1.1 The Need for Guidance on ERP System Validation

There are numerous books that address the topic of computer systems validation in the regulated life sciences industry covering the general topic of computer systems validation. Based upon feedback from industry and the number of questions typically raised when validating (or planning to validate) an ERP system it has become clear that there is a need to address the specific topic of validating ERP and other large configurable systems.

While the life sciences industry has generally done a good job in meeting the regulatory requirement to validate GxP critical systems this is not always the case with large, complex, configurable systems.

There are many reasons for this, and Successfully Validating ERP Systems (and other large configurable applications) is intended to help readers understand:

- The reasons why the validation of such systems often takes longer than expected
- Why the implementation often costs more than originally planned
- Why, in some cases, the validation fails to meet regulatory expectations.

Successfully Validating ERP Systems (and other large configurable applications) has been written specifically to address this topic and provide pragmatic guidance based upon real industry experience. Fundamentally it sets out to answer the question ‘Why is validating ERP systems different, and what is needed to successfully validate such a system?’

In summary, these differences come down to addressing:

- How to position rapid applications development (RAD) in the validation life cycle, including the role of the conference room pilot (CRP)
• The need to integrate the configuration of configurable off-the-shelf software and the
development of custom software within the same project
• How to define appropriate verification activities for all of the multiple, complex and
simultaneous tasks which take place during an ERP implementation project
• The fact that the validation of the software is only part of validating the ERP solution
• How to plan and manage all of these validation activities in the context of a large,
complex, fast moving and often politically charged project environment.

The intended audience for this book is diverse — as indeed is the makeup of a typical ERP
project. While anyone engaged in implementing ERP systems in the regulated life sciences
industries will find sections of the book useful, its primary audience is:

• Those responsible for the validation of ERP systems such as validation workstream
managers and quality or regulatory affairs departments (depending upon the User
organisation)
• Those supporting the validation of an ERP system (members of the validation workstream)
• Those responsible for managing the implementation of ERP systems, including members
of the program steering committee, the program manager, members of the program
management office and workstream managers (see Program Structure, Chapter 2).

The book is not intended to be an introduction to the basics of computer systems validation,
nor a simple ‘how-to’ tutorial. Some knowledge on the topic of computer systems validation is
assumed, but where useful, references are given for further reading.

Unlike many other books on the subject of computer system validation the intent is not to
provide ideal world models that cannot be applied in practice. This book sets out with the
specific intention of recognising and discussing the real world human and contractual issues
that lead to validation issues.

These include:

• The complex interaction of financial constraints
• The fundamentally opposing objectives of many of the parties involved with
implementing such systems
• The unavoidable issue of program politics, problems with training and education
• The basic problem of applying inappropriate validation models to complex program
implementations.

In some User organisations these issues start with the fundamental question of ‘Why do we
need to validate our ERP system?’
1.2 The Need to Validate ERP Systems

Implementing any large, complex system such as an Enterprise Resource Planning (ERP) or other large configurable system always brings its own challenges. The common requirement for the validation of such systems in the regulated life sciences industry (i.e. pharmaceuticals, medical devices, biotechnology etc) brings additional challenges which many individuals or organisations struggle to meet. Note that this book is written specifically in the context of the life sciences industry and although the principles can be broadly applied this does not include the expectations of other regulated industries (i.e. financial services, nuclear engineering etc).

While an ERP system may not be considered as a ‘production system’ by many people in the business, the requirement to validate it is driven by the fact that the ERP system implements or supports production processes that are within the scope of the various regulations. Functions such as goods inwards inspection, quality inspections and product release are included within such regulations, and take the ERP system into the production area. The need to validate systems supporting such processes is clearly defined in regulations and standards:

- US 21CFR Part 211.68
- US 21CFR Part 820.70
- US 21CFR Part 11.10
- Annex 11 of the EU and PIC/S Good Manufacturing Practices
- ISO 13485 part 7.5.2.1.

Additionally, given the broad functional scope of many ERP systems other business processes such as the management of customer complaints, corrective and preventative actions, and product design will also take the ERP system into areas covered by these (or other) regulations, also requiring the system to be validated.

This is evidenced by the fact that most, if not all, regulatory agencies will include the ERP system within the scope of regulatory inspections and will treat a failure to validate the system as a major non-conformance likely to be included as an inspection observation.

