Informed consent and assent guide for paediatric clinical trials in Europe

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ABSTRACT
Objective Clinical trial sponsors spend considerable resources preparing informed consent (IC) and assent documentation for multinational paediatric clinical trial applications in Europe due to the limited and dispersed patient populations, the variation of national legal and ethical requirements, and the lack of detailed guidance. The aim of this study was to design new easy-to-use guide publicly available on European Medicines Agency’s, Enpr-EMA website for all stakeholders.

Methods Current EU legal, ethical and regulatory guidance for paediatric clinical trials were collated, analysed and divided into 30 subject elements in two tables. The European Network of Young Person’s Advisory Group reviewed the data and provided specific comments. A three-level recommendation using ‘traffic light’ symbols was designed for four age groups of children, according to relevance and the requirements.

Results A single guide document includes two tables: (1) general information and (2) trial-specific information. In the age group of 6–9 years old, 92% of the trial-specific subject elements can be or should be included in the IC discussion. Even in the youngest possible age group (2–5 years old children), the number of elements considered was, on average, 52%.

Conclusion The EU Clinical Trial Regulation (2014) does not contain specific requirements exclusively for paediatric clinical trials. This work is the first to extensively collate all the current legal, regulatory and ethical documentation on the IC process, together with input from adolescents. This guide may increase the ethical standards in paediatric clinical trials.

INTRODUCTION
Children represent 20% of the European population (~90 million citizens).1 In 2020, on average, 11% of all clinical trials registered into the European Clinical Trials database (EudraCT) were paediatric clinical trials with investigational medicinal product.2 Due to the EU Paediatric Regulation,3 all new clinical trials with medicinal products must consider if the inclusion of children (0–18 years of age) from all or some age ranges is relevant, which means the number of paediatric clinical trials will increase. The number of participants per country can be very small.

These multicentre and multinational trials create demanding research environment for academic groups and pharmaceutical industry in Europe, as they spend considerable time and resources preparing documentation for clinical trial application. The submission packages for the competent authorities and ethics committees including informed consent (IC) and assent documents are challenging due to non-harmonised legislation and wide variation between national regulatory requirements.4,5

In Europe, the legal age for giving independent IC for participation in a clinical trial varies between 14 and 18 years. In paediatric trials, the consent process includes the child’s own consent, assent or agreement. According to many national laws, a child’s own assent is usually not sufficient alone to allow his/her participation, unless supplemented by the consent of the child’s legally designated representative(s).4,6

Legal, ethical and regulatory framework for paediatric clinical trials in Europe
After the EU Paediatric Regulation (EC No 1901/2006),7 in 2014 the EU Clinical Trials...
Regulation followed\(^6\) the current EU Clinical Trial Directive.\(^3\) The clinical trials regulation will facilitate clinical trials in the EU after it will come into application on 31 January 2022. Under this regulation, ethical review will remain under each EU Member State according to their national laws. In May 2018, the new EU General Data Protection Regulation\(^9\) came into force, impacting clinical trials and consent process, requiring an explanation of the legal basis for the collection and processing personal data on trials. Some countries require even explicit consent for data processing.

The ethical foundation for clinical research is the updated World Medical Association’s Declaration of Helsinki (WMA DoH),\(^10\) incorporated into the International Conference on Harmonisation (ICH) Good Clinical Practice (GCP) guideline.\(^11\) Both principles of the WMA DoH and GCP are evident in the EU Clinical Trial Regulation. In 2017, the EU Commission’s revised version of the EU ethics guideline for paediatric clinical trials was published.\(^5\)

### Children’s right to be involved in clinical trial design and IC process

According to the current legal texts and ethical recommendations, children should be able to take part in the consent process. A child’s participation is derived from the general requirements for the right to express their own opinions from the Fundamental rights of the Convention on the Rights of the Child.\(^12\) 13

