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Published in:
B M J Open

DOI:
10.1136/bmjopen-2017-021358

Publication date:
2018

Document version
Publisher's PDF, also known as Version of record

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Conservative care with or without manipulative therapy in the management of back and/or neck pain in Danish children aged 9–15: a randomised controlled trial nested in a school-based cohort

Kristina Boe Dissing,1 Jan Hartvigsen,1,2 Niels Wedderkopp,3,4 Lise Hestbæk1,2

ABSTRACT

Background A substantial number of children experience spinal pain, that is, back and/or neck pain. Today, no ‘gold-standard’ treatment for spinal pain in children exists, but manipulative therapy is increasingly being used in spite of a lack of evidence of its effectiveness. This study investigates the effectiveness of adding manipulative therapy to other conservative care for spinal pain in a school-based cohort of Danish children aged 9–15 years.

Methods and findings The design was a two-arm pragmatic randomised controlled trial, nested in a longitudinal open cohort study in Danish public schools. 238 children from 13 public schools were randomised individually from February 2012 to April 2014. A text message system and clinical examinations were used for data collection. Interventions included either (1) advice, exercises and soft-tissue treatment or (2) advice, exercises and soft-tissue treatment plus manipulative therapy. The primary outcome was number of recurrences of spinal pain. Secondary outcomes were duration of spinal pain, change in pain intensity and Global Perceived Effect. We found no significant difference between groups in the primary outcome (control group median 1 (IQR 1–3) and intervention group 2 (IQR 0–4), p=0.07). Children in the group receiving manipulative therapy reported a higher Global Perceived Effect: OR 2.22, (95% CI 1.19 to 4.15). No adverse events were reported. Main limitations are the potential discrepancy between parental and child reporting and that the study population may not be comparable to a normal care-seeking population.

Conclusions Adding manipulative therapy to other conservative care in school children with spinal pain did not result in fewer recurrent episodes. The choice of treatment—if any—for spinal pain in children therefore relies on personal preferences, and could include conservative care with and without manipulative therapy. Participants in this trial may differ from a normal care-seeking population.

Trial registration number NCT01504698; Results.

INTRODUCTION

Spinal pain is common in children and adolescents and prevalence rates reach adult levels already around the age of 18.1 For most children, episodes are transient and inconsequential, and therefore, the area has been largely ignored in research. However, some children have frequent, recurrent and bothersome complaints,2–5 impacting their mental well-being6 and with the potential to decrease the level of physical activity. Importantly, these problems seem to track into adulthood, that is, the most affected adolescents grow up to be the most affected adults.7 8 Therefore, proper management at an early stage is essential to improve lifetime trajectories of spinal pain.

Management of children’s musculoskeletal disorders relies to a large extent on parents’ values, preferences and experience, and due to absence of guidelines for the treatment of spinal pain in children, healthcare
professional therapy (MT) is defined as joint manipu-
lation and/or mobilisation with the aim to restore
compromised function of joints. This type of therapy
is increasingly being used in children because it is
generally recommended as a treatment option for adults
with spinal pain, and is delivered by various health
professions, both on its own and in combination with
other types of therapy, such as advice, exercises and soft-
tissue treatment. One study recently demonstrated a
small but statistically significant effect of adding MT to
exercise therapy in adolescents with low back pain.
However, this is the only full-scale randomised controlled
trial (RCT) conducted to date to investigate the effect of
SMT in children with any type of spinal pain.

The aim of this pragmatic RCT was to determine the
effectiveness of adding MT to other conservative care
(advice, exercises and soft-tissue treatment) on the
number of recurrences of spinal pain in children aged
9–15 years who were participating in a school-based open
cohort study. Secondary outcomes included the short-
term effect on duration of spinal pain episodes, pain
intensity and Global Perceived Effect.

METHOD
Study design
A pragmatic parallel observer-blinded RCT nested in a
school-based open cohort.

