MOSAIC – A Modular Approach to Data Management in Epidemiological Studies

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Summary
Introduction: In the context of an increasing number of multi-centric studies providing data from different sites and sources the necessity for central data management (CDM) becomes undeniable. This is exacerbated by a multiplicity of featured data types, formats and interfaces. In relation to methodological medical research the definition of central data management needs to be broadened beyond the simple storage and archiving of research data.

Objectives: This paper highlights typical requirements of CDM for cohort studies and registries and illustrates how orientation for CDM can be provided by addressing selected data management challenges.

Methods: Therefore in the first part of this paper a short review summarises technical, organisational and legal challenges for CDM in cohort studies and registries. A deduced set of typical requirements of CDM in epidemiological research follows.

Results: In the second part the MOSAIC project is introduced (a modular systematic approach to implement CDM). The modular nature of MOSAIC contributes to manage both technical and organisational challenges efficiently by providing practical tools. A short presentation of a first set of tools, aiming for selected CDM requirements in cohort studies and registries, comprises a template for comprehensive documentation of data protection measures, an interactive reference portal for gaining insights and sharing experiences, supplemented by modular software tools for generation and management of generic pseudonyms, for participant management and for sophisticated consent management.

Conclusions: Altogether, work within MOSAIC addresses existing challenges in epidemiological research in the context of CDM and facilitates the standardized collection of data with pre-programmed modules and provided document templates. The necessary effort for in-house programming is reduced, which accelerates the start of data collection.

1. Introduction

The collection and provision of medical data forms the basis for an analytical medical research. Therefore, cohort studies and registries mainly focus on quality-assured collection of primary research data and associated metadata as well as preservation of data interpretability.

When designing cohort studies and registries many research projects are confronted with considerable challenges (▶Figure 1). Considering the phases of the data lifecycle [1], this concerns organisational and technical effort necessary for the realization of a comprehensive data management including collection, processing, long-term storage and provision of data pursuant to recommendations of accredited institutions (e.g. the DFG [2]). One particular challenge is the integration of data from heterogeneous source systems. This includes laboratory information and management systems (LIMS), clinical information systems (CIS), diagnostic equipment and electronic case report forms (eCRF) usually generating a wide variety of data types and formats (e.g. form or image data, CSV exports, GDT files, XML structures, HL7 messages). Consequently, suitable measures must be adopted in order to ensure data quality throughout processing activities, to enable automated validation and to support interactive data correction. In addition, compliance with legislation for data protection must be assured at both federal and state level. For this reason, the ethics model must be built around a central consent management system administrating individual items of consent and authorisation (i.e. informed consent) and permitting revocation cross-checking on a daily basis. Management of...
participants is required in order to aggregate personal medical data within a central data repository while avoiding mistakes due to homonyms or synonyms. Furthermore, the ability to process and provide research data in a pseudonymised form is mandatory. Therefore, the introduction of a central data management (CDM) helps to simplify the process of design and implementation of data management for cohort studies and registries. This is accomplished by providing structured, uniform, and reproducible processes throughout all phases of the lifecycle of research data. Simultaneously, preconditions for long-term data usability and the comparability of derived results are constituted. More precisely, the use of CDM allows for the integration of all core issues for data management into cohort studies and registries as early as the planning stage [3]. In order to reduce effort, the deployment of reusable, modular solutions is encouraged. Thereby, the high effort arising from the repeated procedure of developing concepts and implementing dedicated software solutions for individual projects is avoided.

2. Objectives

Aim of this paper is to highlight typical requirements of CDM for cohort studies and registries, deduced from legal aspects, organisational task areas and technical approaches of data management typically applied within the phases of epidemiological projects. Based on cohort studies and registries, the paper illustrates how MOSAIC provides orientation for CDM by addressing selected data management challenges with a first set of ready-to-use and open source tools. MOSAIC is introduced to support the implementation of CDM for cohort studies and registries and to facilitate reducing in-house programming effort as well as promoting sustainability of existing solutions.