The process of validating any computerised system can be summarised as demonstrating fitness for purpose. This can largely be achieved by user acceptance testing. However, the need to clearly define requirements and produce other objective evidence of the verification activities (as recommended by regulatory guidance on the topic of computer systems validation) also requires careful thought on activities and resulting documentation for ERP implementation projects.

In the context of this book the process of validation includes:

- All activities defined by an appropriate implementation life cycle (including their verification )
- The production and control of appropriate objective evidence of such implementation and verification activities
• Testing to demonstrate fitness for purpose
• The production of other program documentation required to demonstrate an appropriate
degree of control with respect to key activities defined in regulatory guidance.

Where useful, other general industry guidance has been referenced throughout. This includes
many references to GAMP®, (a technical subcommittee of ISPE), which provides a range of
practical guidance on the topic of computer systems validation, and which is referenced by
various regulatory agencies in their own guidance documents.

GAMP® is the most widely referenced industry guidance, and forms the basis for
computer systems validation training for many regulatory inspectors from various agencies. It
is therefore useful to be able to explain the process of validating an ERP system in terms that
are generally understood by the regulators.

It should, however, be stressed that any defined implementation and validation life cycle is
acceptable to regulatory agencies, and in some cases strict adherence to the GAMP® validation
model will cause issues during the implementation of an ERP system. Specific attention is
therefore given to describing the ERP validation process in terms more familiar to regulatory
agencies (i.e. in the context of a GAMP® like validation model). Other GAMP® good practice
guides applicable to many different types of system are also referenced as appropriate.

Reference to GAMP® may perhaps be less useful, depending upon the specific industry
sector or User organisation. (For instance, in the medical devices sector, or where more specific
regulatory guidance on computer systems validation is available.) When defining the validation
process, User organisations are also recommended to refer back to the specific computer
systems validation guidance published by applicable regulatory authorities (see Bibliography)
and ensure that their implementation and validation approach meets the fundamental
requirements of such guidance and delivers the appropriate objective evidence.

1.3 The ERP Implementation Phenomenon

During the 1990s a number of large organisations, both in the life sciences sectors and other
industries, implemented ERP systems. There were a number of reasons for this:

• The move towards business consolidation and globalisation
• The need to drive business improvements
• The need to become ‘Y2K’ compliant
• The need to address 21CFR Part 11 (Electronic Records, Electronic Signatures)
• In the European Union (and other countries), the need to support the new Euro currency
  (many older systems were not capable of this)
• In some cases, the need to address regulatory non-compliances, including a failure to
  validate previous systems.
Many of these initial implementations gained a reputation for cost overruns, late delivery and failure to provide the promised functionality and business process improvements.

Suppliers and system integrators (SIs) responded by developing techniques to implement ERP systems in a faster and less expensive manner that also leveraged ‘industry standard’ solutions, incorporating preconfigured processes. Typically these included some sort of RAD approach to implementation (discussed in further detail in Chapters 4 and 6).

As the IT market slowed down at the start of the 21st century, competitive pressures also meant that SIs felt the need to reduce costs and implementation timescales even further, and to develop their own implementation methodologies. In many cases this was a cutdown version of the Suppliers standard RAD approach.

Such ‘plain vanilla’ or ‘slam-dunk’ implementation approaches often require the client to change their business processes to adopt the industry standard contained in the preconfigured solution.

In many cases these approaches were successful in delivering lower costs and faster implementations. However, in the life sciences industry such an implementation approach led to a number of common problems that underlie many of the current issues associated with successfully validating ERP systems. These are:

- An initial failure to plan for successful validation
- Failure to capture and document the specific requirements of the individual User
- A tendency to implement ‘standard’ business processes that were non-compliant with the User specific interpretation of the applicable regulations
- Failure to adequately address electronic records and electronic signature requirements
- General failure to provide adequate program documentation
- Failure to sufficiently test the solution
- Insufficient user training and a failure to update associated standard operating procedures (SOPs)
- Failure to qualify the associated IT infrastructure
- A failure to sufficiently consider maintaining the system in a compliant manner throughout its operational life.

Many of these problems were exacerbated by Suppliers and SIs who, although experts in their specific field, were usually generalists, and often did not fully understand the validation requirements in the life sciences industry.

As the pace of ERP implementation has slowed its associated problems can now be considered with more perspective. Each of these are discussed in more detail in the following chapters.
1.4 Why Enterprise Resource Planning Systems?

