

Quantitative and Qualitative Analysis of Cell Culture Medium Using SWATH[®] Acquisition

Featuring Data Independent Acquisition on SCIEX TripleTOF[®] 6600 System

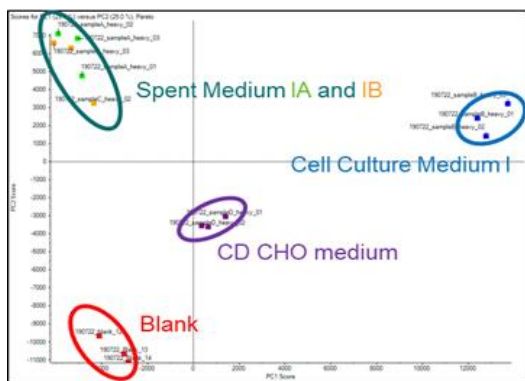
Zuzana Demianova¹, David Cox², Elsa Gorre³, Andrew Mahan³ and Lei Xiong⁴

¹ SCIEX, Brea, CA, USA; ² SCIEX, Concord, CAN;

³ Janssen Research & Development, Spring House, PA, USA; ⁴ SCIEX, Redwood Shore, CA, USA

Biotherapeutic production requires precise monitoring of cell culture medium components across development and manufacturing stages. Cell culture media (CCM) and feeds tailored to specific cell lines maintain or improve the process yield, drug efficiency, safety and quality consistency. While there is a vital need to develop analytical methods for cell culture media (CCM) analysis, the diversity of compound classes and wide dynamic range of their natural abundances make the assay development challenging.

Recently, SCIEX introduced a MRM based workflow to quantify 110 cell culture components with a single LC-MS method.¹ This MRM workflow provides superior quantitation of the targeted CCM analyte in the defined list. Beyond the targeted quantitation workflow, there are also increased discussions in this area of research on simultaneously quantifying and confirming all analytes (expected or unexpected) in the CCM. In this aspect, SWATH[®] acquisition has been attracting significant interests, as it provides high resolution MS/MS level quantification and confirmation/identification of every detectable compound. Herein, a data independent workflow using SWATH[®] acquisition is demonstrated for comprehensive CCM analysis.



Statistical Comparison Between Different Media. PCA plot shows a good separation between blank, cell culture media and spent media.



ExionLC[™] System coupled to TripleTOF[®] 6600 LC-MS/MS system and SCIEX OS 1.5 for data processing

Key Features of Cell Culture Media Analysis using SWATH[®] Acquisition

- SWATH[®] acquisition on the TripleTOF[®] system offers:
 - Wide analyte coverage by quantifying and confirming every ESI MS detectable compound in the medium
 - Superior quantitation quality based on high resolution MS/MS spectra, compared to single MS spectra
 - Ease of method development by using generic method parameters with minimum method optimization requirement
 - Variable Q1 window acquisition by optimizing window width based on m/z density of precursors, allows for greater selectivity in chromatographic periods of high precursor ion density
- Phenomenex Kinetex[®] F5 column provides excellent resolution of target analytes across different chemistries
- The Accurate Mass Metabolite Spectra Library allows compound confirmation with over 550 culture media components and metabolites for biological processes
- Powerful, comprehensive software solution including SCIEX OS-Q and MarkerView[™] software offers versatile qualitative and quantitative workflows with statistical analysis

Table 2. Summary of mass spectrometry parameters.

Parameter	Value	Parameter	Value
MS mass range	50-700 m/z	MS/MS mass range	25-700 m/z
MS accumulation time	100 ms	MS/MS accumulation time	35 ms
Curtain gas:	30 psi	Source temperature:	400 °C
Ion source gas 1:	50 psi	Ion source gas 2:	50 psi
Polarity:	+ or -	Ion spray voltage:	5500 or -4500V

Table 1. The summary of cell culture media component coverage among various compound groups.

