

Development of a core outcome set for clinical trials in childhood constipation: a study using a Delphi technique

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ABSTRACT

Objective Patients, their parents and healthcare professionals (HCPs) have a different perception regarding the symptoms of functional constipation (FC). Consequently, a lack of agreement exists on definitions and outcomes used in therapeutic trials of FC. Therefore, our aim was to develop a core outcome set (COS) for FC for children aged 0–1 year and 1–18 years.

Design and setting Prospective study design: primary, secondary and tertiary care settings.

Methods This COS was developed using a Delphi technique. First, HCPs, parents of children with FC and patients aged ≥12–18 years were asked to list up to five outcomes they considered relevant in the treatment of FC. Outcomes mentioned by >10% of participants were included in a shortlist. In the next phase, outcomes on this shortlist were rated and prioritised by HCPs, parents and patients. Outcomes with the highest scores were included in a draft COS. In a face-to-face expert meeting, the final COS was determined.

Results The first phase was completed by 109 HCPs, 165 parents and 50 children. Fifty HCPs, 80 parents and 50 children completed the subsequent phase. The response rate was between 63% and 100% in both steps. The final COS for all ages consisted of: defecation frequency, stool consistency, painful defecation, quality of life, side effects of treatment, faecal incontinence, abdominal pain and school attendance.

Conclusion The use of this COS for FC will decrease study heterogeneity and improve comparability of studies. Therefore, researchers are recommended to use this COS in future therapeutic trials on childhood FC.

INTRODUCTION

Functional constipation (FC) is a common problem with a worldwide prevalence of 0.7%–29.6%.¹ Approximately 95% of children with constipation has FC, meaning that no organic or anatomic cause can be identified.

Standard definitions and criteria for FC exist but are rarely used in research and clinical practice.² In 1999, the first diagnostic criteria for paediatric functional gastrointestinal disorders were published: the Rome

What is already known about this subject?

- Heterogeneity in defining and measuring outcomes hampers comparison of efficacy trial results.
- A core outcome set (COS) for functional constipation (FC) can solve this issue.

What this study hopes to add?

- Involvement of healthcare professionals, parents and patients in the development of this COS for FC ensured that appropriate outcomes are measured.
- Final COS: defecation frequency, stool consistency, painful defecation, quality of life, side effects of treatment, faecal incontinence (if age appropriate), abdominal pain (if age appropriate) and school attendance (if age appropriate).

II criteria. Several studies showed, however, that these were too restrictive and were therefore modified into the Rome III criteria (2006) and recently into the Rome IV criteria.^{3–8}

Differences in outcome definitions, measurement and reporting across clinical trials make it difficult to pool study results. Moreover, negative results are less likely to be reported, which can cause bias.⁹ These problems can be tackled by developing agreed standardised sets of outcomes: core outcome sets (COS). COS serve as a guide for what should be measured and reported, but measured outcomes do not need to be restricted to the COS. Researchers can examine additional outcomes that might be of interest to them.^{10–12}

COS increase consensus about the efficacy of an intervention and can improve healthcare, but it needs to contain outcomes that really matter to stakeholders. It is important to involve healthcare professionals (HCPs), parents and patients with FC when developing a COS. Especially since the literature

