



# All Ireland NCI Cancer Consortium (AICC) Research & Innovation Grant Scheme 2024

Pre-Application (Expressions of Interest) Guidance Notes



## Guidance Notes

Key Dates & Times	
Application Open	01 March 2024
Application Closing Date	26 April 2024 @13:00

*Applications must be completed and submitted through the HRB online Grant E-Management System (GEMS) (<https://grants.hrb.ie>), and this system will close automatically at the stated deadline and timeline listed above.*

*\*Prior to final submission to the HRB, all applications must first be reviewed and approved within GEMS by the authorized approver at the Host Institution as listed in the application form. It is critical therefore that applicants leave sufficient time in the process for the Research Office (or equivalent) in their nominated Host Institution to review, seek clarifications and approve applications prior to the final submission date. This may involve being aware of and complying with any internal Host Institution deadlines for review and approval, distinct from the HRB deadline.*

## Table of Contents

- 1 Introduction ..... 4
- 2 Aim and Objectives..... 5
- 3 Scope of Call..... 6
- 4 Funding Available, Duration and Start Date ..... 7
- 5 Eligibility Criteria ..... 8
- 6 Host Institution ..... 11
- 7 Application, Review Process and Assessment Criteria ..... 12
- 8 Timeframe..... 14
- 9 Contacts ..... 14
- Appendix I: Detailed Guidance on the Application Form..... 16
- Host Institution ..... 17
- Signatory Notification (within Host Institution) ..... 17
- 1 Proposal Details..... 18
- 2 Application Upload Section – Expression of Interest..... 18
- 3 Lead Applicant Information..... 20
- 4 Co-Applicant Details (to be included for each Co-Applicant) ..... 21
- 5 Collaborators’ Details ..... 21
- Submission of Application..... 21
- Appendix II: HRB Funding Policies and Procedures ..... 23
- Appendix III: Resources/Useful Links ..... 28
- Definitions ..... 37

## 1 Introduction

The Health Research Board's (HRB) Strategy<sup>1</sup> sets out our leading role to invest in research that delivers value for health, the health system, society, and the economy. The HRB achieves this by engaging with partners in the health and social care system and with other funders to facilitate dialogue on key issues and agendas to ensure stronger collaboration, coordination and prioritisation. The HRB collaborates with the Health & Social Care (HSC) Public Health Agency and other stakeholders in Northern Ireland to advance all island activities in areas of mutual interest.

The HSC, established in 2009, is responsible for the administration and coordination of the research & development budget on behalf of Department of Health, Northern Ireland (DoH NI). Through its current strategy, Research for Better Health and Social Care Strategy (2016-25)<sup>2</sup>, it seeks to ensure that research and development is an integral part of ensuring that health and social care services are of the highest quality and informed by the best available and up-to-date evidence.

The Ireland-Northern Ireland-National Cancer Institute Cancer Consortium (AICC) was originally launched in October of 1999, with the aim of reducing cancer incidence and mortality on the island of Ireland through cross-border and transatlantic collaborations in cancer research and education.

Cancer services across the island had undergone major restructuring because of several government reports. Specifically, the National Strategy Document for Cancer proposed that cancer treatment services should be centred around primary care services, regional services, and a national coordinating structure. Therefore, this was an opportune time to bring the National Cancer Institute (NCI) on board in a determined effort to redevelop and significantly improve services and outcomes for cancer patients throughout the island. The Consortium was regarded as a major step toward enhancing diplomatic relations between the United States, Ireland, and Northern Ireland, bringing the governments of the United States and both parts of the island together in the fight against cancer.

In March 2021 a new MoU was signed by Minister for Health Stephen Donnelly TD, Northern Ireland Minister for Health Robin Swann MLA, and Dr Norman E Sharpless MD on behalf of the NCI during a special virtual broadcast ahead of Saint Patrick's Day celebrations.

The MoU serves as an enabling framework, supporting cancer care and cancer research priorities, and offering more opportunities to develop North-South and Ireland-US collaborations. The renewed Consortium builds on previous successes and will provide an excellent framework to support national coordination of cancer research which is one of the key research actions in the National Cancer Strategy 2017-2025.

Priority themes for the All-Ireland NCI Cancer Consortium include prevention and integrated care pathways, research capacity, clinical trials, genomics and precision medicine.

Since 2021, the HRB has worked with the Department of Health (DoH) to establish an implementation mechanism and put in place appropriate structures to oversee the Consortium to ensure ongoing commitment to, and longevity of, the initiative.

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<sup>1</sup> <https://www.hrb.ie/strategy-2025/>

<sup>2</sup> <https://www.health-ni.gov.uk/publications/research-better-health-and-social-care-strategy-2016-25>

The current structures include:

- A Steering group made up of representatives of the three signatory parties.
- An Implementation group, currently chaired by the CEO of the HRB and a researcher from the University of Ulster, with representatives from health agencies (National Cancer Control Programme and equivalent in NI, National Cancer Registries (North and South), HRB, NCI, HSC Public Health Agency (NI) and from the research community (Cancer Trials Ireland, AICRI).
- Working groups (WG) in the three priority areas with membership drawn mainly from the above Implementation group:
  - Cancer Care Delivery and Survivorship
  - Clinical Research and Trials
  - Genomic and Precision Medicine

Both the HRB and HSC R&D Division have long supported the Cancer Consortium in a wide spectrum of its research-related activities and will now launch the AICC Scheme.

This is a dedicated funding stream to support research proposals within the three key priority areas identified by the AICC. It is a timely investment for the AICC's 25<sup>th</sup> Anniversary in September 2024.

## 2 Aim and Objectives

The aim of the AICC scheme is to support innovative, dynamic, and collaborative cancer research and innovation within the three key priority areas identified by the AICC to benefit all-island cancer patients.

The objectives of this scheme are:

- To stimulate, engage, develop and improve all-island and trans-Atlantic multi-disciplinary cancer research collaborations, networking, and engagement.
- Prime cancer research activity within the three key AICC priority areas:
  - 1. Cancer Care Delivery and Survivorship**
  - 2. Clinical Research and Trials**
  - 3. Genomic and Precision Medicine**
- Support high quality, innovative cancer research pilots, mapping activities, projects and/or training initiatives.

The overall goal of the consortium awards is to increase the level of collaboration and innovation amongst cancer researchers and experts across the three jurisdictions of Ireland, Northern Ireland, and the United States.

As such we expect collaboration across the three jurisdictions to form a key component of the research proposal. All applicants will be required to have cross border collaborators as co-applicants to be deemed eligible to apply e.g., a Lead Applicant from the Republic of Ireland requires a Co-Applicant from Northern Ireland and vice versa. Additionally, applicants will be required to have an NCI researcher/research group as a collaborator or other as appropriate. However, research proposals which are solely focused on strengthening and, or improving North-South cancer research

and innovation collaborations may apply with sufficient justification for exclusion of NCI partners in the proposal.

Applications can be submitted for pilot projects, mapping activities, standalone research projects, or the establishment of a training scheme/programme, or similar.

### 3 Scope of Call

As the AICC scheme is intended as a dedicated pilot-& learn funding stream to enhance, extend and strengthen research and innovation capability, across the island of Ireland supported via trans-Atlantic partnerships, collaborative scientific research projects including mapping, networking and pilot activities and/or exchanges and training of researchers and experts, aligned to the HRB's strategic remit, will be supported.

The research proposal must fall under one of three priority themes as identified by the AICC:

- 1. Cancer Care Delivery and Survivorship**
- 2. Clinical Research and Trials**
- 3. Genomic and Precision Medicine**

The AICC award is intended as a strategic and focused investment and must demonstrate added-value above and beyond any research activities, collaboration or networking that is currently taking place.

*Where an application does not address the aims, objectives, and scope of the call, the application will be deemed ineligible and will not be accepted for review.*

#### 3.1 Out of Scope

Out of scope for this scheme:

- Applications which are solely or predominately developing the infrastructure for biobanking, databases or patient registers\*.
- Applications which are solely or predominately health service developments without a predominant research element. The HRB will not fund the cost of providing the service itself, only the research element.
- Applications from individuals applying for, holding, or employed under a research grant from the alcohol industry, tobacco industry or related actors.<sup>3,4</sup>

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<sup>3</sup> Any company, entity, or organisation involved in the development, production, promotion, marketing, or sale of tobacco in any country of the world. The term also includes any companies that are a subsidiary or a holding company or affiliate of the above. This also includes e-cigarette companies and non-tobacco related companies which are fully or partially owned by the tobacco industry.

