Adolescent and young adult patients as co-researchers: A scoping review

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Abstract
Background: As part of a research project aimed at evaluating a hospital-based adolescent transition programme, we asked ourselves what is known about the ethical and methodological challenges of research involving adolescent patients as co-researchers. The aim of our review was to summarize empirical evidence and identify knowledge gaps about the involvement of young patients as co-researchers.

Methods: We conducted a scoping review through searches in MEDLINE, EMBASE, PsychINFO, AMED.

Results: We found reports of young patients being actively engaged as co-researchers in any stage of a research project, although commonly they were not involved in every stage. Including young patients as co-researchers is resource demanding and time-consuming. Involving young patients as co-researchers contributes to the fulfilment of their right to participation and may improve the relevance of research. Benefits for the young co-researcher include empowerment, skills building and raised self-esteem. Few authors go into detail about ethical considerations when involving young co-researchers. None of the included articles discuss legal considerations.

Discussion and conclusion: No lists of recommendations are given, but recommendations can be deduced from the articles. There is need for time, funding and flexibility when including young patients as co-researchers. Knowledge gaps concern legal and ethical dilemmas of including a vulnerable group as co-researchers. More reflection is needed about what meaningful participation is and what it entails in this context.

Patient or Public Contribution: This review is part of a research project where the hospital youth council has been involved in discussions of focus area and methods.

KEYWORDS
adolescent patients, co-research, health services, participation
1 | INTRODUCTION

The idea of citizen participation is a little like eating spinach: no one is against it in principle because it is good for you. What Arnebm pointed out in 1969 was that people were talking the talk, but were not necessarily willing to walk the walk of true citizen participation. It was a question of empty ritual versus real power to influence. Arnebm's original typology of the eight-rung ladder of participation illustrates different forms of participation. The ladder is frequently referenced and has been adapted by many, also to health care. It is now usually referred to by five levels: (a) information, (b) consultation, (c) advice, (d) collaboration and (e) control. Only the two last levels—collaboration and control—refer to approaches where power is truly shared.

We are seeing an increasing focus on patient participation in health research, with an increasing number of national guidelines being issued, for example in the UK, in Norway and in Belgium. However, as Malterud and Elvbakken point out, even though patient involvement in research has increased since the beginning of this decade, '[...] this kind of research remains far from standard'. Engaging patients in research on health care and health system issues is challenging for a number of reasons. Abma states 'social conditions for dialogue between patients and researcher are not given and should be sought for'. In other words, in most cases the process does not spontaneously unfold. In addition, there are differences in the terminology used about participation and a need to clarify what concepts one uses. Abma argues that participatory approaches differ in their conceptualizations of participation, rationales of participation, norms and values, as well as their definitions of who is a legitimate participant. Participation at the level of consultation implies a different role to the role at the level of collaboration and control. The term 'co-researcher' was coined by Smith in 1994 about being fully included in the research team, thus referring to the two upper levels of the participation ladder. There are lessons learned from projects that involved patients as co-researchers, that is, with participation at the levels of collaboration and control. These include a need to clarify the roles of stakeholders, to reflect on what it means to be a patient representative, to choose the most suitable engagement methodologies for the project and to provide training and support to the patients so that they are sufficiently prepared. Furthermore, one should continuously monitor and provide feedback on the interactions within the project and be prepared to refine procedures as necessary.

As part of a research project evaluating a hospital-based adolescent transition programme, we asked ourselves how the ethical and methodological challenges identified in research with adult co-researchers apply when adolescent patients are co-researchers. Children up to 18 years of age should be given the right to participate in research within the overall framework of human rights. Furthermore, the ladder of participation has been adapted also to children's participation. However, although some recommendations have been issued as part of the national INVOLVE initiative in the UK, there is little evidence of the benefits of partnerships with adolescents in research within health care. Indeed, a systematic review by Haijes et al highlighted that participatory research in paediatrics is limited. Moreover, most results reported by Haijes et al concern research projects where young people were consulted or allowed to give advice, thus referring to levels 2 or 3 of the participation ladder and not to co-research. A recent scoping review by van Schelven et al also points out that more research is needed to expand the evidence base of involving young people with chronic conditions in projects regarding their health and social care.

The aim of this review was to summarize empirical evidence and identify knowledge gaps about the involvement of young patients as co-researchers. By co-research, we refer to full participation in the research team, that is levels 4 and 5 of the simplified participation ladder. Our review aimed to address the following questions: 1. At what stages of a research project are adolescent co-researchers engaged? 2. What methodological, legal and ethical issues and challenges are discussed? 3. What benefits and drawbacks for the co-researchers, for the research and for other stakeholders are discussed? 4. What are the recommended strategies for meaningfully involving young patients as co-researchers?