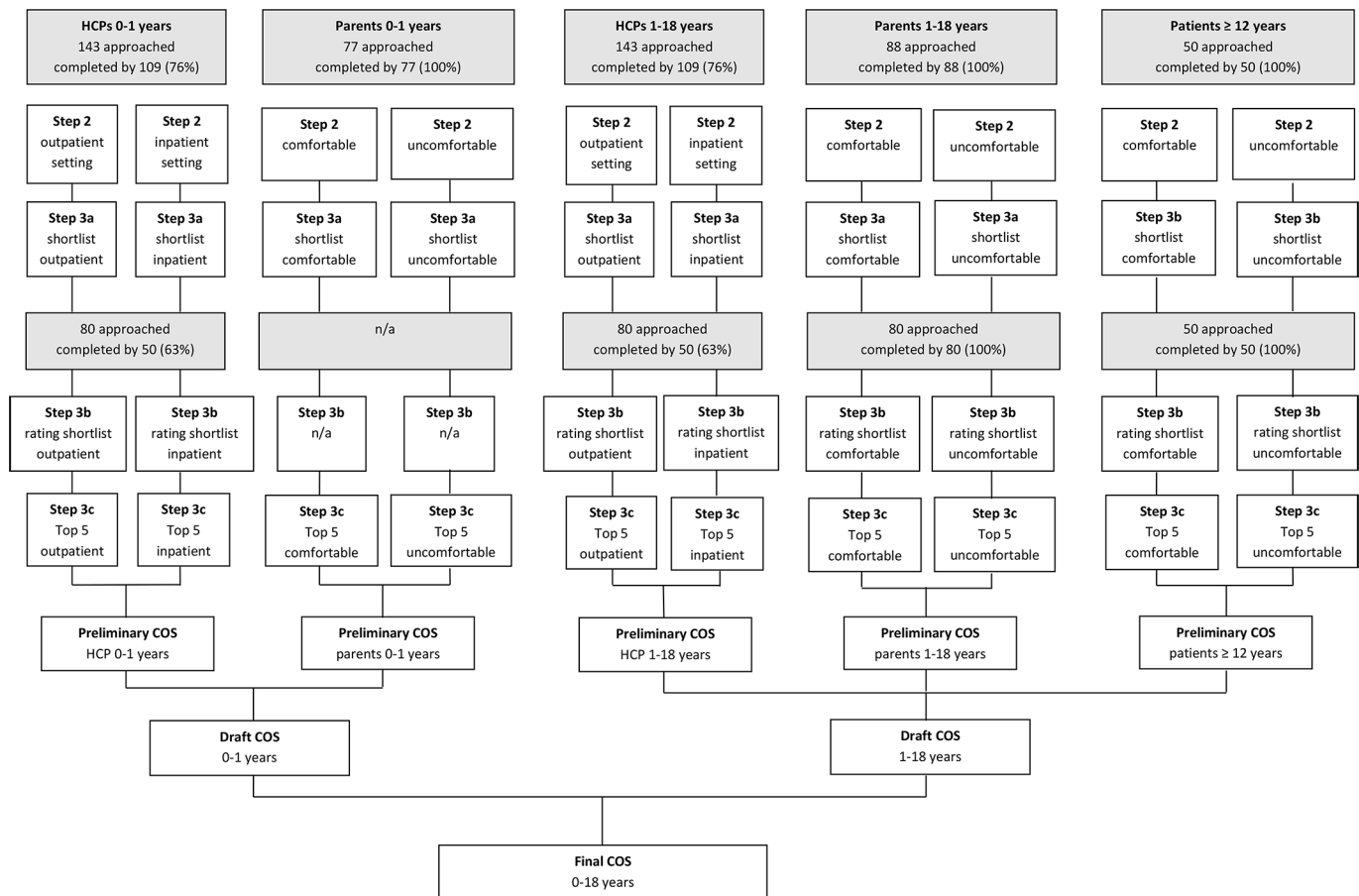


Figure 1 Flow diagram of the core outcome set (COS) development. HCPs, healthcare professionals; n/a, not applicable.

has shown that parents of children with FC have different concerns regarding their child's symptoms than HCPs.¹³

In recent years, several COS were developed for paediatric topics such as acute diarrhoea, infant colic and asthma.¹⁴⁻¹⁶ These studies used the Delphi technique to identify clinically important outcomes for HCPs and parents. Sinha *et al* also involved patients ≥12 years in this process.¹⁶

This work was commissioned by the Consensus Group on Outcome Measures Made in Paediatric Enteral Nutrition Clinical Trials (COMMENT), an initiative of the European Society for Paediatric Gastroenterology Hepatology and Nutrition (ESPGHAN). COMMENT agreed that consensus was needed on core outcomes for FC. Therefore, we aim to develop a COS for therapeutic trials in children with FC, in primary to tertiary care settings.

METHODS

The method for the COS development was based on previous publications.¹⁴⁻¹⁶

The Medical Ethics Review Committee confirmed that the Medical Research Involving Human Subjects Act did not apply to this study and therefore official approval by the committee was not required.

Descriptive statistics were used to summarise results.

