

Faculty of Health Sciences Ethics Office for Research, Training and Support

APPLICATION PROCESSES FOR ETHICS APPROVAL THROUGH HREC AND ANIMCARE

There are five possible application processes:

1. A first time application for a **single study**/a **larger study** with defined postgraduate student projects/a **single study affiliated to a larger study**
2. A **sub-study application** (student) under an approved larger study
3. A **systematic review**
4. An application for an **amendment** to an approved study
5. **Monitoring report**/or request for **extension** of an approved study

Definitions:

- **Single study**
A study consisting of one or more researchers not intended to involve other potential Masters or Doctoral students. The extent of the data/biological samples is limited in nature.
- **Larger study**
A study planning to involve several researchers, Masters and Doctoral students and that clearly identify the objectives and the methodology to be used by the researchers and the potential students. The extent of the data/biological samples is extensive in nature. This study usually runs over a longer period of time e.g. longitudinal or has an intervention as outcome.
- **Single study affiliated to a larger study**
A planned affiliated study that is *similar in purpose* to an ethically approved larger study and that will contribute to the larger study but will *change the original methodology* described in the larger study proposal to analyze data/biological samples generated in the larger project or it will generate new data/biological samples using a different methodology.
The study was not originally identified as a potential Masters or PhD study in a larger study. This could also be the case when *secondary data analysis* is done on data that were previously published but using different methods of analysis or a combination of data sets/biological samples that will provide meaningful new results.
- **Sub-study**
A sub-study that has been identified as a potential Masters or PhD study in an ethically approved larger study that covers a *specific stated objective* of the larger study and uses *identical methodology* or section/s of the methodology than the larger study. It could be that data/biological samples have already been collected or are going to be collected. **NB The sub-study can add no new methodology that was not covered in the larger study.** If the latter is needed the larger study should be amended first.

- **Amendment**

Any change made to the originally planned proposal and that happens while the study is being conducted. No change may be implemented without the necessary approval by HREC or AnimCare.

- **Monitoring**

Monitoring refers to the process of observing quality and conduct of the research while in progress. *Passive monitoring* refers to the compulsory reporting required by HREC or AnimCare (minimum on an annual basis). *Active monitoring* refers to unannounced monitoring visits conducted by HREC or AnimCare to e.g. the research sites or where data is stored.

- **Extension**

A study is approved for a year. Submission and review of the annual monitoring report provides approval to continue with the study. However, if a researcher requires extension for a study not falling in this process, extension can be requested by submitting a monitoring report to HREC or AnimCare.

1. **A first time application for a single study (including an affiliated study to a larger study) or a larger study with defined postgraduate student projects or a single study affiliated to a larger study**

Process:

Conceptualize the research study (Observing problems, reading literature, discussion etc.)



Develop the research proposal and applicable accompanying documentation and enter into negotiations with potential authorities to ensure that they will be open for the research to be conducted.



Request the necessary application documents either from the:

- AnimCare administration for research involving animals (Ethics-AnimCare@nwu.ac.za) or
- HREC administration for research involving humans (Ethics-HRECAppl@nwu.ac.za).

OR

Download it from the ethics website



Submit the proposal to the *scientific/proposal committee* in your research entity for scientific evaluation and approval.

Obtain a letter of approval by them that has to be attached to the application.



Once the proposal has been approved by the scientific/proposal committee submit the title registration request through the Faculty office (this is a process that runs parallel with the research ethics application).



Submit the completed ethics application either to the:

- AnimCare administration for research involving animals (Ethics-AnimCare@nwu.ac.za) or
- HREC administration for research involving humans (Ethics-HRECAppl@nwu.ac.za)

Attach all the required documents *separately* to the e-mail (see attachment checklist below).

Attach a *covering letter* indicating:

- the title of the research
- the researcher/s
- the type of research ethics application
- which documents are attached to the application, and
- add any explanation to clarify your application



Application sent by administration (four working days) to two to three independent reviewers (5 working days for review).



The application is discussed at the appropriate Research Ethics Committee meeting e.g. research involving animals at the AnimCare committee meeting and research involving humans at the HREC meeting.

- Decision process
 - Aggregate individual views
 - Deliberation (debate)
 - Analogue (consensus)
 - Vote if necessary
- Decision
 - Approved
 - Approved with minimal/several changes
 - Deferred (too many changes and committee deliberation needed)
 - Disapproved (have to go back to the drawing board)



Formal letter of decision of the REC with attached independent reviewer reports are sent to the applicant (always the study leader or PI) as soon as possible (approximately three working days) after the meeting by the appropriate administrator.



