INVESTIGATING THE MANAGEMENT OF CARIOUS PRIMARY TEETH IN GENERAL DENTAL PRACTICE: AN OVERVIEW OF THE DEVELOPMENT AND CONDUCT OF THE FICTION TRIAL

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ABSTRACT

The management of carious primary teeth is a challenge for patients, parents and clinicians. Most evidence supporting different management strategies originates from a specialist setting and therefore its relevance to the primary care setting is questionable. The UK National Institute for Health Research (NIHR) Health Technology Assessment (HTA) has commissioned the FICTION (Filling Children’s Teeth: Indicated Or Not?) trial; a multi-centre primary dental care randomised controlled trial (RCT) to determine the most clinically and cost-effective approach to managing caries in the primary dentition in the UK. This large trial began in 2012, is due to be completed in late 2017 and involves 72 practices and 1,124 children initially aged three to seven years with dentine caries, following randomisation to one of three caries management strategies. Clinical, radiographic, quality of life, treatment acceptability and health economics data are collected during the three-year follow up period. This article provides an overview of the development and conduct of FICTION and discusses some approaches adopted to manage challenges and achieve the patient recruitment target.

Introduction

The Filling Children’s Teeth: Indicated Or Not? (FICTION) trial is a randomised controlled trial (RCT) based in primary dental care, comparing alternative methods of managing caries in the primary dentition. This article describes some of the challenges encountered on the way to recruiting the target number of participants needed, and how the FICTION research team has worked along with the FICTION practice teams to overcome these challenges.

Dental care for the vast majority of children in the UK is provided by general practitioners rather than specialists; a situation common to many other countries. However, much of the evidence for the efficacy of restorations in the primary dentition is based on treatment undertaken in secondary care by specialists. This has contributed to uncertainty surrounding the most appropriate approach to managing carious primary teeth, with dental undergraduate teaching historically based on guidance advising that carious primary teeth should be managed by complete removal of carious tissues, followed by placement of a restoration. More recently, research has shown that selective (or partial) removal of carious tissue in primary teeth, prior to placement of a restoration, or indeed no caries removal at all (Hall Technique), can decrease postoperative complications.

Two studies, conducted in specialist paediatric dental practice, have been recently published. A hospital-based RCT in Germany found lower failure rates after one year with the Hall Technique over conventional restorations or prevention alone treatment (0%, 9% and 8% respectively). Additionally, a retrospective case note analysis showed high success rates using preformed metal crowns; conventionally and with the Hall Technique (94% and 97% respectively).

Evidence from specialist practice does not seem to have translated into primary care, with the proportion of primary teeth with visible caries that are restored (the Care

KEY WORDS

FICTION Trial, Dental Caries, Caries Prevention, Primary Teeth, Prevention, Paediatric Dentistry, Restoration, Fillings, RCT, Primary Care

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Dental caries remains the primary reason for children being admitted to hospital for treatment under general anaesthetic in the UK. In 2013/14, over 62,747 children and young people were admitted to hospital in England, Scotland and Wales with a diagnosis of dental caries; the most common age group being five to nine-year-olds (Table 2). This care has high direct costs for the NHS, with over £30 million spent on hospital-based tooth extractions for children aged 18 years and under in England alone in 2013-14 (NHS reference costs 2013/14). Indirect costs also accrue from this treatment (such as time off work, childcare, and non-prescribed medications).

In response to the evidence that dental caries was not being managed optimally and carried significant costs, the UK National Institute for Health Research (NIHR) Health Technology Assessment (HTA) Programme issued a call in 2007 to commission a study to answer the research question: “What is the clinical and cost-effectiveness of fillings in primary teeth, compared to no treatment?” As a result of that call, the FiCITION trial (registration no. ISRCTN77044005), a three-arm, parallel group, patient-level RCT, based in general dental practice was funded; the published protocol can be found at www.biomedcentral.com/1472-6831/13/25. The three interventions being assessed include an intermediate arm and are detailed in Table 3. All children in the trial, irrespective of allocated treatment arm, receive the same level of preventive care, i.e. best practice prevention. The FiCITION collaboration comprises clinicians and researchers from nine dental institutions across the UK with external oversight from a Trial Steering Committee and Independent Data Monitoring Committee.

### Pilot rehearsal trial and feasibility study

Before commencing the main FiCITION trial, a feasibility study and pilot rehearsal trial were conducted. In the feasibility study, a postal questionnaire sent to a randomly selected group of dental practices (n=273) in potential FiCITION trial areas gauged practitioners’ levels of interest and readiness to participate in the study. A total of 97 practices responded, and of these 70 (72%) said they would be prepared to participate in the main FiCITION trial.