Participants and setting
This study was nested in The Childhood Health,
Activity and Motor Performance School Study Denmark
(CHAMPS Study-DK), which is a Danish longitudinal
school-based open cohort study including approximately
1400 children aged 9–15 years from 13 public schools. The
CHAMPS Study-DK was an open cohort study hence chil-
dren could enter or leave the cohort at any time during
the study period. The children were followed weekly with
text messages (SMS) to one of their parents inquiring,
among other things, about any musculoskeletal pain the
child might have had during the past week (questions in
online supplementary file 1). Data collection on musculo-
skeletal complaints for this RCT began in February 2012
and ended at the end of June 2014.

Eligibility determination
All children enrolled in the CHAMPS Study-DK were
invited to participate in the RCT. The complete protocol
for the RCT is described in detail elsewhere. Briefly,
when a parent answered positively on the SMS to the pres-
ence of spinal pain in their child, a member of a screening
team (licensed chiropractors and physiotherapists) tele-
phoned the parent and conducted a standardised inter-
view about the complaint, in order to determine whether
the child was eligible for inclusion in the RCT. Initial
eligibility was based on: (1) the pain was spinal and still
present at the time of the interview, (2) the parent had
agreed, on behalf of the child, to join the RCT and (3) the
child had not had any manual treatment of the spine
during the previous 2 months. Within 2 weeks, the child
was evaluated at the school by a chiropractor from the
RCT team (seven licensed chiropractors) to determine
whether he or she fulfilled the inclusion criteria (table 1).

Table 1 Inclusion and exclusion criteria

<table>
<thead>
<tr>
<th>Inclusion criteria</th>
<th>Exclusion criteria</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pain in neck or back equal to or greater than 3 on an 11-box Numerical Rating Scale for more than 3 days indicated by the child at the first visit.</td>
<td>Serious pathology (cancer, inflammatory diseases, vertebral fractures, cauda equina syndrome).</td>
</tr>
<tr>
<td>Manual treatment for the past 2 months (for this particular complaint).</td>
<td></td>
</tr>
<tr>
<td>Handicaps preventing normal physical activity.</td>
<td></td>
</tr>
</tbody>
</table>

After the evaluation, both the child and his/her parents
were informed about the results and treatment was initi-
ated. The flow from SMS to RCT can be seen in figure 1.

Randomisation
A computer-generated block randomisation was made
with block sizes alternating between two and six at the
time of inclusion, using a 1:1 allocation to the two groups.
The consecutive designations of the two groups were
written on separate pieces of paper and given to the chiro-
practors in the RCT team in sealed opaque envelopes. A
research assistant, who was not otherwise connected to
the study, performed the procedure.

First consultation
At the first consultation, the chiropractor obtained a case
history, including pain intensity on an 11-box Numerical
Rating Scale, performed a clinical examination, and
various baseline data were acquired (online supplemen-
tary file 2). Two weeks after inclusion, the child was asked
about Global Perceived Effect (online supplementary file 3)
and pain intensity.

If a child experienced a recurrence of pain (ie, the
parent-reported pain on the weekly SMS), the proce-
dure was repeated except for randomisation, which was
carried forward throughout the study period regardless
of the body location in which the complaint occurred. All
data were filed in electronic data storage systems.
established specifically for this project and stored on secure servers.

Interventions
The non-MT group received advice, exercises and soft-tissue treatment, and the MT group received advice, exercises and soft-tissue treatment plus MT (table 2).

Both groups were treated by the RCT team consisting of seven chiropractors. MT was defined as high velocity, low amplitude manipulation and/or mobilisation of the joints to restore segmental spinal motion. This was delivered at the discretion of the chiropractor and applied on the basis of a combination of biomechanical dysfunction and pain provocation responses found during the clinical examination of the child, since palpatory findings by itself have been found unreliable. If the child experienced any pain in the extremities during the study period, these were also treated with MT at the discretion of the treating chiropractor. Because of the pragmatic nature of the study, the frequency and content of treatments in both groups was determined by the treating chiropractor at each visit, similar to what is normal in clinical practice. Because the RCT team consisted of seven chiropractors, a child could be treated by different chiropractors during different appointments. Treatments continued until the child no longer had any symptoms related to the musculoskeletal complaint, or until the chiropractor or parent decided that further treatment was not indicated. The child and/or parents could terminate the treatments or drop out of the RCT at any time during the study period, but still stay in the cohort of the CHAMPS Study-DK.

