Roberts, Andrew and Wales, Johanna and Smith, Heather and Sampson, Christopher James and Jones, Peter and James, Marilyn (2014) A randomised controlled trial of laser scanning and casting for the construction of ankle foot orthoses. Prosthetics and Orthotics International . ISSN 0309-3646 (In Press)

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A randomised controlled trial of laser scanning and casting for the construction of ankle foot orthoses

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November 25, 2014

Publication type:
Peer-reviewed journal article

This version:
This is a post-print version of the manuscript, meaning that it has been peer-reviewed.

Published version:

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A randomised controlled trial of laser scanning and casting for the construction of ankle foot orthoses.
ABSTRACT

Study Design: Randomised controlled trial with blinding of orthotists and patients to the construction technique used. Background: Three-dimensional laser scanning has been used for patient measurement for cranial helmets and spinal braces. Ankle foot orthoses are commonly prescribed for children with orthopaedic conditions. This trial sought to compare ankle foot orthoses produced by laser scanning or traditional plaster casting. Objectives: Assessment of the effectiveness and efficiency of using laser scanning to produce AFOs.

Methods: A randomised double blind trial comparing fabrication of AFOs from casts or laser scans. Results: The time spent in the rectification and moulding of scanned AFOs was around 50% less than for cast AFOs. A non-significant increase of 9 days was seen in the time to delivery to the patient for LSCAD/CAM. There was a higher incidence of problems with the scan-based AFOs at delivery of the device, but no difference in how long the AFOs lasted. Costs associated with laser scanning were not significantly different from traditional methods of AFO manufacture. Conclusions: Compared with conventional casting techniques laser scan based AFO manufacture neither significantly improved the quality of the final product nor delivered a useful saving in time.

(CLINICAL RELEVANCE

Ankle foot orthoses (AFOs) are a common requirement for chronic neurological conditions during childhood. Improved efficiency of provision of AFOs would benefit children and families by reducing the delay in provision of devices and would benefit the health service by making best use of valuable orthotist time.

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(word count 47)
BACKGROUND

Orthoses are devices that are applied externally in order to modify structural and functional attributes of the neuromuscular and musculoskeletal systems, for example in cerebral palsy.

The prevalence of cerebral palsy in the UK has been estimated as 2 per 1000 live births for birth years 1986-1996\(^1\).

AFOs have considerable support in the literature in the management of cerebral palsy\(^2\).

Children who are managed in AFOs often require intervention and care from multiple clinicians and agencies so efficient orthotic provision is important in reducing the logistical demands on their families.

The traditional method of producing an AFO involves taking a negative limb cast from the patient and then filling the negative cast with plaster to make a positive cast. The orthosis is then formed by moulding polypropylene sheets over the positive cast. This method can be replaced by laser scanning with computer-aided design and manufacture (LSCAD/CAM), where a positive of the limb is milled in foam directly from the digital image. A recent systematic review suggests that foot orthotics designed from 3D scans are at least comparable with those made through traditional methods, although none of the studies reviewed were randomised controlled trials and further research is required to confirm this \(^3\). One study suggested that plaster casting may be preferable to laser scanning when it is important to capture the forefoot-to-rearfoot relationship \(^4\).

We were able to identify four studies comparing plaster casting with LSCAD/CAM technology in the fabrication of spinal orthoses for scoliosis \(^5\-\(^8\). Radiological improvement of the scoliotic curves was assessed in three of these studies, and LSCAD/CAM was reported to be as effective as the traditional method. The LSCAD/CAM technique was also found to increase
productivity in both of the studies which measured this outcome. None of these studies measured health economic outcomes.

In the NHS, high capital costs and a lack of orthotists with time to learn new technology are likely reasons why LSCAD/CAM technology has not been introduced. If it proved successful, LSCAD/CAM could enable a more efficient process for the provision of a large number of bespoke orthotic devices.

