

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.

LMS

Training records that hold up in an audit.

Role-based learning, assessments and controlled training records for regulated teams.

THE CONTROLLED WORKFLOW

01

Role-based training assignment and curricula

02

Assessments, completion tracking and due-date visibility

03

Current training status for employees and functions

04

Audit-ready records, e-signature and audit trail support

BUILT FOR

Pharma quality, manufacturing, laboratory and support functions.

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.

■

ADOPTION

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

■

VALIDATION

Implementation and validation support from the team that builds it.

See it against your real process.