<sup>4</sup> Including social aspects/public relations organisations (SAPROs) funded by alcohol companies or trade associations in which such companies are members.

- Infrastructure involving building work, fit-out of buildings, or purchase of major pieces of equipment.
- Applications involving basic biomedical research.
- Research intended to create human embryos solely for the purpose of research or for the purpose of stem cell procurement, including by means of somatic cell nuclear transfer.
- Applications using cell lines, animals or their tissue that do not constitute pre-clinical research (see [Appendix II](#) for a definition of pre-clinical research in the context of this scheme).
- Applications which are solely literature reviews, audits, questionnaires/surveys, or technology development (although these elements may be part of an integrated mapping activity, pilot or project/research study).
- Applied research projects to generate evidence for local/national health and social care needs/priorities, where the results are not generalisable or actionable beyond the local/national setting. Such applications may be suitable for submission to the HRB Applied Partnership Awards (APA) scheme.
- Applications seeking to evaluate a definitive intervention or a stand-alone feasibility study<sup>5</sup> for a definitive intervention. Such studies are supported through the HRB Definitive Intervention and Feasibility Awards (DIFA) scheme.

*\*Due to the critical importance of having a sustainable funding model and appropriate governance for patient registries, this scheme should not be used to establish new registries or to subvert the costs of maintenance of existing registries. Consideration can be given, however, to expanding and optimising the use of registries to support cancer research.*

**Where an application is outside the scope of the scheme, the application will be deemed ineligible and will not be accepted for review.**

## 4 Funding Available, Duration and Start Date

The AICC Scheme design is a two-tiered scheme intended to maximise and stimulate potential activity and impact within the three key priority research areas identified by the AICC.

**Tier one** of the scheme is designed to allow for smaller scale applications which involve pilot projects or mapping activities due to occur over a maximum duration of two years to apply for funding, exclusive of overheads, up to a maximum of €100,000.

Mapping activities may include cross-border needs assessments e.g. in relation to education/training or cancer research needs to improve cross-border cancer collaboration and activity. Audits/surveys or questionnaires alone are not eligible unless part of a larger mapping activity project.

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<sup>5</sup> Sandra M. Eldridge et al. *Defining Feasibility and Pilot studies in preparation for Randomised Controlled Trials: Development of a conceptual Framework.* PLoS ONE 11(3): e0150205

**Tier two** is for specific, targeted projects or potential training initiatives which may occur over a maximum 3-year duration to apply for funding awards, exclusive of overheads, up to a maximum of €330,000.

The total funding envelope for the AICC scheme including overhead contribution is €1,200,000. The awards allocated across tiers will be decided by international panel review and scoring.

The AICC 2024 award will provide support for research-related costs including salary for research staff, running costs, FAIR data management costs, equipment and dissemination costs, and overheads contributions. The HRB will calculate overheads during contracting.

Note: The AICC 2024 Award will not fund the salary and related costs of tenured academic staff within research institutions (including buy-out from teaching time etc.).

The budget requested and the award duration must reflect the scale and nature of the proposed research with sufficient justification. Reviewers will thoroughly assess each application, the level of funds and timeframe requested.

Eligible scheme costs include:

- Contributions to personnel costs: for example, AICC Project Research Coordinator (to ensure coordination across Ireland and connectivity with external collaborators).
- Running costs: for example, travel costs, PPI costs, training and exchange bursaries, networking events
- Dissemination and knowledge exchange costs
- Overhead contribution of 25-30% based on the proposed activity.

The earliest start date of the Grant is 01 November 2024.

## 5 Eligibility Criteria

This call is open to Host Institutions from the Republic of Ireland and Northern Ireland.

### 5.1 Applicant Team

Applications should be made on behalf of a team made up of researchers, knowledge user(s) and PPI contributors.

All applicants will be required to have cross border collaborators as co-applicants to be deemed eligible to apply e.g., a Lead Applicant from the Republic of Ireland requires a co-applicant from Northern Ireland and vice versa. Additionally, applicants will be required to have an NCI collaborator, however research proposals which are solely focused on strengthening and, or improving North-South cancer research and innovation collaborations may apply with sufficient justification of NCI exclusion.

The Lead Applicant will serve as the primary point of contact for the HRB during the review process and on the award, if successful. The Lead Applicant will be responsible for the scientific and technical direction of the research project. They have primary fiduciary responsibility and accountability for



carrying out the research within the funding limits awarded and in accordance with the terms and conditions of the HRB.

The Lead Applicant **must**:

- Hold a post (permanent or a contract that covers the duration of the award) in a HRB recognised Host Institution in the island of Ireland (the “Host Institution”) as an independent investigator. For clinicians, an adjunct position in a HRB recognised Host Institution is acceptable.

**OR**

- Be an individual who will be recognised by the Host Institution upon receipt of an award as an independent investigator who will have a dedicated office and research space for the duration of award, for which they will be fully responsible. The Lead Applicant does not necessarily need to be employed by the Host Institution at the time of the application submission.

They **must** show evidence of achievement as an independent researcher in their chosen research field by:

- a) Demonstrating a record of research output, with at least three publications of original research in peer reviewed journals. Where appropriate, they should also provide evidence of other outputs (e.g., published book chapters, reports to government, research data and datasets, research materials, databases, audio/video products, national and/or international reports, patents, models and protocols, software production, evidence of influence on health policy and practice, outreach and/or knowledge translation activities, media coverage or other relevant activities) and/or any other relevant outputs that have resulted in a significant impact in their field.
- b) Demonstrating record of independence by showing that they have secured at least one peer-reviewed research grant for a research project/s, as either the Lead Applicant or a Co-Applicant. Funding received for travel to seminars/conferences and/or small personal bursaries will not be considered in this regard.
- c) Show evidence that they possess the capability and authority to manage and supervise the research team.

**Applicants may apply to one, or both tiers.**

***Where an applicant fails to meet the eligibility criteria, the application will be deemed ineligible and will not be accepted for review. The HRB will contact the Lead Applicant if this situation arises.***

### **5.1.1 Co-Applicants**

All applications must include a cross-border Co-Applicant.

Co-Applicants will be asked to select whether they are a Researcher, Knowledge User, Data Controller/Processor, or PPI contributor Co-Applicant for the purpose of the proposed research.

Up to a maximum of 8 Co-Applicants can be included.

A Co-Applicant has a well-defined, critical, and substantial role in the conduct and steering of the proposed research. Co-Applicants from outside the island of Ireland are welcome where this is appropriately justified in terms of added value for the project. A Co-Applicant may receive funding for items such as running costs and personnel but will not receive support towards their own salary if they are in salaried positions. However, researchers in contract positions/independent investigators, knowledge user and PPI contributor Co-Applicants can request their own salary, depending on their role and percentage of time dedicated to the research for the duration of the award (up to a maximum of 8 Co-Applicants can be listed).

Each Co-Applicant must confirm their participation and is invited to view the application form online. The terms of any co-application should be determined early, and relevant agreements should be in place by the onset of the project. The HRB advise that consideration should be given to issues such as relative responsibilities, governance arrangements, intellectual property rights, reporting and access to data and samples when working up co-application agreements.

### 5.1.2 Collaborators

All applications must include a NCI Collaborator.

The NCI Collaborator may be an intramural or extramural researcher depending on the specific proposal and area of expertise required. Any NCI Collaborator proposed as part of the Applicant Team, would be expected to be involved and well-connected with and part of the wider NCI community actively engaged within NCI structures, committees, working groups, frameworks, networks, designated cancer centres etc. in order to meet eligibility expectations.

If the NCI collaborator proposed is part of a NCI designated cancer centre and connected/participating within the wider NCI networks this meets eligibility expectations. The aim is to increase trans-Atlantic NCI and cross-border collaborations.

