



EQUIPMENT HUB



Streamline Equipment Logging for GMP Compliance

Using paper logbooks can lead to errors that could cost your operations significant compliance issues and expose your teams to risks that could cost you time and money.

Shiftconnector Equipment Hub helps **ensure data integrity**, **simplifies search and retrieval** for audits and inspections and **minimizes** the need for **multiple entries**.

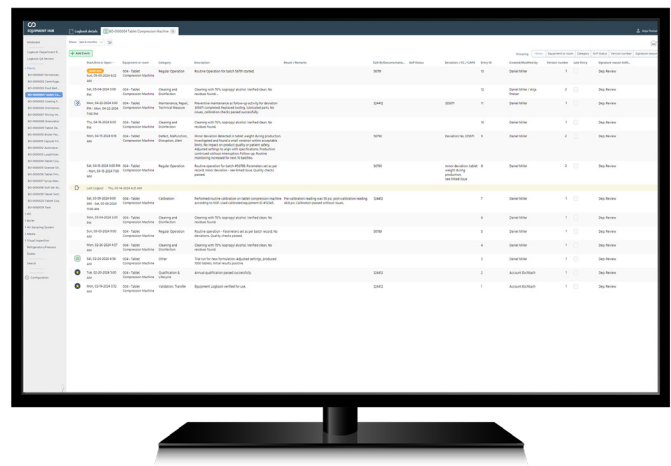
Using **Shiftconnector Equipment Hub**, allows you to

- Add, edit or archive logbooks
- Implement Review Workflows to ensure proper signoffs
- Search across multiple logbooks
- Aggregate data from BDR, LIMS, CMMS and SCADA systems for easier event tracking
- View and search detailed log history records
- Ensure compliance with tailored templates
- Easily create reports
- Have multiple sign offs
- Secure logs with multi-level password authentication

Built specifically with the pharmaceutical industry in mind, it is **fully compliant with GMP and FDA 21 CFR 211.67(a)**. Digital recording of data makes information accessible and transparent, connecting operators, supervisors and quality control in real-time.

Get rid of paper logs and give your teams the digital tools they need to **streamline compliance reporting** and **manage equipment efficiently**.

Contact us to see it in action today!



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