Two hypotheses are explored in this paper. Firstly, that there is a difference in the length of time taken to deliver of a well-fitting AFOs produced by means of LSCAD/CAM compared to AFOs produced by traditional plaster casting. Secondly, that there is a difference in patient benefit in terms of functioning and satisfaction with AFOs produced by means of LSCAD/CAM compared to AFOs produced by traditional plaster casting. In addition the cost-effectiveness of AFOs produced by means of LSCAD/CAM versus traditional plaster casting was estimated using a societal perspective.

**METHODS**

**Design of Study**

A double blind randomised (1:1) controlled trial design in which patients had their AFO(s) fabricated either using LSCAD/CAM technology or by a traditional plaster method.

A positive opinion for the study was received from the xxxxxxx Local Research Ethics Committee on 15th August 2007 (ref: 07/Q2604/85).

**Inclusion and Exclusion Criteria**

All patients up to 18 years of age referred for rigid and hinged AFOs at the Trust were candidates for inclusion. Young adults up to the age of 21 years were also included if they were
still receiving services for children (for example, because they were in education). Exclusion of the patient occurred if informed consent was not obtained; the patient was considered by the orthotist to be un-splintable; a Saltiel style device had been specified or if communication was not possible with the subject or their parent/carer.

**Setting**

The study took place in the xxxxxxx Unit (xxxxx) and Orthotics Departments at the xxxxxxxxxxx NHS Foundation Trust between February 2008 and August 2010. Eligible subjects were contacted by the study nurse who provided an explanation of the study. Those subjects who assented to participate had demographic data and details about diagnosis and condition collected.

**Interventions**

All subjects had both a casting and a scan of each limb requiring an AFO, in the same session, and the order in which the two methods were performed was determined according to a computer-generated randomisation list. The allocation sequence was concealed from both the researcher and the participant in sequentially numbered, sealed envelopes. The study orthotists were coached on how to minimise bias in their interaction with the subject (for example, participants were to receive the same level of encouragement for each method). After experiencing both methods subjects were asked to state whether they would prefer to have casting or scanning next time. The time taken and consumables for each method and session were measured and recorded by the attending clinician. The subjects or their parents/carers completed a questionnaire about travel and associated costs. The subject was given an appointment to return two weeks later to receive their AFO(s). Both the LSCAD/CAM digital
image (scan) and the cast were rectified by an orthotic technician and length of time for each method was recorded. Our usual Standard Operating Procedure was applied to both the cast and scan where 3mm is applied to malleoli/metatarsals and navicular when requested on the specification sheet. A 1% increase in all dimensions was added to the scans to allow for ease of doffing and donning as well as allowing for growth.

A second randomisation was then performed to determine whether the AFO was manufactured from the cast or from the scan. Allocation to a manufacturing method was achieved through a central telephone randomisation system provided by the University of York Trials Centre, and was stratified by age (less than 6 years and 6-21 years), Gillette Functional Walking score (1-5 and 6-10)\(^9\) and required number of AFOs (unilateral or bilateral).

Laser scans were obtained with a Fastscan (Polhemus, 40 Hercules Drv, PO Box 560, Colchester, VT 05446, USA.) and rectified using Rodin4D software (Rodin4D Parc Le Biogalien - Bâtiment B, 27 Allée Charles Darwin, 33600 Pessac, France). Image files were passed to an orthotic milling company (Relief Orthotics, Unit 34B, Viking Road, Airdrie, Scotland ML6 9SE) who milled a positive of the limb in foam that was returned to the Trust for manufacture of the polypropylene device. Where cast manufacture was employed, a plaster male form was cast, rectified and then draped in the usual fashion. The length of time for the moulding of the polypropylene AFO was recorded. There were no differences in the manufacturing process between the two groups once the draping of the polypropylene sheet and curing had been performed.

**Sample size Calculations**

The sample size was based on data which suggest the total time taken in the moulding and rectification process is reduced by approximately 60% when using LSCAD/CAM compared to
casting. The proposed sample size of 150 patients would enable the detection of a proportional reduction in time taken in the moulding and rectification of a cast of 30% with 90% power using a two-sided test at $p=0.05$.