Blinding
Due to the nature of the intervention, blinding of the treating chiropractors was not possible, however, neither parents nor children were informed about group allocation and parents did not attend treatment sessions and answered the SMS without contact with clinicians or researchers. The coding of the intervention group was not revealed to the primary investigator or the statisticians until after the analyses had been completed.

Outcomes
The primary outcome was the number of recurrences as measured via the weekly SMS messages. A recurrence was defined as a new episode of spinal pain (ie, back and/or neck pain) occurring after at least 1 week without spinal pain following the end of the previous episode (see secondary outcomes, table 3).

Sample size
As the study had continuous inclusion, we continued to recruit participants until 3 months prior to the end of data collection in summer 2014, to include as many participants as possible with varying follow-up times. Based on preliminary analyses, this resulted in a power of 76% for the number of recurrences, 20% for episode length and 87% for overall complaint time.

Statistical methods
All analyses used an intention-to-treat approach. Various types of regression analyses were used depending on the type of outcome; follow-up time was included as an exposure time variable; subject was included as a random effect in models with repeated measurements; and class and school were evaluated and included in the models as random effects if their effect was statistically significant (see details, table 3). No effect was seen on any of the outcomes and hence, cluster was not included in the models. For linear models, means and SDs were used if data were normally distributed; otherwise medians and

Table 2 Intervention groups

<table>
<thead>
<tr>
<th>The non-manipulative group</th>
<th>The manipulative group received</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pragmatic advice (activity level, ergonomics, cold packs, etc)</td>
<td>Advice, exercises and soft-tissue treatment</td>
</tr>
<tr>
<td>Exercises (stretching and/or strengthening exercises)</td>
<td>Manipulative therapy: joint manipulation and/or mobilisation</td>
</tr>
<tr>
<td>Soft-tissue treatment (manual trigger point therapy or massage)</td>
<td></td>
</tr>
</tbody>
</table>
Table 3  Outcomes, definitions and statistical methods

<table>
<thead>
<tr>
<th>Definition</th>
<th>Statistical method</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Primary outcome</strong></td>
<td></td>
</tr>
<tr>
<td>Number of recurrences of spinal pain (3–27 months follow-up).</td>
<td>(1) A positive answer on the weekly text message for spinal pain (2) minimum of 1 week without report of spinal pain prior to the recurrence.</td>
</tr>
<tr>
<td><strong>Secondary outcomes</strong></td>
<td></td>
</tr>
<tr>
<td>Average duration of spinal pain episodes.</td>
<td>The number of consecutive weeks the child was affected by spinal pain (response option ‘1’).</td>
</tr>
<tr>
<td>Total duration of complaint time in relation to individual follow-up time.</td>
<td>Total number of weeks a child was affected by spinal pain (response option ‘1’) in the entire follow-up period.</td>
</tr>
<tr>
<td>Global Perceived Effect after 2 weeks.</td>
<td>Dichotomised into two groups: ‘Much better’ and ‘The same or worse’.</td>
</tr>
<tr>
<td>Change in pain intensity after 2 weeks.</td>
<td>Rated on an 11-point Numerical Rating Scale with ‘0’ being ‘no pain’ and ‘10’ being ‘worst pain’.</td>
</tr>
</tbody>
</table>

IQRs were reported. All methods were checked according to fulfilment of other assumptions and changed where appropriate. Due to some missing SMS answers, we imputed missing data as follows: if four or fewer consecutive missing answers were preceded and followed by a ‘1’, this was considered as one continuous episode and the missing values were imputed as ‘1’. Since this type of outcome measure has not been used in previous trials, there is no consensus on how to substitute data. In a previous article, we have described the consequences of different data substitution strategies.

