

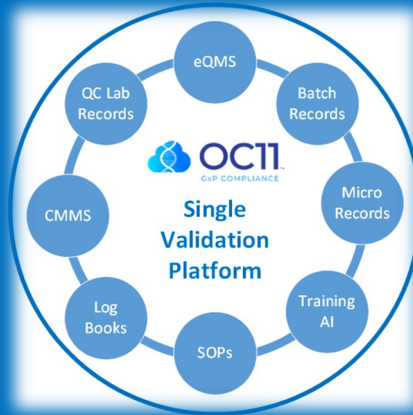
REDUCE COST OF COMPLIANCE

The OC11 Solution System is indeed a proven game-changing platform designed to ensure compliance with GxP and Data Integrity (DI) regulatory requirements. Here are some key points about the OC11 Solution:

Cost-Effective: The OC11 Platform allows adding GxP systems at a fraction of the cost, making it a budget-friendly solution that helps ease the burden on your compliance budget.

GxP and DI Compliance: The system ensures that any CGMP record entered is safeguarded against both intentional and unintentional violations, providing peace of mind for regulatory compliance.

Artificial intelligence (AI): Create the user curricula on the OC11 platform. OC11 will manage the user access based on the curricula and training completion. If you have any questions or need further details, please ask!



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HERE



Reduce Cost of Compliance by 70%



OC11 Platform

CSV is not required for systems developed on the validated OC11 Platform.

Connect:

CPHI Milan, Italy
8-10 Oct 2025
Stand: 6SU83
Hall: 6

Inspected By Global Agencies



Advantages

- Artificial Intelligence (AI) Training Program
- Simple to Use: Works the same as the current Paper System
- No Limit on Number of System Developed at Fraction of cost
- Prevent GxP or Data Integrity Compliance
- Forced Audit Trail Review
- Eliminate Paper Records
- Real-time documents tracking
- Regulatory Inspection or Client Audit ready - 27X7
- Reduce Cost of Compliance

Next Generation Solution



GLOBAL SPEAKER

CPHI North America 2024

In May 2024, at CPHI North America in Philadelphia, USA, the founder and CEO addressed the challenging topic of **"Difficulties Meeting US FDA Regulatory Requirements."**

- Founder & CEO (ADPT)

CPHI MENTOR

CPHI Milan, 2024

ADPT Solution is set to participate in the CPHI mentor program in 2024, offering in-depth knowledge, experience, and practical solutions to support the CGMP within the CPHI community.

SINGLE VALIDATION



Inspected by EU Regulators

Several systems implemented on the Single OC11 Validated Platform underwent inspection and were found to be free of regulatory issues.

- ◆ # of Sites: 4
- ◆ # of Systems: >60
- ◆ # of SOPs: >1000
- ◆ # of Specifications: >2300
- ◆ # of CGMP Records: >2500

Advanced Technologies

- ◆ Training AI
- ◆ CGMP DI logical algorithms

What is Single Validation?

Single validation refers to a system of reusable components that have been validated within the established frameworks of GxP and data integrity algorithms.

Reduce 95% of CSV Cost & Efforts