**Outcome Measures**

1. The primary outcome measure was the length of time taken (in minutes) in the moulding and rectification process.

2. Secondary outcome measures included: (a) Length of time (in minutes) spent with subjects to cast and scan limbs. (b) Length of time (in minutes) to fit the AFO(s). (c) Number of days taken from initial scanning/casting of a subject to the completion of an AFO (or AFOs) that met fitting specifications. Any additional time taken for rescanning or remaking was included. (d) The length of life of the AFO in days.

Patient-focused outcome measures included the “Satisfaction with Device” and “Satisfaction with Service” questionnaires, which were developed in the US as part of the Orthotics and Prosthetics Users’ Survey (OPUS)\textsuperscript{10}. Both questionnaires use a four-point Likert scale ranging from Strongly Agree (highest level of satisfaction) to Strongly Disagree (lowest level of satisfaction). Satisfaction with Device was measured with nine items (e.g. skin abrasions/irritations; whether the orthosis was pain-free to wear; whether it looked good and whether it was easy to put on) and ten items measured Satisfaction with Service (e.g. amount of time waiting for services; whether the orthotist was responsive to concerns; and whether staff co-ordinated services with doctors and therapists). The questions were administered by post at 3, 6, and 12 months follow-up. Non-responders were followed up by telephone if possible. In most cases parents/carers acted as proxies by answering the questions on behalf of their child.
**Statistical Analysis**

All planned analyses were by intention-to-treat. Differences in the primary and secondary outcome measures were analysed using the independent t-test for continuous data and the logrank tests for time to event data. Data from the patient-focused satisfaction questions, using the Likert scale, were analysed individually using the Mann Whitney U test, proportions using chi squared tests and time to event data using the logrank test. All tests were carried out using a 5% level of significance and no adjustment was made for multiple testing and missing values were not imputed.

**Economic Analysis**

In addition to the above, an economic analysis was completed of LSCAD/CAM compared with traditional casting. The work undertook a cost-effectiveness analysis using the time to functional AFO as the primary outcome, and sought to assess costs using a societal perspective. An individual resource collection tool was developed through pilot work with parent-representatives and designed to fully capture all the elements. In addition to intervention costs and health care resource use, this included the cost of patient travel to the hospital and productivity losses associated with time off work due to treatment and care. The time horizon was 12 months; as such, costs and outcomes were not discounted.

Resource use was collected by various means and in most cases national unit costs were assigned. Resource use relating to the scanning and casting sessions and the manufacture of a device were recorded by the attending orthotist. This included data on the time spent carrying out specific tasks, staff attending, the use of consumables and the duration of group sessions. Similarly, these data were collected for fittings and adjustments.
Follow-up data were collected from patients at 3, 6, 9 and 12 months. These included medication, time off work, service use and associated travel costs. Published unit costs from NHS Staff Earnings Estimates\(^\text{11}\), Personal Social Services Research Unit \(^\text{12}\) and British National Formulary \(^\text{13}\) were applied as appropriate. All costings were made at 2011 prices.

An incremental cost-effectiveness ratio (ICER) – the cost per day saved of LSCAD/CAM compared with traditional casting – was estimated and bootstrapping was used to estimate confidence intervals for the ICER. Results are based on a complete case analysis.

RESULTS

Subjects

During the study the provision of orthotic services at the Trust was reorganised following a “sustainable services review” and the study orthotists left the Trust. At this point, 92 (69% of total) subjects had been recruited into the study. New study orthotists were less experienced in laser scanning and had less time to devote to the study so the ability to recruit patients into the trial became more difficult. Despite an extension to the study duration; widening of the inclusion criteria from 18 years to 21 years of age (provided the subject was in education); and a move to bring more AFO moulding ‘in house’ in xxxxxxxx, the sample size target of 150 subjects was not met. One hundred and fifty one individuals were approached and 134 subjects were recruited into the study (two participants were inadvertently randomised twice; in both cases the first allocation was used). The flow of subjects and numbers randomised into each group, including those that received the planned intervention, are presented in Figure 1. The majority of subjects were affected by cerebral palsy with no statistically significant differences between groups (Table 1). Ninety-six (72%) patients required bilateral AFOs while assistive devices were used by 42% of the patients.
Ages ranged from 1.5 to 21.3 years with a mean age of 10.7 years (SD 4.9) and 9.8 years (SD 4.1) for males and females respectively. Fifty-nine girls and 75 boys were evenly distributed between the two allocations.

