## Application summary

**Application title**
This is the title of your proposed project.

**Proposed duration of funding (months)**
This is normally 60 months (FTE). The Fellow's salary and all research costs must end at the same time.

**Proposed start date**
This date must be at least six months after the full application deadline. You should also take into account the date of the Sir Henry Dale Fellowship Interview Committee.

*You can change your start date if your application is successful. All grant expenditure and activities must be within the grant start and end dates.*

**Name of administering organisation**
If your application is successful, this is the organisation that will be responsible for administering the award.

**Lead applicant's address at administering organisation**
If your application is successful, we will use this address in your award letter.

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**Research area**
Select the most relevant area, based on the key aims of the research. This allocates your application to the relevant Grants team. We may reallocate your application to another area if we consider it appropriate.

## Lead applicant
Lead applicant details

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<th>Full Name</th>
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Career history (current/most recent first)

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<th>From</th>
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<th>Position</th>
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Education/training

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<th>From</th>
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Source(s) of personal salary support
State all your sources of salary funding (for example, through your organisation’s block grant from a higher education funding body), and the percentage of your salary they contribute. Answer 'not applicable' if you are not currently employed.

Your source of salary may affect your eligibility - please check the relevant scheme webpage.

If your source of salary places any restrictions on intellectual property rights or publications arising from your research, contact us as this may also affect your eligibility.

Current/last appropriate salary details

If you are currently unemployed give salary details from your most recent employment.

Basic salary (per annum)

Currency
<table>
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<th><strong>Clinical status</strong></th>
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<tr>
<td>Do you have a medical, veterinary, dental or clinical psychology degree?</td>
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<td>Specify</td>
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<th><strong>Are you clinically active?</strong></th>
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<th><strong>What is your specialty?</strong></th>
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<td>If your specialty is not on the list, select ‘Other’ and specify.</td>
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<td>Please specify</td>
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<th><strong>Career breaks</strong></th>
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<tr>
<td>Have you taken any career breaks or periods of part-time work, for example parental, long-term sick leave, carer responsibilities?</td>
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<td>We encourage applications from researchers who have taken career breaks. We want to ensure that any such breaks are taken into account when we consider your track record. State when and for what period you took a break, or were working part-time. We are not asking for the reasons for this break so please do not provide these here, including sharing any sensitive personal health information.</td>
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<td>Provide details.</td>
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<th><strong>Do you wish to undertake this award part time?</strong></th>
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<tr>
<td>If you wish to undertake this award part-time, either from the start or part way through the grant, your host organisation must employ you on a part-time basis during that time.</td>
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<td>We provide flexible research career opportunities. If you’re applying for funding, you can request flexible and part-time working. This could be to help you manage family commitments or if you have individual needs which make undertaking an award full time challenging.</td>
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<tr>
<td>Career contributions</td>
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<td>What are your most important research-related contributions? These may be from any stage of your research career. State what each contribution was, when it came about, why you think it is important and what impact it has had. Examples include publications, patents and impacts on policy. (350 words max.)</td>
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| **Personal statement** |
How will this Fellowship further your research and career aspirations? (300 words max.)

Research environment
What scientific considerations led you to choose this research environment and sponsor for your research? If you have already been based in this research environment for two years or more, you must provide a strong justification for why you are proposing to stay and how you will demonstrate your independence. (400 words max.)

Research outputs
List up to 20 of your most significant research outputs; at least five of these must be from the last five years. For 10 of these outputs, provide a statement describing their significance and your contribution (up to 50 words per output).

Research outputs may include (but are not limited to):
- Peer-reviewed publications and preprints;
-Datasets, software and research materials;
-Inventions, patents and commercial activity.

For original research publications, indicate those arising from the Funders’ grants in bold (indicating whether Royal Society or Wellcome Trust), and provide the PubMed Central ID (PMCID) reference for each of these. You can find more information on this in the guidance to this question.

Give the citation in full, including the title of paper and all authors (unless more than 10, in which case you may use 'et al', ensuring that your position as author remains clear). Citations to preprints must state "Preprint", the repository name and the articles persistent identifier (e.g. DOI).

Include here systematic reviews (e.g. Cochrane Reviews) and meta analyses, but exclude abstracts and literature reviews. We encourage you to include articles published via open research publishing platforms, such as Wellcome Open Research, providing they have passed peer review.

Only include preprints, complete manuscripts that have been submitted to a preprint repository or service (for example, bioRxiv, PeerJ Preprints, arXiv, SocArXiv or PsyArXiv), if they have a permanent identifier such as a DOI or arXiv identifier. Our open access policy requires all original peer-reviewed research papers, supported in whole or in part by our funding, to be made available through PubMed Central (PMC) and Europe PMC as soon as possible and in any event within six months of the journal publisher’s official date of final publication.

The PubMed Central ID (PMCID) is the unique identifier assigned to every full text paper in PubMed Central (PMC) and Europe PMC.

We actively monitor compliance with our open access policy and we ask successful applicants to provide a full list of all their Wellcome-funded research papers, and confirm compliance by providing the PMCID identifier for these, before the award letter can be issued. You can find further guidance in our open access policy statement and authors’ information.

How many peer-reviewed publications have you authored/co-authored? Include systematic reviews and meta analyses but exclude abstracts and literature reviews.

We encourage you to include articles published on open research publishing platforms, such as Wellcome Open Research, providing they have passed peer review.
Recommendation by applicant’s present Head of Department or supervisor

Upload a letter of support from your current Head of Department or supervisor (500 words maximum).

Your current Head of Department or supervisor should give an assessment of your calibre. If they feel that they are not in a position to make an assessment (for example, if you have only recently joined the department), then you can ask an alternative. This should be someone you have worked with in the past twelve months.

