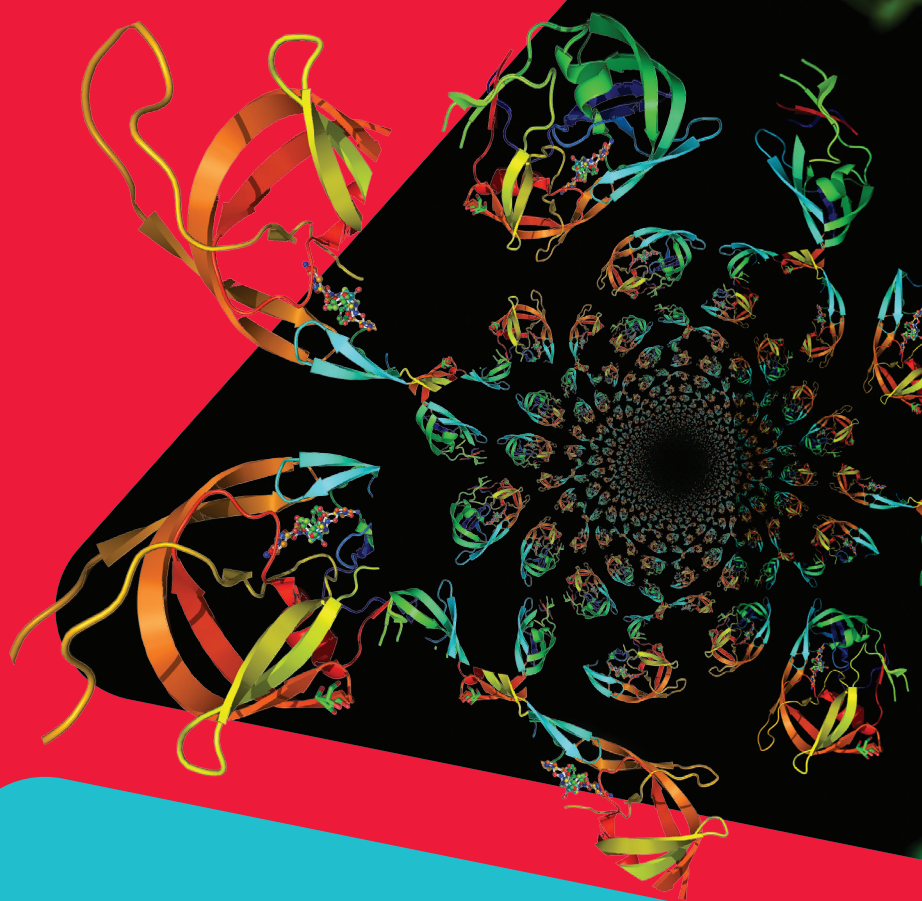
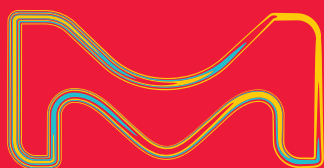


Sigma-Aldrich®

Lab Materials & Supplies

Tools for Mass Spectrometry

Proteomics and Metabolomics



The life science business of Merck operates as MilliporeSigma in the U.S. and Canada.

MERCK

As your partner in discovery, we are here to provide you with the products and information that you need to stay on the leading edge of mass spectrometry (MS) research in proteomics and metabolomics. Our MS portfolio covers workflows from sample preparation to downstream analysis, and allows you to probe biological systems and analyze prospective biomarkers in depth. We are committed to providing you with the broadest range of advanced products for MS workflow applications, backed by unrivaled scientific knowledge, customer service and technical service.

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Metabolomics Tools

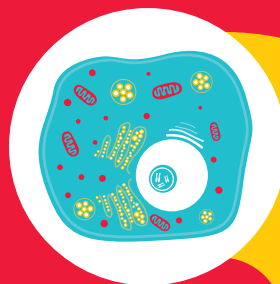
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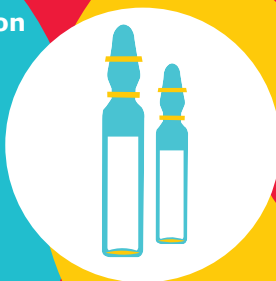
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Solutions Across the Entire MS Workflow

Sample Preparation



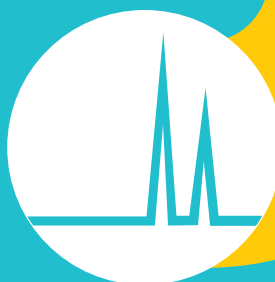
Digestion/Labeling



Standardization / Calibration



Chromatographic Separation



Detection & Analysis

Stable Isotope Labeled Products for Proteomics

Isotec® has the Products and Materials You Need for Stable Isotope Labeling in Cell Culture

ISOTEC Stable Isotopes offers a complete line of labeling materials to facilitate your quantitative proteomics experiments.

- 99% enriched amino acids – ^{13}C and/or ^{15}N Arginine and Lysine to improve overall quantification accuracy

Isotopically Labeled Amino Acids

Cat. No.	Description
608033	L-Arginine- $^{13}\text{C}_6$, $^{15}\text{N}_4$ hydrochloride
643440	L-Arginine- $^{13}\text{C}_6$ hydrochloride
600113	L-Arginine- $^{15}\text{N}_4$ hydrochloride
608041	L-Lysine- $^{13}\text{C}_6$, $^{15}\text{N}_2$ hydrochloride
643459	L-Lysine- $^{13}\text{C}_6$ hydrochloride
608092	L-Isoleucine- $^{13}\text{C}_6$, ^{15}N
605239	L-Leucine- $^{13}\text{C}_6$
608068	L-Leucine- $^{13}\text{C}_6$, ^{15}N
608106	L-Methionine- $^{13}\text{C}_5$, ^{15}N
299154	L-Methionine- <i>methyl</i> - ^{13}C , d_3
300616	L-Methionine- <i>methyl</i> - d_3
749915	L-Leucine- $^{13}\text{C}_6$, ^{15}N , 2, 3, 4, 4, 5, 5, 5- d_7 , 4- <i>methyl</i> - d_3
609021	L-Lysine- $^{15}\text{N}_2$ hydrochloride
643459	L-Lysine- $^{13}\text{C}_6$ hydrochloride
608041	L-Lysine- $^{13}\text{C}_6$, $^{15}\text{N}_2$ hydrochloride
749907	L-Lysine- $^{13}\text{C}_6$, $^{15}\text{N}_2$, 2, 3, 3, 4, 4, 5, 5, 6, 6- d_9 monohydrochloride
609242	L-Methionine- ^{15}N
749893	L-Methionine- $^{13}\text{C}_5$, ^{15}N , 2, 3, 3, 4, 4- d_5 , <i>methyl</i> - d_3
608114	Proline- $^{13}\text{C}_5$, ^{15}N
750018	L-Arginine- $^{13}\text{C}_6$, $^{15}\text{N}_4$, 2, 3, 3, 4, 4, 5, 5- d_7 hydrochloride
749915	L-Leucine- $^{13}\text{C}_6$, ^{15}N , 2, 3, 3, 4, 4, 5, 5, 5- d_7 -4- <i>methyl</i> - d_3
609021	L-Lysine- $^{15}\text{N}_2$ hydrochloride
643459	L-Lysine- $^{13}\text{C}_6$ hydrochloride
608041	L-Lysine- $^{13}\text{C}_6$, $^{15}\text{N}_2$ hydrochloride
749907	L-Lysine- $^{13}\text{C}_6$, $^{15}\text{N}_2$, 2, 3, 3, 4, 4, 5, 5, 6, 6- d_9 monohydrochloride
609242	L-Methionine- ^{15}N
749893	L-Methionine- $^{13}\text{C}_5$, ^{15}N , 2, 3, 3, 4, 4- d_5 - <i>methyl</i> - d_3
608017	L-Phenylalanine- $^{13}\text{C}_9$, ^{15}N
607770	L-Threonine- $^{13}\text{C}_5$, ^{15}N
492868	L-Tyrosine- $^{13}\text{C}_9$
600148	L-Valine- $^{13}\text{C}_5$, ^{15}N

Protected Amino Acids

Protected amino acids provide researchers with the ability to create their own unique peptide standards for MS-based applications. Using a synthetic peptide standard enables researchers to quantify and study the characteristics of proteins across a multitude of applications, including disease biomarkers.¹

ISOTEC Stable Isotopes offers a wide selection of both Fmoc and t-Boc protected labeled amino acids for the synthesis of peptide standards.

Cat. No.	Description
489905	Fmoc-Ala-OH- ^{15}N
605131	Fmoc-Ala-OH- $^{13}\text{C}_3$
667064	Fmoc-Ala-OH, $^{13}\text{C}_3$, ^{15}N
485888	Fmoc-Ala-OH-3, 3, 3- d_3
653659	Fmoc-Arg(Pbf)-OH- $^{13}\text{C}_6$, $^{15}\text{N}_4$
668753	Fmoc-Asn(Trt)-OH- $^{13}\text{C}_4$, $^{15}\text{N}_2$
683639	Fmoc-Asp(OtBu)-OH- $^{13}\text{C}_4$, $^{15}\text{N}_2$
663956	Fmoc-Gln(Trt)-OH- $^{13}\text{C}_5$, $^{15}\text{N}_2$
605182	Fmoc-Gly-OH-1- ^{13}C
578622	Fmoc-Ile-OH- ^{15}N
597228	Fmoc-Ile-OH- $^{13}\text{C}_6$, ^{15}N
485950	Fmoc-Leu-OH- ^{15}N
653632	Fmoc-Lys(Boc)-OH- $^{13}\text{C}_6$, $^{15}\text{N}_2$
651443	Fmoc-Phe-OH- $^{13}\text{C}_9$, ^{15}N
651451	Fmoc-Pro-OH- $^{13}\text{C}_5$, ^{15}N
609145	Fmoc-Ser(tBu)-OH- ^{15}N
658928	Fmoc-Ser(tBu)-OH- $^{13}\text{C}_3$, ^{15}N
658162	Fmoc-Thr(tBu)-OH- ^{15}N
658898	Fmoc-Tyr (t-Bu)-OH- $^{13}\text{C}_9$, ^{15}N
642886	Fmoc-Val-OH- $^{13}\text{C}_5$, ^{15}N
579890	Fmoc-Asn-OH- $^{15}\text{N}_2$
666009	Fmoc-Glu(OtBu)-OH- $^{13}\text{C}_5$, ^{15}N
489530	Fmoc-Glu(OtBu)-OH- $^{13}\text{C}_2$, ^{15}N
653640	Fmoc-Met-OH- $^{13}\text{C}_5$, ^{15}N
694274	Fmoc-Thr(tBu)-OH- $^{13}\text{C}_4$, ^{15}N
489913	Boc-Ala-OH- ^{15}N
603449	Boc-Ala-OH-2- ^{13}C , ^{15}N
596188	Boc-Ala-OH-2- ^{13}C , ^{15}N
588407	Boc-Glu-OBzl- $^{13}\text{C}_5$, ^{15}N
587699	Boc-Glu-OH- ^{15}N
489557	Boc-Gly-OH-2- ^{13}C , ^{15}N
609161	Boc-Lys(Z)-OH- α - ^{15}N
591092	Boc-Tyr-OH- ^{15}N
604976	Boc-Val-OH-1- ^{13}C

Reference

1. Ciccimaro, E. and Blair, I.A. Stable-isotope dilution LC-MS for quantitative biomarker analysis. *Bioanalysis*, 2(2), 311-341(2010).

For a complete listing, visit

[SigmaAldrich.com/protectedaa](https://www.sigmaaldrich.com/protectedaa)

Enzymatic Labeling with Water-¹⁸O

Trypsin-mediated incorporation of ¹⁸O remains an important technique for the exogenous isotopic enrichment of proteins for quantitative proteomics. Two ¹⁸O atoms are introduced into the carboxy terminus of protein fragments during proteolytic cleavage in heavy water. The quantification of protein samples is achieved by combining natural abundance ¹⁶O fragments and ¹⁸O labeled peptide fragments then subjecting the mixture to mass spectrometric analysis to determine the ratio of ¹⁶O/¹⁸O labeled peak pairs.¹⁻² ¹⁸O enzymatic labeling has gained popularity in the examination of differential protein expression in pharmacological and cancer research.³⁻⁴

Cat. No.	Description	Isotopic Purity
487090	Water-18O	99 atom % 18O
329878	Water-18O	97 atom % 18O

References

1. Johnson, K.L., and Muddiman, D.C., (2004) A method for calculating ¹⁶O/¹⁸O peptide ion ratios for the relative quantification of proteomes. *J. Am. Soc. Mass Spectrom.*, **15**, 437-445.
2. Fenselau, C. (2007) A review of quantitative methods for proteomic studies. *J. Chromatography B*, **855**, 14-20.
3. Wang, J., et al., (2007) Integration of ¹⁸O labeling and solution isoelectric focusing in a shotgun analysis of mitochondrial proteins. *J. Proteome Res.*, **6**, 4601-4607.
4. Lane, C.S., et al., (2007) Comparative cytochrome P450 proteomics in the livers of immunodeficient mice using ¹⁸O stable isotope labeling. *Mol Cell Proteomics.*, **6**, 953-962.

Chemical Labeling

Stable Isotope coded labels enable researchers to perform NMR and mass spectrometric-based proteomics studies in the absence of metabolic labeling. Isotope labeling occurs by site-specific incorporation of labeled tags at cysteine residues or the general labeling of amines and carboxylic groups in protein samples. These techniques are particularly useful for applications where metabolic labeling is impractical or undesirable.

ISOTEC® offers a wide variety of labeled reagents that can be used as chemical labels in peptide and protein studies.

Cat. No.	Description
531227	Acetaldehyde- ¹³ C ₂
487821	Acetic anhydride- ¹³ C ₄
633259	N-Acetoxy-d ₃ -succinimide
607517	Acetyl chloride-1- ¹³ C
485681	Benzoic acid-(phenyl- ¹³ C ₆)
366048	Benzyl chloride-d ₅
283835	Bromoacetic acid- ¹³ C ₂
488232	Bromobenzene- ¹³ C ₆
488534	Chlorobenzene- ¹³ C ₆
485500	Dimethyl sulfate- ¹³ C ₂
457833	Ethylene-d ₄ oxide
596388	Formaldehyde- ¹³ C, ₂ solution 20 wt. % in D ₂ O
607312	Guanidine- ¹³ C, ¹⁵ N ₃ hydrochloride
592668	Iodoacetamide- ¹⁵ N
277185	Iodomethane- ¹³ C
640492	2-Nitrobenzenesulfonylchloride- ¹³ C ₆
615692	Propionic anhydride-d ₁₀
578517	Succinic anhydride- ¹³ C ₄
299359	Urea- ¹³ C

For a complete list, visit
SigmaAldrich.com/chemtag

Seppro® Depletion Technology

Simplify Your Sample

- Depletion from a variety of sample types
- Spin-Column, LC, and 96-Well Plate
- Specific capture of target proteins
- Complete solution, reagents, and labware included

The Seppro depletion technology allows for the removal of several highly abundant proteins from a variety of biological samples.

