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SwissPedData: Standardising hospital records for the benefit of paediatric research

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Summary

BACKGROUND: Improvement of paediatric healthcare is hampered by inefficient processes for generating new evidence. Clinical research often requires extra encounters with patients, is costly, takes place in an artificial situation with a biased selection of patients, and entails long delays until new evidence is implemented into health care. Electronic health records (EHR) contain detailed information on real patients and cover the entirety of patients. However, the use of EHR for research is limited because they are not standardised between hospitals. This leads to disproportionate amounts of work for extracting data of interest and frequently data are incomplete and of poor quality.

AIMS: SwissPedData aims to lay the foundation for a paediatric learning health system in Switzerland by facilitating EHR-based research. In this project, we aimed to assess the way routine clinical data are currently recorded in large paediatric clinics in Switzerland and to develop a national EHR-based set of common data elements (CDEs) that covers all processes of routine paediatric care in hospitals.

METHODS: A taskforce of paediatricians from large Swiss children's hospitals reviewed the current status of routine data documentation in paediatric clinical care and the extent of digitalisation. We then used a modified Delphi method to reach a broad consensus on a national EHRbased set of CDEs.

RESULTS: All Swiss children's hospitals use EHR to document some or all aspects of care. One hundred and nineteen paediatricians, representing eight hospitals and all paediatric subspecialties, participated in an extended Delphi process to create SwissPedData. The group agreed on a national set of CDEs that comprises a main module with general paediatric data and sub-modules relevant to paediatric subspecialties. The data dictionary includes 336 CDEs: 76 in the main module on general paediatrics and between 11 and 59 CDEs per subspecialty module. Among these, 266 were classified as mandatory, 52 as recommended and 18 as optional.

CONCLUSION: SwissPedData is a set of CDEs for information to be collected in EHR of Swiss children's hospitals. It covers all care processes including clinical and paraclinical assessment, diagnosis, treatment, disposition and care site. All participating hospitals agreed to implement SwissPedData in their clinical routine and clinic information systems. This will pave the way for a national paediatric learning health system in Switzerland that enables fast and efficient answers to urgent clinical questions by facilitating high-quality nationwide retrospective and prospective observational studies and recruitment of patients for nested prospective studies and clinical trials.

Introduction

The creation of new evidence in medicine and the improvement of patient care are hampered by inefficient and laborious processes [1, 2]. Most evidence is gathered through stand-alone research projects that are costly, timeconsuming, and conducted in an artificial research setting with a selected sample of patients. It also takes a long time for evidence to be implemented in health care [3]. Delays of many years are common, caused by the need to acquire research grants, recruit staff, obtain ethical approval, set up the study, recruit participants, collect and analyse data,

Correspondence: Prof. Dr. med. Claudia Kuehni Institute of Social and Preventive Medicine University of Bern Mittelstrasse 43 CH-3012 Bern claudia.kuehni[at] ispm.unibe.ch write up and publish the results, and integrate these results into current standards of care. Paediatric research lags behind adult research for various reasons, including that the paediatric population is small, many paediatric health conditions are rare and ethical requirements are high. Given these constraints, results from studies in adults are often extrapolated to children [4, 5]. However, because of the important changes that occur during their development, children differ fundamentally from adults in many aspects. These include large age-related differences in susceptibility to environmental influences, in disease manifestations, in the adequacy and performance of diagnostic tests, in drug disposition, and in responses to treatment [6].

The digitalisation of health records could significantly improve the evidence for paediatric medicine and rare diseases as it potentially allows easy and fast access to clinical data from routine patient encounters. It could make clinical research faster and cheaper and make its results more representative of the patients typically seen in health care. Electronic health records (EHR) are widely used in hospitals to document clinical and administrative information about patient encounters. Unfortunately, EHR are rarely standardised within and between institutions and data are often entered into open text fields, resulting in unstructured data. Research on rare diseases relies on data from multiple centres and is limited by the time and costs required to extract and recode these data into a common format. Such data abstraction is particularly challenging when the original data are unstructured [7, 8]. Natural language processing and machine learning methods are increasingly being used to process unstructured data and make them available to research; however, many challenges remain [9]. Furthermore, retrospective standardisation often leads to a loss of information and impairment of data quality. These limitations could largely be circumvented if the original data were recorded in a structured and standardised way [10, 11]. A common EHR architecture allowing structured data capture during routine medical encounters could enable rapid analysis of healthcare data followed by speedy feedback of the knowledge generated into the same health care settings, a process called a learning health system [12, 13]. The aim of our project, which we have named SwissPed-

Data, is to facilitate paediatric clinical research by improv-

LIST OF	ABBREVIATIONS
CDE	Common data element
EHR	Electronic health record
ISPM Berr	1
	Institute of Social and Preventive Medicine, University of Bern
PEDSnet	A multi-specialty network that conducts observational research and clinical trials across multiple children's hospital health systems in the US (www.pedsnet.org)
PECARN	Pediatric Emergency Care Applied Research Network
SPHN	Swiss Personalized Health Network (https://sphn.ch/)
SwissPed	Data
	"Harmonizing the collection of health-related data and biospecimens in pediatric hospitals throughout Switzer- land", an infrastructure development project of the SPHN funded in 2017
SwissPed	Net Swiss Research Network of clinical Pediatric Hubs (www.swisspednet.ch)

ing and standardising the quality of data generated by paediatric health care in Switzerland. To achieve this, we first assessed the status quo, i.e., the relevant aspects of paediatric care for which data are collected, the way these data are recorded, and the data management systems used in the participating paediatric hospitals in Switzerland. Second, we developed and approved a standardised paediatric set of common data elements (CDEs) for EHR across Switzerland by conducting a multi-stage consensus finding process among general paediatricians and paediatric subspecialists of university and cantonal children's hospitals. This paper describes the status quo of the project, the process of standardisation and the resulting set of CDEs: SwissPedData, Version 1.0.

Methods

SwissPedData taskforce

SwissPedNet, the research network of Swiss Children's hospitals (https://www.swisspednet.ch/home/), received an infrastructure grant from the Swiss Personalized Health Network (SPHN) to develop a common data structure in paediatric hospitals and launched SwissPedData with the support of the Swiss Society of Paediatrics (https://www.paediatrieschweiz.ch). SPHN, an initiative of the Swiss Federal Government, aims to achieve a nationwide interoperability of health data produced in university hospitals (https://sphn.ch). SPHN funds the development of infrastructures that make health data shareable for research, following a decentralised approach where data remain in each hospital. Data sharing should become possible either through the direct transfer of individual health data or through distributed analyses, whereby the data do not travel, but are processed decentrally by algorithms and then only data summaries and results are transferred to a central location [14]. SwissPedData is coordinated by a taskforce that consists of a core team at the Institute of Social and Preventive Medicine, University of Bern (ISPM Bern) and representatives from all participating hospitals (fig. 1). All the university hospitals (Basel, Bern, Geneva, Lausanne and Zurich) and three cantonal children's hospitals (Lucerne, St Gallen and Ticino) participated. The clinical directors of each hospital proposed one senior physician to represent the hospital's management board and one junior physician to represent the house officers and registrars who enter the most data into the EHR. The directors also suggested senior physicians representing general paediatrics and all major paediatric subspecialties for collaboration as experts on the Delphi panel. Each hospital suggested at least one expert for general paediatrics and one for each subspecialty. These were then contacted by the core team. Distinct panels were set up for the following subspecialties: paediatric cardiology, endocrinology, gastroenterology, allergy/immunology, infectious diseases, metabolic diseases, nephrology, neurology, pulmonology and rheumatology. Paediatric oncology and neonatology were considered separately because standardised datasets for these subspecialties have already been developed by the Swiss Neonatal Network & Follow-Up Group (Swiss-NeoNet, https://www.neonet.ch/swissneonet) [15] and the Childhood Cancer Registry (https://www.childhoodcancerregistry.ch) [16]. Both datasets have been in use for

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many years and have been continuously refined and thus could be included directly in SwissPedData without further discussions. A related project is developing a set of CDEs for paediatric emergency medicine using the same approach. The results of that effort will be reported separately.

SwissPedData scope

SwissPedData focuses on the standardisation of the documentation of clinical encounters by paediatricians in children's hospitals. This documentation encompasses medical history, physical examination, investigations, diagnosis, treatment and procedures. It excludes laboratory data and biospecimens, as these types of data are usually not entered into EHR by the clinicians themselves. Other SPHN-funded projects are working towards the harmonization of laboratory data in Switzerland (https://sphn.ch/fr/network/project-overview/).

Preparatory steps

To prepare the ground for determining the new set of CDEs, the core team assessed the current status of clinical data documentation during routine encounters in participating hospitals and in ongoing clinical registries and cohort studies. They then searched the literature for other initiatives aiming to standardise paediatric EHR (fig. 2). The core team visited each participating hospital and collected clinical data entry forms and information on the EHR system used and on the degree of digitalisation of health records. The team identified any large existing national or regional clinical paediatric registries and cohort studies via the registry centre (https://www.paediatrieschweiz.ch/ swisspedregistry/) and the clinical hubs of SwissPedNet and through information obtained from the task force members of the participating hospitals. The core team collected metadata describing the datasets collected in these registries and cohort studies and investigated the content and format of the variables.

The core team also conducted a non-systematic, focused literature search to identify approaches to standardising paediatric data across multiple centres in other countries. The reference lists of the relevant publications identified were also scanned.

