

## Key factors in trial design

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.....with thanks to Dr Wendy Baird and Prof Will Steward











## **Maximising Chance of Success**

- Ensure questions meaningful
  - if not, trial will have little/no value
  - Drives interest of investigators/patients
- Think through the questions to be answered
  - Formulate specific aims/trial objectives
  - Integrate potential investigators/teams in the development of questions/objectives
- Decide on biological endpoints and their value early during development of trial
  - Integrated, integral or correlative?

## **Understand the Setting**

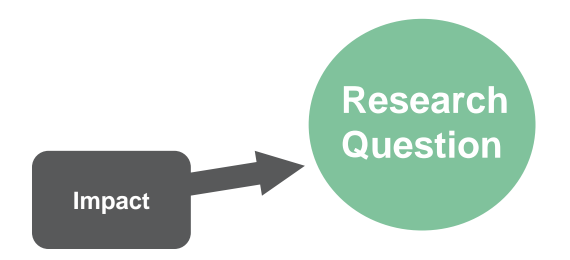
#### The key questions

- how much, how safe, how active, how effective?
- vary depending on the phase of development
- Understand <u>which</u> questions are appropriate for each stage of development
- Recognize that some questions bridge stages

#### Tailor the design of the trial to the therapy

- The <u>standard</u> approach is not the <u>only</u> approach
- Design a trial that incorporates or is even built around measurements best suited to capture the effect of that therapy







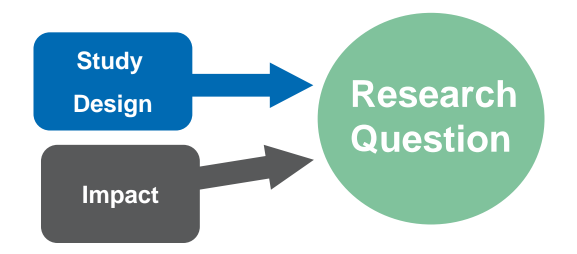
## 1. Impact

- Low risk strategy (incremental advance) e.g. Combining existing therapies
   If it is too modest (low-risk):
  - Will anyone be interested?
  - Does it justify use of resources?
- High risk strategy (breakthrough, paradigm changing advance) e.g. replacing standard therapy with new agent (imatinib in CML or GIST)

If it is too innovative (high-risk):

- Will anyone understand and want to participate?
- Put yourself in the place of a colleague who knows nothing about the background of the trial:
- Would you participate?
- If so, with what level of enthusiasm?



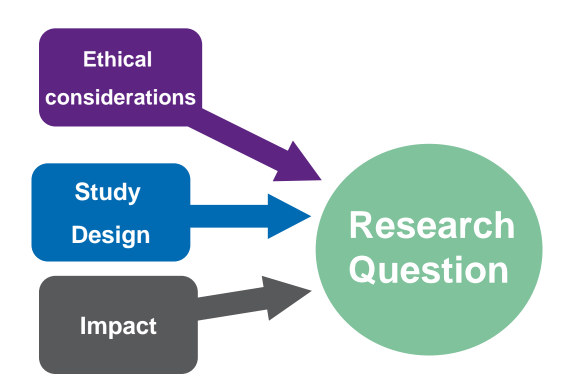




## 2. Study design

- Research approach (qualitative, quantitative or mixed methods)
- Study population (define, inclusion/exclusion criteria, sub-groups)
- Intervention (define intervention and comparator)
- Data collection (volume of data and collection methods)
- Data analysis (meta-analysis, homogeneous, type of study, presentation of results)
- Appropriate primary endpoint (PFS, OS)







#### 3. Ethical Considerations

- NHS patients, data or premises
- Vulnerable participants
- Highly sensitive topics
- Highly sensitive methods
- Patient burden
- Human Tissue Act







## 4. Is the study feasible?

#### Scientifically?

Are the clinical and biological endpoints valid, reliable, and appropriate?