ERP systems are not the only type of large configurable systems implemented across life sciences organisations. While this book specifically addresses issues associated with ERP systems, the principles outlined can be applied to any large configurable system. This is pointed out where relevant throughout the book.

The successful validation of an ERP system depends to some extent on the ERP system chosen. While the author has personal opinions on the relative merits of different ERP systems, no attempt is made to compare, recommend, or endorse specific systems.

It is up to individual Users to determine requirements; conduct their own package assessments and supplier audits, and plan accordingly (see further in Chapters 2, 3 and 4). Almost any ERP system is capable of successful validation, but depending upon the choice of system there are obvious cost and timescale implications.

Many of the technical issues, suggestions and recommendations discussed in this book are clearly included to address validation problems specific to ERP systems. These are based upon real program experience implementing real systems, and the more experienced reader may recognise the particular ERP system concerned.

Other issues discussed are more generic, and appropriate subject-matter experts who are familiar with other large configurable systems will be able to apply the principles to other systems:

- Laboratory Information Management Systems (LIMS)
- Electronic Document Management Systems (EDMS)
- Manufacturing Execution Systems (MES)
- Electronic Batch Record Systems (EBRS).

As many validation issues are related to program, organisational and people issues, they apply directly to other types of system, and readers may well recognise these from their own experience in implementing other types of system.

1.5 Background and Content

The content of this book is based upon practical experience, both with the successful validation of such ERP systems and, in some cases, a few programs that encountered issues with their validation.

The suggestions and recommendations given are also based on practical experience. These can be used in isolation, when particular problems are encountered during the life of the program, or can be combined when planning a program. The latter approach is generally recommended — ‘prior planning prevents poor performance’.
This book does not attempt to mandate ‘good implementation practice’ — the focus is very much on successful validation rather than successful implementation. Generally speaking successful implementation supports a successful validation simply because the focus is on getting things ‘right first time’. Various chapters thus reference appropriate good implementation practice commonly used on successful projects.

For consistency, various chapters assume that each stage of the ERP implementation is conducted by a common program organisation, following a common implementation model. These organisation and implementation models are defined in the early chapters.

However, every program is different, and the suggestions and recommendations for successful validation should be tailored for the specific program. While it is always good to learn from experience (and especially other people’s issues) there is no substitute for thinking and planning. Among other things this tailoring will depend on:

- The size of the program and system implemented
- The duration of the program
- The implementation methodology employed
- The program’s organisational structure
- The key individuals concerned.

If organisations choose to use this book to help in planning the validation of their new ERP system (or the retrospective validation of their existing ERP system) it is recommended that the planning process includes all key stakeholders (discussed in more detail in Chapter 4).

Validation planning by a limited number of individuals, and failure to communicate the validation approach is one of the fundamental problems that should be avoided.

Readers should also note that the use of the capitalised words ‘User’ and ‘Supplier’ have specific significance. These, along with other terms, are defined in Appendix A.

1.6 Chapter by Chapter

The structure of this book generally follows the system development life cycle. However, this is not as simple as it first appears as one of the fundamental challenges with implementing large configurable systems is the complexity of the implementation life cycle. The order of the chapters generally follows the implementation life cycle, but a number address topics and issues that apply to multiple program phases (or in some cases, the entire program).

Structuring the book in such a way facilitates an overview of the process. It also means the book can be used as a reference, re-reading certain chapters when particular program phases or activities approach, or when specific issues arise.
A typical ERP program has many activities happening in parallel so the order of the various chapters may sometimes appear confusing, or even illogical — welcome to the world of ERP system implementations.

The concluding chapter pulls everything together again, and if you get lost along the way remember that it is okay to skip ahead to the final chapter, and see how it all works out in the end.

1.7 Background Reading

This book is not intended to be a tutorial on the general topic of computer system validation. It builds on the principles established in existing regulatory guidance, industry guidelines, and other books addressing similar issues.

In some cases, the reader will encounter new terminology, or will wonder about the context in which particular terms are used in this book. Appendix A provides a list of abbreviations and acronyms as well as a definition of some of the ERP specific terminology used.

Appendix B provides references and a bibliography of suggested reading for those less familiar with the topic of computer systems validation or large, configurable systems. References to RAD are also provided, but it should be remembered that general RAD approaches employed in non-regulated industries may need to be modified to support the validation of such systems.