A sensitivity analysis was conducted to assess the effect of the choice of definitions in relation to recurrence and duration in the present study. In this analysis, a new episode was defined to occur after 4 weeks of ‘no pain’ instead of 1 week before it was considered a new episode.

STATA V.14.2 (StataCorp) was used for data analyses. Significance level was set to 5%.

A child could be withdrawn from the study at any time during the study period and the study was conducted according to the Declaration of Helsinki.

Patient and public involvement
There was no patient involvement in the formulation of the research question, the choice of outcome measures, the design, the recruitment procedures, conduct of the study or assessment of the burden of the intervention.

Parents of the included children will receive information about the study and its results via newsletters and the project’s website.

RESULTS
The inclusion period ran from 1 February 2012 to 1 April 2014, and the follow-up period ended on 27 June 2014 (the end of the school year). Follow-up time was defined as ‘Number of days between inclusion date and last SMS’. Since one child left the study the day after inclusion, this resulted in 1 to 868 follow-up days, (mean 477 days; SD 233). A total of 770 children reported spinal pain on SMS, and after telephone interviews, 483 children were evaluated for eligibility but did not fulfil the inclusion criteria. Additionally, 44 individuals reported pain less than 3 on the Numerical Rating Scale on the day of examination, leaving 243 children randomised and enrolled in the study. During data cleaning, we found five participants had been wrongly included, that is, the SMS answer indicated no spinal pain, and they were excluded from the analyses. Thus, the final cohort for analysis consisted of 238 children with a mean age of 12.6 years: 116 in the non-MT group (49%) and 122 in the MT group (51%), (see figure 2).

Baseline covariates can be seen in table 4, which also reports the amount of missing data for each variable. There was no difference between the groups for any of the covariates indicating randomisation was successful, and therefore, univariate analyses were performed for all analyses.

Primary outcome
During the follow-up period, 175 (74%) of the children had a total of 592 recurrences, ranging from 1 to 21 recurrences per child. The median number of recurrences was 2 (IQR 0–4) for the MT group and 1 (IQR 1–3) for the non-MT group, revealing no statistically significant difference between groups, incidence rate ratio 1.26 (95% CI 0.98 to 1.61), p=0.07.

Secondary outcomes
We found no significant difference in the average episode length, total number of pain weeks or change in pain intensity between the two groups. Children in the group receiving MT reported a higher Global Perceived Effect: OR 2.22, (95% CI 1.19 to 4.15), that was statistically significant. All results are displayed in table 5.
**Figure 2** CONSORT flow diagram. CONSORT, Consolidated Standards of Reporting Trials; MT, manipulative therapy; NRS, Numerical Rating Scale; SMS, text message.
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Table 4  Baseline data and covariates by intervention group

<table>
<thead>
<tr>
<th></th>
<th>Non-MT group (n=116)</th>
<th>MT group (n=122)</th>
<th>Missing non-MT group*</th>
<th>Missing MT group*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sex, female, no (%)</td>
<td>73 (63)</td>
<td>78 (64)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age at inclusion</td>
<td>12.6 (12.4 to 12.9)</td>
<td>12.6 (12.3 to 12.9)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Follow-up time (days)</td>
<td>492 (448 to 536)</td>
<td>463 (423 to 504)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pain intensity at baseline (NRS)</td>
<td>5.3 (5.1 to 5.6)</td>
<td>5.2 (4.9 to 5.5)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Expecations of the clinical course ('Worse')</td>
<td>7.6% (3.4% to 16.1%)</td>
<td>7.6% (3.4% to 16.1%)</td>
<td>32% (37)</td>
<td>35% (43)</td>
</tr>
<tr>
<td>KID physical well-being</td>
<td>44.7 (38.5 to 49.6)</td>
<td>43.8 (40.5 to 49.6)</td>
<td>4% (5)</td>
<td>1% (1)</td>
</tr>
<tr>
<td>KID psychological well-being</td>
<td>49.5 (44.8 to 56.0)</td>
<td>48.5 (44.8 to 56.0)</td>
<td>5% (6)</td>
<td>2% (3)</td>
</tr>
<tr>
<td>KID autonomy and relation</td>
<td>49.5 (45.2 to 55.8)</td>
<td>49.5 (45.2 to 55.8)</td>
<td>4% (5)</td>
<td>2% (3)</td>
</tr>
<tr>
<td>KID social support and peers</td>
<td>53.2 (46.9 to 57.8)</td>
<td>53.2 (46.9 to 57.8)</td>
<td>4% (5)</td>
<td>1% (1)</td>
</tr>
<tr>
<td>KID school</td>
<td>51.1 (45.4 to 58.2)</td>
<td>51.1 (45.4 to 54.4)</td>
<td>4% (5)</td>
<td>1% (1)</td>
</tr>
</tbody>
</table>

