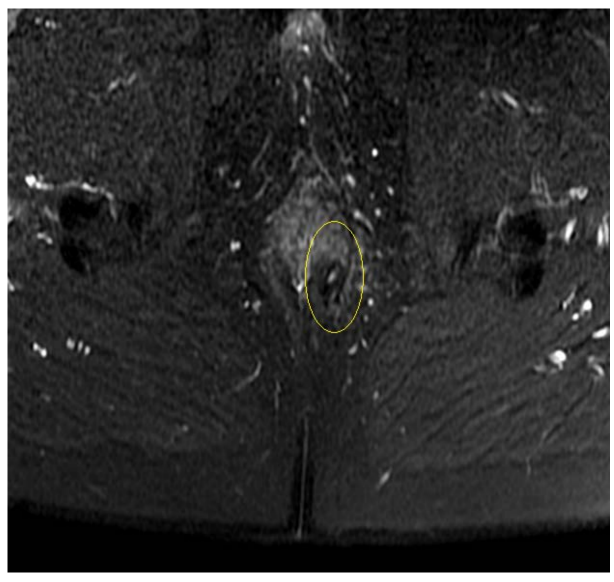


Figure D: Resolution post treatment

MRI of the perineum: Axial STIR image shows residual minimal inflammation with majority of fibrosis as hypointense scar.

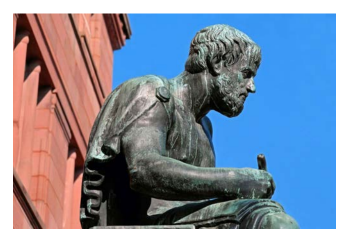
# Global Journals Guidelines Handbook 2020

www.GlobalJournals.org

# **MEMBERSHIPS**

# FELLOWS/ASSOCIATES OF MEDICAL RESEARCH COUNCIL

# FMRC/AMRC MEMBERSHIPS



# INTRODUCTION

FMRC/AMRC is the most prestigious membership of Global Journals accredited by Open Association of Research Society, U.S.A (OARS). The credentials of Fellow and Associate designations signify that the researcher has gained the knowledge of the fundamental and high-level concepts, and is a subject matter expert, proficient in an expertise course covering the professional code of conduct, and follows recognized standards of practice. The credentials are designated only to the researchers, scientists, and professionals that have been selected by a rigorous process by our Editorial Board and Management Board.

Associates of FMRC/AMRC are scientists and researchers from around the world are working on projects/researches that have huge potentials. Members support Global Journals' mission to advance technology for humanity and the profession.

# **FMRC**

# FELLOW OF MEDICAL RESEARCH COUNCIL

FELLOW OF MEDICAL RESEARCH COUNCIL is the most prestigious membership of Global Journals. It is an award and membership granted to individuals that the Open Association of Research Society judges to have made a 'substantial contribution to the improvement of computer science, technology, and electronics engineering.

The primary objective is to recognize the leaders in research and scientific fields of the current era with a global perspective and to create a channel between them and other researchers for better exposure and knowledge sharing. Members are most eminent scientists, engineers, and technologists from all across the world. Fellows are elected for life through a peer review process on the basis of excellence in the respective domain. There is no limit on the number of new nominations made in any year. Each year, the Open Association of Research Society elect up to 12 new Fellow Members.



# BENEFIT

# TO THE INSTITUTION

# GET LETTER OF APPRECIATION

Global Journals sends a letter of appreciation of author to the Dean or CEO of the University or Company of which author is a part, signed by editor in chief or chief author.



# **EXCLUSIVE NETWORK**

# GET ACCESS TO A CLOSED NETWORK

A FMRC member gets access to a closed network of Tier 1 researchers and scientists with direct communication channel through our website. Fellows can reach out to other members or researchers directly. They should also be open to reaching out by other.

Career

Credibility

Exclusive

Reputation



# **CERTIFICATE**

# CERTIFICATE, LOR AND LASER-MOMENTO

Fellows receive a printed copy of a certificate signed by our Chief Author that may be used for academic purposes and a personal recommendation letter to the dean of member's university.

Career

Credibility

Exclusive

Reputation



# **DESIGNATION**

# GET HONORED TITLE OF MEMBERSHIP

Fellows can use the honored title of membership. The "FMRC" is an honored title which is accorded to a person's name viz. Dr. John E. Hall, Ph.D., FMRC or William Walldroff, M.S., FMRC.

Career

Credibility

Exclusive

Reputation

# RECOGNITION ON THE PLATFORM

# BETTER VISIBILITY AND CITATION

All the Fellow members of FMRC get a badge of "Leading Member of Global Journals" on the Research Community that distinguishes them from others. Additionally, the profile is also partially maintained by our team for better visibility and citation. All fellows get a dedicated page on the website with their biography.

Career

Credibility

Reputation



© Copyright by Global Journals | Guidelines Handbook

# **FUTURE WORK**

# GET DISCOUNTS ON THE FUTURE PUBLICATIONS

Fellows receive discounts on the future publications with Global Journals up to 60%. Through our recommendation programs, members also receive discounts on publications made with OARS affiliated organizations.

Career

Financial



# GJ Internal Account

Unlimited forward of Emails

Fellows get secure and fast GJ work emails with unlimited storage of emails that they may use them as their primary email. For example, john [AT] globaljournals [DOT] org.

Career

Credibility

Reputation



# PREMIUM TOOLS

# ACCESS TO ALL THE PREMIUM TOOLS

To take future researches to the zenith, fellows receive access to all the premium tools that Global Journals have to offer along with the partnership with some of the best marketing leading tools out there.