In step 1, we systematically assessed how definitions and outcomes were defined in therapeutic randomised controlled trials of children with FC. Results were published in two manuscripts; one concerning children ≤4 years and another concerning children aged 1-18 years.^{2,17}

Table 1 Country of practice of participating HCPs

Country of practice	No of HCPs
Italy	18
Israel	14
UK	10
Poland	9
The Netherlands	7
Mexico/Turkey	4
Belgium/Ecuador/Germany/Greece	3
Brazil/Czech Republic/New Zealand/Norway/Romania/Sweden/Thailand/USA	2
Australia/Canada/Croatia/France/Japan/Jordan/Spain/Philippines/Russia	1
Not reported	6
Total	109

HCPs, healthcare professionals.

Participants

In step 2, we collected important treatment outcomes among stakeholders.

A survey was conducted among HCPs visiting two international paediatric gastroenterology conferences. Participants were randomly approached and asked to list up to five harmful/beneficial treatment outcomes, which they considered important and guided their clinical decision-making in outpatient and inpatient settings, as outcomes could differ in these settings. Data regarding profession and country of practice were collected. We aimed to include ≥ 100 HCPs.¹⁴

We aimed to include 160 parents in four countries (Belgium, Italy, Poland and the Netherlands). Parents of children with FC (according to the Rome III criteria) were randomly approached by their child's treating physician to participate in this anonymous survey.^{5 6} Parents were asked to list up to five treatment outcomes that made them feel comfortable or made them feel that their child was being treated adequately. The same question was asked for treatment outcomes that made parents feel uncomfortable or made them feel their child was being treated inadequately. Questions were translated into the native language of parents and answers were carefully translated back to English by their HCP.

As an anonymous pilot study, we assessed which treatment outcomes were relevant to patients ≥ 12 years, diagnosed with FC according to the Rome III criteria.^{5 16}

We aimed to include 50 patients at the gastroenterology outpatient clinic of the Emma Children's Hospital/Academic Medical Centre, Amsterdam, the Netherlands. Children were randomly invited by their doctor and questions from the parental survey were adjusted for this population.

Outcomes

In step 3, reported outcomes were classified in predefined domains based on our findings in step 1 (defecation, associated symptoms, use of medication, treatment success, quality of life (QoL) and hospital).^{2 17} Domains functioned as subcategories to group outcomes with similar characteristics. When necessary, we added new domains based on survey results. Similar outcomes were combined and outcomes mentioned by $\geq 10\%$ of participants were included in a shortlist. The 10% threshold was chosen by COMMENT to keep shortlists manageable. Ten separate shortlists were created.

Next, shortlists for HCPs were sent to those that participated in the first survey and agreed to participate in the second survey as well. They were asked to rate outcomes on clinical relevance on a scale of 0–4 (0=not relevant, 4=very relevant) and to prioritise outcomes by selecting five outcomes they thought to be most important in guiding their clinical decision-making.

A new group of parents and patients with FC, from the same four countries, was invited to participate in this phase. They received shortlists for parental and patient outcomes, respectively, and were asked to rank outcomes

on a scale of 0–4 (0=does not make me feel (un)comfortable, 4=makes me feel very (un)comfortable) and to prioritise them by selecting the five outcomes that made them feel most (un)comfortable. Surveys were returned anonymously. We aimed to include 80 parents and 50 patients.^{14 16}

Subsequently, the five outcomes with the highest rank were selected for each group. This resulted in preliminary outcome sets for HCPs, parents and patients, for both age groups.

After combining the preliminary outcome sets for each age group, a draft COS was presented to an expert panel during the COMMENT Working Group (WG) meeting at the ESPGHAN meeting in Athens, Greece (2016). Here consensus (a unanimous decision) regarding the final COS was reached by discussion.

RESULTS

Flow diagram of COS development is shown in [figure 1](#). One hundred and nine out of 143 (76%) HCPs completed the first questionnaire. They originated from 28 countries ([table 1](#)) and included 52 paediatric gastroenterologists, 24 general paediatricians, 17 fellows, 4 residents in paediatrics, 4 paediatric nutritionists, 3 researchers, 3 not specified, 1 intern and 1 general doctor. In some originating countries, healthcare is organised in such a way that paediatricians represent primary and secondary care as well.

For infants aged 0–1 years, 89 and 74 different outcomes were reported for outpatient and inpatient settings, respectively. For children aged 1–18 years, 76 and 72 different outcomes were reported for outpatient and inpatient settings, respectively. [Table 2](#) shows an example of reported outcomes.

