Health Technology Assessment Programme



HTA no 13/144

Standing frames as part of postural management for children with spasticity

Introduction

The aim of the HTA Programme is to ensure that high quality research information on the effectiveness, costs and broader impact of health technology is produced in the most efficient way for those who use, manage, provide care in or develop policy for the NHS. Topics for research identified and prioritised to meet the needs of the NHS. Health technology assessment forms a substantial portfolio of work within the National Institute for Health Research and each year about fifty new studies are commissioned to help answer questions of direct importance to the NHS. The studies include both primary research and evidence synthesis.

Research Question:

What is the acceptability of a trial to determine the efficacy of standing frames?

- **1. Intervention:** Use of standing frames as part of a postural management programme.
- **2. Patient group:** Children with severe spasticity, such as that caused by cerebral palsy (CP) (Gross Motor Function Classification System IV or V GMFCS).
- 3. Setting: Home and community.
- **4. Comparator:** A suitable comparator to determine the efficacy of standing frames will be determined from this research.
- **5. Study design:** A sufficient survey of current use of standing frames (including clinical prescribing and carer reports of actual use); a qualitative assessment of attitudes and the acceptability of randomisation to different management regimes, e.g. no standing frame use, short period in standing frame or long period in standing frame.
 - The survey should involve relevant health care professionals, parents of children with severe spasticity (such as CP) and older children with a range of experiences of use or no use of a standing frame.
- **6. Important outcomes:** The findings of the survey (i.e. current practice standing frame prescribing by physiotherapists; willingness to be randomised); and proposed design of a substantive trial.

NHS decision problem to be addressed by this research:

Cerebral palsy is the most common condition associated with spasticity in children and young people and 1 in every 400 children in the UK is affected by the condition. The impact of spasticity and co-existing motor disorders and their early musculoskeletal complications on the child or young person varies, but the ultimate goal of treatment is to maximise the child or young person's potential and promote independence and quality of life. This may be achieved by improving motor function, relieving pain and preventing secondary musculoskeletal complications.

One of the 'treatments' used widely in the management of children with spasticity is postural management which is an accepted, although very under-researched, aspect of physical therapy. Postural management programmes include the use of standing frames, as well as resting, lying and seating support systems. Standing frames are part of established clinical practice as a means of improving functional ability, slowing or preventing musculoskeletal deformity and facilitating participation in activities. However, there is little evidence supporting their use and current regimes and some argue that they may be associated with discomfort and pain for the child along with

increased care demands on the family, which reduces their ability to enhance the child's participation and their quality of life. It is proposed that initial research is conducted among health care professionals, parents and children to determine what the current standing frame regimes are, the perceived benefits/harms and, ultimately, willingness to be randomised to a full trial to determine the efficacy of standing frame use. This study (survey) should be simple and of modest cost.

Notes to Applicants

The NIHR Health Technology Assessment Programme is funded by the NIHR, with contributions from the CSO in Scotland, NISCHR in Wales, and the Public Health Agency in Northern Ireland. Researchers from Northern Ireland and Scotland for certain NICE related calls should contact NETSCC to discuss their eligibility to apply.

For many of the questions posed by the HTA Programme, a randomised controlled trial is likely to be the most appropriate method of providing an answer. However, there may be practical or ethical reasons why this might not be possible. Applicants proposing other research methods are invited to justify these choices.

Applicants are asked to:

- Follow the Medical Research Council's Good Clinical Practice guidelines (http://www.mrc.ac.uk/Utilities/Documentrecord/index.htm?d=MRC002416) when planning how studies, particularly RCTs, will be supervised. Further advice specific to each topic will be given by the HTA Programme at full proposal and contract stages.
- 2. Note that trials involving medicinal products must comply with "The Medicines for Human Use (Clinical Trials) Regulations 2004". In the case of such trials, the DH expects the employing institution of the chief investigator to be nominated as the sponsor. Other institutions may wish to take on this responsibility or agree co-sponsorship with the employing institution. The DH is prepared to accept the nomination of multiple sponsors. Applicants who are asked to submit a full proposal will need to obtain confirmation of a sponsor(s) to complete their application. The DH reserve the right to withdraw from funding the project if they are not satisfied with the arrangements put in place to conduct the trial.

The MHRA (<u>info@mhra.gsi.gov.uk</u>, <u>http://www.mhra.gov.uk</u>) can provide guidance as to whether your trial would be covered by the regulations. The DH/MRC website (<u>http://www.ct-toolkit.ac.uk/</u>) also contains the latest information about Clinical Trials regulations and a helpful FAQ page.

In line with the government's transparency agenda, any contract resulting from this tender may be published in its entirety to the general public. Further information on the transparency agenda is at: http://transparency.number10.gov.uk/#

Applicants are recommended to seek advice from suitable methodological support services, at an appropriate stage in the development of their research idea and application. It is advisable to make contact at an early a stage as possible to allow sufficient time for discussion and a considered response.

The NIHR Research Design Service (www.nihr.ac.uk/research/Pages/ResearchDesignService.aspx) can advise on appropriate NIHR Programme choice, and developing and designing high quality research grant applications.

Clinical Trials Toolkit

Researchers designing or undertaking clinical trials are encouraged to consult the Clinical Trials Toolkit (www.ct-toolkit.ac.uk). This NIHR resource is a website designed to help researchers navigate through the complex landscape of setting up and managing clinical trials in line with regulatory requirements. Although primarily aimed at those involved in publicly funded Clinical Trials of Investigational Medicinal Products (CTIMPs), the Toolkit will also benefit researchers and R&D staff working on trials in other areas, who will find useful information and guidance of relevance to the wider trials environment.