The use of avian polyclonal IgY (Immunoglobulin Yolk) antibodies provides unique and advantageous features that allow highly-specific partitioning of protein mixtures. As a result, previously masked proteins become more accessible for investigation.

The Seppro platform, incorporating Supermix technology, represents the most complete human protein depletion system available, removing 14 of the most abundant proteins from human serum or plasma, as well as other high and medium abundance proteins. Additional products are available for the depletion of mouse and rat samples, as well as the industry's only depletion system for the removal of Rubisco from plant samples.

The new HT Seppro® IgY14 Plate is specifically designed for high throughput removal of 14 highly abundant proteins from human serum or plasma. This product is based on the same avian antibodies (IgY) immobilized on a resin and incorporated into a 96-well plate. plate. incorporated into a 96 well plate.

Human Depletion Media

Cat. No.	Format	Size	Sample Capacity	Proteins Depleted
SEP010	Seppro IgY 14 Spin Columns	500 µL resin bed volume	15–20 µL (pooled normal human plasma or serum)	HSA, IgG, Fibrinogen, Transferrin, IgA, IgM, Haptoglobin, α ₂ -Macroglobulin, α ₁ -Acid Glycoprotein, Apo A-I HDL, α ₁ -Antitrypsin, Apo A-II HDL, Complement C3, LDL (Apo B)
SEP020	Seppro IgY14 LC2	6.4 × 63.0 mm (2 ml resin bed volume)	40–50 µL (pooled normal human plasma or serum)	HSA, IgG, Fibrinogen, Transferrin, IgA, IgM, Haptoglobin, α ₂ -Macroglobulin, α ₁ -Acid Glycoprotein, Apo A-I HDL, α ₁ -Antitrypsin, Apo A-II HDL, Complement C3, LDL (Apo B)
SEP030	Seppro IgY14 LC5	12.7 × 39.5 mm (5 ml resin bed volume)	100 µL (pooled normal human plasma or serum)	HSA, IgG, Fibrinogen, Transferrin, IgA, IgM, Haptoglobin, α ₂ -Macroglobulin, α ₁ -Acid Glycoprotein, Apo A-I HDL, α ₁ -Antitrypsin, Apo A-II HDL, Complement C3, LDL (Apo B)
SEP040	Seppro IgY14 LC10	12.7 × 79.0 mm (10 ml resin bed volume)	200–250 µL (pooled normal human plasma or serum)	HSA, IgG, Fibrinogen, Transferrin, IgA, IgM, Haptoglobin, α ₂ -Macroglobulin, α ₁ -Acid Glycoprotein, Apo A-I HDL, α ₁ -Antitrypsin, Apo A-II HDL, Complement C3, LDL (Apo B)
S2453	HT Seppro IgY14 96 well plate	350 µL collection well volume	1.5-2.0 µL per well (pooled normal human plasma or serum)	HSA, IgG, Fibrinogen, Transferrin, IgA, IgM, Haptoglobin, α ₂ -Macroglobulin, α ₁ -Acid Glycoprotein, Apo A-I HDL, α ₁ -Antitrypsin, Apo A-II HDL, Complement C3, LDL (Apo B)

Isolate and identify your target protein at

[SigmaAldrich.com/seppro](https://www.sigmaaldrich.com/seppro)

Human Supermix Media

Cat. No.	Format	Size	Sample Capacity	Proteins Depleted
SEP050	Seppro Human Supermix LC2	6.4 × 63 mm (2 mL resin bed volume)	Flow Through volume from IgY14 LC5	Further partitions complex human plasma/serum samples
SEP060	Seppro Human Supermix LC5	12.7 × 39.5 mm (5 mL resin bed volume)	Flow Through volume from IgY14 LC10	Further partitions complex human plasma/serum samples.

Mouse Depletion Media

Cat. No.	Format	Size	Sample Capacity	Proteins Depleted
SEP110	Seppro Mouse Spin Columns	500 µL resin bed volume	15–20 µL (pooled normal mouse plasma or serum)	Mouse serum albumin, IgG, Transferrin, Fibrinogen, IgM, Haptoglobin, α1-Antitrypsin
SEP090	Seppro Mouse LC10	12.7 × 79.0 mm (10 mL resin bed volume)	200–250 µL (pooled normal mouse plasma or serum)	Mouse serum albumin, IgG, Transferrin, Fibrinogen, IgM, Haptoglobin, α1-Antitrypsin

Mouse Supermix Media

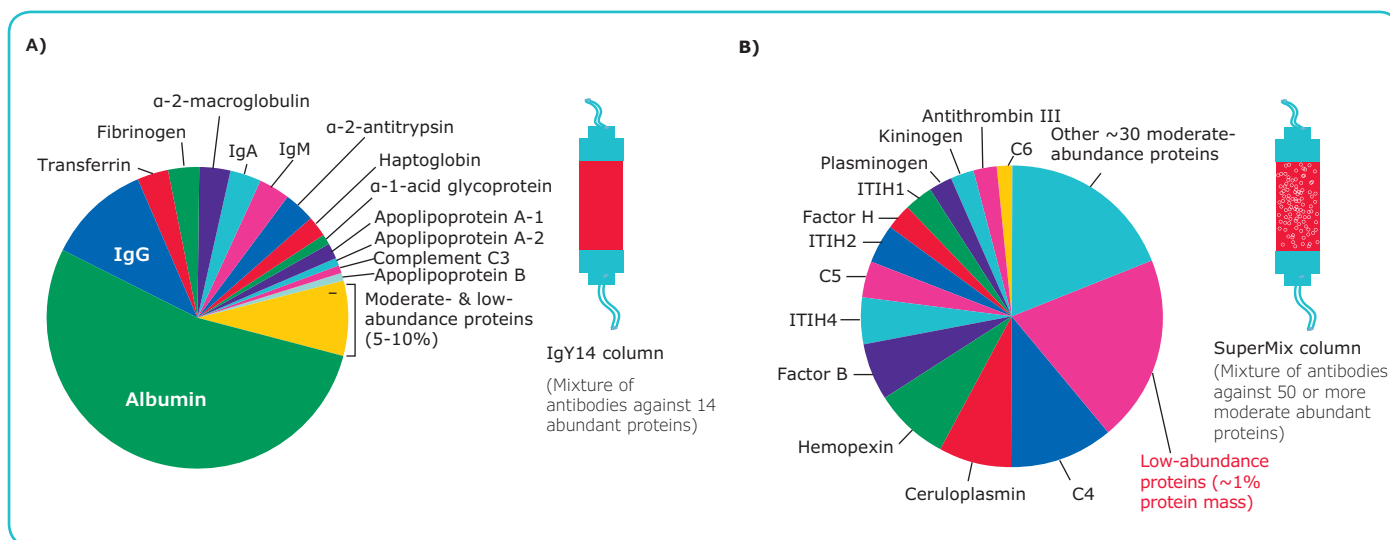
Cat. No.	Format	Size	Sample Capacity	Proteins Depleted
SEP100	Seppro Mouse Supermix LC5	12.7 × 39.5 mm (5 mL bed volume)	Flow Through volume from mouse LC10	Further partitions complex mouse plasma/serum samples.

Rat Depletion Media

Cat. No.	Format	Size	Sample Capacity	Proteins Depleted
SEP130	Seppro Rat Spin Columns	500 µL resin bed volume	15–20 µL (pooled normal rat plasma or serum)	Rat serum albumin, IgG, Fibrinogen, Transferrin, IgM, Haptoglobin, α1-Antitrypsin
SEP120	Seppro Rat LC10	12.7 × 79.0 mm (10 mL resin bed volume)	200–250 µL (pooled normal rat plasma or serum)	Rat serum albumin, IgG, Fibrinogen, Transferrin, IgM, Haptoglobin, α1-Antitrypsin

Rubisco Depletion Media

Cat. No.	Format	Size	Sample Capacity	Proteins Depleted
SEP070	Seppro Rubisco Spin Columns	500 µL resin bed volume	15–20 µL (pooled normal plant sample)	RuBisCO (Ribulose-1, 5-bisphosphate carboxylase/oxygenase)
SEP080	Seppro Rubisco LC2	6.4 × 63.0 mm (2 mL resin bed volume)	40–50 µL (pooled normal plant sample)	RuBisCO (Ribulose-1, 5-bisphosphate carboxylase/oxygenase)



Protein Extraction

Recombinant Protein Extraction

CellLytic™ Lysis Reagent

Cell lysis and high-yield protein extraction are important for quality recombinant protein purification. CellLytic formulations for bacterial, mammalian, yeast, or plant cell lysis result in higher protein yields than traditional physical disruption methods. CellLytic formulations are compatible with most affinity purification techniques, preserving protein function for downstream analysis.

Cat. No.	Product Description	Pack Size/Quantity
Bacterial Cell Lysis		
C8740	CellLytic B Cell Lysis Reagent, for bacterial cell lysis, 10 × concentrate	10 mL, 50 mL, 100 mL
B7310	CellLytic B Cell Lysis Reagent, for bacterial cell lysis, 10 × concentrate	50 mL, 250 mL
B7435	CellLytic B Cell Lysis Reagent, for bacterial cell lysis, standard strength	50 mL, 500 mL
CB0050	CellLytic B Plus Kit, for bacterial lysis	1 kit
C1990	CellLytic Express, for in-culture bacterial cell lysis	25 mL, 10 × 25 mL, 500 mL, 6 × 500 mL
C5491	CellLytic Express, 1 mL tablets, for direct lysis of bacterial cultures and for use in the HIS-Select® iLAP® column	25 each, 100 each
C5236	CellLytic Ib Inclusion Body Solubilization Reagent	25 mL, 100 mL
Mammalian Cell Lysis		
C2978	CellLytic M, Cell Lysis Reagent, suitable for mammalian cell lysis and protein solubilization	10 mL, 40 mL, 50 mL, 60 mL, 250 mL, 1L
CE0050	CellLytic MEM Protein Extraction Kit, for membrane proteins	1 kit
C3228	CellLytic MT Cell Lysis Reagent, for mammalian tissues	50 mL, 500 mL
NXTRACT	CellLytic NuCLEAR™ Extraction Kit, for mammalian tissue or cultured cells	1 kit
R0278	RIPA Buffer, for adherent and suspension cultured mammalian cells	50 mL, 500 mL
Plant Cell Lysis		
C2360	CellLytic P Cell Lysis Reagent, for plant lysis	50 mL, 250 mL
CELLYTPN1	CellLytic PN Isolation/Extraction Kit, for Plant Leaves	1 kit
Yeast Cell Lysis		
C4482	CellLytic Y Cell Lysis Reagent, for Yeast Cells	50 mL, 250 mL, 500 mL
CYP1	CellLytic Y Plus Kit, for Enzymatic Yeast Cell Lysis	1 kit

Native Protein Extraction

ProteoPrep® Lysis Reagent

ProteoPrep kits and individual extraction reagents allow for selective or total protein extracts from cellular samples. The protein extracts obtained with each component can be optimized to meet your individual needs. The reducing and alkylating reagents produce protein samples that exhibit improved focusing and decreased streaking in 2D gels. Enough of each component is provided to process multiple protein samples. For researchers who have optimized

an extraction protocol using one chaotropic extraction reagent, each kit reagent is also available as an individual product.

Cat. No.	Product Description	Pack Size/Quantity
PROTTOT	ProteoPrep Total Extraction Sample Kit	1 kit
PROTTWO	ProteoPrep Universal Extraction Kit	1 kit
PROTMEM	ProteoPrep Membrane Extraction Kit	1 kit
PROTDT	ProteoPrep Detergent Sample Kit	1 kit
PROTPR	ProteoPrep Protein Precipitation Kit	1 kit
PROTRA	ProteoPrep Reduction and Alkylation Kit	1 kit

Digestion and Enrichment

ProteoExtract® Kits

ProteoExtract® kits efficiently extract proteins from cellular samples. The proteins are then tryptically digested or enriched for peptide analysis. Enrichment and digestion kits are fully compatible with MALDI and ESI LC-MS workflows.

Cat. No.	Product Description	Pack Size/Quantity
650212	ProteoExtract® All-in-One Trypsin Digestion Kit	1 kit
72103	ProteoExtract® Glycopeptide Enrichment Kit	1 kit
539722	ProteoExtract® Phosphopeptide Enrichment TiO ₂ Kit	1 kit

Organelle Isolation

Kits and Reagents for Organelle Isolation

For enrichment of functional mitochondria, chloroplasts, nuclei, golgi, and other important organelles.