Financial

# **CONFERENCES & EVENTS**

# ORGANIZE SEMINAR/CONFERENCE

Fellows are authorized to organize symposium/seminar/conference on behalf of Global Journal Incorporation (USA). They can also participate in the same organized by another institution as representative of Global Journal. In both the cases, it is mandatory for him to discuss with us and obtain our consent. Additionally, they get free research conferences (and others) alerts.

Career

Credibility

Financial

# EARLY INVITATIONS

### EARLY INVITATIONS TO ALL THE SYMPOSIUMS, SEMINARS, CONFERENCES

All fellows receive the early invitations to all the symposiums, seminars, conferences and webinars hosted by Global Journals in their subject.

Exclusive

© Copyright by Global Journals | Guidelines Handbook





# PUBLISHING ARTICLES & BOOKS

# EARN 60% OF SALES PROCEEDS

Fellows can publish articles (limited) without any fees. Also, they can earn up to 70% of sales proceeds from the sale of reference/review books/literature/publishing of research paper. The FMRC member can decide its price and we can help in making the right decision.

Exclusive

Financial

# REVIEWERS

# GET A REMUNERATION OF 15% OF AUTHOR FEES

Fellow members are eligible to join as a paid peer reviewer at Global Journals Incorporation (USA) and can get a remuneration of 15% of author fees, taken from the author of a respective paper.

Financial

# ACCESS TO EDITORIAL BOARD

# BECOME A MEMBER OF THE EDITORIAL BOARD

Fellows and Associates may join as a member of the Editorial Board of Global Journals Incorporation (USA) after successful completion of three years as Fellow and as Peer Reviewer.

Career

Credibility

Exclusive

Reputation

# AND MUCH MORE

# GET ACCESS TO SCIENTIFIC MUSEUMS AND OBSERVATORIES ACROSS THE GLOBE

All members get access to 5 selected scientific museums and observatories across the globe. All researches published with Global Journals will be kept under deep archival facilities across regions for future protections and disaster recovery. They get 10 GB free secure cloud access for storing research files.



# **AMRC**

# ASSOCIATE OF MEDICAL RESEARCH COUNCIL

ASSOCIATE OF MEDICAL RESEARCH COUNCIL is the membership of Global Journals awarded to individuals that the Open Association of Research Society judges to have made a 'substantial contribution to the improvement of computer science, technology, and electronics engineering.

The primary objective is to recognize the leaders in research and scientific fields of the current era with a global perspective and to create a channel between them and other researchers for better exposure and knowledge sharing. Members are most eminent scientists, engineers, and technologists from all across the world. Associate membership can later be promoted to Fellow Membership. Associates are elected for life through a peer review process on the basis of excellence in the respective domain. There is no limit on the number of new nominations made in any year. Each year, the Open Association of Research Society elect up to 12 new Associate Members.



# BENEFIT

# TO THE INSTITUTION

# GET LETTER OF APPRECIATION

Global Journals sends a letter of appreciation of author to the Dean or CEO of the University or Company of which author is a part, signed by editor in chief or chief author.



# **EXCLUSIVE NETWORK**

# GET ACCESS TO A CLOSED NETWORK

A AMRC member gets access to a closed network of Tier 2 researchers and scientists with direct communication channel through our website. Associates can reach out to other members or researchers directly. They should also be open to reaching out by other.

Career

Credibility

Exclusive

Reputation



# CERTIFICATE

# CERTIFICATE, LOR AND LASER-MOMENTO

Associates receive a printed copy of a certificate signed by our Chief Author that may be used for academic purposes and a personal recommendation letter to the dean of member's university.

Career

Credibility

Exclusive

Reputation



# DESIGNATION

# GET HONORED TITLE OF MEMBERSHIP

Associates can use the honored title of membership. The "AMRC" is an honored title which is accorded to a person's name viz. Dr. John E. Hall, Ph.D., AMRC or William Walldroff, M.S., AMRC.

Career

Credibility

Exclusive

Reputation

# RECOGNITION ON THE PLATFORM

# BETTER VISIBILITY AND CITATION

All the Associate members of AMRC get a badge of "Leading Member of Global Journals" on the Research Community that distinguishes them from others. Additionally, the profile is also partially maintained by our team for better visibility and citation.

Career

Credibility

Reputation



# **FUTURE WORK**

# GET DISCOUNTS ON THE FUTURE PUBLICATIONS

Associates receive discounts on future publications with Global Journals up to 30%. Through our recommendation programs, members also receive discounts on publications made with OARS affiliated organizations.

Career

Financial



# GJ ACCOUNT

# Unlimited forward of Emails

Associates get secure and fast GJ work emails with 5GB forward of emails that they may use them as their primary email. For example, john [AT] globaljournals [DOT] org.

Career

Credibility

Reputation



# PREMIUM TOOLS

# ACCESS TO ALL THE PREMIUM TOOLS

To take future researches to the zenith, fellows receive access to almost all the premium tools that Global Journals have to offer along with the partnership with some of the best marketing leading tools out there.

Financial

# **CONFERENCES & EVENTS**

# ORGANIZE SEMINAR/CONFERENCE

Associates are authorized to organize symposium/seminar/conference on behalf of Global Journal Incorporation (USA). They can also participate in the same organized by another institution as representative of Global Journal. In both the cases, it is mandatory for him to discuss with us and obtain our consent. Additionally, they get free research conferences (and others) alerts.

Career

Credibility

Financial

# **EARLY INVITATIONS**

# EARLY INVITATIONS TO ALL THE SYMPOSIUMS, SEMINARS, CONFERENCES

All associates receive the early invitations to all the symposiums, seminars, conferences and webinars hosted by Global Journals in their subject.

