Effectiveness of instrumented gait analysis in interdisciplinary interventions on parents' perception of family-centered service and on gross motor function in children with cerebral palsy: a randomized controlled trial

Esmann Fonvig, Christina; Rasmussen, Helle Mätzke; Overgaard, Soren; Holsgaard-Larsen, Anders

Published in:
B M C Pediatrics

DOI:
10.1186/s12887-020-02315-2

Publication date:
2020

Document version
Final published version

Document license
CC BY

Citation for published version (APA):
Esmann Fonvig, C., Rasmussen, H. M., Overgaard, S., & Holsgaard-Larsen, A. (2020). Effectiveness of instrumented gait analysis in interdisciplinary interventions on parents' perception of family-centered service and on gross motor function in children with cerebral palsy: a randomized controlled trial. B M C Pediatrics, 20, [411]. https://doi.org/10.1186/s12887-020-02315-2

Terms of use
This work is brought to you by the University of Southern Denmark through the SDU Research Portal. Unless otherwise specified it has been shared according to the terms for self-archiving. If no other license is stated, these terms apply:

- You may download this work for personal use only.
- You may not further distribute the material or use it for any profit-making activity or commercial gain
- You may freely distribute the URL identifying this open access version

If you believe that this document breaches copyright please contact us providing details and we will investigate your claim. Please direct all enquiries to puresupport@bib.sdu.dk
Effectiveness of instrumented gait analysis in interdisciplinary interventions on parents’ perception of family-centered service and on gross motor function in children with cerebral palsy: a randomized controlled trial

Christina Esmann Fonvig 1,2*, Helle Mätzke Rasmussen 1,2, Søren Overgaard 1,2 and Anders Holsgaard-Larsen 1,2

Abstract

Background: Children with cerebral palsy often exhibit an altered gait pattern; however, it is uncertain whether the use of an instrumented gait analysis in interdisciplinary interventions affects the perceived experience of family-centered service (FCS) and/or gross motor function. The aim of this study is to investigate whether individually tailored interdisciplinary interventions, based on an instrumented gait analysis report, has a superior effectiveness on perceived FCS and gross motor function in children with cerebral palsy, compared to ‘care as usual’ without the use of instrumented gait analysis. Furthermore, to investigate potential associations between perceived FCS and gross motor function improvement with the goal of improving future therapy on gross motor function.

Method: This is a sequel analysis on tertiary outcome measures from a prospective, single blind, randomized, parallel group study including two groups of 30 children aged 5–8 years with spastic cerebral palsy at Gross Motor Function Classification System levels I-II (n = 60). The intervention group underwent a three-dimensional gait analysis, from which a clinical report was written with recommendations on interdisciplinary interventions, such as physical therapy, orthopedic surgery, orthotics or spasticity management.

To assess effectiveness on perceived FCS and gross motor function, at baseline, 26 weeks and 52 weeks, the five domains in the Measure of Processes of Care (MPOC-20) (Enabling and partnership, Providing general information, Providing specific information about the child, Respectful and supportive service, and Coordinated and comprehensive care) and the Gross Motor Function Measurement (GMFM-66) were used as outcome measures.

(Continued on next page)
Background

Cerebral palsy (CP) is an umbrella term for a range of conditions caused by a non-progressive injury in the developing fetal or infant brain, resulting in permanent disorders of the development of movement and posture, causing activity limitations [1]. With a prevalence of 2–3 per 1000 live births, CP is the most common disabling impairment amongst children [2, 3].

Although 50–80% of children with CP are capable of walking independently, gait is usually affected as a result of musculoskeletal deformity, abnormal muscle tone, inadequate balance and impaired motor control [4]. As a result, the following impairments may be observed: drop foot during swing phase, plantarflexion of foot in stance, leg length discrepancy or restricted joint movement due to muscle contracture [5].

To monitor and improve the quality of healthcare for children with CP, standardized clinical examinations are offered throughout childhood as a means of early detection of complications [6]. The hospital pediatric departments offer interdisciplinary consultations, where children with CP, their families and the local healthcare team, consisting of professionals from the municipal and regional healthcare systems, meet and agree on future surveillance, coordinate common goals and plan interdisciplinary interventions for the child [6]. However, the current standard care does not include systematic evaluation of the child’s gait-related impairments.

Family-centered service (FCS) is considered best practice in providing healthcare services to children and their families, and is associated with improved parent satisfaction within healthcare services, strengthening of family empowerment, and improved development and functioning of the child [7, 8]. One possible way to support an FCS approach in children with CP may be to provide the family and the healthcare personnel with a detailed description of the child’s gait-related impairments along with specific recommendations regarding tailored interventions, thus warranting the involvement of the family in the planning of their child’s treatment.

