# Institutional Review Board (IRB)

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Department of Family and Community Medicine
College of Medicine, King Saud University

King Saud University

October 2019



# Learning Objectives



At the end of this session, you will be able to:

- ✓ Identify what is IRB
- ✓ Understand the structure of IRB
- **✓ Complete KSU IRB forms**

# What is IRB?

• An Institutional Review Board (IRB) is an independent body established to protect the rights and welfare of human research participants.

# What is IRB?

 Any clinical investigation must be reviewed and approved by an IRB

 No clinical study may begin enrolling participants until it has received IRB approval

# Main Purpose of IRB

The purpose of an IRB is to <u>safeguard the rights</u>, <u>safety</u>, <u>and well-being of all human research</u> <u>participants</u>.

HOW??

# Main Purpose of IRB

- Reviewing the full study plan for a research study to ensure that the research meets pre-defined criteria.
- Confirming that the research plans do not expose participants to unreasonable risks
- Reviewing and approving proposed payments or other compensation to study participants.
- Ensuring that human participant protections remain in force throughout the research by conducting continuing review of approved research. This continuing review is conducted at intervals appropriate to the degree of risk posed by each study, but not less frequently than once a year.

# Main Purpose of IRB

- Considering adverse events, interim findings, and any recent literature that may be relevant to the research
- Assessing suspected protocol violations, complaints expressed by research participants, or violations of institutional policies
- Reviewing proposed changes to previously approved studies

# The IRB has the authority to:

- Approve, disapprove, or terminate all research activities
- Require modifications in protocols, including protocols of previously approved research.
- Require that participants be given any additional information that will assist them in making an informed decision to take part in research.
- Require documentation of informed consent or allow a waiver of documentation.

# **KSU IRB Forms**



# KSU IRB Meeting Schedule (2019 - 2020)

Kingdom of Saudi Arabia King Saudi University (034) p.o. Box 7805 Riyadh 11472 Tel: +966 11 467 00 11 Fax: +96611 467 19 92

http://medicalcity.ksu.edu.sa

المملكة العربية السعودية جامعة الملك سعود (٣٤) ص.ب ٨٠٥ الرياض ١٤٧٢ هاتف: ١١ . ٢٦٩ الـ٢١٠ فاكس: ١٩٩٢ ٢٩٢ الـ٢٩+



المدينة الطبية الجامعية

## Institutional Review Board Schedule of Meetings Academic Year 1440 – 1441

Meeting	Gregoria	n	Hijra		Submission
No.	Month/ Year	Date	Month/Year	Date	Deadline (by 12:00 pm)
1	September 2019	26	Muharram 1441	27	Sept. 05, 2019
2	October 2019	31	Rabi-I 1441	03	Oct. 10, 2018
3	November 2019	28	Rabi-II 1441	01	Nov. 07, 2018
4	December 2019	26	Rabi-II 1441	29	Dec. 05, 2018
5	January 2020	30	Jumaada-II 1441	04	Jan. 09, 2019
6	February 2020	27	Rajab 1441	03	Feb. 06, 2019
7	March 2020	19	Rajab 1441	24	Feb. 27, 2019
8	April 2020	23	Sha'ban 1441	30	April 09, 2019



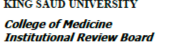
## College of Medicine King Saud University

## Institutional Review Board (IRB)

## Checklists for Submission of a Research Protocol for Review

esea	rch Project Title:
rima	ry Investigator:
ULL	PROTOCOL REVIEW CHECKLIST
	The submitted proposal should be detailed as per IRB requirements.
	$\underline{\mbox{All co-investigators should sign the proposal}}$ (investigator's page) and make sure that their names and titles are correct.
	<u>Informed Consent Form</u> (in accordance with the enclosed Sample Consent Form).
	Signed and dated IRB Declaration of Conflict of Interest (enclosed)
	Signed and dated IRB Certificate of Confidentiality
	Signed and dated CV of Principal Investigator (PI) in IRB CV Template
	For retrospective studies, <u>or</u> department other than the Investigator, <u>approval from the involved Unit</u> . (Approval by the chairman of the department).
	Provide a brief description of the role of each investigator.
	<u>Data Sheet/Questionnaire/Study tools</u> developed by Investigator. If from other author, copyrights/permission to use <u>or</u> confirm 'open access for research & academic purpose).
	For Master student project: Letter from supervisor/institute as evidence of project
	Certificate of Bio-Ethics of each study member completed free online through