Component group	Number of components
Amino acids	39
Vitamins	15
Carbohydrates	4
Fatty acids	5
Nucleic acids	17
Others	32

Methods

Sample preparation: The stock solutions of individual standards (1 mg/ml) from various compound classes (Table 1) were prepared with different solvents depending on compound solubility. The final master mix was prepared by mixing the stock solutions, resulting in final concentration range from 6.67 to 20 µg/ml. CD CHO medium (Gibco), cell culture medium I and its spent media were diluted 5-fold with 0.1% formic acid in 50% acetonitrile, and centrifuged. The supernatants were collected and further diluted 60 fold with 0.1% formic acid prior to LC-MS analysis.

Chromatography: Samples were subjected to LC-MS analysis in triplicate by using a TripleTOF® 6600 LC-MS/MS system coupled with an ExionLC™ system. Analytes were separated using a Phenomenex Kinetex® F5 column (150 mm x 2.1 mm ID, particle size 2.6 µm). The LC run time was 20 min at a flow rate of 200 µL/min. Mobile phase A was 0.1% formic acid in water, and mobile phase B was 0.1% formic acid in acetonitrile. Column oven temperature was 40°C and the injection volume was 5 µL.

Mass Spectrometry: The data were generated with variable window SWATH® acquisition using 20 variable windows. The window widths were determined based on the MS ion intensity

distribution using the SWATH® acquisition Variable Window Calculator.² The method details are listed in table 2.

Data processing: The data were analyzed using SCIEX OS 1.5 software with details listed below.

In the targeted workflow (quantitation and confirmation), an MRM transition library¹ with 196 positive and 71 negative entries (Table 3) was incorporated into SCIEX OS to generate the quantitation method. Peak integration was performed with MQ4 algorithm. For compound confirmation, the acquired MS/MS spectra were matched against MS/MS spectra in the built-in high resolution spectral library. This built-in spectral library contains entries from the SCIEX Accurate Metabolite Spectral library, Antibiotics and NIST library.

In the untargeted compound screening workflow, the LC-MS peak picking option was set as semi-exhaustive. All LC-MS peaks for the same analyte in the forms of different charge states or adducts were grouped together and analyzed as a single entry, which significantly simplified the data process. To achieve the optimal performance of compound identification, three processing functions were used: 1) FormulaFinder: identify possible compound formula based on TOF MS spectra (compound molecular weight); 2) ChemSpider: link compound formula with in silico MS/MS database predicted from compound structures; 3) library search: search MS/MS spectra with built-in spectral library.

Statistical analysis: MarkerView™ software was used for analyzing cell culture media and spent media samples with statistical tools, including principle components analysis (PCA), t-test, volcano plots and differential abundance visualization between media.

Overview of SWATH® acquisition

In this project, SWATH® acquisition was utilized to develop a comprehensive workflow for quantitative compound profiling in the cell culture medium. SWATH® acquisition, also known as data independent acquisition (DIA), collects MS and MS/MS information from all ionized compounds, with no requirement to obtain sample information prior to data generation. Compared to the data dependent acquisition (DDA), SWATH® acquisition workflow demonstrates significant advantages in multiple aspects: 1) analyte coverage: it allows to keep permanent quantifiable digital records of every detectable compound in the medium; 2) quantitation capability: it provides high resolution MS/MS based quantitation; 3) ease of method development: it uses generic method parameters with minimum method optimization requirement.

The SWATH® acquisition workflow is demonstrated in Figure 1. The Q1 quadrupole transmits ions within a wide mass range (m/z isolation window) through to the collision cell. This produces a MS/MS spectrum which is a composite of all the analytes within this wide Q1 m/z window. Because the fragment ions are high resolution, high quality XICs can be generated post-acquisition to produce the MRM-like data. This Q1 window can be stepped across the mass range, collecting full scan composite MS/MS spectra at each step, with an LC compatible cycle time.

