

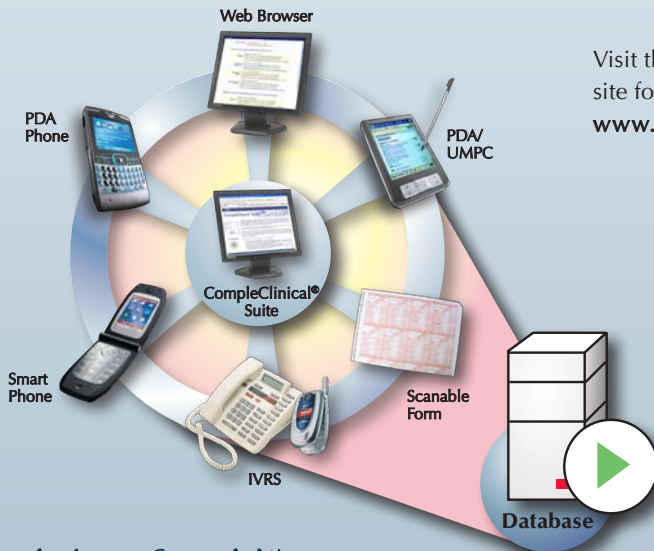
WebEPRO® enables patient reported outcomes data (diaries, quality of life measures, and other PRO end points) to be collected electronically and viewed on the Internet.

WebEPRO®:

- Uses an administrative tool to define protocols - does not require any custom programming allowing rapid setup (1-2 weeks)
- Is platform independent - Web, PDA, telephone, IVRS, scannable form
- Uses CompeWare unified user log on (OneStop™)
- Is a component of the CompleClinical® Suite
- Generates reports and study metrics to follow study progress online
- Is unmatched for simplicity for users and complexity of real time edit checks
- Is compliant with 21 CFR Part 11 requirements for electronic records

web-e-pro (web'e prō)

Computer data collection, data entry, data analysis, data management, data organization and protocol design and monitoring for clinical trial information and data; clinical research software for use in the collection, entry, analysis, management, organization, validation, design, monitoring, randomization and conversion of clinical trial information and data; a trademark/servicemark



Visit the CompeWare Web site for more information:
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