*Number of children with missing data according to intervention group.

MT, manipulative therapy; NRS, Numerical Rating Scale.

Table 5  Results on secondary outcomes

<table>
<thead>
<tr>
<th></th>
<th>MT group</th>
<th>Non-MT group</th>
</tr>
</thead>
<tbody>
<tr>
<td>Length of spinal pain episode</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total no of episodes</td>
<td>456 (55%)</td>
<td>374 (45%)</td>
</tr>
<tr>
<td>Median (IQR) (no of weeks)</td>
<td>2 (1–6)</td>
<td>2 (1–5)</td>
</tr>
<tr>
<td>β-coefficient (95% CI)</td>
<td>0.11 (~0.17 to 0.19)</td>
<td></td>
</tr>
<tr>
<td>P value</td>
<td>0.21</td>
<td></td>
</tr>
<tr>
<td>Total duration of complaint time per child</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total no of pain weeks</td>
<td>1–114</td>
<td>1–111</td>
</tr>
<tr>
<td>Median (IQR)</td>
<td>9 (IQR 4–22)</td>
<td>7 (IQR 4–18)</td>
</tr>
<tr>
<td>IRR (95% CI)</td>
<td>1.16 (0.92 to 1.48)</td>
<td></td>
</tr>
<tr>
<td>P value</td>
<td>0.22</td>
<td></td>
</tr>
<tr>
<td>Global perceived effect</td>
<td></td>
<td></td>
</tr>
<tr>
<td>No of children in analysis*</td>
<td>96 (52%)</td>
<td>86 (48%)</td>
</tr>
<tr>
<td>OR (95% CI)</td>
<td>2.22 (1.19 to 4.15)</td>
<td></td>
</tr>
<tr>
<td>P value</td>
<td>0.01</td>
<td></td>
</tr>
<tr>
<td>NRS change</td>
<td></td>
<td></td>
</tr>
<tr>
<td>No of children in analysis*</td>
<td>112 (50%)</td>
<td>111 (50%)</td>
</tr>
<tr>
<td>Mean (SD)</td>
<td>2.2 (2.5)</td>
<td>2.3 (2.7)</td>
</tr>
<tr>
<td>β-coefficient (95% CI)</td>
<td>0.10 (~0.17 to 0.28)</td>
<td></td>
</tr>
<tr>
<td>P value</td>
<td>0.76</td>
<td></td>
</tr>
</tbody>
</table>

*Number of children in analysis of the first episode due to missing data. IRR, incidence rate ratio; NRS, Numerical rating Scale.

Sensitivity analysis on number of pain free weeks

The number of recurrences declined from a total of 592 to 259 when we defined a new episode to occur after 4 weeks of ‘no pain’ instead of 1 week. This, however, did not change the between-group difference on either the primary outcome or most of the secondary outcomes, but it did result in a statistically significant increased length of episode for the MT group, mean 3.5 (3.0–4.0) vs 4.4 weeks (3.8–5.0) and median 2 (1–4) vs 2 (1–4), p=0.045.