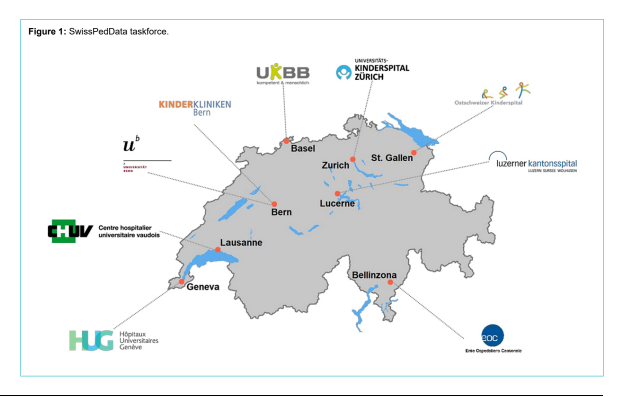
Selection of candidate common data elements for SwissPedData

Based on the information gained in the preparatory phase, the core team defined an initial list of CDEs to be considered for inclusion in the main module (general paediatrics) of SwissPedData. This was done based on an overview of the clinical data routinely documented in the hospitals; the variables collected in ongoing clinical cohort studies and registries; and the datasets of similar international initiatives. The initial list of CDEs was further refined during a two-day retreat held at the ISPM Bern with an interdisciplinary group including six paediatricians, three paediatric epidemiologists and two paediatric registry managers.

For each paediatric subspecialty, the initial list of candidate CDEs was drafted by the core team together with one hospital paediatrician who represented the subspecialty. This first draft was based on existing datasets specific to each subspecialty, such as large cohort studies or clinical registries, and/or on expert opinion (fig. 2, selection of candidate CDEs).

Reaching a consensus: the Delphi process

The consensus finding process aimed to reach agreement on 1) a list of CDEs for SwissPedData, 2) a standardised answer format for each CDE and 3) a classification of each CDE as either mandatory, recommended or optional. Starting with the initial selection of candidate CDEs, we implemented four Delphi rounds, consisting of one face-to-



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face meeting and three online surveys, to obtain a final set of CDEs based on a broad consensus (fig. 2). The Delphi method achieves consensus through a multi-round iterative process that involves eliciting opinions from experts and controlled feedback from the coordinating team [17, 18]. The same basic scheme was followed for the main general paediatric module and for each of the subspecialty modules. All experts were invited to each round, irrespective of whether or not they had given inputs in the previous rounds. For each online survey, the experts were asked to complete the questionnaire within two weeks. Those who had not responded within one week received a reminder email. The online surveys were programmed with the soft-

Figure 2: Consensus finding process followed to define SwissPedData, a set of CDEs for recording routine encounters in children's clinics in Switzerland. CDE: Common Data Element

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I. Source of variables

- a) Clinical forms used currently in the Swiss Children's hospitals
- b) Existing clinical cohort studies and registries in Switzerland
- c) Published datasets from similar initiatives, in particular PEDSnet
- d) Suggestions from taskforce members

II. Selection of candidate CDEs

General paediatrics: Initial list (based on sources a, b and c) (n=150) was refined during a 2-days interdisciplinary retreat oaf paediatricians, epidemiologists and registry managers.

Subspecialty modules: The core team drafted a list of candidate variables using existing datasets specific to each subspecialty (sources a, b and c) in collaboration with one representative per subspecialty.

III. Set up of the Delphi process

Eleven working groups:

- One group for general paediatrics
- One group per subspecialty: cardiology, endocrinology, gastroenterology, immunoallergology, infectiology, metabolic diseases, nephrology, neurology, pulmonology, rheumatology
- Neonatology and paediatric oncology had already defined standard dataset prior to this project
- Paediatric emergency medicine is currently going through the procedure.

Experts were invited to each round, independent of their participation in previous rounds.

Delphi process

Round 1: Online survey (1 per subspecialty)

- Decision for inclusion or exclusion of candidate CDEs, according to relevance for research and clinical work. Inclusion if > 80% agreed, exclusion if >80% agreed. The others were classified as controversial.
- Included variables classified as mandatory, recommended or optional.
- Suggestion of new variables (classified all as controversial)

Round 2: Face-to-face meeting (1 per subspecialty)

- Discussion of all controversial CDEs (< 80% agreement or variables newly suggested in first round) until a group consensus was reached
- · Definition of answer choices of included variables

Round 3: Online survey (1 per subspecialty)

- Experts received excel file containing all the CDEs
- They could propose to add new CDEs, delete CDEs, and define answer choices where they were missing

Round 4: Online survey (all participants)

- All experts received the excel file containing the paediatric core CDEs and all subspecialty CDEs
- last input (minor suggestions) and final approval of entire dataset

SwissPedData, Version 1.0

General paediatrics: 76 CDEs, Cardiology: 11 CDEs; Endocrinology: 59 CDEs; Gastroenterology: 17 CDEs; Allergy/Immunology: 29 CDEs; Infectious diseases: 47 CDEs; Metabolic diseases: 20 CDEs; Nephrology: 34 CDEs; Neurology: 17 CDEs; Pulmonology: 31 CDEs; Rheumatology: 16 CDEs; Neonatology (SwissNeonet); Oncology (Childhood Cancer Registry); Paediatric Emergency care: ongoing

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ware SurveyMonkey Inc., San Mateo, California, USA and analysed using Microsoft Excel.

In the first round, the experts evaluated the candidate CDEs according to their relevance for research and clinical work (fig. 2, round 1). Each expert was asked to vote for the inclusion or exclusion of each candidate CDE and to suggest any additional CDEs. The questions were: "please state for each of the proposed variables (CDEs) below whether you think they should be included in this subspecialty module of SwissPedData" and "would you add other variables (CDEs)?". When opting for inclusion of a CDE, experts were further asked to classify the CDE as "mandatory", "recommended" or "optional". We retained CDEs that reached 80% for inclusion (designated as agreed) and excluded CDEs for which 80% of experts voted for exclusion. All other CDEs, including the additional CDEs suggested by the experts, were classified as "controversial". There is no standard level of consensus in the literature, but levels ranging from 50% to 80% are commonly used [19, 20].

The second round consisted of face-to-face meetings, which were moderated by the core team and held at the ISPM Bern. During the face-to-face meetings, participants discussed all controversial CDEs and the additional CDEs suggested in the first online survey. They also agreed on standardised answer formats for the included CDEs. Eligible answer formats were a date, a date and time, a number, a binary response (e.g., yes/no), standardised response options or free text. When the discussions did not lead to a consensus, we used majority voting. Each face-to-face meeting lasted about three hours.

The third round was another e-Delphi survey, with participants being asked to check if key CDEs for their discipline were missing and to propose standardised answer formats or response options where these were missing.

In the **fourth and final round**, the agreed CDEs and answer formats were sent by email to all the experts for any last inputs and final approval.

Ethical approval was not required for this study, which did not involve the collection or use of patients' data.

Results

Current status of EHR in participating hospitals and existing initiatives aiming to standardize paediatric data

The eight participating hospitals were using different clinical systems for EHR from various vendors (table 1). Their degree of digitalisation varied: while some hospitals were using EHR for all care processes, others were only doing so for some. For example, all hospitals were recording clinical notes relating to inpatients electronically, but only half of them were using electronic drug prescriptions at the time of the survey.

We identified 5 paediatric cohort studies and 25 paediatric clinical registries with a nationwide or multiregional reach (appendix 1). The focused literature search identified four projects with similar goals in other countries, namely PECARN (Pediatric Emergency Care Applied Research Network), PHIS+ (Pediatric Health Information System), PROS (Pediatric Research in Office Settings) and PED-Snet. The initiative most similar to ours was PEDSnet, an American national paediatric learning health system that was founded in 2014 by eight children's hospitals, primarily to obtain child-specific data on the efficacy and safety of new and approved drugs [21] (https://pedsnet.org/data/). Currently, PEDSnet hosts analysis-ready, standardised longitudinal data from the primary, secondary and tertiary care of over 6.5 million patients. PEDSnet uses a common interoperable data platform that optimises the use of EHR, ensuring that data are entered once only. The collected data include demographics, vital status, encounters, diagnoses, vital signs, treatment and immunisations, among others (https://pedsnet.org/data/common-data-model/).

Consensus finding process (Delphi method)

Clinical directors proposed 121 experienced general paediatricians and subspecialists for the Delphi process, of whom 119 agreed to participate. Of these, 73 took part in the first round (online survey), 45 attended the second round (face-to-face meetings), 58 commented in the third round of the Delphi process and 68 gave their final approval of the dataset (appendix 2). The working groups contained between 7 and 14 members. All disagreements could be settled during the process through majority voting or through discussions. Most disagreements were about answer format rather than about which CDEs should be included in SwissPedData.

Table 1:

Electronic health records systems used in Swiss children's hospitals and digitalization of clinical documentation.

Children's hospital	Main IT system	Emergency clinical notes	Outpatient clinical notes	Inpatient	Inpatient			
				Clinical notes	Drug prescription	Vital signs		
Basel	Phoenix	E	E+P	E	P	E		
Bellinzona	DPI	E	E	E	E	E		
Bern	ipdos	E	E + P	E	E	E		
Geneva	DPI	E	E	E	E	E		
Lausanne	Soarian	E	E	E	E	E		
Luzern	Epic/LUKiS	E	E	E	E	E		
St.Gallen	KISIM	E	E	E	P	E + P		
Zürich	Phoenix	E	E	E	E	E		

E: Electronic, P: Paper

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SwissPedData (Version 1.0)

SwissPedData consists of 336 CDEs: 76 in the main module on general paediatrics and between 11 and 59 in each of the 10 subspecialty modules (table 2 and appendix 3). The main module covers aspects concerning all paediatric patients, whether they are outpatients or inpatients. The subspecialty modules cover aspects specific to paediatric subspecialties that are not already covered by the main module. Each module is formally structured into the same nine domains representing all care processes: 1. Care Site, 2. Demographics, 3. Medical History, 4. Physical Examination, 5. Clinical Scores, 6. Investigations, 7. Diagnosis, 8. Treatment, and 9. Equipment and Procedures. These represent domains commonly covered by EHR. The Care Site domain contains administrative data related to the hospital and to patient encounters. It includes type of admission, length of stay and scheduled follow-up. The Demographics domain contains demographic data, for example date of birth, gender, address, and country of birth. The Medical History and Physical Examination domains include clinical information such as birth history, family history, symptoms, medications and vital signs. The Clinical Scores domain contains specific scores, for example triage scale for emergency department patients or developmental tests. The Investigations domain contains data on investigations performed, such as lung function, renal ultrasound or blood glucose monitoring for patients with diabetes. The Diagnosis domain includes diagnosis and date of diagnosis, as well as diagnosis classifications such as Online Mendelian Inheritance in Man (OMIM) codes. The Treatment domain contains data on medications prescribed and administered in hospital, treatment adverse events and reasons for discontinuation of treatment. The Equipment and Procedures domain contains data on procedures performed on the patient, such as dialysis.