One hundred and sixty-five parents (100%) of children with FC completed the first questionnaire. Parents of infants aged 0–1 years reported 28 treatment outcomes that made them feel comfortable and 26 outcomes that made them feel uncomfortable. Parents of children aged 1–18 years reported 52 treatment outcomes that made them feel comfortable and 45 that made them feel uncomfortable.

Fifty patients ≥ 12 years (mean age 14.8 years; response rate 100%) completed the first questionnaire. Patients reported 21 outcomes that made them feel comfortable and 22 that made them feel uncomfortable.

[Table 3](#) shows outcomes that were reported most frequently.

Creating a shortlist and final COS

Fifty out of 80 HCPs (63%) completed the second questionnaire regarding the rating and prioritising of outcomes on the shortlist. Since the top 5 of outcomes for outpatient and inpatient settings were rather similar ([tables 4 and 5](#)), we combined these into one preliminary outcome set per age group.

Table 2 Outcome measures as reported by healthcare professionals (HCPs) for the outpatient setting in children 1–18 years; step 2 (n=109)

Subdomain	Outcome	n
Domain: defecation		
Constipation	Duration of constipation	3
	Constipation	2
	Disappearance of constipation	2
Defecation frequency	Defecation frequency	60
	Regular soft bowel movements	1
Stool consistency	Stool consistency	34
	Quality of stools	3
Painful defecation	Pain with defecation	21
Domain: associated symptoms		
Abdominal pain/discomfort	Abdominal pain	42
	Discomfort	14
	No bowel distention	1
	Abdominal bloating	1
Faecal incontinence	Faecal incontinence	28
Bowel habits—other	Blood in stool	9
	Less anxious to defecate	3
	Normal bowel movement	2
	Less stool withholding	2
	Clarity of bowel movement	1
	Amount of stool	1
	Sensation of emptying	1
	Number of days without stool	1
	Emergency defecation on the way to school	1
	Crying	Crying
Straining	Straining	3
Other gastrointestinal complaints	Less gas	1
Domain: quality of life		
Quality of life	Quality of life of child/parents	18
	Reduce stress in family	4
	Reduce parental anxiety	3
	Behaviour/energy/mood of child	3
	Normal social life	1
	Parent's discomfort	1
Domain: treatment success		
Faecal impaction	Rectal impaction	3
	Faecal impaction resolves	1
	Efficiency of disimpaction	1
	No effect of clean out	1
Satisfaction	Satisfaction of child/parent	4
	Patient satisfaction	1
	Parental satisfaction	1
Treatment compliance	Treatment compliance	5

Continued

Table 2 Continued

Subdomain	Outcome	n
Diet	Diet	3
	Improve appetite/normalisation of food intake	3
	Eating problems	2
	Less milk	1
	More fibre	1
	Nutritional status	1
Education	Parental reassurance	1
	Explanation of normal stool pattern	1
	Explanation of long duration of therapy	1
	Re-education	1
	Explanation about the disease	1
Improvement	Time to improvement of symptoms	2
	Improvement of clinical status	1
Adequate relief	Adequate relief rated by patients and parents	1
Lack of success	Lack of success	1
Domain: use of medication		
Medication	Use of laxatives	8
	Characteristics of medication (like taste and side effects)	8
	Re-education of pharmacological treatment	2
	No medication required	2
	Efficacy of medication	2
	Decrease of harmful medication	1
	Already used medication	1
Domain: hospital		
Hospital visits	Fewer consultations with HCPs	3
	Number of visits to outpatient department	1
Domain: other		
Growth	Thriving/growth	5
	Weight loss	1
Other	School attendance	4
	Manage anal fissure	3
	Sleep	2
	Frequency of voiding	1
	Costs of treatment	1
	Presence of alarm symptoms	1
	Adequate nursing by family	1
	Physical examination	1
	Colonic transit time	1
	Decrease of diameter of the rectum	1
	Control of sphincter	1

Three unclear outcomes such as 'familiar environment' and 'organisation' were not displayed.

In step 2, only five outcomes were mentioned by >10% of the parents of infants 0–1 years. It was unnecessary to approach new parents for step 3, as the rating and prioritising would not affect the top 5 outcomes.