2 | METHODS

To explore these topics, we performed a scoping review following the outline of the process as described by Peterson et al. A scoping review gives an overview of a broad topic and is a good method for exploring new topics.

2.1 | Data collection

A search of databases (Table 1) was performed in September 2018 with the help of a medical librarian. The following databases were searched: MEDLINE, EMBASE, PsychINFO and AMED. The search was limited to articles written in English, Norwegian, Swedish, Danish, French, Italian, Portuguese and Spanish. This yielded 454 results. The search in AMED yielded no results and is therefore not presented in the table.

In stage two of the process, the first and last author screened the abstracts. The following inclusion criteria were used: articles concerning patients 12 years and above; patients are co-researchers; the research topic is in relation to health care and health systems research; describes 4th and 5th level of participation according to the participation ladder as amended by Teunissen and referenced by de Wit et al; and publication date 2000–current. The first author and one co-author performed the full-text review. In case of disagreement, a third author was consulted. All co-authors were asked to consider whether they knew other publications that should be reviewed. The reference lists of the included articles were hand-searched. This did not lead to any further inclusions. The search was updated in June 2020, and an additional 5 articles were included. Thus, we included 14 articles out of 731 (454 + 277) (Figure 1).

The characteristics of the included articles are presented in Table 2.
Although not mandatory for a scoping review, we chose to perform a quality assessment of the included articles. As argued by van Schelven et al, it helps to put the results in context. The quality of reporting was assessed by the last author. Assessment of the article in Spanish was done by a Spanish-speaking co-author. The quality appraisal was intended as indicative and was not part of the selection process. A protocol designed by Kmet et al was used, which includes ten items. Details are outlined in Table 3. Each article was assessed according to the fulfilment of items and was rated as ‘yes’ (=2), ‘partial’ (=1) or ‘no’ (=0) on each item. Each study was assigned a summary score between 0 and 1 (with higher scores indicating better quality of reporting) by adding the scores of all individual items and dividing by the maximum possible score (20 for qualitative studies).
### TABLE 2 Characteristics of studies

| Author(s): | Dunn, 2017 | Edwards et al, 2016 | Flicker, 2008 | Flicker et al, 2005 | Kruzhik & Jivangee, 2011 | Lincoln et al, 2015 | Mitchell et al, 2017 |
|------------|------------|-------------------|-------------|--------------------|---------------------|-------------------|---------------------|
| Language:  | English    | English           | English     | English            | English             | English           | English             |
| Country of study: | UK         | UK                | Canada      | Canada             | US                  | US                | Canada              |
| Aim:       | To coproduce a transition preparation programme. | To develop a framework for a service evaluation measure for young people to use. | To explore who benefits from community-based participatory research in health research. | To investigate what can be done to better support HIV-positive youth | To explore perceptions of barriers and facilitators for community integration of young people with mental health needs. To influence design of an intervention. | To gain understanding of housing support needs of Transition Age Youth living with mental health conditions. To develop a mechanism for this group to take part in design and conduct of mental health research. | To gain understanding of experiences of young adults in Take Home Naloxone (THN) programmes. To get young adults suggestions for improvement of such a programme. |
| Research design/ methods (as labelled by authors): | Participatory Research | Qualitative mixed methods | Qualitative approach to data | Community-based participatory research | Participatory action research | Community-based participatory research | Principals of community-based participatory action research |
| Description of participants: | Age 17-22 Total of 18 | Age 17-25 Total of 12 | Total of 79 | Unspecified number of HIV-positive youth took part in stakeholder group | Age 17-24 | Age 18-25 | Two peer researchers with lived experiences |

### 2.3 Data analysis

An extraction grid was developed. Categories were created to reflect the research questions and the stages of a research process. The extraction grid was tested by the first and last authors. Amendments were made before application to the included articles. An integrated form of analysis was performed with the help of the extraction grid.

### 3 RESULTS

#### 3.1 Characteristics of the included studies

None of our 14 articles were included in the review by Haijes & van Thiel. Four articles were from the mental health context. Three reports included patients with somatic conditions followed up in hospital. Eight articles were from Northern America, three from the UK, two from the Netherlands and one from Colombia. Twelve projects included adolescents aged 17 years and above. Kramer and Schwartz included co-researchers aged 14-21, and Moules had co-researchers as young as 12-16 years of age. All studies reported on projects with a mainly qualitative approach.