However, in the current standard care for children with CP, which does not include systematic evaluation of the child’s gait-related impairments, it is uncertain what the family’s overall perception of FCS is and to which degree interventions are agreed upon in collaboration with the child’s parents at interdisciplinary consultations.

As a means of clarifying the intervention needs of a child with CP, an evaluation of gait-related impairments by an instrumented gait analysis (IGA) can be used to provide objective and valid measures of gait. This analysis identifies features affecting gross motor function and underlying neuro-musculoskeletal impairments [4].

Studies show that surgeons follow the IGA recommendations 42–97% of the time [9–11] and in cases where surgery had already been decided, the addition of an IGA has been shown to change 40–89% of surgical treatment plans [9, 12–14]. Thus, IGA is considered a valuable tool in clinical decision making. However, while it has not been previously investigated, it is also reasonable to expect that the addition of an IGA report could support an FCS approach, as the IGA report supplies the families with detailed information on their child’s gait as well as recommendations (in layman’s terms) for interventions. Thus, this report could allow them to participate in the planning and tailoring of specific interdisciplinary interventions in conjunction with physiotherapists, orthopedic surgeons, neuropediatricians and other healthcare professionals.

Gross motor function can be used as a proxy measure for overall improvement during intervention [15]. A potential association between improvement in parents’ perceptions of the FCS they and their children received during the study period and the child’s gross motor improvement could indicate that gross motor function is related to the parents’ experiences of, and involvement in, the processes of care. Thus, an association could serve as an indicator for healthcare personnel allowing them to detect families who may benefit from enhanced FCS.

This study aimed to test the hypothesis that the use of an IGA report in the decision making of interdisciplinary interventions for children with CP, has a superior effectiveness on the parents’ perceived experience of FSC and
on the child’s gross motor function compared to ‘care as usual’, without the use of an IGA report.

Furthermore, to test if potential associations between perceived experience of FCS and gross motor function exists with the perspective of detecting components of the FCS experience worthwhile targeting to potentially improve gross motor function.

Methods
This study is a sequel analysis (on tertiary outcome measures) of a randomized controlled trial based on a dataset powered for a different question [16, 17]. The study is approved by the Committee for Medical Research Ethics in the Region of Southern Denmark (S-20120162) and the Danish Data Protection Agency (2008-58-0035) and is compliant with the Declaration of Helsinki. Tertiary outcome measures for the present analysis were a priori defined at ClinicalTrials.gov (NCT02160457) [17]. Data collection was performed August 2014 to July 2017. Reporting followed the CONSORT statement for pragmatic trials [18].

One hundred and sixty children with CP from the Regions of Southern Denmark and North Denmark were screened for eligibility and invited to participate, of whom 60 were randomized to either the experimental or the control group after baseline assessment. Randomization and allocation is described in detail elsewhere [16]. In brief, the allocation sequence was computer generated by a researcher with no other involvement in the study. The allocation sequence was concealed in sequentially numbered, opaque, sealed envelopes. After completion of baseline assessment, the project manager of the trial (HMR) opened the envelope and informed the parents and the local team about the allocation [16, 17].

Inclusion criteria were children aged five to 8 years with a diagnosis of spastic CP at Gross Motor Function Classification System (GMFCS) level I-II [19]. Exclusion criteria were 1) orthopedic surgery within 52-weeks prior to baseline assessment, 2) injection with botulinum toxin type A within 12-weeks prior to baseline assessment, 3) inability to participate in examination, or 4) lack of ability to speak and understand Danish. Parents signed informed consent following the receipt of verbal and written information at the Motion Analysis Laboratory at Odense University Hospital [16].

Administration of data collection was performed blinded. The participants (children with CP and their parents) and the local health teams were unblinded to the intervention.

Intervention
The interventions under investigation were: 1) individually tailored interdisciplinary intervention based on recommendations from measures performed as part of standard care as well as from an IGA report and 2) individually tailored interdisciplinary intervention based on clinical examinations WITHOUT an IGA report (‘care as usual’).

The IGA was performed at baseline, and the report was subsequently sent to the families as well as the treating healthcare personnel so that the information was available at the following interdisciplinary consultations held at the pediatric departments.

The interdisciplinary interventions recommended in the IGA report have been described in detail elsewhere [17]; but in brief they were divided into the following four categories: (1) physical therapy, (2) orthotics, (3) spasticity management, and (4) orthopedic surgery. The recommended interventions were provided by the child’s local healthcare team regardless of allocation group. Due to the study’s pragmatic approach, interventions made at multiple departments and/or clinics were not standardized, nor was adherence to the recommended interventions a requirement. For an anonymized example of an instrumented gait analysis report, including interdisciplinary interventions, see Additional file 1: IGA report.

Outcome measures
Height and weight, used for the calculation of Body Mass Index Standard Deviation Score (BMI SDS), were measured at baseline. CP subtype was collected from the local healthcare team.