Cat. No.	Product Description	Pack Size/Quantity
MITOISO1	Mitochondria Isolation Kit, sufficient for 10–20 g (animal tissue), sufficient for 50 assays (2 mL), isolation of enriched mitochondrial fraction from animal tissues	1 kit
MITOISO2	Mitochondria Isolation Kit, sufficient for 50 applications (2–5 × 10 ⁷ cells), isolation of enriched mitochondrial fraction from cells	1 kit
NUC101	Nuclei Isolation Kit: Nuclei EZ Prep, sufficient for 25 nuclei preparations (~1–10 × 10 ⁷ cells/preparation)	1 kit
NUC201	Nuclei Isolation Kit: Nuclei PURE Prep, sufficient for 15 nuclei preparations (~1–10 × 10 ⁷ cells or 1 g of tissue per preparation)	1 kit
MITOISO3	Yeast Mitochondria Isolation Kit, sufficient for 40 applications (using 20 OD culture preparations), isolation of an enriched mitochondrial fraction of yeast cells	1 kit

To view all products in this range, visit [SigmaAldrich.com/organelle](https://www.sigmaaldrich.com/organelle)

For more information on Protein Extraction, visit [SigmaAldrich.com/extraction](https://www.sigmaaldrich.com/extraction)

MSSAFE: MS-SAFE Protease and Phosphatase Inhibitor Cocktail

- The MS-SAFE Protease and Phosphatase Inhibitor Cocktail (Cat. No. MSSAFE) is Sigma's new combination protease inhibitor and phosphatase inhibitor cocktail that is designed to be totally compatible with downstream mass spectrometry applications.
- MSSAFE is the only commercial inhibitor cocktail completely free of any inhibitors that can potentially modify proteins.
- MSSAFE is also fully compatible with IMAC, as MSSAFE completely omits any metal chelators.
- Available in 1VL and 5 × 1VL quantities

Light: Heavy Ratio

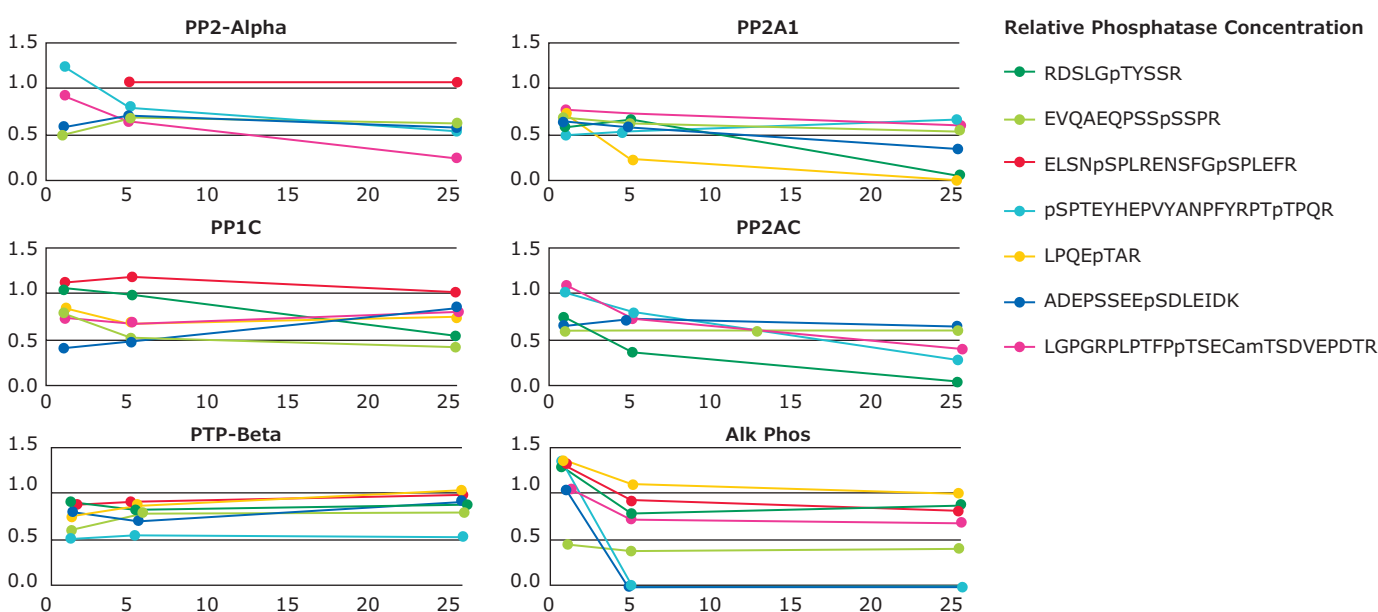


Figure 1. Demonstration of phosphatase specificity and inhibition with the Sigma PhosphoMix phosphopeptide standard against different phosphatases. A decrease in the light:heavy ratio of the phosphorylated peptide at higher phosphatase concentrations indicates that the phosphopeptide is a substrate for the phosphatase, and additionally that MSSAFE is properly inhibiting the phosphatase in question.






MSSAFE Inhibitor Components

Protease Inhibitor	Specific Inhibitory Target of Component
Bestatin hydrochloride	Aminopeptidases (e.g. leucine aminopeptidase, alanyl aminopeptidase)
Leupeptin	Serine and cysteine proteases (e.g. trypsin, plasmin, trypsinogen, urokinase, kallekrein)
Phosphoramidon sodium salt	Thermolysin, collagenase
Pepstatin A	Acid proteases (e.g. pepsin, renin, cathepsin D, many microbial aspartic proteases)
Elastatinal	Elastase
Aprotinin	Serine proteases (e.g. chymotrypsin, trypsin, elastase)
Nafamostat mesylate	Serine proteases, kallikrein

Protease Inhibitor	Specific Inhibitory Target of Component
Antipain	Serine/cysteine proteases, some trypsin-like serine proteases
Okadaic acid	Type 2A protein phosphatases
Sodium fluoride	Serine phosphatases, threonine phosphatases
Sodium orthovanadate	ATPases, protein tyrosine phosphatases, other phosphate-transferring enzymes
Bromotetramisole oxalate	L-isomers of alkaline phosphatases

For more information, visit SigmaAldrich.com/mssafe

Trypsin and Protease Selection Guide

	Trypsin Proteomics Grade	Trypsin Singles Proteomics Grade	Trypsin Spin Column Proteomics Grade	Trypsin Profile In-Gel Digest Kit	Protease Profiler Kit
					
Features and Benefits	<ul style="list-style-type: none"> • Reductively methylated to minimize autolytic activity • TPCK treated to quench chymotryptic activity • Highly purified 	<ul style="list-style-type: none"> • All the advantages of Proteomics Grade Trypsin in a convenient, single-use 1 µg package • Eliminates repetitive pipetting 	<ul style="list-style-type: none"> • 15 minute protein digestion • Eluted peptides are ready for MS analysis – no additional sample preparation required 	<ul style="list-style-type: none"> • Fast, efficient, complete in-gel tryptic digestion from PAGE to MALDI • Digested proteins are ready for MALDI-MS – no additional sample preparation required 	<ul style="list-style-type: none"> • Five proven proteases for detailed characterization of proteins of interest • Perform double enzymatic digests
Cat. No.	T6567	T7575	TT0010	PP0100	PP0500 (Individual components also available separately, see below)
Package Size	<ul style="list-style-type: none"> • 20 µg (sufficient to digest 400 µg – 2 mg of sample) • 5 × 20 µg (sufficient to digest 2–10 mg of sample) • 1 mg (sufficient to digest 20–200 mg of sample) 	96 × 1 µg (sufficient to digest 96 samples, 20–100 µg each)	10 columns (sufficient to digest 10 samples, 10–100 µg each)	1 kit (sufficient for up to 100 excised protein spots)	1 kit (sufficient to digest up to 5,900 µg of sample)
Components	Trypsin, Proteomics Grade	<ul style="list-style-type: none"> • Trypsin, Proteomics Grade (T6567) • Trypsin Solubilization Buffer • Enzyme Reaction Buffer • Biotech Grade Acetonitrile 	<ul style="list-style-type: none"> • Trypsin Spin Columns containing 75 mg of solid support in a 50% acidic glycerol suspension • Collection Tubes • Enzyme Reaction Buffer 	<ul style="list-style-type: none"> • Trypsin, Proteomics Grade (T6567) • Destaining Solution • Enzyme Reaction Buffer • Biotechnology Grade Acetonitrile • Trypsin Solubilization Reagent • Peptide Extraction Solution 	<ul style="list-style-type: none"> • Trypsin, Proteomics Grade (T6567) • Asp-N Protease (P3303) • Lys-C Protease (P3428) • Glu-C Protease (P6181) • Arg-C Protease (P6056) • Enzyme Solubilization Reagent • Enzyme Reaction Buffer
Suggested Sample Size	20–100 µg of sample per 1 µg of trypsin	20–100 µg of sample per 1 µg Trypsin Single	10–100 µg of sample per spin column	One excised protein spot or band	<ul style="list-style-type: none"> • 20–100 µg of sample per 1 µg of Trypsin • 50–200 µg of sample per 1 µg of Asp-N • 20–100 µg of sample per 1 µg of Lys-C • 20–100 µg of sample per 1 µg of Glu-C • 20–100 µg of sample per 1 µg of Arg-C
Digest Time	Overnight	Overnight	15 min	Overnight	Overnight
Digest Types	Solution or In-Gel	Solution or In-Gel	Solution	In-Gel	Solution or In-Gel
Enzyme Purity	>98%	95–98%	>98%	>98%	95–98%
Chymotryptic Activity	None Detected	None Detected	None Detected	None Detected	None Detected
Downstream Applications	HPLC-MS, MALDI-MS	HPLC-MS, MALDI-MS	HPLC-MS, MALDI-MS	HPLC-MS, MALDI-MS	HPLC-MS, MALDI- MS

Additional Proteases

Protease	Cat. No.	Package Size	Cleavage Specificity
Alpha-Lytic Protease (ALP)	A6362	20 µg, 5 × 20 µg	C-terminal to: Thr (T) Ala (A) Ser (S) Val (V)
Alpha-Lytic Protease (ALP) W190A Mutant	A6487	20 µg, 5 × 20 µg	C-terminal to: Met (M) Phe (F) Leu (L)
Endoproteinase Arg-C	P6056	1 VL	C-terminal to: Arg (R)
Endoproteinase Asp-N	P3303	1 VL	N-terminal to: Asp (D) Cysteic acid
Endoproteinase Glu-C	P6181	50 µg	C-terminal to: Glu (E) Asp (D)
Endoproteinase Lys-C	P3428	1 VL	C-terminal to: Lys (L)
Endoproteinase Pro-C	45167 E7656	1 mg 1 mg	C-terminal to: Pro (P)

For a complete listing of proteomics grade proteases, visit [SigmaAldrich.com/proteasefinder](https://www.sigmaaldrich.com/proteasefinder)

Introducing Solution Stable Trypsin

Our Advanced Proteomics Grade SOLu-Trypsin (EMS0004) is an exclusive, solution stable enzyme for mass spectrometry. Designed to be stable in solution when refrigerated, SOLu-Trypsin can be used immediately without preparation. Other forms of trypsin require thawing or reconstitution, and must be discarded if not used immediately. SOLu-Trypsin allows excess product to be saved for future use, thus eliminating unnecessary waste and cost. SOLu-Trypsin is formulated with a high-purity recombinant trypsin, free of chymotryptic activity, to ensure high fidelity digestion.

Key benefits of SOLu-Trypsin include:

- Ready to use - no preparation, such as reconstitution or thawing, is required
- Fits seamlessly into established workflow - no need to modify protocols
- Eliminates waste - remains stable in the refrigerator after use so there is no need to discard excess product

- Recombinant, porcine sequence - no chymotryptic activity
- Stable for short-term use at room temperature in an autosampler or on a liquid handling robot

For more information, visit [SigmaAldrich.com/solu-trypsin](https://www.sigmaaldrich.com/solu-trypsin)

SCIEX iChemistry™ Solutions

Tagging Chemistries For Quantitative Protein Mass Spectrometry

We have committed with SCIEX to cooperate in the global distribution of SCIEX's iChemistry™ Solutions portfolio of products. This partnership enables global access to unique MS Reagents for Quantitation across a variety of applications.

SCIEX iChemistry Solutions are MS-based tagging reagents designed to improve sensitivity, productivity, and data precision. The synergy of SCIEX with Sigma-Aldrich products allows users to dig deeper into biological systems, analyze prospective biomarkers, and deliver results.

iTRAQ® Reagents (Isotopic Tags for Relative and Absolute Quantification)

The iTRAQ Reagents are the first set of multiplexed, amine-specific, stable isotope reagents that can label all peptides in up to eight different biological samples, enabling simultaneous identification and quantitation, both relative and absolute, while retaining important post-translational modifications (PTMs).

Benefits

- Flexible – multiplex up to 8 different biological samples in a single experiment
- Simple workflow – no sample fractionation for reduced-complexity samples, such as affinity pull-downs
- Increased protein and proteome coverage – labels all peptides, including those with PTMs

Cat. No.	Description
4352135	iTRAQ Reagents Multiplex Kit
4352160	iTRAQ Reagents Methods Development Kit
4369561	iTRAQ Reagents 3-Assay Duplex Trial Kit
4370280	iTRAQ Reagents Application Kit – Plasma
4374321	iTRAQ Reagents Application Kit – Protein
4381664	iTRAQ Reagent – Multiplex Buffer Kit
4381662 (4390811)	iTRAQ Reagent – 8Plex One Assay Kit
4381663 (4390912)	iTRAQ Reagent – 8Plex Multiplex Kit
4390733 (4393528)	iTRAQ Reagent – 8Plex 25 U bulk pack
4390731 (4393529)	iTRAQ Reagent – 8Plex 50 U bulk pack
4466096	iTRAQ Reagent 25U Kit

mTRAQ® Reagents (Mass differential Tags for Relative and Absolute Quantification)

The mTRAQ Reagents are amine-specific, stable isotope-labeled reagents, available in duplex or triplex format, designed for high-confidence protein and peptide biomarker verification using multiple reaction monitoring (MRM) analysis.

Benefits

- Established chemistry – >95% labeling efficiency
- High selectivity – MRM-based relative quantitation of proteins/peptides/PTMs
- Supports reproducible quantitative MRM assays in biomarker verification
- Compatible with fractionation
- Easy transition from iTRAQ reagent-based discovery to verification
- Economical compared to stable isotope-labeled synthetic peptide approaches when larger numbers of peptides need to be monitored

Cat. No.	Description
4374771	mTRAQ Reagent 10 Assay Kit (10 assays of each reagent Δ0, Δ4, and Δ8)
4381664	iTRAQ Reagent-Multiplex Buffer Kit
4427698	mTRAQ Reagent Δ4 50 Unit Pack
4427700	mTRAQ Reagent Δ8 50 Unit Pack
4440015	mTRAQ Reagent Δ0 50 Unit Pack

Cleavable ICAT® Reagents (Isotope Coded Affinity Tag)

Cleavable ICAT reagents are a cysteine-specific, protein-based labeling strategy designed to compare two different sample states.

Benefits

- Provides more complete protein identification and quantification data than is possible with 2-D gels
- Simplifies the peptide pool for MS analysis
- Enables analysis of larger peptides

Cat. No.	Description
4339035	Cleavable ICAT Reagent – Methods Development Kit
4339036	Cleavable ICAT Reagent – 10 Assay Kit
4339038	Cleavable ICAT Reagent – Bulk Kit (10 Units)
4339039	Cleavable ICAT Reagent – Bulk Kit (100 Units)

CYP450 Protein Assay – (Cytochrome P450 Kit)

The CYP450 Protein Assay – Human Induction Kits use an LC/MS workflow to enable direct measurement of protein expression changes of individual CYP450 isoforms with high specificity, sensitivity, and accuracy.