Exclusive

© Copyright by Global Journals | Guidelines Handbook





# Publishing Articles & Books

# EARN 60% OF SALES PROCEEDS

Associates can publish articles (limited) without any fees. Also, they can earn up to 30-40% of sales proceeds from the sale of reference/review books/literature/publishing of research paper

Exclusive

Financial

# REVIEWERS

# GET A REMUNERATION OF 15% OF AUTHOR FEES

Associate members are eligible to join as a paid peer reviewer at Global Journals Incorporation (USA) and can get a remuneration of 15% of author fees, taken from the author of a respective paper.

Financial

# AND MUCH MORE

# GET ACCESS TO SCIENTIFIC MUSEUMS AND OBSERVATORIES ACROSS THE GLOBE

All members get access to 2 selected scientific museums and observatories across the globe. All researches published with Global Journals will be kept under deep archival facilities across regions for future protections and disaster recovery. They get 5 GB free secure cloud access for storing research files.



Associate	Fellow	Research Group	BASIC
\$4800 lifetime designation	\$6800 lifetime designation	\$12500.00 organizational	APC per article
Certificate, LoR and Momento 2 discounted publishing/year Gradation of Research 10 research contacts/day 1 GB Cloud Storage GJ Community Access	Certificate, LoR and Momento Unlimited discounted publishing/year Gradation of Research Unlimited research contacts/day 5 GB Cloud Storage Online Presense Assistance GJ Community Access	Certificates, LoRs and Momentos Unlimited free publishing/year Gradation of Research Unlimited research contacts/day Unlimited Cloud Storage Online Presense Assistance GJ Community Access	<b>GJ</b> Community Access

# Preferred Author Guidelines

# We accept the manuscript submissions in any standard (generic) format.

We typeset manuscripts using advanced typesetting tools like Adobe In Design, CorelDraw, TeXnicCenter, and TeXStudio. We usually recommend authors submit their research using any standard format they are comfortable with, and let Global Journals do the rest.

Alternatively, you can download our basic template from https://globaljournals.org/Template

Authors should submit their complete paper/article, including text illustrations, graphics, conclusions, artwork, and tables. Authors who are not able to submit manuscript using the form above can email the manuscript department at submit@globaljournals.org or get in touch with chiefeditor@globaljournals.org if they wish to send the abstract before submission.

# Before and During Submission

Authors must ensure the information provided during the submission of a paper is authentic. Please go through the following checklist before submitting:

- 1. Authors must go through the complete author guideline and understand and *agree to Global Journals' ethics and code of conduct,* along with author responsibilities.
- 2. Authors must accept the privacy policy, terms, and conditions of Global Journals.
- 3. Ensure corresponding author's email address and postal address are accurate and reachable.
- 4. Manuscript to be submitted must include keywords, an abstract, a paper title, co-author(s') names and details (email address, name, phone number, and institution), figures and illustrations in vector format including appropriate captions, tables, including titles and footnotes, a conclusion, results, acknowledgments and references.
- 5. Authors should submit paper in a ZIP archive if any supplementary files are required along with the paper.
- 6. Proper permissions must be acquired for the use of any copyrighted material.
- 7. Manuscript submitted *must not have been submitted or published elsewhere* and all authors must be aware of the submission.

### **Declaration of Conflicts of Interest**

It is required for authors to declare all financial, institutional, and personal relationships with other individuals and organizations that could influence (bias) their research.

# Policy on Plagiarism

Plagiarism is not acceptable in Global Journals submissions at all.

Plagiarized content will not be considered for publication. We reserve the right to inform authors' institutions about plagiarism detected either before or after publication. If plagiarism is identified, we will follow COPE guidelines:

Authors are solely responsible for all the plagiarism that is found. The author must not fabricate, falsify or plagiarize existing research data. The following, if copied, will be considered plagiarism:

- Words (language)
- Ideas
- Findings
- Writings
- Diagrams
- Graphs
- Illustrations
- Lectures



© Copyright by Global Journals | Guidelines Handbook

- Printed material
- Graphic representations
- Computer programs
- Electronic material
- Any other original work

# AUTHORSHIP POLICIES

Global Journals follows the definition of authorship set up by the Open Association of Research Society, USA. According to its guidelines, authorship criteria must be based on:

- Substantial contributions to the conception and acquisition of data, analysis, and interpretation of findings.
- Drafting the paper and revising it critically regarding important academic content.
- 3. Final approval of the version of the paper to be published.

# **Changes in Authorship**

The corresponding author should mention the name and complete details of all co-authors during submission and in manuscript. We support addition, rearrangement, manipulation, and deletions in authors list till the early view publication of the journal. We expect that corresponding author will notify all co-authors of submission. We follow COPE guidelines for changes in authorship.

# Copyright

During submission of the manuscript, the author is confirming an exclusive license agreement with Global Journals which gives Global Journals the authority to reproduce, reuse, and republish authors' research. We also believe in flexible copyright terms where copyright may remain with authors/employers/institutions as well. Contact your editor after acceptance to choose your copyright policy. You may follow this form for copyright transfers.

# **Appealing Decisions**

Unless specified in the notification, the Editorial Board's decision on publication of the paper is final and cannot be appealed before making the major change in the manuscript.

# **Acknowledgments**

Contributors to the research other than authors credited should be mentioned in Acknowledgments. The source of funding for the research can be included. Suppliers of resources may be mentioned along with their addresses.