To assess the degree of perceived FCS, responses to the Danish version of the Measure of Processes of Care (MPOC-20) questionnaire [20], were obtained at baseline, 26 weeks and 52 weeks post start of intervention. The primary endpoint of the present sequel analysis was the change-score in the perception of FSC from baseline to 52 weeks.

The MPOC-20 is a self-report measure of the parents’ perceptions of the extent to which the health services their child receive are family-centered [21]. The 20 items cover issues of five selected domains; Enabling and partnership, Providing general information, Providing specific information about the child, Respectful and supportive service and Coordinated and comprehensive care. This tool is widely used in pediatric rehabilitation to evaluate FCS [7]. The clinometric properties of the MPOC-20 have shown solid internal consistency, test-retest reliability and good evidence of construct validity [21].

To assess gross motor function, the Gross Motor Function Measure (GMFM-66), a gold standard measure of functional ability in the area of CP [22], was performed at baseline and at 52 weeks post start of intervention. A minimum requirement of 13 relevant items of the GMFM-66 were used for a valid calculation of the GMFM-66 using the Gross Motor Ability Estimator.
Scoring Software [23]. The GMFM-66 is valid and reliable (ICC = 0.99) [23] and is sensitive enough to document clinically meaningful improvement, including responsiveness [24].

**Statistical analysis**

Study power considerations for this tertiary analysis were evaluated a posteriori and showed that if at least 56 participants completed the trial, it would yield a power between 65 and 92% to detect a between-group difference \( p < 0.05 \) of 1 MPOC point with a SD of 1.19–1.82 on the MPOC scale ranging from 0 to 7 (where higher scores indicate that the parents perceive that their needs are being better met). Since no anchor-based minimum clinically important difference (MCID) is available for the MPOC-20, the present study determined one MPOC point as being clinically relevant, as it would increase the scale score of the present sample to what has been seen in international references [7].

The statistical analysis was a priori defined at Clinical-Trials.gov (NCT02160457) [17] and was performed according to intention-to-treat and, due to low expectation of drop-out, with last-observation-carried-forward for missing observations. Descriptive statistics were summarized as appropriate. Evaluating superiority was estimated with a linear model, with between-group change differences as dependent variable and randomization group as independent variable, adjusted for relevant baseline score. Furthermore, categorical analyses investigating the risk difference (RD) between groups in scoring above an MCID for the MPOC-20 and GMFM-66 were used to calculate a possible Number Needed to Treat (NNT). Finally, associations between the MPOC-20 and GMFM-66 were evaluated using multiple linear regressions with A) the MPOC-20 domains at 52–week follow-up as the dependent variable, and the GMFM-66 change score at the 52–week follow-up as the independent variable, adjusted for baseline GMFM-66 score, sex, age, BMI SDS, CP spastic subtype, and GMFCS level, as well as with B) the GMFM-66 change score at the 52–week follow-up as the dependent variable and the MPOC-20 domains at the 52–week follow-up as the independent variable, adjusted for baseline MPOC-20 domain score, sex, age, BMI SDS, CP spastic subtype and GMFCS level. Effect sizes are reported as partial Eta2 for the independent variable.

Statistical analyses were performed using Stata/IC 14.2 or later for Windows (StataCorp LLC, College Station TX, USA). Results are presented with an alpha of 0.05 and a 95% confidence interval.

**Results**

Of the 160 children invited to participate in this study, 83 were screened for eligibility, of which 60 were randomized to either the experimental group \( n = 30 \) or the control group \( n = 30 \). For the primary outcome, complete assessments were available from 56 participants at baseline, 46 participants at the 26-week follow-up, and 54 participants at the primary endpoint of 52 weeks (Fig. 1).

The 60 included children had a median age of 6 years and 10 months (Table 1). The CP subtypes were 43 children with unilateral spastic CP (experimental \( n = 21 \) / control \( n = 22 \)), and 17 children with bilateral spastic CP (experimental \( n = 9 \) / control \( n = 8 \)). GMFCS levels were 42 children at level I (experimental \( n = 20 \) / control \( n = 22 \)), and 18 children at level II (experimental \( n = 10 \) / control \( n = 8 \)) (Table 1).

As previously stated [16], no serious adverse events were reported during the study period.