The increased patient and parent involvement have evolved after the European Young Person’s Advisory Groups (eYPAGnet) was established in 2017.\(^14\) The eYPAGnet is a consortium of Young People’s Advisory Groups (YPAGs) across Europe, supporting the development of new national YPAGs. The YPAG provide a platform for children to have a voice, share their opinions and apply their experience to a variety of issues in clinical trials, such as ICs and assents. These groups include young people aged between 8 and 19 years (even up to 21 years) who are patients and/or healthy children having an interest in science, healthcare and children’s rights.\(^15\)

Comprehensive trial information must always be presented to the child’s legal representative(s) both orally and in writing.\(^5\) Trial information should be also provided to the child participant, but it should be adapted to the child’s language skills and understanding, the child’s developmental stage, intellectual capacity, medical condition, previous life/disease experience and other circumstances. Chronological age only partly correlates with maturity. The maturity is evaluated by the investigator. Where appropriate, translation should be arranged. It is very important to identify a child’s potential dissent or disagreement to participate, which should always be respected.\(^9\)

### Enpr-EMA facilitates the main objective of the paediatric regulation

In 2013, the European Network of Paediatric Research at the European Medicines Agency (Enpr-EMA) initiated various multistakeholder working groups to find solutions to emerging medical needs related to paediatric clinical trials.\(^16\) 18 The ethics working group has focused on ethical issues in paediatric clinical trials and published first ‘Tool Kit’ of European informed consent and assent requirements in December 2015\(^19\) and related article later in May 2016.\(^4\)

### The aim of the new guide for paediatric ICs and assents

Some EU-funded projects and learnt societies have developed new practical tools, guidance, design methods and training programmes for supporting paediatric clinical trials.\(^20\) 25 The aim of this new guide was to provide more practical easy-to-use tool for designing the content of IC and assent documents to enhance the high-quality ethical standards of paediatric clinical trials. This guide can be adapted to all types of paediatric clinical trials on a case-by-case basis, and it can also be used to promote the involvement of children in the consent process. This guide was initially published on Enpr-EMA website on January 2021.\(^20\)

### METHODS

The working group collated all current EU legislative, ethical and regulatory legal texts specific to paediatric clinical trials via literature search and used these as the primary data source. A total of 30 identified main subject elements required for the consent process in legal, ethical or regulatory texts were divided into two tables based on the nature of the requirement. The first table is applicable to all trials (5 general elements) and the second table includes trial-specific topics (25 elements), which can vary between trial design (table 1). All subject elements were considered for four paediatric age groups (0–2; 2–5; 6–9; 10–18) and legal representative(s) as defined in the EU ethics guideline.\(^6\)

A three-level recommendation was established as ‘traffic light’ symbols. These were used to separate subject elements to be considered, not considered or categorised as optional according to the age group per each subject element (table 2).

The three-level recommendation was reviewed against each subject element (5 + 25 elements) and across all age groups. All elements were sorted with the symbols to separate each subject element according to suitability for each age group and this resulted in recommendations per age group.

All subject elements were listed separately as these are unrelated to each other. The data were then arranged into five vertical main sections (columns) (I–V): (I) age group in years, (II) legal representative(s), (III) elements to consider/information which must be included in the consent/assent document, (IV) questions to be addressed, and (V) notes and example methods/texts to be used. The first heading (I) includes four age group columns. One additional sixth section (column VI) was added to table 4 for the element numbering (tables 3 and 4). Both tables include text partly written in bold, emphasising the most important aspects, or requirement, need to be considered.

After structural design, the eYPAGnet members of three national YPAGs (the UK, Scotland and Spain) were requested to review all 30 subject elements in order to identify important information, any issues or missing information and the preferred format of given information for a practical consent discussion. All comments were collected by monographic sessions led by the group facilitators. All feedback was collated into the report. The report data were incorporated into the guide’s section IV, as questions, example methods or texts to be used when designing consent/assent documents, or when recruiting paediatric participants to trials.

The last review for the complete guide was done by the Office of the Paediatric Medicines (Scientific Evidence Generation Department) at the European Medicines Agency. The guide document was finalised according to these comments (figure 1).