Protocol Deviations
A variety of factors including inability to scan a child’s limb; technical problems with the scanning equipment and errors in allocating the manufacturing method correctly in the factory contributed to a total of 15 crossovers from one arm of the study to the other (Table 2 and Figure 1, Consort diagram). The likelihood of crossover occurring was uninfluenced by the need for a bilateral device; sex; previous serial casting; botulinum toxin treatment or calf surgery. All analyses have been performed on an intention to treat basis.

Time Saving and Quality
1. Rectification and moulding of AFO (primary outcome measure)
Data were available for 83 of the 109 AFOs manufactured using the scanning method, and 110 of the 121 AFOs manufactured using the casting method. The mean rectification and moulding time for cast AFOs was 55.1 [SD 26.0] minutes and for LSCAD/CAM AFOs was 26.9 [SD 12.2] minutes. This represented a reduction in time by 28.2 min or 51% in the LSCAD/CAM group (p<.001, two sample t test), 95% CI of difference = 22.6 to 33.8.

2(a). Time taken to scan and cast limbs
All subjects were both cast and scanned and data was available for all 134 subjects. There was no significant difference in the time taken to cast or scan the limbs (p=0.056, paired t test) with the mean times being 12.5 minutes (SD 4.9) and 11.1 minutes (SD 9.5) for cast and scan
respectively, and a mean difference of 1.33 [SD 10.07], 95% CI of paired difference = -0.4 to 3.1.

Evaluation of average casting time throughout the study revealed no training effect or trend in casting time. However, a striking difference was seen over time in the time taken to scan as a result of unplanned staff turnover (Figure 2). If we only include scan times for the two orthotists who had received more training and were already familiar with the technique prior to the start of recruitment, a statistically significant difference (p<.001, two sample t-test) was then seen with average times 13.1 (SD 4.4) minutes for casting and 8.9 minutes (SD 2.9) for scans.

2(b) Supply and fit of AFO(s)

At the second visit the AFO(s) were provided to the patient. A significant proportion (75%) of patients guessed their actual method of fabrication correctly (p<.001, Chi sq test). There were no significant differences in delays in supply between the two methods (p=0.711, logrank test). A significantly higher proportion of scan-based AFOs failed to meet the specification stipulated by the scanning orthotist, see Table 3 (p=0.004, Chi Sq test ). Remaking was required in 17% of scan-based AFOs and 3% of cast-based AFOs (p<.001, Chi Sq test). Whilst none of the cast patients required recasting, 4 scanned patients required a rescan, 1 scanned patient was subsequently cast, and 8 scanned patients required an adaptation of the original scan. Although there was an increased incidence of problems with the scan-based AFOs, the time taken to fit the device was not significantly different between the groups.

2(c) Time to Deliver Device

Data were available for 68 of the subjects allocated to the cast group and 62 subjects allocated to the scanned group. No significant effects were seen (p=.12 logrank test) of allocated group
on the time taken from casting/scanning to delivery of a well-fitting AFO/AFOs. The mean (SD) number of days was 46.9 (45.3) and 37.9 (31.9) for scan and cast respectively with 95% CI for the difference (scan – cast) of -4.5 to 22.5. The time from receipt of the order in the manufacturing unit to the date the order was completed was close in the two groups (13 days ±6 in the cast group, 15 days ±8.2 in the scanned group).

2(d). AFO Length of Life

The comparison of length of AFO life was carried out using Kaplan-Meier survival plots (Figure 3) and a logrank test. This is due to the fact that times to failure are only recorded up to one year and hence those surviving beyond one year are censored. 29 of the 70 subjects allocated to casting and 29 of the 64 subjects allocated to scanning returned to clinic for a new AFO within the first year, and around half of these were as a result of growth (see Table 4). No significant difference was seen in the length of life of AFOs manufactured using the casting or scanning method (p=.57, logrank test).