#### 3.2 Quality of reporting

Quality assessment scores ranged between 0.65 and 0.95. All scores but three were over 0.8, which indicates a high quality of reporting for most articles (Table 3).

#### 3.3 Participation may happen at any stage, but seldom at every stage

There is evidence of adolescent co-researchers being involved in all six stages of research (Table 4), but only two projects report on co-researchers being part of all stages. In the preparatory phase, there is focus on research training. Van Staa et al did a field trip with the co-researchers to a newspaper to gain insight and discuss the interview protocol, while Dunn reports on formal literature search training. Some do not go into details on the training given while Lincoln and colleagues report on extensive training and formal testing of knowledge. It is argued that research training addresses the imbalance of power between researcher and co-researcher and that the training provides the adolescents with a wider knowledge base.

Four articles report on involvement in the recruitment phase. This is the area with least reported participation. These articles...
| Study/Author | Country | Language | Description of Research Design/Aim |
|-------------|---------|----------|-----------------------------------|
| Valencia et al, 2010 | Colombia | Spanish | To identify perceptions, experiences and expectations about health services for youth and health-care agents from the Zona de Ladera in the city of Cali. |
| van Staa et al, 2010 | Netherlands | English | To develop a strategy to improve health services under the ‘Friendly Youth Services’ (SAJ abbreviation in Spanish) guide. |
| Cleverley et al, 2020 | Canada | English | To evaluate a participatory research (PR) project involving chronically ill adolescents as ‘co-researchers’. |
| Kramer & Schwartz, 2018 | US | English | To explore qualitatively the experiences of youth transitioning from Child and Adolescent Mental Health Services to Adult Mental Health Services. |
| Moules, 2009 | UK | English | To develop the Pediatric Evaluation of Disability-Patient Reported Outcome (PEDI-PRO) for 14-21-year-olds. |
| Pullmann et al, 2013 | US | English | To explore quality of care in hospital. |
| Wintels et al, 2018 | Netherlands | English | To develop conceptual definitions of engagement in adolescent substance abuse services. |

### Action Research

| Age | Total | Study Participation |
|-----|-------|---------------------|
| 10-19 | 100 | 26, 29 |
| 15-17 | 9 | 0-22, 24-29 |
| 16-18 | 9 | 0-22, 24-29 |
| Team of 8 youths with DD, Age 14-21 | | |
| 9 youth Age 12-16 | | |
| 3 youths Age 17-19 | | |
| 12 ambassadors | | |

Report on co-creation of the recruitment tool or participation in the design and dissemination of the recruitment material. Lincoln et al report that the recruitment strategy was discussed by ‘the whole research team’. Most articles report on involvement in the **design phase** including project development meetings, protocol writing, discussions of research questions, development of questions for interviews, focus groups or preparation of workshops. Nine articles report on participation in the **data collection phase**. Young researchers were involved in data collection such as interviews or focus groups. In the case of focus groups, they either led or co-hosted the group. All articles report on involvement in the **analysis phase**. Some report briefly that the analysis was done by the team, that adolescents were involved in ‘continuous review of all work’, or that the results were discussed. Others are more specific, describing that the co-researchers transcribed their own interviews after which the group as a whole performed the analysis. Mitchell et al describe a process of sharing transcripts and analysing the data as a group. Wintels et al write that adolescents were involved in coding and member check. Flicker et al report on weekly stakeholder group meetings for discussions. When it comes to the **dissemination stage**, some simply state that dissemination was done ‘as determined by the team’ or was shared. Others report co-presentations in different forums, including co-design of conference posters or conference presentations. Co-writing publications ranged from co-writing the published article to popular or ‘unspecified’ publications.

### 3.4 | Methodological, legal and ethical issues and challenges

The reasoning behind choice of methods revolves around the importance of involvement to better meet the needs of adolescents and arguing the need to do research with rather than on patients. It is argued that participatory or qualitative methods are suitable because they give rich opportunities for adolescents to voice their opinion and share their experience. Challenges of participatory methods are raised when it comes to having co-researchers follow all steps of the research process. Van Staa et al struggled with co-researcher engagement in the analysis stage. Others such as Edwards et al had some adolescents that followed all stages, while others took part in a selection of stages. The additional burden that participation puts on young and sick co-researchers is problematized. Lincoln et al also address the challenge of defining who ‘the community’ in a community-based participatory research approach actually is, thus problematizing representativeness of the co-researchers.
Few authors go into detail on ethical considerations. Most state that an ethics committee evaluated their project. Safeguarding confidentiality and informed consent is touched upon. Lincoln et al.22 describe ethical considerations when developing what they call a human research protocol and that the institutional review boards raised questions related to, for example, the ability of the young co-researcher to judge the informant’s capacity to consent, and whether the young researcher could be faced with their own personal problems while interviewing. To address this, the research team developed a self-care plan. Flicker30 addresses the risk of disclosure and stigma that a young co-researcher faces, that is, that by getting involved as co-researchers they also make public their lived experience of the addressed challenge. No article discusses legal challenges or the role of parents in co-deciding or giving permission for participation as co-researchers.