3. Methods

3.1 Central Data Management in Epidemiological Research

In cohort studies and registries many organisational and technical tasks must be completed prior to actual operation. Figure 2 depicts both organisational and technical measures and parameters that need to be considered during the preparation phase and the subsequent phases of preparation and usage. Starting with the specification of research questions and variables to be surveyed, the ethical framework (including determination of patient details and informed consent (e.g. paper-based or as digital document)) for the study must be established first (preparatory phase). Additionally, appropriate strategies for implementing authorisations and revocations, and the definition of workflows to support these activities are determined. In order to guarantee the consistent participant management and the necessary privacy an unambiguous identification of persons and the pseudonymisation of personal data must be ensured [4]. Subsequently, the necessary techniques for data collection, processing, archiving and provisioning must be defined. Furthermore, the technical environment necessary to ensure protection of data privacy and information security must be specified. Specifications must also be drawn up for separation and storage of personal data and medical data. This comprises a data model capable of mapping the collected information, a data repository for information storage and a role-based management system for access authorisation, to name but a few. Moreover, a data protection concept must be specified considering organisational, technical and personnel-related issues. This concept requires review and approval by the responsible data protection officer.

For the collection of research data, appropriate electronic case report forms (eCRFs) must be designed reverting to standard data collection tools including the data validation and possible data correction (automated and interactive). Amongst others, required interfaces, expected data formats, necessary metadata and data transfer protocols must be defined for data extraction from diagnostic equipment and other sources. In addition, measures to support record linkage, in order to minimize synonym and homonym errors during the data merging process [5], as well as to secure the data transfer between the study centres and the target storage system require specification.

During the subsequent acquisition phase, the focus is on ensuring high quality of the collected research data and on guaranteeing its security and long-term provisioning. Related tasks include the unobstructed operation of all systems required for error-free data collection as well as the definition and implementation of quality
assurance measures (e.g. instructing study personnel or source data checking and correction). At the same time, a high level of data integrity needs to be secured (i.e. by matching procedures for aggregation of personal data from multiple sources). Mechanisms for historisation and version control – aided by continuous monitoring and documentation of all data-handling processes – guarantee the necessary auditability of the data processing systems. In addition, backup strategies and rules regulating data access guarantee data security during this phase. To facilitate follow-up data collection, CDM also supports the recontacting of study participants.

During the usage phase the collected medical data is provided to researchers while complying with stated requirements for data privacy and protection. This comprises well-defined procedures for use and access and study-specific pseudonymisation. Furthermore, free specification of variable sets and export formats is provided to researchers. A preferably automated procedure is applied to determine and request the associated pseudonymised data, following prior verification of the necessary consent. Subsequently, the requested data is transmitted via a dedicated transfer unit. On completion of a study, a similar approach must be used to re-integrate the research results into the data pool. This includes derived variables, generated scores or variable coding work, for example. In the case of subsequent consent revocation or incidental findings subject to reporting obligations, the system must be capable of re-establishing the link from the data to individuals by a defined de-pseudonymisation procedure.

In summary, the study lifecycle requires a wide range of processes and measures that are not limited to separate phases within an individual study. On account of the considerable technical effort required to realize the necessary systems and functionalities, implementation should commence at the earliest possible juncture. However, the system architecture must exhibit sufficient versatility to satisfy supplementary requirements, changes and extensions in the study design.

Epidemiological studies and registries show that individual research projects exhibit particular requirements and settings [6–9]. Nevertheless, implementing a data management system as part of a study generates a recurrent set of similar issues. Such issues can be categorized to functional aspects, non-functional requirements and establishing compliance with legal frame-works. Fundamentally, CDM is targeted on satisfying each of these requirements.

Meyer et al. [10] define non-functional requirements for a CDM system. The authors are particularly focussing on data management processes. Reverting to this, their implementation is described by suitable technical measures with focus on quality assurance concentrating on the processed data.

In the context of epidemiological cohort studies these technical, organisational and staffing measures required to implement data management can be grouped into primary task areas. Combined with the non-functional requirements, they constitute the most important core elements of CDM. Details on the individual measures that each core element typically involves are provided in ▶Table 1.