These kits have been developed to employ MRM analysis to quantify the protein levels of seven key CYP450 protein isoforms for induction studies: 1A2, 2B6, 2C9, 2C19, 2E1, 3A4, and 3A5. Both four isoform (1A2, 2B6, 3A4, and 3A5) and seven isoform (1A2, 2B6, 2C9, 2C19, 2E1, 3A4, and 3A5) kits are available.

Benefits

- Allows for direct measurement of CYP450 induction at the protein level
- Specific, sensitive, and accurate – measures individual CYP450 isoforms
- Easy-to-use kits – method employs a common MRM workflow
- Gives equivalent results to traditional mRNA or activity assays

Cat. No.	Description
4445252	P450 Human Induction 100 Assay Kit
4445494	P450 Human Induction Starter Kit
4465863	CYP450 Peptide Standards Extended Panel Human Induction 100 Assay Kit
4466004	CYP450 Peptide Standards Extended Panel Human Induction Starter Kit

Amplifex™ Reagents

The Amplifex Reagents are designed for derivatization and mass spectrometry analysis of small molecule biomarkers that are otherwise very difficult to study by MS. Amplifex treatment forms positively charged derivatives of the target molecules with strongly improved ionization efficiency and fragmentation. This greatly lowers the limits of target detection and quantitation.

- The Amplifex Diene Reagent reacts with any molecule with a cis-diene group, such as Vitamin D3, Vitamin D2, and analogs of each
- The Amplifex Keto Reagent reacts with the carbonyl groups in keto-or aldehyde-containing species, such as ketosteroids

Benefits

- Greatly enhanced sensitivity and signal strength
- Substantially reduced sample sizes and analysis times
- Derivatization protocols that are simple (one-step) and fast (< 1 hr)
- Potential to prepare internal standards and to do multiplexing

Cat. No.	Description
5037804	Amplifex Diene Reagent Kit
4465962	Amplifex Keto Reagent Kit

For more information, visit SigmaAldrich.com/sciex

Universal Proteomics Standard (UPS)

Standardize Your Proteomics Research

We offer the Universal Proteomics Standard and the Proteomics Dynamic Range Standard as complex, well-defined, well-characterized reference standards for mass spectrometry. Both standards contain the same 48 human proteins ranging in molecular mass from 6,000 to 83,000 Daltons. Each constituent protein has been HPLC purified and AAA quantitated prior to formulation.

Universal Proteomics Standard

UPS1

Developed in collaboration with the Association of Biomolecular Resource Facilities (ABRF) Proteomics Standards Research Group (sPRG), the Universal Proteomics Standard contains 48 human proteins (5 pmoles of each) ranging in molecular mass from 6,000 to 83,000 Daltons.

Proteomics Dynamic Range Standard

UPS2

This standard is an enhancement of our original Universal Proteomics Standard (UPS1). The same complex mixture of 48 human proteins has been formulated into a dynamic range of concentration levels, ranging from 50 pmoles to 0.5 fmoles.

- Troubleshoot and optimize your analytical protocol
- Confirm system suitability before analyzing critical samples
- Normalize analytical results day-to-day or lab-to-lab
- Determine your limit of detection

Ordering Information

Cat. No.	Description	Package Size
UPS1	Universal Proteomics Standard Set*	1KT
UPS2	Proteomics Dynamic Range Standard Set*	1Set

* Each set contains one vial of Standard and one vial (20 µg) of Proteomics Grade Trypsin.

Criteria Used to Develop and Produce the Universal Proteomics Standards Line of Products

Criteria	Verified By
Proteins	More than 175 proteins were screened by SDS-PAGE and LCMS to be considered suitable for use in the standard
Minimal PTMs	LCMS of the intact proteins to verify homogeneity of the protein population
Diversity	Proteins are of wide-ranging molecular masses, hydrophobicities, isoelectric points, etc.
High Purity	Proteins are purified to be single-banded by SDS-PAGE and then further purified by HPLC
Accurate Quantitation	Amino Acid Analysis in triplicate

To learn more, visit [SigmaAldrich.com/ups](https://sigmaaldrich.com/ups)

UniProt Accession Number	UniProt Protein Name [Synonymx]	UPS1 Amount (fmol)	UPS2 Amount (fmol)
P00915	Carbonic anhydrase 1	5,000	50,000
P00918	Carbonic anhydrase 2	5,000	50,000
P01031	Complement C5[Complement C5a]	5,000	50,000
P69905	Hemoglobin alpha chain	5,000	50,000
P68871	Hemoglobin beta chain	5,000	50,000
P41159	Leptin	5,000	50,000
P02768	Serum Albumin	5,000	50,000
P62988	Ubiquitin	5,000	50,000
P04040	Catalase	5,000	5,000
P00167	Cytochrome bs	5,000	5,000
P01133	Epidermal Growth Factor	5,000	5,000
P02144	Myoglobin C	5,000	5,000
P15559	NAD(P)H dehydrogenase [quinone] 1 [DT Diaphorase] C	5,000	5,000
P62937	Peptidyl-prolyl cis-trans isomerase A [Cyclophilin A]	5,000	5,000
Q06830	Peroxiredoxin 1	5,000	5,000
P63165	Small ubiquitin-related modifier 1 [SUMO-1]	5,000	5,000
P00709	Alpha-lactalbumin	5,000	500
P06732	Creatine kinase M-type [CK-MM]	5,000	500
P12081	Histidyl-tRNA synthetase [Jo-1]	5,000	500
P61626	Lysozyme C	5,000	500
Q15843	Neddylin [Nedd8]	5,000	500
P02753	Retinol-binding protein	5,000	500
P16083	Ribosylidihydronicotinamide dehydrogenase [quinone] [Quinone oxidoreductase 2] [NQO2]	5,000	500
P63279	SUMO-conjugating enzyme Ubch9	5,000	500
P01008	Antithrombin-III	5,000	50
P61769	Beta-2-microglobulin	5,000	50
P55957	BH3 Interacting domain death agonist [BID]	5,000	50
O76070	Gamma-synuclein	5,000	50
P08263	Glutathione S-transferase A1 [GST A1-1]	5,000	50
P01344	Insulin-like growth factor II	5,000	50
P01127	Platelet-derived growth factor B chain	5,000	50
P10599	Thioredoxin	5,000	50
P01112	GTPase HRas	5,000	5
P99999	Gelsolin	5,000	5
P06396	Glutathione S-transferase P [GST]	5,000	5
P09211	GTPase HRas [Ras protein]	5,000	5
P01579	Interferon gamma (IFN-gamma)	5,000	5
P02787	Serotransferrin [Apotransferrin]	5,000	5
O00762	Ubiquitin-conjugating enzyme E2 C [UbcH10]	5,000	5
P51965	Ubiquitin-conjugating enzyme E2 E1 [UbcH6]	5,000	5
P08758	Annexin A 5	5,000	0.5
P02741	C-reactive protein	5,000	0.5
P05413	Fatty acid-binding protein	5,000	0.5
P10145	Interleukin-8	5,000	0.5
P02788	Lactotransferrin	5,000	0.5
P10636	Microtubule-associated protein tau [Tau protein]	5,000	0.5
P00441	Superoxide dismutase [Cu-Zn]	5,000	0.5
P01375	Tumor necrosis factor / Tumor necrosis factor, soluble form	5,000	0.5

MSRT1 – MS Retention Time Calibration Mix

Calibrate LC-MS the Right Way

The MS RT Calibration Mix (**Cat. No. MSRT1**) is an injection-ready mixture of 14 stable isotope-labeled peptides that is designed to assess LC-MS platform performance. MSRT1 enables users to translate retention times between platforms. These peptides span the normal elution profile of complex proteomic samples, with a readily visualized and well-separated series of peptide peaks.

Application

MSRT1 allows you to assess such characteristics as:

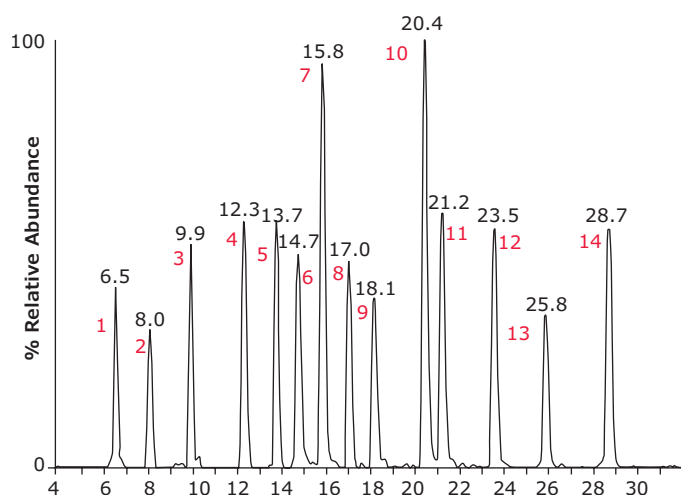
- LC resolution
- Peptide elution profile
- Retention time translation

Features and Benefits

- Well-separated series of peptide peaks in LC-MS analysis
- Easily visualized peaks of comparable intensity
- Injection-ready to save on sample preparation time

Peptide No.	Peptide Sequence (#)	Monoisotopic MW	Retention Factor
1	RGDSPASSP[K]	1008.5080	0.0
2	GLV[K]	423.2937	3.1
3	LGGNETQV[R]	982.5071	13.9
4	AEFAEVS[K]	887.4480	22.7
5	SGFSSVSVS[R]	1021.5068	30.3
6	ADEGISF[R]	903.4325	32.8
7	DISLSDY[K]	947.4691	39.0
8	LVNEVTEFA[K]	1156.6219	45.7
9	DQGGELLSL[R]	1096.5752	49.0
10	GLFIIDD[K]	927.5157	58.9
11	LGEYGFQNA[L]	1117.5517	63.1
12	YWGVASFLQ[K]	1205.6324	75.2
13	TDELFAQIEGLKEELAYL[R]	2176.1291	86.7
14	AVQQPDGLAVLGIFL[K]	1675.9752	100.0

(#) Amino acid in [brackets] denotes site of heavy label incorporation.



Peaks are labeled with Peptide # and retention time. LC-MS was performed on an Acquity-LCT platform with ~8 pmols injected onto a 1 mm I.D. BIOshell A160 Peptide ES C18 column (**Cat. No. 67099-U**) at 90 μ L/min, using a linear organic gradient modified with 0.1% formic acid.

For more information, visit

[SigmaAldrich.com/msrt](https://www.sigmaaldrich.com/msrt)

MSRT2 – SigmaProt Intact Protein LC-MS Standard

Proper Protein LC-MS Performance Verification from the Top Down

The SigmaProt Intact Protein LC-MS Standard (Cat. No. MSRT2) is an injection-ready mixture of 9 proteins that is designed to act as an LC-MS platform standard and to assess LC-MS platform performance. These 9 proteins cover a broad range of hydrophobicity and were chosen for ease of electrospray ionization (ESI). MSRT2 can be used as a performance standard for intact protein analysis, such as in top-down proteomics.

Application

MSRT2 allows you to assess such properties as:

- LC resolution
- Protein elution profile
- Electrospray source conditions
- Deconvolution parameters
- Comparison of LC gradients and columns
- Monitor column and system changes

Features and Benefits

- Well-separated and easily visualized series of protein peaks in LC-MS analysis
- Ready to use after reconstitution, to save on preparation time

Proteins in MSRT2

Protein	RT Order*	Modification Information	Average Mass (Da)*
Ribonuclease B	1	Man ₅ GlcNAc ₂	14,899
		Man ₆ GlcNAc ₂	15,062
		Man ₇ GlcNAc ₂	15,224
		Man ₈ GlcNAc ₂	15,386
		Man ₉ GlcNAc ₂	15,548
Insulin	2	—	5,808
Lysozyme	3	—	14,305
Transferrin	4	—	79,569
BSA	5	BSA	66,430
		BSA-Cysteinylated	66,549
		BSA-Glycated	66,592
Trypsin inhibitor	6	Mature Sequence	20,091
		C-Terminal Leu Truncation	19,978
β-lactoglobulin A	7	—	18,363
Carbonic anhydrase	8	N-Acetylserine	29,025
Lactic dehydrogenase	9	C Chain	36,160

*Confirmed using ESI mass spectrometry following C4 chromatography as described in the Figure “UV Chromatogram of MSRT2”.

UV Chromatogram of MSRT2

UV₂₁₅ chromatogram of MSRT2 using a Waters M-Class Acquity UPLC® and Xevo® G2S mass spectrometer.

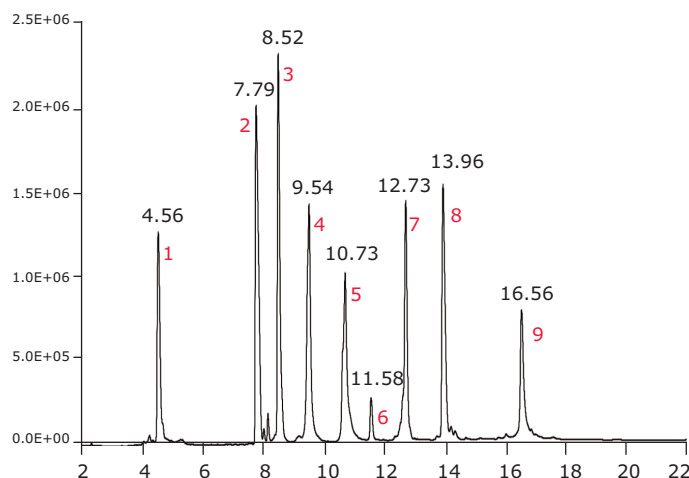
Column Supelco BIOshell® A400 Protein C4, 150 × 1.0 mm, 3.4 μm (Cat. No. 67045-U) at 65 °C.

Flow rate 70 μL/min with Solvent A = Water, 0.1% Trifluoroacetic Acid

Solvent B Acetonitrile, 0.1% Trifluoroacetic Acid.

Injection 1 μL for column load of 1 μg each protein.