3.5 Benefits and drawbacks of involving young patients as co-researchers

Benefits for the young co-researchers, the professional researcher and the research process are discussed. Arguments used are primarily for the research itself, including relevance of questions, methods and findings, as well as recruitment.22,23,25,27,30,32 Benefits for the young co-researcher include empowerment, skills building and raised self-esteem.24,30 The amount of time and the resources required from the participative research processes are problematized.24,29,30 Van Staa et al.24 question whether participation automatically adds value. Lincoln et al.22 raise the challenge that disclosing one’s membership to a community might lead to stigmatization. Getting ethics boards to understand what participatory research entails was pointed out as another challenge. The discussion of benefits and drawbacks is more reflections on the participation processes, than a report of structured evaluations of the process.

3.6 Recommended strategies for meaningful involvement

No article draws up a list of recommendations, but the importance of clarifying roles has been emphasized.30 especially as this may be an unfamiliar situation for both adult researchers and young co-researchers. Adequate funding is also important30 as these are processes that are generally more time and resource demanding than more traditional forms of research. A second recommendation is to address, in advance, questions of who will benefit from the research.24,30 There are three particular recommendations to safeguard the interests of the young co-researcher. The first is proper training.25 The second is to develop a self-care plan so that possible challenges are discussed and addressed ahead of time.22

### Table 3 Quality of reporting

| Question/objective sufficiently described? | Dunn, 2017 | Edwards et al, 2016 | Flicker, 2008 | Flicker et al, 2005 | Kruzich & Jivangee, 2011 | Lincoln et al, 2015 |
|------------------------------------------|----------|-----------------|-------------|-----------------|-----------------|-----------------|
| Study design evident and appropriate?    | 2        | 2               | 2           | 2               | 2               | 2               |
| Context for the study clear?             | 2        | 2               | 2           | 2               | 2               | 2               |
| Connection to a theoretical framework/wider body of knowledge? | 1        | 1               | 2           | 1               | 2               | 2               |
| Sampling strategy described, relevant and justified? | 2        | 1               | 1           | 2               | 2               | 2               |
| Data collection methods clearly described and systematic? | 2        | 2               | 2           | 2               | 2               | 2               |
| Data analysis clearly described and systematic? | 2        | 2               | 2           | 2               | 2               | 2               |
| Use of verification procedure(s) to establish credibility? | 1        | 1               | 2           | 2               | 2               | 2               |
| Conclusions supported by the results?    | 2        | 2               | 2           | 2               | 2               | 2               |
| Reflexivity of the own account?          | 1        | 1               | 2           | 0               | 1               | 1               |
| Total score per article                  | 0.85     | 0.8             | 0.95        | 0.85            | 0.95            | 0.8             |

| Criterion fulfilled (score 2) | Partially fulfilled or don’t know (score 1) | Not fulfilled (score 0) |
|-------------------------------|---------------------------------------------|-------------------------|
|                               |                                             |                         |
Finally, it is recommended to be flexible, that is, that the young co-researchers should be made aware of the possibility to step out of the research, and/or choose which stages of the research to participate in.

4 | DISCUSSION

This is the first scoping review of adolescent patients participating at level 4 or 5 according to the simplified ladder of participation. We identified only 14 articles matching the inclusion criteria. We find it interesting that we do find evidence of young co-researchers being part of all stages of the research process. This is a similar to findings reported in the review by Van Schelven et al. However, what topics are reported on still raises a number of questions for further discussion.

4.1 | Ethical issues

It is our opinion that the included articles would have benefited from a more in-depth portrayal of ethical challenges of involving young co-researchers. This would have provided important learning points for other researchers. Most mention that they have gone through ethics committee approval, but with one exception, they do not go into detail about what issues they highlighted in their reporting, nor what concerns the committees raised in their responses. Gilchrist et al argue ‘Ethical issues to be considered when carrying out research or service evaluation with children may include: power relationships, consent issues, confidentiality and dissemination of results’. They acknowledge that these issues are not unique to research with children (meaning up to 18 years of age), but argue that there may be a need to pay particular attention to them in this context. Furthermore, Gilchrist and colleagues state that we live in an adult-centred society and that the power adults have can be carried over into research. This is an argument also made in a literature review by Kirk.