Corrections are done by the applicant and are sent back as soon as possible to either:

- the AnimCare administrator for research involving animals (Ethics-AnimCare@nwu.ac.za) or
- the HREC administrator for research involving humans (please note that the corresponding person for HREC now changes to Ethics-HRECProcess@nwu.ac.za if corrections are needed).

A rebuttal letter should be included indicating *what, how* and *where* in the documentation the corrections were addressed (Corrections should be **highlighted** in the various documents as well).

The **total set of new documentation** should be included as this will then be the set used for monitoring purposes as required by the NHREC.



The updated application is re-sent to the same independent reviewers for the review of the corrections (three working days).



Corrections are either approved by reviewers or further corrections are requested.

If additional corrections are requested they should be corrected (as previously indicated) and re-submitted by the applicant to either:

- the AnimCare administration for research involving animals (Ethics-AnimCare@nwu.ac.za) or
- the HREC administration for research involving humans (please note that the corresponding person for the HREC remains Ethics-HRECProcess@nwu.ac.za during this reviewing process).



If approved, a letter of approval is sent to the researcher/s by either:

- the AnimCare administrator for research involving animals (Ethics-AnimCare@nwu.ac.za) or
- the HREC administrator for research involving humans (Ethics-HRECApply@nwu.ac.za).

The letter will either indicate **final approval** or **conditional approval** (Conditional approval is given when there are certain processes that have to occur before final approval can be given e.g. approval of a study from the Department of Health can only be applied for after the HREC gives approval however the HREC cannot approve the study without receiving the permission letter from the DoH, therefore

conditional approval is granted; or where interview schedules will be developed as the study unfolds. **These conditions will be clearly stated**).

Once the English version of the informed consent form has been finally approved, the applicants can have the form translated into the culturally relevant languages. This is to ensure that the applicants only have to translate the informed consent documentation once.



If a project has been conditionally approved, please send any other outstanding documents e.g. permission letters from authorities (e.g. Department of Health) that could only be obtained after ethical approval was obtained, to the appropriate administration in the Ethics Office as soon as possible (if applicable).

If the *conditions associated with the approval are process-linked* e.g. development of an interview schedule for phase two of a project is based on the results obtained during phase one of the project, then the research can continue until that point e.g. the end of phase one, where after the applicant **must submit the required documentation for approval before the study can continue**.

This documentation must be submitted to:

- the AnimCare administration for research involving animals (Ethics-AnimCare@nwu.ac.za) or
- the HREC administration for research involving humans (Ethics-HRECAppl@nwu.ac.za).

For research involving humans, please bring the approved informed consent documentation as well as the translated versions of the informed consent documents to be **stamped** by the Ethics Office before they are photocopied and used in the research (Contact Carolien van Zyl x 99 1206 for an appointment).



Research can begin as soon as the researcher has received the ethics approval letter.



The ethics certificate is only issued by the Institutional Office once all conditions are met.



If needed, send any **future amendments** of the proposal or the rest of the documentation to the appropriate administration:

- the AnimCare administration for research involving animals (Ethics-AnimCare@nwu.ac.za)
or
- the HREC administration for research involving humans (Ethics-HRECAppl@nwu.ac.za)

(See Section 3 (amendments) for the process)



For **minimal risk studies involving humans**, an **annual monitoring report** must be submitted for the duration of the study *at least two months before expiry* and annually until it has been completed. For **medium or high risk studies**, a **monitoring report must be submitted six monthly** for the duration of the study. Ensure that the monitoring report submitted for the end of the annual term, is submitted *at least two months before expiry* of the ethics approval of the project (**See Section 4 (monitoring reports) for the process**). For studies involving animals, *Category 0 to 4 studies* require an **annual monitoring report** to be submitted for the duration of the study which should be submitted *at least two months before expiry* of the ethics approval of the project. For *Category 5 studies* a **monitoring report must be submitted six monthly** for the duration of the study. Ensure that the monitoring report submitted for the end of the annual term, is submitted *at least two months before expiry of the ethics approval of the project*.

Note: Only one year ethics approval of projects may be granted due to legal requirements.

It must be indicated in the monitoring report whether the study is *completed or not*. If the study is completed, the monitoring report acts as a *final report*. If the study is not complete, the monitoring report acts as a request to *extend the study*.

NB If a study is **terminated** please immediately notify the appropriate administration.



Research dissemination/publication.