The full set of CDEs is shown in appendix 3, which provides a complete list of all agreed CDEs along with their description, answer format and standardised response options, and importance (mandatory, recommended or optional). Answer choices are number, binary or standardised options, or free text. When the "standardised option" format is used, specific value sets are defined. The CDEs will be implemented in children's hospital EHR depending on their importance, categorised as mandatory, recommended or optional. Mandatory CDEs must be implemented in EHR by all participating hospitals. Recommended CDEs should be implemented and optional CDEs may be implemented at the discretion of each hospital.

Examples of mandatory CDEs are vital parameters in the main module (general paediatrics) or "route of feeding" in the gastroenterology module. In the latter case, "route of feeding" will be recorded with standardised response options (oral, gastrostomy, naso/orogastric tube, intravenous, other). An example of a recommended CDE is "seizure type according to the ILEA 2017 classification of seizures" in the neurology module. "Opening pressure at lumbar puncture" is an optional CDE in the same module (appendix 3).

Discussion

We developed SwissPedData, a standardised national set of CDEs designed to collect clinical data during paediatric routine encounters in a harmonised way. It is the result of a broad consensus between general paediatricians and paediatric subspecialists from eight university and cantonal children's hospitals in Switzerland. It describes all processes of paediatric medical care including clinical and paraclinical assessment, diagnosis, treatment, disposition and care site. Each part of the dataset follows the usual structure of the EHR to allow easy implementation.

Clinical data standardisation for a Swiss paediatric learning healthcare system

SwissPedData aimed to standardise items up-front at the point of data entry. Prospective, standardised recording of routine clinical encounters avoids duplicate entry into research databases. However, this should not happen at the expense of an increase in documentation time by clinicians, a concern raised during our Delphi process. To avoid this pitfall, we focused primarily on data elements that are not only useful for research, but also for clinical work, and included CDEs that are routinely documented in paediatric EHR. SwissPedData is not comprehensive and much of the clinical documentation will remain unstandardised to preserve the rich narrative details that are difficult to capture in standardised fields but are nevertheless important for daily clinical work. These narrative data could be used by researchers applying text-mining approaches. SwissPedData could also be supplemented by questionnaires to patients and their families. The implementation of SwissPedData in EHR will include careful attention to clinician workflow to minimise potential negative consequences of standardisation.

Table 2:

Examples of common data elements (CDEs) of the core module (general paediatrics) of SwissPedData.

Common data element	Format	Standardized response options	Importance	Comment / description
Follow-up after discharge / consul- tation	Standardised op- tions	General paediatrician, General practitioner, Subspecialist, Nurse, None	Mandatory	Scheduled follow-up at discharge
Country of birth	Standardised op- tions	Swiss Federal Statistical Office: ISO code of the country of ori- gin	Mandatory	Country of birth of the patient
Birth weight	Number		Mandatory	Weight at birth in kg
Heart rate	Number		Mandatory	Heart rate in beats per minute
Glasgow Coma Scale	Number		Mandatory	
Indication for imaging study	Free text		Mandatory	Medical reason for the radiological study
Drug name	Standardised op- tions	International non-proprietary name	Mandatory	Name of the drug(s) received as in- patient
Equipment date of insertion	Date	YYYY-MM-DD	Mandatory	

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SwissPedData is designed to provide a basis for a paediatric learning health system in Switzerland in which clinical data from different children's hospitals can be combined to rapidly generate new knowledge relevant for day-to-day practice and translate it into improved health care for children. Existing learning health systems in other countries, such as PEDSnet in the US, have demonstrated that a paediatric learning health system can improve the health outcomes of children [22, 23]. Examples include the rapid identification of children suffering from glomerular diseases for clinical trials [24], comparing weight loss and safety among bariatric procedures using EHR data [25] and, recently, describing the epidemiology of paediatric patients infected by SARS-CoV-2 [26].

Strengths and limitations

The main strength of SwissPedData is that it is based on broad agreement between paediatricians from all university and cantonal paediatric clinics in Switzerland. The project received strong support from all clinical directors of Swiss children's hospitals, from the paediatric research network SwissPedNet and from more than 100 experienced paediatricians who participated in its development. SwissPedData emphasises the prospective collection of standardised data, which can greatly reduce the time and costs needed for data preparation and analysis as it avoids the need for retrospective standardisation or double entry. Our consensus finding approach could be adapted for use by other medical specialties that wish to define CDEs in the future.

SwissPedData has a number of omissions that are intentional. First, we focused on standardising a minimal set of items that are particularly relevant and specific to paediatric routine care. SwissPedData will thus not replace existing terminologies for clinical health care such as SNOMED-CT. Rather, standardised data from SwissPed-Data can in the future be mapped to SNOMED-CT. Second, SwissPedData does not include laboratory data or detailed radiological data. However, other projects within the SPHN are working on the standardisation of these domains. The goal is to link the standardised paediatric data extracted from EHR with laboratory data standardised thanks to other SPHN projects like L4CHLAB. Such linkage can be done through hospital patient IDs, or with birth dates and names. Third, SwissPedData will need to be translated into the Swiss national languages before implementation in children's hospital EHR.

SwissPedData is adapted to the Swiss context

The Swiss healthcare system is decentrally structured, with cantons being responsible for the organisation of local health care, and therefore is highly heterogeneous. As a consequence, children's clinics are relatively small, with catchment areas of a few 100,000 children. Obtaining sufficient patient samples for research is only possible by combining data from multiple hospitals, especially for rare conditions. However, given the differences in EHR and IT systems between hospitals, this results in long delays and huge costs for obtaining, extracting, standardising and cleaning the heterogeneous data. SwissPedData, once implemented in all children's clinics, will allow researchers to identify and recruit patients for clinical trials in real time,

to conduct retrospective studies with high-quality data, and to conduct nested prospective studies. As examples, participants of the "Clinical Data for Paediatric Research: the Swiss Approach" symposium held in 2019 drafted sketches of the following research projects based on SwissPed-Data: a diagnostic study on the validity of the tests used for auditory screening in newborns; a benchmarking study assessing the quality of treatment for bronchiolitis across different children's hospitals; a cohort study on the incidence of hearing loss after treatment with aminoglycosides in infancy; a cohort study on kidney injury after treatment with acyclovir; and a randomised clinical trial comparing the effectiveness of different treatment regimens for type 1 diabetes. Some of these project sketches suggested complementing the hospital dataset with available data from other sources such as the federal statistical office or laboratory data, or through the collection of additional data through questionnaires or specific examinations.

Comparison with other projects

SwissPedData is closely aligned with PEDSnet, a USbased paediatric clinical data research network [21]. PED-Snet includes eight children's hospitals that provide care for 2.8% of the paediatric population in the USA (2.1 million patients) [21]. The database contains standardised clinical data from EHR covering 6.5 million children (https://pedsnet.org/) and forms the basis of a high-quality research programme and learning health system. Studies based on PEDSnet data cover a wide range of research topics and study designs in paediatrics, including descriptive epidemiology [27], computable phenotyping [24], longitudinal observational studies [28] and comparative effectiveness [29]. PEDSnet established a common data model (PEDSnet CDM) from the beginning of their network, based on the Observational Health Data Sciences and Informatics collaborative's OMOP common data model. With SwissPedData, we defined a list of priority CDEs that can be mapped to SNOMED-CT in the future.

PEDSnet may also serve as a role model for the implementation of SwissPedData and has already demonstrated its usefulness for observational and interventional research and for the standardisation of care processes. Each hospital that participates in PEDSnet regularly extracts the standardised data from its EHR in a predefined way [21].

Another notable example of harmonised clinical datasets in paediatrics is the Pediatric Emergency Care Applied Research Network (PECARN), an EHR-based registry that has harmonised data in the paediatric emergency setting in seven American paediatric emergency departments to make it usable for paediatric research. PECARN uses data resources from seven paediatric emergency departments of four hospitals [30].

Outlook and next steps

All participating hospitals are committed to implementing SwissPedData in their EHR by 2024. A committee of clinicians and IT specialists in each hospital will supervise the implementation process. The EHR of children's hospitals will be restructured at the front-end to include SwissPed-Data CDEs. Practically, this means that EHR as seen by their users (physicians) will include the CDEs of Swis-

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sPedData. For some hospitals, where this is not possible in the short term, we will also offer the possibility of transforming the source data to the CDEs and contributing it to the common dataset. SwissPedData is intended to evolve and be adaptive to existing needs. The set of CDEs can be expanded to cover more domains or to include more CDEs per domain. Temporary CDEs can be added for nested research projects. Self-completed or parent-completed questionnaires can add information relating to a child's family and home environment, which is not routinely recorded in EHR. Data from primary care encounters could also be integrated in the future.