Gradient	Time (Min)	%B
	0.0	20
	20.0	60
	20.5	80
	22.5	80
	23.0	20
	30.0	20



MS PhosphoMix Phosphopeptide Standards

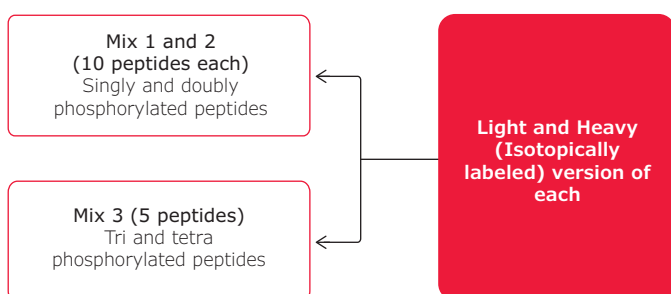
The MS PhosphoMix line of products allows for the testing of the strengths and weaknesses of phosphopeptide sample processing, mass spectrometry analysis, and instrument configurations. The mixes are produced from synthetic phosphopeptides with sequences derived from naturally occurring peptides as identified in HeLa cells.¹ Because the sequences are derived from mammalian cells, many natural phosphorylation motifs, such as those that present an abundance of proline, are represented.² Additionally, the phosphopeptide distribution in each mix has been chosen to present a broad range of characteristics, including ionizability, LC retention time, charge state, and isoelectric point. Finally, MS PhosphoMix-1, 2, and 3 were designed in a complementary fashion, as highlighted on the following page. For example, all three mixes contain peptides of the same sequence with different sites of phosphorylation (Figure 2).

Each of the three phosphopeptides mixes are available in their naturally occurring isotopic abundances (light) or as stable isotope enriched versions (heavy), making the set of products highly amenable to quantitative analyses, allowing users to compare recovery between workflows or techniques.

Features

- Naturally occurring peptide sequences
- Broad range of peptide characteristics
- Complementary produce designs
- Available in light and heavy versions

MS Phosphomix Product Design



Ordering Information

Cat. No.	Description	Amount/vial	Amount/peptide
MSP1L-1VL	MS PhosphoMix 1 Light	200 pmol	20 pmol/peptide
MSP1H-1VL	MS PhosphoMix 1 Heavy	200 pmol	20 pmol/peptide
MSP2H-1VL	MS PhosphoMix 2 Heavy	200 pmol	20 pmol/peptide
MSP2L-1VL	MS PhosphoMix 2 Light	200 pmol	20 pmol/peptide
MSP3H-1VL	MS PhosphoMix 3 Heavy	200 pmol	40 pmol/peptide
MSP3L-1VL	MS PhosphoMix 3 Light	200 pmol	40 pmol/peptide

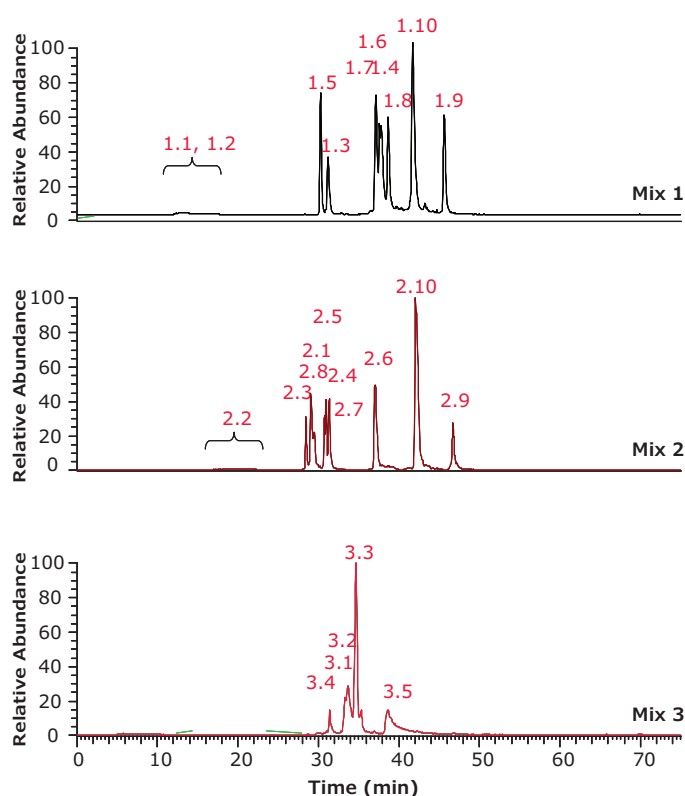


Figure 1. HPLC chromatograms of the 3 phosphopeptide mixtures using reverse phase (C18) stationary phase. The broad range of elution times as well as signal strength following electrospray (shown) or MALDI ionizations were taken into account during product design.

Reference:

1. J. V. Olsen, et al. *Cell*, 2006, **127**: 635-648
2. D. Schwatz, et al. *Nat. Biotech.* 2005, **23**(11): 1391-1398.

Peptide ¹	FASTA Abbreviation ²	Complementary Peptide	No. of Phosphates	Light MW (Monoisotopic)	Heavy MW (Monoisotopic)	Relative Signal Intensity ³	No. of Phosphates per amino acid		
							S	T	Y
PhosphoMix-1									
VLHSGpS[R]	1.1	—	1	834.37	844.38	Weak	1	—	—
RSpYpSRS[R]	1.2	2.2	2	1070.41	1080.41	Weak	1	—	1
RDSLGPpTYSS[R]	1.3	—	1	1220.52	1230.53	Medium	—	1	—
pTKLIpTQLRDA[K]	1.4	—	2	1445.70	1453.72	Strong	—	2	—
EVQAEQPSSpSSP[R]	1.5	—	1	1480.62	1490.63	Medium	1	—	—
ADEPpSSEESDLEID[K]	1.6	1.7, 2.6, 3.5	1	1742.68	1750.69	Strong	1	—	—
ADEPpSSEpSDLEID[K]	1.7	1.6, 2.6, 3.5	2	1822.64	1830.66	Medium	2	—	—
FEDEGAGFEESpSETGDYEE[K]	1.8	—	1	2333.84	2341.85	Strong	1	—	—
ELSpSPLRENSFGpSPLF[R]	1.9	2.9	2	2338.00	2348.01	Medium	2	—	—
SPTEYHEPpYANPFYRPTpTPQ[R]	1.10	—	2	2809.19	2819.20	Strong	—	1	1
PhosphoMix-2									
LPQEpTA[R]	2.1	—	1	893.40	903.40	Weak	—	1	—
RSYpSpSRS[R]	2.2	1.2	2	1070.41	1080.41	Weak	2	—	—
EpTQSPEQV[K]	2.3	—	1	1124.48	1132.49	Weak	—	1	—
VIEDNEpYTA[R]	2.4	—	1	1288.53	1298.54	Medium	—	—	1
pSRSpSSELNN[K]	2.5	—	2	1474.59	1482.60	Medium	2	—	—
ADEPpSSEpSDLEID[K]	2.6	1.6, 1.7, 3.5	1	1742.68	1750.69	Strong	1	—	—
HQYSDYDpYHSSpSE[K]	2.7	—	2	1904.63	1912.64	Medium	1	—	1
NTPpSQSHSpSIQHSPE[R]	2.8	—	2	2000.79	2010.80	Medium	2	—	—
ELSpSPLRENSFGpSPLF[R]	2.9	1.9	2	2338.00	2348.01	Medium	2	—	—
LGpGRPLTFpTSE(CAM) TSDVEPDT[R]	2.10	—	1	2708.22	2718.22	Strong	—	1	—
PhosphoMix-3									
SLpSpYpSP[V]ER	3.1	—	3	1276.42	1282.43	Weak	2	—	1
LQGpSGVpS[L]ApSK	3.2	—	3	1285.48	1292.49	Medium	3	—	—
PPpYpSRV[I]pTQR	3.3	—	3	1455.57	1462.59	Strong	1	1	1
pSRS[R]pSYpTPEpYR	3.4	—	4	1720.54	1730.55	Weak	2	1	1
ADEPpSpSEEpSDLE[I]DK	3.5	1.6, 1.7, 2.6	3	1902.61	1909.63	Medium	3	—	—

1. Amino acid in [brackets] denotes site of label incorporation for heavy mixes

2. A FASTA file with all of the phosphopeptide sequences in the PhosphoMix product line is available for free download on the product display page at sigma-aldrich.com

3. As determined using electrospray ionization following standard reverse phase chromatography

Figure 2. MS Phosphomix product design demonstrating the diversity of the three products. In addition to the broad characteristics, several peptides were chosen to present several phosphorylation patterns as shown in the complementary peptide series.

For more information, visit

SigmaAldrich.com/phosphomix

The SILu™ MAb Collection

Stable Isotope Labeled Monoclonal Antibody Standards

SILuMab is a Critical Tool for Assessment of the Pharmacokinetic Properties of Biotherapeutics

SILuMab is a highly purified stable isotope-labeled monoclonal antibody expressed in a proprietary Sigma-Aldrich CHO cell line grown in serum-free $^{13}\text{C}_6$ $^{15}\text{N}_4$ Arg / $^{13}\text{C}_6$ $^{15}\text{N}_2$ Lys enriched media.

SILuMab design is optimized to be used as an internal standard for quantitation of monoclonal antibodies as well as Fc-fusion therapeutics. Due to overlap with the common sequences in the Fc and light chain regions with candidate antibodies, SILuMab provides universal utility, thus eliminating the need for production of candidate-specific internal standards.

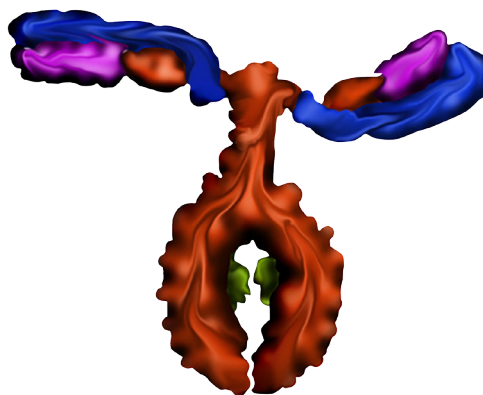
Quantitative LC/MS assays utilizing SILuMab offer advantages over traditional ELISA-based methods because of their superior specificity, sensitivity, and reduced matrix effects.

Vertical integration of SILuMab into the LC/MS analytical workflow yields universal stable isotope-labeled tryptic peptides that are utilized as internal standards.

As a full-length protein standard, SILuMab is superior to horizontally-integrated labeled peptide standards in that it reduces errors associated with fractionation, enrichment, and proteolysis.

SILuMab has been validated as an internal standard for quantitation of relevant biotherapeutics in a complex biological matrix by MRM-based LC-MS/MS.

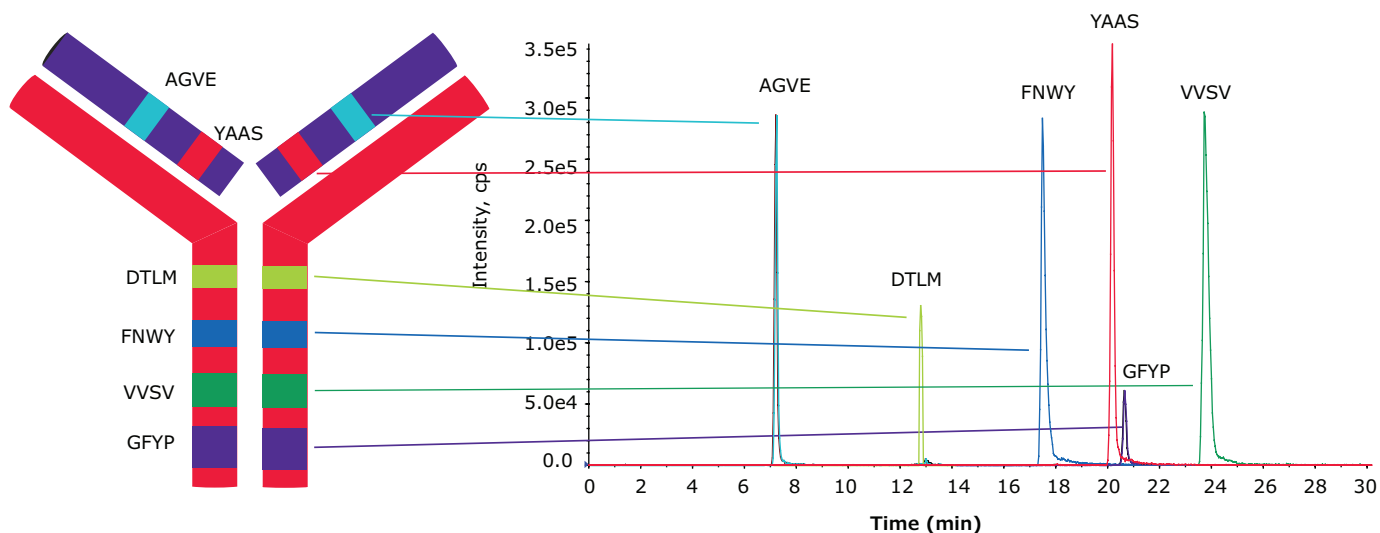
- SILuMab yielded reproducible, linear curves from 0.1 $\mu\text{g}/\text{mL}$ to 1,000 $\mu\text{g}/\text{mL}$ without enrichment or depletion.
- Excellent agreement was observed between multiple peptides derived from the same target.
- SILuMab has been highly characterized.
- Label incorporation was determined to be >98% by mass spectrometry.
- Sequence was confirmed by peptide mapping and intact mass analysis
- Purity has been determined to be $\geq 90\%$ by SDS-PAGE



We also offer SILuLite, which is the same recombinant monoclonal antibody sequence as SILuMab, in unlabeled form.

