

Ennov PV-Works

Human Pharmacovigilance Software

A flexible software system designed to support pharmacovigilance business processes and technical services case handling practices, while meeting your company safety and world-wide regulatory reporting requirements.

The Pharmacovigilance Challenge

Developing or implementing a system to collect, manage, report and evaluate patient safety data can be time consuming and expensive. Companies of all sizes require a system that is compliant with global regulations, easy to access and use, fast to implement, and simple to maintain.

An ideal system should be flexible, should integrate with other databases, and should be designed to manage not only adverse events from clinical trials and those occurring spontaneously in the field but can also manage medical inquiries and product quality complaints for all medical products, including traditional pharmaceuticals, biomedical products and medical devices.

Comprehensive PV Data Management

The Ennov PV-Works database has been designed around the ICH E2B guideline such that every E2B data field maps to a specific database field. Coding to MedDRA is simple. In addition to paper reporting outputs such as CIOMS and MedWatch 3500s, an electronic reporting function is included. Aggregate listing outputs like PSUR, DSUR, and PBRER are simple to generate using built-in tools. In addition to simple yet comprehensive data entry and reporting functions, the fully integrated workflow functionality will ensure SOP compliance and that critical reporting deadlines are met. Regulatory compliance is afforded with the adherence to key requirements such as FDA 21 CFR Part 11 and the provision of a full audit trail.

Core capabilities

ICH E2B compliant safety data collection and reporting
Coding of cases against current MedDRA dictionary
Extensive data validation, cross-field checks and use of pick lists
Duplicate check functionality
Integrated spell checker
Integrated query tool
Automated letter generation
Case data export

Key Features

Manages spontaneous and clinical trial adverse events, technical inquiries and product complaints
Compatibility with with third-party query and reporting tool
Easy to configure workflows that mirror existing business processes
Streamlined case review and data approval concept
Supports pharmaceutical, biological, medical device, and cosmetic vigilance
Available as cloud-based or on-premises deployment
FDA 21 CFR Part 11 compliance



Cloud-based or On Site



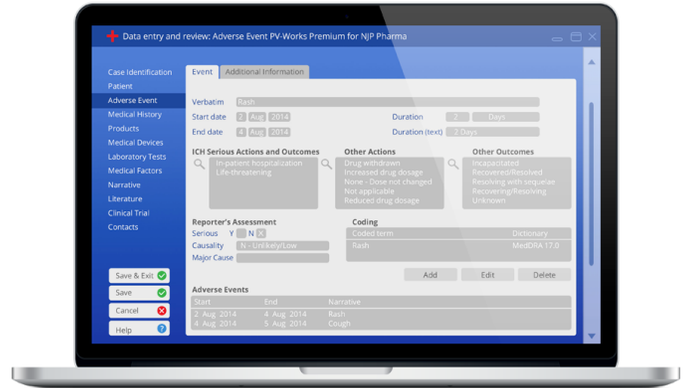
Multi-Platform



ISO 9001:2015 Certified

Advanced Capabilities Ensure Compliance

PV-Works is a process driven system. A purpose-built workflow engine is integrated with comprehensive safety functionality to provide management control of pharmacovigilance processes. The software also includes a wide range of powerful querying functionality, allowing business teams to monitor case handling, track compliance, and execute in-depth trend analysis. PV-Works submits electronic adverse event (AE) reports in full compliance with the ICH E2B standards. Furthermore, electronic medical device AE reports may be submitted to FDA CDRH in eMDR format. The system is easily integrated with standard AS2 gateway software to manage electronic submissions.



PV-Works Human – Part of the Ennov Pharmacovigilance Suite



PV-Works Human



PV-Works Vet



PV-Analyzer



PV-Signal Manager



PV-Entry



PV-Express

Why Choose Ennov?

Hundreds of companies trust Ennov

- **Over 15 years of experience providing PV solutions**
200+ life sciences customers, many more in other industries.
- **Modern architecture and user interface**
100% web-based, Highly scalable. User-centric design.
- **Our commitment to your success**
Very high customer satisfaction. 98.5% of projects delivered on time and within budget.

Providing you freedom of choice

- **Available as cloud-based or on-premises deployment**
You can switch between deployment options at any time.
- **We make you autonomous**
System configuration and management require no IT skills.
- **Improved security and optimized performance**
Data is hosted locally for total flexibility. Single tenancy minimizes business interruptions

Learn more about our unified content and information management platform to support the entire Life Sciences product development continuum at www.ennov.com



Quality



Clinical



Regulatory



Pharmacovigilance



Commercial