



GUIDANCE FOR LNR APPLICATIONS – FORMS AND DOCUMENTS

Below is a list of forms/documents required for LNR research applications reviewed by PH HREC.

New Applications

	Form/Document Name	Mandatory/Project Dependent	Guidance
Ethics Application	LNR VIC Application Form	Mandatory for all applications	Complete through ERM. First time users will need to create an account.
	Ethics Payment Form	Mandatory for all applications	Refer to the payment form and complete relevant section.
Ethics Supporting Documents	Protocol	Mandatory for all applications	Must match the protocol submitted as part of the Low Risk Confirmation. Attach as a supporting document to LNR Application Form.
	Participant Information and Consent Form (PICF)	Project dependent	Templates available from DHHS website. Other PICF formats are not accepted. PICF Health & Social Science for Self PICF Health & Social Science for Parent & Guardian PICF Health & Social Science for Person Responsible
	Implied consent wording for anonymous survey introduction	Project dependent	Template available
	Data Collection Tools	Project dependent	This includes questionnaires, surveys, focus group questions/themes, telephone questionnaires. For telephone questionnaires a Telephone Script Template is available.
	Advertising / Recruitment material	Project dependent	This includes letter or email invitations, posters, brochures or leaflets, content for media (including radio, print and digital/social media). See Advertising Guidelines
	Investigator CV for each researcher	Mandatory when a CV has not been submitted in the previous 3 years	Investigator CV Template available Full CV is not required
Research Governance Application	LNR VIC Site Specific Assessment (SSA) Form	Mandatory for all applications	Create SSA form as a sub-form of the LNR VIC form using ERM
Research Governance Supporting Documents	Research Agreement	Project dependent	Refer to Monash Partners Legal Document Repository Contact the Manager Office for Research if you are unsure if this is required for your project



Post Approval

Below is a list of the post-approval sub-forms which can be created from the **LNR VIC application** in ERM. Not all available forms are applicable. Please refer to the guidance below:

Form Type	Guidance from form	Use for
Amendment Request	<p>Once a research project has been ethically approved, any change to its design or conduct must be approved by the reviewing HREC or ethics review body.</p> <p>An amendment to a research project may also impact research governance/site specific assessment (SSA). The Research Governance Officer (RGO) at each affected site must be notified of the amendment by the site Principal Investigator (PI), in order to determine if research governance/SSA amendment is required.</p> <p>An amendment must not be implemented at a site until the HREC or ethics review body has granted approval of the amendment and (if applicable) the site RGO has granted authorisation of research governance/SSA amendment.</p>	<ul style="list-style-type: none"> • Add or amend documents • Change investigator or personnel • Request extension of HREC approval • Reactivate approval • Other (must specify) • Add site <p>*NOTE: Must tick 'Add or amend documents' when adding a new investigator so CV and Change to research personnel form can be uploaded to form.</p>
Safety Report	<p>The sponsor is responsible for reporting a safety event to the reviewing HREC, in accordance with Safety Monitoring and Reporting in Clinical Trials Involving Therapeutic Goods (NHMRC, 2016).</p>	Not required for LNR projects
Serious Breach Report	<p>A serious breach is a breach of GCP or the protocol that is likely to affect to a significant degree the safety or rights of a research participant of the reliability and robustness of the data generated in the research project. A serious breach must be notified to the reviewing HREC. The form must be completed by the sponsor. It may be used for reporting a serious breach to the HREC or for providing additional/follow-up information following notification by an individual/institution of a confirmed serious breach. Information on reporting breaches is available in Reporting Serious Breaches of GCP or the Protocol for Trials Involving Therapeutic Goods (NHMRC, 2018).</p>	Not required for LNR projects
Suspected Breach Report	<p>A suspected breach is a report that is judged by the reporter as a possible serious breach but has yet to be formally confirmed as a serious breach by the sponsor.</p> <p>This form must be completed when a third party (e.g. individual or institution) wishes to report a suspected breach of Good Clinical Practice (GCP) or the protocol. This should be reported directly to the reviewing HREC without reporting through the sponsor.</p> <p>Information on reporting breaches is available in Reporting of Serious Breaches of Good Clinical Practice (GCP) or the Protocol for Trials Involving Therapeutic Goods (NHMRC, 2018).</p> <p>A serious breach must be notified to the reviewing Human Research Ethics Committee (HREC) using the <i>Serious Breach Report</i>.</p>	Not required for LNR projects