### RESULTS

#### Subject elements adaptable and suitable for various age groups is very high

After all the subject elements were collated, it clearly illustrated the complexity of IC forms and the consent process in the paediatric population. Many items must be discussed in an adapted
manner according to their age and maturity representing all possible therapeutic areas and diseases, from healthy subjects (eg, vaccine trials) to extremely rare disease patient groups.

Listing trial-specific subject elements (25 elements of table 4) relevant to all paediatric clinical trials resulted in all elements (100%) being relevant and to be considered when designing and conducting clinical trials in the oldest age group. Further, when looking at children in the age group 6–9 years, 23 out of 25 (92%) elements can be optionally included or should be included in adapted ways in the consent process. Even in the youngest (92%) elements can be optionally included or should be included in the consent process for this age group. The specific questions addressed to adolescents in the consent discussion

The comments received from the eYPAGnet members resulted in a high number of recommended questions (81) to adolescents per each subject element. The total number of questions in both tables (3 and 4) is 86, and the number of questions varies between one and six per subject element. The first five questions to the table 3 were generated by the working group.

**DISCUSSION**

The practical challenges in the implementation of paediatric multicentre clinical trial

Paediatric clinical trial design and recruitment need specific expertise and experienced, qualified and trained personnel. Similar experience is required from the people reviewing and assessing the trial protocols, patient information, consent and the assent or agreement forms.

### Table 1 Subject elements of IC and assent guide tables 3 and 4

| No of subject element | Subject element to consider, information which must be included in the IC/assent document |
|-----------------------|-----------------------------------------------------------------------------------------|
| Guide table 3 General |                                                                                         |
| 1.1                   | Language, translations, visual materials, methods for giving information, time used for IC process, unbiased approval, competent personnel for the process and the documentation |
| 1.2                   | Concept of the clinical trial methodology                                                |
| 1.3                   | Dissent, refusal, disagreement, voluntariness, free decision and respect for autonomy    |
| 1.4                   | Legal representative(s) roles, sensitive issues for adolescents                          |
| 1.5                   | IC/assent/agreement signatures and re-consenting (long-term studies)                     |
| Guide table 4 Trial specific |                                                                                           |
| 2.1                   | Clinical trial introduction; title, topic, purpose, size (no of participants)             |
| 2.2                   | Protocol introduction; duration, visits, procedures (trial plan)                          |
| 2.3                   | Participant selection and recruitment process                                            |
| 2.4                   | Information about the institution/organisation (ie, hospital) and the personnel involved in the trial |
| 2.5                   | Introduction of the investigational medicinal product(s) and placebo, or device(s) used in trial |
| 2.6                   | Introduction of other treatment options and alternatives for trial                       |
| 2.7                   | Possible benefits or expected benefits of the clinical trial for the participant         |
| 2.8                   | Information about procedures, tests, samples, measurements and possible pain or discomfort included to the trial conduction, how these are prevented and minimised |
| 2.9                   | Detailed information about trial time; visits, timing, place, schedules                  |
| 2.10                  | Introduction of possible risks, disadvantages, side effects or other inconveniences of the trial medication or the trial procedures |
| 2.11                  | Information about genetic testing if included to the trial                              |
| 2.12                  | Use of ionising radiation if used in trial procedures                                    |
| 2.13                  | Biological samples; handling, storage, retention, banking, data use if included to the trial |
| 2.14                  | Possible future effect to fetus/sperm                                                    |
| 2.15                  | Special conditions; emergency, emancipated minor, pregnancy, breast feeding, unexpected problems, change of the legal representative(s) or other changes during the trial conduction |
| 2.16                  | Confidentiality and Data Protection (GDPR)                                              |
| 2.17                  | Information about discontinuation and right to withdraw, discontinuation for medical or safety reasons, adverse effect reporting |
| 2.18                  | Information about trial costs, medicine costs and expenses and allowed compensation NOTE: incentives or inducements are forbidden by EU law |
| 2.19                  | Information about patient insurance and other applicable damage compensations            |
| 2.20                  | After trial measurements and follow-up period if included                                |
| 2.21                  | Results of the trial                                                                    |
| 2.22                  | Information about the Competent Authorities (ethics, medicine agency) and appropriate expertise needed for trial authorisation |
| 2.23                  | Information about clinical trial sponsor and trial funding                                |
| 2.24                  | Contact information for trial personnel                                                  |
| 2.25                  | Confirmation of understanding—verification with some method                              |