The letter of recommendation should be uploaded to the system and show clearly the head’s (or supervisor’s) name, position and address.

### Sponsors

**Primary sponsor**

A sponsor should be able to reassure the Funders that the Fellow will be welcomed into the host department as an independent researcher.

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The primary sponsor must be based at the administering organisation.

**Title of current post**

**Source(s) of personal salary support**

State all your sources of salary funding (for example, through your organisation’s block grant from a higher education funding body), and the percentage of your salary they contribute.

**Recommendation**

Upload your letter of recommendation (500 words maximum).

The sponsor should give an assessment of the calibre of the applicant and an overview of how he/she would complement the on-going activities of the host environment. The sponsor should guarantee in this letter that the applicant will be supported in developing an independent research career and ensure that the Fellow be granted equal status to
other academic staff of similar seniority.

Are further sponsors required for your application? If there is more than one sponsor for the proposal, each must provide their details and answer the related questions.

You should normally only identify an additional sponsor if you propose to undertake fieldwork or research in a low- or middle-income country. In these cases, you should identify an appropriate Head of Department or equivalent (for example, Director of Wellcome Trust Major Overseas Programme) in your overseas organisation.

Additional sponsors

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Title of current post

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State all your sources of salary funding (for example, through your organisation’s block grant from a higher education funding body), and the percentage of your salary they contribute.

If your source of salary places any ties on intellectual property rights or restrictions on publications arising from your research, you must contact us as this may affect your eligibility.

Collaborators

Are any collaborations essential for this proposal? This could be through sharing facilities, providing access to resources (essential reagents, samples, data) or sharing subject-specific knowledge and guidance.

If the answer is “Yes”, you will be asked to provide information about these collaborators and to confirm their willingness to participate in the proposed research.
List any key collaborators (name and organisation) and provide a very brief outline of their role in the proposed research.

You can replace the collaborators named here with suitable alternatives if it is necessary or appropriate to do so.

I confirm that the collaborators named above have agreed to be involved, as described, in the proposed research and are willing for their details to be included as part of this application.

Related applications

Is this or a similar application for funding currently under consideration elsewhere?

We'll consider your application even if you have a similar application being considered by another funder. If the other funder offers you funding, please tell us immediately. We will usually ask you to decide on that offer within one month.

If you decide to apply to another funder with a similar application after you have applied to us, please also let us know.

Provide the name(s) of the funder(s) and the expected decision date(s).

Is this a resubmission of an application submitted to the Wellcome Trust or Royal Society within the last 24 months?

Contact us before resubmitting an application.

By whom and when was it originally considered?

How is this application different?
(200 words max.)

Research summary

Research summary
Provide a summary of your proposed research, including key goals, for an expert audience.
(200 words max.)
The summary should be as complete as possible within the word limit. Include key words that best describe the proposal to enable text searching.

We will use this as a short abstract and to classify your proposal by subject. We may edit your summary and then use it to describe your research on our website and elsewhere (we publish summary details of all our awards).

Lay summary
Provide a summary of your proposed research for a non-specialist audience. You don’t need to oversimplify your research, but try to explain it as clearly as possible. Write in the first person (“I” and “we”) and structure your summary in this order: background to the research problem, your approach and expected impact of your work.

We may edit your summary and then use it to describe your research on our website and elsewhere (we publish summary details of all our awards).

Example of a lay summary:
In response to stress and environmental changes, bacteria generate hundreds of small RNA molecules that have key roles in regulating gene expression. This process of riboregulation involves chaperone proteins that facilitate the actions of regulatory RNAs, and enzymes that affect RNA transcript lifetimes. I aim to understand the molecular basis of riboregulation by taking a multidisciplinary approach. I will use biochemical and structural analyses, including cryoEM and cryoET, to visualise how RNA transcripts are captured and channelled to active sites, either for degradation or processing. I will also identify the RNA targets of chaperone proteins and the degradative machinery, and explore whether the patterns change with physiological state or during the cell cycle, and why. These studies will help to explain how small RNAs enhance the speed and accuracy of bacterial genetic regulation, enriching the capacity of the simplest organisms to exhibit complex behaviour.

Details of research project
Describe your research project. This must include:
(a) Aims and research questions;
(b) Work which has led up to the project;
(c) Approach and methods you will use;
(d) Timetable and milestones, if appropriate.

You should not use more than 2,800 words.

Provide all relevant information within the application form; do not refer to additional unpublished information on personal websites. If you do not understand any part of this guidance, contact us for advice.

The word count must not exceed 2,800 words in total, excluding graphs, figures. You may provide your answer to this question in text entry format or as a PDF attachment. If you are uploading your research vision, the uploaded document must be in 11 point Arial font and portrait format.

If you plan to work in more than one department during the Fellowship (for example, spending a period abroad) you should make clear here which parts of the project you will carry out in each location.

Research questions
State the key question(s) that is/are being addressed by your proposal. If your research is not driven by an underlying hypothesis, state the impact of the proposed studies.

Approach and methods to be used
Details of experimental design for animal studies should be provided as part of the justification for animals in the 'Proposals involving animals' section of the form.

For epidemiological, demographic, case control, cohort and related studies, give a detailed analysis of the study design, including details of any validation already undertaken or rationale for using standard protocols. You should give particular attention to power calculations, sample size justification and, where appropriate, case definition and inclusion/exclusion criteria.
If you are requesting support for a clinical trial, provide full details, including study design, in the clinical trial section of the form.

**Additional information**
You may provide up to the equivalent of two A4 pages of additional information. Embed it in your upload for your research vision or upload it under 'Additional information'. If you choose to embed this information, any text present (such as legends, labels, or captions) can be excluded from the word count. If the embedded information exceeds two pages of A4 we will return your application to you to reduce the amount of information.

