The Only Software That Has It All!

- Calculations and reporting meet all current FDA/ICH/USP validation guidances – including the new USP <1210>!
- Can be used for LC and Non-LC methods (e.g. GC, CE, Q-NMR)!
- Automates LC method validation experiments on multiple instruments and CDS systems!
- Regulatory accepted validation for both Small & Large Molecules!
- Statistically rigorous and defensible robustness testing!
- Handles multiple compounds – creates complete reports for each!
- Shortens your LC method validation time by as much as 75%!
Automated Experimentation for LC Method Validation

The objective of Method Validation is to provide documented evidence and a high degree of assurance that an analytical method employed for a specific test is suitable for its intended use. Method Validation is a regulatory requirement as much as a scientific necessity.

Key Benefits

- Full Automation for LC Method Validation – multiple LCs and CDS systems
- Aligned with FDA and ICH guidances
- 21 CFR II compliance support toolset –
  - Including E-records and E-signatures, full audit logging
  - Workflow management system with E-review and E-approve loops
- Easy setup of experiments –
  - Create standardized workflow templates
  - Facilitate rigorous practice and defensibility
- Simple documentation review and reporting –
  - Easy to defend and communicate
  - Reports meet all FDA and ICH guidelines

Method Validation Experiment Suite

- Analytical Capability and System Suitability
- Specificity
- Filter Validation
- Sample Solution Stability (stability for a given time period under prescribed conditions)
- Accuracy
- Linearity and Range
- Repeatability (intra-assay precision)
- Accuracy/Linearity and Range/Repeatability – Combined Design
- [ICH-Q2(R1) – Accuracy, Linearity, and Repeatability can be done together as a single combined experiment]
- LOQ, LOD
- Intermediate Precision and Reproducibility (USP Ruggedness)
- Robustness – done the right way!

Non-LC Method Validation Experiments

Used successfully for Non-LC methods such as GC, CE, Q-NMR, as well as hyphenated methods (e.g. LC-MS). Accepted in customer regulatory submittals.
Automated LC Method Validation – Five Step Workflow

1. You complete a simple experiment setup template.
2. Fusion QbD creates the Validation Experimental Design and exports it to the CDS.
3. The CDS runs the validation experiment sequence.
4. Fusion QbD imports and analyzes the results.
5. Fusion QbD automatically creates final reports and graphs.

Example Workflow – Combined Accuracy / Linearity / Repeatability

Step 1 – You Complete the Simple Template

Fusion LC Method Validation Software (FMV) has simple experiment setup templates for each type of validation experiment. The simple Linearity and Range template is shown below with user definable settings:

User-definable Settings – Basic Setup

- No. of Compounds
- No. of Levels per Compound
- 100% Standard Level
- Compound Name, Units, and Levels

User-definable Settings – Standards Setup

FMV has a flexible Standards Setup wizard which enables you to select your desired standards strategy for results quantitation within the CDS:

- Bracketing – Overlap
- Bracketing – Non-overlap
- Grand Average
- Calibration and Check Standards
- Multi-level Bracketing – Overlap

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Step 2 – Fusion QbD Creates the Validation Experimental Design and Exports it to the CDS

FVM automatically constructs the validation experiment designs within the CDS as ready-to-run sequences/samples with the proper Vial No. and Injection Type designations for Samples, Standards, and Blanks.

Step 3 – CDS runs the Validation Experiment

FVM sequences run automatically on the CDS. FVM even enables you to include a Shutdown method as the last method run so that you can execute FVM sequences overnight while you sleep!

Step 4 – Fusion QbD Imports and Analyzes the Chromatogram Results

FVM automatically imports the required peak result data from the CDS, and re-maps the results to the design for automated analysis, graphing, and reporting. This is a key feature ensuring quality, as manual transcription is a common source of error and risk.
Flexible Data Analysis Setup Wizard

- Associate different responses with different analyses – e.g.
  - Associate **Amount** results data with analysis of Accuracy
  - Associate **Area** results data with analysis of Linearity
- Include LOD and LOQ and select Calculation Method(s)
- Set Global and Level-specific Acceptance Criteria
- Include Level-specific Spec Limits for Raw Data

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**Step 5 – Fusion QbD Automatically Creates Final Reports and Graphs**

**ICH Q2(R1). LINEARITY**

... If there is a linear relationship, test results should be evaluated by appropriate statistical methods, for example, by calculation of a regression line by the method of least squares...