Harms

Adverse events can be defined as the sequelae following MT to the spine that are medium to long term in duration, with moderate to severe symptoms, and of a nature that is serious, distressing and unacceptable to the patient and requires further treatment. To our knowledge, no adverse events following MT have been reported in children of this age group.25 However, it is common to experience transient side effects such as temporary reddening or soreness in the area being treated after both soft-tissue treatment and MT.28 Treating chiropractors recorded transient side effects if the child stated these at the consultation, but none were reported and no child was referred to other healthcare providers, including general practitioners, because of adverse events.

DISCUSSION

Adding MT to other conservative care for children reporting spinal pain did not result in fewer recurrences in a school-based cohort of Danish children aged 9–15.
years. Furthermore, the average episode length, total number of pain weeks and change in pain intensity were no different between the groups. However, in the sensitivity analyses, filtering out the frequently recurring episodes, the difference for episode length did become statistically significant. Children randomised to the MT group reported a higher Global Perceived Effect that was statistically significant. Thus, no increased effectiveness was evident and no harm was detected.

To our knowledge, this is the first RCT evaluating the added benefit of MT in children with spinal pain (ie, back and/or neck pain). Michaleff et al. found only four RCTs dealing with conservative interventions for low back pain in children and all had a high risk of bias. Only one of these included manual therapy combined with exercise, but it had only 45 participants.

Because this study was a two-armed parallel trial with MT as an addition to other conservative care, it is probably not surprising that we did not find a large difference between the two groups. This RCT was nested in a large cohort study, and hence we could not prolong the study period to increase the sample size; however, given the small absolute differences found on both primary and secondary outcomes, this is unlikely to have changed our conclusions.

Choice of outcomes

We originally intended to analyse the three spinal regions separately, however, the pain site could change within the same individual during follow-up, and many individuals reported pain from several regions. Therefore, the interpretation of our results relate to ‘spinal pain’ as a coherent entity. We could not determine by the SMS answers whether recurrences were actual recurrences of the same problem at the same location in the spine, but simply conclude that there was subsequent spine-related pain. This can be considered a weakness as we cannot determine true recurrences; however, it can also be considered a strength because pain in this age group appears to demonstrate a shift between regions of the spine over time, indicating that there is not independence between pain in the three regions.

The Numerical Rating Scale has been shown to be a valid tool for assessing pain in children, and in this study, the children also appeared to be able to rate their pain on the scale quite easily. However, when analysing the data, we found that Numerical Rating Scale ratings were not always in accordance with Global Perceived Effect ratings, that is, some children would say they felt better, although reporting a higher score on the Numerical Rating Scale at follow-up than at baseline. This noise may be caused by variation in cognitive abilities and maturity between the children, and is probably equally distributed between groups. Regardless, we did not find statistically significant differences between the groups on change in Numerical Rating Scale scores, and both achieved a mean change of ±2.3, which can be regarded as a clinically meaningful change, as studies have shown a minimal clinically important change to be ±1.32 33

We could not find any literature supporting the validity of measures of Global Perceived Effect in children, but validity of this measure has been shown to be good in adults34 35 and we therefore included it as a measure of the child’s own perception of improvement. We would have expected that statistically significant differences between the groups would follow the same pattern for the Numerical Rating Scale and the Global Perceived Effect, but this was not the case. Therefore, the validity of both of these as outcome measures in clinical trials involving children should be further explored.

Strengths and weaknesses

The principal strength of this study was the school-based design, which had a number of advantages: the logistical burden for the parents was reduced because the treatment took place during school time, social bias was likely to be minimal or absent because everybody was invited to participate in the study, and there was equal access because all treatment in the trial was free. Also, this design allowed for a long follow-up period for most children. By nesting this RCT in a school-based cohort, we may however have included children who would not normally have sought care, that is, likely to have had subclinical pain. The inclusion criterion of a Numerical Rating Scale score of 3 or more on the day of examination is probably also below the normal pain intensity threshold for seeking treatment and many parents would probably have waited until the pain had become worse or lasted longer before seeking care. On the other hand, the number and duration of spinal pain episodes were higher in the study sample than in the full cohort (mean number 3.5 vs 2, mean duration 4.6 vs 2.8), suggesting that the children enrolled in this study were more affected by pain than their non-participating peers.