All 80 parents of children 1–18 years (100%) completed step 3. Tables 4 and 5 show the shortlists that were rated and prioritised by parents. Since outcomes that made parents feel comfortable and uncomfortable were rather

Table 3 Outcomes most often reported in step 2

Infants 0–1 years		
HCP	Outpatient setting	Inpatient setting
	Defecation frequency (51%)	Defecation frequency (46%)
	Abdominal pain (32%)	Stool consistency (28%)
	Stool consistency (30%)	Abdominal pain (20%)
Parents	Comfortable	Uncomfortable
	Stool consistency (39%)	Stool consistency (56%)
	Defecation frequency (38%)	Straining (38%)
	Crying (17%)	Defecation frequency (34%)
Children 1–18 years		
HCP	Outpatient	Inpatient
	Defecation frequency (55%)	Defecation frequency (46%)
	Abdominal pain (39%)	Abdominal pain (27%)
	Stool consistency (30%)	Stool consistency (26%)
Parents	Comfortable	Uncomfortable
	Defecation frequency (41%)	Abdominal pain (40%)
	Abdominal pain (39%)	Faecal incontinence (25%)
	Stool consistency (25%)	Defecation frequency (25%)
Patients	Comfortable	Uncomfortable
	Abdominal pain (66%)	Abdominal pain (58%)
	Defecation frequency (40%)	Defecation frequency (35%)
	Faecal incontinence (32%)	Faecal incontinence (28%)

HCP, healthcare professional.

similar for this age group, we combined these into one preliminary outcome set.

Fifty patients (mean age 13.8 years; response rate 100%) completed the rating and prioritising of outcomes (table 6). Outcomes that made patients feel comfortable and uncomfortable were rather similar and we combined these into one preliminary outcome set.

The preliminary COS of HCPs and parents of infants 0–1 years old were comparable. Therefore, these were combined into the following draft COS: defecation frequency, stool consistency, painful defecation, discomfort, crying, abdominal pain, treatment success, straining and QoL of parents.

We repeated this process for children 1–18 years but added the outcomes mentioned by patients aged ≥ 12 years. The preliminary COS of HCPs, parents and

patients appeared to be comparable as well, except for the following outcomes: ‘abdominal pain’ (mentioned by parents and patients), ‘treatment success’ (parents) and ‘school attendance’ (patients). We combined outcomes into the following draft COS: defecation frequency, stool consistency, painful defecation, QoL (child and parents), faecal incontinence, abdominal pain, school attendance and treatment success.

The draft COS for both age groups was presented in a COMMENT WG meeting. Regarding the COS for 0–1 years, it was unanimously decided to exclude ‘discomfort’, ‘abdominal pain’ and ‘crying’ in the final COS, since these outcomes are non-specific and cannot with certainty be attributed to FC or the treatment effect. Additionally, it was decided to exclude ‘straining’, since ‘painful defecation’ was already included in the COS. ‘Treatment success’ was excluded as it is assumed to be a combination of the included outcomes.

Regarding the COS for 1–18 years, it was unanimously decided to add two outcomes mentioned by parents/patients: ‘abdominal pain’ and ‘school attendance’. ‘Treatment success’ was excluded for the same reason as above.

Since the applicability of a treatment depends on a balance between benefits and side effects of that treatment, it is recommended to assess ‘side effects’ as part of the COS.¹⁸ Although ‘side effects of treatment’ were mentioned by <10% of respondents, we added this outcome to the COS. The COS for infants 0–1 years therefore includes: defecation frequency, stool consistency, painful defecation, QoL of the parents and side effects of treatment. The COS for children 1–18 years includes: defecation frequency, stool consistency, painful defecation, QoL of patients and parents, faecal incontinence, abdominal pain, school attendance and side effects of treatment.

For practical reasons, we combined these into one final COS for all children aged 0–18 years (box 1).

DISCUSSION

HCPs, parents and patients ≥ 12 years were involved in a Delphi technique to develop a COS for children with FC aged 0–18 years. Stakeholders largely agreed on which outcomes were most important, but some discrepancies existed. Parents and patients found ‘abdominal pain’ more important than HCPs. Although it is not a characteristic feature of FC, 10%–70% of children with FC are suffering from abdominal pain which can have a large impact on their well-being.¹⁹ Abdominal pain can lead to school absenteeism and reduce participation in social activities. Another discrepancy regarded ‘defecation frequency’: depending on the age group, 46%–55% of HCPs found this the most important treatment outcome, whereas parents and patients mentioned this less often as an important outcome.