In ongoing work, other prerequisites for the implementation of SwissPedData are being put into place: a general consent form for use of the data from patients and caregivers, a data transfer and use agreement (DTUA) between the clinics, and protocols for obtaining ethics approval for SwissPedData overall and for individual research projects. Some aspects are being dealt with within other infrastructure development projects of the SPHN network (www.sphn.ch), namely the C3-Study (citizen centred consent) project and the E-General Consent project. Furthermore, the SPHN provides legal agreement templates, including a DTUA and an ethical framework for all its projects. It is important to stress that only data useful for the clinical management of the patient will be recorded and that these data will always be stored by each children's hospital as part of the patient's file. The only difference to the previous procedure is that some of these clinical data will be recorded in a standardised way. To have access to these data for research, researchers will have to get ethical approval as usual.

It is planned that SwissPedData will be implemented as a project on the SPHN infrastructure for data exchange, so that data can in future be accessed through a central portal. The SPHN Data Coordination Centre and BioMedIT (https://sphn.ch/network/projects/biomedit/) can provide assistance and the infrastructure for this. The aim is to keep SwissPedData CDEs harmonized with the future releases of the SPHN dataset (https://sphn.ch/services/documents/ technical-documents/). An additional central coordination center for paediatric research should facilitate communication between children's clinics, international research partners and funders, and also assist researchers in writing grant applications, obtaining ethical approval and accessing the necessary datasets. The resources needed to maintain SwissPedData will require the support of a central coordination center encompassing an experienced researcher ideally with a background in paediatrics, an IT specialist, andlocal support of the responsible clinicians and IT specialists in each hospital. Funding for the implementation and maintenance of SwissPedData will need to be secured. Potential funding sources are participation in suitable calls for proposals, charging cost-covering fees for services provided by SwissPedData and collaboration with industry, for example for post-marketing studies. Collaborations with international partners such as PEDSnet are foreseen, and first exchanges have occurred.

In conclusion, SwissPedData defines a set of common data elements (CDEs) for clinical paediatric care based on a broad agreement among university and cantonal paediatric hospitals in Switzerland. With SwissPedData, Swiss children's hospitals will be able to provide researchers with standardized, high-quality routine clinical paediatric data in the near future. SwissPedData will provide the basis for a learning health system for paediatric care in Switzerland.

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Appendix

Appendix 1: Paediatric registries and cohort studies in Switzerland

Registry / Cohort Study	Coverage
Childhood Cancer Registry ChCR	National
Swiss Primary Ciliary Dyskinesia Registry (SPCDR)	National
Swiss Cerebral Palsy Registry (Swiss-CP-Reg)	National
Swiss Growth Registry (SGR)	National
Swiss Paediatric Airway Cohort (SPAC)	National
Swiss Paediatric Renal Registry (SPRR)	National
Swiss Rare Disease Registry (SRDR)	National
Swiss Registry for Neuro-Muscular Disorders (Swiss-Reg-NMD)	National
Cystic Fibrosis (CF) newborn screening	National
Iuvenile Inflammatory Rhumatism cohort (JIRcohorte)	European
SwissNeoNet Minimal Neonatal Data Set (MNDS)	National
SwissNeoNet National Asphyxia and Cooling Registry (ASP)	National
SwissNeoNet Follow-Up (FU)	National
Swiss NeuroPaediatric Stroke Registry (SNPSR)	National
Swiss Congenital Lung Anomalies (CLA) Registry	National
Swiss Mother and Child HIV Cohort Study (MoCHiV)	National
Swiss Cystic Fibrosis Infant Lung Development (SCILD) cohort	National
Swiss Pediatric Surveillance Unit (SPSU)	National
Swiss Hemophilia Registry (SHN)	National
COST Action BM1105 Patient Registry - GnRH Network	European
Registry of congenital anomalies in the canton of Vaud	National
Swiss Cleft lip and Palate Registry	National

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Swiss Biliary Atresia Registry	National
Swiss registry on Autoimmune Hepatitis	National
European Registry for Primary Immunodeficiencies (ESID registry)	European
European Cystic Fibrosis Patient Registry (ECFSPR)	European
European Childhood Interstitial Lung Disease (chILD-EU) Registry	European
Swiss Inflammatory Bowel Disease Pediatric Cohort Study (Swiss IBD Pediatric Cohort Study)	National
Splenectomy Registry	Global
Pediatric and Adult Intercontinental Registry on Chronic ITP (PARC-ITP registry)	Global
Diabetes Patienten Verlaufsdokumentation Registry (DPV)	European

Paediatric specialty	Number of experts		Delphi process, numb	er of experts involved	
	invited	1st round	2nd round	3rd round	4th round
General paediatrics	14	8	4	4	5
Cardiology	13	10	4	7	8
Endocrinology	12	7	6	8	9
Gastroenterology	10	8	4	4	6
Allergy/Immunology	12	6	4	8	7
Infectiology	11	8	5	6	9
Metabolic diseases	8	7	2	4	3
Nephrology	12	3	4	5	4
Neurology	14	5	4	3	5
Pulmonology	11	8	5	4	6
Rheumatology	8	3	3	5	6

Appendix 3 : SwissPedData Common Data Model (CDM), Version 1.0

Module(s)	Common Data Element	Format	Standardized response options	Importance	Comment / Description
Domain: Care site					
General paediatrics	Type of admission	standardized options	Elective admission Emergency admission	Mandatory	
General paediatrics	Provenance	standardized options	Other hospital Emergency department Home Other	Mandatory	
General paediatrics	Care Handling Type	standardized options	Inpatient Outpatient	Mandatory	
General paediatrics	Visit start date and time	datetime	YYYY-MM-DD hh:mm:ss	Mandatory	Datetime at which the interaction between individual and the care provider institute started
General paediatrics	Visit end date and time	datetime	YYYY-MM-DD hh:mm:ss	Mandatory	Datetime at which the interaction between individual and the care provider institute stopped
General paediatrics	Datetime of admission	datetime	YYYY-MM-DD hh:mm:ss	Mandatory	Datetime of patient's admission to the care provider institute
General paediatrics	Discharge destination	standardized options	Home Other hospital Institution Other	Mandatory	Location to which the patient is discharged
General paediatrics	Follow-up after discharge / consultation	standardized options	General paediatrician General practitioner Subspecialist Nurse None	Mandatory	Scheduled follow-up at discharge
General paediatrics	Translator needed	standardized options	Yes No Unknown	Recommende d	Translator needed for communication between patient and healthcare team
General paediatrics	Hospital	standardized options	See comments	Mandatory	Standardized response options will be name of participating children's hospitals

Module(s)	Common Data Element	Format	Standardized response options	Importance	Comment / Description
General paediatrics	Department	standardized options	See comments	Mandatory	Standardized response options will be name of departments of participating children's hospitals
General paediatrics	Unit	standardized options	See comments	Mandatory	Standardized response options will be name of units of participating children's hospitals
Infectious diseases	If coming from another hospital: Country	standardized options	Swiss Federal Statistical Office: ISO code of the country of origin	Mandatory	Country of originating hospital
Domain: Demographics				-	
General paediatrics	Patient Datetime of birth	datetime	YYYY-MM-DD hh:mm:ss	Mandatory	Datetime of birth of the patient
General paediatrics	Country of birth	standardized options	Swiss Federal Statistical Office: ISO code of the country of origin	Mandatory	Country of birth of the patient
General paediatrics	Place of birth (CH)	number	Postal code (PLZ/NPA)	Mandatory	Municipality of birth of the patient if in Switzerland, coded by postal codes (PLZ/NPA).
General paediatrics	Patient administrative gender	standardized options	Male Female Other	Mandatory	
General paediatrics	Address (postal code)	number	Postal code (PLZ/NPA)	Mandatory	Current address of the patient, coded by postal codes (PLZ/NPA). Exact address should also be recorded
General paediatrics	Nationality	standardized options	Swiss Federal Statistical Office: ISO code of the country of origin	Mandatory	Current nationality of the patient
General paediatrics	Date of immigration	date	YYYY-MM-DD	Mandatory	Date of first immigration to Switzerland if born abroad

Module(s)	Common Data Element	Format	Standardized response options	Importance	Comment / Description
Infectious diseases	If immigrant: Type of residency permit	standardized options	B C G L F N S undocumented	Optional	
Metabolic diseases	Ethnicity of the mother	standardized options	See comments	Optional	Standard classification to be defined
Metabolic diseases	Ethnicity of the father	standardized options	See comments	Optional	Standard classification to be defined
Rheumatology Pulmonology	Ethnicity of the patient	standardized options	See comments	Optional Recommende d	Standard classification to be defined. Optional for rheumatology, recommended for pulmonology.
Domain: Medical history		I	- L	i	1 ·
General paediatrics	Reason for consultation / for admission	free text		Mandatory	Main reason for consultation or for admission. Standard classification not defined.
General paediatrics	Current medications: Drug name	standardized options	International non-proprietary name	Mandatory	Name of the drug(s) received as inpatient
General paediatrics	Current medications: Route of administration	standardized options	Oral Intravenous Subcutaneous Intramuscular Intrathecal Rectal Inhalation Cutaneous Ocular Nasal Otic Other	Mandatory	
General paediatrics	Current medications: Frequency of administration	number		Mandatory	Number of administrations per 24 hours