In health care, this is influenced by the authority that adolescents are used to impart to their health-care providers. They are used to assigning themselves a subordinate role as patients and of lesser knowledge and age. Whether adolescents and their health-care providers are used to seeing young patients as valid knowledge holders in the context of research will play an important role in their involvement. Are they judged by health professionals as mature enough to take part? Or are the rights to participation lagging behind because health professionals are used to thinking in terms of the right to protection and that safeguarding the best interest of the child is more or less equivalent to protection? The power relation between young patient and health professional is such that in most cases the adolescent would need an invitation from the adult researcher to join research. They would most likely not be the one to take the initiative.
Thus, it is essential that the adult researcher see participation in research as relevant. These ethical considerations should be discussed to advance participation.

A related issue is research ethics boards’ project assessment. How can we ensure that they properly assess not just the research process the co-researchers are to be involved in, but also the role as co-researcher? Researchers wanting to involve young co-researchers should consider and have to account for ethical questions suitable for their projects. This was exemplified by Lincoln et al who developed a self-care plan for the young co-researchers’ well-being.22 Similarly, Kirby13 highlights the importance of recognizing that taking part in research may impact on emotional well-being and that one should discuss this with the young co-researchers at an early stage. The INVOLVE guide13 also addresses issues of safety and well-being when adolescents take part in fieldwork.

4.2 What rungs of the ladder do we aim for?

The discussion on ethics ties into reflections we should have when utilizing the ladder of participation. It is easy to get the impression that participation at the higher rungs of the ladder should be preferred. However, is this really the case? Hart states that his adaptation of the participation ladder, contrary to how it has been used by some, was not meant to be an evaluation tool. He intended his adaptation to be used as a way of portraying different forms of
| Mitchell et al, 2017 | Valencia et al, 2010 | van Staa et al, 2010 | Cleverley et al, 2020 | Kramer & Schwartz, 2018 | Moules, 2009 | Pullmann et al, 2013 | Wintels et al, 2018 |
|---------------------|---------------------|---------------------|---------------------|---------------------|----------------|---------------------|---------------------|
| Two peer researchers recruited. Research training with co-investigators. | Co-researchers invited through nurses. Introduction to research techniques. | | | Interviews done with co-researchers. Discussions on progress. Methods training. | | | |
| | | | | | | | |
| Shared decision making. Development of semi-structured interview guide. | Preparation of interview protocol through discussion of drafted protocol. | Reviewed interview guide. | Develop focus group. | Deciding on research questions. Vignette construction. | Discussion of research questions. Discussion of focus group questions. | Gave input on protocol. Discussion on content of interviews. | |
| Shared Focus groups. Individual interviews. | Performed interviews. | Collected data from own life (written, photo, video and field note). Co-facilitated focus groups. | Took part as storytellers of vignettes. | Took part in focus groups. | | | |
| Read transcripts and analysed data as group. | Involved in discussion of results. | Invited to discussion but unable. Consulted on draft through email. | Participated in team. | Analysed as a group. | Analysed interviews. Analysed vignette data. | Done by the whole group. | Involved in coding. Member check. |
| 'Shared' – Not specified. | Co-develop strategy. Joint design of space. Recommendations to management. | Media activities e.g. interviews. Popular article. Invited to conference | Participated in writing up publication. | | | Conference presentations. |

participation and as a point of departure for reflexion. Are there situations where we should not aim for the top rung of the ladder, for example if aiming for the 5th level of the ladder entails leaving the adolescents on their own in a research process where they should have support? Jones argues that a healthy work environment is the responsibility of the adult researcher. Furthermore, she argues that this should be safeguarded in the process where one considers barriers and boundaries of participation in research. In this process, one should not only look at what barriers to overcome, but also what boundaries to establish for the safety of the adolescent co-researcher. Furthermore, Hart states that ‘It is not appropriate that some children feel that they must always only follow the initiative of others any more than it is good for any child to feel that they should always be a leader’. 