Checklist for attachments for a single study research ethics approval applications to the HREC:

Document		Tick if attached	Comment
1	Cover letter that indicates the title, researcher/s, the type of research ethics application, which documents are attached, and adds any explanations to clarify your application		
2	Executive summary of the project		
3	Proposal approved by a scientific/proposal committee		
4	An ethics application form to provide additional information not covered in the proposal		
5	Informed consent documentation and checklist (if collaborative study, informed consent from all the centers OR if an affiliated study, the original informed consent documentation of the original study)		
6	Advertisements or recruitment materials		
7	Questionnaire/s; interview schedule for interviews or focus groups		
8	Approval letter of the study by the scientific committee		
9	2-page narrative CVs of all the researchers in the project		
10	Proof of ethics training over the past three years for all the researchers in the project		
11	Permission letters from governing bodies to conduct the research		
12	Goodwill permission letters		

13	Any other applicable documentation e.g. MOU, contracts with collaborators/laboratories, permits etc.		
14	Signed NWU code of conduct for researchers for each team member		
15	Signed statistical consultation form		
16	Submitted as hard or scanned copies: Printed and signed pages of the ethics application form for the declarations by the project leader, statistical consultation services, director of the research entity		
17	Checklist of attachments		
	If applicable:		
18	Confidentiality agreement		
19	Indemnity form		
20	Form A for delegated ministerial consent in the case of greater than minimal risk research in children with no prospect of direct benefit to them		
21	Permission letter of the chairs of the HREC and HHREC if the study is an affiliated study or sub-study under a larger study falling on another campus than that where the student is registered		
22	Permission from the project leader if a study is done as an affiliated study under a larger study		
23	If any non-registered medication are used approval letter by the Medical Control Council		
24	If radio-active substances are used letter from the radiation control officer		

Attachments for applications to AnimCare:

	Document	Tick if attached	Comment
1	Cover letter that indicates the title, researcher/s, the type of research ethics application, which documents are attached, and adds any explanations to clarify your application		
2	An ethics application form to provide additional information not covered in the proposal		
3	Project proposal as approved by the Scientific Committee		
4	Scientific Committee's signed letter of approval of the project		
5	Monitoring sheets to observe any undue pain and suffering, and to manage (alleviate) pain and suffering when humane endpoints are reached		
6	Narrative CVs for each member of the project team (<i>project head, supervisor, researchers, students, co-workers, assistants, etc.</i>) and professional supervisors		
7	Copy of the signed NWU Code of Conduct for Researchers for each researcher in the project team (<i>project head, researchers, students</i>)		
8	Proof of ethics training (<i>minimum SANS training during the last 3 years</i>)		
9	Vivarium authorisation (following animal handling course and SAVC authorisation)		
10	Proof of SAVC authorisation (<i>included in Vivarium authorisation</i>)		
11	Proof of training in animal handling (<i>included in Vivarium authorisation</i>)		
12	Animal holding facility's certificate of SAVC registration		

13	Other permission letters, informed consent, permits and contracts as received from relevant governing bodies, collaborators, sponsors or owners		
14	Submitted as scanned copies: Printed and signed pages of the ethics application form for the declarations by the project head, statistical consultation services, directors of the school and the research entity e-mailed as scanned copies		
15	Checklist of attachments		

2. A research ethics approval application for a sub-study under an approved larger study

Process:

Conceptualize the sub-study and how it will fall within the approved larger study (Observing the specific problems, reading focused literature, discussion etc.).



Enter into negotiations with the project leader of the larger study, to ensure that he/she will be open for the sub-study to be conducted under the larger study.

Develop the research proposal for the sub-study and get the applicable accompanying documentation ready.



Submit the proposal to the *scientific/proposal committee* in your entity for scientific evaluation and approval.

Obtain a letter of approval by them that has to be attached to the application.



Once the proposal has been approved by the scientific/proposal committee submit the title registration request through the Faculty office (this is a process that runs parallel with the research ethics application).



Submit the new sub-study **proposal and the additional required documentation** either to:

- AnimCare administration for research involving animals (Ethics-AnimCare@nwu.ac.za) or
- HREC administration for research involving humans (Ethics-HRECAApply@nwu.ac.za).

Attach all the required documents *separately* to the e-mail (see attached checklist below)

Attach a covering letter indicating:

- the title of the research
- the researcher/s

- the type of research ethics application
- which documents are attached to the application, and
- add any explanation to clarify your application



Application sent by administration (four working days) to two to three independent reviewers (5 working days for review).