Module(s)	Common Data Element	Format	Standardized response options	Importance	Comment / Description
General paediatrics	Current medications: Dose	number		Mandatory	Dose given at each administration of the drug
General paediatrics	Current medications: Dose unit	standardized options		Mandatory	List of possible units to be defined
General paediatrics	Use of complementary medicine	yes/no		Optional	Patient treated with complementary medicine at home or in hospital
General paediatrics	Birth weight	number		Mandatory	Weight at birth in kg
General paediatrics	Birth length	number		Mandatory	Length at birth in cm
General paediatrics	Birth's head circumference	number		Mandatory	Head circumference at birth in cm
General paediatrics	Delivery mode	standardized options	Caesarean section Instrumental vaginal delivery Spontaneous vaginal delivery	Mandatory	Birth delivery mode
General paediatrics	Gestational age	number		Mandatory	Post-menstrual age at birth in week and days
General paediatrics	Apgar score 1 min	number		Recommende d	Apgar score 1 min after birth
General paediatrics	Apgar score 5 min	number		Recommende d	Apgar score 5 min after birth
General paediatrics	Apgar score 10 min	number		Recommende d	Apgar score 10 min after birth
General paediatrics	Mother's year of birth	number		Mandatory	Year of birth of the mother
General paediatrics	Father's year of birth	number		Mandatory	Year of birth of the father
General paediatrics	Year(s) of birth of sibling(s)	number		Mandatory	Year of birth of sibling(s) if any
General paediatrics	Drug allergies	standardized options	International Nonproprietary Name of drug	Mandatory	Known drug allergies
General paediatrics	Documented food allergies	yes/no		Mandatory	Presence of any documented food allergy
Endocrinology	Age at menarche	number		Mandatory	Age at menarche in years
Endocrinology	Age at thelarche	number		Mandatory	Age at thelarche in years
Endocrinology	Age at pubarche	number		Mandatory	Age at pubarche in years

Module(s)	Common Data Element	Format	Standardized response options	Importance	Comment / Description
Endocrinology	Single/Multiple birth	number		Recommende d	Number of children born from the same pregnancy as the patient's
Endocrinology	Neonatal hypoglycaemia	standardized options	No Yes, confirmed Yes, reported by patient/family	Recommende d	History of hypoglycaemia in the neonatal period
Endocrinology	Neonatal hyperbilirubinemia	standardized options	No Yes, confirmed Yes, reported by patient/family	Recommende d	History of hyperbilirubinemia in the neonatal period (only hyperbilirubinemia treated with phototherapy)
Endocrinology Nephrology	Mother's height	number		Mandatory	Height of the mother in cm
Endocrinology Nephrology	Father's height	number		Mandatory	Height of the father in cm
Endocrinology	Mother's age at menarche	number		Mandatory	Age of the mother at menarche in years
Endocrinology	Father's puberty	standardized options	Normal Early Late	Mandatory	
Endocrinology	Diabetes in first degree relatives	standardized options	No Yes, Type 1 Yes, Type 2 Yes, Monogenic Unknown	Mandatory	Any type of diabetes in a first degree relative
Endocrinology	Thyroid disorder in first degree relative	yes/no		Mandatory	Presence of thyroid disorder in a first degree relative
Endocrinology	Other auto-immune disorders in first degree relative	yes/no		Recommende d	Presence of auto- immune disorder in a first degree relative. With added box for free text to specify the disease.
Endocrinology	Other endocrinopathy in first degree relative	yes/no		Mandatory	Presence of endocrinopathy in a first degree relative. With added box for free text to specify the disease.

Module(s)	Common Data Element	Format	Standardized response options	Importance	Comment / Description
Endocrinology	Fertility problems in first degree relative	yes/no		Recommende d	Presence of fertility problems in first degree relatives. With added box for free text to specify the disease.
Endocrinology	Severe hypoglycaemia (requiring assistance OR coma)	number		Mandatory	Number of events since last visit
Endocrinology	Mild hypoglycaemia (BG < 3.9mmol/l)	number		Mandatory	Number of events per month
Endocrinology	Ketoacidosis	standardized options	No Yes, managed ambulatorily Yes, with hospitalization	Mandatory	History of ketoacidosis
Endocrinology	Diagnostic of obesity in first degree relative	yes/no		Recommende d	Diagnostic of obesity in a first degree relative
Gastroenterology	Nutrition habits	standardized options	No specific diet Vegetarian Vegan Other	Recommende d	Nutrition habits of the patient
Gastroenterology Metabolic diseases	Route of feeding	standardized options	Oral Gastrostomy Naso/orogastric tube Intravenous Other	Mandatory	The route(s) by which the patient is fed
Allergy/Immunology	History of rhinoconjonctivitis	standardized options	Yes, reported Yes, documented No	Mandatory	
Allergy/Immunology	History of atopic dermatitis	standardized options	Yes, reported Yes, documented No	Mandatory	
Allergy/Immunology	History of wheezing	standardized options	Yes, reported Yes, documented No	Mandatory	
Allergy/Immunology	History of asthma	standardized options	Yes, reported Yes, documented No	Mandatory	
Allergy/Immunology	Respiratory support during first hours of life	standardized options	Yes, reported Yes, documented No	Recommende d	Presence of any kind of respiratory support (non- invasive and invasive ventilation) during first hours of life

Module(s)	Common Data Element	Format	Standardized response options	Importance	Comment / Description
Allergy/Immunology	Supplemental O2 during first hours of life	standardized options	Yes, reported Yes, documented No	Recommende d	Supplemental oxygen administered during first hours of life
Allergy/Immunology	Chronic diarrhea	yes/no		Mandatory	
Allergy/Immunology	Number of hospitalisations for IV antibiotherapy in life	number		Mandatory	
Allergy/Immunology	Maximal number of otitis media in one year	number		Mandatory	
Allergy/Immunology	Number of pneumonias in life	number		Mandatory	
Allergy/Immunology	Number of sinusitis in life	number		Mandatory	
Allergy/Immunology	Number of meningitis in life	number		Mandatory	
Allergy/Immunology	Family history of atopic diseases	yes/no		Mandatory	Presence of atopic diseases in a first degree relative
Allergy/Immunology	Family history of immunodeficiency	yes/no		Mandatory	Presence of immunodeficiency in a first degree relative
Allergy/Immunology	Family history of auto-immune disease	yes/no		Mandatory	Presence of auto- immune disease in a first degree relative
Allergy/Immunology	Family history of angioedema	yes/no		Mandatory	Presence of angioedema in a first degree relative
Allergy/Immunology Gastroenterology	Documented food allergy by oral food challenge	yes/no		Mandatory	Presence of any documented food allergy (diagnosed by physician)
Allergy/Immunology	Hymenoptera venom allergies	standardized options	Yes, reported Yes, documented No	Mandatory	Known documented hymenopter allergies
Allergy/Immunology	History of anaphylaxis	standardized options	Yes, reported Yes, documented No	Mandatory	History of anaphylaxis
Allergy/Immunology	Autoimmune or inflammatory diseases in the patient	yes/no		Mandatory	Classification for type of autoimmunity (organ specific or systemic) and organ(s) involved will be further defined.
Infectious diseases	History of fever (>38°C)	yes/no		Mandatory	
Infectious diseases	If history of fever: Number of days with fever	number		Mandatory	
Infectious diseases	History of cough	yes/no		Mandatory	

Module(s)	Common Data Element	Format	Standardized response options	Importance	Comment / Description
Infectious diseases	History of running nose	yes/no		Mandatory	
Infectious diseases	History of diarrhea	yes/no		Mandatory	
Infectious diseases	History of vomiting	yes/no		Mandatory	
Infectious diseases	History of headache	yes/no		Mandatory	
Infectious diseases	Travel history in the last 6 months	standardized options	Swiss Federal Statistical Office: ISO code of the country of origin (selection of >1 possible)	Mandatory	Country(ies) visited in the last 6 months
Infectious diseases	History of tick bite	yes/no		Recommende d	
Infectious diseases	If history of tick bite: Month of tick bite	date	YYYY-MM	Recommende d	
Infectious diseases	History of contact with animals	yes/no	No Yes	Optional	Standard animal list to be defined
Infectious diseases	Pertussis immunization during pregnancy	yes/no		Mandatory	For patients under 6 months of age
Infectious diseases	Influenza immunization during pregnancy	yes/no		Mandatory	For patients under 6 months of age
Infectious diseases	Prolonged rupture of membranes	yes/no		Mandatory	For patients under 1 month of age. Prolonged rupture defined as longer than 18h
Infectious diseases	Maternal GBS colonization	standardized options	Positive Negative Unknown	Mandatory	For patients under 1 month of age
Infectious diseases	Maternal HIV serology	standardized options	Positive Negative Unknown	Mandatory	For patients under 1 month of age
Infectious diseases	Maternal HBsAg	standardized options	Positive Negative Unknown	Mandatory	For patients under 1 month of age
Infectious diseases	Maternal HBsAb	standardized options	Positive Negative Unknown	Mandatory	For patients under 1 month of age
Infectious diseases	Maternal HBcAb	standardized options	Positive Negative Unknown	Mandatory	For patients under 1 month of age
Infectious diseases	Maternal HBeAg	standardized options	Positive Negative Unknown	Mandatory	For patients under 1 month of age

