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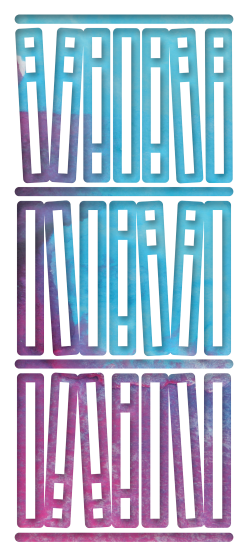
# Adis Pharmacovigilance

Get ahead of the game

The expert solution for local literature monitoring

Tightening safety regulations and a heightened focus on local literature in inspections and audits mean it is increasingly important for pharmaceutical companies to demonstrate that their local literature monitoring is performed in a consistent, complete and timely manner.

Wider global market distribution, increasing drug portfolios and variability in the monitoring process between individual offices have made outsourcing of this resource-consuming task an attractive alternative for many drug companies.



## The challenge for pharma: timely and efficient workflows in the face of increased scope and responsibility

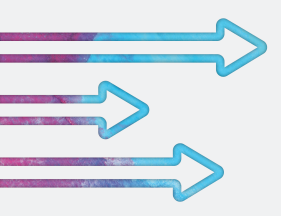
There is much variation in the extent and specificity of local literature monitoring requirements between local regulatory authorities, making it challenging to determine the quantity and quality of efforts required. With the increasing pressure on resources, you need to balance doing enough to satisfy an inspection against the burden of performing this laborious and often manual task.

In addition, the nature of local journals themselves is diverse in many aspects, such as online accessibility, full text availability, frequency and reliability of publication schedules as well as type and presentation of content.

Challenges faced with performing this activity may include determining which journals to monitor in the first place, ensuring timely identification and reporting, and keeping a sufficient audit trail.

### Adis Pharmacovigilance – the solution for compliant and flexible local literature monitoring

- Currently monitoring more than 500 journals in over 30 countries, covering multiple languages
- Identification and recommendation of relevant journals, according to the countries and therapeutic areas of your interest
- Flexibility to expand the journals monitored on request
- Timely and compliant delivery of relevant safety reports, in the format of your choice (including E2B-compatible XML for ICSR)



#### Contact us

Please contact us to learn more about how you can partner with Adis Pharmacovigilance, the experts in regulatory literature monitoring.

We can provide you with a solution that delivers reliable and timely results – and is easily scalable to suit your evolving needs.

Email pharmacovigilance@adis.com