### 4.3 How do we prepare participants for co-research?

A question that arises after reviewing the included articles is what is relevant or adequate research training for young co-researchers? It has been suggested by others that fully involving children as co-researchers can be problematic due to their lack of the theoretical knowledge needed. The way research training was approached, differed greatly in the reported projects. Therefore, the included articles do not provide a clear roadmap for other researchers to follow. The larger the training package, the more time and resource demanding the process will be. This is not an argument against involvement and training of the adolescents, but an argument for thinking this through and accounting for it before endeavouring on
such a process. There is a fine balance between providing research training to the extent that the adolescents feel comfortable to take part and ‘demystifying’ research for them, while still preserving their commitment to the required time and effort.\textsuperscript{38} Interestingly, few of the included articles problematizes what kind of training professional researchers might need to endeavour into co-research with young patients.\textsuperscript{12}

It is important to keep in mind that research training is not the only way to tackle the issue of power balance in the research group. This could also be addressed through creating spaces and opportunities for meaningful participation,\textsuperscript{39} giving each other room, being curious and valuing different kinds of knowledge. One of the most valuable insights that can be gained from adolescent co-researchers is their lived experience as patients, of the care they received and the impact their illness has had in their lives. This can benefit the research in what focus it takes and what questions it investigates. Abma argues ‘People understand the world in different ways but we do not know what those differences are until we have an opportunity to share and discuss’.\textsuperscript{40} If we are to tease out the benefits of participatory research, we need to acknowledge that academic knowledge does not have primacy over other forms of knowledge.\textsuperscript{40} Creating a space where other forms of knowledge emerge may be more important than expecting academic rigour from the young people involved. In fact, Jones argues that children (up to the age of 18 years of age as per the CRC) cannot be held responsible for research,\textsuperscript{37} this responsibility lies with the adult researcher. Research should take into account the skills of the person facilitating the interviews, focus group and overall participation process. Even with education and training, we cannot expect young co-researchers to inhabit the same skills as an experienced adult researcher.

The question remains what all this entails for the training of young co-researchers. We find the example of Jones\textsuperscript{37} interesting. She reports on a project where the training involved learning not only about research methods, but about protecting the rights of the researched, how to ensure own health and safety, techniques to use in interviews, how to logistically get to interviews, planning for things that can go wrong, how to record interviews and also how to do preliminary data analysis.\textsuperscript{37}

### 4.4 Challenges and drawback

Van Schelven et al report in their review that some of their included articles problematize that lack of research experience can lead to lack of depth in for instance interviews.\textsuperscript{15} This is also a concern raised in the article by Van Staa et al included in this review.\textsuperscript{24} Again, we see the need for a discussion on what the top rung of the participation ladder and co-research really entails. Does it mean that young co-researchers take complete lead and are left on their own in all stages of the project? Or, is it more fruitful to think in terms of partnership where the experienced researcher and the young co-researcher, and the research for that matter, would benefit from partnering in the data collection? It all comes back to what the goal of the process is. Is it as Hart puts it liberation of the young co-researcher from adults, or is it to recognize the rights of others to participation and therefore involve them?\textsuperscript{12}

### 4.5 Learning points for future research projects

Finally, what can be learned from the included articles is the need for flexibility. The need for flexibility was also pointed out by van Schelven and colleagues.\textsuperscript{15} A young person may not be able to commit for a long period for different reasons. This brings us back to the discussion of what we aspire to achieve through the participatory process. Is it to have the same adolescents follow the whole process or is it having someone follow each step? And if adolescents are invited to full participation, are well informed, but opt for a lower level of participation such as being consulted, does that mean that the project is not truly participatory? We argue that research would benefit from a mix of forms of participation.\textsuperscript{36} Young people themselves state that there is not one ‘right’ way of involvement, because preferences vary from person to person.\textsuperscript{41} Similarly, the INVOLVE guide cautions against assuming that young people cannot be involved in certain stages of the research, but also against assuming that they wish to be involved in all the stages of the research.\textsuperscript{13} Furthermore, there is a need to acknowledge that desire to participate might shift through time.

### 4.6 Strengths and limitations

One limitation of the review could be specific databases indexing that may have prevented us from identifying relevant publications. Van Schelven et al\textsuperscript{15} problematize that there is a lack of conceptual clarity of PPI resulting in different definitions of the same concepts. This could also have contributed to potential challenges in identifying all relevant studies.

### 5 Conclusion

No lists of recommendations are given in the included articles, but recommendations can be deduced from the articles. There is need for time, funding and flexibility when including young patients as co-researchers. We would argue that research involving adolescents co-researchers should be based on and would benefit from proceeding in line with a rights based framework. We miss a more thorough discussion of legal and ethical dilemmas when including a particularly vulnerable group as co-researchers. More reflection is needed about what meaningful participation is and what it entails in the context of research. Proceeding in line with a child/human rights framework could form the basis for adopted recommendations on how to involve young co-researchers, as well as the development of self-care plans. Research ethics boards need the competencies to properly assess not just the research
process that the adolescents are to be involved in, but their role as co-researchers.