Patient Preference / Satisfaction

After experiencing both casting and scanning (in random order) 70% of the patients said they preferred being scanned to having the limbs cast in plaster.

A Mann Whitney U test to evaluate differences in responses to the four-point Likert scale (Strongly Agree – Strongly Disagree) found no significant differences between the allocated groups when applied individually to the nine “Satisfaction with Device” questions and the 10 “Satisfaction with Service” questions at each time point (3, 6, & 12 months follow-up). “Agree” was the predominant category selected by respondents for all questions thus indicating a reasonable level of satisfaction with the AFOs and services.
Economic Results

The economic results for plaster versus scan show no advantage of scan over plaster in terms of cost-effectiveness. The average total direct cost of providing an AFO (excluding capital costs) was £586 in the scanning group and £416 in the casting group. This includes consumables, overheads and staff costs for assessments (scanning=£98, casting=£86), manufacture (£364, £206) and fitting and adjustments (£120, £117).

The cost of the scanner was taken as £15,000. We assumed that the scanner would completely depreciate in value over 8 years, and that it would be used for the production of around 775 AFOs per year, with 96% of the scanner's use attributable to AFOs. A discount rate of 3.5% was applied. This gives an equivalent annual capital cost per AFO produced of just £2.69. The standard cost of manufacture for a single AFO was £168.66 for scan-based devices and £90.90 for cast-based devices, though some individuals required multiple AFOs and re-manufacturing. The average total societal cost for individuals in the scanning group was £2859, and for those in the casting group it was £2824. The observed incremental cost of scanning for individuals with complete cost and outcome data is £70. For individuals with complete data, the incremental effect of scanning is an increase in the time to delivery of 10 days. Traditional casting is found to strongly dominate laser CAD/CAM. The observed ICER is therefore negative, with a 95% bias-corrected and accelerated bootstrapped confidence interval of -£128 to £1.

DISCUSSION

Synopsis of key findings

The study does not support the introduction of laser scanning and LSCAD/CAM manufacture for AFO provision in children and young people on the basis of time savings or improved quality of
device. Despite a reduction of around 50% in the time spent in the rectification and moulding processes, this benefit was not transferred to the patient where an average increase in the time to delivery of the devices of 9 days was seen for LSCAD/CAM. This delay was probably due to the proportion of devices failing to meet the required specification being considerably higher after manufacture by LSCAD/CAM than after plaster-casting, and thus requiring a re-make and lengthening the time to delivery. However, there is potential for improvement in the design of AFOs manufactured by LSCAD/CAM, and this study has highlighted the importance of training for orthotists in this technique.

**Pros and Cons of LSCAD/CAM compared to plaster casting**

Anecdotally several patients reported that they believed their AFO was produced by a scan because it fitted so well. Conversely, excessively intimate fits, particularly around the malleoli, make donning difficult and spasticity or calf tightness complicates the process. An intimate fit also presents problems with the longevity of the device in the presence of growth. During the study a policy was adopted of enlarging the scan by 1% in all dimensions; this was found to give a closeness of fit comparable to a conventionally fabricated device to give room for growth.

There is clear potential for improvements in the design of AFOs manufactured by LSCAD/CAM. The inability of the orthotist to correctly position the patient during the scanning process makes the correction of postural deformity totally dependent on the rectification process whereas in the process of casting the foot can be held in a corrected position rendering the rectification process less complex and critical.

Re-making is a costly process. Given that 17% of the scanned patients required a re-scan, re-cast or adaptation of the original scan, versus 3% of the plastered patients, plastering is again
the favourable option. Where a re-make is required the hospital bears the staff, capital and consumable costs of this, whilst the families incur increased travel costs and productivity losses.

In addition, there is a large difference in the manufacturing costs between AFOs made by scanning and by casting, with scanning being over £70 more expensive.

Being able to store the scan for enlargement in response to growth does offer a potential advantage in terms of a reduction in return visits for families if a previous scan can be stretched and an AFO fabricated. However, given the increased frequency of rescans and defects seen in this study, the cost of such a strategy would be currently unacceptable and the time saving for families would be reduced.