Module(s)	Common Data Element	Format	Standardized response options	Importance	Comment / Description
Infectious diseases	Maternal HCV serology	standardized options	Positive Negative	Mandatory	For patients under 1 month of age
		options	Unknown		month of upe
Infectious diseases	Maternal CMV serology (IgG / IgM)	standardized	Positive	Optional	For patients under 1
		options	Negative		month of age
			Unknown		_
Infectious diseases	Maternal syphilis serology	standardized	Positive	Mandatory	For patients under 1
		options	Negative		month of age
			Unknown		
Infectious diseases	Maternal rubella serology	standardized	Positive	Mandatory	For patients under 1
		options	Negative		month of age
			Unknown		
Infectious diseases	Maternal toxoplasmosis serology	standardized	Positive	optional	For patients under 1
		options	Negative		month of age
			Unknown		
Infectious diseases	Maternal Chagas serology	standardized	Positive	Mandatory	For patients under 1
		options	Negative		month of age
		,	Unknown		
Metabolic diseases	Self-monitoring of blood glucose	yes/no		Optional	Regular self-
					monitoring of blood
					glucose done at
Metabolic diseases	Self-monitoring of ketone bodies	yes/no		Optional	home Regular self-
wielabolic uiseases	Self-monitoring of ketone bodies	yes/10		Optional	-
					monitoring of ketone bodies done at home
Nephrology	Prenatal ultrasound	standardized	Normal	Mandatory	
Nephrology		option	An-/Oligohydramnios	Walluatory	
		option	Polyhydramnios		
			Megacystis		
			Megaureter		
			Bilateral renal pelvis dilatation > 10 mm		
			Bilateral renal pelvis dilatation < 10 mm		
			Unilateral renal pelvis dilatation > 10 mm		
			Renal cysts		
			Renal agenesis or ectopia		
			Multicystic-dysplastic kidney and bladder extrophy		
Nephrology	Family history of renal disease (1st-2nd degree)	yes/no		Mandatory	
Neurology	Seizure type (ILEA 2017 Classification of Seizures)	standardized	Focal Onset	Recommende	Seizure type
		options	Generalized Onset	d	according to the ILEA
			Unknown Onset		2017 classification of
			Unclassified		seizures
Neurology	Family history of neurological diseases	yes/no		Recommende	Family history of any
5.				d	type of neurological
		1			diseases

Module(s)	Common Data Element	Format	Standardized response options	Importance	Comment / Description
Pulmonology	Cough	standardized options	No Yes, acute and dry Yes, acute and wet Yes, chronic and dry Yes, chronic and wet	Recommende d	Cut-off for acute/chronic 4 weeks
Rheumatology	Recurrent fever	yes/no		Mandatory	History of recurrent fever
Rheumatology	History of uveitis	yes/no		Mandatory	Presence of active uveitis
Rheumatology	History of inflammatory skin disease	yes/no		Mandatory	Presence of skin involvement
Rheumatology	Family history of inflammatory rheumatic disease	standardized options	No Yes, without spondyloarthropathy Yes, with spondyloarthropathy	Mandatory	Presence of any rheumatic disease in the family
Rheumatology	Family history of inflammatory skin disease	standardized options	No Yes, without psoariasis Yes, with psoriasis	Mandatory	Presence of any skin disease in the family
Rheumatology	Family history of chronic intestinal diseases	yes/no		Mandatory	Presence of any chronic intestinal disease in the family
Rheumatology	Family history of recurrent fever	yes/no		Mandatory	Presence of recurrent fever in the family
Domain: Physical exami	nation		- +	I	
General paediatrics	Heart rate	number		Mandatory	Heart rate in beats per minute
General paediatrics	Systolic blood pressure	number		Mandatory	Value of the systolic blood pressure in mmHg
General paediatrics	Diastolic blood pressure	number		Mandatory	Value of the diastolic blood pressure in mmHg
General paediatrics	Respiratory rate	number		Mandatory	Respiratory rate in breaths per minute
General paediatrics	Oxygen saturation	number		Mandatory	Measured oxygen saturation in %
General paediatrics	Temperature	number		Mandatory	Measured temperature of the patient in Celsius degrees
General paediatrics	Weight	number		Mandatory	Measured weight of the patient in kg

Module(s)	Common Data Element	Format	Standardized response options	Importance	Comment / Description
General paediatrics	Height	number		Mandatory	Measured height of the patient in cm
General paediatrics	Head circumference	number		Mandatory	Measured head circumference of the patient in cm
Endocrinology	Sitting height	number		Recommende d	Sitting height measured sitting with straight back in in cm
Endocrinology	Arm span	number		Recommende d	Arm span: arms streched horizontally, measurement from fingertip to fingertip in cm
Endocrinology	Waist circumference	number		Recommende d	In cm
Endocrinology	Hip circumference	number		Recommende d	In cm
Endocrinology	Goiter	yes/no		Recommende d	Presence of goiter
Endocrinology	Gynecomastia	standardized options	No Yes, unilateral Yes, bilateral	Recommende d	Presence of gynecomastia
Endocrinology Metabolic diseases	Dysmorphic signs	yes/no		Recommende d	Presence of dysmorphic features. If answer is yes, specification with standardized classification to be defined.
Endocrinology	Cryptorchidism	standardized options	No Yes, unilateral Yes, bilateral	Mandatory	Presence of cryptorchidism
Endocrinology	Insulin injection site	standardized options	Normal Abnormal, lipoatrophy Abnormal, lipohypertrophy	Optional	Inspection of insulin delivery sites
Endocrinology	Retinopathy screening	normal/abnormal		Optional	
Endocrinology	Neuropathy screening performed	standardized options	No Yes, vibration Yes, monofilament	Optional	
Endocrinology	Testis volume right side	number		Mandatory	Volume of right testis in ml
Endocrinology	Testis volume left side	number		Mandatory	Volume of left testis in ml

Module(s)	Common Data Element	Format	Standardized response options	Importance	Comment / Description
Endocrinology	Tanner breast stage	number		Mandatory	
Endocrinology	Tanner pubic hair stage	number		Mandatory	
Endocrinology	Tanner axillary hair stage	number		Mandatory	
Endocrinology	Tanner genital stage	number		Recommende d	
Endocrinology	Breast size	number		Optional	In cm
Endocrinology	Female genital examination	normal/abnormal		Mandatory	
Endocrinology	Penis length	number		Recommende d	In cm
Endocrinology	Chovstek sign	yes/no		Recommende d	Twitching of facial muscles in response to tapping over the area of the facial nerve
Endocrinology	Trousseau sign	yes/no		Recommende d	Carpopedal spasm that results from ischemia
Endocrinology	Thyroid nodule	yes/no		Recommende d	Presence of thyroid nodule
Infectious diseases Metabolic diseases Rheumatology	Hepatomegaly noted at physical examination	yes/no		Mandatory	
Infectious diseases Metabolic diseases Rheumatology	Splenomegaly noted at physical examination	yes/no		Mandatory	
Infectious diseases	Meningeal signs noted at physical examination	yes/no		Mandatory	
Infectious diseases Rheumatology	Skin lesion noted at physical examination	yes/no		Mandatory	
Infectious diseases	Irritability noted during physical examination	yes/no		Mandatory	
Infectious diseases Rheumatology	Adenopathy noted at physical examination	standardized options	No Yes, localized Yes, generalized	Mandatory	
Infectious diseases	Respiratory distress noted at physical examination	yes/no		Mandatory	
Infectious diseases	Conjunctivitis noted at physical examination	yes/no		Mandatory	
Infectious diseases	Prolonged capillary refill time (> 2 sec) noted at physical examination	yes/no		Mandatory	

Module(s)	Common Data Element	Format	Standardized response options	Importance	Comment / Description
Infectious diseases	Signs of dehydration noted at physical examination	standardized options	No Yes, < 5% Yes, 5-10% Yes, >10%	Mandatory	
Metabolic diseases	Skin abormalities	yes/no		Mandatory	Presence of skin abnormalities
Metabolic diseases	Abnormal body proportions	yes/no		Recommende d	Presence of abnormal body proportions
Nephrology	Average 24-hour arterial pressure, systolic	number		Mandatory	In mmHg
Nephrology	Average 24-hour arterial pressure, diastolic	number		Mandatory	In mmHg
Nephrology	Average daytime systolic BP	number		Mandatory	In mmHg
Nephrology	Average daytime diastolic BP	number		Mandatory	In mmHg
Nephrology	Average night-time systolic BP	number		Mandatory	In mmHg
Nephrology	Average night-time diastolic BP	number		Mandatory	In mmHg
Nephrology	Mean Arterial Pressure (MAP)	number		Mandatory	Measured MAP in mmHg
Nephrology	Blood pressure dipping pattern	number		Mandatory	Difference between daytime mean systolic pressure and night-time mean systolic pressure expressed as a percentage of the day value
Neurology	Walking ability	standardized options	Community ambulator Household ambulator Non-ambulatory	Mandatory	
Pulmonology	Auscultation	normal/abnormal		Mandatory	
Pulmonology	Thorax shape	normal/abnormal		Mandatory	The shape of the thorax
Rheumatology	Active arthritis	yes/no		Mandatory	Presence of active arthritis
Rheumatology	If active arthritis: number of joints involved	number		Mandatory	Number of joints involved in active arthritis
Rheumatology	Maximal mouth opening	number		Mandatory	Maximal mouth opening in mm
Rheumatology	Muscle strength	normal/abnormal		Recommende d	Overall muscle strength

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Module(s)	Common Data Element	Format	Standardized response options	Importance	Comment / Description
Domain: Clinical scores					
General paediatrics	Triage scale (ED), type	standardized options	Australasian Triage Scale Canadian Triage Scale Other	Mandatory	Name of the triage scale used
General paediatrics	Triage scale (ED), value	number		Mandatory	Value of the triage scale
General paediatrics	AVPU score	standardized options	Alert Voice Pain Unresponsive	Mandatory	
General paediatrics	Glasgow Coma Scale	number		Mandatory	
Cardiology	Modified Ross heart failure classification for children	standardized options	Class I Class II Class III Class IV	Mandatory	Class I: Asymptomatic Class II: Mild tachypnea or diaphoresis with feeding in infants, dyspnea on exertion in older children Class III: Marked tachypnea or diaphoresis with feeding in infants, marked dyspnea on exertion, prolonged feeding times with growth failure Class IV: Symptoms such as tachypnea, retractions, grunting or diaphoresis at rest

