

On-campus course 'Improving health by improving trials'

Programme 19th September – 23rd September 2016

Location: University of Manchester

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| Monday 19 th | | Registration and coffee | |
| | | Welcome, introductions and overview of course | |
| | AM SESSION | What do you need to do to design a trial? <i>Professor Paula Williamson, University of Liverpool</i> | Topics: Identifying, defining and justifying the question; the importance of the protocol |
| | PM SESSION | General design issues <i>Dr Susanna Dodd, University of Liverpool</i> <i>Professor Carrol Gamble, University of Liverpool</i> | Topics: Feasibility, external and internal pilot studies; early and later phase trials, pragmatic and explanatory designs, internal and external validity; sample size considerations |
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| Tuesday 20 th | AM SESSION | Introduction to different designs <i>Dr Chris Sutton, Lancashire Clinical Trials Unit</i> <i>Dr Thomas Jaki, Lancaster University</i> | Topics: Adaptive designs; trials of complex interventions; cluster randomised trials |
| | PM SESSION | Recruitment of trial participants <i>Professor Bridget Young, University of Liverpool</i> <i>Dr Emma Bedson, Clinical Trials Research Centre</i> | Topics: Barriers and facilitators; effective recruitment and retention strategies; recruitment monitoring |
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| Wednesday 21 st | AM SESSION | Trial conduct (part 1) <i>Ms Helen Hickey, Clinical Trials Research Centre</i> <i>Dr Duncan Appelbe, Clinical Trials Research Centre</i> <i>Ms Katie Neville, Clinical Trials Research Centre</i> <i>Miss Joanne Eatock, Clinical Trials Research Centre</i> | Topics: Ethical, legal and regulatory requirements; barriers and facilitators to setting up sites; data sources; information systems and data management |
| | PM SESSION | Public and Patient Involvement <i>Professor Peter Bower, University of Manchester</i> <i>Dr Claire Planner, University of Manchester</i> <i>Ailsa Donnelly</i> | Topics: Basic principles of patient centred trials; evidence of benefit |
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| Thursday 22 nd | AM SESSION | Trial conduct (part 2) <i>Dr Catrin Tudur Smith, University of Liverpool</i> <i>Ms Helen Hickey, Clinical Trials Research Centre</i> Keynote talk 'How to be a good Chief Investigator' <i>Professor Peter Sandercock, University of Edinburgh</i> | Topics: Risk assessment; risk based monitoring; pharmacovigilance and safety monitoring; trial oversight committees |
| | PM SESSION | Visit from Clinical Trials Research Centre staff | |
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| Friday 23 rd | AM SESSION | Analysis and reporting (part 1) <i>Dr Richard Emsley, University of Manchester</i> | Topics: Key principles of trial analysis; intention to treat analysis; adjustment for compliance; efficacy and mechanism evaluation designs; stratified medicine trials; causal modelling; setting results in context |
| | PM SESSION | Analysis and reporting (part 2) <i>Professor Dyfrig Hughes, Bangor University</i> <i>Dr Jamie Kirkham, University of Liverpool</i> | Topics: Basic principles of health economics; methods of economic evaluation; economic outcomes; good practice in trial reporting |