

IQVIA Regulatory PDF Tools

Ease your regulatory publishing burden

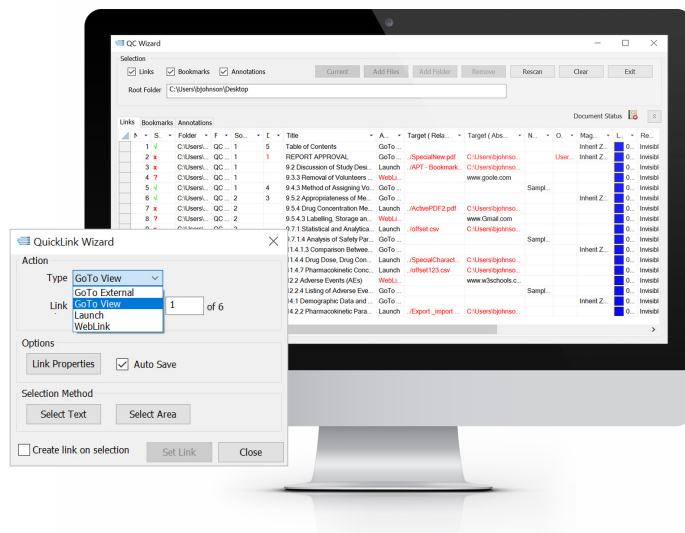
Increase your productivity

Staying compliant with agency requirements demands significant resources. To succeed in today's rapidly changing regulatory ecosystem, companies must find new technology-enabled approaches—large and small—to increase their efficiency, productivity and speed of implementing these mandatory activities.

PUBLISH SUBMISSION-READY PDF DOCUMENTS MORE QUICKLY AND EASILY

Ninety to ninety-five percent of regulatory e-submissions are converted to PDF documents for dispatch to the 150+ regulatory agencies around the world. **IQVIA Regulatory PDF Tools (IRPT)** – specifically designed for the life sciences industry – provide regulatory staff with a simple, powerful way to prepare, publish and deliver PDF documents.

- Assists publishers and reviewers working with regulatory documents
- Addresses specific needs in accordance with regulatory agency and industry requirements
- Employs intuitive user interface and built-in intelligence to maximize efficiencies



IRPT is an Adobe Acrobat plug-in consisting of ~ 30 tools or “wizards”, each with a distinct use. A few examples include:

- When a document delivered in PDF format requires content changes, fix many things at once – such as links – with a single click
- QC wizard will identify and report areas that are out of compliance with agency requirements before you finalize the documents for submission
- Verify that the link or bookmark is going to the intended document location
- Easily add bookmarks and tables of contents to documents received from authors
- Automatically save documents according to pre-set agency standards using the easy auto-save wizard

FIT-FOR-PURPOSE WIZARDS

Each wizard is designed to meet a specific need according to regulatory agency requirements and industry standards

STREAMLINED, COST-EFFECTIVE PROCESSES

IQVIA Regulatory PDF Tools are powerful, fast, reliable – and importantly – affordable

BASED ON DOMAIN EXPERTISE

Our IRPT architects have decades of experience and knowledge of PDF specifications and regulatory requirements from agencies across the globe

RESULTS YOU CAN COUNT ON

Improved submission quality, simplified review processes, increased speed of submission and higher probability of technical approval

IQVIA	Help
Bookmark Wizards	▶
Hyperlink Wizards	▶
Bookmark and Hyperlink Wizards	▶
PDF Wizards	▶
Batch Wizards	▶

IQVIA REGULATORY PDF TOOLS TAB

IQVIA Regulatory PDF Tools are easy to navigate. Once the user has installed IRPT and opens Adobe Acrobat, the 'IQVIA' tab will appear in the Acrobat menu bar and in **Tools** → **IQVIA** as shown in this figure.

Tool (“Wizard”) Categories

Bookmark Wizards

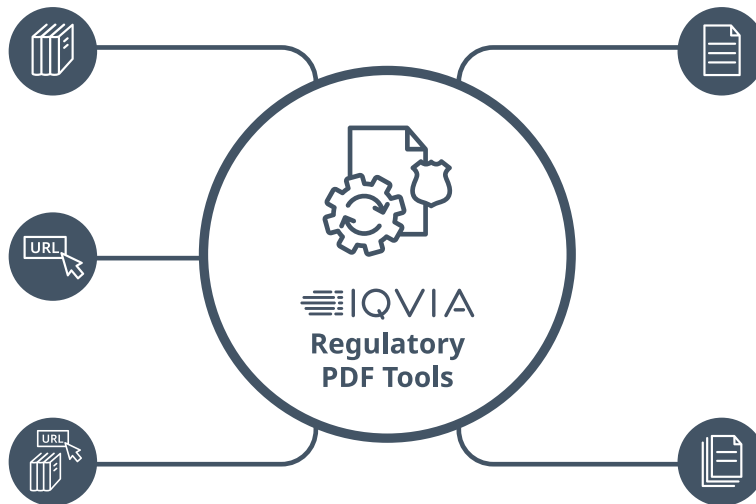
- By font
- From link
- Bookmark
- Swap bookmarks

Hyperlink Wizards

- Doc Link
- Link
- Missing Link
- Quick Link
- Smart Link
- Text Link

Bookmark and Hyperlink Wizards

- Import/export
- QC
- Quick Tools
- TOC



PDF Wizards

- Auto Save
- Redact
- Content
- Merge
- Link Info
- Page Stamp
- Page Size
- Review
- Split
- TOC Generator

Batch Wizards

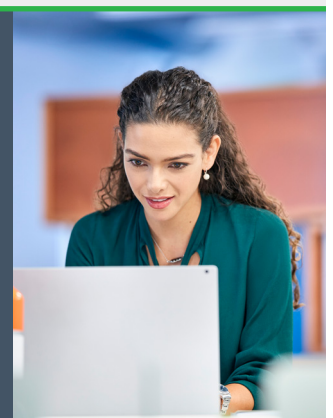
- Batch AutoSave
- Doc Info
- Page

ASK ABOUT OTHER IQVIA REGULATORY PRODUCTIVITY TOOLS

REGULATORY TEMPLATES	eSUBMISSION VALIDATOR
320+ authority-defined regulatory shell documents	Verify tech compliance ahead of submission
Menu-driven template selection	Minimize refusal to file or rejection risk
Fit-for-purpose regulatory-specific toolbar	Ensure compliance with regional and ICH requirements

TRY BEFORE YOU BUY

Test drive IRPT before making a commitment. Access IQVIA Productivity Tools in a complimentary trial. Discover how they can simplify your regulatory staff's daily activities, improving cycle times and minimizing risk.



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