**Solely, being a current or previous grant holder from NCI without ability to further demonstrate connectivity within the wider NCI community and structures would not meet eligibility requirements.**

A Collaborator is an individual or an organisation who will have an integral and discrete role in the proposed research and is eligible to request funding from the award when properly justified. Named collaborators may include investigators or organisations from outside the Republic of Ireland, but an individual or organisation should only be named as Collaborator if they are providing specific contributions (either direct or indirect) to the activities. A collaborator may provide training, supply samples or kits, provide access to specific equipment, specialist staff time, staff placements, access to data and/or patients, instruments or protocols, industry know-how, or may act in an advisory capacity. Collaborators can come from a range of backgrounds such as academia, the private sector, a healthcare organisation, the charity sector, or a patient group (up to a maximum of 10 Collaborators can be listed).

Profile details must be provided for ALL collaborators. In addition, each collaborator must complete a Collaboration Agreement Form. A template Collaborator Agreement form will be made available on GEMS for download.

If access to samples, vulnerable population groups, healthy volunteers or patients, data, databases, or a link to an existing national or international study (e.g., an existing cohort or longitudinal study) are an integral part of the proposed project, evidence of commitment and access must be demonstrated by having the [Data controller](#) or key Gatekeeper of a study included as a Collaborator.

The applicant team will be asked to describe any relevant agreements that they have entered into to facilitate their partnership working. The terms of any collaboration should be determined early, and relevant agreements should be in place by the onset of the project.

The HRB advise that consideration should be given to issues such as relative responsibilities, governance arrangements, ownership and copyright, access and sharing of data and samples etc. when working up Partnership proposals.

### 5.1.3 Funded Personnel

Applicants must demonstrate that the level, expertise, and experience of proposed research personnel matches the ambition and scale of the project and that they possess the necessary breadth and skills in all methodological areas required to deliver the proposed programme of work. Alignment between personnel requested and the proposed project should be demonstrated. Roles and responsibilities of funded personnel must be differentiated and clear.

This scheme is not framed as a training initiative for PhD candidates.

## 6 Host Institution

A HRB Host Institution is a research-performing organisation approved by the HRB for the purpose of receiving and administering HRB grant funding and is responsible for compliance with all general and specific terms and conditions of awards. HRB Host Institution status is a requirement to submit an application under all HRB award schemes. The **Host Institution for the award** is normally that of the **Lead Applicant** but it may be another organisation/institution designated by the research team, where it is clearly justified. In order to be eligible to apply for funding, an Institution must be an **approved** HRB Host Institution no later than two calendar months before the closing date of a call. A list of currently approved HRB Host Institutions and information on the application process for research performing organisations to be approved as HRB Host Institutions can be found on the HRB website<sup>6</sup>.

Please note that this call is open to Host Institutions from **Republic of Ireland and Northern Ireland**. **If invited to Full Application stage, a Host Institution Letters of Support** must be provided for **(1) all Lead Applicants in a contract position and (2) Co-Applicants in a contract position who are seeking their own salary**. The formal letter on headed notepaper, dated and signed by the Head of School/Research Centre/Hospital must include the following information; [*Host Institution – insert name*] which is the host institution of [*applicant – insert name*] confirms that [*applicant – insert name*]: (i) holds an employment contract which extends until [*insert date*] or will be recognized by the host institution upon receipt of the HRB All Ireland Cancer Consortium (AICC) 2024 award as a

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<sup>6</sup> <http://www.hrb.ie/funding/funding-schemes/before-you-apply/all-grant-policies/hrb-policy-on-approval-of-host-institutions/>

contract researcher; (ii) has an independent office and research space/facilities for which they is fully responsible for at least the duration of the award, and (iii) has the capability and authority to mentor and supervise the research team. Electronic signatures are acceptable for letters that are uploaded on the HRB GEMS system.

## 7 Application, Review Process and Assessment Criteria

### 7.1 Grant E-Management System (GEMS)

Applications must be completed and submitted through the HRB online Grant E-Management System (GEMS) (<https://grants.hrb.ie/>).

The application must have been reviewed and approved by the signatory approver at the research office (or equivalent) in the Host Institution before it is submitted to the HRB. Therefore, applicants should ensure that they give the signatory approver sufficient time before the scheme closing date to review the application and approve it on GEMS. Please note that many host institutions specify internal deadlines for this procedure.

The HRB is committed to an open and competitive process underpinned by international peer review. To ensure the integrity of the assessment process, conflict of interest and confidentiality are applied rigorously in each stage of the process.

### 7.2 Review Process

The AICC 2024 will use a two-stage application process consisting of:

#### **Stage 1 – Open call for Expressions of Interest submitted as Pre-Applications from both tiers**

The pre-application consists of an Expression of Interest (EOI) which will focus on project strategy, added value of the project, governance, project management and project budget.

Pre-applications will be reviewed by the HRB and HSC R&D +/- NCI to review alignment with cancer consortium research areas, cancer strategies and proposed collaborations. Applicants will be shortlisted based on these criteria. A brief feedback document will be provided to all applicants.

Shortlisted applicants will then be invited to Stage 2 to submit a Full Application (Stage 2). Full applications will be reviewed by an international Panel according to the assessment criteria for Stage 2.

#### **Stage 2 – Full Application (by invitation only)**

Information provided in the HRB online Grant E-Management System (GEMS) (<https://grants.hrb.ie/>) at Pre-Application stage will feed automatically into the invited full application forms.

The Lead Applicant (LA) will have the opportunity to make revisions from pre-application to full application stage (e.g., addition of expertise/partner, revision of targeted profession/disciplines for training, strengthening the stakeholder participation, etc.). However, full applications should reflect a development of the relevant pre-applications rather than a radically different approach.

Full applications, once submitted, will undergo a **two-step** assessment process as follows:

- Applications will be initially checked for eligibility by HRB staff members.
- Following the initial eligibility check, each eligible application submitted to this scheme will undergo an international panel review process.

## Panel Review

An international grant selection Panel will be convened. Panel members are selected based on the range of applications received and the expertise and skillset needed (e.g., research area and methodological and analytical approaches, coaching and mentoring, knowledge translation/applied health research, etc.). Panel members are assigned as lead and secondary reviewers to specific applications.

Panel members have access to the application. HRB and HSC R&D staff members are present at the meeting to clarify any procedural aspects for the Chair or Panel members and to take notes for the feedback process.

The Panel will review the strengths and weaknesses of the application relating to the assessment criteria detailed [below](#). Successful applicants are expected to score well in all review criteria. While PPI is not a stand-alone assessment criterion, it may influence scores under any criterion as relevant to the application.

At the end of the panel meeting, a final score is collectively agreed for each application and then they will be ranked according to score.

Gender balance of the Lead Applicant will be considered where required to prioritise proposals with the same scores in the Panel ranking list.

The recommendations of the Review Panel will be presented for approval at the next scheduled HRB Board meeting. When the Board of the HRB has approved the process and recommendations, HRB staff will contact the Lead Applicants and Host Institutions to notify them of the outcome. A summary of Panel Member's comments and the panel discussion comments will be issued to the Lead Applicant following the Board approval stage.

## 7.3 Assessment Criteria

The following assessment criteria will be used to assess applications by the panel reviewers. Successful applications will be expected to achieve **high performance in all criteria**.

### Added value of investment

- Proposed investment has clear added value above and beyond any research activities, collaboration or networking currently taking place.
- HRB support complements and leverages other investments.
- Existing cancer research capabilities and collaborations in Ireland harnessed to ensure impact for patients.

### Quality and relevance of proposed award activities

- Investment is strategic and targeted to boost all island and trans-Atlantic collaboration and partnerships in cancer research and innovation.
- Spectrum of activities/supports is appropriate to increase cancer research in areas of need.
- Coordinated approach to engagement with the NCI evident.
- Proposed work is prioritised and phased appropriately.
- Budget and resource details proposed are clear and appropriate

### Strength of collaboration and team

- Suitable applicant team, with complementary expertise and experience including appropriate stakeholders to maximise impact for patients.
- Roles and responsibilities of proposed team are clear.
- Proposed team can deliver on objectives of the award.
- Representation from Ireland, Northern Ireland and NCI Contributors as appropriate.
- Clear, strong links with relevant HIs/Hospitals/research groups/patient organisations evident in governance arrangements.
- Accessibility and suitability of facilities, infrastructure and other supports, as appropriate.