#### Pragmatically?

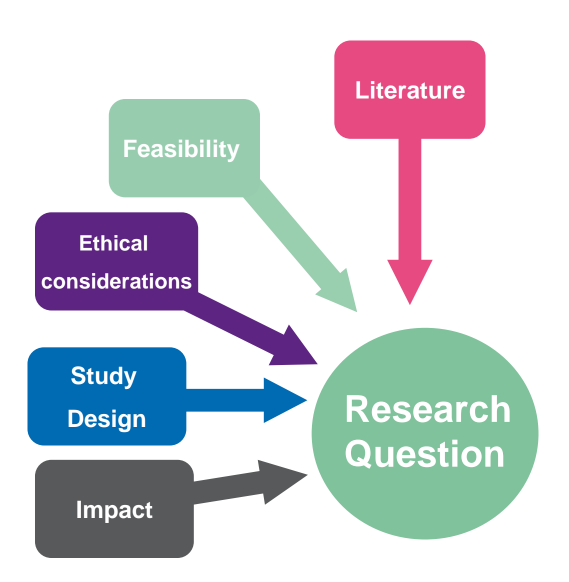
- Is it reasonable to expect that the RR will double or that the relapse rate will be reduced to zero?
- A question that is relevant today may not remain relevant if the study takes 10 years to complete
- Do you have access to an adequate number of patients to complete the trial in a reasonable period of time?
- Do you have the time to devote to all aspects of study development, recruitment and completion?

#### Financially?

Who will pay for extra tests, data collection, and follow-up?

#### Ethically?

Are you asking patients to forego "effective" treatment to participate in your trial?

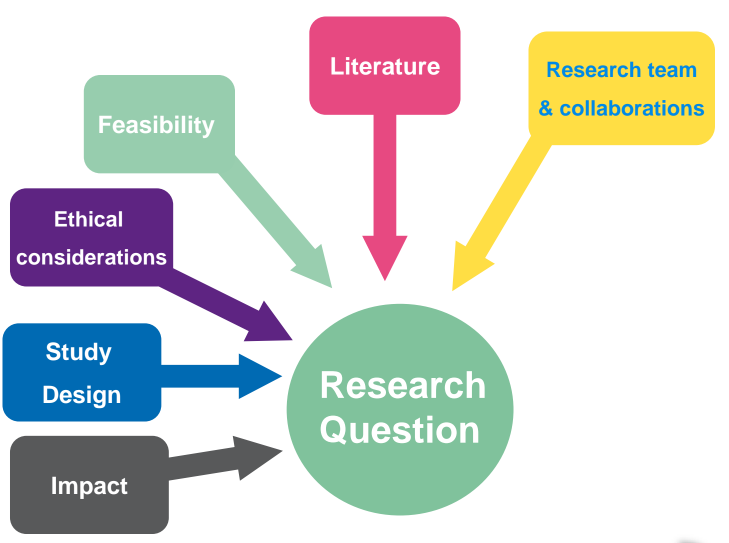




### 5. A literature review will help...

- Establish how good the pre-clinical data is
- Demonstrate a research need/ gap
- Find similar study designs approaches to help demonstrate feasibility/ expected outcomes
- Further develop the research question







#### 6. Research team and collaborations

- Integrate potential investigators/ teams in the development of questions and study objectives
- Ensure centres have experience and track record of quality
- Seek support from other departments early:
  - Surgery
  - Radiology
  - Research nurses
  - Statisticians
  - Data managers
  - Pathology
  - Laboratories or Clinicians