This form asks for all the information we require to consider your application. Do not provide additional information, such as letters of support, unless specifically requested in the form.

(2800 words max.)

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<th>Does your proposal involve human participants?</th>
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<th>Does your proposal involve a clinical trial?</th>
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<td>Yes</td>
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**Wellcome adheres to the World Health Organization definition of a clinical trial as: “any research study that prospectively assigns human participants or groups of humans to one or more health-related interventions to evaluate the effects on health outcomes. Interventions include, but are not restricted to, drugs, cells and other biological products, surgical procedures, devices, behavioural treatments, process-of-care changes, preventive care, etc.”**

You can find further information in our clinical trials policy statement.

If your proposal involves a clinical trial, you should provide details of it within this application form. If your proposal involves more than one clinical trial, you should contact us for guidance.

**Details of studies involving human participants including clinical trials**

Describe the study design. This should include, as applicable:
- number of participants in each group;
- type, frequency and duration of interventions and/or health outcome measures;
- frequency and duration of planned follow up;
- any other activity with potential significant risks to participants;
- details of any investigational product, focusing on manufacture, quality and consistency.

(300 words max.)

**Types of health outcomes or interventions can include but are not restricted to:**
- screening procedures
- collection of biological samples
- biometric and clinical data
- experimental challenges
- behavioural treatments
- process-of-care-changes

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<th>What are the primary and secondary outcome measures, and how will you assess these?</th>
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Outline the strategy for recruitment and describe the inclusion/exclusion criteria for study participants (if applicable). What are the proposed arrangements for allocating participants to study groups?
Detail and justify the power calculation, sample size and proposed statistical analysis, including any interim analyses and/or subgroup analyses. What are the proposed methods for protecting against sources of bias?
(200 words max.)

Describe the supporting personnel and infrastructure that will be used to deliver the proposed research (for example key support staff, roles of team members, participating centre(s) or facilities). Detail any activity a third party is undertaking, and explain what agreements or formal contracts will be in place.
(200 words max.)

How have you involved patients, participants, patient advocacy groups or communities in developing this proposal?
(200 words max.)

You may submit up to two A4 pages of additional information (such as graphs, figures, tables and essential unpublished data).

The additional information (such as graphs, figures, tables and essential unpublished data) provided in support of the research proposal may be embedded in the text of your file upload or attached here as a separate file. If you choose to embed this information, any text present (such as legends, labels or captions) can be excluded from the word count.

Please note that this form asks for all the information we require to consider your application. You should not provide additional information (e.g. letters of support) unless specifically requested in the form.

References
You should give the citation in full, including title of paper and all authors.

You may provide up to the equivalent of two A4 pages of references. Please ensure that all references included are pertinent to your research proposal and are cited in full, including all authors, the full title of each publication, journal title, year, volume and pages. Citations to preprints should state “Preprint”, the repository name and the article persistent identifier (e.g DOI).

References with more than 10 authors may be shortened to et al, but please ensure that your position as author (if applicable) remains clear.

Have you listed any "in press" papers in your References section that you want to submit to us?

Upload papers "in press" as a single PDF
Will the proposed research generate outputs of data, software, materials or intellectual property that hold significant value as a resource for the wider research community?

Our Data, Software and Materials Management and Sharing Policy states that all Wellcome-funded researchers must manage their research outputs in a way that will achieve the greatest health benefit, maximising the availability of research data, software and materials with as few restrictions as possible. If your proposed research is likely to generate significant outputs - data, software, materials and/or intellectual property - that will hold clear value as a resource for others in academia or industry, you are required to provide an outputs management plan.

Select the approach you will use to maximise the impact of your significant research outputs to improve health and benefit the wider research community.

Provide an outputs management plan. Our guidance on developing an outputs management plan sets out where such a plan is required and gives an overview of what you must consider. (700 words max.)

Your plan must be clear, concise, proportionate and focus specifically on how you will identify outputs, manage these and then use them to advance potential health benefits. Set out and address clearly the following:

1) For significant data, software and materials outputs
   (i) What significant outputs will your research generate?
   (ii) When do you intend to share these outputs?
   (iii) Where will you make these outputs available?
   (iv) How will they be discovered and accessed by others?
   (v) Are limits on sharing required?
   (vi) How will these outputs be preserved?

2) For intellectual property outputs
   (i) What IP will your research generate?
   (ii) How will you protect this IP?
   (iii) How will the IP be used to achieve health benefits?
   (iv) Provide the name and contact details for the person in your organisation (e.g., Technology Transfer Officer or Business Development executive) who can act as a point of contact for Wellcome in connection with the protection and commercialisation of this IP.

3) Describe any resources that you will need to deliver your outputs management plan. Regardless of whether or not a plan is required, you must ensure that data and original software underlying published research findings are accessible at the point of publication.

Public engagement

Will you engage with the non-academic public about your work?

We want to foster a culture that values, recognises and better supports public engagement. We encourage researchers we fund to inform, consult and collaborate with the non-academic public. Further information is available on the Wellcome Trust website and the Royal Society website.

Provide a brief outline of your plan to engage with the non-academic public (beyond press and media activity). Do not include engagement that is essential for the ethical conduct of your research, such as patient information leaflets or community advisory boards. This should be part of your research methodology and you can include costs for this within your main research costs.
We may provide additional support during the lifetime of the research grant, with a focus on developing the researcher’s practice in Public Engagement, Diversity & Inclusion, or Open Research through our Research Enrichment scheme. Further details on the scheme, including how to apply, are available on our website.

Work abroad

Are you intending to work abroad during your fellowship?

If you are requesting support for research abroad you will need a sponsor both in your host organisation (the administering organisation) and in the overseas organisation or research facility.