Opposed to the hypothesis, the present study found no superior effectiveness in FSC evaluated by MPOC-20 domains at 52-weeks follow-up (Enabling and partnership: \( p = 0.02 \) [95%CI: \( -0.88 \) to \( -0.92 \)], \( p = 0.97 \), Providing general information: \(-0.16 \) [95%CI: \(-1.00 \) to \(-0.69 \)], \( p = 0.71 \), Providing specific information about the child: -0.39 [95%CI: \(-1.33 \) to \(-0.54 \)], \( p = 0.40 \), Respectful and supportive service: 0.14 [95%CI: \(-0.56 \) to \(-0.83 \)], \( p = 0.70 \), Coordinated and comprehensive care: 0.04 [95%CI: \(-0.62 \) to \(-0.69 \)], \( p = 0.92 \)) (Table 2). Furthermore, no between-group changes were observed at 26-weeks follow-up (p-values ranging from 0.26–0.97 for the five domains) (Table 2). Statistical analyses between weeks 26 and 52 were calculated post hoc and showed no significant difference when looking at within or between group changes (data not shown). The categorical analysis investigating the number of participants scoring above the a posteriori set MCID of 1 MPOC point at the 52-week follow-up, showed no significant differences between the two groups in any of the five domains (Table 3).

A statistically significant between-group change score in the GMFM-66 in favor of the experimental group (3.05 [95%CI: 1.12–4.98], \( p = 0.003 \)) was observed at the 52-week follow-up (Table 2). On the individual level, the number of children that had a gross motor function improvement above the defined MCID of 1.7 [24] was 16/30 (53%) in the experimental group and 6/30 (20%) in the control group; this difference was in favor of the experimental group (RD = 0.33 [95%CI: 0.1–0.56], \( p = 0.004 \)) with an NNT of 3.

The association analyses showed no indication of either a uni- or bidirectional relationship between the parents’ general perceived experiences of FCS over the past year and the child’s gross motor function improvement (Tables 4 and 5).

**Discussion**

In this secondary analysis on tertiary data from a randomized controlled trial, we observed no superior
effectiveness in parents’ perceived experience of FSC at the 52-week follow-up (primary endpoint) or the 26-week follow-up, when providing an IGA report as part of the interdisciplinary interventions for children with CP. Hence, the results cannot support an implementation of IGA as part of the standard care for children with CP at GMFCS level I–II solely with the intention of improving perceptions of FCS. However, a superior improvement in gross motor function was observed in the experimental group, suggesting that the use of an IGA in addition to ‘care as usual’, may benefit gross motor function. However, these findings need to be confirmed in a priori designed studies with the specific purpose of detecting changes in gross motor function.

Although studies have shown, that the use of IGA can alter the clinical decisions about planned surgical interventions [9, 26], the present study, on a sample of children with CP showed that it did not alter the parents’ perceived experiences of FCS.

The idea of adding an IGA report with specific recommendations for interventions for the child was to supply the parents, as well as the healthcare personnel, with a detailed, objective explanation of the impairments present in order to improve the observed gait difficulties. However, to comply with current clinical practice, the present pragmatic study design did not include a thorough oral explanation of the IGA report itself; thus, we do not know whether the parents and/or healthcare personnel understood or even read the report.

Generally, all MPOC-20 domains observed in the present study were evaluated one point below international reference scores [7], indicating generally low perceptions of FCS in the treatment of children with CP in Denmark. Therefore, there is room for improvement for the present healthcare personnel using an FCS...
approach, and the lack of effectiveness observed in the present study cannot be attributed to a ceiling effect. A previous study investigating parents’ perceived experiences of FCS in three municipalities in Denmark [27] reported similar low scores in three of the five domains. This emphasizes the necessity to involve the families in the decision-making process for their child’s care and also highlights the parents’ strong desire for both specialized knowledge about children with CP in general and specific knowledge concerning their own child.

### Table 1 Demographics and baseline data of included children

|                          | Experimental group |                       | Control group |                       |
|--------------------------|--------------------|-----------------------|---------------|-----------------------|
|                          | n = 30             |                       | n = 30        |                       |
| Sex boys/girls, n        | 21/9               |                       | 18/12         |                       |
| CP spastic subtype       | 21/9               |                       | 22/8          |                       |
| GMFCS level I/II, n      | 20/10              |                       | 22/8          |                       |
| Age years, months, median (IQR) | 6y 6 m (2y 8 m) |                       | 6y 11 m (1y 10 m) |                       |
| Height m                 | 1.12 (0.08)        |                       | 1.24 (0.11)   |                       |
| Weight kg, median (IQR)  | 22 (6)             |                       | 21 (11)       |                       |
| BMI SDS                  | −0.32 (1.46)       |                       | −0.62 (1.42)  |                       |

Measure of Processes of Care - 20 domains

|                                      | Experimental group |                       | Control group |                       |
|--------------------------------------|--------------------|-----------------------|---------------|-----------------------|
|                                      | n = 28             |                       | n = 29        |                       |

