

**Bimonthly newsletter of XML4Pharma,
Schlossbergstrasse 20, DE-78224 Singen, Germany
Phone: +49 7731 975044
Web : www.XML4Pharma.com
Mail: Info@XML4Pharma.com
July-August 2008**



Finally, our newsletter is there again! Due to family reasons and a full agenda, I have not been able to send out the newsletter for the last months, but I now finally found the time again to write down the latest news and opinions.

First of all, there is some great news! Our ODMDesigner software has received CDISC ODM certification! Furthermore in this newsletter, you will find news on the European CDISC Interchange in Copenhagen in April, the forthcoming CDISC German CDISC User Group in Basel, and much more.

Furthermore, as some of you may know, I am concerned about some evolutions within CDISC, i.e. the development of HL7-XML messages for some of our standards. Especially worrying is the idea to develop an HL7-XML message for SDTM submissions. Therefore, I wrote an article “Ten good reasons why an HL7-XML message is not always the best solution as a format for a CDISC standard (and especially not for submission data)”. You can find a draft of it as an appendix to this newsletter.

XML4Pharma's ODMDesigner receives CDISC ODM Certification

We are proud to announce that our CDISC ODM Study Designer (ODMDesigner) has now received full ODM Certification from the CDISC organization. The ODMDesigner has been certified for all four use cases of metadata (i.e. study design). You can find the press release [here](#).

The ODMDesigner is not only the first study design software that can write and read Study Design files (others only write ODM), but also the only one that fully supports all vendors extensions according to the ODM Vendor Extension mechanism (ODM 1.2.1 and 1.3). As such, it is the ideal software tool for collaboration.



More information about the ODMDesigner can be found on [our website](#).

The full list of ODM-certified software offerings can be found on the [CDISC website](#).

CDISC European Interchange 2008

We were again present at the European CDISC Interchange in Copenhagen in April. On day 1 and 2, I contributed to the “end-to-end” workshop, where a good number of vendors demonstrated the use of CDISC standards from study design up to SDTM and ADaM submission to the FDA. In between, I also contributed to the SDTM course.

The conference itself was on Wednesday and Thursday. For the first time, there were parallel sessions. For me, that was a disadvantage – sometimes two interesting presentations were at the same time. Furthermore, I believe that having parallel sessions is not a good way to bring the message that all CDISC models interact and are being integrated.

During the main conference, I gave a presentation on “Bridging the gap between ODM and SDTM: Tools for creating SDTM Datasets from ODM Data”. During the presentation, the [SDTM-ETL software](#) was presented. The presentation can be downloaded [here](#). For those that could not attend, you can find the presentations on the [CDISC website](#).

We also had a booth at the exhibition, which was well visited, and allowing us to inform the participants about our services and software offerings.

All together, the European Interchange was again a great event, also with a great conference boating tour and dinner. Over 300 participants attended the meeting, which is once again a new record.

CDISC German-speaking User Group Meeting Basel, September 2, Basel

The German-speaking (well, they all do also speak English too) User Group Meeting will take place in Basel at the Roche facilities on September 2. A final program has not been released yet, so please keep an eye on the [bulletin board](#) of the User Group.

CDISC and HL7-XML messages

Those following the technological development within CDISC may have become confused by some messages that CDISC is considering HL7-XML messages from some of its standards and especially for SDTM submissions to the FDA.

I am regularly contacted by people from pharma companies, CROs and vendors with questions like “will we once again need to switch technology?”.

The idea for the development of HL7-XML messages is driven by the idea of integration with healthcare, and with some ideas within the FDA by people who do not have any understanding of XML.

As I strongly believe that HL7-XML is not the way to go (at least not for CDISC standards), I have written an article “Ten good reasons why an HL7-XML message is not always the best solution as a format for a CDISC standard (and especially not for submission data)”. You can find the second draft of this article (which I intend to publish in a scientific journal or on an external website), as an appendix to this newsletter.

CDISC and electronic signatures and certificates

CDISC has recently started working together with SAFE (Signatures and Authentication For Everyone – www.SAFE-BioPharma.org), a non-profit organization erected by the pharma industry. One of the goals of SAFE is to provide doctors and investigators in hospitals with token-based (e.g. USB-stick, smartcard) electronic certificates, which they can use for as well signing off electronic health records (or other documents), as for signing off electronic Case Report Forms (eCRFs). The token used for that can than also replace the multitude of username and passwords investigators currently need to use when doing clinical studies.

