

Clarmon Corporation introduces QValid 2.0 with Excel support, which will provide an efficient solution for spreadsheet control to mitigate business and compliance risks

The newly added Excel module extends QValid's scope, allowing users not only to create and manage validation records but also to control Excel templates or workbooks and meet GxP and 21 CFR Part 11 compliance.

Life Science companies focused on improving spreadsheet compliance in an efficient manner can opt to implement only the Excel solution. The solution enables users to continue working in Excel but eliminates the significant business and compliance risks of unauthorized access or modification of critical data.

The weaknesses of using Excel to manage critical quality and compliance records in the Life Science Industry are addressed by the QValid™ Add-in for Excel. The solution enforces GxP and Part 11 compliance by providing automated audit trails at a cell level and supporting automated versioning and change control for Excel workbooks. Furthermore, access to view or modify critical spreadsheets can be restricted or controlled by assigning different rights within the system. Expensive manual procedures, which are routinely needed to maintain spreadsheets compliant, are automated using QValid™, providing significant cost reduction. By supporting electronic signatures, the system eliminates the need for hand signed paper printouts which require significant time and effort to manage. Users can perform electronic execution of spreadsheets in the familiar Microsoft Excel environment and still meet all applicable regulatory requirements.

As a real-life alternative to paper or manual systems, the QValid™ software suite is the optimal solution for Life Science companies focused on standardizing their validation and compliance documentation activities across multiple sites and reducing the cost of compliance. QValid™ ensures that all documents are maintained consistent and controlled and enables users to focus exclusively on critical tasks by automating recurring documentation and compliance processes. Unlike traditional document management and compliance systems, QValid™ allows users to work directly in the familiar Microsoft® Office interface and thanks to its unique Linksense™ technology still ensures full traceability to all related documents, drastically reducing the time needed to manage documents.

The recent QValid™ 2.0 extension to control and manage Excel content is aligned with Clarmon's strategy to reduce the cost of compliance for its customers while allowing users to continue working in the familiar systems. The enhancement extends QValid's capabilities for managing and automating validation and compliance processes, increasing the systems' overall benefits and cost saving potential.

About Clarmon

Clarmon Corporation provides advanced software solutions for quality and compliance management. The majority of Clarmon's customers are Life Science companies that operate inside highly regulated environments. Using Clarmon's Web and Microsoft® Office based solutions these companies have been able to replace paper and manual systems with a fully electronic, automated environment that reduces the efforts and costs of demonstrating compliance.

QValid™, Clarmon's validation and compliance management suite, provides a simple method to centrally manage and control documents without requiring users to re-enter information.

About QValid™

QValid™ is a Web and Microsoft® Office based system which supports centralized management and control of quality and compliance data as electronic records. This enables users to continue working in the familiar document and spreadsheet systems while ensuring that 21 CFR Part 11 requirements are met. Unique Linksense traceability, automated workflows and reporting features further reduce the efforts and costs of managing records.