Table 2 Within- and between-group differences at the 26-week and 52-week follow-ups

|                                      | 26-weeks follow-up (mean change from baseline) |                  | p-value | Effect size (Eta²) | 52-weeks follow-up (mean change from baseline) | p-value | Effect size (Eta²) |
|--------------------------------------|-----------------------------------------------|------------------|---------|--------------------|-----------------------------------------------|---------|--------------------|
|                                      | EXPERIMENTAL (mean ± SE) (within-group)       | CONTROL (mean ± SE) (within-group) | Between-group difference in change (mean [95% CI]) |         | EXPERIMENTAL (mean ± SE) (within-group)       | CONTROL (mean ± SE) (within-group) | Between-group difference in change (mean [95% CI]) |         |
| enabling and partnership             | −0.03 ± 0.22                                   | 0.27 ± 0.30      | −0.17 [−0.84–0.50] | 0.43 | 0.004                                 | −0.14 ± 0.36                                   | 0.08 ± 0.39      | 0.02 [−0.88–0.92] | 0.97 | 0.000                                 |
| providing general information        | −0.06 ± 0.33                                   | 0.04 ± 0.33      | 0.09 [−0.80–0.97] | 0.83 | 0.001                                 | −0.41 ± 0.33                                   | 0.03 ± 0.35      | −0.16 [−1.00–0.69] | 0.71 | 0.002                                 |
| providing specific information about the child | −0.40 ± 0.28                                   | 0.06 ± 0.30      | −0.10 [−0.92–0.71] | 0.26 | 0.001                                 | −0.80 ± 0.43                                   | 0.18 ± 0.31      | −0.39 [−1.33–0.54] | 0.40 | 0.014                                 |
| respectful and supportive service    | 0.23 ± 0.21                                    | 0.04 ± 0.24      | 0.03 [−0.57–0.64] | 0.97 | 0.000                                 | −0.09 ± 0.27                                   | −0.16 ± 0.26     | 0.14 [−0.56–0.83] | 0.70 | 0.003                                 |
| coordinated and comprehensive care   | 0.16 ± 0.20                                    | 0.09 ± 0.19      | 0.13 [−0.42–0.69] | 0.79 | 0.004                                 | −0.11 ± 0.21                                   | −0.05 ± 0.29     | 0.04 [−0.62–0.69] | 0.92 | 0.000                                 |
| GMFM-66                               | −3.32 ± 0.84*                                   | 0.16 ± 0.48      | 3.05 [1.12–4.98] | 0.003 | 0.145                                 |

Values are reported as mean and standard deviation (SD) unless stated otherwise

Abbreviations: CP Cerebral Palsy, GMFCS Gross Motor Function Classification System, BMI SDS Body Mass Index Standard Deviation Score showing the BMI percentile and z-score [25], n number, IQR interquartile range, m meters, kg kilogram

Explanations: †Primary outcome measure, *Within-group difference in change score p < 0.001, a Experimental group, n = 28 and control group, n = 28, b Experimental group, n = 28 and control group, n = 29, c Experimental group, n = 30 and control group, n = 30

Between-group differences are adjusted for relevant baseline score

Abbreviations: MPOC-20 Measure of Processes of Care, GMFM-66 Gross Motor Function Measure

Explanations: †Primary outcome measure, *Within-group difference in change score p < 0.001, a Experimental group, n = 28 and control group, n = 28, b Experimental group, n = 28 and control group, n = 29, c Experimental group, n = 30 and control group, n = 30
The present lack of change in the perception of FCS could be because gait function was predefined without the involvement of the families. The parents may have preferred to focus on other areas, e.g., general health, self-care, communication, or caregiver issues [15]. Thus, the inclusion of an IGA report with the intention of improving gait may not necessarily have been a prioritized need of the families.

A clinically relevant between-group change score in the GMFM-66 was detected in favor of the experimental group (3.05 [95% CI: 1.08–5.02] at the 52-week follow-up, resulting in an NNT of 3. However, this is in divergence to the main study of the present trial, which found no improvement in overall gait function [16]. The calculation of the GMFM-66 in the present study was based upon a minimum of items; consequently, the results should be interpreted with caution, taking into consideration the risk of a Type I error. Furthermore, as with most clinical trials, a potential selection bias could influence the results.

The present study found no association between the measured change score in the GMFM-66 and the follow-up score in the MPOC-20. Hence, the hypothesis that an association between these items could be an indicator of families that would benefit from enhanced FCS as a means of improving the child’s gross motor function could not be confirmed. However, the present results from this singular, tertiary trial on the potential improvement in perceived FCS by adding IGA to the decision making should not be considered an exhaustive conclusion on the topic.

**Strengths and limitations**

The external validity of the study is strong due to the pragmatic study design with generalizability to the Danish population of 5–8-year-old children with CP at GMFCS level I–II. However, the results may not be generalizable to older children with CP or to children with more profound disabilities. Furthermore, the risk of selection bias is unknown, as data on non-participants (n = 84) is unavailable. This should be taken into account.