The correlation coefficient, y-intercept, slope of the regression line, and residual sum of squares should be submitted. A plot of the data should be included...:

- Correlation Coefficient
- Y Intercept
- Slope of the Regression Line
- Residual Sum of Squares
- Linear Regression Plot
- Residuals Data Table and Plot

---

**FMV** also enables you to include images of representative chromatograms into your final reports.

**ICH Q2(R1):**

For chromatographic procedures, representative chromatograms should be used to demonstrate specificity, and individual components should be appropriately labeled...
Robustness Validation – DONE RIGHT!

Experiment Setup – LC Robustness Example

You select the parameters to include in the FMV robustness experiment. FMV will automatically generate the robustness design, reconstruct it in the CDS as ready-to-run methods and sequence, import the chromatogram results directly from the CDS, re-map them to the robustness study, and instantly analyze, graph, and report the results.

FMD supports Validation Robustness studies for:
- Isocratic Methods
- Gradient Methods
- Reversed Phase
- Normal Phase
- Chiral
- Ion Exchange
- HILIC
- Size Exclusion

FMD provides visual displays to simplify setup for complex settings such as required pump program conditions and key settings for each included column such as pH upper limit and conditioning time.

The FMV Difference Lowers your Field Failure Risk

FMV robustness analysis wizard lets you set:
- expected parameter variation ranges
- acceptable performance limits for each key response

The wizard then accurately determines and reports the method’s true robustness.
Robustness Validation – Statistical Significance Testing – Model Coefficients

Robustness Report - API - Area (A)
Coded Variable Name - Key

<table>
<thead>
<tr>
<th>Coded Variable Name</th>
<th>Actual Variable Name</th>
<th>A</th>
<th>Moist % Organic</th>
<th>B</th>
<th>Oven Temperature</th>
<th>C</th>
<th>AI</th>
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<tbody>
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</table>

Variable Effects Table - Statistical Significance

<table>
<thead>
<tr>
<th>Model Term</th>
<th>Robustness Testing Range (Coded)</th>
<th>Coefficient Value</th>
<th>Predicted Effect</th>
<th>Effect Standard Error</th>
<th>Effect Statistic</th>
<th>Effect Fail</th>
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<tbody>
<tr>
<td>B</td>
<td>0.0000</td>
<td>4.5203</td>
<td>69.9836</td>
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<tr>
<td>A</td>
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Robustness Validation – Practical Significance Testing – Effects Magnitude

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Automated LC Method Validation – Proven ROI

International Pharma Co. Benchmarking Project
Realized Time Savings = 85%.

Using historical records* and adjusting for project complexity
Minimum Expected Time Savings = 60%.
S-Matrix Software Products and Support

S-Matrix Corporation develops advanced Design of Experiment based-software that automates R&D experimental work according to Quality-by-Design principles and methodologies. S-Matrix’s Fusion QbD platform automates and redefines experimentation in Analytical R&D, Chemical and Process R&D, Formulation, and Product R&D.

Fusion QbD Software System Product Suite

■ Fusion LC Method Development
  Fully automated QbD experimenting on your LC system, integrated DOE, automated robustness simulation & chromatography data modeling. Chemistry screening without the need for peak tracking.

■ Fusion Analytical Method Validation
  Meet regulatory guidelines with a best-practices approach toward LC method validation with comprehensive reporting. Also supports formal validation of Non-LC methods (e.g. GC, CE, Q-NMR).

■ Fusion Inhaler Testing
  Create sampling plans, export and import data from your CDS via validated data exchange, calculate particle size distribution results, and generate reports according to USP 601, Ph.Eur. 2.9.18, and ISO 27427.

■ Fusion Product Development
  The perfect QbD software for formulation & product development – automated experimental design selection, sophisticated analysis tools, including automated modeling and simulation, comprehensive reporting, with a full 21 CFR 11 compliance toolset.

Sales and Support

Sales: Tel: 800-336-8428 (Outside the USA: 707-441-0406). Email: Sales@smatrix.com
Customer Support: Tel: 707-441-0407. Fax: 707-441-0410. Email: Support@smatrix.com

On-site and Web Training

S-Matrix offers on-site training programs for installed systems. Training includes experiment strategies, experimental design (DOE), data analysis, graphical visualization and ranking of effects, numerical and graphical optimization, and QbD Reporting.

S-Matrix also offers interactive web training which covers software features and operation, along with general principles of DOE and QbD. Web training programs can be tailored to suit your individual focus and information requirements.

To arrange an on-site or web-based training program, call 707-441-0406.