Reduction in cost due to time-savings in rectification and moulding may be negated by an increase in costs due to problems at delivery of device (e.g. re-scanning).

Technical difficulties with the hardware and software led to occasional failures of scanning, particularly early on in the study when the technology was relatively new. Whilst the trained orthotists were able to scan in a consistently short period of time, the locums and new staff members frequently had prolonged setup times and were uncomfortable with the process. A study by Telfer et al. demonstrated that only the more experienced CAD operator was able to achieve excellent inter- and intra-operator reliability. However, in this study the imperfections in the resulting AFOs were seen equally in scans performed by “trained” orthotists as well as those not previously trained in the use of the technology. The consistent times for casting across all orthotists and throughout the study demonstrated the ubiquitous nature of the manual skill of casting instilled in trainee orthotists in the United Kingdom.

Limitations of the study
The study had a number of limitations resulting from factors associated with the pragmatic nature of the trial and unexpected difficulties encountered during the study. Whilst the personnel disruption encountered during the study might seem to detract from the information derived, ABLE was a pragmatic trial of a technology intended to evaluate a ‘real-life’ situation for immediate use in patients.

Since the study started in 2008, technology has progressed and more advanced scanners and fabrication methods are available. The study failed to recruit the target number of 150 participants. However, the eventual sample size was sufficient to achieve 80% power.

A high number of participants successfully guessed the manufacturing method applied to their AFOs (75%). The majority of participants in this study were used to having cast AFO's; thus, many could detect the subtle differences in intimacy of fit of AFOs produced by LSCAD/CAM. The digital nature of the scanning process allows the initial assessment of the patient to occur remote to the manufacturing facility with the time and expense of transport of fragile casts being eliminated by electronic transmission of the digital mesh. This cost is not applicable at our centre due to the availability of an on-site facility, but would be more relevant where peripatetic orthotic services are in place.

CONCLUSIONS
This study demonstrated that compared with conventional casting techniques laser scan based AFO manufacture neither significantly improved the quality of the final product nor delivered a useful saving in time. Time savings of around 30 minutes were achieved during the rectification and moulding processes in the LSCAD/CAM group, and although this may have implications for staff costs, this would be of no real benefit to the patient. Significant improvements to the quality
of devices manufactured using the LSCAD/CAM method would be required before any overall benefit would be seen.

ACKNOWLEDGEMENTS
All authors contributed equally in the preparation of this manuscript. xxxxxxxxxxx coordinated subject recruitment and undertook the data collection and collation. xxxxxxxxxxx and xxxxxxxxxxx managed the orthotic workshop and implemented the randomisation provided by the York Trials Unit. The orthotists taking part in the trial included xxxxxxxx, xxxxxxxxxxxxxx, xxxxxxxxxxxxxx, xxxxxxxxxx, xxxxxxxxxx, xxxxxxxxxx and xxxxxxxxxx. xxxxxx staff, xxxxxxxxx and xxxxxxxxxx, offered xxxxxxx facilities to support the study.

DECLARATION OF CONFLICTING INTERESTS
The Authors declare that there is no conflict of interest.

FUNDING ACKNOWLEDGEMENT
This work was supported by the National Institute for Health Research reference number PB-PG-0706-10626.

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REFERENCES


FIGURES

Figure 1. Consolidated Standards of Reporting Trials (CONSORT) flow chart. [insert figure 1, pdf]

Figure 2. Trends in first visit scan time during study. Δ indicates the point during the study when the orthotists experienced in the use of the scanner left the Trust and were replaced with orthotists inexperienced in the use of the scanner. [insert figure 2]

Figure 3. Kaplan-Meier plot of time to failure (days) of the device for the two groups. Grey line = AFOs manufactured by casting method, black line = AFOs manufactured using LSCAD/CAM [insert figure 3]
TABLES

Table 1: Recruitment allocation by diagnostic category

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<th>Main Diagnosis</th>
<th>Cast</th>
<th>Scan</th>
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Table 2: Crossovers from allocated method to other method