Our interest in SAFE is its use in EDC for signing off eCRFs. As many of you know, the CDISC ODM standard already allows to sign off a complete ODM file using XML-Signature and electronic certificates. My personal view is that this should also be possible with individual eCRFs in ODM-XML format. Technically seen, there is no problem to do this already. There are however some other problems: for example, the XML-Signature standard does not comply to EU legal requirements (e.g. a

datetimestamp is missing). An extension to XML-Signature (XAdES – XML Advance Electronic Signature) however does. Furthermore, investigators in hospitals do not usually already use electronic certificates, as the current procedures for obtaining one are currently complicated and costly. SAFE is therefore developing simple, but safe procedures to allow investigators to obtain an electronic certificate, e.g. via the sponsors. Essentially, the idea is that the electronic can be used independently on which clinical study the investigator is currently running, i.e. it should be universal.

For EDC systems, we do see that completed eCRFs are already sent by many EDC systems as XML to the server or database, or even as ODM-XML. These XML submissions can have an XML-Signature with or without an electronic certificate attached. Therefore, SAFE will have its impact on the further development of the ODM standard.

Personally, I volunteered to act as a liaison between the CDISC ODM team and SAFE. The reason is that in order to support XAdES and SAFE certificates, the ODM standard will need to be further developed. However, I strongly believe this is the way to go.

Initiative for an ODM extension for ePRO

Due to discussions with a number of ePRO vendors at the last CDISC Interchange, it became clear to me that the time is right for starting working on an ODM-extension for ePRO (electronic Patient Report Outcome, or “electronic diaries”), especially for the study design part.

ePRO typically has some features that are not covered yet by ODM, such as complex conditional navigation (though ODM 1.3 has “skip rules”), and timing rules (e.g. “display a specific form 1 hour after the subject has indicated to have pain”).

As I am working on such extensions for a few customers at this moment, why not try to come to a standard?

Therefore, I am currently trying to set up an “interest group” that may then become a subteam of the CDISC ODM working team.

If you or your company is interested in contributing to the development of such a standardized extension of ODM for ePRO, please let me know. I will then try to get this started after the summer holidays.

Ten good reasons why an HL7-XML message is not always the best solution as a format for a CDISC standard (and especially not for submission data)

Jozef Aerts, XML4Pharma
Version 0.2, July 2008

Introduction

CDISC standards have traditionally been based on XML technology, developed by teams of volunteers including XML experts¹. Examples include the CDISC Operational Data Model (ODM) standard, the Case Report Tabulation Data Definition Specification Standard (CRT-DDS), better known as define.xml, and the CDISC Lab standard, which has an ASCII implementation, a SAS implementation and an XML implementation.

Not very long ago, CDISC decided to develop the technical implementation of all new standards as extensions to the ODM standard. The most obvious reason for this is that the ODM standard is very successful in keeping as well metadata (such as study design) as clinical data². As submission data (standardized according to the Submission Data Tabulation Model (SDTM) **is** clinical data, and a standard format has already been established (as an ODM extension) to hold the submission metadata (define.xml standard), the most logical step was therefore to also develop an ODM-based format for holding the submission data itself. This format has been developed by a group of volunteers, consisting of as well submission specialists as highly skilled XML specialists, but has not been published by CDISC yet.

A current tendency however is being observed for wanting to format all new CDISC standards as HL7-XML messages. This tendency is driven by the desire for integration with the healthcare world, where HL7 is well-established, and by the FDA, who already uses a number of HL7-XML based standards, such as the annotated ECG standard (aECG). This idea is especially supported by people that have never been actively involved in XML development.

To my personal opinion, and to the opinion of many other XML specialists, this is a tendency that is at least concerning. This article will list and discuss ten good reasons why HL7-XML messages should in many cases not be considered as being the best format for new CDISC standards, and especially cannot be a good format for holding CDISC submission (SDTM) data.

1 Several CDISC volunteer teams have “XML-gurus” in their ranks, with considerable records of service in XML technologies
2 But also reference data and clinical trial administrative data.