Module(s)	Common Data Element	Format	Standardized response options	Importance	Comment / Description
Cardiology	NYHA classification for adults	standardized options	Class I Class II Class III Class IV	Mandatory	Class I: No symptoms and no limitation in ordinary physical activity Class II: Mild symptoms (mild shortness of breath and/or angina) and slight limitation during ordinary activity. Class III: Marked limitation in activity due to symptoms, even during less- than-ordinary activity. Comfortable only at rest. Class IV: Severe limitations. Experiences symptoms even while at rest. Mostly bedbound patients.
Endocrinology	Endocrinology clinical score type	standardized options	Crook score Billewicz score Ferriman-Gallway score Prader stage External genitalia score	Optional	Type of score
Endocrinology	Endocrinology clinical score result	number	number	Optional	Result of score
Gastroenterology	PCDAI	number		Mandatory	Paediatric Crohn's Disease Activity Index
Gastroenterology	PUCAI	number		Mandatory	Paediatric Ulcerative Colitis Activity Index
Gastroenterology	PYMS score	number		Mandatory	Paediatric Yorkhill Malnutrition Score
Gastroenterology	Bristol stool scale	number		Mandatory	
Allergy/Immunology	SCORAD index	number		Mandatory	SCORing Atopic Dermatitis Index
Metabolic diseases Neurology	Developmental test: Type	standardized options	Bayley II Bayley III Griffith Other	Mandatory	Type of developmental test performed

Module(s)	Common Data Element	Format	Standardized response options	Importance	Comment / Description
Metabolic diseases Neurology	Development test: Results	normal/abnormal		Mandatory	Result of developmental test performed
Metabolic diseases	Developmental delay	yes/no		Mandatory	Developmental delay as assessed by treating physician
Nephrology	CKD stage	number		Mandatory	Chronic Kidney Disease stage
Pulmonology	Epworth Sleepiness Scale	number		Mandatory	
Pulmonology	Lung-to-Head-Ratio	number		Mandatory	Congenital diaphragmatic hernia
Pulmonology	PICADAR	number		Mandatory	PrImary CiliARy DyskinesiA Rule
Domain: Investigations					
General paediatrics	Type of radiological study (detailed)	standardized options	See comments	Mandatory	Standard classification to be defined
General paediatrics	Date and time of imaging study	datetime	YYYY-MM-DD hh:mm:ss	Mandatory	Date and time of the radiological study
General paediatrics	Radiation dose	number		Mandatory	If applicable, dose of radiation in mSv
General paediatrics	Indication for the imaging study	free text		Mandatory	Medical reason for the radiological study
Cardiology	ECG performed	yes/no		Mandatory	Date of study should be recorded
Cardiology	Holter-ECG	yes/no		Mandatory	Date of study should be recorded
Cardiology	Ergometry	yes/no		Mandatory	Date of study should be recorded
Cardiology	Echocardiography performed	yes/no		Mandatory	Detailed standardized echo measurements will be discussed in the future. Date of study should be recorded
Cardiology	Cardiac electrophysiology study performed	yes/no		Mandatory	Date of study should be recorded
Cardiology	Diagnostic cardiac catheterization (hemodynamic study) performed	yes/no		Mandatory	Date of study should be recorded

Module(s)	Common Data Element	Format	Standardized response options	Importance	Comment / Description
Endocrinology	Bone age: method	standardized options	Greulich & Pyle BoneXpertR Tanner Whitehouse	Mandatory	Method used to assess radiographic bone age. Date of study should be recorded
Endocrinology	Bone age: result	number		Mandatory	Bone age result in years
Endocrinology Metabolic diseases	Use of continuous glucose monitoring	yes/no		Recommende d	Use of glucose sensor
Endocrinology	Number of days per week with continuous glucose monitoring	number		Mandatory	Days per week
Endocrinology	Continuous glucose monitoring: Device	standardized options	Freestyle libre Freestyle libre 2 Dexcom G5 Dexcom G6 Medtronic Guardian Medtronic Enlyte	Mandatory	
Endocrinology	Blood glucose self-measurement	number		Mandatory	Number of measures per week
Endocrinology	Scans per day	number		Mandatory	If Flash Glucose Monitoring (FGM) is used
Endocrinology	Blood ketone measurement	number		Mandatory	Number of measures per week
Endocrinology	Mean glucose	number		Mandatory	mmol/l
Endocrinology	Glucose variability	number		Mandatory	%
Endocrinology	Time in range	number		Mandatory	Time between 4.0 and 10.0 mmol/l in %
Endocrinology	Time in hypoglycemia	number		Mandatory	Time < 3.9mmol/l in %
Gastroenterology	Type of gastrointestinal endoscopy	standardized options	Upper Lower Upper and lower Other	Mandatory	Date of study should be recorded
Gastroenterology	Indication for gastrointestinal endoscopy	standardized options	Rectal bleeding Abdominal pain Dysphagia Diarrhea Other	Mandatory	Medical reason for the endoscopic study. Other include for example oesophageal atresia or other anatomical abnormality, food impaction

Module(s)	Common Data Element	Format	Standardized response options	Importance	Comment / Description
Gastroenterology	Gastrointestinal endoscopic biopsy	yes/no		Mandatory	Gastrointestinal endoscopic biopsy performed. Date of study should be recorded
Gastroenterology	Impedance-pHmetry	yes/no		Mandatory	Date of study should be recorded
Gastroenterology	Type of breath test	standardized options	Lactose Lactulose Fructose Urea Other	Mandatory	Type of breath test. Date of study should be recorded
Gastroenterology	Capsule endoscopy	yes/no		Mandatory	Date of study should be recorded
Gastroenterology	Endoscopic ultrasound	yes/no		Mandatory	Date of study should be recorded
Gastroenterology	Liver biopsy	yes/no		Mandatory	Date of study should be recorded
Allergy/Immunology	Prick-test performed	yes/no		Mandatory	Date of study should be recorded
Allergy/Immunology Pulmonology	sigE performed	yes/no		Mandatory	sigE stands for specific serum immunoglobulin E. Date of study should be recorded
Allergy/Immunology	Result of sigE	positive/negative		Mandatory	sIgE stands for specific serum immunoglobulin E
Allergy/Immunology	Result of prick-test	positive/negative		Mandatory	Date of study should be recorded
Allergy/Immunology	Allergen challenge performed	yes/no		Mandatory	Date of study should be recorded
Allergy/Immunology	Allergen challenge result	positive/negative		Mandatory	
Infectious diseases	Urine collection method	standardized options	Urethral catheterization Clean catch void Urine collection bag Mid-stream urine Suprapubic aspiration	Mandatory	Method of collection of urine for culture. Date of study should be recorded
Infectious diseases	Mantoux test	number		Mandatory	In mm. Date of study should be recorded
Infectious diseases	Mantoux test: interpretation	standardized options	Positive Negative Doubtful Unknown	Mandatory	Healthcare provider's interpretation of Mantoux test

Module(s)	Common Data Element	Format	Standardized response options	Importance	Comment / Description
Infectious diseases	IGRA result	standardized options	Positive Negative Indeterminate	Mandatory	IGRA stands for Interferon-Gamma Release Assay. Date of study should be recorded
Nephrology	Renal ultrasound result	normal/abnormal		Mandatory	Date of study should be recorded
Nephrology	Renal MRI result	normal/abnormal		Mandatory	Date of study should be recorded
Nephrology	Voiding cystourethrography or kidney microbubble ultrasound results	standardized options	No vesicoureteral reflux Vesicoureteral reflux, unilateral – Grade I Vesicoureteral reflux, unilateral – Grade II Vesicoureteral reflux, unilateral – Grade III Vesicoureteral reflux, unilateral – Grade IV Vesicoureteral reflux, bilateral – Grade I Vesicoureteral reflux, bilateral – Grade II Vesicoureteral reflux, bilateral – Grade II Vesicoureteral reflux, bilateral – Grade III	Mandatory	Date of study should be recorded
Nephrology	Posterior urethral valves	yes/no			
Nephrology	Renal scintigraphy results	standardized options	Normal Hypoplasia Scars Other	Mandatory	Date of study should be recorded
Nephrology	Estimated GFR by Schwartz formula	number		Mandatory	GFR [ml/min]
Nephrology	Proteinuria	number		Mandatory	In mg/mmol (spot- urine) or mg/m2/h for 24h Urine
Nephrology Neurology Pulmonology	Genetic test performed	yes/no		Mandatory Recommende d	Mandatory for nephrology, recommended for neurology and pulmonology
Neurology	Neurologic electrophysiologic study: Type	standardized options	EEG EMG AEP SEP VEP Other	Mandatory	EEG: electroencephalogra m, EMG: electromyography, AEP: auditory evoked potentials, SEP: somatosensory evoked potentials, VEP: visual evoked potential Date of study should be recorded

Module(s)	Common Data Element	Format	Standardized response options	Importance	Comment / Description
Neurology	Neurologic electrophysiologic study: Result	normal/abnormal		Mandatory	
Metabolic diseases Neurology	Hearing test: Type	standardized options	OAE AEP Pure tone audiometry	Mandatory	OAE: otoacoustic emissions, AEP: auditory evoked potentials Date of study should be recorded
Metabolic diseases Neurology	Hearing test: Result	normal/abnormal		Mandatory	
Neurology	Vision test: Performed by	standardized options	Ophtalmologist Optometrist Peadiatrician Other	Recommende d	Health professional who tested vision
Neurology	Vision test: Result	normal/abnormal		Recommende d	
Neurology	Lumbar puncture performed	yes/no		Mandatory	Date of study should be recorded
Neurology	Opening Pressure at Lumbar Puncture	number		Optional	Opening pressure in cmH2O
Pulmonology	Spirometry performed	yes/no		Mandatory	Date of study should be recorded
Pulmonology	Lung function: RV	number		Recommende d	RV: Residual volume. In L
Pulmonology	DLCO	number	Diffusion capacity of the lung for carbon monoxide	Recommende d	DLCO: diffusing capacity of the lungs for carbon monoxide. In ml CO/min/mmHg
Pulmonology	Lung function: Bronchodilator administered	yes/no		Mandatory	
Pulmonology	Bronchoscopy performed	yes/no		Mandatory	Date of study should be recorded
Pulmonology	Lung function: Challenge test performed (treadmill, methacholine challenge test)	yes/no		Mandatory	
Pulmonology	Broncho-alveolar lavage performed	yes/no		Mandatory	
Pulmonology	Sweat test results	standardized options		Mandatory	Chloride in mmol/l (Macroduct) Conductivity in mmol/l eq NaCl (Nanoduct) Date of study should be recorded