SMS is a very efficient way of collecting frequent data over a long time.57 58 In this study, the SMS responses were a reflection of how often the parents reported on their child’s pain and might not have been a true reflection of how the child actually felt. We know that there is a discrepancy between parent and child reporting of spinal pain.59 60 Parents appear to under-report compared with their child when pain is at a low level, whereas concordance is higher when the pain is more severe. Thus, it is possible that the parents stopped reporting pain because they assumed the complaint to be minor, even though the child might still have had pain. This could explain some of the difference between outcomes reported by the children (Global Perceived Effect) and outcome reported by the parents (SMS).

Using different practitioners prevents a potential patient–practitioner relationship and is considered a strength; however, the more people involved, the more irregularities and mistakes are likely to occur. One example of this is the poor response rate to the measures.
collected by the clinicians, for example, Numerical Rating Scale and Global Perceived Effect scores.

Missing data
The amount of missing data was substantial for some of the secondary outcomes, and therefore, we analysed only those for the first spinal pain episode. However, there was no difference in response rates between groups, and it was assumed that data were missing completely at random and not due to any underlying confounding factors or bias. Possible reasons for missing data could be practitioners’ forgetfulness or an electronic system defect resulting in missing data. Because of missing data, we cannot say anything valid about the course of pain, for example, whether there is a learning effect over time or whether expectations of treatment differ over time between the two groups.

Future research
Since the inclusion criteria in this study were very broad, subgroup analyses would be valuable to inform future studies, that is, if there are subgroups of children who respond better or worse to MT than to other treatments. Future RCTs should include care-seeking children who self-report their response to treatment in order to evaluate effectiveness in that population. In addition, inclusion of an untreated group would elucidate the effect of treating these children, whether MT is included or not.

CONCLUSION
We found no significant difference in the number of recurrences of episodes of spinal pain in a school-based cohort of children when adding MT to advice, exercises and soft-tissue therapy. The study population may not be comparable to a normal care-seeking population, and therefore, the results may not be directly transferrable.

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Acknowledgements
The authors gratefully acknowledge the Nordic Institute of Chiropractic and Clinical Biomechanics for providing office space and support. Furthermore, we would like to thank Suzanne Capell for proof reading the manuscript. Finally, we would like to thank the participants and their parents and the participating schools, and Werner Vach and Eleanor Boyle for advice in matters relating to sample size calculations and description of the analysis. In addition, we acknowledge all the members of the CHAMPS Study-DK and the clinicians taking part in this study for making it possible.

Contributors
All authors (KBD, JH, NW and LH) participated in the design and interpretation of analyses of this study. KBD was project manager for the trial and drafted the manuscript. All authors (KBD, JH, NW and LH) contributed with revisions and approved the final version of the manuscript.

Funding
This work was supported by The IMK Foundation, The Danish Chiropractic Research Foundation, The Nordea Foundation and The TRYG Foundation, who funded the data collection as well as salaries and equipment for examination and treatment of the children in the RCT. The salary of the first author (KBD) was funded by the Danish Chiropractic Research Foundation and the University of Southern Denmark, in order to complete this project. The other authors did not receive specific grants for this study.

Disclaimer
The funders had no role in the study design, data analysis, decision to publish or preparation of this paper.

Competing interests
None declared.

Patient consent
Not required.

Ethics approval
The project was approved by The Regional Committee on Health Research Ethics (#5-20110042) and data were handled according to the regulations set by the Danish Data Protection Agency (#2013-41-1738).

Provenance and peer review
Not commissioned; externally peer reviewed.

Data sharing statement
Data are from the Childhood Health, Activity and Motor Performance School Study (CHAMPS Study-DK) and are available on request from the project manager Niels Wedderkopp.

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