Ten good reasons

1. An HL7-XML message is not compatible with define.xml

The wonderful thing about the ODM standard is that it works as a “framework”. It is perfectly suited to hold as well study metadata as clinical data, in such a way that the clinical data can easily be validated against the metadata. Also define.xml, the well-established (and by the FDA embraced) standard for SDTM submission metadata is based on this concept. Developed as an ODM extension, it holds the metadata of the SDTM submission. The submission data themselves still need to be delivered in the legacy SAS Transport 5 format. If the submission data itself however comes as an HL7-XML message (which has completely different concepts than define.xml), how will it be possible to validate the submission data against the metadata? Of course one can argue that it is always possible to compare apples to oranges, but it will considerably more difficult. Writing software to validate ODM clinical data against their metadata is pretty easy, especially when using modern (but easy-to-learn) XML-languages such as XPath and Schematron. With an HL7-XML message however, these technologies will make little chance, and other, much more complicated technologies will need to be used.

2. HL7-XML messages take years to develop

CDISC volunteers (under which a few XML-veterans) have already developed an ODM-based extension to hold SDTM submission data that easily validate against define.xml metadata. This model could in principle be put into operation within a year, thus replacing the legacy SAS Transport 5 format. The latter still stems from the seventies, and is a binary format based on IBM mainframe technology, also completely outdated.

The latter is a binary format from the seventies, based on IBM mainframe, so completely outdated.

It is well known that HL7 messages take years to develop and to get approved. The FDA itself has foreseen that, in case an HL7-XML message is being developed for transporting SDTM data, the SAS Transport format will at least remain in place until 2015 or even later³. With all the problems the outdated SAS Transport format delivers, this is a timescale the industry cannot afford.

3. Many HL7-XML messages are overcomplicated

Those who have ever inspected an HL7 aECG-XML file in detail, may have been surprised by the enormous complexity of the XML. One of the reasons for this is that the XML structure (as defined in its XML-Schema) is not developed by XML specialists, but is derived from UML diagrams.

I have been teaching a lot of XML in the last ten years and have experienced that CDISC ODM can be learned in a one day course. No chance however to accomplish this with aECG. Therefore, the amount of people that really understand aECG-XML is very limited, this in contradiction to the amount of people that understand ODM-XML.

Similarly, though XML is usually defined as being “as well machine-readable as human-readable”, one may question whether the latter is applicable to some HL7-XML messages such as aECG: not only the complexity is overwhelming, but it also uses a lot of coded, ununderstandable for the human reader.

In 2006, Gartner issued a Note entitled "HL7 V3 Messages Need a Critical Midcourse Correction", stating that "*HL7 must act vigorously to make Version 3 messages easier to use and more compact*" [see e.g. <http://2006.xmlconference.org/programme/presentations/141.html>].

The direct consequence of this overcomplexity is that it is much harder (and thus much more expensive) to develop software to read and write HL7-XML than it is for ODM-XML. From my personal experience (30 years) in software development, I estimate that the cost of developing software for a complex HL7-XML message is at least the twentyfold than it is for ODM-XML.

Fortunately, not all HL7-XML messages are overcomplicated. A good example is the XML implementation of the CDISC Lab standard. Although it has some minor design errors, it is pretty easy to handle and to understand. So, if it is really necessary to develop an HL7-XML message for SDTM, it should not be more complicated than the CDISC Lab-XML standard.

³ My personal estimate is 2020.

4. XML is not UML

HL7-XML messages are developed in a somewhat curious way: first of all one or more UML diagrams are developed, and then the XML-Schema is derived from the UML.

Though this may be the perfect and well-established way for translating a software design to software classes, it is considered bad practice by XML specialists. Transformation of UML to XML-Schema in general leads to “spaghetti XML”, introducing unnecessary complexity. Of course it is an “easy” way: the world has much more UML specialists than it has XML-Schema specialists. Personally, I would consider transformation of UML to XML-Schema the “lazy man's way”. The result can however be catastrophic.

By the way, none of the most popular XML-based standards, such as MathML, VoiceXML, XHTML etc. have ever been developed using UML.

5. HL7-XML is nearly used in the industry

HL7 messages are pretty successful in the healthcare industry, especially in Northern America. However, when one looks in more detail, it is observed that over 99% of the implementations are HL7 version 2 implementations (not using XML), and that less than 1% are version 3 implementations (using XML). So the “market share” of HL7-XML within HL7 is less than 1%.