Panel members will be advised to take PPI approaches into consideration under any of the assessment criteria if considered relevant.

## 8 Timeframe

Date	
01 March 2024	Call Opening Pre-Application Stage
26 April 2024 @13:00	Call Closing Pre-Application Stage
May 2024	Shortlisting Review Meeting
04 June 2024	Call Opening Full-Application Stage
02 Aug 2024 @13:00	Call Closing Full-Application Stage
Sept 2024	Panel Review Meeting
Sept 2024	Panel recommendations presented to HRB Board
Oct 2024	Contracting stage (subject to approval)
01 Nov 2024	Earliest start date

## 9 Contacts

For further information on the AICC 2024 scheme contact:

**Dr Chiara Mizzoni**

email: [cmizzoni@hrb.ie](mailto:cmizzoni@hrb.ie)

Project Officer

Research and Innovation Infrastructures

Research Strategy and Funding

**The HRB reserves the right to reject any application that does not meet the terms of this call. The HRB's Policy on Appeals on funding decisions is available at <https://www.hrb.ie/funding/funding-schemes/before-you-apply/all-grant-policies/hrb-policy-on-appeals/>.**

## Appendix I: Detailed Guidance on the Application Form

Only registered users of the GEMS system can apply for grants. In order to submit an online application to the HRB, applicants are required to register at the following address:

<https://grants.hrb.ie>

Please refer to the GEMS Technical Guidance Notes, available on the left-hand column of your GEMS profile homepage, for further information.

The Lead Applicant must create the application, and it can then be jointly completed with named co-applicants or on behalf of a group/organisation.

- Lead Applicants can register on GEMS and they will receive an email to confirm their registration and log in details. The Lead Applicant can then add information on their contact and CV details in the 'Manage My Details' section of their GEMS account.
- Lead Applicants previously registered on GEMS can login to their GEMS account and update any information regarding their basic CV details in the 'Manage my details' section.

Once logged in to GEMS applicants are taken directly to the Home Page which is the starting point to create a new Grant Application.

When the Lead Applicant opens a new application in GEMS, they will be asked to complete a check list of mandatory questions. In order to access the application form, the Lead Applicant must satisfy the conditions of this check list. The checklist for the All-Ireland Cancer Consortium 2024 is as follows:

Requirements	
I have read the Guidance Notes for the AICC 2024 call.	✓
I am clear about the role of the authorised signatory in the nominated Host Institution, and I am aware that I need to build sufficient time into the application process for the HI to access, review and approve my final proposal for submission to the HRB through the GEMS system.	✓
By submitting this application, I agree to (a) sharing of my data outside of the European Economic Area (EEA) for the purpose of international peer review, and (b) the use of my data for assessment of my application; monitoring of successful awards; and evaluation of HRB's approach to funding and investment in research, in line with HRB policies and as detailed in the AICC 2024 Call Guidance Notes.	✓
I confirm I hold a post in a HRB recognised Host Institution in the island of Ireland as an independent investigator or (2) I am a clinician who holds an adjunct position in a HRB recognised Host Institution, or (3) I am an individual who will be recognised by the Host Institution upon receipt of an award as an independent investigator who will have a dedicated office and research space for the duration of award.	✓



The Lead Applicant will be then able to select the Host Institution and Notify the Authorised Signatory before starting the application. Further details for completing each of the main sections of the application are provided below.

## Host Institution

A **HRB Host Institution** is research performing organisation that is approved by the HRB for the purpose of receiving and administering HRB grant funding and is responsible for compliance with all general and specific terms and conditions of awards. HRB Host Institution status is a requirement to submit an application under all of the HRB's award schemes. The **Host Institution for the award** is normally that of the Lead Applicant but it may be another organisation/institution designated by the research team, where it is clearly justified. Information is available on the HRB website on the current approved Host Institutions and on the application process for research performing organisations to be approved as HRB Host Institutions<sup>7</sup>.

In GEMS you will be asked to identify a Host Institution (from this list) and type it in full (do not use acronyms such as UCD, TCD, UG). Once you have entered the first 3-4 characters of the Host Institution, you will be assisted with auto-select options. It is important that the HI name is entered accurately and in full as an incorrect entry may result in delays in attaining Host Institution approvals.

It is important to note that the Host Institution for the award is the body in charge of the financial and administrative co-ordination of the AICC grant award. This must be one of the collaborating academic institutions.

## Signatory Notification (within Host Institution)

Once the Host Institution is selected at the initial stages of application creation this will allow the Lead Applicant to notify the **authorised signatory** (Dean of Research or equivalent person authorised to endorse research grant applications for the Host Institution) in that Host Institution of the Lead Applicant's intention to submit an application to this scheme. The signatory's details are pre-populated in the system, so the applicant just needs to click 'NOTIFY' within GEMS. **We recommend that you notify the HI signatory of your intention to apply for the full application as soon as possible in the application process.** The signatory will receive an email from GEMS with the name and email details of the Lead Applicant and if they have any queries or clarifications, they can engage directly with the applicant to resolve them. The HI signatory must confirm their willingness to participate as HI for the application through GEMS and once they do this a PDF of the application will be available for them to review with a view to them ultimately approving the final version of the proposal for submission to the HRB.

Only the Host Institution for the award, which is the body in charge of the financial and administrative co-ordination of the AICC grant award is required to authorise the submission of the application to the HRB.

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<sup>7</sup> <https://www.hrb.ie/funding/funding-schemes/before-you-apply/all-grant-policies/approval-of-host-institutions>.

# 1 Proposal Details

## 1.1 Title

This should be descriptive and concise and should reflect the aim of the project.

## 1.2 Title Acronym

This is optional.

## 1.3 Abstract

This should be a succinct summary of the proposed research project. The aims and main objectives should be conveyed with clarity and what the programme is expected to establish should be described. Ideally the abstract provides a clear synopsis of your proposal. The word limit is **300 words**.

## 1.4 Type of Funding Requested

**AICC 2024 is a two-tier scheme.**

**Tier one** of the scheme is designed to allow for smaller scale applications which involve *pilot projects or mapping activities* up to a maximum individual award, exclusive of overheads totalling **€100,000** over a two-year duration.

**Tier two** is for specific, targeted *projects or potential training initiatives* which may occur over a 3-year period. This is exclusive of overheads and should not exceed **€330,000**.

Overheads are calculated at 25% for desk-based research or 30% for all other research.

Please select which tier you wish to apply for:

**Tier 1**

**Tier 2**

# 2 Application Upload Section – Expression of Interest

Section two of the application form is an upload option for an expression of interest letter. The purpose of the upload format is to allow the applicants to collate and include the required information as they see fit.

The word limit of this entire section is **1500 words** (excluding references) you may also include figures and/or tables as appropriate.

The upload file size is unlimited and must be uploaded in pdf format. The following items should be included. Applicants have the flexibility to include additional items which may not be listed.

## 2.1 Proposal Strategy

- Outline the **national context and need for this research project** in relation to the Irish health research and services system North and South
- Alignment with [HRB Strategy 2021 – 2025](#)
- Alignment with cancer strategies for both/either the Republic of Ireland and/or Northern Ireland.
- Alignment with one of the AICC research priority areas:
  - Cancer care delivery and survivorship *or*
  - Clinical research *or*
  - Genomics and precision medicine
- The **ambition, vision, aims and objectives** of the programme.

## 2.2 Added Value of the Proposal

Please clearly describe the added value of this proposal in the context of the All-Ireland(north/south) cancer environment.

## 2.3 Governance and Management (roles, responsibilities and memberships)

It is expected that Applicant team will put appropriate management and governance structures in place to ensure the efficient operation of the team and the delivery of the main objectives of the research project.

## 2.4 Budget

Please provide a brief summary of your funding request and justification of the costs. Tier one applications can apply for a maximum total award €100,000, exclusive of overheads. Tier two applicants can apply for a maximum total award value of €330,000 exclusive of overheads.