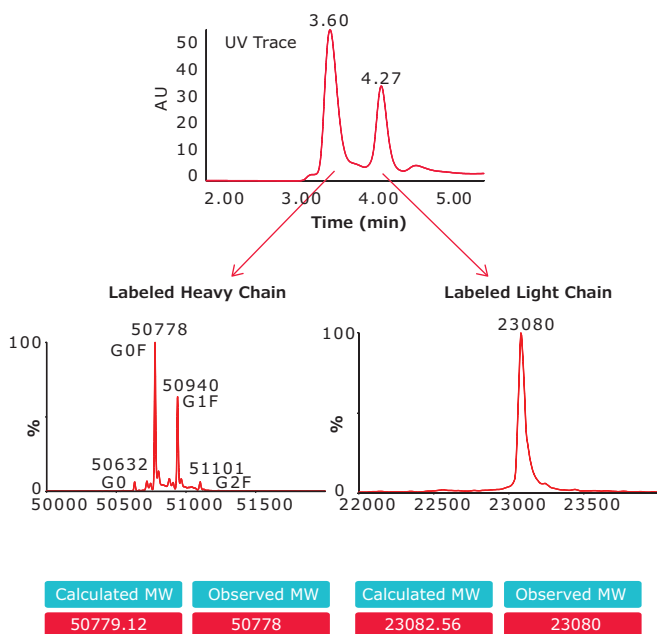
For more information or to place an order, call or contact your Sigma-Aldrich Sales Representative or visit [SigmaAldrich.com/silumab](https://www.sigmaaldrich.com/silumab)

Universal MRM Utility



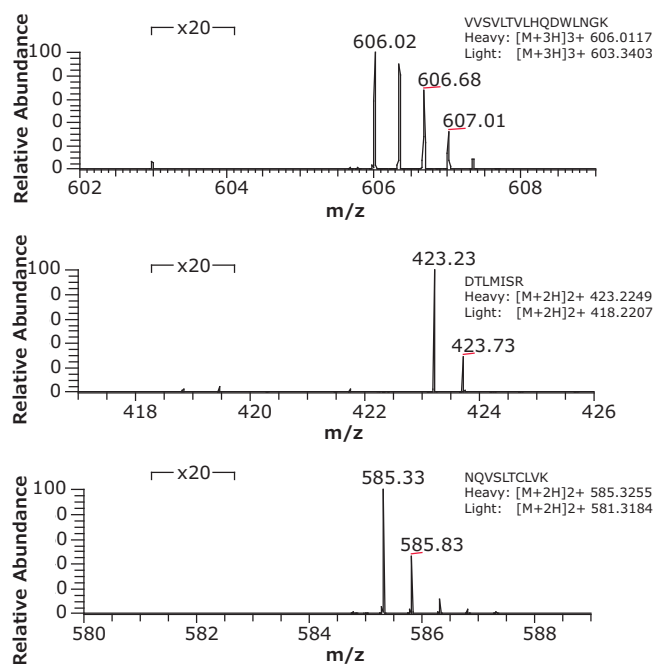
Extracted ion chromatogram (XIC) of representative peptides from the digested SILuMab. Using optimized overlap with common sequences in the Fc region of candidate antibodies, SILuMab provides universal utility, thus eliminating the need for production of candidate-specific internal standards.

Highly Characterized



SEC-UV and deconvoluted spectra resulting from intact mass analysis of the SILuMab standard. Calculations were based on the assumption that 99% label incorporation was achieved. Excellent agreement was seen between the calculated and observed molecular weight values.

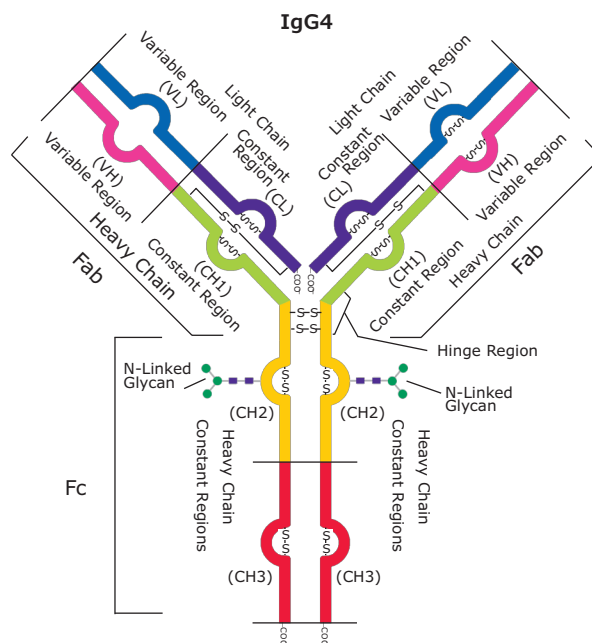
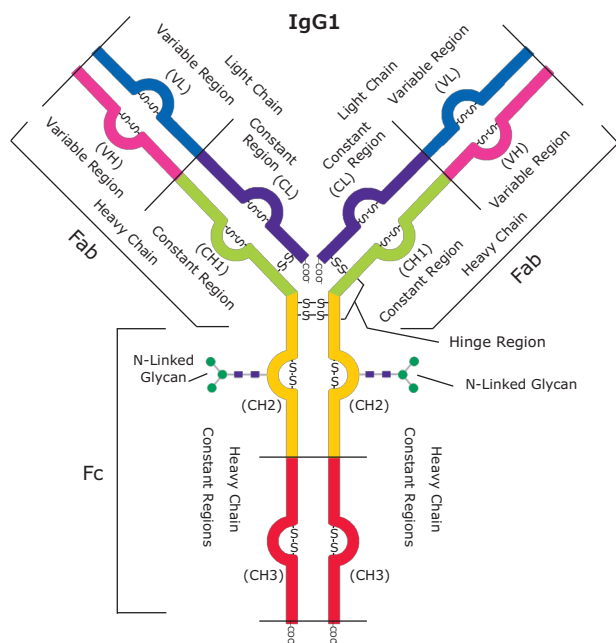
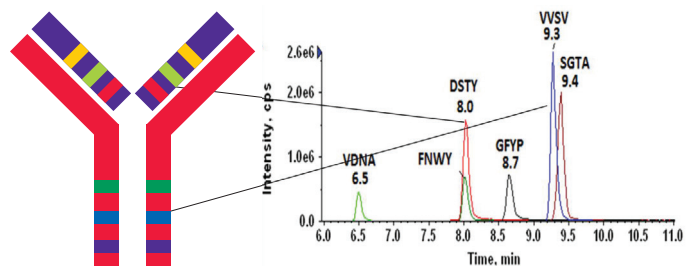
>98% Isotopically Labeled



Incorporation of stable labeled isotopes for representative SILuMab peptides. No unlabeled peptides were detectable. Therefore incorporation was considered to be >98%.

SILu™MAB K1 & K4

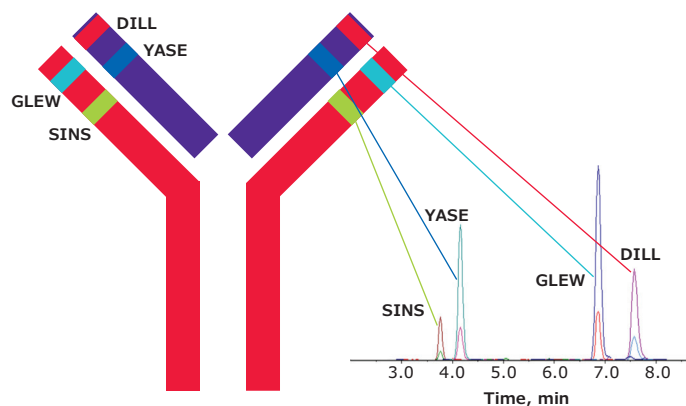
Stable-Isotope Labeled Universal Monoclonal Antibody (SILuMAB) K1 and K4 are the two new internal standards in the SILuMAB collection. These standards serve as a reference point in mass spectrometry analysis; by comparing the similar peaks of SILuMAB and a sample, the concentration of a target protein in the sample can be determined. SILuMAB K1 and SILuMAB K4 are highly purified IgG1 and IgG4 antibodies, respectively, that possess kappa peptide chains. These peptides create a stronger resemblance between SILuMAB and drugs in the pharmaceutical industry.



SILu™MAB Infliximab

SILu™MAB Infliximab (MSQC9) is a recombinant, stable isotope-labeled, monoclonal antibody which incorporates [¹³C₆, ¹⁵N₄]-Arginine and [¹³C₆, ¹⁵N₂]-Lysine. Expressed in CHO cells, it is designed to be used as an internal standard for the quantitative mass spectrometry analysis of Infliximab in human serum.

Infliximab is a chimeric monoclonal antibody biologic drug that works against tumor necrosis factor alpha (TNF-α) and is used to treat autoimmune diseases. Infliximab has been approved for the treatment of Crohn's disease, ulcerative colitis, psoriasis, psoriatic arthritis, ankylosing spondylitis, and rheumatoid arthritis. SILu™MAB Infliximab is for research use only and is not intended for diagnostic or therapeutic use.



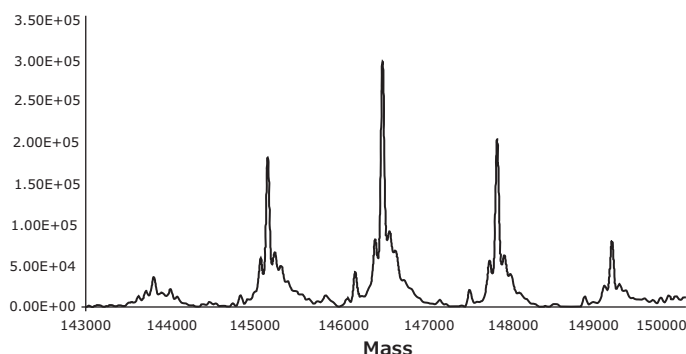
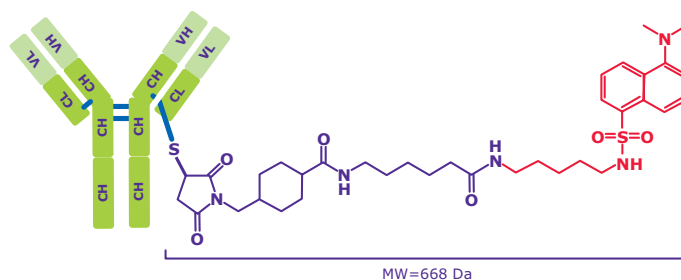
SigmaMAB ADC Mimic

The SigmaMAB Antibody Drug Conjugate (ADC) Mimic, the newest addition to the SILu™MAB collection, is a non-toxic drug mimic that is utilized as a standard for mass spectrometry and high performance liquid chromatography. The ADC Mimic is SigmaMAB (MSQC4), an IgG1 monoclonal antibody, conjugated to dansyl fluorophores via a LC-SMCC crosslinker. This dansyl-LC-SMCC attachment serves as the substitution for the toxic payload of a real ADC.

The ADC Mimic is affordable and allows for safer and more consistent testing, making it the optimal product over real ADCs. With the help of this non-toxic product, LC-MS processes can be developed in a safer environment.

For more information or to place an order for any of the SILu™Mab products, visit

SigmaAldrich.com/silumab



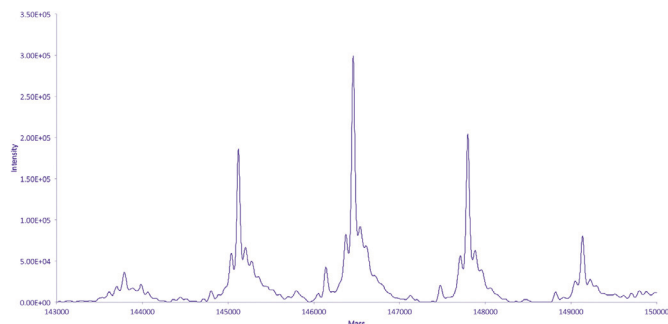
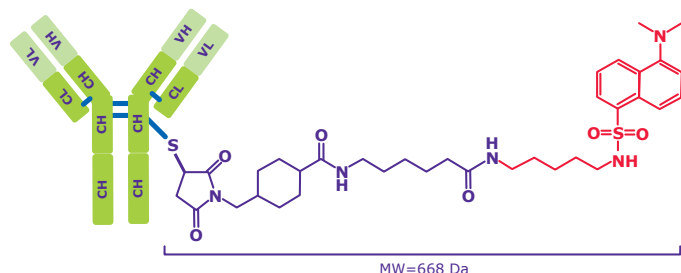
Description	Pk. Size	Cat. No.
SILu™Mab Stable-Isotope Labeled Universal Monoclonal Antibody Standard	100 µg	MSQC3
SigmaMAB SILu™Lite Monoclonal Antibody Standard (Unlabeled)	1 mg	MSQC4
SILu™Mab Stable Isotope-Labeled Monoclonal Antibody Glycan Standard	100 µg	MSQC5
SILu™MAB K1 Stable-Isotope Labeled Universal Monoclonal Antibody Standard	100 µg	MSQC6
SILu™MAB K4 Stable-Isotope Labeled Universal Monoclonal Antibody Standard	100 µg	MSQC7
SigmaMAB Antibody Drug Conjugate (ADC) Mimic	500 µg	MSQC8
SILu™MAB Infliximab Stable-Isotope Labeled Monoclonal Antibody Standard	100 µg	MSQC9
SILu™MAB Mouse Stable-Isotope Labeled Monoclonal Antibody Standard	100 µg	MSQC10
SILu™MAB Adalimumab Stable-Isotope Labeled Monoclonal Antibody Standard	100 µg	MSQC11

SILu™Prot and SILuLite

Full-Length Protein Standards for Mass Spectrometry

SILu™Prot is a collection of stable isotope-labeled (SIL) full-length proteins designed to be used as internal standards for quantitative proteomics. By adding a full-length SIL recombinant protein — which is equivalent to the native target protein — to the sample before the enzymatic digestion, the reproducibility and accuracy of protein quantification using mass spectrometry is optimized. Each SILu™Prot product serves as a standard for a different protein, which are all listed below.

In addition to the heavy labeled full-length protein standards, we have a line of unlabeled full-length protein standards, SILu™Lite. Each SILu™Prot product has its own corresponding SILu™Lite product. These full-length protein standards are intended to be used as the starting material for preparation of calibrators and controls in LC-MS application.



Product Name	Pk. Size	Cat. No.
SigmaMAb Antibody Drug Conjugate Mimic	500 µg	MSQC8

For more information or to place an order for SILu™Prot, visit [SigmaAldrich.com/siluprot](https://www.sigmaaldrich.com/siluprot)

MSQC1 – MS Qual/Quant QC Mix

Better than BSA for Standardizing MS Proteomic QC Runs

Experience the newest product available in proteomic analysis. MS Qual/Quant QC Mix (**Cat. No. MSQC1**) allows you to benchmark and monitor the daily performance of both qualitative and quantitative proteomic platforms.