IC, informed consent.
Table 3  General information for informed consent and assent (agreements)

| Age group in years | Legal representative(s) | Elements to consider/information which must be included into the assent/consent document | Questions to be addressed | NOTES and example methods/techniques to be used |
|--------------------|-------------------------|-----------------------------------------------------------------------------------------|--------------------------|-----------------------------------------------|
| 0-2                |                         | - Language/translations/methods used for providing information/time used for information (paper or electronic) Language must be concise, clear, relevant and understandable, adapted to the age and maturity of the recipient of the information: The assent/consent agreement (agreement) should be documented, and in case it is not possible to seek assent from the child the reasons for this should be explained. The target is to obtain credible informed assent/consent with unbiased approval, understanding and willingness or refusal by both parties (child and legal representative). The personnel providing the information should be competent in communicating and working with children and families, providing time and space for the discussion and decision. Ensure that children are properly informed. | - Do the legal representative(s) and the child/adolescent understand the information given? - Is there enough time to provide relevant information, for discussion and to answer questions? - Are the facilities suitable/safe for discussion? - Have the child/adolescent and legal representative(s) had time to read the information beforehand (eg, at home) prior to the discussion? - Have you used correct grammar? Do you have the correct form of address (per age) for the addressee (child/adolescent) to avoid infantilisation? | Use visual and informative materials/sources (IT based or manual) to increase understanding (if appropriate) such as: - Videos, DVDs - Pictogram - Pictures - Drawings - Cartoons - Photographs - Diagrams/charts/tables - Social media contents, www-links - Computer programmes - physical instrument/tool/device mimics of methodology - Glossary/dictionary of terms eYPAGnet® notes: Max. 3 pages/3 screen views (total) for the assent/consent document. Large enough font. Clear layout. No bullet points. No pictures from 'Clip Art'. Different colours to highlight important info. Chart for visits is helpful. Add list of additional resources if some of the information cannot fit on three pages. Support to create text to documents: 1. Readability: Flesch–Kincaid Readability Score testing tool. Available at: https://www.webfx.com/tools/read-able/flesch-kincaid.html 2. Health literacy: Quick Guide to Health Literacy. Available at: https://health.gov/communication/literacy/quickguide/default.htm |  |
| 2-5                |                         | - Explanation of the concept of a clinical trial and the methodology used. | - What is a clinical trial? - How does the clinical trial differ from normal care? - What is randomisation/double-blind/open label? etc? | NOTE: Explain only the relevant methodology—a short version—used according to the current protocol. Avoid complex terms and flowcharts with too much detail. |  |
| 6-9                |                         | - Dissent/refusal/disagreement/respect for autonomy Voluntariness/right to refuse/right to dissent/free decisions | - What is the explicit wish of the child/adolescent (capable of forming an opinion and assessing the information)? - Has the child/adolescent understood that they may refuse participation or withdraw at any time during the trial? - Is the child/adolescent’s free wish/decision respected (according to age/maturity) by the investigator/trial personnel? | NOTE: The agreement of a child should be requested systematically, even if the assent is not legally required. Children should be provided with age-appropriate information (with supplementary visual information where appropriate) and have the opportunity to form an opinion or decision. Their refusal or dissent should be respected, objections should be analysed (reason), and possible help sought for anticipated burden (fear, distress, etc). Resistance of very young children should be identified and discussed with legal representatives. |  |
| 10-18              |                         | - Legal representative/role/sensitive issues of adolescents (not discussed with parents/legal rep(s)) | - Is there an opportunity for older adolescents to have a private conversation (without parents) with the trial personnel about confidential/sensitive issues? | NOTE: Legal reps, roles as empowered for decision making should be recognised, but there should be an additional option for the adolescent to express any concerns or worries so as to respect their autonomy (eg, an independent person or mailbox or other method). NOT the trial personnel: as this may create conflict. |  |