SWATH® acquisition also offers variable window acquisition strategy³ by allowing the user to vary the size of the Q1 isolation window based on the density of analyte precursor masses (Figure 2). In this project, an optimal method with 20 variable isolation windows was built to allow full mass range coverage and high quantitation performance

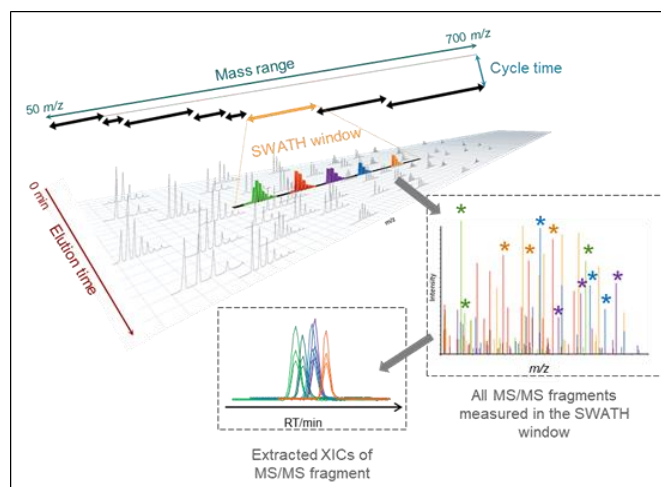


Figure 1. The general workflow of SWATH® acquisition.

Cell culture media analysis workflow using SWATH® acquisition

The above described SWATH® acquisition workflow was applied for both qualitative and quantitative analysis of cell culture medium (Figure 3). In the targeted (quantitation and confirmation) workflow, an MRM transition library¹ was incorporated into SCIEX OS to generate quantitation method. Peak integration was performed with the autopeak algorithm. The abundant fragment ions of the target analyte were automatically extracted by software for peak integration. Figure 4 showed an example of choline quantification and confirmation in master mix, in which the calibration curves (Figure 4A) and XICs (Figure 4B) of multiple fragment ions, TOF MS spectrum (Figure 4C) and MS/MS library matching (Figure 4D) were presented. Figure 5 demonstrated the quantification of selected analytes in serial dilution samples with positive or negative ionization. Another example (Figure 6) was shown as the quantification of

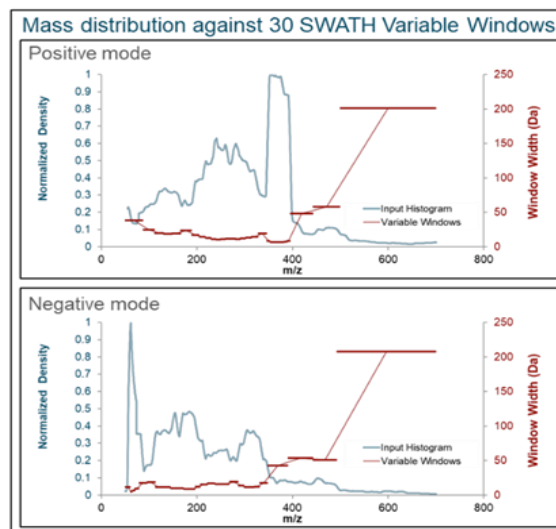


Figure 2. Graphic representation of windows size variation. The size of the Q1 isolation window (red line) is determined by the density of component precursor masses (blue line).