The application is discussed at the appropriate Research Ethics Committee meeting e.g. research involving animals at the AnimCare committee meeting and research involving humans at the HREC meeting

- Decision process
 - Aggregate individual views
 - Deliberation (debate)
 - Analogue (consensus)
 - Vote if necessary
- Decision
 - Approved
 - Approved with minimal/several changes
 - Deferred (too many changes and committee deliberation needed)
 - Disapproved (have to go back to the drawing board)



Formal letter of decision of the REC with attached independent reviewer reports are sent to the applicant (always the study leader or PI) as soon as possible (approximately three working days) after the meeting by the appropriate administrator.



Corrections are done by the applicant and are sent back as soon as possible to either:

- the AnimCare administration for research involving animals (Ethics-AnimCare@nwu.ac.za) or
- the HREC administration for research involving humans (please note that the corresponding person for HREC now changes to Ethics-HRECProcess@nwu.ac.za if corrections are needed).

A rebuttal letter should be included indicating *what, how* and *where* in the documentation the corrections were addressed (Corrections should be highlighted in the various documents as well).

The **total set of new documentation** should be included as this will then be the set used for monitoring purposes as required by the NHREC.



The updated applications are re-sent to the same independent reviewers for the review of the corrections (three working days).



Corrections are either approved by reviewers or further corrections are requested.

If additional corrections are requested they should be corrected (as previously indicated) and re-submitted by the applicant to either:

- the AnimCare administration for research involving animals (Ethics-AnimCare@nwu.ac.za)
or
- the HREC administration for research involving humans (please note that the corresponding person for the HREC remains Ethics-HRECProcess@nwu.ac.za during this reviewing process).



If approved a letter of approval is sent to the researcher/s by either:

- the AnimCare administration for research involving animals (Ethics-AnimCare@nwu.ac.za)
or
- the HREC administration for research involving humans (Ethics-HRECApply@nwu.ac.za)

The letter will either indicate *final approval* or *conditional approval* (Conditional approval is given when there are certain processes that have to occur before final approval can be given e.g. approval of a study from the Department of Health can only be applied for after the HREC gives approval however the HREC cannot approve the study without receiving the permission letter from the DoH, therefore *conditional approval* is granted. **These conditions is clearly stated**).

Once the English version of the informed consent form has been finally approved, the applicants can have the form translated into the culturally relevant languages. This is to ensure that the applicants only have to translate the informed consent documentation once.



If a project has been conditionally approved, please send any other outstanding documents e.g. permission letters from authorities (e.g. Department of Health) that could only be obtained after ethical approval was obtained, to the appropriate administration in the Ethics office as soon as possible (if applicable).

If the *conditions associated with the approval are process-linked* e.g. development of an interview schedule for phase two of a project is based on the results obtained during phase one of the project, then the research can continue until that point e.g. the end of phase one, where after the applicant **must submit the required documentation for approval before the study can continue**.

This documentation must be submitted to:

- the AnimCare administration for research involving animals (Ethics-AnimCare@nwu.ac.za)
or
- the HREC administration for research involving humans (Ethics-HRECAppl@nwu.ac.za).

For research involving humans, please bring the approved informed consent documentation as well as the translated versions of the informed consent documents to be *stamped* by the Ethics office before they are photocopied and used in the research (contact Carolien van Zyl x 99 1206 for an appointment).



Research can begin as soon as the researcher has received the ethics approval letter.



The ethics certificate is only issued by the Institutional Office once all conditions are met.



If needed, send any *future amendments* of the proposal or the rest of the documentation to the appropriate administrator:

- the AnimCare administration for research involving animals (Ethics-AnimCare@nwu.ac.za)
or
- the HREC administration for research involving humans (Ethics-HRECAppl@nwu.ac.za)

(See Section 3 (amendments) for the process)



For ***minimal risk studies involving humans***, an ***annual monitoring report*** must be submitted for the duration of the study *at least two months before expiry* and annually until it has been completed. For ***medium or high risk studies***, a ***monitoring report must be submitted six monthly*** for the duration of the study. Ensure that the monitoring report submitted for the end of the annual term, is submitted *at least two months before expiry of the ethics approval of the project* (See Section 4 (monitoring reports) for the process). For studies involving animals, *Category 0 to 4 studies require an annual monitoring report* to be submitted for the duration of the study which should be submitted *at least two months before expiry of the ethics approval of the project*. For *Category 5 studies* a ***monitoring report must be submitted six monthly*** for the duration of the study. Ensure that the monitoring report submitted for the end of the annual term, is submitted *at least two months before expiry of the ethics approval of the project*.

