

KING KHALID UNIVERSITY COLLEGE OF DENTISTRY SCIENTIFIC RESEARCH COMMITTEE



OFFICIAL USE ONLY		
REGISTRATION NUMBER: SRC/ETH/20 /	REGISTRATION DATE:	ا لا ــــ ـــ ــــ ـــــــــــــــــــــ
APPLICATION FOR ETHICAL	CLEARANCE: (CLINICAL TR	<u>IALS)</u>
Research Title	Principal Investig	ator:
	Name:	
	Position :	
	Tel no: Email:	
Dessearch Trues		
Research Type	Co-Investigato	rs
Staff research: Internship research:	1	
Student research: Funded research:	3.	
	4.	
Research Details	Participant Check	x List
Research site:	Will informed consent be	Yes No
Multi-site study: Yes: No:	obtained? (attach the copy)	
Proposed start date: Anticipated study duration:	Will questionnaire be used? (attach the copy)	Yes No
Anticipated study duration.	Relationship between	
Participant detail:	investigator and participant:	
Male: Female: Total:	Does the research include	Yes No
	procedures on	
Age range: years to	Under-age subjects?	
	Does the study involve	Yes No
Method of participant selection:	Vulnerable groups? (Mentally/physically challenged)	
	Is there any financial /other	Yes No
Are there any perceived potential risks for	inducement:	
participants/ researchers? Yes: No:	Are any participants or	Yes No
If Yes Explain:	research related work from	
	outside KSA?	
Data Collection & Handling	Action of SRC-0	COD
Form of data collection;		
Identified: Potentially identifiable:		
De-identified:		
Data storage:		
Identified: Potentially identifiable:	L	
Is collected data secured? Yes : No:		
Are the participants identifiable in publication	Signature of Chairperson	-SRC-COD
/output ?: Yes : No:		

1. Mandatory for all investigations involving human participants with Clinical trials.

2. Please submit additional documents if required /asked to confirm the ethical practice in research activity and data handling.



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Fill appropriate sections applicable for your research

Administration Of Substance/ Drugs	Bodily Tissue Collection
Name of substance:	The bodily tissue collection is:
Dosage per administration:	Prospective : Retrospective:
	What tissue/ fluid :
Frequency of administration:	
Total amount of administration:	Frequency and volume
Is your study a non-interventional trial?	
Yes: No:	Method of storing /disposing
Please register with Saudi food and drug agency (MDCI@sfda.gov.sa)	
Is your study a Post-Authorisation safety trial?	
Yes: No:	Who will collect and expertise in doing so:
Please register with Saudi food and drug agency	
(MDCI@sfda.gov.sa)	
Anticipated effects:	The collection of biological material is for :
	Clinical care : Research purpose:
Radioactive Material	Medical Devices
Exposure to Radioactive material: Yes No	This study on medical device is:
Exposure to Radioactive material: Yes No	
Exposure to Radioactive material: Yes No Therapeutic : Diagnostic: Does the trial involve additional radiation	This study on medical device is: Trial investigating : Evaluating :
Exposure to Radioactive material: Yes No Therapeutic : Diagnostic: Does the trial involve additional radiation exposure? Yes: No:	This study on medical device is:
Exposure to Radioactive material: Yes No Therapeutic : Diagnostic: Does the trial involve additional radiation	This study on medical device is: Trial investigating : Evaluating : Name of the Instrument/ Device:
Exposure to Radioactive material: Yes No Therapeutic : Diagnostic: Does the trial involve additional radiation exposure? Yes: No:	This study on medical device is: Trial investigating : Evaluating : Name of the Instrument/ Device: Is the instrument approved for safety standards?
Exposure to Radioactive material: Yes No Therapeutic : Diagnostic: Does the trial involve additional radiation exposure? Yes: No: State the associated risk: Explain about clinical monitoring and safety	This study on medical device is: Trial investigating : Evaluating : Name of the Instrument/ Device:
Exposure to Radioactive material: Yes No Therapeutic : Diagnostic: Does the trial involve additional radiation exposure? Yes: No: State the associated risk:	This study on medical device is: Trial investigating : Evaluating : Name of the Instrument/ Device: Is the instrument approved for safety standards?
Exposure to Radioactive material: Yes No Therapeutic : Diagnostic: Does the trial involve additional radiation exposure? Yes: No: State the associated risk: Explain about clinical monitoring and safety	This study on medical device is: Trial investigating : Evaluating : Name of the Instrument/ Device: Is the instrument approved for safety standards? Provide details: Is the study within the terms of its safety
Exposure to Radioactive material: Yes No Therapeutic : Diagnostic: Does the trial involve additional radiation exposure? Yes: No: State the associated risk: Explain about clinical monitoring and safety	This study on medical device is: Trial investigating : Evaluating : Name of the Instrument/ Device: Is the instrument approved for safety standards? Provide details: Is the study within the terms of its safety standards or to obtain the safety standard
Exposure to Radioactive material: Yes No Therapeutic : Diagnostic: Does the trial involve additional radiation exposure? Yes: No: State the associated risk: Explain about clinical monitoring and safety	This study on medical device is: Trial investigating : Evaluating : Name of the Instrument/ Device: Is the instrument approved for safety standards? Provide details: Is the study within the terms of its safety standard certificate? Provide explanation:
Exposure to Radioactive material: Yes No Therapeutic : Diagnostic: Does the trial involve additional radiation exposure? Yes: No: State the associated risk: Explain about clinical monitoring and safety	This study on medical device is: Trial investigating : Evaluating : Name of the Instrument/ Device: Is the instrument approved for safety standards? Provide details: Is the study within the terms of its safety standards or to obtain the safety standard
Exposure to Radioactive material: Yes No Therapeutic : Diagnostic: Does the trial involve additional radiation exposure? Yes: No: State the associated risk: Explain about clinical monitoring and safety	This study on medical device is: Trial investigating : Evaluating : Name of the Instrument/ Device: Is the instrument approved for safety standards? Provide details: Is the study within the terms of its safety standards or to obtain the safety standard certificate? Provide explanation: Please register with Saudi food and drug agency
Exposure to Radioactive material: Yes No Therapeutic : Does the trial involve additional radiation exposure? Yes: No: State the associated risk: Explain about clinical monitoring and safety measures of the participants:	This study on medical device is: Trial investigating : Evaluating : Name of the Instrument/ Device: Is the instrument approved for safety standards? Provide details: Is the study within the terms of its safety standards or to obtain the safety standard certificate? Provide explanation: Please register with Saudi food and drug agency
Exposure to Radioactive material: Yes No Therapeutic : Does the trial involve additional radiation exposure? Yes: No: State the associated risk: Explain about clinical monitoring and safety measures of the participants: Qualification and competence of Principal	This study on medical device is: Trial investigating : Evaluating : Name of the Instrument/ Device: Is the instrument approved for safety standards? Provide details: Is the study within the terms of its safety standards or to obtain the safety standard certificate? Provide explanation: Please register with Saudi food and drug agency

I declare that the above project has been developed and will be conducted in accordance with relevant SRC-COD standards, policies, and codes of practice including any standard or special condition for ongoing ethics clearance. In case, there will be any changes in the procedures for obtaining the subjects, or if there will be some physical or emotional harm to the subjects, I shall report these to the SRC-COD.

Signature of Principal Investigator(Including on behalf of all Co-Investigators)

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