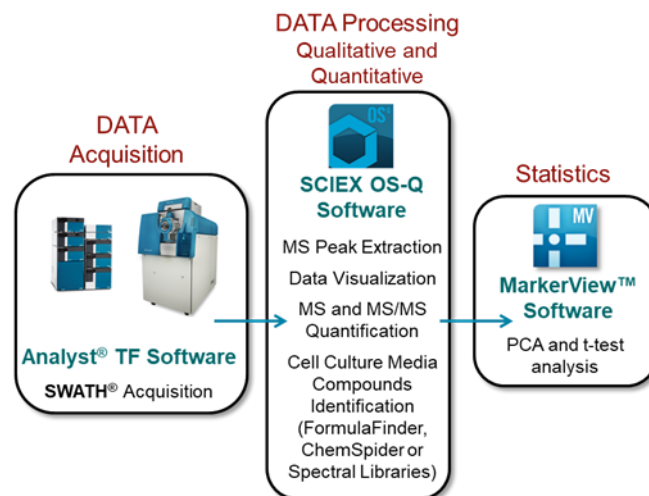


Figure 3. Cell Culture media workflow: from data acquisition to trends visualization. Data are acquired in SWATH® acquisition mode, qualitatively and quantitatively processed with SCIEX OS-Q, and statistically analyzed with MarkerView™.

multiple components in the Gibco's CD CHO cell medium, in which the analyte abundances covered over four orders of magnitudes.

In the untargeted compound screening workflow, both MS and MS/MS spectra were processed by SCIEX OS-Q for compound identification (Figure 6). The TOF MS spectrum was linked with predicted compound formula by FormulaFinder and searched against ChemSpider for structure matching and fragment ion prediction. The MS/MS spectrum is searched against MS/MS spectral library wherein multiple libraries (the SCIEX accurate

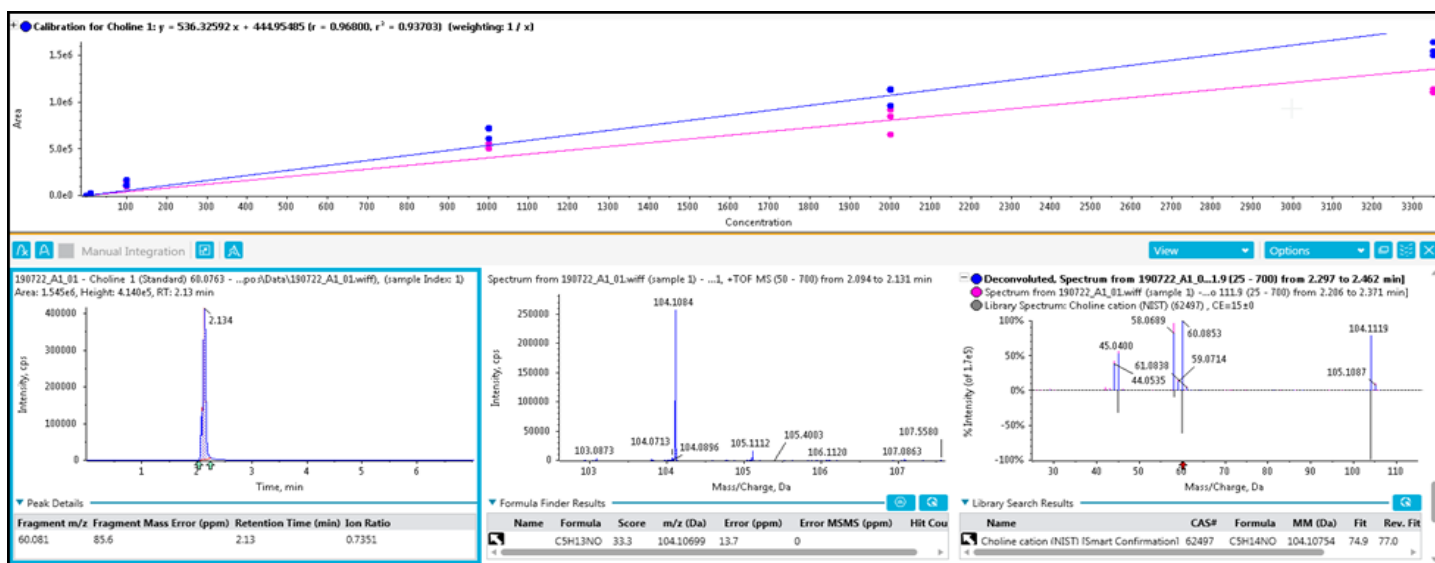


Figure 4. Choline quantification and confirmation in master mix. A) calibration curves; B) (left) XICs of multiple fragment ions; C) (middle) TOF MS spectrum with FormulaFinder matching; D) (right) MS/MS spectrum with library matching.