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<th>Allocated scan</th>
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Table 3: Incidence of problems at delivery of first device

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<td>%</td>
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<td>%</td>
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<td>Total number of subjects for which data is available for</td>
<td></td>
</tr>
<tr>
<td>Remake needed</td>
<td>2/69</td>
<td>3</td>
<td>11/63</td>
<td>17</td>
</tr>
<tr>
<td>Recast / Rescan</td>
<td>0/69</td>
<td>0</td>
<td>5/63</td>
<td>8</td>
</tr>
<tr>
<td>Lack of intimate fit</td>
<td>7/69</td>
<td>10</td>
<td>13/62</td>
<td>21</td>
</tr>
<tr>
<td>Difficulty donning</td>
<td>3/69</td>
<td>4</td>
<td>10/62</td>
<td>16</td>
</tr>
<tr>
<td>Poor quality finish</td>
<td>1/69</td>
<td>1</td>
<td>5/62</td>
<td>8</td>
</tr>
<tr>
<td>Trim lines incorrect</td>
<td>8/68</td>
<td>12</td>
<td>15/62</td>
<td>24</td>
</tr>
<tr>
<td>Footplate incorrect</td>
<td>40/69</td>
<td>58</td>
<td>47/63</td>
<td>75</td>
</tr>
<tr>
<td>Height incorrect</td>
<td>5/69</td>
<td>7</td>
<td>6/62</td>
<td>10</td>
</tr>
<tr>
<td>Pitch incorrect</td>
<td>0/69</td>
<td>0</td>
<td>5/62</td>
<td>8</td>
</tr>
<tr>
<td>Adjustments needed</td>
<td>34/69</td>
<td>49</td>
<td>37/62</td>
<td>60</td>
</tr>
</tbody>
</table>

Table 4: Reasons for replacement of first AFO

<table>
<thead>
<tr>
<th>Reason for Replacement</th>
<th>Cast</th>
<th>Scan</th>
</tr>
</thead>
<tbody>
<tr>
<td>Outgrown</td>
<td>18</td>
<td>13</td>
</tr>
<tr>
<td>Needs different design</td>
<td>4</td>
<td>6</td>
</tr>
<tr>
<td>Poor fit</td>
<td>4</td>
<td>6</td>
</tr>
<tr>
<td>Too uncomfortable</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Broken</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Worn out</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Not made to specification</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Total</td>
<td>29</td>
<td>29</td>
</tr>
</tbody>
</table>
Enrollment
Assessed for eligibility, n = 153
Excluded, n = 17
- Not meeting inclusion criteria, n = 3
- Declined to participate, n = 5
- Other reasons, n = 9
  (n=3: staffing issues, n=1: patient un-cooperative unable to scan, n=4: MKS not needed for the patient, n=1: patient did not arrive for appi)
Randomized (n=136)

Allocation
Allocated to cast, n = 71
- Received allocated intervention, n = 67
- Randomisation twice, thus excluded, n = 1
- Did not receive allocated intervention, n = 2
  (crossed over into scan group)
  (Reason - n=2: Human error)

Allocated to scan, n = 55
- Received allocated intervention, n = 51
- Randomisation twice, thus excluded, n = 1
- Did not receive allocated intervention, n = 13
  (crossed over into cast group)
  (Reason - n=2: Insufficiency of randomisation to interpret the scan, n=6: Scan poor quality due to patient movement, n=1: Software/protocol training issue, n=4: Loss of scan data, n=3: Unknown reason)

Lost to follow-up, n=2
3m Follow-Up
Lost to follow-up, n=2
Data missing for this time-point, n=1
6m Follow-Up
Lost to follow-up, n=2
Withdrawn, n=1
9m Follow-Up
Lost to follow-up, n=3
Withdrawn, n=1
Withdrawn, n=1
12m Follow-Up
Lost to follow-up, n=5
Withdrawn, n=1
Figure 2:
Figure 3

Cumulative probability of survival of device

Scan group

Cast group

Time (days)