Note: Only one year ethics approval of projects may be granted due to legal requirements.

It must be indicated in the monitoring report whether the study is *completed or not*. If the study is completed, the monitoring report acts as a *final report*. If the study is not complete, the monitoring report acts as a request to *extend the study*.

NB If a study is *terminated* please immediately notify the appropriate administration.



Research dissemination/publication.

Checklist for attachments for a sub-study under a larger study research ethics approval applications to the HREC:

Document		Tick if attached	Comment
1	Have the data/biological samples already been gathered or is it in a process of longitudinal gathering or part of an intervention?	If yes: If no:	Continue Make sure the larger study truly qualifies as a larger study by completing the attached evaluation form .
2	Is the study clearly stated as an objective in the larger study	If yes: If no:	Continue Make sure the larger study truly qualifies as a larger study by completing the attached evaluation form .
3	Cover letter that indicates: <ul style="list-style-type: none"> Title of the larger study Title of the sub-study Student information Study-leader/s What the sub-study is about and how it fits into the larger study What documents are attached Detailed description of any outstanding issues of the larger study identified during the evaluation of the larger project (see evaluation form below) done by the project leader and how it will be addressed (Note: <i>This should be handled as a separate amendment to the larger study if it involves changes that will still take place in future and should be done before the sub-study is submitted for ethics approval</i>). 		
4	Executive summary of the sub-study		
5	Original proposal of the larger study		
6	Original informed consent documentation of the larger study		
7	Copy of the ethics approval certificate of the larger study		
8	Letter from the project leader clearly indicating <i>what can be done</i> as a sub-study under the larger project, as well as clearly specifying <i>what part of the previously collected data/biological samples can be used and for what</i>		
9	Approval letter of the sub-study by the scientific/proposal committee		
10	New proposal of the sub-study		
11	2-page narrative CVs of all the researchers in the sub-study		
12	Proof of ethics training over the past three years for all the researchers involved in the sub-study		
13	Signed NWU code of conduct for researchers for each team member		
14	Signed statistical consultation form		
15	Submitted as hard or scanned copies:		

	Printed and signed pages of the ethics application form for the declarations by the project leader, statistical consultation services, director of the research entity		
16	Checklist of attachments		
	If applicable:		
17	Confidentiality agreement		
18	Indemnity form		
19	Form A for delegated ministerial consent in the case of greater than minimal risk research in children with no prospect of direct benefit to them		
21	Permission letter of the chairs of the HREC and HHREC if the study is an affiliated study or sub-study under a larger study falling on another campus than that where the student is registered		
22	Evaluation form to see if the larger study qualifies as a larger study, completed by the project leader		

Checklist for attachments for a sub-study research ethics approval application to AnimCare:

	Document	Tick if attached	Comment
1	Cover letter that indicates: <ul style="list-style-type: none"> Title of the larger study Title of the sub-study Student information Study-leader/s What the sub-study is about and how it fits into the larger study What documents are attached Detailed description of any outstanding issues of the larger study identified during the evaluation of the larger project (see evaluation form below) done by the project leader and how it will be addressed (Note: <i>This should be handled as a separate amendment to the larger study if it involves changes that will still take place in future and should be done before the sub-study is submitted for ethics approval</i>).		
2	Larger project proposal as approved by the Scientific Committee		
3	Sub-study proposal as approved by the Scientific Committee		
4	Copy of the ethics approval certificate of the larger project		
5	Project head's permission letter that the sub-study may fall under the large project		
6	Scientific Committee's signed letter of approval of the sub-study		
7	2-page narrative CVs of all the researchers in the sub-study		
8	New monitoring sheets (<i>only if not included in the approved large project application</i>) to observe any undue pain and suffering, and to manage (alleviate) pain and suffering when humane endpoints are reached		
9	Copy of the signed NWU Code of Conduct for Researchers of new members only		
10	Proof of ethics training of new members only (minimum SANS training during the last 3 years)		
11	Vivarium authorisation of new members only (following animal handling course and SAVC authorisation)		
12	Proof of SAVC authorisation of new members only (included in Vivarium authorisation)		
13	Proof of training in animal handling of new members only (included in Vivarium authorisation)		

14	Other new permission letters, informed consent, permits and contracts as received from relevant governing bodies, collaborators, sponsors or owners		
15	If any non-registered medication are used approval letter by the Medical Control Council		
16	Submitted as scanned copies: Printed and signed pages of the ethics application form for the declarations by the project head, statistical consultation services, directors of the school and the research entity scanned and emailed		

3. Systematic review

In the case of a systematic review it *might/might not* have ethical implications when the study involves research with humans e.g. deciding on an intervention or leading to guidelines. When a minimal risk (or higher) exists, ethics approval is required. In most cases the journal expects an ethics approval number. To obtain such a number the research proposal needs to be evaluated by HREC.