Interestingly, most Rome IV criteria are represented in our COS, except for ‘presence of a large faecal mass

Table 4 Shortlist outcome measures (rated and prioritised) for infants 0–1 years old

Rank	Outcome measures	Domain	Percentage	Average rating
Healthcare professionals				
Outpatient setting				
1	Defecation frequency	Defecation	96	3.14
2	Painful defecation	Defecation	94	2.98
3	Stool consistency	Defecation	82	3.20
4	Discomfort	Associated symptoms	74	2.28
5	Quality of life (parents)	Quality of life	64	1.76
6	Abdominal pain	Associated symptoms	36	1.44
7	Crying	Associated symptoms	36	0.18
8	Faecal incontinence	Associated symptoms	18	0.76
Inpatient setting				
1	Defecation frequency	Defecation	92	3.18
2	Stool consistency	Defecation	88	3.24
3	Painful defecation	Defecation	84	2.94
4	Discomfort	Associated symptoms	72	2.30
5	Quality of life (parents)	Quality of life	62	2.31
6	Time of hospitalisation	Hospital	52	1.82
7	Abdominal pain	Associated symptoms	28	1.50
8	Faecal incontinence	Associated symptoms	22	0.80
Parents				
Comfortable				
1	Stool consistency	Defecation	N/A	N/A
2	Defecation frequency	Defecation		
3	Crying	Associated symptoms		
4	Abdominal pain	Associated symptoms		
5	Treatment success	Treatment success		
Uncomfortable				
1	Stool consistency	Defecation	N/A	N/A
2	Straining	Associated symptoms		
3	Defecation frequency	Defecation		
4	Crying	Associated symptoms		
5	Painful defecation	Defecation		

N/A, not applicable.

Outcomes ranked 1 to 5 no background colour and outcomes ranked 6 to 8 (if applicable a darker colour to visualise they were excluded from the top 5).

in the rectum', 'retentive posturing' and 'large diameter stools' which were mentioned by 3%, 2% and 0% of HCPs, respectively. Furthermore, 'faecal incontinence' was mentioned as an important outcome for infants aged 0–1 years. A potential explanation for this could be that this was a Rome III criterion for children aged 0–4 years. Since this is very difficult to assess in children wearing diapers, the Rome IV criterion 'faecal incontinence' is only used as an additional criterion to diagnose FC in toilet-trained children.⁸

One of the strengths of this study is the high response rate. Furthermore, participation of physicians around

the globe, involvement of parents from four European countries and inclusion of patients increase the generalisability of our COS. This study has some limitations. First, our study focused on children whose FC is managed by hospital paediatricians. In some participating countries, however, healthcare is organised in such a manner that paediatricians represent primary and secondary care as well. Although we included participants from around the globe, developing countries were under-represented which may limit the external validity of this COS in a developing country setting. Data collection from HCPs occurred at international paediatric gastroenterology

Table 5 Shortlist outcome measures (rated and prioritised) for children 1–18 years old

Rank	Outcome measures	Domain	Percentage	Average rating
Healthcare professionals				
Outpatient setting				
1	Faecal incontinence	Associated symptoms	94	3.63
2	Defecation frequency	Defecation	84	3.50
3	Painful defecation	Defecation	78	3.24
4	Stool consistency	Defecation	78	3.16
5	Quality of life (child/parents)	Quality of life	76	3.31
6	Abdominal pain	Associated symptoms	66	2.60
7	Discomfort	Associated symptoms	24	1.76
Inpatient setting				
1	Faecal incontinence	Associated symptoms	88	3.59
2	Defecation frequency	Defecation	86	3.48
3	Stool consistency	Defecation	78	3.24
4	Painful defecation	Defecation	74	3.18
5	Quality of life (child/parents)	Quality of life	70	2.98
6	Abdominal pain	Associated symptoms	52	2.54
7	Time of hospitalisation	Hospital	40	2.28
8	Discomfort	Associated symptoms	12	1.72
Parents				
Comfortable				
1	Quality of life (child and parents)	Quality of life	83	3.41
2	Abdominal pain	Associated symptoms	77	3.30
3	Faecal incontinence	Associated symptoms	76	3.24
4	Painful defecation	Defecation	66	2.96
5	Stool consistency	Defecation	65	3.16
6	Treatment success	Treatment success	64	3.08
7	Defecation frequency	Defecation	47	2.67
8	Complete information given by doctor	Other	36	2.81
Uncomfortable				
1	Lack of treatment success	Treatment success	88	2.40
2	Abdominal pain	Associated symptoms	84	2.95
3	Stool consistency	Defecation	78	2.57
4	Defecation frequency	Defecation	68	2.40
5	Faecal incontinence	Associated symptoms	62	2.28
6	Lack of information given by doctor	Other	44	1.92