Continued
Pharmaceutical companies and Contract Research Organisations have their own document templates, resulting in hundreds of different consent and assent documents submitted to European ethics committees, which repeatedly receive incomplete or inappropriate documentation for paediatric clinical trials. The most problematic area is patient information and consent forms, as these are often too complex for children to understand, and too extensive to read and comprehend. The language is often targeted for adults. The child’s participation cannot rely on the adult IC, which applies to decisions made by those with the legal and intellectual capacity to make such choices. Children usually lack such capacity, and they need adapted information.

Repeated amendments and committee’s re-evaluations prolong ethics approvals and cause delays to the start of clinical trials, causing unnecessary loss of time and money. To avoid such pitfalls, consent and assents need to be designed according to high ethical standards.6 27–31

The age groups of children can be used for different purposes depending on scope

The International Conference of Harmonisation (ICH) E11 Guideline32 lists five age groups (preterm newborn infants, term newborn infants; 0–27+ days, infants and toddlers; 28 days to 23 months, children; 2–11 years and adolescents; 12 to 16–18 years dependent on the region). This ICH guideline is scoped for 23 months, children; 2–11 years and adolescents; 12 to 16–18 newborn infants; 0–27+ days, infants and toddlers; 28 days to appropriate involvement of children.

Instructions for planning the consent process and the appropriate involvement of children.

ICH guideline, as the purpose and scope is to provide general

For the first time, the EU Clinical Trial Regulation makes the involvement of children in the consent process mandatory, for younger children, parents can explain by using “story telling” method.

The young people provide important expertise to paediatric trials

For this guide, the young person’s groups provided important input. This represents the direct voice of young people and is extremely valuable in the consent discussion. The importance of the opinions and experiences of children and their families is now becoming a crucial part in the development of new medicinal products, as recognised by many stakeholders.34–36

Current existing guidance is not enough to design consents and assents for children

To date, there are only a limited number of public ethical and practical guidance documents regarding consent and assent design specifically for paediatric clinical trials. Some learnt societies have provided articles, books, videos, and recommendations, practical guidance and training programmes for all stakeholders.20–25 However, none of these existing materials provide similarly detailed instructions for the design of the paediatric consent, assent or agreement document structure, and guide for the whole consent process, including the voice of children.

Practical limitations and implementation of this guide

Each Member States of the EU and European Economic Area has its own national legislation and detailed requirements for consent documents, and these vary among countries. Therefore, this guide should be used together with the local or national requirements. It is also important to acknowledge that while this guide contains instructional text (ie, ‘should be’), these are not official requirements or legal guidance, but suggestions serving as an example for every age group. All the subject elements need to be adapted case-by-case according to the nature of the clinical trial protocol.

This guide is designed to support the consent process for paediatric clinical trials in Europe across all age groups. It can be used when preparing submission packages to competent

However, it is important to distinguish between the requirement for consent for clinical trial participation and the requirements for lawful processing of personal data under the GDPR. The standards of quality (reliability of data) and patient safety for medicinal products fall under different legal bases. Therefore, the requirement for IC by the EU Clinical Trial Regulation is a safeguard, not a legal basis for data processing, and must not be confused with consent as a legal ground for processing personal data set out in Article 6(1)(a) of the GDPR.31

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### Table 4: Trial specific information for informed consent and assent (agreements)