If invited to full application stage, a detailed breakdown of costings and justification for all funding will be required. The budget requested must reflect the scale and nature of the proposed project and reviewers will thoroughly assess the level of funding when reviewing the proposal.

Eligible scheme costs include:

- **Contributions to Personnel costs:** *for example*, AICC Project Research Coordinator (to ensure coordination across Ireland and connectivity with external collaborators).
- **Running costs:** *for example*, travel costs, PPI costs, training and exchange bursaries, networking events
- **Dissemination and knowledge exchange costs**
- **Overhead contribution** of 25-30% based on the proposed activity. Please indicate the overhead % requested based on the proposed work.

## 2.5 References

A full description of the references cited should be provided. Please include a **maximum of 10 publications**.

### For peer-reviewed publications:

Gallagher PA, Shoemaker JA, Wei X, Brockhoff-Schwegel CA, Creed JT. Extraction and detection of arsenicals in seaweed via accelerated solvent extraction with ion chromatographic separation and ICP-MS detection. *Fresenius J Anal. Chem.* 2001 Jan 1;369(1):71-80. PMID: 11210234.

### For book and printed source citations:

Farrell M, Gerada C and Marsden J (2000) External review of drug services for the Eastern Health Board. London: National Addiction Centre.

### For data citation:

Authors, year, article title, journal, publisher, DOI

Author(s), year, dataset title, data repository or archive, version, global persistence identifier

## 3 Lead Applicant Information

### GEMS Profile Details – Basic CV information

The Lead Applicant's CV details (Name, ORCID iD, institution, profession, education and employment history) are managed under the "Manage my Details" section of their GEMS account.

**Note:** The HRB is an ORCID member. Leadership team members are encouraged to include an ORCID iD by updating their GEMS profile under 'Manage my Details' and this will feed automatically into the application form. You have also the option to import your publication record from ORCID iD in addition to PubMed. Please note this is not a mandatory field for submitting the application. For more information and to register please see <https://orcid.org/>.

### ORCID iD

To include an ORCID iD in your application please update your GEMS profile under 'Manage my Details'. For more information and to register please see <https://orcid.org/>.

### Gender

Please select:

- Man
- Woman
- Nonbinary
- Another gender identity
- Prefer to not disclose

This question is included with the application form in light of the **HRB Gender Policy**. The HRB has the responsibility to support everyone to realise their full potential in order to ensure equality of

opportunity and to maximise the quantity and the quality of research. The information is for HRB internal use only.

## 4 Co-Applicant Details (to be included for each Co-Applicant)

Co-applicant CV details (name, institution, profession, education and employment history) and funding records are managed under the “Manage my Details” section of your GEMS account.

**Please note** you do not need to complete or update your publications under ‘Manage my Details’ as they will not feed through to this application and you will be asked to enter them manually in the section below.

### ORCID iD

To include an ORCID iD in your application please update your GEMS profile under ‘Manage my Details’. For more information and to register please see <https://orcid.org/>.

### Gender

This question is included with the application form in light of the HRB Gender Policy. The HRB has the responsibility to support both women and men to realise their full potential in order to ensure equality of opportunity and to maximise the quantity and the quality of research. **The information is for HRB internal use only.**

Please select:

- Man
- Woman
- Nonbinary
- Another gender identity
- Prefer to not disclose

## 5 Collaborators’ Details

The Lead Applicant can add **up to 10 collaborators** per application. Unlike Co-Applicants, the information for Collaborators **is not** automatically drawn from the ‘Manage my Details’ section of GEMS but must be entered by the Lead Applicant. The Lead Applicant must enter **contact and CV details** for all Collaborators including name, contact information, institution or organisation and present position.

In addition, for each Collaborator a signed **Collaboration Agreement Form** must be provided. A template Collaboration Agreement Form is available for download from GEMS. Forms must be completed, signed, dated, and uploaded where indicated on HRB GEMS. Please label each form with the name of the relevant Collaborator. Electronic signatures are acceptable on letters/forms that are uploaded on GEMS.

## Submission of Application

**The deadline for submission of complete application is 26<sup>th</sup> April 2024 at 13:00.**

1. After successful validation the Lead Applicant may submit the application. It will then be routed to the designated signatory at the Host Institution for their approval.
2. If a signatory rejects the application the Lead Applicant will be notified, along with any feedback the signatory has supplied.
3. The application can then be re-submitted; it will be returned to the signatory and will continue through the approval process as before.
4. On completion of the final approval by the Host Institution signatory the application automatically gets submitted to the HRB through GEMS for consideration for funding.
5. Upon submission to the HRB a grant application number is assigned to the application.

**The HRB reserves the right to reject any application that does not meet the terms of this call.**

## Appendix II: HRB Funding Policies and Procedures

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### Access and support from research infrastructures

Applications availing of the advice, research design, data management services and/or other forms of support from a Clinical Research Facility/Centre (CRF/CRC), other infrastructure (e.g., Centre for Applied Medical Imaging, CAMI, Centre for Support and Training in Analysis and Research (CSTAR), HRB-Trials Methodology Research Network (HRB-TMRN), Cancer Groups, National clinical trials office (NCTO), CTNs or a biobank) are required to provide additional information detailing the scope and nature of the engagement (this includes national and international facilities, Units, and networks where justified).

An **Infrastructure Agreement form** will be requested as part of the application addressing the nature/scope of the service or collaboration, the rationale behind the choice of facility/centre/network and any costs associated with the project (including those provided as in-kind contributions). Applications proposing research with patients which do not detail advice and/or support from a CRF/CRC/CTU will be asked to justify why they have not done so.

### Public and Patient Involvement (PPI) in Research

The HRB promotes the active involvement of members of the public and patients in the research that we fund<sup>8</sup>. Public and patient involvement in research means that the public and patients are involved in planning and doing research from start to finish and help tell the public about the results of research. PPI, as defined here, is distinct from and additional to activities which raise awareness, share knowledge, and create a dialogue with the public, and it is also distinct from recruitment of patients/members of the public as participants in research.

PPI represents an active partnership between members of the public and patients and researchers in the research process. This can include, for example, involvement in the selection of research topics, assisting in the design, advising throughout or at specific decision points of the research project or in carrying out the research.

PPI contributors should be actively involved and part of decision making. Involving members of the public in research can improve quality and relevance of research. It can:

- Provide a different perspective – even if you are an expert in your field, your knowledge and experience will be different to the experience of someone who is using the service or living with a health condition.
- Help to ensure that the research uses outcomes that are important to the public.
- Identify a wider set of research topics than if health or social care professionals had worked alone.

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<sup>8</sup> <https://www.hrb.ie/funding/funding-schemes/public-and-patient-involvement-in-research/>

- Make the language and content of information such as questionnaires and information leaflets clear and accessible.
- Help to ensure that the methods proposed for the study are acceptable and sensitive to the situations of potential research participants.
- Help you increase participation in your research by making it more acceptable to potential participants.

In addition to improving relevance and quality of research, it ensures that research is influenced by broader principles of citizenship, accountability, and transparency. PPI is an ethos as well as a practice. It should be context-specific and should aim to ensure that all voices are heard. Where members of the public or patients are involved, they must be compensated for their time and contributions.

**In the application, you are asked to describe any public involvement in your research throughout the various stages of identifying and prioritising the research question, the research design, conduct, analysis, and dissemination. We recognise that the nature and extent of active public involvement is likely to vary depending on the context of each study or award. PPI contributors should be named as Co-applicants where justified by their level of involvement.**

**For guidance and support on PPI in your research please consult with the PPI Ignite Network Ireland or your Host Institution. The PPI Ignite Network Ireland has offices located in the following seven Host Institutions: DCU, NUIG, RCSI, TCD, UCC, UCD, UL.**

## FAIR Data Management and Stewardship

Data management/stewardship plans (DMP) are nowadays widely accepted as part of good research practice. The HRB support [open research](#)<sup>9</sup> and open publishing directly through the [HRB Open Research platform](#)<sup>10</sup>. The HRB is driving the making of research data **FAIR** (Findable, Accessible, Interoperable and Re-usable) in order to benefit science by increasing the re-use of data and by promoting transparency and accountability.