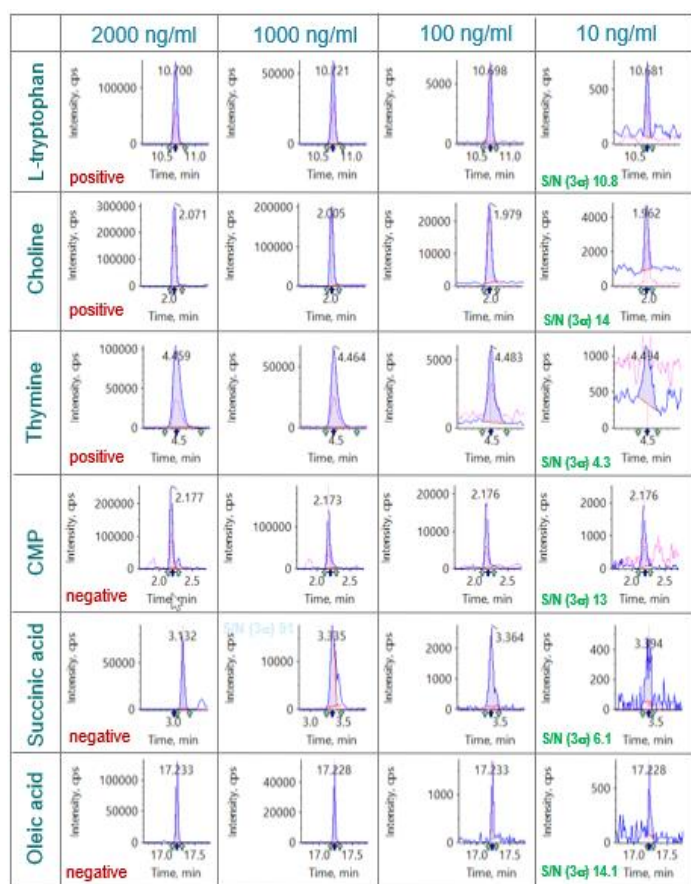


Figure 5. XICs of representative compounds in serial dilution master mix (10, 100, 1000 and 2000 ng/mL) with positive or negative ionization.

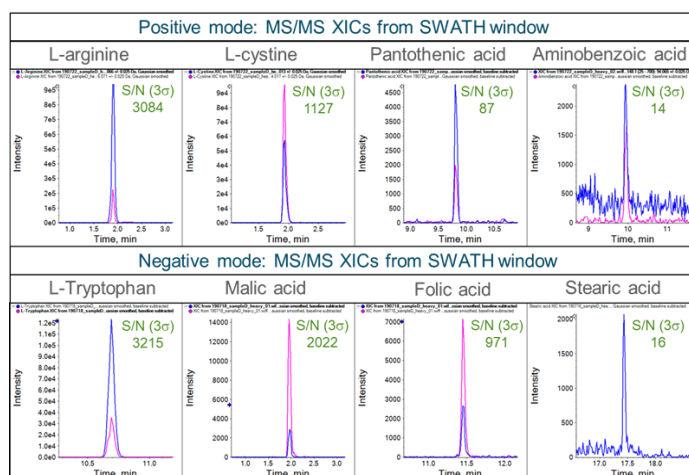


Figure 6. XICs of representative compounds from Gibco's CD CHO medium. Selected analytes abundances covered four orders of magnitudes. The medium sample was diluted 300 fold prior to analysis.