You may normally spend up to 12 months of the Fellowship outside your host organisation during the period of award.

You must make clear in the description of your research proposal (‘Details of research project’) which parts of the project are to be carried out in each research facility.

Work abroad

1

Name of host overseas organisation

Address of host overseas organisation

Dates of travel and duration of trip(s)
Specify weeks, months.

Purpose of the visit(s)
(100 words max.)

Upload a letter of recommendation from your sponsor at the host overseas organisation (300 words maximum).

The letter should explain why the visit is essential for the proposed research, and in developing the skills and career aspirations of the applicant.

The letter of recommendation should show clearly the name, position and address of the sponsor.

Location of activity

Will the funded activity take place at more than one location?
List any locations outside of your administering organisation where you will be
conducting research or redirecting funds. This includes, but is not limited to, anywhere in receipt of indirect funding, fieldwork sites, and time spent working in another organisation/laboratory. This does not include conference attendance.

For each location, select the country and, where applicable, state the organisation. You must include the administering organisation.

Enter the approximate percentage of the total funds that will be spent in each location. Enter zero for locations where activity will take place but no significant funds will be spent. If you are requesting salary costs, attribute them to the employing organisation.

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<th>Country</th>
<th>Organisation</th>
<th>Percentage of funds</th>
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Costs requested and justification

Salaries
Are you requesting salaries?
For details of what staff costs you can request, refer to the relevant scheme page and check the guidance notes for this question.

Detail salaries requested for all staff, including the applicant, to be funded on the grant.

The host organisation will determine the appropriate basic salary for the applicant, in line with the skills, responsibilities and expertise necessary to carry out the role. The total costs of the post are to be detailed here. These costs should include the applicant's basic salary, employer's contributions, incremental progression and any locally-recognised allowances (e.g. London allowance), as applicable. Any previous Wellcome Trust supplement paid to an individual should also be excluded.

Figures for Year 1 may include known pay awards that will take place during the first year (or an assumed percentage, equivalent to our current inflation rate, as set out on the scheme webpage, where the scheduled pay award has not yet been confirmed). Inflation for Year 2 onwards will be based on our current inflation allowance rates and should not be factored into the requested salary.

Fellows are not expected to undertake more than 6 hours of non-research activities per week.

Provide the names of individuals for posts involving the handling of, and research on, non-human primates. Whilst your application is being considered, you must notify the Funders of any change to the individual(s) named in the application.

Definition of terms

Staff category: For example: Postgraduate research assistant; Postdoctoral research assistant; Technician; Fieldworker. Specify the level of seniority of the post where relevant, e.g. Junior postdoctoral research assistant, Senior postdoctoral research assistant.

Salary grade/scale: The national or local salary grade/scale on which the individual will be employed.

Basic starting salary: Annual salary to be paid to the individual upon their appointment to the post, exclusive of any allowances for which the individual is eligible. If the post is part time, the annual salary must be quoted on a pro rata basis.

Total cost on grant: Total cost of the post, inclusive of any locally-recognised allowances (for example, London allowance), employer's contributions and increments, over the period of the grant. Employer's contributions should include any statutory obligations (e.g. for the UK, National Insurance contributions) and contributions towards an organisational pension scheme.

Salaries / Stipends
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<th>Staff category</th>
<th>Name (if known)</th>
<th>Basic starting salary (p.a.)</th>
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</table>

**Materials and consumables**
Are you requesting materials and consumables?

**Materials and consumables**

<table>
<thead>
<tr>
<th>Description</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
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</tbody>
</table>

**Animals**
Are you requesting animals?

In order to ensure animal experimentation costs are accurate, you must complete this section after consultation with your animal house or biological services manager. Your organisation must apply a consistent costing methodology when presenting cost details.

If appropriate, costings can be clarified under ‘Justification for resources requested’.

We may ask for more detailed costing information where a large number of animals and/or substantial costs are involved.

<table>
<thead>
<tr>
<th>Animal species</th>
<th>Total no. to be purchased</th>
<th>Total purchase cost</th>
<th>Total maintenance and procedures cost</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
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</table>

**Associated animals costs**

<table>
<thead>
<tr>
<th>Description</th>
<th>Total</th>
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<td></td>
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</tbody>
</table>

These costs cover specific and relevant training and environmental enrichment, including training for animal husbandry, welfare and associated training for animal technicians, and the cost of animal licences.

**Equipment**
Are you requesting equipment or equipment maintenance?

The organisation’s Director of Procurement/Head of Purchasing (or equivalent) must be aware of all potential capital purchases and we require organisations to use best procurement practice when purchasing equipment with Wellcome Trust funds.

**Equipment to be purchased**

We expect you to consider the cost-effectiveness of the proposed purchase of equipment. The estimated price of the equipment must cover all aspects including delivery, installation, maintenance and training, where appropriate. We expect discounts to be negotiated and included in quoted prices.

If there is a preferred manufacturer for certain items of equipment, you can explain this in the ‘Type of equipment’ field.

We expect that the equipment you request will be covered by the manufacturer’s warranty for the first year after it is purchased. We will fund reasonable maintenance costs for four years after the initial period of warranty on all equipment (irrespective of the length of award made), where this is negotiated as part of the capital purchase cost. We will also
consider costs for the maintenance of equipment over 5 years old if you can demonstrate that it is cost-effective.

Value Added Tax (VAT)
For grants to be held in the UK, the costs of all equipment to be used for medical and veterinary research must be quoted exclusive of VAT. For equipment that does not fall within this definition, VAT costs should be shown.