### Kingdom of Saudi Arabia KING SAUD UNIVERSITY





المملكة العربية السعونية جامعة الملك سعود كلية الطب مجلس أخلاقيات البحوث الطبية

# نموذج طلب موافقة مجلس أخلاقيات البحوث الطبية رقم البحث:

	عنوان البحث :
	الباحث الرنيس والقسم الذي ينتمي إليه : الباحثون المشاركون والأقسام التي ينتمون إليها :
	البخلول المساركون والرصام التي يتمون إليها :  مدة البحث : شهراً المقترحة :
	·
	Summary: (150 – 200 words)
*	Research Title: (عنوان البحث)
	Research Problem: (مثنكلة البحث وأهميت)
	Research Significance: (أهمية البحث)
	Research Objectives: (أهداف البحث)
-	Research Methodology: (منهجية البحث) Study design; study setting; sample size; sampling technique; data collection methods (e.g. questionnaire; lab investigations; measurements; data analysis plan (including software; techniques, etc)
	This is only one page, not more
	2.178
	توقيع الباحث الرنيس : التاريخ :

ملاحظة: لا يقبل أي ملخص ما لم يقدم مطبوعاً باللغة العربية و موقعاً من الباحث الرنيس.

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College of Medicine Institutional Review Board (IRB)



المملكة العربية السعودية جامعة الملك سعود كلية الطب مجلس أخلاقيات البحوث الطبية

# مقترح مشروع بحثي Research Project Proposal

Please, type either in English

فضلاً، ثنم الطباعة إما باللغة الحربية أو الإنجليزية

or Arabic

<i>التوقيع</i> Signature	الكلية/القسم	الرتبة العلمية	أسماء الباحثين *
Signature	College/Department	Academic Title	*Investigators Names
			-1
			-2
			-3
			-4
			-5
			-6
			-7
			-8
			-9

\* الاسم الأول: الباحث الرئيس.

الاسم الثاني: الباحث المشارك الذي يرشحه الباحث الرئيس ليتولى القيام بالبحث في حال تغيب الباحث الرئيس أو نخليه عن البحث.

#### NOTE:

Please provide contact details where we can contact you in case of incomplete/missing documents.

Office Tel. No.	Department Tel. No.
Mobile No. (optional)	Email:

<sup>\* &</sup>lt;u>First name</u> indicates the Principal Investigator (PI).

<u>Second name</u> is the co-investigator designated by the PI to assume all responsibilities, in case of the absence of the PI.

# مشكلة البحث و أهميته Research Problem and Significance

[Type here]
This document is copyright © (KSU-REC) King Saud University, 2017. No part may be reproduced in any form or by any means or transmitted, or published without prior written consent from King Saud University. Form #KSU-IRB 019-E, Version 2.0. Last updated 02 Oct. 2017.

- Background of the topic
- Rationale
- Significance
- Research question

# أهداف البحث

# Research Objectives

Overall and specific SMART\* objectives, including person, place and time (all objectives)

\* SMART:

Specific

Measurable

Achievable

Relevant

Time-bound

Hypotheses

# Literature Review

Latest relevant literature (international, regional, national, local as available and appropriate) using suitable search engines and methods

This could be 2-3 pages maximum, with suitable referencing

## منهجية البحث Research Methodology

(Please tick box [] if there is standard deviation to this form).

Study Design: (تصميم الدراسة)

(quantitative [observational, experimental]; qualitative, mix, etc)

Study Duration (مدة الدراسة) (When the study will be conducted and its duration)

Study Setting: (اعداد الدراسة) (where will the study be carried out, including secondary data collection places)

Target Population/Sample Size: (المكان المستهدفين / حجم العينة) ( Mention about the target population and narrative of sample size estimation with all assumptions and by using standard equations or soft wares)

Inclusion Criteria: (معايير الاشتمال) (Inclusion criteria for the study subjects, if any)

Exclusion Criteria: (معايير الاستبعاد) (Exclusion criteria for the study subjects, if any).

Study Variables: (متغيرات الدراسة) (All study variables such as: Age, gender,..... and outcome variables of the study:.....)

Ethical Considerations: (الاعتبارات الاخلاقية) consent form to be attached, choose from the following as applicable to study design / objectives):

- The informed consent will be clear and indicates the purpose of the study and the right of the
  participant to withdraw at any time without any obligation towards the study team.
- Participants anonymity will be assured by assigning each participants with a code number for the purpose of analysis only.
- No incentives or rewards will be given to participants. Snack / refreshments may be provided to
  establish a bond with participants, with no obligation to participate.