Process:

Conceptualize the research study (Observing problems, reading literature, discussion etc.)



Develop the research proposal and applicable accompanying documentation.



Request the necessary application documents from the:

- HREC administration for research involving humans (Ethics-HRECAApply@nwu.ac.za).

OR

Download it from the ethics website



Submit the proposal to the *scientific/proposal committee* in your research entity for scientific evaluation and approval.

Obtain a letter of approval by them that has to be attached to the application.



Once the proposal has been approved by the scientific/proposal committee submit the title registration request through the Faculty office (this is a process that runs parallel with the research ethics application).



Submit the completed ethics application either to the:

- HREC administration for research involving humans (Ethics-HRECAppl@nwu.ac.za)

Attach all the required documents *separately* to the e-mail (see attachment checklist below).

Attach a *covering letter* indicating:

- the title of the research
- the researcher/s
- the type of research ethics application
- which documents are attached to the application, and
- add any explanation to clarify your application



Application sent by administration (four working days) to two to three independent reviewers (5 working days for review).



The application is discussed at Health Research Ethics Committee.

- Decision process
 - Aggregate individual views
 - Deliberation (debate)
 - Analogue (consensus)
 - Vote if necessary
- Decision
 - Approved
 - Approved with minimal/several changes
 - Deferred (too many changes and committee deliberation needed)
 - Disapproved (have to go back to the drawing board)



Formal letter of decision of the REC with attached independent reviewer reports are sent to the applicant (always the study leader or PI) as soon as possible (approximately three working days) after the meeting by the appropriate administrator.



Corrections are done by the applicant and are sent back as soon as possible to:

- the HREC administrator for research involving humans (please note that the corresponding person for HREC now changes to Ethics-HRECProcess@nwu.ac.za if corrections are needed).

A rebuttal letter should be included indicating *what*, *how* and *where* in the documentation the corrections were addressed (Corrections should be **highlighted** in the various documents as well).

The **total set of new documentation** should be included as this will then be the set used for monitoring purposes as required by the NHREC.



The updated application is re-sent to the same independent reviewers for the review of the corrections (three working days).



Corrections are either approved by reviewers or further corrections are requested.

If additional corrections are requested they should be corrected (as previously indicated) and re-submitted by the applicant to:

- the HREC administration for research involving humans (please note that the corresponding person for the HREC remains Ethics-HRECProcess@nwu.ac.za during this reviewing process).



If approved, a letter of approval is sent to the researcher/s by:

- the HREC administrator for research involving humans (Ethics-HRECAppl@nwu.ac.za).

The letter will either indicate **final approval** or **conditional approval** (Conditional approval is given when there are certain processes that have to occur before final approval can be given. **These conditions will be clearly stated**).



If a project has been conditionally approved, please send any other outstanding documents to the administration in the Ethics Office as soon as possible (if applicable).

This documentation must be submitted to:

- the HREC administration for research involving humans (Ethics-HRECAppl@nwu.ac.za).



Research can begin as soon as the researcher has received the ethics approval letter.



The ethics certificate is only issued by the Institutional Office once all conditions are met.



If needed, send any **future amendments** of the proposal or the rest of the documentation to the appropriate administration:

- the HREC administration for research involving humans (Ethics-HRECAppl@nwu.ac.za)

(See Section 3 (amendments) for the process)



For **minimal risk studies** involving humans, an **annual monitoring report** must be submitted for the duration of the study *at least two months before expiry* and annually until it has been completed. For **medium or high risk studies**, a monitoring report must be submitted **six monthly** for the duration of the study. Ensure that the monitoring report submitted for the end of the annual term, is submitted *at least two months before expiry* of the ethics approval of the project (See Section 4 (monitoring reports) for the process).

Note: Only one year ethics approval of projects may be granted due to legal requirements.

It must be indicated in the monitoring report whether the study is *completed or not*. If the study is completed, the monitoring report acts as a *final report*. If the study is not complete, the monitoring report acts as a request to *extend the study*.