Equipment

<table>
<thead>
<tr>
<th>Type of equipment</th>
<th>No. of items</th>
<th>Cost per item</th>
<th>Cost of maintenance contract</th>
<th>Contribution from other sources</th>
<th>Total</th>
</tr>
</thead>
</table>

Maintenance for existing equipment

<table>
<thead>
<tr>
<th>Details of equipment/ facility</th>
<th>Wellcome Trust grant number</th>
<th>Date of purchase</th>
<th>End date of current contract</th>
<th>Total cost of contract</th>
<th>% time on project</th>
<th>Total</th>
</tr>
</thead>
</table>

We consider requests for maintenance of existing equipment if the grant that funded its purchase has ended. We only provide maintenance costs for equipment more than five years old if it is cost-effective to keep maintaining it.

- Are you requesting a piece of equipment with a list price of £100,000 or more?

- Please upload a copy of at least one formal quote
  
  *If there is more than one quote, please submit these as a single PDF.*

Synchrotron radiation sources

Will you need access to a synchrotron source?

We collect data on access to synchrotron sources for information purposes.

Apply directly to the synchrotron facility you want to use.

Which source(s) will you be applying to? (Select all that apply)

Specify:

Are you requesting costs from the Funders?

These facilities are normally free at the point of access for researchers who are prepared to publish their results in the public domain. If this is not the case for the facility that you have selected, please provide details of the costs in the ‘Access charges’ section.

In instances where the costs of travel and subsistence will not be met by the facility, they may be requested from us. Please provide details of these costs in the ‘Travel and subsistence’ costs section.

Access charges
Are you requesting access charges?

If the equipment/facility was funded by the Funders or other sources, you can only request access charges if the original grant has ended.

The use of the equipment/facilities must be essential to the proposed research.

Please refer to scheme webpage for information on allowable access charges.

### Access charges

<table>
<thead>
<tr>
<th>Details of equipment/facility</th>
<th>Original source of funding</th>
<th>Wellcome Trust grant number, if applicable</th>
<th>Standard access charge per unit</th>
<th>Specify unit</th>
<th>No. of units to be used for this project</th>
<th>Total</th>
</tr>
</thead>
</table>

### Travel and subsistence

Are you requesting travel and subsistence?

Include here conference attendance and collaborative visits. Where necessary, please state the host organisation.

**Conference attendance**

The lead applicant and any research staff to be employed on the grant can request costs to attend academic/scientific conferences, including conference registration fees, up to the maximum annual amount specified on the scheme page. Specify the amount being requested per person.

**Collaborative visits**

If you are requesting costs for collaborative visits, state the host organisation and provide a detailed breakdown of the travel and subsistence costs. Justify the need for the visit, and its duration, in your application.

### Travel and subsistence

<table>
<thead>
<tr>
<th>Description</th>
<th>Total</th>
</tr>
</thead>
</table>

### Miscellaneous costs

Are you requesting miscellaneous costs?

We require a detailed breakdown of the miscellaneous costs requested. Costs that do not fall under any other category should be entered in this section. These may fall under specific subheadings (such as ‘Overseas allowances’ and ‘Provision for Public Engagement’); where they do not, please select ‘Other’ and type a description of the item.

**Working abroad**

If costs are requested for the applicant(s) and/or research staff to be employed on the grant to carry out any of the proposed research abroad, please state the overseas host organisation, and detail the travel costs and other overseas allowances. Allowances should be itemised (e.g. “baggage/freight”, “medical insurance”). Further guidance can be found on the scheme webpage.

**Personal removal expenses**

We will consider providing a contribution towards your personal removal expenses if you will be relocating to take up the award. For further information on the amount that can be requested, please see the scheme webpage. A justification for the expenses must be provided in the application, together with an estimate of the costs.

### Miscellaneous other
### Justification for resources requested

Justify these costs under each of the above headings (for example, "Salaries"; "Equipment"; "Miscellaneous").
(1000 words max.)

Before completing this section, check the Sir Henry Dale Fellowship scheme webpage on our website for more information on costs we will cover and disallowed costs.

For staffing, justify the type and seniority of each post sought. If you are requesting funds for staff based in different locations, tell us where they will be working.

Explain the need for any collaborative/overseas visits and their duration.

Justify all animal and animal associated costs. You do not need to justify animal numbers required here; you can include this in the 'Proposals involving animals' section.

If you are requesting equipment which costs more than £100,000, provide details of:
- similar equipment in your department and adjacent departments
- the reasons why it cannot be used for this particular project
- any other individuals likely to benefit from use of the equipment.

### Summary of financial support requested

<table>
<thead>
<tr>
<th>Type</th>
<th>Description</th>
<th>Total</th>
</tr>
</thead>
</table>

### Full economic costing

Is your organisation calculating the full economic cost of this proposal?

What is the total full economic cost of your research proposal (£)?
Include inflation in your costs at the percentage rate currently used by your administering organisation.

### Research involving human participants, human biological material and identifiable data

Does your project involve human participants, human biological material, or identifiable/potentially identifiable data?

The following notes relating to 'Research involving human participants, human biological material and identifiable data’ are intended to provide guidance and advice in completing the form, rather than a comprehensive review of the legal and regulatory environment in which the application is made.

The World Health Organization defines research with human subjects as “any social science, biomedical, behavioural, or epidemiological activity that entails systematic collection or analysis of data with the intent to generate new knowledge, in which human beings i) are exposed to manipulation, intervention, observation, or other interaction with investigators either directly or through alteration of their environment, or ii) become individually identifiable through investigator’s collection, preparation, or use of biological material or medical or other records.”
Our policy position on research involving human participants can be found on our website.