Application

MS Qual/Quant QC Mix is an injection ready standard, optimized to assess platform characteristics including:

- Repeatability between runs
- System stability (drift, chromatography, signal intensity, sensitivity, etc.)
- Inter- and intra-platform and lab comparisons

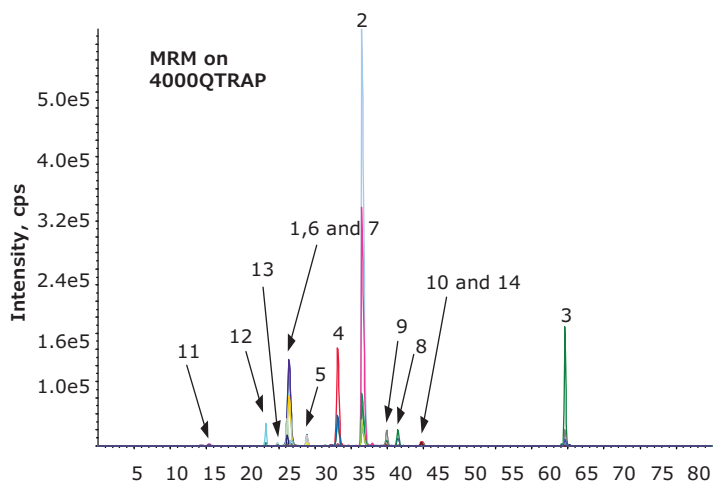
Features and Benefits

- **Complexity** – defined mixture gives confidence in your instruments analysis
- **Dynamic range** – generate data points that simulate the range of real world conditions
- **Predigested** – eliminate variability
- **Injection ready** – decrease prep time

Qualitative Benefits

Mixture of 6 tryptically digested human proteins

- Consistent, defined mixture
- 25-fold concentration range
- C18 purified



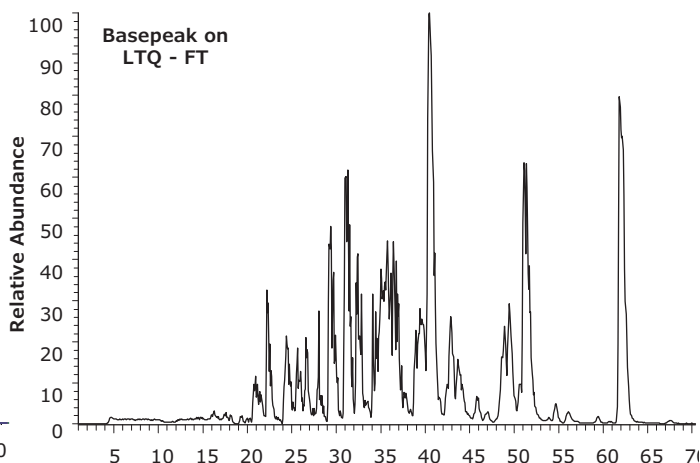
Quantitative Benefits

14 stable isotope labeled (SIL) peptides corresponding to 2–3 tryptic peptides for each protein have been incorporated into the protein digest

- Concentration range greater than three orders of magnitude
- MRM settings and transitions provided
- Light:Heavy ratios from 50:1 to 1:5






Protein	Peptide No.	Peptide Sequence*
Carbonic Anhydrase I	1	GGPFSDSY[R]
	2	VLDALQAI[K]
Carbonic Anhydrase II	3	AVQQPDGLAVLGIFL[K]
	4	SADFTNFDP[R]
NAD(P)H dehydrogenase	5	ALIVLAHSE[R]
	6	EGHLSDPDIAEQ[K]
C-reactive Protein	7	ESDTSYVSL[K]
	8	GYSIFSAT[K]
Peptidyl-Prolyl cis-trans isomerase A	9	FEDENFIL[K]
	10	VSFELFAD[K]
	11	TAENF[R]
Catalase	12	FSTVAGESGSADTV[R]
	13	NLSVEDAA[R]
	14	GAGAFGYFEVTHDIT[K]

*Amino acid in [brackets] denotes site of heavy label incorporation



For more information, visit SigmaAldrich.com/msqc

ProteoMass™ Selection Guide

	ProteoMass Peptide and Protein Calibration Kit	ProteoMass Peptide Calibration Kit	ProteoMass Protein Calibration Kit	ProteoMass vMALDI Calibration Kit	ProteoMass Guanidination Kit
Primary Application(s)	 Calibration and sensitivity testing of MALDI-MS instruments.	 Calibration and sensitivity testing of MALDI-MS instruments.	 Calibration and sensitivity testing of MALDI-MS instruments.	 Calibration and sensitivity testing of Thermo Scientific Exactive Series MS, LTQ XL and LTQ Hybrids with MALDI ion source	 Enhancement of MALDI signal strength and sequence coverage through efficient conversion of C-terminal lysine residues to homoarginine.
Mass Range	757 to 66,000 Daltons	757 to 3,500 Daltons	5,700 to 66,000 Daltons	523 to 3,675 Daltons	—
Cat. No.	MSCAL1	MSCAL2	MSCAL3	MSCAL4	MS0100
Package Size	1 kit	1 kit	1 kit	1 kit	1 kit
Components	<p>Calibrants</p> <ul style="list-style-type: none"> • Bradykinin Fragment 1–7, 10 nmols • Angiotensin II, 10 nmols • P14R, 10 nmols • ACTH Fragment 18–39, 10 nmols • Insulin Oxidized B chain, 10 nmols • Insulin, 10 nmols • Cytochrome c, 10 nmols • Apomyoglobin, 10 nmols • Aldolase, 10 nmols • Albumin, 10 nmols <p>Matrices</p> <ul style="list-style-type: none"> • α-cyano, 4 x 10 mg • sinapinic acid, 4 x 10 mg <p>Solvents</p> <ul style="list-style-type: none"> • 0.1% TFA – 30 mL • 1% TFA – 4 mL • Acetonitrile – 30 mL 	<p>Calibrants</p> <ul style="list-style-type: none"> • Bradykinin Fragment 1–7, 2 x 10 nmols • Angiotensin II, 2 x 10 nmols • P14R, 2 x 10 nmols • ACTH Fragment 18–39, 2 x 10 nmols • Insulin Oxidized B chain, 2 x 10 nmols <p>Matrix</p> <ul style="list-style-type: none"> • α-cyano, 8 x 10 mg <p>Solvents</p> <ul style="list-style-type: none"> • 0.1% TFA – 30 mL • 1% TFA – 4 mL • Acetonitrile – 30 mL 	<p>Calibrants</p> <ul style="list-style-type: none"> • Insulin, 2 x 10 nmols • Cytochrome c, 2 x 10 nmols • Apomyoglobin, 2 x 10 nmols • Aldolase, 2 x 10 nmols • Albumin, 2 x 10 nmols <p>Matrix</p> <ul style="list-style-type: none"> • sinapinic acid, 8 x 10 mg <p>Solvents</p> <ul style="list-style-type: none"> • 0.1% TFA – 30 mL • 1% TFA – 4 mL • Acetonitrile – 30 mL 	<p>Calibration Mix, Standard Range</p> <ul style="list-style-type: none"> • 5 vials, each containing MRFA, Bradykinin 1–7, Angiotensin 1, Neurotensin, Renin Substrate, and Bradykinin <p>Calibration Mix, High Range</p> <ul style="list-style-type: none"> • 5 vials, each containing MRFA, Bradykinin, ACTH 1–16, Melittin, and ACTH 7–38 <p>Sensitivity Standard</p> <ul style="list-style-type: none"> • Angiotensin II – 2 x 500 pmols <p>Matrix</p> <ul style="list-style-type: none"> • α-cyano, 5 x 5 mg <p>Solvents</p> <ul style="list-style-type: none"> • 1% TFA – 4 mL • Acetonitrile – 30 mL • Ethanol, 200 Proof, Molecular Biology Grade – 10 mL 	<ul style="list-style-type: none"> • O-Methylisourea hemisulfate • Base reagent • Stop Solution • Control Peptide

For more information, visit
SigmaAldrich.com/mscal

MALDI Matrices Selection Table

Matrix-assisted laser desorption/ionization (MALDI) has expanded MS into the analysis of high molecular mass, non-volatile, and thermally labile compounds, such as intact proteins and oligonucleotides. Moreover, it has become an important technique in proteomics research.¹⁻³ Further significant applications of MALDI-MS include the analysis of polymers, glycans, lipids, and metabolites.

A typical MALDI matrix substance is an aromatic acid with a chromophore that absorbs strongly at the wavelength of the incident laser. The MALDI technique generally involves mixing the sample with a matrix substance, followed by crystallization by different techniques on the MALDI sample plate. The crystallized sample-matrix mixture is irradiated by laser light, usually UV. As the matrix absorbs the light energy, it vaporizes into the gas phase, resulting in an indirect ionization of the sample molecules.⁴⁻⁶

Choosing a suitable matrix of high quality is the key to the success of a MALDI-MS experiment. Organic impurities can lead to extraneous peaks, especially in the low mass range. Trace levels of ions, especially Na⁺ and K⁺, form adducts with sample molecules. These adducts differ in mass according to the number of positive ions and complicate the MS spectrum. Since the matrix substance is generally applied in large excess to the sample, a very high purity is even more crucial.

The MALDI Matrices Selection Table below facilitates choosing the appropriate matrix for the use in proteomics and metabolomics.

Features and Benefits

- High chemical purity
- Low trace metal content to minimize adduct formation and simplify the resulting MS spectrum
- Ultra pure grades of the most popular matrix substances with extremely strict specifications concerning purity, trace metal content, appearance, and solubility

References

1. Karas, M., *et al.*, Matrix-assisted ultraviolet laser desorption of non-volatile compounds. *Int. J. Mass Spectrom. Ion Proc.*, **78**, 53-68 (1987).
2. Hillenkamp, F., and Peter-Katalinic, J. (eds.), *MALDI MS. A Practical Guide to Instrumentation, Methods and Applications*, Wiley-VCH (2007).
3. Aebersold, R., and Mann, M., Mass spectrometry-based proteomics. *Nature*, **422**, 198-207 (2003).
4. Dreisewerd, K., The desorption process in MALDI. *Chem. Rev.*, **103**, 395-425 (2003).
5. Karas, M., and Krüger, R., Ion formation in MALDI. *Chem. Rev.*, **103**, 427-439 (2003).
6. Knochenmuss, R., and Zenobi, R., MALDI ionization: The role of in-plume processes. *Chem. Rev.*, **103**, 441-452 (2003).

Cat. No.	Description	Purity	Abbreviation	Proteins	Peptides	Glycans	Oligonucleotides	Polymers	Lipids	Other Analytes	Note	Pack Sizes
92817	9-Aminoacridine	≥99.5%	9-AA							• Metabolites		1 g
89063	4-Bromo- α -cyanocinnamic acid	≥95%	BrCCA		•				•	Phosphopeptides, phospholipids, chlorinated lipids, drugs, fragile analytes, ionic liquids (quantification), ME-SIMS, CID-MS/MS		100 mg
68914	4-Bromo- α -cyanocinnamic acid - 4-Chloro- α -cyanocinnamic acid mixture	≥95%	BrCCA:CICCA		•				•	Phosphopeptides, phospholipids, chlorinated lipids, drugs, fragile analytes, ionic liquids (quantification), ME-SIMS, CID-MS/MS		100 mg
05851	4-Aminoquinaldine	≥99.0%					•			amino acids		1 g
56229	9-Nitroanthracene	≥98.5%	9-NA					•		fullerenes, humic acids		100 mg 1 g
69028	4-Phenyl- α -cyanocinnamamide	≥98.5%							•	MALDI imaging		100 mg
76884	Anthranilamide	≥99.0%		•	•	•						1 g
78246	Curcumin	≥99.5%							•	pharmaceuticals, drugs, MALDI imaging		100 mg
83788	(2E)-3-(9-Anthryl)-2-cyanoacrylic acid	≥97.0%								low molecular weight compounds		100 mg
87884	<i>trans</i> -2-[3-(4-tert-Butylphenyl)-2-methyl-2-propenylidene]malononitrile	≥99.0%				•		•		Gold nanoparticles, fullerenes, organometallics, macrocycles		250 mg 1 g

Cat. No.	Description	Purity	Abbreviation	Proteins	Peptides	Glycans	Oligonucleotides	Polymers	Lipids	Other Analytes	Note	Pack Sizes
94477	(E)-2-Cyano-3-(2-naphthyl) acrylic acid	≥98.0%								low molecular weight compounds		100 mg
55841	4-Bromo- α -cyanocinnamic acid - α -Cyano-2,4-difluorocinnamic acid mixture	≥95%	BrCCA:DiFCCA		•				•	Phosphopeptides, phospholipids, chlorinated lipids, drugs, fragile analytes, ionic liquids (quantification), ME-SIMS, CID-MS/MS		100 mg
60018	Caffeic acid	≥99.0%		•	•							1 g, 5 g
94141	4-Chloro- α -cyanocinnamic acid	≥95%	CICCA		•				•	Phosphopeptides, phospholipids, chlorinated lipids, drugs, fragile analytes, ionic liquids (quantification), ME-SIMS, CID-MS/MS		100 mg
39379	4-Chloro- α -cyanocinnamic acid - α -Cyano-2,4-difluorocinnamic acid mixture	≥95%	CICCA:DiFCCA		•				•			100 mg
77646	α -Cyano-2, 4-difluorocinnamic acid	≥95%	DiFCCA		•					Phosphopeptides, phospholipids, chlorinated lipids, drugs, fragile analytes, ionic liquids (quantification), ME-SIMS, CID-MS/MS		100 mg
77081	α -Cyano-4-fluorocinnamic acid	≥95%	FCCA		•					Phosphopeptides, phospholipids, chlorinated lipids, drugs, fragile analytes, ionic liquids (quantification), ME-SIMS, CID-MS/MS		100 mg
70990	α -Cyano-4-hydroxycinnamic acid	≥99.0%	CHCA	•	•	•						250 mg, 1 g
39468	α -Cyano-4-hydroxycinnamic acid	≥99.5%, Ultra pure	CHCA	•	•	•						10x10 mg
03841	α -Cyano-4-hydroxycinnamic acid - α -Cyano-2, 4-difluorocinnamic acid - α -Cyano-2, 3, 4, 5, 6-pentafluorocinnamic acid mixture	≥95%	CHCA:DiFCCA: PentaFCCA		•					Phosphopeptides, phospholipids, chlorinated lipids, drugs, fragile analytes, ionic liquids (quantification), ME-SIMS, CID-MS/MS		100 mg
38419	α -Cyano-2, 3, 4, 5, 6-pentafluorocinnamic acid	≥95%	PentawFCCA		•					Phosphopeptides, phospholipids, chlorinated lipids, drugs, fragile analytes, ionic liquids (quantification), ME-SIMS, CID-MS/MS		100 mg
56451	1,5-Diamino naphthalene	≥99.0%	1,5-DAN		•		•			In-Source-Decay		250 mg
37468	2',6'-Dihydroxy acetophenone	≥99.5%	2,6-DHAP	•	•	•			•			1 g, 5 g
85707	2, 5-Dihydroxybenzoic acid	≥99.0%	DHB	•	•	•			•	Organic molecules		10 mg, 250 mg, 1 g
39319	2, 5-Dihydroxybenzoic acid	≥99.5%, Ultra pure	DHB	•	•	•			•	Organic molecules		10x10 mg
46278	trans-Ferulic acid	≥99.0%	FA	•	•							1 g, 5 g
54793	2-(4-Hydroxy phenylazo) benzoic acid	≥99.5%	HABA	•	•	•			•			1 g, 5 g
56197	3-Hydroxypicolinic acid	≥99.0%	3-HPA				•			Oligosaccharides		250 mg, 1 g
73148	3-Nitrobenzyl alcohol	≥99.5%										5 g
80362	3-Nitrobenzonitrile	≥99.0%	3-NBN							Tissues via MAIV		1 g
84228	Salicylamide	≥99.0%					•					1 g
85429	Sinapic acid	≥99.0%	SA	•	•					Dendrimers, Fullerenes		1 g, 5 g
49508	Sinapic acid	≥99.5%	SA	•	•					Dendrimers, Fullerenes		10 x 10 mg
50862	Super-DHB BioReagent		Super-DHB	•	•	•						10 x 10 mg, 1 g, 5 g
91928	2', 4', 6'-Trihydroxy acetophenone monohydrate	≥99.5%	THAP	•	•	•	•					1 g, 5 g