Data Collection/Data Source: (جمع البيانات المصدر) (e.g. questionnaire [sections / variables mentioned, with copy attached]; biochemical measurements [with references of techniques / kits]; physical measurements [with description of method / reference, as applicable])

Questionnaires/data Sheets from other authors (Copyrights or permission to use, or open access for academic and research purpose)

Statistical Analysis: (حطيل المصانية) Data will be analyzed using SPSS 24.0 version statistical software. Descriptive statistics ( mean, standard deviation, frequencies and percentages) will be used to describe the quantitative and categorical variables. Bivariate statistical analysis will be carried out using appropriate ( Chi-square, student's t test, one-way analysis of variance and Pearson's correlation ) statistical tests, based on the type of study and outcome variables. A p-value of <0.05 and 95% CI will be used to report the statistical significance and precision of results.

# المراجع References

Using standard methods of citing references, e.g. Vancouver style (New England Journal of Medicine) as

دور المحققين								
Investigators Names	Brief Description of the Role of Each Investigator							
	e.g. Review of literature, study design, data collection, Data management and analysis, progress report, data analysis, final report and manuscript writing (as applies to each investigator)							

# الخطة الزمنية للبحث RESEARCH TIME SCHEDULE

starting Date : / / 142 H. ما يخ بداية البحث: / / 142 هـ

البند Items	النوالي المخطط للأعمال الرئيسة		First Year السنة الأولى Second Year					ear	السنة التانية																
Iter	Planned sequence of major tasks	1	2	3	4	5	6	7	8	9	10	11	12	1	2	3	4	5	6	7	8	9	10	11	12
	Literature Review																								
	Questionnaire Design																								
	Data collection																								
	Data management and analysis																								
	Final report writing																								
	Presentation and manuscript writing																								



King Saud University Medical City

Institutional Review Board (IRB)

#### **CONFLICT OF INTEREST**

Projec	ct Title:
Princi	pal Investigator:
	searcher(s) must declare any potential conflict of interest that could affect the outcome of posed research in any form like:  Any financial or other ties of the investigator(s) or a member of his/her family to any
	party directly or indirectly involved in the field of study such as:
	Holding stocks or shares     Receiving educational or other research grant
	3- Employment opportunity
	4- Any gifts of any kind
-	Career development opportunity
-	Speaking arrangements
-	Publication opportunity
-	Providing advisory and consultancy services
-	Board memberships
-	Personal considerations or relationships
-	Promises of any of the above
	If there is any Conflict of Interest, please report:
0	No Conflict of Interest to report.
Ir	nvestigator Name & Signature Date

جامعة الملك سعود King Saud University

وكلّة الجامعة للدارسات العلما والبحث العلمي Vice Rectorate for Graduate Studies & Scientific Research عمادة البحث العلمي Deanship of Scientific Research

Research Ethics Committee لجنة اخلاقيات البحوث

## نموذج موافقة مسبقة للمشاركة في دراسة قائمة على الاستبيان رمز النموذج أ -KSU-REC 006QS جامعة الملك سعود، الرياض، المملكة العربية السعودية

عنوان مشروع البحث: (يرجى إدراج عنوان الدراسة بالكامل)

اسم الباحث الأساسي:

اسم و عنوان الجامعة:

رقم الاتصال:

أعزائى المشاركين،

أود أن انتهاز هذه الفرصة و سؤالك إذا كنت ترغب المشاركة في هذه الدراسة القائمة على الاستبيان. يرجى الإجابة على الأسئلة إلى حد معرفتك. جميع المعلومات المطلوبة في استبيان هذه الدراسة سوف يتم التعامل معها بكل سرية. و اذا كنت ترغب في المشاركة تطوعياً في هذه الدراسة، يرجى التوقيع على هذا النموذج و سوف تحصل على نسخة و تحفظ في السجلات الخاصة بك.

## تم التوقيع بواسطة:

اسم الباحث بالكامل:
اسم الدراسة:
التوقيع:
التاريخ (يوم/شهر/سنة):

- ( ) انا أوافق على المشاركة في هذه الدراسة و على استخدام المعلومات لأغراض البحث العلمي.
- ( ) أوافق على السماح للباحثين بوضع صوت صوتي في المقابلة لأغراض البحث (إذا كان قابل للتطبيق)
- ( ) انا أوافق على السماح للباحثين بالوصول إلى سجلاتي الطبية الموجودة، سواء كانت إلكترونية أو ورقية لدراستهم، وجمع البيانات بأثر مستقبلي، واستخراجها من الممارسات والإجراءات الروتينية .