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CONFLICT OF INTEREST
The authors declare no conflict of interest.

AUTHOR CONTRIBUTIONS
Kjersti J. Ø. Fløtten designed and conducted the work presented in this paper, analysed the data and wrote the paper. Isabelle Aujoulat designed the work, analysed the data and has co-written the final version of the manuscript. Ana Isabelle F. Guerreiro has participated in the design, offered advice on data collection and has co-written the final manuscript. Ilaria Simonelli has participated in the design, offered advice on data collection and has co-written the final manuscript. Anne Lee Solevåg participated in the design, offered advice on data collection and has co-written the final version of the manuscript.

DATA AVAILABILITY STATEMENT
Data sharing is not applicable to this study as no new data were created or analysed.

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REFERENCES
1. Arnstein SR. A ladder of citizen participation. J Am Inst Plann. 1969;35(4):216-224.
2. Abma TA, Nierse CJ, Widdershoven GA. Patients as partners in responsive research: methodological notions for collaborations in mixed research teams. Qual Health Res. 2009;19(3):401-415.
3. de Wit MP, Kvien TK, Gossec L. Patient participation as an integral part of patient-reported outcomes development ensures the representation of the patient voice: a case study from the field of rheumatology. RMD Open. 2015;1(1):e000129.
4. National Institute for Health Research. INVOLVE. 2019; https://www.involve.org.uk/. Accessed December 18, 2019.
5. Helse Vest, Helse Midd-Norge, Helse Nord, Helse Sør-Øst. Veiledere for brukermedvirkning i helseforsknings i spesialhelse-setjeningen. Helse Vest, Helse Midd-Norge, Helse Nord, Helse Sør-Øst; 2018.
6. Cleemput I, Dauvain M, Kohn L, Mistiaen P, Christiaens W, Léonard C. Position of KCE on patient involvement in health care policy research. Brussels, Belgium: Belgian Health Care Knowledge Centre; 2019.
7. Malterud K, Elvangen KT. Patients participating as co-researchers in health research: a systematic review of outcomes and experiences. Scand J Public Health. 2020;48(6):617-628.
8. Abma TA. Patients as partners in a health research agenda setting: the feasibility of a participatory methodology. Eval Health Prof. 2006;29(4):424-439.
9. de Wit M, Cooper C, Tugwell P, et al. Practical guidance for engaging patients in health research, treatment guidelines and regulatory processes: results of an expert group meeting organized by the World Health Organization (WHO) and the European Society for Clinical and Economic Aspects of Osteoporosis, Osteoarthritis and Musculoskeletal Diseases (ESCEO). Aging Clin Exp Res. 2019;31(7):905-915.
10. de Wit M, Cooper C, Reginster J-Y. Practical guidance for patient-centred health research. Lancet. 2019;393(10176):1095-1096.
11. Sirevag K, Flotten KJO, Nakstad B, Hvideberg AI, Odden JP, Roy BV. From child to grown up in a medical world: developing an adolescent transition programme at a Norwegian University hospital. Int J Adolesc Med Health. 2017;31(4):1-8.
12. Hart R. Stepping Back From “The Ladder”: Reflections on a Model of Participatory Work with Children. In: Reid A, Jensen BB, Nikel J, Simovska VE, eds. Participation and Learning. Perspectives on Education and the Environment, Health and Sustainability. Dordrecht, Netherlands: Springer; 2008:19-31.
13. Kirby P. A guide to actively involving young people in research for researchers, research commissioners and managers; 2004.
14. Haines HA, van Thiel GJ. Participatory methods in pediatric participatory research: a systematic review. Pediatr Res. 2016;79(5):676-683.
15. van Schelven F, Boeije H, Marien V, Rademakers J. Patient and Public Involvement of young people with a chronic condition in projects in health and social care: a scoping review. Health Expect. 2020;23(4):789-801.
16. Petersen J, Pearce PF, Fergusson LA, Langford CA. Understanding scoping reviews: definition, purpose, and process. J Am Assoc Nurse Pract. 2017;29(1):12-16.
17. Sucharew H, Macaluso M. Progress notes: methods for research evidence synthesis: the scoping review approach. J Hosp Med. 2019;14(7):416-418.
18. Kmet LM, Lee RC, Cook LS. Standard Quality Assessment Criteria for Evaluating Primary Research Papers from a Variety of Fields. Edmonton, AB: Alberta Heritage Foundation for Medical Research; 2004.
19. Bradley EH, Curry LA, Devers KJ. Qualitative data analysis for health services research: developing taxonomy, themes, and theory. Health Serv Res. 2007;42(4):1758-1772.
20. Dunn V. Young people, mental health practitioners and researchers co-produce a Transition Preparation Programme to improve outcomes and experience for young people leaving Child and Adolescent Mental Health Services (CAMHS). BMC Health Serv Res. 2017;17(1):293.
21. Kruzich JM, Jivanjee P. Using evidence from youths to guide improvements in the mental health system. In Roberts-DeGennaro M, Fogel SJ eds. Using evidence to inform practice for community and organizational change. Chicago, IL: Lyceum Books. 2011:157-176.
22. Lincoln AK, Borg R, Delman J. Developing a community-based participatory research model to engage transition age youth using mental health service in research. Fam Community Health. 2015;38(1):87-97.
23. Cleverley K, Lelters M, McCann E. “Objectively terrifying”: a qualitative study of youth’s experiences of transitions out of child and adolescent mental health services at age 18. BMC Psychiatry. 2020;20(1):147.
24. van Staa A, Jedeloo S, Latour JM, Trappenburg MJ. Exciting but exhausting: experiences with participatory research with chronically ill adolescents. Health Expect. 2010;13(1):95-107.
25. Edwards M, Lawson C, Rahman S, Conley K, Phillips H, Uings R. What does quality healthcare look like to adolescents and young adults? Ask the experts! Clin Med (Northfield Il). 2016;16(2):146-151.
26. Moules T. ‘They wouldn’t know how it feels …’: characteristics of quality care from young people’s perspectives: a participatory research project. J Child Health Care. 2019;13(4):322-332.
27. Kramer JM, Schwartz AE. Development of the pediatric disability inventory-patient reported outcome (PEDI-PRO) measurement conceptual framework and item candidates. Scand J Occup Ther. 2018;25(5):335-346.
28. Mitchell K, Durante SE, Pellatt K, Richardson CG, Mathias S, Buxton JA. Naloxone and the Inner City Youth Experience (NiCYE): a community-based participatory research study examining young people’s perceptions of the BC take home naloxone program. Harm Reduction J. 2017;14(1):34.