| Element number | Age group in years | Legal representative(s) | Elements to consider/information which must be included into the assent/consent document | Questions to be addressed | Notes and example methods/taxs to be used |
|----------------|-------------------|-------------------------|------------------------------------------------------------------------------------------|--------------------------|-------------------------------------------|
| 1              | 0–2               | ✓                       | Trial/topic of the trial/introduction/purpose of the trial/size of the trial (how many patients/participants, how many sites) | ► What is the purpose of this trial? ► Why is this trial needed? ► Is there information about this trial available somewhere? | NOTE: EUCTR number and trial protocol code must be added. The trial must be registered in the official Trial Registry (EudraCT) before the start of the trial. Additionally, in other registries or websites according to national requirements. |
| 2              | 2–5               | ✓                       | Introduction of the protocol (trial plan)                                                 | ► What data is needed and why? ► What will be done? | NOTE: Use a glossary of definitions or dictionary of terms if the protocol is very complex to explain. |
| 3              | 6–9               | ✓                       | Participation/recruitment/child/adolescent selection                                      | ► Why has the child/adolescent been invited to participate in this trial? | |
| 4              | 10–18             | ✓                       | Information about the institution/ organisation (eg, hospital) and doctor(s)/investigator(s)/nurse(s)/trial personnel conducting the trial | ► Who are they? (eg, family GP) ► Do they have required professional expertise? ► Will the doctor be the same throughout the trial? | NOTE: Include relevant information about all relevant partners and personnel involved in the conduct of the trial, including their expertise. |
| 5              |                   | ✓ ✓                     | Introduction of the investigational products/placebo/device used in the trial            | ► What medicines/devices is it? ► What (pharmaceutical) form is it? ► How does it work and are there side effects? What are the effects on contraception/use of alcohol/ smoking? ► How will it be administered/taken? ► Has it been tested in children before? ► What does placebo mean? | |
| 6              |                   | ✓ ✓                     | Other treatments/alternative/options for clinical trials or diagnosis                     | ► What are the current/existing alternative methods/treatments if the child/adolescent does not want to take part in this trial? | NOTE: The risks of premature termination need to be explained to legal reps. (and children if they have capacity to understand). Reassurance should always be given that withdrawal will not affect normal treatment (alternative, standard) which should be available. |
| 7              |                   | ✓ ✓                     | Possible benefits/expected benefits                                                       | ► Why might it be beneficial for the child/adolescent to be in this trial? | NOTE: Explain the possible benefits (direct or same benefit for the population). |
| 8              |                   | ✓ ✓                     | Procedures/tests/measurements according to the trial protocol                            | ► What will happen? ► What tests will be done and when? ► Why are these needed? ► How does the procedure is part of normal daily life, food/drink/sports and hobbies? ► How does it differ from current standard care? ► Does it mean taking any time off school? | NON-Europe procedures should be preferred, where possible. Legal reps. and children should be informed whether the procedure is part of usual standard of care or if it is part of the trial (extra), and whether there is direct benefit or not. An explanation about the procedure in honest (not frightening) language must be offered to both the legal representative and the child prior to the procedure. |
| 9              |                   | ✓ ✓                     | Time/timing/trial duration (whole trial, at specific site) Visits/trial schedule          | ► How long will the trial take? ► When are the trial visits? ► How will it affect school/vacations/ holiday/travel? | NOTE: Use flowcharts. The expected duration of participation must be stated. |
| 10             |                   | ✓ ✓                     | Risk/diadvantages/side effects of the trial procedures/investigational medicinal products/trial conduct/diagnosis | ► What might inconvenience the child/adolescent? ► Will there be extra pain/burden or side effects—related to either procedures/tests or to the trial medication? | NOTE: Risk/burden assessments must be done prior to the trial (via scientific medical and ethical assessment conducted by competent authorities). Possible realistic/anticipated risk/burden during the trial must be described at a reasonable level during the assessment process. |
| 11             |                   | ✓ ✓                     | Genetic testing                                                                          | ► What kind of test? Blood/urine/ other type? Why is it taken? ► How will it be taken? ► How will the information be collected from the genetic test? ► How will the information be used? ► Who will be informed of the test results? ► What is the duration of sample storage? | A separate consent/assent must be required for genetic tests. Disclosure of genetic information to the child/adolescent requires parent information or expert counselling (must be known what information is collected, how the information will be used, and how it is interpreted.) Explain the reasons/rationale and the procedure for giving the results in practice, if planned to do so in a trial. NOTE: Only if relevant (trial includes genetic testing). |
| 12             |                   | ✓ ✓                     | Use of ionising radiation                                                                 | Is it used? How? Are there any side effects? | NOTE: Only if relevant (trial includes ionising radiation) |
| 13             |                   | ✓ ✓                     | Biological samples/handling/ storing/banking                                             | ► What kind of samples? Tissue/blood/ saliva/epithelial fluid/other? ► Why are they taken? ► Where are the samples stored and for how long? ► Is it possible to cancel/withdraw consent for future use? | A separate consent/assent must be required for biological samples. Must comply with General Data Protection Regulation (GDPR2016) and national legislation. The collection and use of samples must be described in consent/assent forms. NOTE: Only if relevant (trial includes biological samples). |