We require ethical approval (usually from the appropriate National Health Service (NHS) research ethics committees) for all research we fund involving human participants, biological samples or personal data. Personal data (as defined in the Data Protection Act 2018) is any information relating to an identified or identifiable living person i.e. a person who can be identified either directly from that information or indirectly by combining it with other available information. Any use of personal data or biological samples, relating to living or dead persons, should conform to MRC guidelines available at: http://www.highlights.rsc.mrc.ac.uk/PIHR/index.html#?_k=opxohv and https://mrc.ukri.org/publications/browse/human-tissue-and-biological-samples-for-use-in-research.

You should seek approval from other regulatory bodies such as the Human Fertilisation and Embryology Authority (HFEA) or the Gene Therapy Advisory Committee in the UK where necessary, e.g. research involving human embryos may require a licence from the HFEA (please refer to www.hfeagov.uk for more information). If your proposal involves research on gene therapy which requires regulatory approval, you should seek approval from your Local Research Ethics Committee, the University’s Genetic Manipulation Committee, the Gene Therapy Advisory Committee and the Medicines and Healthcare products Regulatory Agency (MHRA).

The organisation must ensure that ethical approval is in place at all relevant times during the project. For research carried out at multiple sites, ethics committee approval must cover each site.

Where the research, or part of the research, is to be performed outside the UK, independent ethics review must be obtained. For research involving people living in low and middle income countries, see the Wellcome Trust’s website.

Confirm that you have read our guidance on the feedback of health-related findings in research (available on our website) and that you are in the process of considering your approach to this.

Who has, or will, review the ethics of the project and when? Detail any other regulatory approvals you have obtained, or will seek.
We reserve the right to see relevant approval documents at any point during the lifetime of the grant, in accordance with our policy position on research involving human participants.

Is the proposed clinical trial covered by The Medicines for Human Use (Clinical Trials) Regulations in the UK?

Under the Medicines for Human Use (Clinical Trials) Regulations 2004, applicants must identify a sponsor (which will normally be either a university or NHS Trust) who fully understands the responsibilities and costs associated with assuming this role. Please note that the Wellcome Trust cannot act as sponsor.

Please confirm that the trial will be registered on the International Standard Randomised Controlled Trial Number Register (ISRCTN), ClinicalTrials.gov, or on another register listed on the WHO International Clinical Trials Registry Platform (ICTRP).

Do you propose to use facilities within the National Health Service (NHS) in the UK or to involve patients being cared for by the NHS?

By agreeing to fund work which requires NHS support, the Wellcome Trust is agreeing to abide by the Statement of Partnership on Non-commercial R&D in the NHS in England (and the corresponding statements in Northern Ireland, Scotland, and Wales). Researchers must therefore meet the obligations of the Partnership and may not carry out any research until the NHS has given its consent.

The Research Governance Framework for Health & Social Care, published by the Department of Health in England can be downloaded from the Department of Health website http://www.dh.gov.uk/health/category/research. The Wellcome Trust cannot act as sponsor.
<table>
<thead>
<tr>
<th>Do you propose to use NHS England facilities, staff or patients?</th>
</tr>
</thead>
<tbody>
<tr>
<td>Have you completed a Schedule of Cost Attribution Tool?</td>
</tr>
<tr>
<td>This must be signed off by a local Clinical Research Network AcoRD specialist. Download a template SoECAT here. See our webpage on Excess Treatment Costs for more information.</td>
</tr>
<tr>
<td>Explain why you have been unable to complete a Schedule of Cost Attribution Tool. During NHSE’s pilot phase (October 2018 – April 2019) you can submit your SoECAT whilst we are reviewing your application but Wellcome cannot make a funding decision without it. If you do not have a signed off SoECAT form your research will not receive HRA approval. (100 words max.)</td>
</tr>
<tr>
<td>Is a formal sponsor required for the project, for example under the Medicines for Human Use (Clinical Trials) Regulations or the Research Governance Framework for Health and Social Care and equivalent guidance?</td>
</tr>
<tr>
<td>Please indicate which organisation(s) has/have agreed to fulfil this role. Please note that the Wellcome Trust cannot act as sponsor.</td>
</tr>
<tr>
<td>Confirm you have in place or you will seek appropriate informed consent to use any potentially commercially exploitable results from tissues or samples derived from human participants.</td>
</tr>
<tr>
<td>Please answer ‘not applicable’ if no potentially commercially exploitable results (based on human tissues or samples) will be produced during the course of the proposed research.</td>
</tr>
<tr>
<td>Describe the oversight arrangements for the clinical trial (e.g. membership of Trial Steering Committee, Data Monitoring Board etc.)</td>
</tr>
</tbody>
</table>

**Proposals involving animals**

Select any of the following that apply to your proposed work: *(Proposal involves the use of animals, Proposal involves the use of animal tissue, Neither of the above)*

The following notes relating to ‘Proposals involving animals’ are intended to provide guidance and advice in completing the form, rather than a comprehensive review of the legal and regulatory environment in which the application is made.

Applicants must refer to the Funders’ policy on the use of animals in medical and veterinary research.

In all animal experiments we support, the principles of reduction, replacement and refinement will apply. In all experimental studies, applicants must actively consider:
Refined methods in animal research are those which alleviate or minimise any adverse effects for the animals involved, and/or enhance animal welfare. Refinements may be applied at any stage in the life of an animal. Thus, refinement encompasses all aspects of a procedure, including:

- the source, transport, husbandry and environment of the animals involved;
- the experimental design (for example, the choice of species and the group size employed);
- the techniques applied;
- the end points of the procedures; and
- care of the animals before, during and after a procedure.

For further information regarding the National Centre for the Replacement, Refinement and Reduction of Animals in Research (NC3Rs), see www.nc3rs.org.uk

Monoclonal antibodies
The use of ascitic animals for monoclonal antibodies (mAb) production in vivo may only be proposed when in vitro attempts at mAb production have failed or the use of animals is considered justified for specific diagnostic or therapeutic products. You must give a full explanation if in vitro production methods are not considered to be suitable.

