Ludwig Institute for Cancer Research Turns to Medidata and EXTEDO for End-to-End Pharmacovigilance Solution

The Challenge
The Ludwig Institute for Cancer Research is a nonprofit research organization committed to improving the control of cancer through integrated laboratory and clinical research, and novel therapeutic strategies based on the emerging understanding of cancer. Ludwig runs clinical oncology trials based on discoveries made by its researchers that are focused on exploring the therapeutic modalities of cancer vaccines, targeted antibodies and small molecule inhibitors. During these trials, Ludwig captures data from trial participants, including serious adverse events (SAEs), which must be understood, distributed, triaged and reconciled in order to maintain a high level of patient safety.

Faced with the standard, paper-based SAE reporting process and a growing number of international studies, Ludwig sought to automate and optimize their system. An automated system would eliminate manual data entry and reconciliation of the safety and clinical databases.

The Solution
The Ludwig Institute chose to create a complete, end-to-end SAE solution by enabling and configuring Rave Safety Gateway, Medidata Rave EDC-to-safety-system interface, with EXTEDO’s PcVmanager, a drug safety management solution built specifically with open data standards in mind. This cross-vendor solution leverages the E2B standard for SAEs using an AS2-compliant electronic data interchange (EDI) service, allowing a fully electronic workflow from data entry to Medwatch/CIOMS creation, complete with an integrated Rave EDC query system for resolving any questions between Ludwig and the clinical sites. PcVmanager’s gateway can also be configured to submit electronically to the FDA and EMA in addition to partners. With EXTEDO providing a fully centralized safety database, this integration allows all medical coding to occur directly in Rave EDC. Any user with coding questions can utilize Rave EDC’s flexible query work flows before sending the coded safety data on to PcVmanager.

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One Fully Electronic Workflow

- From adverse event data entry to medical coding, triage, analysis and agency communication, all safety events follow a common electronic workflow.
- Duplicate data between EDC and Pharmacovigilance (PV) forms is not necessary. All SAE data is entered directly into Rave EDC.
- Both Rave EDC and PcVmanager provide reports with consolidated views of all SAEs available in a fully transparent fashion.
- PcVmanager continues the thread of electronic workflow for each SAE with the ability to recode, classify, review, categorize and submit.

SAE Reconciliation Reduced

- SAE queries can be resolved quickly using the standard Rave EDC query workflows.
- Affiliates and large partners are able to use a familiar query interface in Rave EDC, and are able to see at all times the latest version of the safety data.
- Little to no reconciliation effort is necessary at the end of each trial, saving hours to weeks in reconciliation time.
- There are significant savings in time and resources with the implementation of a seamless, electronic SAE reporting process.

The Platform of Choice for Clinical Research

The Medidata Rave Clinical Cloud is the cutting-edge platform that transforms the clinical trial experience for patients, sponsors, CROs, and research sites. Designed with a unified data platform, the Rave Clinical Cloud creates a single source of truth for all study-related data. Simply put, enter data once and let the platform master and populate it throughout the end-to-end suite of Rave applications. Optimize operational execution, decrease the data entry and maintenance burden, and reduce the number of clinical systems across your study teams. Throw away your list of passwords and excel sheets, you are now on a truly unified platform.