### **Declaration of funding sources**

Global Journals is in partnership with various universities, laboratories, and other institutions worldwide in the research domain. Authors are requested to disclose their source of funding during every stage of their research, such as making analysis, performing laboratory operations, computing data, and using institutional resources, from writing an article to its submission. This will also help authors to get reimbursements by requesting an open access publication letter from Global Journals and submitting to the respective funding source.

# Preparing your Manuscript

Authors can submit papers and articles in an acceptable file format: MS Word (doc, docx), LaTeX (.tex, .zip or .rar including all of your files), Adobe PDF (.pdf), rich text format (.rtf), simple text document (.txt), Open Document Text (.odt), and Apple Pages (.pages). Our professional layout editors will format the entire paper according to our official guidelines. This is one of the highlights of publishing with Global Journals—authors should not be concerned about the formatting of their paper. Global Journals accepts articles and manuscripts in every major language, be it Spanish, Chinese, Japanese, Portuguese, Russian, French, German, Dutch, Italian, Greek, or any other national language, but the title, subtitle, and abstract should be in English. This will facilitate indexing and the pre-peer review process.

The following is the official style and template developed for publication of a research paper. Authors are not required to follow this style during the submission of the paper. It is just for reference purposes.



# Manuscript Style Instruction (Optional)

- Microsoft Word Document Setting Instructions.
- Font type of all text should be Swis721 Lt BT.
- Page size: 8.27" x 11'", left margin: 0.65, right margin: 0.65, bottom margin: 0.75.
- Paper title should be in one column of font size 24.
- Author name in font size of 11 in one column.
- Abstract: font size 9 with the word "Abstract" in bold italics.
- Main text: font size 10 with two justified columns.
- Two columns with equal column width of 3.38 and spacing of 0.2.
- First character must be three lines drop-capped.
- The paragraph before spacing of 1 pt and after of 0 pt.
- Line spacing of 1 pt.
- Large images must be in one column.
- The names of first main headings (Heading 1) must be in Roman font, capital letters, and font size of 10.
- The names of second main headings (Heading 2) must not include numbers and must be in italics with a font size of 10.

# Structure and Format of Manuscript

The recommended size of an original research paper is under 15,000 words and review papers under 7,000 words. Research articles should be less than 10,000 words. Research papers are usually longer than review papers. Review papers are reports of significant research (typically less than 7,000 words, including tables, figures, and references)

A research paper must include:

- a) A title which should be relevant to the theme of the paper.
- b) A summary, known as an abstract (less than 150 words), containing the major results and conclusions.
- c) Up to 10 keywords that precisely identify the paper's subject, purpose, and focus.
- d) An introduction, giving fundamental background objectives.
- e) Resources and techniques with sufficient complete experimental details (wherever possible by reference) to permit repetition, sources of information must be given, and numerical methods must be specified by reference.
- Results which should be presented concisely by well-designed tables and figures.
- g) Suitable statistical data should also be given.
- h) All data must have been gathered with attention to numerical detail in the planning stage.

Design has been recognized to be essential to experiments for a considerable time, and the editor has decided that any paper that appears not to have adequate numerical treatments of the data will be returned unrefereed.

- i) Discussion should cover implications and consequences and not just recapitulate the results; conclusions should also be summarized.
- j) There should be brief acknowledgments.
- k) There ought to be references in the conventional format. Global Journals recommends APA format.

Authors should carefully consider the preparation of papers to ensure that they communicate effectively. Papers are much more likely to be accepted if they are carefully designed and laid out, contain few or no errors, are summarizing, and follow instructions. They will also be published with much fewer delays than those that require much technical and editorial correction.

The Editorial Board reserves the right to make literary corrections and suggestions to improve brevity.



# FORMAT STRUCTURE

It is necessary that authors take care in submitting a manuscript that is written in simple language and adheres to published guidelines.

All manuscripts submitted to Global Journals should include:

### Title

The title page must carry an informative title that reflects the content, a running title (less than 45 characters together with spaces), names of the authors and co-authors, and the place(s) where the work was carried out.

### **Author details**

The full postal address of any related author(s) must be specified.

### **Abstract**

The abstract is the foundation of the research paper. It should be clear and concise and must contain the objective of the paper and inferences drawn. It is advised to not include big mathematical equations or complicated jargon.

Many researchers searching for information online will use search engines such as Google, Yahoo or others. By optimizing your paper for search engines, you will amplify the chance of someone finding it. In turn, this will make it more likely to be viewed and cited in further works. Global Journals has compiled these guidelines to facilitate you to maximize the webfriendliness of the most public part of your paper.

# Keywords

A major lynchpin of research work for the writing of research papers is the keyword search, which one will employ to find both library and internet resources. Up to eleven keywords or very brief phrases have to be given to help data retrieval, mining, and indexing.

One must be persistent and creative in using keywords. An effective keyword search requires a strategy: planning of a list of possible keywords and phrases to try.

Choice of the main keywords is the first tool of writing a research paper. Research paper writing is an art. Keyword search should be as strategic as possible.

One should start brainstorming lists of potential keywords before even beginning searching. Think about the most important concepts related to research work. Ask, "What words would a source have to include to be truly valuable in a research paper?" Then consider synonyms for the important words.

It may take the discovery of only one important paper to steer in the right keyword direction because, in most databases, the keywords under which a research paper is abstracted are listed with the paper.

# **Numerical Methods**

Numerical methods used should be transparent and, where appropriate, supported by references.

# **Abbreviations**

Authors must list all the abbreviations used in the paper at the end of the paper or in a separate table before using them.

