Seven Habits of Highly Effective Issue Managers
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ABSTRACT

Pinnacle 21 Validator identifies problems in data; however, diagnostics, assessment and resolution of reported validation issues may feel like a complicated, never-ending process. In this presentation, we will discuss common challenges in managing data validation issues and how to handle them effectively. We will show you how to identify the source of validation issues, and how to classify them to understand when to fix or when to explain. We will also discuss cross-team collaboration, ways to improve your process, and habits that lead to faster issue resolution.

INTRODUCTION

All issues for a CDISC deliverable are not created equal: they come from different sources, have different impacts, and require different means of resolution. Programmers rely on Pinnacle 21 Validator to identify problems with data but resolving issues is often difficult - especially when there are several stakeholders to satisfy including your manager, the study statistician, the sponsor, and ultimately FDA and PMDA reviewers. An issue manager is usually a programmer, but it could be anyone at your organization; this paper talks about how you can be an effective issue manager that satisfies stakeholders by ensuring issues are resolved quickly and correctly.

Briefly, the seven habits are:

1. Validates early
2. Gathers all relevant info about issues
3. Identifies the source(s) of issues
4. Tracks changes between validation reports
5. Communicates issues to others
6. Knows when to fix validation issues
7. Knows when to explain validation issues

HABIT ONE: VALIDATES EARLY

Resolving validation issues is part of the CDISC process for ongoing studies. CDISC data are maintained many years in long clinical trials and there is a growing demand to have SDTM sooner than ever before, often leading programmers to have specifications and programs in place by the time a database goes live. When data conversion starts early, so should validation. Early validation can actually save time by acting as a first-pass QC to find and fix programming and specification-related issues. Using validation as a first-pass QC method will cut down on the amount of back-and-forth between primary and validation programmers later.

HABIT TWO: GATHER ALL RELEVANT INFO ABOUT ISSUES

After you run a validation, it’s important to be able to look-up erred records in more detail, sometimes even tracing data all the way back to the raw EDC data. One reason to do this is so that you can accurately identify the source of each issue. Another reason is because you need to communicate with data management about some issues and data managers often need more information about the record than what is provided on the Details tab of a Pinnacle 21 Validation Report. One way to quickly get more information about erred records is to use a SAS® macro like the one presented in The Devil is in the Details – Reporting from Pinnacle 21 (OpenCDISC) Validation Report (Garrett and Whalen, 2015). This macro creates a report of each issue on a separate excel tab and provides complete
information about each erred record. Alternatively, Pinnacle 21 Enterprise provides full details about erred records directly within the system (see Figure 1).

**Figure 1.** Example of record details from Pinnacle 21 Enterprise.

**HABIT THREE: IDENTIFIES THE SOURCE(S) OF ISSUES**

In order to understand where quality problems originate, it’s important to identify the source of each issue and categorize it. Categories can also be used as a way to determine which issues need to be delegated to other people for resolution (for example, data collection issues should be assigned to data management). One way this is achieved is by assigning a primary source to each issue. In Pinnacle 21 Enterprise, users can make use of tags to categorize each issue by its source; the tag(s) assigned will be automatically exported to the validation report. Figure 2 is an example of tagging issues in Pinnacle 21 Enterprise. Alternatively, you can directly type the issue source into Excel validation report. Most issues will fall into one or more of the following categories:

- data collection error
- programming/spec error
- metadata (define.xml)
- sponsor-defined addition
- study is ongoing
- false positive

**Figure 2.** Example of Pinnacle 21 Enterprise issue table with tags applied to denote the issue source.

**HABIT FOUR: TRACKS CHANGES BETWEEN REPORTSEARLY**

Validation is an ongoing process and it’s important to track the delta from one validation report to the next so that you can determine when new issues are occurring and how quickly known issues are being
resolved. There are three types of issues to track between validations: new issues, resolved issues, and issues that are on both reports (possibly affecting a different number of records). When the same issue is present in both reports, prior comments should be copied from one report to the next, as applicable. One way to track the delta between validation reports is to use a program (SAS macro or other application) to compare the two reports together and produce a consolidated report with comments carried forward. Another option is Pinnacle 21 Enterprise, which automatically copies issue comments from one validation report to the next and has a feature that allows users to easily compare any two versions of a validation report (see Figure 3).

Figure 3. Example validation report comparison report filtered to only show new issues.

HABIT FIVE: COMMUNICATES ISSUES TO OTHERS

The Pinnacle 21 Validation Report can serve as a communication device and be distributed to peers, managers and other stakeholders. Since all stakeholders may not be well-versed in CDISC, you should mark-up the validation report for better consumption. You should clearly state the source of an issue, who is responsible for resolving it, and provide any extra information that may be needed about an issue. Pinnacle 21 Enterprise allows users to assign issues to users, tag issues with their source, and provide comments about an issue, all of which are automatically exported to the Validation Report as depicted in Figure 4.

Figure 4. Sample Pinnacle 21 Enterprise Validation Report with comments, assignee, tags, and explanations pre-populated based on information provided in the Enterprise system.