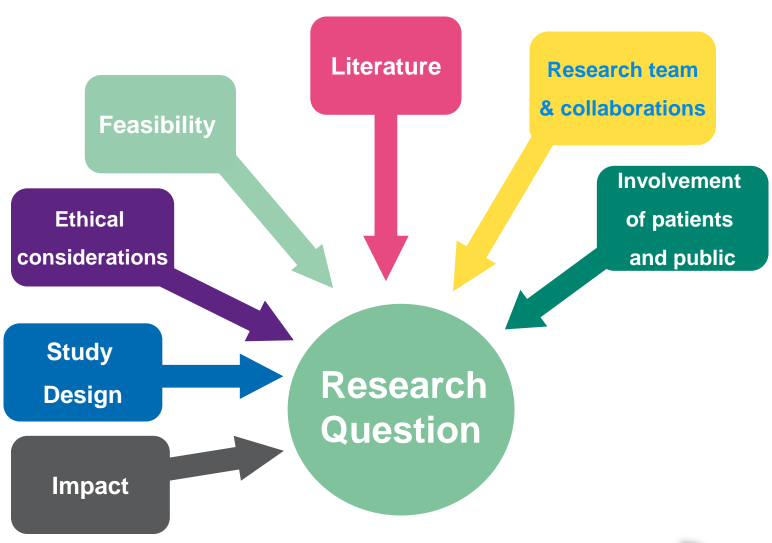


#### 6. Research team

#### In your Grant application you should highlight:

- Relevant skills and experience of the team that make them well placed to carry out the work
- ensure all components of the project have an appropriate person listed to complete the work
- 'sell' previous experience, such as working on other research projects
- if you have limited experience of running research then emphasise links with organisations that will guide you through – R&D, CTU, NIHR networks





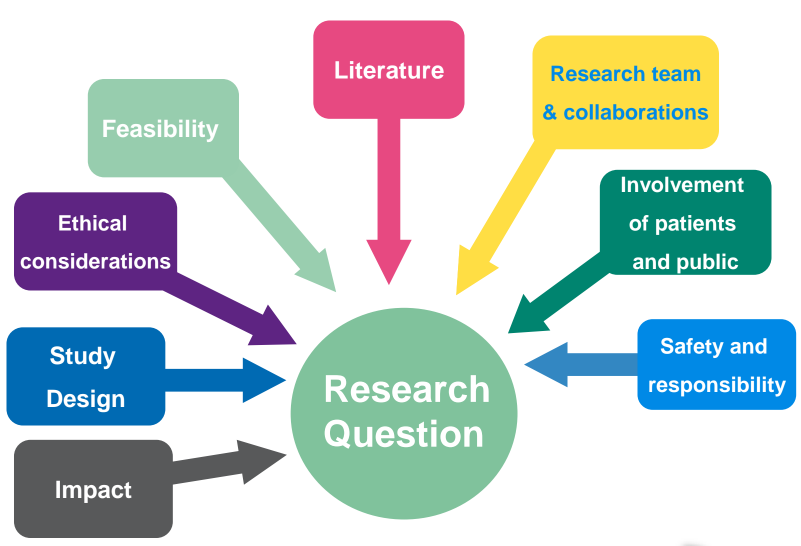


## 7. Involvement of patients and public

In your Grant application you should highlight where patients and public have been involved with:

- Design of the research
- management of the research (e.g. steering group membership)
- developing participant information resources
- undertaking/analysing the research (e.g. member of the research team)
- contributing to the drafting of the study report
- dissemination of the research







## 8. Safety and responsibility

- Don't underestimate adverse event reporting
- Ensure training of teams for data processing
- Inform the patient
- Establish a process to report toxicity
- Ensure data timeliness

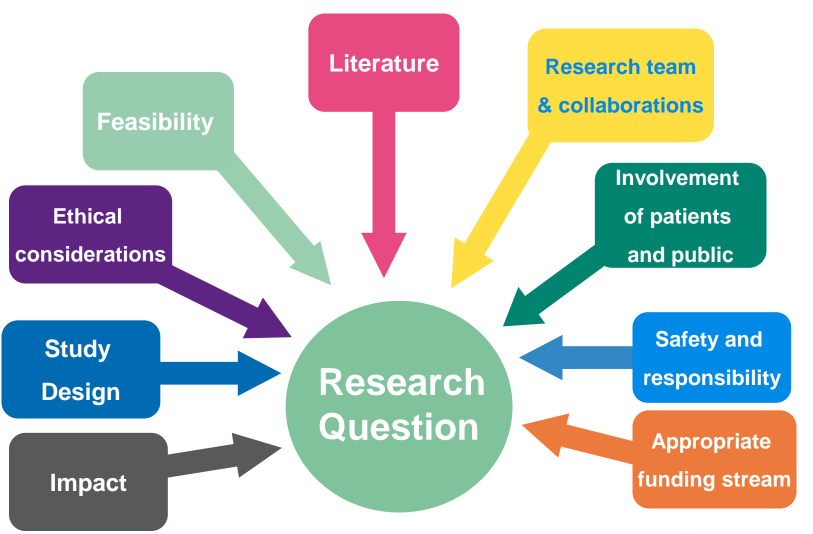
#### **Sponsor's responsibilities**

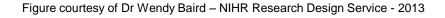
- Submission of SAEs as reported by the investigator
- Assessment of expectedness (SUSAR)
- Reporting SUSARs to Authorities
- Annual safety reports