The present study design did not allow the parents to attend a consultation where they were individually informed about the content of the IGA report by the researchers who evaluated and recommended the interventions. Doing so could potentially have helped the parents understand the contents of the report, thereby providing them with specific information about their child and increasing their ability to be critically involved in the planning of their child’s treatment. No data of the parents’ psychological health was collected, and thus it could not be evaluated.

Another limitation is that an anchor-based MCID for the MPOC-20 is not available, and, consequently, the cut-off point of one MPOC-20 point may be considered

| Table 3 | Categorical analysis of participants scoring above an MCID of 1 MPOC-20 point |
|-----------------|-----------------|-----------------|-----------------|-----------------|
| MPOC-20 domains | EXPERIMENTAL | CONTROL | RD | [95%CI] | p-value |
| enabiling and partnership | 6/28 | 7/28 | −0.04 | [−0.26–0.19] | 0.75 |
| providing general information | 7/28 | 4/29 | 0.11 | [−0.09–0.32] | 0.28 |
| providing specific information about the child | 4/28 | 8/29 | −0.13 | [−0.34–0.07] | 0.21 |
| respectful and supportive service | 6/28 | 2/29 | 0.15 | [−0.03–0.32] | 0.11 |
| coordinated and comprehensive care | 3/28 | 5/28 | −0.07 | [−0.07–0.11] | 0.44 |

Values reported as amount in numbers, risk difference (RD), and 95% confidence interval (CI)

Abbreviations: MCID Minimal Clinically Important Difference, MPOC-20 Measure of Processes of Care

| Table 4 | Associations between MPOC-20 and GMFM-66 |
|-----------------|-----------------|-----------------|-----------------|-----------------|
| Independent variable: MPOC-20 52-weeks follow-up domain scores | Independent variable: ΔGMFM较差 scores | (mean ± SE) | (95%CI) | p-value | Effect size (Eta²) |
| enabling and partnership | 0.10 ± 0.05 | [−0.01–0.20] | 0.066 | 0.044 |
| providing general information | 0.01 ± 0.06 | [−0.10–0.12] | 0.816 | 0.001 |
| providing specific information about the child | 0.002 ± 0.06 | [−0.12–0.12] | 0.975 | 0.000 |
| respectful and supportive service | 0.07 ± 0.04 | [−0.02–0.15] | 0.106 | 0.032 |
| coordinated and comprehensive care | 0.05 ± 0.04 | [−0.03–0.13] | 0.214 | 0.016 |

Dependent variable: MPOC-20 domains at the 52-week follow-up
Independent variable: GMFM-66 change score at the 52-week follow-up
Adjusted for baseline GMFM-66 score, sex, age, BMI SDS, CP spastic subtype, and GMFCS level

Abbreviations: GMFM-66 Gross Motor Function Measure, GMFCS Gross Motor Function Classification System, MPOC-20 Measure of Processes of Care, CI confidence intervals, SE standard error
Table 5 Associations between MPOC-20 and GMFM-66

| Dependent variable: ΔGMFM-66 score | Independent variable: MPOC-20 52-weeks follow-up domain scores | (mean ± SE) | [95%CI] | p-value | Effect size (Eta²) |
|------------------------------------|---------------------------------------------------------------|-------------|---------|---------|-------------------|
| enabling and partnership           | 0.66 ± 0.39                                                   | [−0.14−1.45] | 0.103   | 0.070   |
| providing general information      | 0.27 ± 0.41                                                   | [−0.56−1.10] | 0.511   | 0.003   |
| providing specific information about the child | −0.27 ± 0.41                                             | [−1.10−0.55] | 0.510   | 0.002   |
| respectful and supportive service  | 0.61 ± 0.44                                                   | [−0.28−1.50] | 0.175   | 0.055   |
| coordinated and comprehensive care | 0.23 ± 0.34                                                   | [−0.46−0.91] | 0.512   | 0.034   |

Adjusted for relevant MPOC-20 domain score, sex, age, BMI SDS, CP spastic subtype, and GMFCS level

Abbreviations: GMFM-66 Gross Motor Function Measure, GMFCS Gross Motor Function Classification System, MPOC-20 Measure of Processes of Care, CI confidence intervals, SE standard error

arbitrary despite being based upon reference values in the literature [7]. Furthermore, three out of five post hoc power analyses for MPOC-20 domains were underpowered for effects in between-group FCS change. Additionally, statistical results were not corrected for multiple testing, as the study is based on tertiary outcomes [28]. However, no statistical differences in between-group change scores were observed, and the observed wide 95% confidence intervals demonstrated with a clear statistical certainty that the intervention does not provide superior effectiveness on parents’ perception of FCS. A primary analysis on an anchor-based MCID for the MPOC-20 is needed to evaluate clinically relevant changes in perception of FCS over time.