spectra metabolite and antibiotics library plus a subset of the NIST spectral library) could be combined to achieve optimal compound coverage. This workflow significantly reduced false discovery rate by providing MS/MS level confirmation in addition to the retention time and precursor mass information. Figure 7 shows an example of unknown compound identification in the medium sample. The compound was identified as Spermine which is used in serum free cell culture medium to improve expression levels recombinant protein, cell growth and viability.⁴

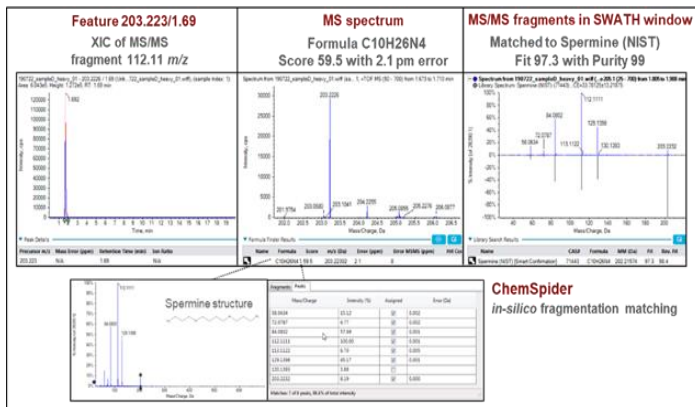


Figure 7. Untargeted compound screening using SCIEX OS-Q software. Spermine was identified as an addition to known cell culture medium compound. The TOF MS spectrum $[M+H]^+$; 203.2226 m/z) was matched to compound formula (C10H26N4) and searched against ChemSpider. The MS/MS spectrum was searched against MS/MS spectral library, showing a good matching (Fit 98.3, Purity 99%) between measured MS/MS spectrum (blue MS/MS spectrum) and library spectrum (grey MS/MS spectrum).

Statistical Analysis

SCIEX MarkerView™ software was applied in cell culture media analysis workflow for statistical evaluation. The data generated from different cell culture media and spent media samples were analyzed in SCIEX OS-Q and imported to MarkerView™ software for principal components analysis and analyte abundance visualization among multiple samples. As shown in Figure 8A, two spent media were grouped together and well separated from the “fresh” cell culture medium (cell culture medium I.). This suggested that there were significant variations in component regulation between start and end point of biopharmaceuticals production. As shown in Figure 8B, levels of L-serine varied among media samples (decreased during

production). L-Serine (a non-essential amino acid) stimulates cell growth and prolong cell viability.⁵ The decrease of its level could be related to its involvement in the nucleic acid precursor metabolism to support cell proliferation, through the folate (THF) cycle and the methionine cycle.⁵

Conclusions

A powerful data independent workflow using SWATH® acquisition was demonstrated for comprehensive cell culture media analysis. It provided 1) the high resolution MS/MS based quantitation and confirmation of every detectable compound in media, 2) the full data processing software solution including library search and statistical analysis. Combined with the previously reported MRM workflow,¹ a versatile application solution for cell culture media analysis was offered serving different needs during biopharmaceuticals manufacturing.

References

1. Quantitative LC-MS Solution for Targeted Analysis of Cell Culture Media, SCIEX Technical Note RUO-MKT-02-9746-A.
2. SWATH® acquisition Variable Window Calculator - Excel tool. Download from <http://sciex.com/support/software-downloads>.
3. Improved Data Quality Using Variable Q1 Window Widths in SWATH® Acquisition, SCIEX Technical Note RUO-MKT-02-2879-B.
4. Method for culturing mammalian cells to improve recombinant protein production, Patent US20100221823
5. One-Carbon Metabolism in Health and Disease, Cell Metabolism, 25, 1, 27-42, 2017