NB If a study is **terminated** please immediately notify the appropriate administration.



Research dissemination/publication.

Checklist for attachments for a single study research ethics approval applications to the HREC:

Document		Tick if attached	Comment
1	Cover letter that indicates the title, researcher/s, the type of research ethics application, which documents are attached, and adds any explanations to clarify your application		
2	Executive summary of the project		
3	Proposal approved by a scientific/proposal committee		
4	A systematic review ethics application form to provide additional information not covered in the proposal		
5	Approval letter of the study by the scientific committee		
6	2-page narrative CVs of all the researchers in the project		
7	Proof of ethics training over the past three years for all the researchers in the project		
8	Signed NWU code of conduct for researchers for each team member		
9	Submitted as hard or scanned copies: Printed and signed pages of the ethics application form for the declarations by the project leader, statistical consultation services, director of the research entity		
10	Checklist of attachments		

4. Application for an amendment to an approved study

Process:

Decide what the required amendments are for the present study (*It might be that amendments require speedy approval*).



Review and update the proposal and any study documentation and indicate clearly where the possible changes have been made in order to amend the existing study (**using yellow highlight**).

Formulate a clear and systematic cover letter guiding the appropriate ethics committee e.g. AnimCare for research involving animals or HREC for research involving humans, through the amendments that have been made:

- the title of the research
- the researcher/s
- that it is an amendment request
- the nature of the amendment (Indicating what changes have been made and where)
- which documents are attached to the application, and
- add any explanation to clarify your application



Submit the amended ethics application either to the:

- AnimCare administration for research involving animals (Ethics-AnimCare@nwu.ac.za) or
- HREC administration for research involving humans (Ethics-HRECAppl@nwu.ac.za).

Attach all the required documents separately to the e-mail (see document checklist below).



Application sent by administration (within three days) to two to three independent reviewers (3 working days for review).



The application is handled as expedited (changes not of a large nature) or discussed at the next appropriate Ethics Committee meeting (if large changes are made) e.g. research involving animals at the AnimCare committee meeting and research involving humans at the HREC meeting if of a larger nature.

- Decision process
 - Aggregate individual views
 - Deliberation (debate)
 - Analogue (consensus)

- Vote if necessary
- Decision
 - Approved
 - Approved with minimal/several changes
 - Deferred (too many changes and committee deliberation needed)
 - Disapproved (have to go back to the drawing board)

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Formal letter of decision of the REC with feedback is sent to the applicant (always the study leader or PI) as soon as possible (approximately three working days) after the meeting by the appropriate administration or sooner if expedited.

↓

Corrections are done by the applicant and are sent back to either:

- the AnimCare administration for research involving animals (Ethics-AnimCare@nwu.ac.za) or
- the HREC administration for research involving humans (please note that the corresponding person for HREC now changes to Ethics-HRECProcess@nwu.ac.za).

A rebuttal letter should be included indicating *what, how* and *where* in the documentation the corrections were addressed (Corrections should be **highlighted** in the various documents as well).

The **total set of new documentation** should be included as this will then be the set used for monitoring purposes as required by the NHREC.

↓

The updated application is re-sent to the same independent reviewers for the review of the corrections (three working days).

↓

Corrections are either approved by reviewers or further corrections are requested.

If additional corrections are requested they should be corrected (as previously indicated) and re-submitted by the applicants to either:

- the AnimCare administration for research involving animals (Ethics-AnimCare@nwu.ac.za) or
- the HREC administration for research involving humans (please note that the corresponding person for the HREC remains Ethics-HRECProcess@nwu.ac.za during this reviewing process).

↓

If approved a letter of approval is sent to the researcher/s by either:

- the AnimCare administration for research involving animals (Ethics-AnimCare@nwu.ac.za)
or
- the HREC administration for research involving humans (Ethics-HRECAppl@nwu.ac.za)



Research can continue with the amended approach and documentation as soon as the researcher has received the ethics approval letter from the appropriate REC for the amendments.