Since the primary task of CDM is to aggregate data from heterogeneous and federated sources and to transform these data into suitable structures, the focus is on ensuring that the collection, processing, storage and provisioning of data is standardized and homogenized and coordinated from a single central point.

Therefore, the successful implementation of CDM in an epidemiological context is conditional on expert support (i.e. by com-

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**Organizational issues**

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<th>Technical measures</th>
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<td>Proband management</td>
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<td>Data formats</td>
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<td>Encryption</td>
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<td>Authorization management</td>
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| Data protection |
| Informed consent |
| Data types/sources |
| ID management |
| Security measures |
| Authorization management |
| ID management |
| Consent |
| Pseudonymization |
| Re-identification |
| Trusted Third Party |

| Acquisition |
| Examinations |
| Surveys |
| Analyses |

| Follow-up 1:N |
| (Communication of incidental findings) |

| Usage |
| Data processing |
| Data provisioning |

| Quality assurance |
| Versioning |
| Monitoring |
| Reporting |
| Controlling |

| Data sharing & Re-integration |
| Requesting process |
| Specific pseudonymization |
| Data transfer |

| Interfacing |
| ETL |
| Standards |
| Data mapping |

| Storage |
| Availability |
| Maintenance |
| Historization |
| Backup |
| Preservation |
| Technical monitoring |
| Data recovery |

**Figure 2** Typical phases of a cohort study: Organisational issues and technical measures concerning CDM regarding the phases of a cohort study
puter scientists and IT personnel) for the research. In addition, data management must be included in study planning at an early stage. Ideally, implementation of the technical and organisational requirements should be completed before data collection starts.

4. Results

4.1 A Modular Approach to Support the Implementation of CDM

In the context of the MOSAIC project (DFG program “Information infrastructure for research data”) a set of mutually independent tools is developed, each of them addressing a practical requirement of data management in cohort studies and registries (Figure 2).

Primarily targeting newly initiated cohort studies and registries, the MOSAIC web-portal (mosaic-greifswald.de) presents guidance and insights to legal, organisational and technical aspects of CDM. Scientists’ attention is drawn to relevant literature, existing solutions (provided and recommended e.g. by the TMF [12]) and current issues, assisting and encouraging them to specify the individual requirements for their study setting. Addressing these requirements and in accordance to the rules the TMF consented upon with the German protection officer [12], MOSAIC provides a portfolio of independent software modules, templates, checklists and recommendations in order to facilitate reuse of existing solutions. For already established cohort studies and registries, some MOSAIC tools might be of interest as well, e.g. if previously defined requirements have changed and new technical ways for participant management or pseudonymisation of existent medical data have to be identified.

The aim of MOSAIC is to provide support for the responsibilities of CDM through modular solutions (Figure 3). This approach allows for focusing on selected challenges instead of establishing complete solutions for a data management lacking adaptability, interoperability and flexibility.

A particular benefit of modularity is enhanced re-usability. The tools might be utilized separately in one epidemiological project to address a specific issue or used in combination in another project to satisfy a set of requirements. Nevertheless, MOSAIC does not intend to provide a complete software suite serving as a stand-alone solution for CDM. Each tool is or will be developed based on long-time experience in data management and in close cooperation with potential users. For this purpose, MOSAIC seeks to cooperate with newly initiated and already existing cohort studies and registries to acquire knowledge about specific needs and perceptions resulting from individual settings. The first set of tools, already available from the MOSAIC portal, is derived from solutions developed within existing projects [6–9]. Taking usability improvements for individual software tools as one example, the target was to reduce the number of configuration steps and to simplify the integration into existing infrastructures. Ease-of-use is improved by deploying web-based graphical user interfaces and by adding documentation (e.g. quick start guides, developer documentation and brochures).

To feature a high-level overview necessary for planning, designing and implementing CDM, a web-based reference platform [13] is offered. Depicting the necessary steps and typical issues occurring during the individual phases, the interactive reference platform aims to concentrate existing topical knowledge from the research community. For this purpose, based

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<th>Functional and non-functional elements of CDM</th>
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<td><strong>Core Element</strong></td>
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<td><strong>Functional requirements</strong></td>
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<tr>
<td>Acquisition</td>
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on structured questions and answers, users are encouraged to rate existing solutions and to share their issues and experiences. The reference platform is based on the open source question-and-answer framework Question2Answer [14] and is completely integrated into the MOSAIC webportal.