Module(s)	Common Data Element	Format	Standardized response options	Importance	Comment / Description
Pulmonology	Sleep studies	standardized options	Polysomnography Respiratory Polygraphy Oximetry	Mandatory	Sleep studies performed Date of study should be recorded
Pulmonology	Lung function: FEV1	number		Mandatory	FEV1: forced expiratory volume- one second. pre/post absolute number in L
Pulmonology	Lung function: FVC	number		Mandatory	FVC: forced vital capacity. Pre/post. In L
Pulmonology	Lung function: TLC	number		Recommende d	TLC: Total lung capacity. In L
Pulmonology	Lung function: LCI	number		Recommende d	LCI: Lung clearance index. Equipment / gas currently in use in each center
Pulmonology	Lung function: Nasal NO	number		Recommende d	Nasal NO: Nasal nitric oxide measurement. in ppb or nl/mn
Pulmonology	Lung function: FeNO	number		Recommende d	FeNO: exhaled nitric oxide test. Online or Off-line method. absolute number in ppb
Pulmonology	Lung function: CPET performed	yes/no		Recommende d	CPET: Cardiopulmonary Exercise Testing
Pulmonology	Lung function: FEF 25-75	number		Recommende d	FEF25-75: Forced expiratory flow over the middle one half of the FVC (force vital capacity). in L/s
Pulmonology	Lung function: FEV 0.75	number		Recommende d	FEV 0.75: forced expiratory volume in 3/4 of a second. pre/post. absolute number in L
Pulmonology	Lung function: sRaw	number		Recommende d	sRaw: specific airway resistance. kPa/sec
Pulmonology	Lung function: FRC	number		Recommende d	FRC: functional residual capacity. in L
Pulmonology	Lung function: FRC: Test	standardized options	Bodyplethysmography MBW	Recommende d	MBW: multiple breath washout

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Module(s)	Common Data Element	Format	Standardized response options	Importance	Comment / Description
Domain: Diagnosis					
General paediatrics	Diagnosis	See comments	See comments	Mandatory	Inpatients diagnosis are ICD10 coded and outpatients diagnosis are free text.
General paediatrics	Date of diagnosis	date	YYYY-MM-DD	Mandatory	
General paediatrics	Cause of death	See comments	See comments	Mandatory	Standard classification to be defined
General paediatrics	Date of death	date	YYYY-MM-DD	Mandatory	
Cardiology	IPCCC diagnosis	standardized options	IPCCC Code	Mandatory	IPCCC: International Paediatric and Congenital Cardiac Code
Allergy/Immunology	Allergic disease confirmation	standardized options	Skin prick test Allergen challenge slgE	Mandatory	
Infectious diseases	If infectious diagnosis: Type of documentation	standardized options	Clinically documented infection Microbiologically documented infection	Mandatory	
Infectious diseases	If infectious diagnosis: Nosocomial	yes/no		Mandatory	
Infectious diseases	If nosocomial infection: Date of first symptom	date	YYYY-MM-DD	Mandatory	
Infectious diseases	If nosocomial infection: Site of infection	standardized options	Respiratory tract Gastro-intestinal tract Urinary tract Surgical site Other	Mandatory	
Metabolic diseases	Diagnosis confirmation	standardized options	Clinical Biochemical Enzymatic Genetic	Mandatory	The way diagnosis has been confirmed
Metabolic diseases	Diagnosis suspicion	standardized options	Prenatal Newborn Selective	Mandatory	The type of screening that led to the diagnosis
Neurology	OMIM code	standardized options	OMIM code	Recommende d	OMIM: Online Mendelian Inheritance in Man
Neurology	HPO code	standardized options	HPO code	Optional	HPO: Human Phenotype Ontology
Domain: Treatment					

Module(s)	Common Data Element	Format	Standardized response options	Importance	Comment / Description
General paediatrics	Drug name	standardized options	International non-proprietary name	Mandatory	Name of the drug(s) received as inpatient
General paediatrics	Prescribed drug at discharge	standardized options	International non-proprietary name	Mandatory	Name of the drug(s) prescribed at discharge
General paediatrics	Route of administration	standardized options	Oral Intravenous Subcutaneous Intramuscular Intrathecal Rectal Inhalation Cutaneous Ocular Nasal Otic Other	Mandatory	
General paediatrics	Date and time of first administration	datetime	YYYY-MM-DD hh:mm:ss	Mandatory	Time of first administration of the drug
General paediatrics	Date and time of last administration	datetime	YYYY-MM-DD hh:mm:ss	Mandatory	Time of last administration of the drug
General paediatrics	Frequency of administration	number		Mandatory	Number of administrations per 24 hours
General paediatrics	Dose	number		Mandatory	Dose given at each administration of the drug
General paediatrics	Dose unit	standardized options		Mandatory	List of possible units to be defined
General paediatrics	Reason for discontinuation of treatment	standardized options	Recovery Change to another medication No effect observable Adverse events Reducing polypharmacy Other	Mandatory	Reason why a treatment is stopped
General paediatrics	Adverse events	standardized options	MedDRA classification	Mandatory	MedDRA: Medical Dictionary for Regulatory Activities
General paediatrics	Supplemental O2: Date and time of start	datetime	YYYY-MM-DD hh:mm:ss	Mandatory	Time at starting oxygen therapy
General paediatrics	Supplemental O2: Date and time of interruption	datetime	YYYY-MM-DD hh:mm:ss	Mandatory	Time at stopping oxygen therapy

Module(s)	Common Data Element	Format	Standardized response options	Importance	Comment / Description
General paediatrics	Supportive services: Type	standardized options	Physiotherapy Ergotherapy Social assistance Other	Mandatory	
Endocrinology	Type of insuline therapy	standardized options	MDI CSII	Mandatory	MDI: Multiple dose injection. CSII: Continuous subcutaneous insulin infusion
Endocrinology	Total daily dose of insuline (long and short acting)	number		Mandatory	units per kg per day
Endocrinology	Basal insuline	number		Mandatory	Percentage of basal insuline (%)
Gastroenterology Metabolic diseases	Therapeutic diet	yes/no		Mandatory	Therapeutic diet prescribed by physician
Metabolic diseases	Type of therapeutic diet	standardized options	Low-protein Ketogenic Low-fat Frequent meals Nocturnal feed Medical food Other	Mandatory	Type of therapeutic diet prescribed
Allergy/Immunology	Epinephrine Pen prescribed	yes/no			
Infectious diseases	BCG immunization	standardized options	Yes No Unknown	Mandatory	
Neurology	Rehabilitation supportive devices: Type	standardized options	Upper limb orthoses Lower limb orthoses Corset Standing frame Walking aid (crutches NF-walker, rollator etc.) Wheelchair: Manual Wheelchair: Electric powered Other	Recommende d	
Pulmonology	Pulmonary rehabilitation	yes/no		Recommende d	
Domain: Equipment and	procedures	•			•
General paediatrics	Equipment type	standardized options	See comments	Mandatory	Standard classification to be defined
General paediatrics	Equipment date of insertion	date	YYYY-MM-DD	Mandatory	
General paediatrics	Equipment date of withdrawal	date	YYYY-MM-DD	Mandatory	
	•		•		

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Module(s)	Common Data Element	Format	Standardized response options	Importance	Comment / Description
Cardiology	Cardiac procedures	standardized options	IPCCC Code	Mandatory	IPCCC: International Paediatric and Congenital Cardiac Code
Cardiology	Date of cardiac procedure	date	YYYY-MM-DD	Mandatory	Date of intervention
Gastroenterology	Therapeutic gastrointestinal endoscopic procedures	standardized options	Haemostasis Oesophageal dilatation (Balloon/Savary) Percutaneous endoscopic gastrostomy (PEG) Endoscopic retrograde cholangiopancreatography (ERCP) Other	Mandatory	
Nephrology	Type of dialysis (1)	standardized options	Acute Chronic	Mandatory	
Nephrology	Type of dialysis (2)	standardized options	Haemodialysis Peritoneal dialysis Hemodiafiltration	Mandatory	If peritoneal dialysis, type of catheter and number of peritonitis should be specified
Nephrology	Date of dialysis initiation	datetime	YYYY-MM-DD hh:mm:ss	Mandatory	
Nephrology	Date of dialysis termination	datetime	YYYY-MM-DD hh:mm:ss	Mandatory	
Nephrology	Dialysis: vascular access type	standardized option	Central venous catheter Arteriovenous fistula Arteriovenous graft	Mandatory	If central venous catheter, its localization should be specified
Nephrology	Renal transplantation, graft (1)	standardized options	Deceased donor Living donor	Mandatory	
Nephrology	Renal transplantation, graft (2)	standardized options	Related donor Unrelated donor	Mandatory	
Nephrology	Renal transplantation	standardized options	Preemptive transplantation Nonpreemptive transplantation	Mandatory	
Nephrology	Renal transplantation: Number of received grafts	number		Mandatory	Number of grafts received including present one
Nephrology	Plasmapheresis performed	yes/no		Mandatory	
Nephrology	Renal biopsy performed	standardized options	No Yes, without complication in the following 24 hours Yes, with complications in the following 24 hours	Mandatory	
Nephrology	Cystoscopy performed	yes/no		Mandatory	
Nephrology	Angiography performed	yes/no		Mandatory	