Select which of the following species you will use:
(Primate, Cat, Dog, Equidae, Pig, Genetically Altered Animals, Other animals)

The NC3Rs will review all applications involving the use of primates, cats, dogs and equidae animals, or their tissue/data.

All proposed research projects involving genetically altered mice are expected to consider the principles of welfare assessment set out on the NC3Rs website: https://www.nc3rs.org.uk/generation-and-breeding-genetically-altered-mice.

Select 'Add...' to enter the animal species and total numbers required (this may differ from the number to be purchased, maintained).

<table>
<thead>
<tr>
<th>Animal species</th>
<th>Strain (if appropriate)</th>
<th>Total number required to carry out proposed work</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
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</table>

Provide a justification of the proposed sample size alongside details of the planned statistical analyses. Describe experimental design, including any plans to reduce bias such as blinding or randomisation if appropriate. You must include power calculations if appropriate. (1,000 words max.)

You may provide your answer to this question in the field provided (text entry format) or as a PDF attachment (upload format). If you are uploading your answer, the uploaded document must be in 11 point Arial font and portrait format.

For each species, you must ensure that adequate experimental detail is provided to justify both the use and number of animals. This should include:

- definition of unit of analysis (i.e. N referring to animal or sample number);
- means of avoidance of bias (e.g. blinding or randomisation);
- statistical analysis to be used and explanation of how sample and/or group size was derived;
- where repeated measures are used, the number of time points;
- an indication of number of replications of each experiment to mitigate spurious non-replicable results.

You may include tables and figures in this section to help justify animal numbers. You can find additional guidance on designing animal experiments through the NC3Rs Experimental Design Assistant.

(1000 words max.)
<table>
<thead>
<tr>
<th>Why are the species to be used the most appropriate? (250 words max.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>It is particularly important to justify the species when an animal is being used as a model for a human physiological or pathological condition.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Does your proposal include procedures to be carried out on animals in the UK which require a Home Office licence? Your organisation must ensure that research involving the use of animals complies at all times with UK laws and regulations.</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Is there a current Home Office Personal Project Licence (PPL) that authorizes the proposed procedures to be carried out in the UK?</th>
</tr>
</thead>
<tbody>
<tr>
<td>Provide the name of the licence holder and the PPL number.</td>
</tr>
</tbody>
</table>

<table>
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<tr>
<th>Detail your plans and timelines for acquiring the appropriate licence.</th>
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</thead>
</table>

<table>
<thead>
<tr>
<th>If your proposal involves the use of animals, what would be the severity of the procedures? You can find guidance on assessing the severity of a procedure on the Home Office website.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Provide details of any moderate, severe or non-recovery procedures. (250 words max.)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Does your proposal involve the use of animals or animal tissue outside the UK?</th>
</tr>
</thead>
<tbody>
<tr>
<td>Confirm that the proposed animal work outside of the UK will comply with the principles of UK law. Animal research conducted outside the UK must, as a minimum standard, be carried out in accordance with the principles of UK law and regulation. (1000 words max.)</td>
</tr>
</tbody>
</table>

| Law and regulatory standards often vary from country to country. If experiments are to be carried out on animals outside the UK, the experiments proposed must be performed to standards which accord with the principles of UK legislation. Furthermore, the housing and care of animals must similarly accord with the standards and principles of UK legislation. You should refer to NC3Rs guidance on carrying out work abroad and choosing contractors. |

<table>
<thead>
<tr>
<th>Why is animal use necessary: are there any procedures of less severity that could be used? (250 words max.)</th>
</tr>
</thead>
</table>
Non-human primates If you are exclusively using already generated tissue or data in the proposed experiments answer all relevant questions as thoroughly as possible, in particular the questions relating to housing and procedures. If a question is not relevant you can answer N/A, but we may request additional information if necessary for assessment.

Do the facilities and practices, and the proposed research comply with the principles set out in the ‘National Centre for the Replacement, Refinement and Reduction of Animals in Research (NC3Rs) Guidelines: Primate accommodation, care and use’?

Please explain why not

Will it be necessary to transport the non-human primates (i.e. from breeding facility and within the host organisation environment)?

Indicate approximate journey times and the measures that will be taken to minimise the potential stress during transport.

Please provide details of the housing for the animals, e.g. enclosure size, environmental enrichment.

Will single housing of the non-human primates be necessary at any time?

Provide a justification for single housing, its duration, and explain what additional resources you will provide to the animals to minimise the impact on animal welfare.

Describe the experimental procedures involved and how any pain, suffering, distress and/or lasting harm will be minimised. Have the procedures been recently reviewed by the Named Veterinary Surgeon (NVS), Named Animal Care and Welfare Officer (NACWO) and Animal Welfare and Ethical Review Body (AWERB)?

Will any of the experimental procedures involve food and/or water restriction?

Justify why this is necessary and outline what alternatives have been considered.

Will any of the experimental procedures involve restraint?

What alternatives have been considered? Describe the nature of the restraint, its duration and frequency, and what will be done to avoid distress.
What prior experience and training in non-human primate use, care and welfare will you require of the staff named in the application? What provision are you making for continuing professional development in these areas?

Will any of the staff involved require specific training for any of the procedures concerned?

Provide details of the training needed and where it will be undertaken.

**Cats, Dogs, Equidae and Pigs** If you are exclusively using already generated tissue or data in the proposed experiments answer all relevant questions as thoroughly as possible, in particular the questions relating to housing and procedures. If a question is not relevant you can answer N/A, but we may request additional information if necessary for assessment.

From where will the animals be sourced?

Will it be necessary to transport the animals?