### Formulas and equations

Authors are advised to submit any mathematical equation using either MathJax, KaTeX, or LaTeX, or in a very high-quality image.

### **Tables, Figures, and Figure Legends**

Tables: Tables should be cautiously designed, uncrowned, and include only essential data. Each must have an Arabic number, e.g., Table 4, a self-explanatory caption, and be on a separate sheet. Authors must submit tables in an editable format and not as images. References to these tables (if any) must be mentioned accurately.



# **Figures**

Figures are supposed to be submitted as separate files. Always include a citation in the text for each figure using Arabic numbers, e.g., Fig. 4. Artwork must be submitted online in vector electronic form or by emailing it.

# Preparation of Eletronic Figures for Publication

Although low-quality images are sufficient for review purposes, print publication requires high-quality images to prevent the final product being blurred or fuzzy. Submit (possibly by e-mail) EPS (line art) or TIFF (halftone/ photographs) files only. MS PowerPoint and Word Graphics are unsuitable for printed pictures. Avoid using pixel-oriented software. Scans (TIFF only) should have a resolution of at least 350 dpi (halftone) or 700 to 1100 dpi (line drawings). Please give the data for figures in black and white or submit a Color Work Agreement form. EPS files must be saved with fonts embedded (and with a TIFF preview, if possible).

For scanned images, the scanning resolution at final image size ought to be as follows to ensure good reproduction: line art: >650 dpi; halftones (including gel photographs): >350 dpi; figures containing both halftone and line images: >650 dpi.

Color charges: Authors are advised to pay the full cost for the reproduction of their color artwork. Hence, please note that if there is color artwork in your manuscript when it is accepted for publication, we would require you to complete and return a Color Work Agreement form before your paper can be published. Also, you can email your editor to remove the color fee after acceptance of the paper.

# TIPS FOR WRITING A GOOD QUALITY MEDICAL RESEARCH PAPER

- 1. Choosing the topic: In most cases, the topic is selected by the interests of the author, but it can also be suggested by the guides. You can have several topics, and then judge which you are most comfortable with. This may be done by asking several questions of yourself, like "Will I be able to carry out a search in this area? Will I find all necessary resources to accomplish the search? Will I be able to find all information in this field area?" If the answer to this type of question is "yes," then you ought to choose that topic. In most cases, you may have to conduct surveys and visit several places. Also, you might have to do a lot of work to find all the rises and falls of the various data on that subject. Sometimes, detailed information plays a vital role, instead of short information. Evaluators are human: The first thing to remember is that evaluators are also human beings. They are not only meant for rejecting a paper. They are here to evaluate your paper. So present your best aspect.
- 2. Think like evaluators: If you are in confusion or getting demotivated because your paper may not be accepted by the evaluators, then think, and try to evaluate your paper like an evaluator. Try to understand what an evaluator wants in your research paper, and you will automatically have your answer. Make blueprints of paper: The outline is the plan or framework that will help you to arrange your thoughts. It will make your paper logical. But remember that all points of your outline must be related to the topic you have chosen.
- **3.** Ask your guides: If you are having any difficulty with your research, then do not hesitate to share your difficulty with your guide (if you have one). They will surely help you out and resolve your doubts. If you can't clarify what exactly you require for your work, then ask your supervisor to help you with an alternative. He or she might also provide you with a list of essential readings.
- **4.** Use of computer is recommended: As you are doing research in the field of medical research then this point is quite obvious. Use right software: Always use good quality software packages. If you are not capable of judging good software, then you can lose the quality of your paper unknowingly. There are various programs available to help you which you can get through the internet.
- 5. Use the internet for help: An excellent start for your paper is using Google. It is a wondrous search engine, where you can have your doubts resolved. You may also read some answers for the frequent question of how to write your research paper or find a model research paper. You can download books from the internet. If you have all the required books, place importance on reading, selecting, and analyzing the specified information. Then sketch out your research paper. Use big pictures: You may use encyclopedias like Wikipedia to get pictures with the best resolution. At Global Journals, you should strictly follow here.



- 6. Bookmarks are useful: When you read any book or magazine, you generally use bookmarks, right? It is a good habit which helps to not lose your continuity. You should always use bookmarks while searching on the internet also, which will make your search easier.
- 7. Revise what you wrote: When you write anything, always read it, summarize it, and then finalize it.
- 8. Make every effort: Make every effort to mention what you are going to write in your paper. That means always have a good start. Try to mention everything in the introduction—what is the need for a particular research paper. Polish your work with good writing skills and always give an evaluator what he wants. Make backups: When you are going to do any important thing like making a research paper, you should always have backup copies of it either on your computer or on paper. This protects you from losing any portion of your important data.
- **9. Produce good diagrams of your own:** Always try to include good charts or diagrams in your paper to improve quality. Using several unnecessary diagrams will degrade the quality of your paper by creating a hodgepodge. So always try to include diagrams which were made by you to improve the readability of your paper. Use of direct quotes: When you do research relevant to literature, history, or current affairs, then use of quotes becomes essential, but if the study is relevant to science, use of quotes is not preferable.
- **10.** Use proper verb tense: Use proper verb tenses in your paper. Use past tense to present those events that have happened. Use present tense to indicate events that are going on. Use future tense to indicate events that will happen in the future. Use of wrong tenses will confuse the evaluator. Avoid sentences that are incomplete.
- 11. Pick a good study spot: Always try to pick a spot for your research which is quiet. Not every spot is good for studying.
- 12. Know what you know: Always try to know what you know by making objectives, otherwise you will be confused and unable to achieve your target.
- **13.** Use good grammar: Always use good grammar and words that will have a positive impact on the evaluator; use of good vocabulary does not mean using tough words which the evaluator has to find in a dictionary. Do not fragment sentences. Eliminate one-word sentences. Do not ever use a big word when a smaller one would suffice.

