



Job title	Trial Administrator	Job family and level	Administrative, Professional and Managerial Level 2
School/ Department	School of Medicine, Nottingham Clinical Trials Unit (NCTU)	Location	The East Midlands Campuses of the University of Nottingham

Purpose of role

The Nottingham Clinical Trials Unit (NCTU) is seeking to recruit a Clinical Trial Administrator with a strong background in administrative support. You will work within our multidisciplinary team to support the successful delivery of trials.

NCTU is a UK Clinical Research Collaboration registered Clinical Trials Unit (<http://www.ukrcctu.org.uk>) based in the School of Medicine at the University of Nottingham. The unit's mission is to conduct high quality, high impact multicentre trials to improve the nation's health and well-being.

NCTU has a broad portfolio of trials including studies evaluating medical devices, medicines and complex interventions. The unit currently has around 100 staff, and with the recent investment of strategic development funding from the University this is anticipated to expand to 120 staff over the next 3 to 5 years.

For further information see: www.nctu.ac.uk

The School of Medicine recognise the importance of continuous professional development and therefore the importance of providing opportunities, structured support and encouragement to engage in professional development each year.

To find out more about the School of Medicine, its values, vision, teaching and research, please see our [further information leaflet](#).

	Main responsibilities (Primary accountabilities and responsibilities expected to fulfil the role)	% time per year
1	<p>Administrative Support To provide administrative support to the trial team, primarily, to help ensure the smooth running of the trials, as well as support for the Unit Administrators:</p> <ul style="list-style-type: none"> ▪ Printing and sending out materials to study sites and pharmacy ▪ Sending questionnaires or other research related materials to participants ▪ Maintenance & filing to Trial Master File, local site files and central files ▪ Planning travel arrangements 	80%

	<ul style="list-style-type: none"> ▪ Helping to arrange and minute research related meetings ▪ General administrative duties ▪ Entering participants tracking data into trial management database (as required) ▪ Tracking sent and received documents ▪ Liaising with third party providers ▪ Keeping finance spreadsheets up to date and checking income and outgoings against the University finance system ▪ Undertaking training to ensure familiarity with Clinical Trial Regulations ▪ Undertaking training to ensure familiarity with University systems 	
2	<p>Contacting participants / sites:</p> <ul style="list-style-type: none"> ▪ Making regular contact, including calls, with participants in the study and recruiting sites ▪ Interpreting information given by participants and entering data to the database as appropriate ▪ Referring any issues arising as a result of the contact to the clinicians or the trial manager as appropriate 	10%
3	<p>Data entry:</p> <ul style="list-style-type: none"> ▪ Entering participant data onto the database in an accurate and timely manner ▪ Performing initial quality review of the CRFs for missing data and erroneous data, as required ▪ Highlighting data queries to the trial manager and liaising with Data Management and study site staff to resolve, if deemed appropriate by the trial manager. Where appropriate, queries are then followed up by the post holder with either the doctor or the participant, to ascertain the correct information 	10%
4	<p>Other</p> <ul style="list-style-type: none"> ▪ Any other duties appropriate to the grade and level of the role 	N/A

Person specification

	Essential	Desirable
Skills	<ul style="list-style-type: none"> ▪ High level of IT skills, particularly in Microsoft applications (Excel, Word, Outlook and PowerPoint) ▪ Excellent written and verbal communication skills and interpersonal skills ▪ Good organisational skills ▪ Capable of working without close supervision ▪ Ability to plan and prioritise work activities ▪ Attention to detail ▪ Ability to work as part of a team as well as individually ▪ Must have a good telephone manner ▪ Adaptable and flexible attitude 	<ul style="list-style-type: none"> ▪ Initiative
Knowledge and experience	<ul style="list-style-type: none"> ▪ Previous experience working in a busy environment 	<ul style="list-style-type: none"> ▪ Clinical trial administration experience ▪ Experience of working in a healthcare/research environment ▪ Experience of working as part of a team across multiple projects ▪ Experience contacting participants and doctors ▪ Clinical trial data entry experience ▪ Knowledge of databases ▪ Understanding of research approvals (ethics and Trust approval)
Qualifications, certification and training (relevant to role)	<ul style="list-style-type: none"> ▪ GCSE's grade C or above in English and Mathematics (or equivalent qualifications or experience) <p style="text-align: center;">OR</p> <p>Relevant vocational qualifications (e.g. NVQ2/3 administration, City & Guild) or equivalent, plus some experience in a relevant role</p>	
Other	<ul style="list-style-type: none"> ▪ Willingness to adopt the vision and values of the School of Medicine 	



Expectations and behaviours

The University has developed a clear set of core expectations and behaviours that our people should be demonstrating in their work, and as ambassadors of the University's strategy, vision and values. The following are essential to the role:

- Valuing people** Is friendly, engaging and receptive, putting others at ease. Actively listens to others and goes out of way to ensure people feel valued, developed and supported.
- Taking ownership** Is clear on what needs to be done encouraging others to take ownership. Takes action when required, being mindful of important aspects such as Health & Safety, Equality, Diversity & Inclusion, and other considerations.
- Forward thinking** Drives the development, sharing and implementation of new ideas and improvements to support strategic objectives. Engages others in the improvement process.
- Professional pride** Is professional in approach and style, setting an example to others; strives to demonstrate excellence through development of self, others and effective working practices.
- Always inclusive** Builds effective working relationships, recognising and including the contribution of others; promotes inclusion and inclusive practices within own work area.

Key relationships with others