29. Pullmann MD, Ague S, Johnson T, et al. Defining engagement in adolescent substance abuse treatment. Am J Community Psychol. 2013;52(3-4):347-358.

30. Flicker S. Who benefits from community-based participatory research? A case study of the positive youth project. Health Educ Behav. 2008;35(1):70-86.

31. Wintels SC, Smits D-W, van Wesel F, Verheijden J, Ketelaar M. How do adolescents with cerebral palsy participate? Learning from their personal experiences. Health Expect. 2018;21:1024-1034.

32. Flicker S, Skinner H, Read S, et al. Falling through the cracks of the big cities: who is meeting the needs of HIV-positive youth? Can J Public Health. 2005;96(4):308-312.

33. Valencia CP, Canaval GE, Molina AP, et al. Health care services for young people: a joint construction among youth and health personnel. [Spanish]. Colomb Med. 2010;41(1):26-34.

34. Gilchrist F, Rodd HD, Deery C, Marshman Z. Involving children in research, audit and service evaluation. Br Dent J. 2013;214(11):577-582.

35. Kirk S. Methodological and ethical issues in conducting qualitative research with children and young people: a literature review. Int J Nurs Stud. 2007;44(7):1250-1260.

36. Guerreiro A, Fløtten K. Article 12: The Translation into Practice of Children’s Right to Participation in Health Care. In: Liefaard T, Sloth-Nielsen J, ed. The United Nations Convention on the Rights of the Child. Taking Stock after 25 Years and Looking Ahead. Leiden: Brill Nijhoff; 2016:681-707.

37. Jones A. Involving Children and Young People as Researchers. In Fraser S, Lewis V, Ding S, Kellett M, Robinson C, eds. Doing Research with Children and Young People. London, UK: SAGE Publications; 2004.

38. LoIacono Merves M, Rodgers CR, Silver EJ, Sclafane JH, Bauman LJ. Engaging and sustaining adolescents in community-based participatory research: structuring a youth-friendly community-based participatory research environment. Fam Community Health. 2015;38(1):22-32.

39. Lundy L. ‘Voice’ is not enough: conceptualising Article 12 of the United Nations Convention on the Rights of the Child. Br Educ Res J. 2007;33(6):927-942.

40. Abma T, Banks S, Cook T, et al. Participatory Research for Health and Social Well Being. Cham, Switzerland: Springer Nature Switzerland AG; 2019.

41. Lightfoot J, Sloper P. Having a say in health: involving young people with a chronic illness or physical disability in local health services development. Child Soc. 2003;17(4):277-290.

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