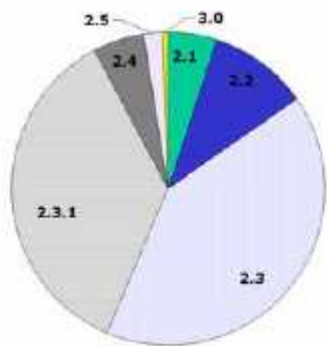


Figure 1: Approximate real-world usage of HL7 messaging standards. The vast majority of HL7 messaging is done using messages that approximate HL7 2.3 or HL7 2.3.1. Newer releases of HL7 (2.5, 3.0, and soon 2.6) represent a very small portion of real-world interfaces.

[REFERENCE: <http://www.neotool.com/pdf/HL7-Version-3-with-HL7-Version-2-History.pdf>].

Reasons mentioned for this relative non-success of HL7 version 3 are the enormous cost and lacking backward compatibility with version 2. Software developers and system architects that have successfully implemented one the version 2 HL7 standards, have great difficulty with understanding and implementing version 3 HL7-XML [REFERENCE].

The “market share” of HL7-XML might further decrease, as many new HL7 messages are developed and further maintained as HL7 v.2 messages (non-XML).

6. In many cases HL7-XML ignores existing XML standards

When developing the CDISC ODM standard and its extensions, the CDISC ODM team has always looked at other XML-based standards, and has always tried to prevent reinventing the wheel. An example is the implementation of the XML-Signature standard within the ODM, but also the fact that all ODM data types are based on native XML-Schema data types. For example, an ODM “date” is based on the native XML-Schema datatype “xs:date”, which is based on a subset of the ISO-8601 standard. Similar for time, datetime, incomplete-datetime, partial-datetime, and durations. In too many HL7-XML messages however (there are exceptions), all these are defined as being of data type “text”, making it very hard to validate instance data against the XML-Schema.

For example, whereas a date “2008-02-30” will immediately fail when validating against the ODM XML-Schema (without writing any special software), a date “20080230” (observed as date format in many HL7 XML-messages) is much more harder to detect as being an invalid date (special software is required).

ODM however has even defined some other ISO-8601 based XML data types, such as incompleteDataTime and partialDateTime, which can be of extreme importance for SDTM submissions in XML. In HL7-XML, these are based on string patterns, with the same limitations in validation.

In some HL7-XML messages we have even observed that two different ways of representing dates are being used in the same file – so the implementation is even not consequent.

```
- <AnnotatedECG xmlns="urn:hl7-org:v3" xmlns:voc="urn:hl7-org:v3/voc" xmlns:xsi="http://
  xsi:schemaLocation="urn:hl7-org:v3 /HL7/aECG/2003-12/schema/PORT_MT020001.
  <id root="ADB8E58B-1B4F-4F42-B9CE-AD3DDF55E57F" />
  <code code="93000" codeSystem="2.16.840.1.113883.6.12" codeSystemName="CPT-4"
  <text />
- <effectiveTime>
  <low value="20040115102010.000" inclusive="true" />
  <high value="20040115102020.000" inclusive="false" />
</effectiveTime>
```

Incorrect formatting of date-time
extensive use of codes
human-readable ?

This ignorance of existing XML standards may be a consequence of the fact that HL7-XML schemas are “derived” from UML, rather than developed from the ground up by XML-Schema specialists.

7. Why make it complex when it can be simple ?

There is no need for having SDTM submission data in a complex data structure. The SDTM standard describes all submission data as simple two-dimensional tables, not as complex objects. So a relative simple ODM-based data format (similar to define.xml and ODM-ClinicalData) is more than sufficient as a vehicle for porting SDTM data to the FDA. Arguments for an HL7-XML message for SDTM are based on the desire for integration with the healthcare world, where patient data are usually exchanged using HL7 messages. SDTM data are however much more simple, and can easily be transformed (if necessary at all) from complicated HL7 messages with patient data to the much more simple ODM format using XSLT stylesheets.

SDTM does not have the concept of “actors”, “roles”, ... SDTM data are simple two-dimensional tables with some table and column attributes (the metadata in the define.xml). So designing an HL7-XML message for transporting is a complete overkill.

8. Complex data structures can be transformed to simple datastructures by XSLT, the reverse is much more difficult

Operational clinical data are usually exchanged using the CDISC ODM, XML-based standard. This standard is now in its version 1.3, and is continuously improved by an enthusiastic group of volunteers, containing as well clinical data specialists, as XML specialists. This standard is embraced by the FDA, as it is fully CFR Part 11 compliant, including audit trails and signatures. The ODM structure is however, compared to the HL7-XML structure, uncomplicated and not complex at all.