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Project Progress Report	Information on the progress of an approved research project must be provided to the reviewing the Human Research Ethics Committee (HREC) or ethics review body, in accordance with the National Statement on Ethical Conduct in Human Research (NHMRC, 2007).	<ul style="list-style-type: none"> • See form guidance • Submit annually by 1 September • Report should cover the reporting period 1 July – 30 June each year • The Self Audit Tool (LNR VIC SSA sub-form – see below) must be submitted at the same time as the progress report.
Site Closure Report	<p>If an individual site closes from within an ongoing multi-site research project, the reviewing HREC or ethics review body must be notified.</p> <p>This <i>Site Closure Report</i> must be used for an individual site closure in a multi-site research project. If the research project is completed at all sites approved by the reviewing HREC, use the <i>Project Final Report</i> instead.</p>	<ul style="list-style-type: none"> • See form guidance • Not for single-site projects • For our site if we close prior to lead site
Project Final Report	<p>When a research project is completed at all approved sites, the reviewing HREC or ethics review body must be notified.</p> <p>This <i>Project Final Report</i> must be used when the research project is completed at all sites approved by the reviewing HREC.</p>	<ul style="list-style-type: none"> • See form guidance • For single-site studies that are either abandoned or completed
Project Notification Form	This notification form should be used to inform the reviewing Human Research Ethics Committee (HREC) or ethics review body of pertinent matters for which there is not a dedicated reporting form available.	<ul style="list-style-type: none"> • See form guidance • Report an adverse event/incident occurring to a participant (incidence resulting in harm must be notified to the Office for Research within 24 hours). Reporting in VHIMS may be required

Below is a list of the post-approval sub-forms which can be created from the **LNR VIC SSA application** in ERM. Not all available forms are applicable. Please refer to the guidance below:

Form Type	Guidance from form	Use for
Complaint Report	<p>If a complaint is made about a research project, the site Principal Investigator must report it to the site Research Governance Officer (RGO).</p> <p>The site RGO will advise whether the complaint should also be sent to the reviewing Human Research Ethics Committee (HREC).</p>	<ul style="list-style-type: none"> • See form guidance
Non-Serious Breach/Deviation Report	<p>A deviation is any breach, divergence or departure from the requirements of Good Clinical Practice (GCP) or the protocol that does not have a significant impact on the continued safety or rights of participants or the reliability and robustness of the data generated in the research project. If a deviation is considered to be a serious breach it should be reported using the <i>Serious Breach Report</i>.</p> <p>To fulfil GCP requirements, any deviation must be reported to the sponsor. The sponsor, in collaboration with the site Principal Investigator, should complete this <i>Non-serious</i></p>	<ul style="list-style-type: none"> • Not required for LNR Projects



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	<p><i>Breach/Deviation Report</i> form to inform the site Research Governance Officer (RGO) of a non-serious breach/deviation.</p> <p>Some deviations may require reporting to the reviewing Human Research Ethics Committee (HREC). The RGO will advise whether this is required and, if so, the form should be forwarded to the reviewing HREC. For a multi-site project, the Coordinating Principal Investigator should be informed if HREC reporting is required.</p>	
Site Audit Report	If the site Research Governance Officer (RGO) requests a self-audit report for a research project, the site Principal Investigator should complete this <i>Site Audit Report</i> .	<ul style="list-style-type: none">• This should be submitted annually at the same time as the annual progress report
Site Notification Report	This notification form should be used to inform the site Research Governance Officer (RGO) of site matters for which there is not a dedicated reporting form available.	<ul style="list-style-type: none">• Not required for LNR projects
Site Progress Report	The site Principal Investigator should report to the site Research Governance Officer (RGO) according to site policy.	<ul style="list-style-type: none">• Not required for LNR projects