Finally, as the present study is not powered to detect a between-group difference in GMFM-66, it is necessary to confirm the above-mentioned results in an a priori designed study with the primary purpose to investigate the use of an IGA report in interdisciplinary intervention on gross motor function improvement.

Conclusions
In this secondary analysis on tertiary data from a randomized controlled trial, the addition of an IGA in an individually tailored interdisciplinary intervention showed no superior effectiveness in improving the parents’ perceived experiences of FCS in a relatively young group of children with CP at GMFCS level I-II. The results can therefore not support an implementation of IGA as part of standard care with the sole intention of improving the parents’ perception of FCS. However, the addition of IGA to ‘care as usual’ may benefit the child’s gross motor function. These findings need to be confirmed in a priori designed studies with the specific purpose of detecting change in gross motor function. The study could not detect an association between perception of FCS and gross motor function.

In perspective, the results from the present study warrant further investigation into the reasons behind the relative parental dissatisfaction with FCS, and into which steps can be addressed in the clinical decision making process to improve parents’ perception of FCS when planning interdisciplinary approaches for improved gait function. This would provide clinical knowledge about how to ensure that the parents’ needs for specialized information and involvement are more appropriately met in the future.

Supplementary information
Supplementary information accompanies this paper at https://doi.org/10.1186/s12887-020-02315-2.

Additional file 1.

Abbreviations
BMI SDS: Body Mass Index Standard Deviation Score; CP: Cerebral palsy; FCS: Family-centered service; GDI: Gait Deviation Index; GMAE-2: Gross Motor Ability Estimator Scoring Software; GMFCS: Gross Motor Function Classification System; GMFM-66: Gross Motor Function Measure; MCID: Minimum clinically important difference; MPOC-20: Measure of Processes of Care; IGA: Three-dimensional instrumented gait analysis.

Acknowledgements
Niels Wisbech Pedersen is acknowledged for his contribution to the academic and logistic setup of the study as well as his great influence on the gait analysis reports.

Authors’ contributions
Conceptualization and design of the study: HMR, AHL & SO. Collection and assembly of data: HMR & CEF. Analysis of data: CEF, HMR & AHL. First draft of manuscript: CEF. Critical revision of manuscript for important intellectual content and approval of final version: all authors.

Funding
This study was supported by grants from the University of Southern Denmark, Odense University Hospital Research grants (2012), the Region of Southern Denmark Research grants and Ph.D. grants (2012), the Physiotherapy Practice Foundation, Ludwig and Sara Elsass Foundation, the Danish Physiotherapy Research Fund (2013) and the Linex Foundation (2015). The funders had no role in the design of the study, or with regards to collection, analysis or interpretation of data or in the writing of the manuscript.

Availability of data and materials
The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Ethics approval and consent to participate
The study is approved by the Committee for Medical Research Ethics in the
Region of Southern Denmark (S-20120162). Parents signed informed consent
following the receipt of verbal and written information at the Motion
Analysis Laboratory at Odense University Hospital [16].

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

Received: 11 May 2020 Accepted: 24 August 2020
Published online: 01 September 2020