Outcomes ranked 1 to 5 no background colour and outcomes ranked 6 to 8 (if applicable a darker colour to visualise they were excluded from the top 5).

conferences. This may have caused bias since these HCPs often treat patients with severe and long-lasting FC, and they could therefore have chosen different outcomes compared with primary care physicians. Also, patients ≥ 12 years were recruited in a tertiary care setting as a pilot study. Only recruiting patients in such a specialised setting might have led to biased results.

Furthermore, since answers were collected anonymously in both Delphi rounds, we were unable to

assess any potential differences between respondents and non-respondents. The Delphi technique is well suited as a method for consensus building, but it is a time-consuming process, that requires active participation throughout the Delphi process, which can result in high dropout rates.²⁰ However, our response rates were 63%–100% in both steps.

Another potential limitation is the use of questionnaires that were developed in English and thereafter translated into

Table 6 Shortlist outcome measures of patients ≥ 12 years old

Rank	Outcome measures	Domain	Percentage	Average rating
Comfortable				
1	Abdominal pain	Associated symptoms	86	3.52
2	School absence	Treatment success	74	3.14
3	Faecal incontinence	Associated symptoms	66	2.86
4	Quality of life (child)	Quality of life	54	2.94
5	Defecation frequency	Defecation	52	2.58
6	Stool consistency	Defecation	42	2.46
7	Use of medication	Use of medication	16	1.92
Uncomfortable				
1	Abdominal pain	Associated symptoms	86	3.38
2	School absence	Treatment success	70	3.08
3	Faecal incontinence	Associated symptoms	68	3.06
4	Quality of life (child)	Quality of life	50	2.74
5	Defecation frequency	Defecation	48	2.48
6	Stool consistency	Defecation	40	2.42
7	Use of medication	Use of medication	22	2.18

Outcomes ranked 1 to 5 no background colour and outcomes ranked 6 to 8 (if applicable a darker colour to visualise they were excluded from the top 5).

Box 1 Final core outcome set for functional constipation in children 0–18 years

- ▶ Defecation frequency
- ▶ Stool consistency
- ▶ Painful defecation
- ▶ Quality of life of parents and patients
- ▶ Side effects of treatment
- ▶ Faecal incontinence, if age appropriate
- ▶ Abdominal pain, if age appropriate
- ▶ School attendance, if age appropriate

four different languages. Although this was done carefully, it is possible that this resulted in subtle changes in questions or answers. Some responses in step 2 were somewhat open to interpretation; therefore, we may have misinterpreted and incorrectly combined outcomes. Furthermore, not all outcomes mentioned by parents or patients were included in the final COS. The WG unanimously decided that these outcomes were non-specific for FC (crying, discomfort) or already included in the COS (straining, treatment success).

As mentioned before, a COS is the basis for what should be measured and reported in clinical trials, but researchers can add more outcomes if needed.²¹ Selective reporting should be avoided by presenting results for both core outcomes and additional outcomes.

To enable comparison of trial results, there is not only a need for consensus regarding outcomes, we also need homogeneity in used definitions of FC. We

recommend to use the recently published and internationally accepted Rome IV criteria for FC in future trials.⁷

COS have the potential to improve evidence-based healthcare, but their creation does not automatically lead to its implementation and it would therefore be valuable to assess how widely this COS is being implemented.

In conclusion, the COS for childhood FC (0–18 years) consists of defecation frequency, stool consistency, painful defecation, QoL and side effects. Faecal incontinence, abdominal pain and school attendance should be measured if age appropriate. We recommend researchers to use this COS to decrease outcome heterogeneity and improve comparability of study results.

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Competing interests None declared.

Patient consent Obtained.

Provenance and peer review Commissioned; externally peer reviewed.

Data sharing statement Tables with reported outcome measures of step 2 are available from the authors.

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