**FAIR data principles**<sup>11</sup> provide a guideline for those wishing to enhance the re-usability of their data holdings: these principles put specific emphasis on enhancing the ability of machines to automatically find and use the data, in addition to supporting its re-use by individuals. For researchers, the move to FAIR and open data, where applicable, means researchers should consider data management issues and find suitable data repositories at the research planning stage. Applicants will have to provide information about their plans for data management and data sharing at application stage.

In line with the HRB's policy on management and sharing of research data<sup>12</sup>, all successful applicants are required to submit a completed data management plan (DMP) to the HRB on or before three

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<sup>9</sup> <http://www.hrb.ie/funding/policies-and-principles/open-research/>

<sup>10</sup> <https://hrbopenresearch.org/>

<sup>11</sup> <https://www.nature.com/articles/sdata201618>

<sup>12</sup> [https://www.hrb.ie/fileadmin/user\\_upload/HRB\\_Policy\\_on\\_sharing\\_of\\_research\\_data.pdf](https://www.hrb.ie/fileadmin/user_upload/HRB_Policy_on_sharing_of_research_data.pdf)



months after the award start date, and a final updated version of the DMP with the last annual report.

The DMP will need to be submitted alongside a certification of completion from the designated representative(s) within the Host Institution e.g. data steward

Applicants will have to provide an outline of their plans for data management and data sharing in the application inclusive of the costs associated to the plan.

The timing for completion and submission of the DMPs must be also included among the objectives and deliverables of the programme. Please submit a draft DMP within the first three months of the grant start date to [DMP@hrb.ie](mailto:DMP@hrb.ie). This should be followed by a final DMP version submitted at the end of the grant.

## General Data Protection Regulation

The **General Data Protection Regulation (GDPR)** came into force on 25 May 2018. As a result, the applicant team will be asked through the HRB online grant management system GEMS to **confirm you understand** that personal data provided as part of this application, including but not limited to CV information, may be shared with person(s) based outside of the European Economic Area (EEA) for the specific purpose of obtaining peer reviews of this application. International reviewers play a vital role for the HRB in setting standards and in benchmarking our scientific community to enable them to operate in a global context. Individual peer reviewers are selected for their specific expertise in relation to submitted applications and can be based anywhere in the world.

Furthermore, by confirming participation, you will be asked to confirm you understand that HRB uses the information you provide (regarding all applicant team members) to consider your application, contact you about your application, and if you are successful, to manage your grant throughout its lifetime in accordance with HRB general T&C for research awards. This will include contacting you with regard to monitoring of progress through written reporting and other means e.g., interim review. We will publish some basic information on successful awards including PI, Host Institution, amount awarded and lay summary on our website and may highlight individual awards or researchers in more detail (with specific consent). We will also use the information you have provided to generate general statistics around our current funding portfolio, and to evaluate our funding mechanisms and investment. After your grant has ended, we will continue to keep your information on file (in accordance with HRB policies) to allow us to evaluate the outcomes, outputs and impacts of HRB investment in your research.

Please note that we will also use information associated with *unsuccessful* applications for a number of the purposes outlined above such as generating general statistics around our current funding portfolio, and to evaluate our funding mechanisms and investment e.g., demographics of applicants, research areas of applicants. Similarly, we will use the information provided about people employed on awards to help evaluate our career support and capacity building initiatives.

## The Health Research Regulations

Following the implementation of GDPR, a regulation for health research known as the Health Research Regulations 2018 (S.I. 314) has been implemented, with further amendments made in 2019 (S.I. 188) and 2021 (S.I. 18)<sup>13</sup>. These regulations outline the mandatory suitable and specific measures for the processing of personal data for the purposes of health research. They further set out that explicit consent is a mandatory safeguard that must be obtained from individuals when using their personal data for health research. Where it is not feasible to obtain explicit consent, an application for a consent declaration can be made to the Health Research Consent Declaration Committee<sup>14</sup>.

## Research on Research

The HRB is developing its approach to research on research (RoR) with the aim of enhancing the evidence base for HRB research funding practices. We may also collaborate with researchers on request regarding specific RoR questions. Should your application become of interest to such a study, the HRB will seek your consent for the use of your information.

## HRB Gender Policy

In line with international best practice, the **HRB Gender Policy**<sup>15</sup> recognises the responsibility of the HRB to support everyone to realise their full potential in order to ensure equality of opportunity and to maximise the quantity and the quality of research. To ensure fairness and equality to all applicants, each funding application received will be assessed as outlined in the call guidance documentation for that particular funding round. To ensure gender balance in decision-making, the HRB aims to reach the international best practice target of 40% of the under-represented gender in all HRB panels where possible. Gender will also be considered when appointing the position of Panel Chair.

## Conflict of Interest

Conflict of interest rules *are applied rigorously*. Where a conflict of interest exists, the reviewer is requested to inform the HRB immediately so that an alternative reviewer may be appointed. International peer reviewers will not provide comments or scores on any application on which they have a conflict of interest.

Reviewers must adhere to high standards of integrity during the peer review process. They must respect the intellectual property of applicants and may not appropriate and use as their own, or disclose to any third party, ideas, concepts, or data contained in the applications they review.

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<sup>13</sup> <http://www.irishstatutebook.ie/eli/2021/si/18/made/en/pdf>

<sup>14</sup> <https://hrcdc.ie/>

<sup>15</sup> <http://www.hrb.ie/funding/funding-schemes/before-you-apply/all-grant-policies/hrb-policy-on-gender-in-research-funding/>

## Appeals Procedure

The HRB's Policy on Appeals on funding decisions is available at <https://www.hrb.ie/funding/funding-schemes/before-you-apply/all-grant-policies/hrb-policy-on-appeals/>.

## Privacy Policy and Retention Policy

To understand why we collect the information we collect and what we do with that information, please see our Privacy<sup>16</sup> and Retention Policies<sup>17</sup>.

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<sup>16</sup> <https://www.hrb.ie/about/legal/privacy-policy/>

<sup>17</sup> [https://www.hrb.ie/fileadmin/user\\_upload/HRB\\_Document\\_retention\\_policy..docx](https://www.hrb.ie/fileadmin/user_upload/HRB_Document_retention_policy..docx)

## **Appendix III: Resources/Useful Links**

### **STUDY DESIGN FOR INTERVENTIONS**

**“Defining Feasibility and Pilot Studies in Preparation for Randomised Controlled Trials: Development of a Conceptual Framework”** by Eldridge S. *et al.*

<http://journals.plos.org/plosone/article?id=10.1371/journal.pone.0150205>

**“The PRECIS-2 tool: designing trials that are fit for purpose”** by Loudon *et al.*

<http://dx.doi.org/10.1136/bmj.h2147>

**“A process for Decision-making after Pilot and feasibility Trials (aDePT): development following a feasibility study of a complex intervention for pelvic organ prolapse”** by Bugge C *et al.*

<http://trialsjournal.biomedcentral.com/articles/10.1186/1745-6215-14-353>

**“Developing and Evaluating Complex Interventions”** by MRC, UK

[www.mrc.ac.uk/complexinterventionsguidance](http://www.mrc.ac.uk/complexinterventionsguidance)

**“Process evaluation of complex interventions: Medical Research Council guidance”** by Moore GF. *et al.*

<http://dx.doi.org/10.1136/bmj.h1258>

**“Using natural experiments to evaluate population health interventions: Guidance for producers and users of research evidence”** by MRC, UK

[www.mrc.ac.uk/naturalexperimentsguidance](http://www.mrc.ac.uk/naturalexperimentsguidance)

**Consort 2010 Statement:** updated guidelines for reporting parallel group randomised trials

[www.consort-statement.org](http://www.consort-statement.org)

**SQUIRE Guidelines:** provides a framework that authors can use when developing applications or writing research articles about quality improvement

[www.squire-statement.org](http://www.squire-statement.org)

**HIQA Guidelines** for the Economic Evaluation of Health Technologies in Ireland (2018)

<https://www.hiqa.ie/reports-and-publications/health-technology-assessment/guidelines-economic-evaluation-health>