#### Investigator's responsibilities

- Causality of AEs
- Reporting all adverse events in the source documents and CRFs
- Reporting SAEs within time period specified in the protocol
- Notifying Ethics committee







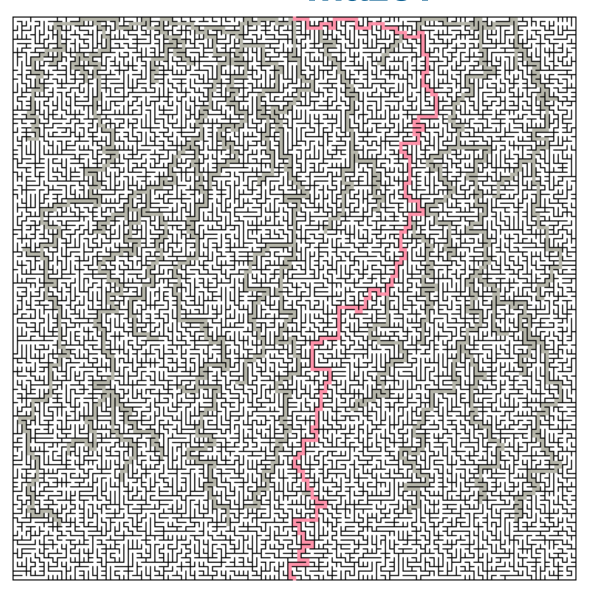


## 9. Appropriate funding streams

- Ensure your research question is within scope of the funding stream you are applying to
- If you are unsure ask *before* submitting an application



## How to find our way through the biomarker maze?





## **Categories of Biomarkers**

- Intended Use in the Trial
  - Integral
  - Integrated
  - Correlative
  - See definitions at <a href="http://biqsfp.cancer.gov/">http://biqsfp.cancer.gov/</a>
- EU Commission

http://ec.europa.eu/health/medicaldevices/documents/revision/index en.htm



# REporting recommendations for tumor MARKer prognostic studies (REMARK). Breast Cancer Res Treat. 2006 Nov;100(2):229-35.

McShane LM, Altman DG, Sauerbrei W, Taube SE, Gion M, Clark GM; Statistics Subcommittee of NCI-EORTC Working Group on Cancer Diagnostics

Cancer Research UK Biomarker Roadmaps: http://www.cancerresearchuk.org/science/funding/funding-committees/science-committee/biomarker-research/

Hall JA, Brown R, 2013, Developing translational research infrastructure and capabilities associated with cancer clinical trials, *Expert Reviews in Molecular Medicine*, Vol:15, ISSN:1462-3994



# Why do biomarker proposals not reach funding cut-off (a personal perspective)?

- No or flawed scientific hypothesis
- Stamp collecting
- Kitchen sink science
- Statistically underpowered
- Samples not fit for purpose
- Assay not fit for purpose
- Analysis not fit for purpose
- Committee didn't understand the proposal and they are all a bunch of idiots







**Cancer Research Network** 

#### NCRI Biomarkers and Imaging CSG

- To promote high quality translational (correlative) science within the NCRN portfolio of clinical trials in cancer through the following activities
- Identifying and monitoring strengths, weaknesses, opportunities and barriers
- Methodology harmonisation, design of generic protocols and education
- Interactions with tumour specific Clinical Studies Groups





**Cancer Research Network** 

#### **BICSG Work-Streams**

- Imaging integration and harmonisation (Fiona Gilbert)
- Biomarker technologies and applications (Craig Robson)
- Bioinformatics and biostatistics in biomarker study design (Expert Working Group)
- Education in biomarkers and personalised medicines