Continued
| Element number | Age group in years | Legal representative(s) | Elements to consider/information which must be included into the assent/consent document | Questions to be addressed | Notes and example methods/texts to be used |
|----------------|-------------------|-------------------------|--------------------------------------------------------------------------------|--------------------------|------------------------------------------|
| 14             | 0-2, 2-5, 6-9, 10-18 | ✔️ ✔️ ✔️                  | Possible future effects (infertility, birth defects, miscarriage) | ✚ Can the trial medication or treatments have some effect on the fetus via mother or father, in case of the child/adolescent's pregnancy? <br>✠ What happens if this situation occurs? <br>✠ Who will be told about this? | <br>NOTE: Information about the potential teratogenic risks during pregnancy (fertility (both females and males)) should be discussed, and also the possibility to use contraception, and what type of contraception should be used if it is required. Explain what should happen if pregnancy arises during the trial. |
| 15             |                    | ✔️                       | Special situations: Emergency/emancipated minor/pregnancy/breast feeding/unexpected problems | ✚ How will these situations be handled if they occur? <br>✠ Who will be contacted? <br>✠ Is there a need for prior consent? | <br>NOTE: The protocol should define emergency situations and conditions (eg, time lag until consent is signed) for deferred consent (can be delayed and sought as soon as possible after inclusion). Consent may be deferred in certain emergency situations. Emergency: Shortened time for decision—full explanation should follow later. |
| 16             |                    | ✔️ ✔️ ✔️                  | Confidentiality/personal data protection | ✚ What data from patient files and what additional information is needed and why? <br>✠ How will the data be used? <br>✠ Who has access to this data? Must be clearly stated (using names, if required) who has access to data, how it is used and how the anonymisation is done. | <br>NOTE: Regulatory Authorities have legal permission to have access to research documents and data during inspections and audits. The data will be stored anonymously, and the researcher will have access to it. Explain what anonymisation means. |
| 17             |                    | ✔️ ✔️ ✔️                  | Discontinuation by child/adolescent/wright to withdraw/discontinuation for medical reasons (safety) by the sponsor | ✚ How it may happen/when/for what reasons? <br>✠ What happens later? <br>✠ What are the options for further care/medication? | <br>NOTE: Identification of possible risks, minimising and monitoring the risks—assent/consent should include an explanation about the probability and magnitude of harm anticipated in the trial and how this will be minimised, followed and handled. Stopping rules must be included in the protocol. NOTE: Check with the relevant national authorities regarding the right (or not) to have information already collected deleted and samples already collected destroyed. |
| 18             |                    | ✔️ ✔️ ✔️                  | Compensation/expenses/NO incentives/ inducements used | ✚ What expenses are expected during the trial? (exact level) <br>✠ What costs will be covered by the trial sponsor/hospital? <br>✠ How will expenses be compensated? | <br>NOTE: No financial contribution/ inducements should be offered, except compensation for the parents’ legal representatives’ expenses and loss of earnings directly related to the child’s participation. A small token of appreciation may be acceptable—but needs to be approved by the Ethics Committee (EC) through review/assessment. Children do not need detailed reimbursement information if not directly related to personal compensations (eg, travels, meals, etc). Legal reps. need all information of the compensations and reimbursements. |
| 19             |                    | ✔️ ✔️ ✔️                  | Insurance | ✚ What possible damages could be expected during the trial (ie, what may go wrong and how are these covered by insurance)? Rerally. | <br>NOTE: Damage compensation is mandatory as per CTR and should be ensured by Member States. Insurance should not waive liabilities regarding long-term effects (delayed effects are typical in children when they grow up). |
| 20             |                    | ✔️ ✔️ ✔️                  | After trial/follow-up measurements | ✚ Will the trial medication be available after the trial? <br>✠ What happens after the trial? | |
| 21             |                    | ✔️ ✔️ ✔️                  | Results of the trial | ✚ When is it expected to have trial results? <br>✠ Will child/adolescent/legal rep. be informed of the results? | <br>NOTE: Information about availability of trial results must be stated. The information should include a summary of results presented in terms understandable to a layperson and must be available through EU database (patient). |
| 22             |                    | ✔️ ✔️ ✔️                  | Information about the Competent Authority (CA) authorizations and Ethics Committee (EC) review/approvals including the appropriate expertise used in the assessment. The reviewers should be independent of the sponsor, the investigator and the trial. | ✚ Who has reviewed/approved the trial? <br>✠ Has the review been undertaken by people who have official authority, and expertise for the assessment? Is this legal and what? | <br>NOTE: Should be explained who has reviewed/approved the protocol, but they have not decided on behalf of potential child/adolescents. NO need to include information about EC approval in the young child’s assent, as they do not understand the concept of ‘ethics’. |
| 23             |                    | ✔️ ✔️ ✔️                  | Information about the sponsor/funding of the trial | ✚ Who will fund this trial? <br>✠ Do they pay the trial personnel directly? | <br>NOTE: Should be stated that the hospital receives money not the investigator directly. Need to be clear interest differentiation between the investigator and sponsor. |
| 24             |                    | ✔️ ✔️ ✔️                  | Contact information | ✚ Who can be contacted at any time/ for any reason during the trial? <br>✠ How should they be contacted? | <br>NOTE: Always give contact information to all participants. NO need to have contact information in assent/consent for very young children. |
| 25             |                    | ✔️ ✔️ ✔️                  | Confirmation of understanding | ✚ Were answers provided to all the questions asked by the child/adolescent and/or legal rep.? <br>✠ Are there still issues that are unclear and need to be resolved? | <br>NOTE: You may use an additional summary leaflet, but it depends on the complexity of the trial. |
The guide data was collected by literature search. The legal, ethical and regulatory requirements for paediatric clinical trials and consent process were analysed, structured and divided to table format of 30 separate elements (rows). All elements were further divided into 5 sections; I-V (columns). All relevant references were included.