**HIQA Guidelines** for the budget Impact Analysis of Health Technologies in Ireland (2015)

[https://www.hiqa.ie/system/files/Guidance\\_on\\_Budget\\_Impact\\_Analysis\\_of\\_Health\\_Technologies\\_in\\_Ireland.pdf](https://www.hiqa.ie/system/files/Guidance_on_Budget_Impact_Analysis_of_Health_Technologies_in_Ireland.pdf)

**HIQA Guidelines for Evaluating the Clinical Effectiveness of Health technologies in Ireland (2011)**

<http://www.hiqa.ie/system/files/HTA-Clinical-Effectiveness-Guidelines.pdf>

## STUDY REGISTRATION

**International Clinical Trials Registration Platform** (run by the WHO)

<http://apps.who.int/trialsearch/Default.aspx>

**European Clinical Trials Database** (EudraCT): database of all regulated clinical trials which commenced in the EU from 1 May 2004

<https://eudract.ema.europa.eu/results-web/>

**US National Library of Medicine database:** database of privately and publicly funded clinical studies – regulated and unregulated – conducted around the world

<https://www.clinicaltrials.gov/>

## REPORTING

**COMET (Core Outcome Measures in Effectiveness Trials) Initiative:** development and application of agreed standardised sets of outcomes, known as ‘core outcome sets’

<http://www.comet-initiative.org/>

**EQUATOR Network Library for health research reporting:** an international initiative that seeks to improve reliability and value of health research literature by promoting transparent and accurate reporting of research studies

<https://www.equator-network.org/library/>

**Registry of Research Data Repositories**

<http://www.re3data.org/>

**Zenodo Data Repository (OpenAIR)**

<https://zenodo.org/about>

<https://zenodo.org/>

## EVIDENCE SYNTHESIS

**Evidence Synthesis Ireland:** aims to build evidence synthesis knowledge, awareness and capacity among the public, health care institutions and policymakers, clinicians, and researchers on the Island of Ireland.

<https://evidencesynthesisireland.ie/>

**The Cochrane Library:** online collection of databases in medicine and other healthcare specialties which summarise and interpret the results of medical research.

[www.thecochranelibrary.com](http://www.thecochranelibrary.com)

**The Campbell Collaboration:** promotes positive social and economic change through the production and use of systematic reviews and other evidence synthesis for evidence-based policy and practice.

<https://www.campbellcollaboration.org/>

**The Campbell Collaboration UK & Ireland:** hub at Queens University Belfast.

<https://www.qub.ac.uk/research-centres/CampbellUKIreland/>

**EQUATOR Network Library for health research reporting:** an international initiative that seeks to improve reliability and value of health research literature by promoting transparent and accurate reporting of research studies.

<http://www.equator-network.org/resource-centre/library-of-health-research-reporting/>

## CLINICAL RESEARCH INFRASTRUCTURES

**All Ireland Hub for Trials Methodology Research**

<http://www.qub.ac.uk/research-centres/CentreforPublicHealth/Research/TheAll-IrelandHubforTrialsMethodologyResearch/>

**Centre for Advanced Medical Imaging, St James' Hospital Dublin**

<http://www.3tcentre.com/>

**Centre for Support and training Analysis and Research (CSTAR)**

<http://www.cstar.ie>

**Children's Clinical Research Unit**

<https://www.nationalchildrensresearchcentre.ie/childrens-clinical-research-unit/apply-for-support/>

**Clinical Research Support Unit, Limerick**

<https://www.ul.ie/hri/clinical-research-support-unit>

**Clinical Research Centre, Royal College of Surgeons in Ireland**

<https://www.rcsi.com/dublin/research-and-innovation/research/resources-and-facilities/clinical-research-centre>

**Clinical Research Facility, University College Dublin**

<http://www.ucd.ie/medicine/ourresearch/researchenvironment/ucdclinicalresearchcentre/>

**Clinical Research Support Centre (Northern Ireland)**

<http://www.crsc.n-i.nhs.uk/>

**HRB Clinical Research Facility, Cork (HRB CRFC)**

<http://www.ucc.ie/en/crfc/>

**HRB Clinical Research Facility, Galway (HRB CRFG)**

[http://www.nuigalway.ie/hrb\\_crfg/](http://www.nuigalway.ie/hrb_crfg/)

**HRB Critical Care Clinical Trials Network (HRB Critical Care CTN)**

[ICC-CTN \(iccctn.org\)](http://www.iccctn.org)

**HRB Irish Network for Children’s Clinical Trials (in4kids)**

[In4kids](#)

**HRB Primary Care Clinical Trials Network (HRB Primary Care CTN)**

[Primary Care Clinical Trials Network Ireland – HRB PC CTNI \(primarycaretrials.ie\)](#)

**HRB Trials Methodology Research Network (TMRN)**

<http://www.hrb-tmrn.ie>

**The National Clinical Trials Office (NCTO)**

Email [trials-ireland@ucc.ie](mailto:trials-ireland@ucc.ie)

<https://ncto.ie/>

**Wellcome Trust-Health Research Board Clinical Research Facility, St James’s Hospital (WT-HRB CRF SJH)**

<http://www.sjhcrf.ie/>

## **BIOBANKING**

**Council of Europe Recommendation of the Committee of Ministers to member States on research on biological materials of human origin (2016)**

[https://search.coe.int/cm/Pages/result\\_details.aspx?ObjectId=090000168064e8ff](https://search.coe.int/cm/Pages/result_details.aspx?ObjectId=090000168064e8ff)

**BBMRI-ERIC is a European research infrastructure for biobanking**

<https://www.bbmri-eric.eu/>

**OECD Guidelines on Human Biobanks and Genetic Research Databases**

<http://www.oecd.org/science/biotech/44054609.pdf>

**ISBER Best Practices for Repositories**

<https://www.isber.org/page/BPR>

**Molecular Medicine Ireland Biobanking Guidelines**

<http://www.molecularmedicineireland.ie/resources/biobanking-guidelines/>

**NCI Best Practices for Biospecimen Resources (2016 version)**

<https://biospecimens.cancer.gov/bestpractices/2016-NCIBestPractices.pdf>

## **PUBLIC AND PATIENT INVOLVEMENT IN RESEARCH & RESEARCH PRIORITIES**

**The National PPI Ignite Network**

<https://ppinetwork.ie/>

**NIHR PPI resources**

<https://www.nihr.ac.uk/documents/ppi-patient-and-public-involvement-resources-for-applicants-to-nihr-research-programmes/23437>

**Patient-Centred Outcomes Research Institute (PCORI)**

<http://www.pcori.org>

**Public Involvement Impact Assessment Framework:** Provides tools for successful involvement of members of the public in research projects and for assessment of impacts.

<http://piiaf.org.uk/>

**NIHR Payment guidance for researchers and professionals**

<https://www.nihr.ac.uk/documents/payment-guidance-for-researchers-and-professionals/27392>

**European Patient Forum Value + Handbook:** For Project Co-ordinators, Leaders and Promoters on Meaningful Patient Involvement.

[http://www.eu-patient.eu/globalassets/projects/valueplus/doc\\_epf\\_handbook.pdf](http://www.eu-patient.eu/globalassets/projects/valueplus/doc_epf_handbook.pdf)

**The James Lind Alliance Priority Setting Partnerships:** Research priorities in disease areas set jointly by patients, clinicians, and researchers.

<http://www.jla.nihr.ac.uk/>

**Campus Engage:** Supporting Irish HEIs to embed civic engagement in their work. Includes resources, how-to-guides, and case studies for engaged research.

<http://www.campusengage.ie/what-we-do/publications/>

**UK Standards for Public Involvement:** The six UK Standards for Public Involvement provide clear, concise statements of effective public involvement against which improvement can be assessed.

<https://sites.google.com/nihr.ac.uk/pi-standards/home>

**The Involvement Matrix:** A tool for researchers/project leaders to promote collaboration with patients in projects and research.

<https://www.kcrutrecht.nl/involvement-matrix/>

**The Evaluation Toolkit:** is a resource designed for practitioners of the health sector, produced after the completion of a rigorous systematic review of patient and public engagement evaluation tools.