Verbs have to be in agreement with their subjects. In a research paper, do not start sentences with conjunctions or finish them with prepositions. When writing formally, it is advisable to never split an infinitive because someone will (wrongly) complain. Avoid clichés like a disease. Always shun irritating alliteration. Use language which is simple and straightforward. Put together a neat summary.

- **14. Arrangement of information:** Each section of the main body should start with an opening sentence, and there should be a changeover at the end of the section. Give only valid and powerful arguments for your topic. You may also maintain your arguments with records.
- **15. Never start at the last minute:** Always allow enough time for research work. Leaving everything to the last minute will degrade your paper and spoil your work.
- **16. Multitasking in research is not good:** Doing several things at the same time is a bad habit in the case of research activity. Research is an area where everything has a particular time slot. Divide your research work into parts, and do a particular part in a particular time slot.
- 17. Never copy others' work: Never copy others' work and give it your name because if the evaluator has seen it anywhere, you will be in trouble. Take proper rest and food: No matter how many hours you spend on your research activity, if you are not taking care of your health, then all your efforts will have been in vain. For quality research, take proper rest and food.
- 18. Go to seminars: Attend seminars if the topic is relevant to your research area. Utilize all your resources.
- 19. Refresh your mind after intervals: Try to give your mind a rest by listening to soft music or sleeping in intervals. This will also improve your memory. Acquire colleagues: Always try to acquire colleagues. No matter how sharp you are, if you acquire colleagues, they can give you ideas which will be helpful to your research.



- **20.** Think technically: Always think technically. If anything happens, search for its reasons, benefits, and demerits. Think and then print: When you go to print your paper, check that tables are not split, headings are not detached from their descriptions, and page sequence is maintained.
- 21. Adding unnecessary information: Do not add unnecessary information like "I have used MS Excel to draw graphs." Irrelevant and inappropriate material is superfluous. Foreign terminology and phrases are not apropos. One should never take a broad view. Analogy is like feathers on a snake. Use words properly, regardless of how others use them. Remove quotations. Puns are for kids, not grunt readers. Never oversimplify: When adding material to your research paper, never go for oversimplification; this will definitely irritate the evaluator. Be specific. Never use rhythmic redundancies. Contractions shouldn't be used in a research paper. Comparisons are as terrible as clichés. Give up ampersands, abbreviations, and so on. Remove commas that are not necessary. Parenthetical words should be between brackets or commas. Understatement is always the best way to put forward earth-shaking thoughts. Give a detailed literary review.
- **22. Report concluded results:** Use concluded results. From raw data, filter the results, and then conclude your studies based on measurements and observations taken. An appropriate number of decimal places should be used. Parenthetical remarks are prohibited here. Proofread carefully at the final stage. At the end, give an outline to your arguments. Spot perspectives of further study of the subject. Justify your conclusion at the bottom sufficiently, which will probably include examples.
- **23. Upon conclusion:** Once you have concluded your research, the next most important step is to present your findings. Presentation is extremely important as it is the definite medium though which your research is going to be in print for the rest of the crowd. Care should be taken to categorize your thoughts well and present them in a logical and neat manner. A good quality research paper format is essential because it serves to highlight your research paper and bring to light all necessary aspects of your research.

# INFORMAL GUIDELINES OF RESEARCH PAPER WRITING

### Key points to remember:

- Submit all work in its final form.
- Write your paper in the form which is presented in the guidelines using the template.
- Please note the criteria peer reviewers will use for grading the final paper.

# **Final points:**

One purpose of organizing a research paper is to let people interpret your efforts selectively. The journal requires the following sections, submitted in the order listed, with each section starting on a new page:

The introduction: This will be compiled from reference matter and reflect the design processes or outline of basis that directed you to make a study. As you carry out the process of study, the method and process section will be constructed like that. The results segment will show related statistics in nearly sequential order and direct reviewers to similar intellectual paths throughout the data that you gathered to carry out your study.

# The discussion section:

This will provide understanding of the data and projections as to the implications of the results. The use of good quality references throughout the paper will give the effort trustworthiness by representing an alertness to prior workings.

Writing a research paper is not an easy job, no matter how trouble-free the actual research or concept. Practice, excellent preparation, and controlled record-keeping are the only means to make straightforward progression.

# General style:

Specific editorial column necessities for compliance of a manuscript will always take over from directions in these general guidelines.

To make a paper clear: Adhere to recommended page limits.



### Mistakes to avoid:

- Insertion of a title at the foot of a page with subsequent text on the next page.
- Separating a table, chart, or figure—confine each to a single page.
- Submitting a manuscript with pages out of sequence.
- In every section of your document, use standard writing style, including articles ("a" and "the").
- Keep paying attention to the topic of the paper.
- Use paragraphs to split each significant point (excluding the abstract).
- Align the primary line of each section.
- Present your points in sound order.
- Use present tense to report well-accepted matters.
- Use past tense to describe specific results.
- Do not use familiar wording; don't address the reviewer directly. Don't use slang or superlatives.
- Avoid use of extra pictures—include only those figures essential to presenting results.

# Title page:

Choose a revealing title. It should be short and include the name(s) and address(es) of all authors. It should not have acronyms or abbreviations or exceed two printed lines.

**Abstract:** This summary should be two hundred words or less. It should clearly and briefly explain the key findings reported in the manuscript and must have precise statistics. It should not have acronyms or abbreviations. It should be logical in itself. Do not cite references at this point.

