

Performance in Initiating and Delivering Clinical Research – NCH&C Q3 2016/17

The Government wishes to see a dramatic and sustained improvement in the performance of providers of NHS services in initiating and delivering clinical research. The aim is to increase the number of patients who have the opportunity to participate in research and to enhance the nation's attractiveness as a host for research.

The Government's Plan for Growth, published in March 2011, announced the transformation of incentives at local level for efficiency in initiation and delivery of clinical research.

As part of this incentivisation, the Department of Health requires, via the new National Institute for Health Research (NIHR) contracts with providers of NHS services, the publication on a quarterly basis of information regarding: the 70-day benchmark for clinical trial initiation; and the recruitment to time and target for commercial contract clinical trials.

Providers of NHS services are required to publish the following information for **Initiating Clinical Research** (i.e. the 70-day benchmark) on a publicly available part of their website:

- The name of the trial;
- The Research Ethics Committee reference number and IRAS number
- The date of receipt of a Valid Research Application;
- The date of the recruitment of first patient; and
- Where the benchmark has not been achieved for a particular clinical trial, the reason for not doing so.

From Q4 2015-16, for trials which are subject to HRA approval rather than NHS approval, providers must publish the following information:

- The Research Ethics Committee (REC) reference number
- The Integrated Research Application System (IRAS) Number
- The name of the clinical trial
- The date the site was invited
- The date the site was selected
- The HRA Approval date
- The date the site was confirmed by the sponsor
- The date the site was confirmed
- The date when the site is ready to start
- The date of the recruitment of first patient, and
- Reasons for delay in recruiting the first patient into a trial.

Plus, in the event that a trial initiation did not proceed to confirmation:

- The non-confirmation status.

Providers of NHS services are also required to publish the following information regarding commercial contract clinical trials, to meet the transparency commitment for delivering clinical research to time and target on a publicly available part of their website:

From Q4 2015-16, providers are required to publish the follow information for **Performance in Delivering** Clinical Research:

- The Research Ethics Committee (REC) reference number
- The Integrated Research Application System (IRAS) Number
- The name of the clinical trial
- Whether or not a target number of patients was agreed
- The minimum number of patients agreed to be recruited (if a range has been agreed; this will be the same as the maximum if a fixed number has been agreed)
- The maximum number of patients agreed to be recruited (if a range has been agreed; this will be the same as the minimum if a fixed number has been agreed)
- Whether or not a target date to recruit patients was agreed
- The date agreed to recruit the target number of patients
- The total number of patients recruited at the agreed target date
- The date that the trial closed to recruitment
- The reason for the closure of the trial .

Norfolk Community Health & Care NHS Trust published this data for the first time in Q3 2014/15

Table 1 - Performance in Initiating Research Q3 2016/17

- NHS permission given:**

Name of Trial	Research Ethics Committee Reference Number	IRAS number	Date of receipt of a Valid Research Application	Date of Recruitment of First Patient	70 day Benchmark Met?	Reasons for not meeting benchmark
ATILLA - Assistive Technology and Telecare to Maintain Independent Living At Home for People with Dementia	12/LO/1816	111293	18/02/2016	n/a	No	There were staff availability issues coming from the NHS Provider. The study is now closed to recruitment

- HRA Approval given:**

Name of Trial	Research Ethics Committee Reference Number	IRAS number	Date Site Invited	Date site selected	HRA Approval date	Date site confirmed by Sponsor	Date site confirmed	Date Site Ready to Start	Date of Recruitment 1st Patient	Reasons for delay in recruiting the first patient into the Trial
FeSTivAPP – an innovative method for delivering Functional Strength Training exercises for the Upper Limb in people after stroke – a feasibility study	16/WM/0481	159752	01/08/2016	16/12/2016	06/01/2017	06/01/2017	17/01/2017	n/a	n/a	70 days have not elapsed since the date the site was selected

For this Q3 16/17 there are no commercial clinical trials that fulfil the criteria for the PID reporting and there is a nil return for Performance in Delivering