<https://ceppp.ca/en/evaluation-toolkit/>

**GRIPP2 reporting checklists:** Tools to improve reporting of patient and public involvement in research

<https://researchinvolvement.biomedcentral.com/articles/10.1186/s40900-017-0062-2#Tab1>

## USE OF ANIMALS IN RESEARCH

**EU Reference Laboratory for alternatives to animal testing (EURL ECVAM)** (reviews of available non animal models)



[https://joint-research-centre.ec.europa.eu/eu-reference-laboratory-alternatives-animal-testing-eurl-ecvam\\_en](https://joint-research-centre.ec.europa.eu/eu-reference-laboratory-alternatives-animal-testing-eurl-ecvam_en)

**Experimental Design Assistant (EDA)** (online tool for design of animal experiments)

<https://eda.nc3rs.org.uk/>

**PREPARE (Planning Research and Experimental Procedures on Animals: Recommendations for Excellence) guidelines**

<https://norecopa.no/prepare>

**ARRIVE (Animal Research: Reporting of In Vivo Experiments) guidelines**

<https://arriveguidelines.org/>

**SYRCLE (Guidance and training on systematic review of animal studies)**

<https://www.syrclenetwork/>

**PROSPERO (Register for systematic reviews including animal studies)**

<https://www.crd.york.ac.uk/PROSPERO/>

## **GENDER AND/OR SEX ISSUES IN RESEARCH**

**Examples of case studies in Health & Medicine where gender/sex in research matters**

<http://genderedinnovations.stanford.edu/case-studies-medicine.html>

**Gender Toolkit in EU-funded research for examples and guidance**

[http://www.yellowwindow.be/genderinresearch/downloads/YW2009\\_GenderToolKit\\_Module1.pdf](http://www.yellowwindow.be/genderinresearch/downloads/YW2009_GenderToolKit_Module1.pdf)

**Sex/Gender Influences in Health and Disease**

<https://orwh.od.nih.gov/sex-gender/sexgender-influences-health-and-disease>

**Methods and Techniques for Integrating Sex into Research**

<https://orwh.od.nih.gov/sex-gender/methods-techniques-integrating-sex-research>

**NIH Policy on Sex as a Biological Variable**

<https://orwh.od.nih.gov/sex-gender/nih-policy-sex-biological-variable>

## **DATA MANAGEMENT AND SHARING AND FAIR PRINCIPLES**

**Digital Curation Centre:** How to develop a data management and sharing plan and examples DMPs.

<http://www.dcc.ac.uk/resources/data-management-plans/guidance-examples>

**FAIR data principles FORCE 11**

<https://www.force11.org/fairprinciples>

**UK Concordat on Open Research Data (July 2016)**

<https://www.ukri.org/wp-content/uploads/2020/10/UKRI-020920-ConcordatonOpenResearchData.pdf>

**Guidelines on FAIR data management plans in Horizon 2020**

[http://ec.europa.eu/research/participants/data/ref/h2020/grants\\_manual/hi/oa\\_pilot/h2020-hi-oa-data-mgt\\_en.pdf](http://ec.europa.eu/research/participants/data/ref/h2020/grants_manual/hi/oa_pilot/h2020-hi-oa-data-mgt_en.pdf)

**FAIR at the Dutch centre for Life sciences**

<https://www.dtls.nl/fair-data/>

**Registry of Research Data Repositories**

<http://www.re3data.org/>

## RESEARCH DATA MANAGEMENT PLANS

**Data Stewardship Wizard created by ELIXIR CZ and NL**

<https://dmp.fairdata.solutions/>

**DMPonline of the Digital Curation Centre (DCC), UK**

<https://dmponline.dcc.ac.uk/>

**DMPTool of University of California Curation Center of the California Digital Library (CDL), USA**

<https://dmptool.org/>

**RDMO Research Data Management Organiser of the German Research Foundation, Germany**

<https://rdmorganiser.github.io/en/>

**Guidelines on FAIR data management plans in Horizon 2020**

[http://ec.europa.eu/research/participants/data/ref/h2020/grants\\_manual/hi/oa\\_pilot/h2020-hi-oa-data-mgt\\_en.pdf](http://ec.europa.eu/research/participants/data/ref/h2020/grants_manual/hi/oa_pilot/h2020-hi-oa-data-mgt_en.pdf)

## KNOWLEDGE TRANSLATION RESOURCES

**Health Service Executive Research & Development Main Page**

<https://hseresearch.ie/research-dissemination-and-translation/>

**Health Service Executive Research & Development: Knowledge Translation, Dissemination, and Impact A Practical Guide for Researchers**

<https://hseresearch.ie/wp-content/uploads/2021/04/Tools-and-templates-.pdf>

**Integrated Knowledge Translation (iKT) NUI Galway**

<https://www.nuigalway.ie/hbcrg/ikt/>

**The Canadian Institutes of Health Research: Guide to Knowledge Translation Planning**

<https://cihr-irsc.gc.ca/e/45321.html>

**Training Institute for Dissemination and Implementation Research in Health:** Open Access Course

<https://cancercontrol.cancer.gov/is/training-education/TIDIRC-open-access>

## IMPLEMENTATION SCIENCE RESOURCES

**Centre for Effective Services**

<https://www.effectiveservices.org/resources/implementation>

**UCC Implementation Science Training Institute**

<https://www.ucc.ie/en/cpd/options/medhealth/cpd1778uccimplementationsciencetraininginstitute/>

**European Implementation Collaborative**

<https://implementation.eu/resources/>

## CO-CREATION RESOURCES

**ACCOMPLISSH Guide to impact planning**

<https://www.accomplish.eu/publications-and-deliverables>

**Working together to co-create knowledge: A unique co-creation tool – Carnegie UK Trust**

<https://www.carnegieuktrust.org.uk/publications/working-together-to-co-create-knowledge-a-unique-co-creation-tool/>

## INFORMATION ON PERSISTENT IDENTIFIERS

**DOI:** List of current DOI registration agencies provided by the International DOI Foundation

[http://www.doi.org/registration\\_agencies.html](http://www.doi.org/registration_agencies.html)

**Handle:** Assigning, managing and resolving persistent identifiers for digital objects and other Internet resources provided by the Corporation for National Research Initiatives (CNRI)

<http://www.handle.net/>

**PURL:** Persistent Identifiers developed by the Online Computer Library Center (OCLC). Since 2016 hosted by the Internet Archive

<https://archive.org/services/purl/>

**URN:** List of all registered namespaces provided by the Internet Assigned Numbers Authority (IANA)

<https://www.iana.org/assignments/urn-namespaces/urn-namespaces.xml>

## DATA REPOSITORIES

**Registry of Research Data Repositories**

<http://www.re3data.org/>

**Data centers accredited by the German Data forum according to uniform and transparent standards (Germany)**

<https://www.ratswd.de/forschungsdaten/fdz>

**Zenodo Data Repository (OpenAIR)**

<https://zenodo.org/>

## **OTHER USEFUL LINKS**

**Tool that helps to select and apply a license to a resource, provided by Creative Commons**

<https://creativecommons.org/choose/>

## Definitions

### Data controller

A '**controller**' refers to a person, company, or other body that decides how and why a data subject's personal data are processed. If two or more persons or entities decide how and why personal data are processed, they may be 'joint controllers', and they would both share responsibility for the data processing obligations<sup>18</sup>. Data Controllers from the provider organisation should be named as Co-applicants where justified by their level of involvement. Otherwise, they should be named as Collaborators.

### Knowledge user

A **knowledge user** is defined as one in a position of authority to influence and/or make decisions about health policy or the delivery of services and can act to ensure that the findings of the research will be translated to influence decision making and change within their (or other) organisations. This is typically managers, policy makers, clinicians, health professionals or others who are in a position to make significant changes to policy or practice. Knowledge user organisations may be Government departments, HSE, other agencies, hospitals or hospital groups, community healthcare organisations, local government, voluntary organisations, research charities, patient/consumer groups or other organisations involved in making decisions regarding the management, structuring and/or delivery of practice or policy in the Irish health and social care system.

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<sup>18</sup> <https://www.dataprotection.ie/sites/default/files/uploads/2019-07/190710%20Data%20Protection%20Basics.pdf>