All 30 elements were sub-divided and analysed per 4 age groups according the EU Ethics Guideline (R1). The 30 elements were further divided to two tables: 1) General information for Informed Consent and Assent (Agreements), and 2) Trial Specific Information for Informed Consent and Assent (Agreements). For table 2, one additional sixth section (column VI) was added for the element numbering. A 3-level "traffic light" recommendation per each age group was designed.

Three national groups of European Young Person’s Advisory Groups Network (eYPAGnet) reviewed the data of tables 1) and 2) and provided feedback for all 30 elements by monographic sessions led by the group facilitators. The report of collated data of three groups was integrated to the table section IV: "Questions to be addressed".

The relevant introduction with methods and list of abbreviations and definitions was added to the guide tables. The complete guide was reviewed by the Office of the Paediatric Medicines (EMA).

The guide document with all references was finalized according EMA’s comments and published on Enpr-EMA website.

Figure 1 The flowchart of the development process of the informed consent and assent guide.

CONCLUSION

This work is the first to extensively collate and document all the current legal, regulatory and ethical guidelines on the consent and assent process for paediatric clinical trials in Europe, together with input from adolescents. It is a single easy-to-use publicly available guide for all stakeholders and the contents can be adapted to trials in case-by-case.

Although the new EU Clinical Trial Regulation will harmonise Clinical Trial Application practices in Europe, it does not contain specific guidance for the paediatric clinical trial consent process. As there are still many national legal differences in the requirements for consent and assent documents, this guide may increase the level of ethical standards and facilitate the harmonisation of paediatric consent and assent documents in Europe.

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