An abstract is a brief, distinct paragraph summary of finished work or work in development. In a minute or less, a reviewer can be taught the foundation behind the study, common approaches to the problem, relevant results, and significant conclusions or new questions.

Write your summary when your paper is completed because how can you write the summary of anything which is not yet written? Wealth of terminology is very essential in abstract. Use comprehensive sentences, and do not sacrifice readability for brevity; you can maintain it succinctly by phrasing sentences so that they provide more than a lone rationale. The author can at this moment go straight to shortening the outcome. Sum up the study with the subsequent elements in any summary. Try to limit the initial two items to no more than one line each.

Reason for writing the article—theory, overall issue, purpose.

- Fundamental goal.
- To-the-point depiction of the research.
- Consequences, including definite statistics—if the consequences are quantitative in nature, account for this; results of any numerical analysis should be reported. Significant conclusions or questions that emerge from the research.

# Approach:

- Single section and succinct.
- An outline of the job done is always written in past tense.
- o Concentrate on shortening results—limit background information to a verdict or two.
- Exact spelling, clarity of sentences and phrases, and appropriate reporting of quantities (proper units, important statistics) are just as significant in an abstract as they are anywhere else.

### Introduction:

The introduction should "introduce" the manuscript. The reviewer should be presented with sufficient background information to be capable of comprehending and calculating the purpose of your study without having to refer to other works. The basis for the study should be offered. Give the most important references, but avoid making a comprehensive appraisal of the topic. Describe the problem visibly. If the problem is not acknowledged in a logical, reasonable way, the reviewer will give no attention to your results. Speak in common terms about techniques used to explain the problem, if needed, but do not present any particulars about the protocols here.



The following approach can create a valuable beginning:

- o Explain the value (significance) of the study.
- o Defend the model—why did you employ this particular system or method? What is its compensation? Remark upon its appropriateness from an abstract point of view as well as pointing out sensible reasons for using it.
- Present a justification. State your particular theory(-ies) or aim(s), and describe the logic that led you to choose them.
- Briefly explain the study's tentative purpose and how it meets the declared objectives.

# Approach:

Use past tense except for when referring to recognized facts. After all, the manuscript will be submitted after the entire job is done. Sort out your thoughts; manufacture one key point for every section. If you make the four points listed above, you will need at least four paragraphs. Present surrounding information only when it is necessary to support a situation. The reviewer does not desire to read everything you know about a topic. Shape the theory specifically—do not take a broad view.

As always, give awareness to spelling, simplicity, and correctness of sentences and phrases.

# Procedures (methods and materials):

This part is supposed to be the easiest to carve if you have good skills. A soundly written procedures segment allows a capable scientist to replicate your results. Present precise information about your supplies. The suppliers and clarity of reagents can be helpful bits of information. Present methods in sequential order, but linked methodologies can be grouped as a segment. Be concise when relating the protocols. Attempt to give the least amount of information that would permit another capable scientist to replicate your outcome, but be cautious that vital information is integrated. The use of subheadings is suggested and ought to be synchronized with the results section.

When a technique is used that has been well-described in another section, mention the specific item describing the way, but draw the basic principle while stating the situation. The purpose is to show all particular resources and broad procedures so that another person may use some or all of the methods in one more study or referee the scientific value of your work. It is not to be a step-by-step report of the whole thing you did, nor is a methods section a set of orders.

# **Materials:**

Materials may be reported in part of a section or else they may be recognized along with your measures.

### Methods:

- Report the method and not the particulars of each process that engaged the same methodology.
- Describe the method entirely.
- o To be succinct, present methods under headings dedicated to specific dealings or groups of measures.
- Simplify—detail how procedures were completed, not how they were performed on a particular day.
- o If well-known procedures were used, account for the procedure by name, possibly with a reference, and that's all.

### Approach:

It is embarrassing to use vigorous voice when documenting methods without using first person, which would focus the reviewer's interest on the researcher rather than the job. As a result, when writing up the methods, most authors use third person passive voice.

Use standard style in this and every other part of the paper—avoid familiar lists, and use full sentences.

# What to keep away from:

- o Resources and methods are not a set of information.
- o Skip all descriptive information and surroundings—save it for the argument.
- o Leave out information that is immaterial to a third party.



© Copyright by Global Journals | Guidelines Handbook

### **Results:**

The principle of a results segment is to present and demonstrate your conclusion. Create this part as entirely objective details of the outcome, and save all understanding for the discussion.

The page length of this segment is set by the sum and types of data to be reported. Use statistics and tables, if suitable, to present consequences most efficiently.

You must clearly differentiate material which would usually be incorporated in a study editorial from any unprocessed data or additional appendix matter that would not be available. In fact, such matters should not be submitted at all except if requested by the instructor.

### **Content:**

- Sum up your conclusions in text and demonstrate them, if suitable, with figures and tables.
- o In the manuscript, explain each of your consequences, and point the reader to remarks that are most appropriate.
- o Present a background, such as by describing the question that was addressed by creation of an exacting study.
- Explain results of control experiments and give remarks that are not accessible in a prescribed figure or table, if appropriate.
- Examine your data, then prepare the analyzed (transformed) data in the form of a figure (graph), table, or manuscript.

# What to stay away from:

- Do not discuss or infer your outcome, report surrounding information, or try to explain anything.
- Do not include raw data or intermediate calculations in a research manuscript.
- o Do not present similar data more than once.
- o A manuscript should complement any figures or tables, not duplicate information.
- Never confuse figures with tables—there is a difference.