Setting up a new epidemiological project typically includes drafting a data protection concept. To provide a starting point and to prevent missing essential data protection issues, MOSAIC shares a document template (DOCX-Format) to simplify the process of writing. Structured questions and recommendations guide potential authors towards the integration of study-specific attributes into a pre-defined, pre-formatted concept paper. Alongside a conventional and directly deployable version of the document, an additional “generator” version enables interactive modification of the document template structure regarding mandatory, selectable and optional components beforehand. The document-generator utilizes several preformatted Microsoft Word building blocks, macros and configuration scripts optimized for the use with Microsoft Office 2010.

To avoid duplicate participant entries, the ID Management solution E-PIX (Enterprise Identifier Cross Referencing) applies the Fellegi-Sunter algorithm [15] and the Levenshtein distance. The independent software module allows for unambiguous participant management and efficient aggregation of medical research data from federated study centres. Additionally, the correction of potential synonym errors is supported (i.e. false-negative record linkage). The E-PIX, as well as the subsequent tools, applies a service-oriented architecture to provide all functionalities via web services. It was developed using Java EE and several development frameworks, including PrimeFaces [16] for a web-based user-interface.

Before research data can be collected or provided within a cohort study or registry, legal conditions require checking for available consents or revocations from the specific participant. To manage both digital and paper-based informed consents, MOSAIC offers the Consent Management...
The growing relevance of large-scale research networks in the scientific community makes the deployment of uniform methodologies and the generation of homogeneous data essential, since this allows for efficient data pooling and facilitates comparability. Consequently, work on the MOSAIC project complies with the recommendations of the German Research Foundation (DFG) for secure storage and provisioning of digital research data [2].

Previous work by Fraser et al. [24] has shown that the re-use of specially developed tools is capable of cutting costs and effort involved for data collection in separate studies. Moreover, utilizing the proposed set of MOSAIC tools increases the efficiency and standardisation of individual work within cohort studies and registries resulting from methodology streamlining. This standardisation facilitates an increased data quality [25].

The MOSAIC project’s approach to provide modular solutions for specific requirements addresses the need for practical tools, in terms of readily deployable templates and software, in order to support planning, design and implementation of CDM for cohort studies and registries. Especially in all cases where the scientific environment offers only marginal experience in data management, lacks access to IT personnel or suffers from insufficient resources in terms of software development.

Implementation of the respective tools demands precise knowledge of developments in the research community in order to avoid conducting primary development work in parallel to existing solutions. In case proven solutions existed for well-defined issues, the extent to which these solutions meet the requirements of epidemiological cohort studies and registries and the degree of applicability was assessed.

The TMF already provides a Guideline for Data Protection in Medical Research Projects (published in 2006, updated in 2014) [12]. Though it presents an introduction, guidance and recommendations to all aspects of data protection and ethical issues, it lacks an easy to use document template, which actually supports the respective author to write a data protection concept note. This aspect is well addressed by the MOSAIC template (published 2013), providing a structured starting point for essential typical data protection issues to be answered in the latter process. Using a question-based approach supports the application in various study or registry scenarios. Applying pre-worded text-blocks would narrow the scope of potential users, e.g. the data protection template of the Open source Registry System for Rare Diseases in the EU (published 2014 within OSSE, [26]) fits only for registries using the OSSE Software, but actually accelerates drafting a data protection concept note.

Several tools for the management of participants exist in the scientific community. For example the TMF PID-Generator (published 2005) [27] allows for merging participant identifying data from federated study sites even if the data sets are incomplete or faulty. The Java-based Mainzel List (published 2013) [28] aims for a more contemporary approach. It comes with a likely set of functionality, but with an easy to use REST-interface, which facilitates simplified system integration. However, unlike the E-PIX (published 2014), both systems are not yet capable of managing multiple local identifiers and identities for each participant and lack a graphical user interface to support the respective user in detecting and solving possible synonym errors.