Several vendors⁴ have already developed ODM to SDTM mapping tools, and have implemented an XML-based SDTM format, based on the ODM. As such, transforming operational clinical data (ODM) to such (ODM-based) SDTM-XML data is uncomplicated and straightforward. These vendor tools are even able to generate the XSLT transformation code to execute the transformation in an automated way!

XSLT transformation from a non-complex format to another non-complex format is easy. XSLT transformation from a complex format (such as HL7-XML) to an uncomplicated XML format is also pretty easy.

⁴ Including XML4Pharma, XClinical and Formedix

Transformation from a non-complex format (such as the ODM) to a complex format (HL7-XML) however is very complicated. Automated generation of XSLT code to transform operational clinical data (ODM) to an complex HL7-XML message is probably a programmer's nightmare, meaning that mapping software will be very complex and very expensive.

9. CDISC standards are owned by CDISC, HL7 messages by HL7

The SDTM standard is a development by CDISC, not by the FDA nor by HL7. Though CDISC and HL7 have an “Associate Charter Agreement” (meaning a strong relationship), there may be ownership issues when an HL7 message is developed for carrying SDTM data. Who would be the owner of such a format? Would it be CDISC, who clearly is the owner of SDTM, or would it be HL7? Probably the latter. Or would it even be the FDA? This would lead to the strange situation that the owner of the content standard and the owner of its transport format are not the same⁵.

10. There is little or no XML knowledge at the FDA

The desire to have an HL7-XML message for submission (SDTM) data clearly comes from the FDA [http://www.fda.gov/CDER/REGULATORY/ersr/2003_06_17_XML/sld001.htm]. The amount of knowledge of XML technology (and especially XML-Schema technology) at the FDA is however very limited. Therefore we must suppose that the choice for an HL7 message is a political one rather than a technical or pragmatic one.

This means that the FDA will essentially obtain a “black box” from HL7, with the danger that they will need to rely heavily on external parties for the development and deployment of software tools for working with the SDTM HL7-XML message⁶. For example, for viewing SDTM data in a tabular way, they will need extremely complicated stylesheets, or even very special software.

With ODM-based technology however, this danger is much smaller. For example, the CDISC define.xml team has developed several uncomplicated stylesheets to view the SDTM metadata in a standard browser. These stylesheets are being used by the FDA, but have also been refined by many sponsors. Similarly, several vendors have developed viewers for inspecting ODM clinical data, based on stylesheets. If the SDTM data themselves come as an ODM-extension, the same technology can be used to visualize SDTM datasets, and the wheel does not have to be reinvented or redeveloped. Additionally, the metadata and data can be visualized together (define.xml and SDTM-ODM-XML) as they are based on the same base standard. Establishing the same for HL7-XML based SDTM datasets is probably an illusion.

What is then the value of HL7-XML messages?

There is certainly a great value in HL7-XML messages. HL7 messages try, with a good amount of success, to capture the whole healthcare information and data streams. The healthcare world is a complex world, one of the reasons why HL7 messages are so complex. Very probably, electronic health records (EHRs) can never be described using a non-complex format like the one used in the ODM standard. For this, an HL7-XML message may be ideal. Assuming however that an HL7-XML message is therefore also necessary for operational or submission clinical data is an error. Operational (ODM) and submission (SDTM) data in XML format can better be transformed from EHRs using a complex-to-simple transformation (probably using XSLT), rather than using a complex-to-complex transformation⁷.

Also HL7 is very important to CDISC for semantic interoperability. If information from EHRs need to be transformable into clinical data, it is very important the EHR and CDISC standards “speak the same language”. This however does not mean that they need to use the same format.

⁵ This is not the case for the CDISC Lab standard which was developed by CDISC as an HL7 message. As it was developed by CDISC, the owner is CDISC.

⁶ This was exactly the case with the aECG HL7-XML message. It was developed by B.D.Brown from Mortara, and F.Badilini from AMPS llc. The input of the FDA was limited to writing the user requirements.

⁷ I doubt whether EHR patient data will ever have to be directly transformed into SDTM data. Transformation to operational data (e.g.ODM) is straightforward, but transformation to SDTM data always involves an interpretation step.