The pilot rehearsal trial allowed the research processes (e.g. screening, recruitment, data collection) to be assessed prior to commencing the main trial and is discussed later. In parallel, a qualitative investigation of the views and opinions of both service providers (dentists and the practice team) and participants (children and their parents) on participating in FiCITION processes was undertaken. The findings of these studies were used to modify practice training, improve research processes and increase the required screening numbers for the main FiCITION trial.

### The main study Design

The main FiCITION trial commenced in 2012 (Figure 1). The recruitment phase is now complete and participant follow-up is underway. The primary outcome for the trial is the incidence of pain and/or sepsis for the three different caries management approaches carried out over a three-year period. Secondary clinical outcomes include; incidence of caries in primary and secondary teeth, quality of life, acceptability of treatment experiences to children and parents, and dentists’ treatment preferences.

### The main economic objective of FiCITION

Is to determine the incremental cost per
episode of pain and/or sepsis for the different treatment approaches. To allow a full understanding of cost-effectiveness and add value to the analysis, two different ways of measuring incremental costs will be compared:

1. Time/material-based costs (cost at the practice level).
2. The current cost to the NHS for treating children with caries in primary teeth incorporating national funding arrangements.

Dental practices across five main regions of the UK (London, North-East England, Scotland, Wales and Yorkshire) with the later addition of Liverpool and Manchester have been recruited and trained for the study. Each centre has a Clinical Lead who is a Specialist in Paediatric Dentistry. Eligibility for Fitness4Children participation depended on practices caring for children under NHS contracts with participating practices inviting children to be screened for potential participation in Fitness4Children against eligibility criteria (Table 4) as part of their routine check-up.

Progress to date

The target sample size required for the Fitness4Children trial was initially calculated based on pain and/or sepsis rates recorded in the literature. Assuming three-year rates of 20%, 10% and 3% for prevention alone, conventional with prevention and biological with prevention arms respectively, and allowing for loss to follow up over the three years of 25%, it was calculated that a total of 1,461 child participants would provide 90% power for the comparisons of the differences in rates between the groups.
Based on the FiCTION pilot rehearsal trial, we predicted that 50 dental practices (80-100 dentists), screening a total of approximately 12,000 children and each recruiting to a target of 30 participants, would be required to meet the overall recruitment target within 12 months (Figure 2).

Practice recruitment and training for the main FiCTION trial commenced in summer 2012 (Figure 1). The first practices began to recruit participants in October 2012, although it took until spring 2013 to complete training and site visits for the first cohort (Phase 1; n=44) of practices. FiCTION-trained GDPs in each participating practice were responsible for child participant recruitment to the study.

By early 2013, it was evident that the child participant recruitment rate was lower than initial projections (Figure 3). Dental practices, initially confident of recruiting sufficient participants, reported difficulties in finding enough children who met the inclusion criteria. The numbers of children judged to be clinically free of active caries appeared higher than predicted from the results of the pilot rehearsal trial. Additionally, 17 practices (39% of Phase 1 practices) became unable to continue with the trial for a variety of reasons.

In response to the shortfall in recruitment, additional dental practices were recruited through Phases 2-5 (Table 5). The impacts of the unanticipated training of new practice staff, plus 28 additional practice setup and associated monitoring visits, placed pressure on financial, time and staff resources. To date, 72 recruiting practices (209 dentists; 228 team members) have been trained.

By the end of 2013 it had become apparent, with the recruitment rate dwindling, that achieving the target of 1,461 participants was unlikely (Figure 3). To address this, a contract variation was submitted to the HTA in August 2014 in which it was projected that, with a 12-month extension, the trial could recruit 1,113 child participants by June 2015. Of these, the 996 randomised by June 2014 will have the originally proposed three-year follow-up, while the remaining 117 participants will have a variable follow-up of between two and three years, with follow-up complete by June 2017. These measures would allow for 82% power to detect a difference between the groups and was accepted by the HTA. Additional practices were then recruited in Manchester and Liverpool to help reach the revised target (Table 5).

**What have we learnt so far**

There is a drive to improve efficiency within trials and although FiCTION is ongoing, valuable lessons have already been learnt; most notably through the conduct of the pilot rehearsal trial and feasibility study, summarised above, and supplemented more recently by a mid-recruitment survey of dental practices to determine the main barriers and facilitators to participant recruitment.