Also for pseudonymisation and de-pseudonymisation a well-established tool exists. The TMF Pseudonymisation Service (PDS, published 2010, [29]) generates pseudonyms with a fixed length and alphabet (depending on the selected encryption method) using a synchronous algorithm. Thus the storage of associated value-pairs (original value and associated pseudonym) is not necessary. As a consequence the anonymisation of a medical dataset by simply deleting the association, which connects original value and pseudonym, is not possible. However gPAS (published 2013) [18] provides this mechanism and additionally facilitates the creation of pseudonym hierarchies. The generation of multiple pseudonyms combined with organisational and technical requirements for pseudonym administration (gPAS) [20] is also addressed.

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donyms for one participant allows a con-
text-related pseudonymisation e.g. to use
specific pseudonyms for different data
sources (eCRF, specimen, MRI) or for data
provision during the use and access pro-
cess.

Unlike the TMF Informed Consent
Wizard (published 2007) [30], gICS does
not provide textual support for drafting a
document. Neither gICS focusses clinical
practise nor it is limited to a simplified file-
based approach of HL7-documents like the
Consent Management Suite (COMS, pub-
lished 2011) [31]. gICS (published 2014)
focusses on the management of informed
consent documents using re-usable policies
to build modules, providing the necessary
service functionalities to support automatic
checks within seconds, e.g. whether a par-
ticipants allows to share his specimen or
has revoked his consent.

The developed MOSAIC tools will be
evaluated in cooperation with the users in
order to assess their efficacy and accept-
ance. One particular evaluation element is
a dynamic review from a functional per-
spective. This review is conducted by de-
ploying the tools to current and future ex-
ternal and internal projects enabling direct
tool modifications and enhancements. An-
other element is tool testing conducted by
external partners. The template for drafting
data protection concepts has already been
used in several research projects such as
the TORCH registry of the DZHK or the
MonDAFIS Study of the Centre for Stroke
Research Berlin at the Charité (CSB). The
modules for ID-Management, pseudonym
administration and informed consent admin-
istration have recently been used to set
up trusted third parties as a substantial part
of CDM in compound projects such as the
German Centre for Cardiovascular Re-
search [9] or the German National Cohort
[8]. In addition the MonDAFIS Study of
the CSB uses the ID-Management solution
E-PIX altogether with the open source
EDC-Solution REDCap [32].

6. Conclusions

Applying common phases of a cohort
study, this paper has deduced core ele-
ments of CDM for cohort studies and
registries in terms of functional and non-
functional requirements. Associated chal-
enges were highlighted, including require-
ments for IT security and legal data pro-
tection regulations.

In summary, CDM comprises the im-
plementation of extensive technical and or-
ganisational means for the collection, pro-
cessing, storage and provisioning of medi-
cal research data in accordance with the in-
dividual requirements specified by an epi-
demiological cohort study. Early incorpo-
ration of CDM features in study planning
permits successful implementation before
data collection commences. Furthermore
CDM facilitates a process of systematic im-
provements in terms of data quality and
availability while compliance with statu-
ory legal frameworks is assured.

This paper has introduced the MOSAIC
project as a potential resource to support
the implementation of CDM in epidemi-
ological research with a first set of open
source tools. Initial responses from the
scientific community document how the
concept of tool re-usability can be imple-
mented while preserving associated bene-
fits (cf. [24]). An important lesson learned
is that the simplest solutions (e.g. a docu-
ment template) awake great public interest.

Nonetheless, the MOSAIC tools will be
unable to cover each requirement for CDM
from an epidemiological perspective. Ini-
tially, no support for specimen manage-
ment or for the linkage, import and pro-
cessing of secondary data will be offered. In
addition, no provision of servers or facili-
ties for long-term data preservation is in-
tended at present. MOSAIC also does not
act as the commissioning instance for the
services of a Trusted Third Party. On the
other hand, technical support requests re-
lated to the tools as provided can be sub-
mitted via the project portal at any time.

For the remainder of its term, the MO-
SAIC project will focus on developing ad-
ditional tools, which will continuously be
made available to the research community
via the MOSAIC project portal (mosaic-
greifswald.de).

Acknowledgments

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