

VALIDATION + GXP SOFTWARE

Reduce compliance risk. Remove operational friction.

PharmaEddge helps regulated teams validate computerized systems, improve inspection readiness and replace fragile manual workflows with purpose-built software.

RISK-BASED

Validation effort matched to system risk

AUDIT-READY

Evidence written for inspection

FLEXIBLE

Remote or on-site, worldwide

WHAT WE DELIVER

Validation & Compliance

- CSV and CSA
- IQ/OQ/PQ protocols
- Data integrity assessments
- Audit readiness

Custom GxP Software

- Regulated workflow design
- Controlled records and approvals
- Access control and audit trails
- Validation support

Consultation & Advisory

- Fractional QA support
- Remediation roadmaps
- Inspection coaching
- Documentation strategy

COMMON REGULATORY FRAMEWORKS

GAMP 5

ALCOA+

21 CFR PART 11

PIC/S

EU ANNEX 11

CSA

 Start with the pressure point.

DOCUMENT MANAGEMENT SYSTEM

Controlled documents, from draft to master copy.

Document workflows, approvals, master copies and version management across different document types.

THE CONTROLLED WORKFLOW

01

Configurable drafting, review and approval workflows

02

Controlled master copies and effective versions

03

Document-type-specific numbering and metadata

04

Traceable revision history and access control

BUILT FOR

Quality units replacing manual or fragmented document control.

WHY PHARMAEDDGE SOFTWARE

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CONTROL

Replace side spreadsheets and informal tracking with one governed workflow.

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TRACEABILITY

Keep status, history, decisions and evidence connected.

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ADOPTION

Give users a practical interface aligned to the work they perform.

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VALIDATION

Implementation and validation support from the team that builds it.

See it against your real process.