References
1. Rosenbaum P, Paneth N, Leviton A, Goldstein M, Bax M, Damiano D, et al. A
report: the definition and classification of cerebral palsy April 2006. Dev
Med Child Neurol Suppl. 2001;1098–14.
2. Ulldal P, Michelsen S, Topp M, Madsen D. The Danish cerebral palsy
registry. A registry on a specific impairment. Dan Med Bull. 2001;48(3):161–3.
3. Osikou M, Coutinho F, Dykeman J, Jette N, Pringsheim T. An update on the
prevalence of cerebral palsy: a systematic review and meta-analysis. Dev
Med Child Neurol. 2013;55(6):509–19.
4. Gage JR, Schwartz MH, Koop SE. Identification and treatment of gait
problems in cerebral palsy. London: Mac Keith Press; 2009.
5. Armand S, Decoulon G, Bonnefoy-Mazure A. Gait analysis in children with
cerebral palsy. EJORT Open Rev. 2016;1(2):48–60.
6. Rasmussen HM, Nordbye-Nielsen K, Møller-Madsen B, Johansen M,
Ellitsgaard N, Pedersen CR, et al. The Danish cerebral palsy follow-up
program. Clin Epidemiol. 2016;8:457–60.
7. Almasri NA, An M, Palisano RJ. Parents’ perception of receiving family-
centered Care for Their children with physical disabilities: a meta-analysis.
Phys Occup Ther Pediatr. 2018;38(4):427–43. https://doi.org/10.1080/
01942638.2017.1337646.
8. King G, Ciarello L. Family-centered care for children with cerebral palsy:
conceptual and practical considerations to advance care and practice. J
Child Neurol. 2014;29(8):1046–54.
9. Loftroed B, Terjesen T, Skareet I, Huse AB, Jahnsen M. Preoperative gait
analysis has a substantial effect on orthopedic decision making in children
with cerebral palsy: comparison between clinical evaluation and gait
analysis in 60 patients. Acta Orthop. 2007;78(1):74–80.
10. Wren TA, Wooff K, Kay RM. How closely do surgeons follow gait analysis
recommendations and why? J Pediatr Orthop B. 2005;14(3):202–5.
11. Wren TA, Otsuka NY, Bowen RE, Scaduto AA, Chan LS, Dennis SW, et al.
Outcomes of lower extremity orthopedic surgery in ambulatory children
with cerebral palsy with and without gait analysis: results of a randomized
controlled trial. Gait Posture. 2013;38(2):236–41.
12. DeLuca PA, Davis RB 3rd, Ounpuu S, Rose S, Sikrin R. Alterations in surgical
decision making in patients with cerebral palsy based on three-dimensional
gait analysis. J Pediatr Orthop. 1997;17(5):608–14.
13. Kay RM, Dennis S, Rethfilsen S, Reynolds RA, Skaggs DL, Tolo VT. The effect
of preoperative gait analysis on orthopaedic decision making. Clin Orthop
Relat Res. 2000;372:217–22.
14. Cook RE, Schneider I, Hazlewood ME, Hillman SJ, Robb JE. Gait analysis alters
decision-making in cerebral palsy. J Pediatr Orthop. 2003;23(3):292–5.
15. Vargus-Adams JN, Martin LK. Measuring what matters in cerebral palsy: a
breath of important domains and outcome measures. Arch Phys Med
Rehabil. 2009;90(12):2089–95.
16. Rasmussen HM, Pedersen NW, Overgaard S, Hansen LK, Dunkhase-Heini U,
Petkovic Y, et al. Gait analysis for individually tailored interdisciplinary
interventions in children with cerebral palsy: a randomized controlled trial.
Dev Med Child Neurol. 2019;61(10):1189–95.
17. Rasmussen HM, Pedersen NW, Overgaard S, Hansen LK, Dunkhase-Heini U,
Petkovic Y, et al. The use of instrumented gait analysis for individually
tailored interdisciplinary interventions in children with cerebral palsy: a randomised
controlled trial protocol. BMC Pediatr. 2015;15(1):202.
18. Zwartenstein M, Treweek S, Gagnier JJ, Altman DG, Tunis S, Haynes B.
Improving the reporting of pragmatic trials: an extension of the CONSORT
statement. BMJ. 2008;337:a2590.
19. Palisano R, Rosenbaum P, Bartlett D, Livingston M. Gross Motor Function
Classification System Expanded and Revised: CanChild Centre for Childhood
Disability Research Institute for Applied Health Sciences, McMaster
University; 2007 [Available from: https://canchild.ca/system/tenon/assets/
attachments/000/000/058/original/GMFM-ERO_English.pdf].
20. CanChild. MPOC–20 2019 [Available from: https://www.canchild.ca/en/
shop/1-m poc–20].
21. King S, King G, Rosenbaum P. Evaluating health service delivery to children
with chronic conditions and their families: development of a refined
measure of processes of care (MPOC–20). Child Health Care. 2004;33(1):35–7.
22. Lundkvist Josenby A, Jamali G-B, Gummesson C, Nordmark E. Longitudinal
construct validity of the GMFM-88 Total score and goal Total score and the
GMFM-66 score in a 5-year follow-up study. Phys Ther. 2009;89(4):342–50.
23. Russell DJWM, Rosenbaum PL, Avery LM. Gross motor function measure
(GMFM–66 & GMFM–88) User’s manual. 2nd ed. London: Mac Keith Press;
2013.
24. Alobaidi M, Long T, Kennedy E, Bavischi S. The efficacy of GMFM–88 and
GMFM–66 to detect changes in gross motor function in children with
cerebral palsy (CP): a literature review. Disabil Rehabil. 2014;36(6):617–27.
25. Philadelphia TCsHo. Pediatric Z-score Calculator 2018 [Available from:
https://zscore.research.chop.edu/].
26. Loftroed B, Terjesen T. Results of treatment when orthopaedic surgeons
follow gait-analysis recommendations in children with CP. Dev Med Child
Neurol. 2008;50(7):503–9.
27. Nielsen K, Sørensen SH, Dunkhase-Heini U, Rasmussen HM. Intern
evaluating. Opfølgningsprogram og Indsatser for cerebral parese, CPOP–t:
Udvikling af tilbud om tværfagligt og tværs境界lt koordineret indsats til
børn med cerebral parese. 2012.
28. Ranstam J. Multiple P-values and Bonferroni correction. Osteoarthr Cartil.
2016;24(5):763–4.

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published
maps and institutional affiliations.