Research Project Title:

Name of Principal Investigator:

#### King Saud University ركالة الجامعة للدارسات العليا والبحث العلمي Vice Rectorate for Graduate Studies & Scientific Research Deanship of Scientific Research Research Ethics Committee

جامعة الملك سعود لجنة أخلاقيات البحوث

For REC use only:				
Expedited [	]			
Proposal No				

## INFORMED CONSENT FOR QUESTIONNAIRE-BASED STUDY Form # KSU-REC 006QS-E

King Saud University, Riyadh, Kingdom of Saudi Arabia

[please type the complete title of the study]

Name and address of Institution: Contact no:	
Dear Participants,	
Please answer the questions to	ity if you are willing to take part of this questionnaire-based survey. the best of your knowledge. All information asked in this study infidential. If you are willing to participate voluntarily in this study, in a copy for your own records.
Signed by:	
Investigator's Complete Name:	
Study Designation:	
Signature:	
Date (dd/mmm/yyyy):	
purposes.	s study survey, and to utilize the information for scientific research rchers to audiotape my voice as in interview for research purpose, (if



جامعة الملك سعود

## King Saud University

Vice Rectorate for Graduate Studies & Scientific Research وكالة الجامعة للدارسات العليا والبحث العلمى

Deanship of Scientific Research عمادة البحث العلم

Study Delegation Log Form # KSU-IRB 0023 E

## **Delegation of Authority Log**

Investigator's Name:	Protocol Title:	IRB Project No.

List staff to whom the Principal Investigator (PI) has delegated significant study-related duties.

Name	Responsibilities	Initials	Signature	Start Date	End Date	PI Initials/Date

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Form # KSU-REC 0023-E, Version 1.0, Dated 02 Oct. 2017.

By initialing above, I, the PI, declare that during the conduct of the above study, I have delegated the following study-related activities:

Responsibilities Legend				
1. Obtain Consent	7. Laboratory processing	16. Complete Source Documents		
2. Screen Subjects	8. Shipping of samples	17. Complete Case Report Forms		
3. Obtain Medical History	9. Review Safety parameters	18. Provide Discharge Instructions		
4. Perform Physical Exam	10. Randomize Subjects	19. Make Follow-up Phone Calls		
5. Determine Eligibility	11. Dispense Study Drug 20. Others, specify			
6. Obtain study samples	12. Administer IMP			
	13. Administer IDP			
	14. Drug Accountability			

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Form # KSU-REC 0023-E, Version 1.0, Dated 02 Oct. 2017.

Signature of Principal Investigator: Date:				
For more information, please visit the website of Research Ethics Committee in King Saud University (http://dsrs.ksu.edu.sa/ar/comm_Policies)  This document is copyright © (KSU-REC) King Saud University, 2017. No part may be reproduced in any form or by any means ,or transmitted, or published without prior written consent from King Saud University.				



## **Certificate of Confidentiality**

This document certify that the Principal Investigator.
With Study protocol titled
Ensure the confidentiality of research participant's data and biological material obtained either prospectively or from existing record, under strict privacy and security throughout the study duration, publication and at any public presentation.
Principal Investigator will disclose the custodianship of the study material, with (first) and (second) or if applicable, (third) party, who shares the same, or a part, inside or outside the Kingdom of Saudi Arabia, as stated in the submitted protocol. Patient Information Sheet/Informed Consent Form, should have clear description of the information shared.
The privacy must ensure:
<ul> <li>Secured Access to data and bio samples of research subject</li> <li>Security on subject's data and bio samples</li> <li>Secured electronic data access with user password</li> <li>De-identification of research subjects personal information at publication or public presentation</li> <li>Compliance on 'duration' of archiving and storage location specified in protocol, agreed and signed</li> </ul>
The signatory acknowledge that s/he have read and understood this agreement and agree to be bound by its terms.
Signed by:
Principal Investigator
Signature:
Name: Date:



#### **CURRICULUM VITAE**

Name:					
Present appointment: (	lob title, department, and organi	zation)			
Address: (Full work addres	es)				
•					
Telephone number:	KSU Email address:	Personal Email address: (optional)			
Qualifications:					
Professional Registration	on: (Name of body, registration	number and date of registration.)			
Previous and other appointments.)	ointments: (Include previous a	ppointments in the last 5 years and other			
		nary of research experience, including the esearch experience relevant to the current			
.,,					
		ning in the design or conduct of research,			
for example in the Clinical Trials Regulations, Good Clinical Practice, consent or other training appropriate to non-clinical research. Give the date of the training.)					
Relevant Publications: publications relevant to the		ations in the last two years plus other			
Signature:	Date:				