Indicate approximate journey times and the measures that will be taken to minimise the potential stress during transport.

Are animals to be imported?

Where animals are to be imported, what journey times have been agreed with the Home Office? Describe the conditions for the animals at the breeding establishment and how the potential stress during transport will be minimised.

Provide details of the housing for the animals, e.g. enclosure size, environmental enrichment.

*Please see the NC3Rs guidance on animal housing and husbandry for further details: https://www.nc3rs.org.uk/3rs-resources/housing-and-husbandry*

Will single housing of the animals be necessary at any time?

Provide a justification for single housing, its duration, and explain what additional resources you will provide to the animals to minimise the impact on animal welfare.
Describe the experimental procedures involved and how you will minimise any pain, suffering, distress and/or lasting harm. Have the procedures been recently reviewed by the Named Veterinary Surgeon (NVS), Named Animal Care and Welfare Officer (NACWO) and Animal Welfare and Ethical Review Body (AWERB)?

What adverse effects might the animals experience? List the clinical and other signs that will be monitored, the frequency of monitoring and, where relevant, the humane endpoint criteria established for the study. (1000 words max.)

Will any of the experimental procedures involve restraint?

What alternatives have been considered? Describe the nature of the restraint, its duration and frequency, and what will be done to avoid distress.

What prior experience and training in animal use, care and welfare will you require of the staff named in the application? What provision are you making for continuing professional development in these areas?

Will any of the staff involved require specific training for any of the procedures concerned?

Provide details of the training needed and where it will be undertaken.

### Risks of research misuse

Confirm that you have considered whether your proposed research could generate outcomes that could be misused for harmful purposes.

Wellcome encourages applicants and their host organisations to consider carefully any risks that the potential outcomes (information, products or technologies) of the research could be misused for harmful purposes. These include actions that pose a significant threat to humans, animals, plants or the environment - including terrorist misuse.

Examples of possible research areas that are associated with dual-use risks of this type, include (but are not restricted to) research that aims to:

- demonstrate how to render a vaccine ineffective
- confer resistance to a therapeutically useful antibiotic or antiviral agent
- enhance the virulence of a pathogen or renders a non-pathogen virulent
- increase the transmissibility or alter the host range of a pathogen
- enable the evasion of diagnostic and detection methods
- enable the weaponisation of a biological agent or toxin
- generate or reconstitute an eradicated or extinct agent or toxin
Have you identified any tangible risks of this type?

Briefly describe these risks and explain how you and your organisation will manage them. (250 words max.)

Where you judge there are tangible (real and non-hypothetical) risks that the proposed research will itself generate outcomes that could be misused to cause harm, you (and your fellow researchers and host organisations) must take appropriate steps to monitor the research as it proceeds and minimise these risks. Risk mitigation could include establishing a process to review dual use risks on an on-going basis through the project and to gain independent expert advice as appropriate. You must also ensure that all members of your team are aware of these risks in progressing their research, and receive appropriate education and training on these issues.

The identification of tangible risks in a research project should be clearly balanced against the benefits and value that is to be gained for health, science and society. We recognise that most research could conceivably generate results that might hypothetically be misused at some point in the future, and we are not asking applicants to appraise these kinds of remote and hypothetical risks.

Refer to the joint BBSRC, MRC and Wellcome policy and position statement on managing risks of research misuse, and our guidelines on good research practice.

**Freedom to operate/conflicts of interest**

Describe any freedom to operate or other intellectual property related issues that might affect your ability to carry out the proposed research and/or to use, share or commercialise the research outputs. Explain how you will address these.

In particular, consider the following:

- Will your research use technology, software, databases, materials or patented inventions that are owned or controlled by others and which you do not already have written permission to use?
- Will the ownership, use, commercialisation and/or sharing of research outputs with the wider research community, be subject to agreements with commercial, academic or other organisations? This includes arrangements with collaborators named in this application.

(250 words max.)

Refer to Clause 8 of our Grant Conditions at www.wellcome.ac.uk/funding/managing-grant/grant-conditions.

Disclose all relevant information pertinent to your grant proposal, including proprietary information where appropriate, to provide the most comprehensive picture of how any commercial/IP matters may affect the delivery of your proposed research and the subsequent use, commercialisation and/or sharing of your research outputs.

If you are satisfied that there are no issues, enter N/A. If you have fully addressed such issues in your outputs management plan under the question on “Outputs management and sharing”, then you may refer to that answer.

Describe any conflicts of interest which might affect your ability to carry out the proposed research and/or to share or commercialise the research outputs. Explain how you and your organisation will manage these and how you will comply with your organisation’s requirements in relation to conflicts of interest.

In particular, consider the following: Does anyone involved in your project hold any consultancies, advisory roles, or equities in, or directorships of, companies or other organisations that might have an interest in the results of your proposed research?
Confirm in each case whether the conflict has been disclosed to your organisation. (250 words max.)

Refer to our policy on conflicts of interest related to Wellcome-funded researchers and commercial organisations: www.wellcome.ac.uk/funding/managing-grant/policy-relationships-between-trust-funded-researchers-and-commercial-organisations.

If you are satisfied that there are no issues, enter N/A.

<table>
<thead>
<tr>
<th>Wellcome Trust supported facilities</th>
</tr>
</thead>
<tbody>
<tr>
<td>Will the project be based in one of the following Wellcome Trust supported facilities:</td>
</tr>
<tr>
<td>- the Wellcome Trust Sanger Institute</td>
</tr>
<tr>
<td>- a Wellcome Trust Centre</td>
</tr>
<tr>
<td>- an Africa and Asia Programme</td>
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<tr>
<td>- the Francis Crick Institute?</td>
</tr>
<tr>
<td>Specify</td>
</tr>
</tbody>
</table>

Sample