# Approach:

As always, use past tense when you submit your results, and put the whole thing in a reasonable order.

Put figures and tables, appropriately numbered, in order at the end of the report.

If you desire, you may place your figures and tables properly within the text of your results section.

# Figures and tables:

If you put figures and tables at the end of some details, make certain that they are visibly distinguished from any attached appendix materials, such as raw facts. Whatever the position, each table must be titled, numbered one after the other, and include a heading. All figures and tables must be divided from the text.

# Discussion:

The discussion is expected to be the trickiest segment to write. A lot of papers submitted to the journal are discarded based on problems with the discussion. There is no rule for how long an argument should be.

Position your understanding of the outcome visibly to lead the reviewer through your conclusions, and then finish the paper with a summing up of the implications of the study. The purpose here is to offer an understanding of your results and support all of your conclusions, using facts from your research and generally accepted information, if suitable. The implication of results should be fully described.

Infer your data in the conversation in suitable depth. This means that when you clarify an observable fact, you must explain mechanisms that may account for the observation. If your results vary from your prospect, make clear why that may have happened. If your results agree, then explain the theory that the proof supported. It is never suitable to just state that the data approved the prospect, and let it drop at that. Make a decision as to whether each premise is supported or discarded or if you cannot make a conclusion with assurance. Do not just dismiss a study or part of a study as "uncertain."



Research papers are not acknowledged if the work is imperfect. Draw what conclusions you can based upon the results that you have, and take care of the study as a finished work.

- o You may propose future guidelines, such as how an experiment might be personalized to accomplish a new idea.
- o Give details of all of your remarks as much as possible, focusing on mechanisms.
- o Make a decision as to whether the tentative design sufficiently addressed the theory and whether or not it was correctly restricted. Try to present substitute explanations if they are sensible alternatives.
- One piece of research will not counter an overall question, so maintain the large picture in mind. Where do you go next? The best studies unlock new avenues of study. What questions remain?
- o Recommendations for detailed papers will offer supplementary suggestions.

# Approach:

When you refer to information, differentiate data generated by your own studies from other available information. Present work done by specific persons (including you) in past tense.

Describe generally acknowledged facts and main beliefs in present tense.

# THE ADMINISTRATION RULES

Administration Rules to Be Strictly Followed before Submitting Your Research Paper to Global Journals Inc.

Please read the following rules and regulations carefully before submitting your research paper to Global Journals Inc. to avoid rejection.

Segment draft and final research paper: You have to strictly follow the template of a research paper, failing which your paper may get rejected. You are expected to write each part of the paper wholly on your own. The peer reviewers need to identify your own perspective of the concepts in your own terms. Please do not extract straight from any other source, and do not rephrase someone else's analysis. Do not allow anyone else to proofread your manuscript.

Written material: You may discuss this with your guides and key sources. Do not copy anyone else's paper, even if this is only imitation, otherwise it will be rejected on the grounds of plagiarism, which is illegal. Various methods to avoid plagiarism are strictly applied by us to every paper, and, if found guilty, you may be blacklisted, which could affect your career adversely. To guard yourself and others from possible illegal use, please do not permit anyone to use or even read your paper and file.



# CRITERION FOR GRADING A RESEARCH PAPER (COMPILATION) BY GLOBAL JOURNALS

Please note that following table is only a Grading of "Paper Compilation" and not on "Performed/Stated Research" whose grading solely depends on Individual Assigned Peer Reviewer and Editorial Board Member. These can be available only on request and after decision of Paper. This report will be the property of Global Journals.

Topics	Grades		
	A-B	C-D	E-F
Abstract	Clear and concise with appropriate content, Correct format. 200 words or below	Unclear summary and no specific data, Incorrect form Above 200 words	No specific data with ambiguous information Above 250 words
Introduction	Containing all background details with clear goal and appropriate details, flow specification, no grammar and spelling mistake, well organized sentence and paragraph, reference cited	Unclear and confusing data, appropriate format, grammar and spelling errors with unorganized matter	Out of place depth and content, hazy format
Methods and Procedures	Clear and to the point with well arranged paragraph, precision and accuracy of facts and figures, well organized subheads	Difficult to comprehend with embarrassed text, too much explanation but completed	Incorrect and unorganized structure with hazy meaning
Result	Well organized, Clear and specific, Correct units with precision, correct data, well structuring of paragraph, no grammar and spelling mistake	Complete and embarrassed text, difficult to comprehend	Irregular format with wrong facts and figures
Discussion	Well organized, meaningful specification, sound conclusion, logical and concise explanation, highly structured paragraph reference cited	Wordy, unclear conclusion, spurious	Conclusion is not cited, unorganized, difficult to comprehend
References	Complete and correct format, well organized	Beside the point, Incomplete	Wrong format and structuring



# INDEX

Therapeutic · 9, 11

V A Aesthetic · 19, 24 Virulent · 36  $\text{Ailments} \cdot 9$ Alleviate  $\cdot$  36 Anthropometric · 7, 8, 9, 11 Aqueous · 19 Aspirate · 36 Autonomic · 8 Incubator · 1, 13 Μ Meditative · 7, 9 P Precedence · 7 Q Quaternary · 20 R Recuperation · 31 Rejuvenation · 9, 22, 23 S Sterilization  $\cdot$  1, 5, 13 Sympathetic  $\cdot$  37 **T** 



# Global Journal of Medical Research

Visit us on the Web at www.GlobalJournals.org | www.MedicalResearchJournal.org or email us at helpdesk@globaljournals.org





122N 9755896