Research networks

The HTA Programme expects, where appropriate, that applicants will work with the relevant research network.

Making an application

If you wish to submit an outline proposal on this topic, complete the on-line application form at www.nets.nihr.ac.uk/funding/hta-commissioned and submit it on line by **7 Jan 2014**. Applications will be considered by the HTA Commissioning Board at its meeting in **March 2014**. IMPORTANT: For outline applications, if shortlisted, investigators will be given a minimum of **eight**

weeks to submit a full proposal. The full proposal will be considered at the Commissioning Board in July 2014.

Applications received electronically after <u>1300 hours</u> on the due date will not be considered.

Please see GUIDANCE ON APPLICATIONS overleaf.

Guidance on applications

Required expertise

HTA is a multidisciplinary enterprise. It needs to draw on the expertise and knowledge of clinicians and of those trained in health service research methodologies such as health economics, medical statistics, study design and qualitative approaches. The HTA Programme expects teams proposing randomised controlled trials to include input from an accredited clinical trials unit, or one with equivalent experience. Applicants are also expected to engage a qualified Trial Manager for appropriate projects. A commitment to team working must be shown and applicants may wish to consider a collaborative approach between several institutions.

Public involvement in research

The HTA Programme recognises the benefit of increasing active involvement of members of the public in research and would like to support research projects appropriately. The HTA Programme encourages applicants to consider *how* the scientific quality, feasibility or practicality of their proposal *could* be improved by involving members of the public. Examples of how this has been done for health technology assessment projects can be found at www.nets.nihr.ac.uk/ppi. Research teams wishing to involve members of the public should include in their application: the aims of active involvement in this project; a description of the members of the public (to be) involved; a description of the methods of involvement; and an appropriate budget. Applications that involve members of the public will not, for that reason alone, be favoured over proposals that do not but it is hoped that the involvement of members of the public will improve the quality of the application.

Outcomes

Wherever possible, the results of HTA should provide information about the effectiveness and costeffectiveness of care provided in its usual clinical setting and for the diverse subjects who would be
eligible for the interventions under study. The endpoints of interest will in most cases include disease
specific measures, health related quality of life and costs (directly and indirectly related to patient
management). Wherever possible, these measurements should be made by individuals who are
unaware of the treatment allocation of the subjects they are assessing. We encourage applicants to
involve users of health care in the preparation of their proposal, for instance in selecting patientoriented outcomes. Where established Core Outcomes exist they should be included amongst the list
of outcomes unless there is good reason to do otherwise. Please see The COMET Initiative website
at www.comet-initiative.org to identify whether Core Outcomes have been established. A period of
follow up should be undertaken which is sufficient to ensure that a wider range of effects are identified
other than those which are evident immediately after treatment. These factors should guide applicants
in their choice of subjects, settings and measurements made.

Sample size

A formal estimate should be made of the number of subjects required to show important differences in the chosen primary outcome measure. Justification of this estimate will be expected in the application.

Communication

Communication of the results of research to decision makers in the NHS is central to the HTA Programme. Successful applicants will be required to submit a single final report for publication by the HTA Programme. They are also required to seek peer-reviewed publication of their results elsewhere and may also be asked to support NETSCC, HTA in further efforts to ensure that results are readily available to all relevant parties in the NHS. Where findings demonstrate continuing uncertainty, these should be highlighted as areas for further research.

Timescale

There are no fixed limits on the duration of projects or funding and proposals should be tailored to fully address the problem (including long-term follow-up if necessary). Applicants should consider however that there is a pressing need within the NHS for this research, and so the duration of the research needs to be timely.

Feasibility and Pilot studies

We expect that when pilot or feasibility studies are proposed by applicants, or specified in commissioning briefs, a clear route to the substantive study will be described. This applies whether the brief or proposal describes just the preliminary study or both together. Whether preliminary and main studies are funded together or separately may be decided on practical grounds.

Feasibility Studies are pieces of research done before a main study. They are used to estimate important parameters that are needed to design the main study. Feasibility studies for randomised controlled trials may not themselves be randomised. Crucially, feasibility studies do not evaluate the outcome of interest; that is left to the main study. If a feasibility study is a small randomised controlled trial, it need not have a primary outcome and the usual sort of power calculation is not normally undertaken. Instead the sample size should be adequate to estimate the critical parameters (e.g. recruitment rate) to the necessary degree of precision.

Pilot studies are a version of the main study that is run in miniature to test whether the components of the main study can all work together. It is focused on the processes of the main study, for example to ensure recruitment, randomisation, treatment, and follow-up assessments all run smoothly. It will therefore resemble the main study in many respects. In some cases this will be the first phase of the substantive study and data from the pilot phase may contribute to the final analysis; this can be referred to as an internal pilot. Or at the end of the pilot study the data may be analysed and set aside, a so-called external pilot.

For a full definition of the terms 'feasibility study' and 'pilot study' visit the NETSCC website glossary page www.nets.nihr.ac.uk/glossary

Diagnostics and Imaging

In evaluating diagnostic and imaging techniques, the emphasis of the HTA Programme is to assess the effect on patient management and outcomes (particularly where changes in management can be shown to have patient benefits). Improvements in diagnostic accuracy, whilst relevant, are not the primary interest of this commissioned research programme. Applicants should justify where they consider improvements in diagnostic accuracy to be relevant to these objectives. Where there is poor evidence to link diagnostic improvements to patient benefits, part of the primary research may be to assess the effects of such changes on patient outcome.

An assessment should also be made of changes in other resources (particularly other subsequent therapies) used as a result of changes in diagnostic methods.