**TABLE 3**

| Best practice prevention alone | Biological with best practice prevention | Conventional with best practice prevention |
|------------------------------|----------------------------------------|------------------------------------------|
| Dental caries is managed with best practice preventive care\textsuperscript{5,34} to arrest carious lesion progression, with teeth left to exfoliate naturally in due course. | Local anaesthetic injections are not usually required. Dental caries is either partially removed or left in situ, and then sealed from the oral environment with adhesive restorative materials or a preformed crown. Best practice prevention is carried out in line with current guidelines. | Dental caries is completely removed, generally under local anaesthesia, and the tooth restored with a restorative material in line with current evidence.\textsuperscript{35} Best practice prevention is carried out in line with current guidelines. |
From the pilot rehearsal trial
Conducted in three regions of the UK (Scotland, North-East England and Sheffield), the pilot rehearsal trial involved 11 practices (20 dentists) and ran from January 2010 to October 2011 (Figure 1). While uptake amongst eligible participants was high (80%), by the end of this 18-month pilot, fewer than 50% of the expected number of patients had been recruited. This was valuable information which, along with the comments and suggestions of stakeholder groups, (Table 6) helped inform main trial planning in a number of ways, including a revision of the planned numbers of children to be screened.

Table 4: Child Participant Eligibility Criteria

| Inclusion                                                   | Exclusion                                                   |
|-------------------------------------------------------------|-------------------------------------------------------------|
| • Aged three to seven years at enrolment                    | • Lack of accompanying adult legally                        |
| • Willing to be dentally examined                           | • able to consent                                           |
| • Caries into dentine in $\geq 1$ primary molar teeth       | • Pain due to caries                                         |
| • Likely to return for follow-up                            | • Sepsis                                                    |
|                                                            | • Medical condition requiring treatment in secondary care   |
|                                                            | • Involvement in competing study                            |
|                                                            | • Likely to leave the practice catchment area during study  |

Table 5: Distribution of Practices Who Have Recruited at Least One Participant to Fiction According to Region and Phase of Site Entry

| Region               | London | North-East England | Scotland | Wales | Yorkshire | Liverpool & Manchester | Total |
|----------------------|--------|--------------------|----------|-------|-----------|------------------------|-------|
| Phase 1 (Sep 12-Jun 13) | 9      | 11                 | 12       | 4     | 8         | -                      | 44    |
| Phase 2 (Jul 13-Aug 13) | -      | 2                  | 2        | -     | -         | -                      | 4     |
| Phase 3 (Sep 13-Nov 13) | -      | 6                  | 5        | -     | 1         | -                      | 12    |
| Phase 4 (Feb 14-Mar 14) | 2      | 1                  | 5        | -     | 2         | -                      | 10    |
| Phase 5 (Jan 15)      | -      | -                  | -        | -     | 2         | 2                      | 2     |
| Overall              | 11     | 20                 | 24       | 4     | 11        | 2                      | 72    |

*Month of site initiation visit

On the basis of this feedback, changes made to the main trial included:
• Paperwork refined to allow more efficient completion.
• FiCTION-branded merchandise was created for participants: fluffy bugs, colouring-in pages, brushing timers, bookmarks.
• Colouring-in competitions.
• Quarterly newsletter for distribution to practices.

These modifications aimed to make participation more straightforward for practices and dental teams, and proactively engage participating children in the research.24 The introduction of FiCTION merchandise was well received by the children as a gesture of thanks for their efforts in the study (Figure 4).

From the mid-recruitment questionnaire
Having initially recruited strongly (Figure 3), the rate of participants joining FiCTION began to slow around May 2014. This is well recognised as a common pattern and many trials in different settings struggle to recruit to target.25-27 In order to understand and try to address the barriers to recruitment, a web-based questionnaire study23 was conducted to allow practitioners to comment on participant recruitment processes.

An existing clinical trial survey tool was adapted to reflect the major differences in contextual settings of FiCTION (i.e. primary dental environment).28 This modified survey was distributed online and explored practitioners’ and dental
team members’ views on their experiences of participating in the trial, and allowed them to suggest improvements to facilitate participant recruitment.

Although some proposals involved aspects of the trial that were inflexible at this stage—for example increasing eligibility criteria (age range), additional remuneration for practices and simplifying the paperwork—other proposals were amenable to intervention, for example increasing the number of sites, acknowledgement of the effort of teams and participants and offering training to additional team members (Table 7).