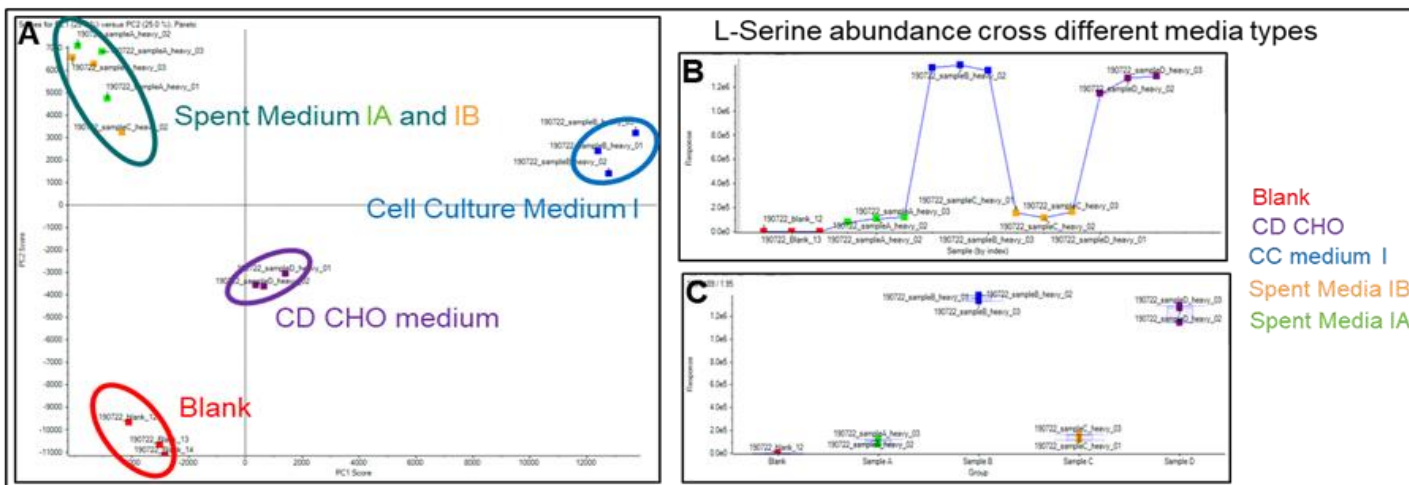


Figure 8: Statistical comparison between different media and L serine uptake during biopharmaceutical production. A) PCA plot shows a good separation between different media, B) quantitative trend of L-serine between different samples and C) Box plots of L-serine in different samples. Media are color coded listed on the left side of figure.

Table 3. List of components in the MRM library for the targeted workflow.