If needed, send any **future amendments** of the proposal or the rest of the documentation to the appropriate administration:

- the AnimCare administration for research involving animals (Ethics-AnimCare@nwu.ac.za)
or
- the HREC administration for research involving humans (Ethics-HREC@nwu.ac.za)

Checklist for attachments for an amendment to a study to the HREC:

Document		Tick if attached	Comment
1	Cover letter that indicates the title, researcher/s, the nature of the amendment, and what has been changed where in the various attached documents (NB highlighted)		
2	Adjusted proposal		
3	Adjusted documentation (if applicable)		

Checklist for attachments for an amendment to a study to AnimCare:

Document		Tick if attached	Comment
1	Cover letter that indicates the title, researcher/s, the nature of the amendment, and what has been changed where in the various attached documents (NB highlighted)		
2	The updated ethics application form		
3	Amended project proposal (as approved by the Scientific Committee) with changes highlighted		
If applicable:			
4	Scientific Committee's signed letter of approval of the project		
5	Scientific Committee's signed letter of approval of the project amendment		
6	Any new/amended monitoring sheets (<i>to observe any undue pain and suffering, and to manage (alleviate) pain and suffering when humane endpoints are reached</i>)		
7	Narrative CVs of all <u>new</u> members of the project team (<i>not included in the original application</i>)		
8	Proof of ethics training for all <u>new</u> members (minimum SANS training during the last 3 years)		
9	Vivarium authorisation for all <u>new</u> members (following animal handling course and SAVC authorisation)		
10	Proof of SAVC authorisation for all <u>new</u> members (included in Vivarium authorisation)		

11	Proof of training in animal handling for all <u>new</u> members (included in Vivarium authorisation)		
12	Animal holding facility's certificate of SAVC registration for any <u>new</u> facilities (<i>not included in the original application</i>)		
13	Project head's and professional supervisor declarations forms (<i>as applicable to the amendment</i>)		
14	Other <u>new</u> permission letters, informed consent, permits and contracts as received from relevant governing bodies, collaborators, sponsors or owners		

5. Monitoring report/or request for extension of the study

A **compulsory** annual (in the case of minimal risk studies) and six monthly (in the case of medium and high risk studies) monitoring report of approved projects is required. This should be submitted at least *two months before the expiry date* of the study. The monitoring report requests a clear indication of the status of the study:

Status of study	Yes	No	NA
Has the study been completed and does this serve as your final report?			
Has this project been terminated? If so, please indicate the date, reason for termination and when HREC was notified:			
Does the project have to continue in the following year?			

If the study has not been completed an *extension* will automatically be granted for the project if the monitoring report is approved.

Note: Should you require an extension for the study at a time which does not fall within the required monitoring report period you can use the same process to request for an extension by completing the monitoring report. A cover letter should clearly indicate that this is what you require.

Monitoring report process:

For minimal risk studies, an annual monitoring report must be submitted for the duration of the study until it has been completed. For medium or high risk studies, a monitoring report must be submitted six monthly for the duration of the study.

Two months before the end of the ethics approval period indicated for the different risk level studies, the researcher needs to complete a monitoring report to be obtained from Ethics-Monitoring@nwu.ac.za or from the webpage.



Complete the monitoring report ensuring that all appropriate sections are completed.

It must be indicated in the monitoring report whether the study is *completed or not*. If the study is completed, the monitoring report acts as a *final report*. If the study is not complete, the monitoring report acts as a *request to extend the study*.



Submit your completed monitoring report to Ethics-Monitoring@nwu.ac.za (Please indicate your choice of option in the report).



The monitoring report is sent (within three working days) to two independent reviewers (5 days to review).



Feedback from the monitoring reports is consolidated and discussed at the appropriate Ethics Committee meeting e.g. research involving animals at the AnimCare committee meeting and research involving humans at the HREC meeting.

- Decision
 - Clarification
 - Completion
 - Suspended
 - Continuation
 - Termination



A formal letter of decision is sent to applicants as soon as possible by the administration.

If any clarification or feedback is requested, the applicants should send the required information within a week to Ethics-Monitoring@nwu.ac.za.

Clarifications are sent back to the same independent reviewers.



Clarifications are either approved by reviewers or further clarification is requested.

If additional clarification is requested, it should be corrected (as indicated) and re-submitted within a week by the applicant to Ethics-Monitoring@nwu.ac.za.

A letter will be sent to the applicant stating the status of the research. If it is a continuation, it will state the date for the next monitoring report.



The researcher can continue with the research as soon as he/she has received the letter indicating continuation.

NB Please notify the administration at Ethics-Monitoring@nwu.ac.za as soon as possible if the study is terminated unexpectedly.

Note: Extension request not falling in the monitoring report cycle:

If a researcher wants to **extend an approved research project at any other time other than the compulsory monitoring times** i.e. annually for minimal risk studies and six monthly for a medium or high risk study, the researcher can do so by submitting the same monitoring report with a very clear cover letter indicating that extension is requested that falls outside the monitoring cycle.