**Ongoing changes**

Recruitment challenges in clinical trials are not limited to FICTION or to dentistry\(^\text{22-27, 30}\), and later recruitment difficulties were managed with a flexible approach iteratively. Practices joining later were asked to carry out a database search for patients of appropriate age to ensure an eligible child participant cohort. Practices were also encouraged to send the trial team a copy of their letterhead and lead dentist’s CV to be gauged.

Review and refinement of the training process for FICTION practices throughout the trial has enabled new practitioners and teams to be recruited and trained with minimal disruption to clinics. This has been particularly valuable when bringing new dentists into existing FICTION practices.

In terms of communication with practices, we have responded to practitioners’ preferences for contact by the trial team. Initially, evening meetings were held for FICTION practices but, as the trial has progressed, we have moved to lunchtime teleconferences in some areas. This has improved attendance and allowed more practitioners to share their thoughts on how trial processes could be further improved. Feedback suggests that the changes actioned over the course of the study have positively contributed to reaching our revised target of 1,113 participants.

Given the work required to achieve this target, the FICTION trial represents a rich source of information about the design and conduct of research trials in primary dental care.

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**TABLE 6**

COMMENTS & SUGGESTIONS OF STAKEHOLDER GROUPS

| Group          | Comments and suggestions                                                                 |
|----------------|-----------------------------------------------------------------------------------------|
| Participants   | • Stickers and colouring-in materials should be available                                 |
|                | • Some children didn’t understand or remember the trial                                  |
| Parents        | • Some felt they would like more information on the trial                                  |
|                | • Paperwork could be refined                                                               |
|                | • The appointments seemed to be longer                                                   |
|                | • Rewards for the children should be introduced                                           |
| Dental teams   | • Identifying enough suitable patients with dental caries was more challenging than expected |
|                | • Low response rates from invitation pack                                                  |
|                | • Unprepared for some treatment arms; conventional arm, Hall Technique, radiographs      |
|                | • Rigidity of study protocol was unfamiliar                                                 |

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**INVESTIGATING THE MANAGEMENT OF CARIOUS PRIMARY TEETH IN GENERAL DENTAL PRACTICE: AN OVERVIEW OF THE DEVELOPMENT AND CONDUCT OF THE FICTION TRIAL**
Conclusion

The outcomes of the Fiction trial will be known in early 2018. It is not possible, nor appropriate, to hypothesise at this time which treatment arm might be the most clinically and cost-effective. One treatment arm may be superior for the primary outcome with a reduction in pain/infection incidence. However, should there be no clear difference, the secondary outcomes (costs, anxiety, provider preferences, quality of life, etc.) will help inform the overall trial conclusions and recommendations. For example, if children find it difficult to tolerate a clinically effective strategy, then its success in the practice setting could be reduced. The secondary outcomes will also inform decision-making about implementation of future clinical guidance into general practice.

This randomised controlled trial is being conducted on a scale not previously attempted in paediatric primary dental care. Research is not easy, particularly in the busy primary care setting and the Fiction trial has been an ambitious project from the start. Whilst there have been difficulties, a successful participant recruitment phase has now moved into the equally important follow-up phase although the transition from recruitment to retention will no doubt hold new challenges. A flexible and pragmatic approach has been adopted throughout, while maintaining the rigorous principles required in research.

The progression of the trial has only been possible because of the goodwill and commitment of all the Fiction dental practice teams across the UK, who continue to treat their Fiction children. Additional thanks must go to the children and families participating in the trial. The findings will inform the clinical management of the primary teeth of children in the UK through general dental practitioners, evidence-based recommendations, commissioning of services and policy and ultimately will positively impact upon the health and experiences of our young patients.

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TABLE 7

| Suggestions                                      | Action taken                                                                 |
|--------------------------------------------------|-----------------------------------------------------------------------------|
| Acknowledgment of efforts of teams and families of participants | Practices successful in recruitment were sent tea-break sets with Fiction-branded mugs as a small token of the trial team's appreciation of their efforts. Participants already received Fiction merchandise as part of their recall appointments. |
| Increase recruitment of practices                | Additional sites were recruited from the Manchester and Liverpool regions. Some additional practices had prior hands-on experience of research projects. |
| Training and delegation to involve full practice team | The training syllabus for Fiction was modified to make the process simpler and more effective to allow the entire practice team to help and increase awareness of Fiction. |
| Positive support from central Fiction Team       | Additional intensive support was offered mostly on site to new practices in the early stages to assist in the transition from training to recruitment. |

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