HABIT SIX: KNOWS WHEN TO FIX VALIDATION ISSUES

An effective issue manager is one that knows when and how to fix each issue. You should understand the risk of each issue and prioritize fixing issues that have the highest impact on regulatory review. The FDA is no longer publishing severity, but it can still be used as a proxy for impact level; Errors and Reject rules usually have the highest impact on review. Pinnacle 21 Enterprise provides review impact for each issue,
as shown in Figure 5, and recommends that you fix issues with high review impact first. In general, data collection errors, programming/spec errors, and issues related to the define.xml (issue with DD prefix) should always be fixed while issues due to an ongoing study should resolve naturally over time.

Figure 5. Sample issue summary table with Review Impact highlighted.

Knowing how to fix each issue is challenging for any one individual, leading some companies to maintain a document with instruction guidance for how to fix the most common issues. This kind of document is especially helpful to more junior members of the team. An example of one such document can be found in the paper entitled Common Pinnacle 21 Report Issues: Shall we Document or Fix (Gupta, 2018). Pinnacle 21 Enterprise provides fix-tips for issues as well and can be customized for an organization (see Figure 6.)

Figure 6. Sample issue detail with Pinnacle 21 Fix tips provided.

HABIT SEVEN: KNOWS WHEN TO EXPLAIN VALIDATION ISSUES

No study is perfect and there will always be a subset of pesky validation issues just won’t go away. Any issue that cannot be fixed, even false positives, should be explained in the Reviewer’s Guide. Best
Practice for Explaining Validation Results in the Study Data describes a good explanation as “one that conveys transparency about the study data and increases the reviewability” (Kelly, 2018).

It’s a best practice to explain issues consistently across an organization. To keep explanations consistent, some issue managers keep a list of standard explanations with bracket placeholders for study specific information and then copy these explanations into the Reviewer’s Guide either manually or with a merge process. Another option is to use Pinnacle 21 Enterprise which allows users to upload standard explanations, apply them to issues and modify them as necessary. These explanations will automatically be exported to the Reviewer’s Guide. Figure 7 shows an organization’s standard explanation (right) and how it was applied to this study (left). Figure 8 shows how the explanation would appear in the Reviewer’s Guide.

Figure 7. Example of Pinnacle 21 Enterprise Suggested Explanation and how a standard explanation can be customized with study-specific information.

<table>
<thead>
<tr>
<th>Check ID</th>
<th>Diagnostic Message</th>
<th>FDA Severity</th>
<th>Dataset</th>
<th>Count (Issue Rate)</th>
<th>Explanation</th>
</tr>
</thead>
<tbody>
<tr>
<td>SD0011</td>
<td>ARM is not 'Screen Failure', when ARMCD equals 'SCRNF' or vice versa</td>
<td>Error</td>
<td>DM</td>
<td>25 (100.00%)</td>
<td>&lt;Explanation&gt;</td>
</tr>
<tr>
<td>SD0012</td>
<td>AESDLY is after AEENDY</td>
<td>Error</td>
<td>AE</td>
<td>5 (0.28%)</td>
<td>&lt;Explanation&gt;</td>
</tr>
<tr>
<td>SD0013</td>
<td>AESDTC is after AEENDTC</td>
<td>Error</td>
<td>AE</td>
<td>1 (&lt; 0.1%)</td>
<td>&lt;Explanation&gt;</td>
</tr>
<tr>
<td>SD0015</td>
<td>Negative value for SUDUR</td>
<td>Error</td>
<td>SU</td>
<td>3997 (96.76%)</td>
<td>&lt;Explanation&gt;</td>
</tr>
<tr>
<td>SD0017</td>
<td>Invalid value for RPTEST variable</td>
<td>Error</td>
<td>RP</td>
<td>53 (25.24%)</td>
<td>&lt;Explanation&gt;</td>
</tr>
<tr>
<td>SD0019</td>
<td>Invalid value for TSPARM variable</td>
<td>Error</td>
<td>TS</td>
<td>1 (3.70%)</td>
<td>&lt;Explanation&gt;</td>
</tr>
<tr>
<td>SD0021</td>
<td>Missing End Time-Point value</td>
<td>Warning</td>
<td>AE</td>
<td>576 (24.23%)</td>
<td>Subject XYZ-0001 did not have end dates or references collected and so could not populated in SDTM</td>
</tr>
</tbody>
</table>

Figure 8. Example of a Reviewer’s Guide generated with Pinnacle 21 Enterprise.
CONCLUSION

Validation is a process and the individuals tasked with creating and interpreting the validation report need to be organized and use the tools at their disposal to ensure an efficient end-to-end process. SAS macros can be utilized to make some process tasks easier; users also have technologies like JIRA to create and manage issues. Pinnacle 21 Enterprise offers several features specifically designed for issue management such as standard issue explanations and the ability to assign a validation issue to a specific individual. Regardless of the technology that is used, a successful validation issue manager knows the importance of validating early, tracking the validation results, and communicating findings to the right individuals.

REFERENCES


Kristin Kelly. Best Practice for Explaining Validation Results in the Study Data. Proceedings of the PharmaSUG 2018 Conference.

RECOMMENDED READING


CONTACT INFORMATION

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