Cell Culture compound	Group		
L-Alanine	Amino acid	D-Pantothenic acid hemicalcium	Vitamin
Beta-Alanine	Amino acid	Folic acid	Vitamin
L-Arginine	Amino acid	L-Ascorbic acid	Vitamin
L-Aspartic Acid	Amino acid	L-Ascorbic acid 2-phosphate sesquimagnesium salt	Vitamin
L-Cystine	Amino acid	Niacinamide	Vitamin
L-Glutamic acid	Amino acid	Nicotinic acid (niacin)	Vitamin
Glycine	Amino acid	Pyridoxal hydrochloride	Vitamin
L-Histidine	Amino acid	(-)-Riboflavin	Vitamin
L-Isoleucine	Amino acid	ergocalciferol	Vitamin
L-Leucine	Amino acid	sodium ascorbate	Vitamin
L-Lysine	Amino acid	pyridoxine	Vitamin
L-Methionine	Amino acid	(-)-alpha-lipoic acid	Vitamin
L-Phenylalanine	Amino acid	Taurine	Others
L-Proline	Amino acid	2-isopropyl Malic acid	Others
L-Serine	Amino acid	2-oxovaleric acid	Others
L-Threonine	Amino acid	Citric acid	Others
L-Tyrosine	Amino acid	DL-A-Keto-B-methyl-n-valeric acid sodium	Others
L-Valine	Amino acid	DL-Isocitric acid trisodium hydrate	Others
L-Tryptophan	Amino acid	DL-P-Hydroxyphenyllactic acid	Others
gamma-Amino-n-butyric acid	Amino acid	Fumaric acid	Others
L-alpha-Amino-n-butyric acid	Amino acid	Lactic acid	Others
DL-beta-Aminoisobutyric acid	Amino acid	Pyruvic acid	Others
L-Carnosine	Amino acid	Succinic acid	Others
L-Citrulline	Amino acid	malic acid	Others
L-Cystathionine	Amino acid	D-gluconic acid sodium	Others
Ethanolamine	Amino acid	L-Asparagine	Others
L-Homocystine	Amino acid	ALA-GLN	Others
delta-Hydroxylysine	Amino acid	gly-gln monohydrate	Others
Hydroxy-L-proline	Amino acid	4-Aminobenzoic acid	Others
1-Methyl-L-histidine	Amino acid	Choline chloride	Others
3-Methyl-L-histidine	Amino acid	L-2-Aminoadipic acid	Others
L-Ornithine	Amino acid	L-Pipecolic acid	Others
L-Sarcosine	Amino acid	Uric acid	Others
L-Asparagine	Amino acid	Folinic acid calcium salt hydrate	Others
L-Glutamine	Amino acid	Penicillin G sodium	Others
L-Methionine sulfoxide	Amino acid	2-Aminoethanol (monoethanolamine)	Others
L-pyroglutamic acid	Amino acid	Ethylenediamine	Others
N-Acetyl-L-aspartic acid	Amino acid	Histamine free base	Others
N-Acetyl-L-cysteine	Amino acid	O-Phosphoethanolamine	Others
trans-4-hydroxy-L-Proline	Amino acid	Putrescine	Others
L-Norvaline	Amino acid	Phosphocholine chloride calcium salt tetrahydrate	Others
sarcosine	Amino acid	Glutathione oxidized	Others
L-Kynurenine	Amino acid	L-glycerophosphocholine	Others
linolenic acid	Fatty acid	D-Threonic acid lithium salt	Others
linoleic acid	Fatty acid		
oleic acid	Fatty acid		
stearic acid	Fatty acid		
palmitic acid	Fatty acid		
Adenine	Nucleobase		
Guanine	Nucleobase		
thymine	Nucleobase		
Uracil	Nucleobase		
hypoxanthine	Nucleobase		
Xanthine	Nucleobase		
2'-Deoxycytidine	Nucleoside		
Adenosine free base	Nucleoside		
Adenosine 5'-monophosphate	Nucleoside		
Cytidine	Nucleoside		
Cytidine 5'-monophosphate	Nucleoside		
Guanosine	Nucleoside		
Guanosine 5'-monophosphate disodium hydrate	Nucleoside		
Inosine	Nucleoside		
Thymidine	Nucleoside		
Uridine	Nucleoside		
Xanthosine dihydrate	Nucleoside		
Sucrose	Saccharides		
D-(+)-glucose	Saccharides		
D-(+)-glucosamine hydrochloride	Saccharides		
(-)-Tocopherol acetate	Vitamin		
Biotin	Vitamin		
Cyanocobalamin	Vitamin		

For Research Use Only. Not for use in Diagnostic Procedures. Trademarks and/or registered trademarks mentioned herein are the property of AB Sciex Pte. Ltd., or their respective owners, in the United States and/or certain other countries.

AB SCIEX™ is being used under license. Beckman Coulter® is being used under license. ICAT™ is a trademark of the University of Washington and is exclusively licensed to AB Sciex Pte. Ltd.

© 2019 DH Tech. Dev. Pte. Ltd. RUO-MKT-02-10242-A



Headquarters

500 Old Connecticut Path | Framingham, MA 01701 USA
Phone 508-383-7700
sciex.com

International Sales

For our office locations please call the division headquarters or refer to our website at sciex.com/offices