Supel™-Select Polymeric SPE Products

Key Features and Benefits

- Hydrophilic-modified styrene resin extracts and recovers a broad range of analytes (polar to nonpolar, acidic to basic) using a single sorbent
- Generic methodology saves time, money, and headaches during method development
- Greater capacity allows for smaller bed weights = smaller elution volumes = time savings in sample processing
- Resistant to over-drying allowing for more robust methodology

Versatile and Simple Sample Cleanup by SPE

Supel-Select SPE phases are ideal for the solid phase extraction (SPE) of a broad range of compounds from aqueous samples. While reversed-phase interactions dominate retention on the Supel-Select HLB, and the retention mechanisms of the Supel-Select SAX and SCX are predominately based on ion-exchange, the hydrophilic modifications of the styrene-based polymer backbone allow for retention and recovery of more polar compounds.

Supel-Select Properties

HLB Phase Chemistry	Hydrophilic-modified styrene polymer
SAX Phase Chemistry	Quaternary amine-functionalized hydrophilic-modified styrene polymer
SCX Phase Chemistry	Sulfonic acid-functionalized hydrophilic modified styrene polymer
Suitable for MS Detection?	Yes
pH Compatibility	0 – 14
Particle Size	50 – 70 µm
Surface Area	160 – 420 m ² /g
Pore Volume	0.8 – 1.2 mL/g
Pore Size	80 – 200 Å

Description	Qty.	Cat. No.
Supel-Select HLB 96-well SPE		
10 mg/ well	1	Inquire
30 mg /well	1	575661-U
60 mg/ well	1	575662-U
Supel-Select SAX 96-well SPE		
10 mg/well	1	Inquire
30 mg/well	1	575660-U
60 mg/well	1	575663-U
Supel-Select SCX 96-well SPE		
10 mg/well	1	Inquire
30 mg/well	1	575664-U
60 mg/well	1	575665-U
Supel-Select HLB SPE		
30 mg/1 mL	100	54181-U
60 mg/3 mL	50	54182-U

LC-MS Analysis of Illicit Bath Salts in Urine on Ascentis Express HILIC with and without Supel-Select SCX SPE Cleanup

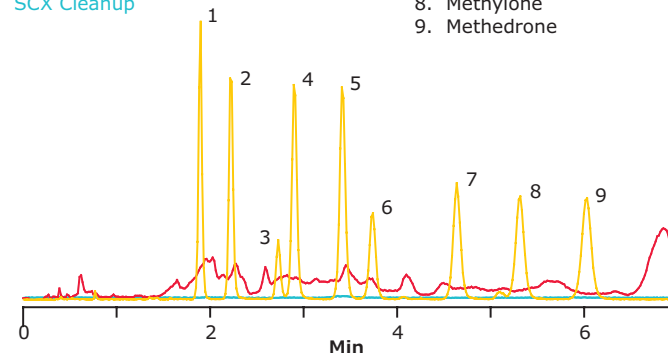
SPE tube	Supel-Select SCX, 30 mg/1 mL (54240-U)
column	Ascentis Express HILIC, 10 cm x 2.1 mm I.D., 2.7 µm (53939-U)
mobile phase	(A) 5 mM ammonium formate acetonitrile; (B) 5 mM ammonium formate water; (98:2, A:B)
flow rate	0.6 mL/min
pressure	127 bar
column temp	35 °C
detector	MS, ESI+, 100-1000 m/z
injection	1 µL
sample	200 ng/mL in acetonitrile (standards from Cerilliant)

Spiked Urine Sample Monitored Ions After Supel-Select SCX Cleanup

Diluted Spiked Urine Monitored Ions Without SPE Cleanup

Urine Blank Monitored Ions After Supel-Select SCX Cleanup

1. 3,4-Methylenedioxypropylrovalerone (MDPV)
2. Buphedrone
3. 3-Fluoromethcathinone
4. Butylone
5. Ethylone
6. 4-Fluoromethcathinone
7. Mephedrone
8. Methyone
9. Methedrone



Description	Qty.	Cat. No.
200 mg/6 mL	30	54183-U
500 mg/12 mL	20	54184-U
1 g/20 mL	20	54186-U
Supel-Select SAX SPE		
30 mg/1 mL	100	54231-U
60 mg/3 mL	50	54233-U
200 mg/6 mL	30	54235-U
500 mg/12 mL	20	54236-U
1 g/20 mL	20	54237-U
Supel-Select SCX SPE		
30 mg/1 mL	100	54240-U
60 mg/3 mL	50	54241-U
200 mg/6 mL	30	54242-U
500 mg/12 mL	20	54243-U
1 g/20 mL	20	54245-U

For more information, visit [SigmaAldrich.com/supel-select](https://www.sigmaaldrich.com/supel-select)

HybridSPE®-Phospholipid Products for Consistent LC-MS Ionization

Key Features and Benefits

- Maximize sensitivity by minimizing ion-suppression
- 100% removal of phospholipids and precipitated proteins
- 2-3 step generic procedure
- Ideal for high-throughput pre-clinical and clinical studies

Ion-Suppression and Phospholipid Contamination

When analyzing a compound and its metabolites in biological fluids, such as plasma or serum, one frequently deals with interference from the complex sample matrix. Ion-suppression of the mass spec signal due to contaminants in the matrix often limits our ability to properly identify and quantify the analytes of interest. The presence of phospholipids in biological fluids is one of the major causes of ion-suppression in LC-MS when using positive ion electrospray mode (+ESI). Removing phospholipids with HybridSPE-Phospholipid is a rapid and reliable means to improve identification and quantification of compounds in biological matrices using LC-MS.

How Does HybridSPE-Phospholipid Work?

Sample preparation with HybridSPE-Phospholipid is a very rapid and simple procedure. Proteins in the sample are precipitated by addition of acetonitrile containing

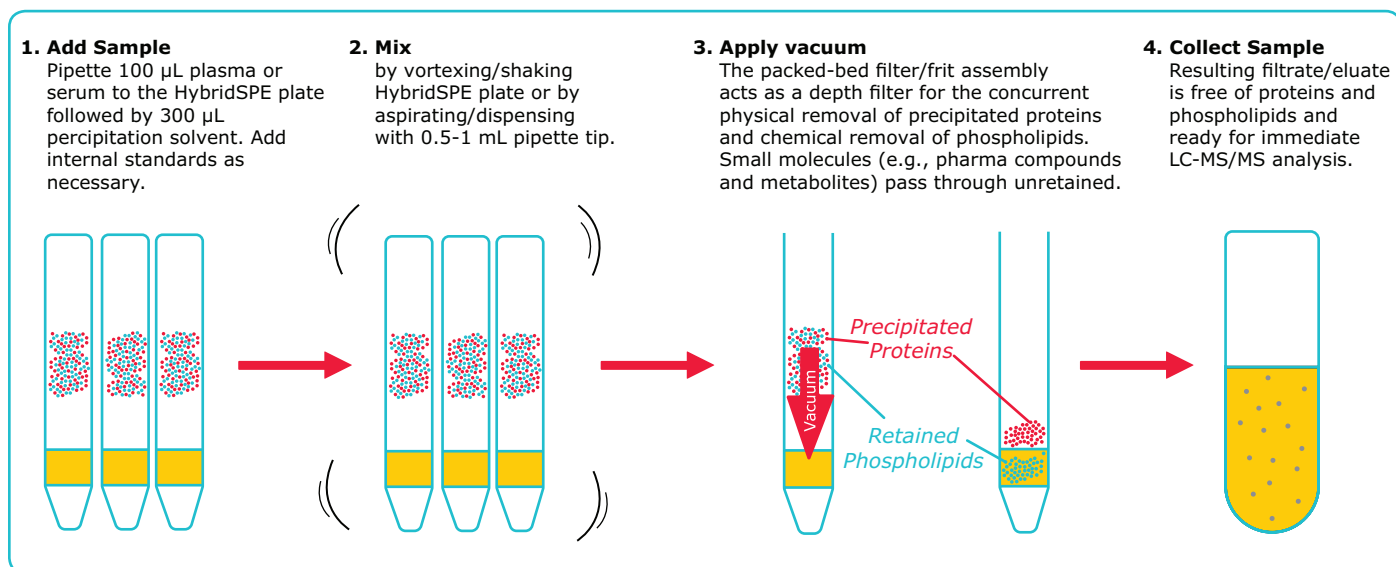
1% formic acid. The sample is then added to the HybridSPE-Phospholipid packed bed, either well plate or tube format. As shown in the accompanying figure, the bed consists of proprietary zirconia-coated silica particles. The zirconia sites exhibit Lewis acid (electron acceptor) properties that will interact strongly with Lewis bases (electron donors).

Phospholipids structurally consist of a polar head group (zwitterionic phosphonate moiety) and a large hydrophobic tail (two hydrophobic fatty acyl groups). The phosphonate group acts as a very strong Lewis base that interacts strongly with zirconia. Formic acid in the precipitation solvent is a critical modifier used to improve the recovery of many analytes of interest (particularly acidic compounds) by preventing analyte retention, while not affecting phospholipid removal.

The HybridSPE-Phospholipid sample preparation products are available in several configurations.

- Two 96-well plate formats for sample volumes of ~100 μ L and 20-40 μ L. Both formats allow for in-well precipitation.
- Three SPE tube formats; the ultra version allows for in-tube protein precipitation.

For more information and to view a video of HybridSPE-Phospholipid in operation, visit SigmaAldrich.com/hybridspe

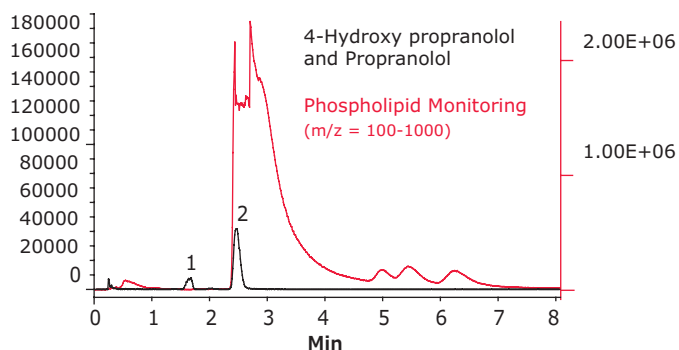


Ion-Suppression from Phospholipids: Standard Protein Precipitation vs. HybridSPE-Phospholipid

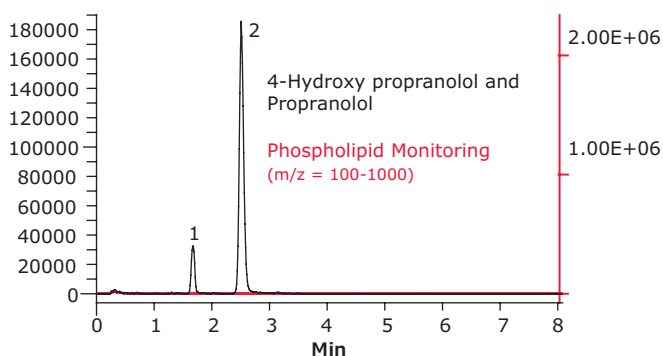
Sample prep	standard protein precipitation or HybridSPE-Phospholipid (575656-U)
Column	Ascentis Express F5, 5 cm x 2.1 mm I.D., 2.7 µm (53567-U)
Mobile phase	(A) 2 mM ammonium formate in acetonitrile; (B) 2 mM ammonium formate in water; (90:10, A:B)
Flow rate	0.4 mL/min
Pressure	1073 psi
Column temp	35 °C
Detector	MS, ESI(+) TOF, m/z = 100-1000
Injection	2 µL
Sample	Agilent 1200SL Rapid Resolution; 6210 Time of Flight (TOF) MS
System	Agilent 1200SL Rapid Resolution; 6210 Time of Flight (TOF) MS

Standard Protein Precipitation Technique

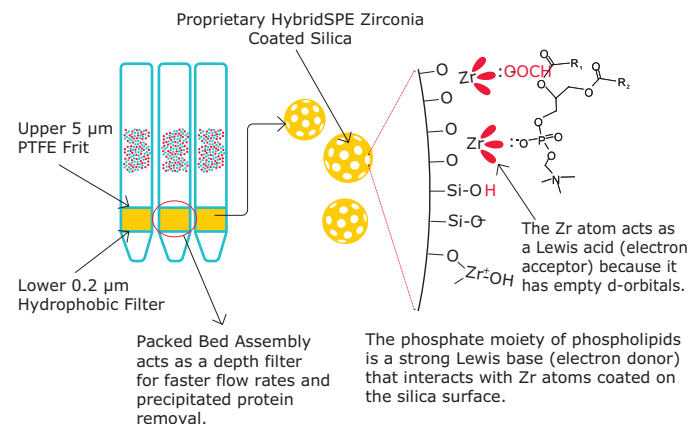
(Note suppression of propranolol signal)



HybridSPE-Phospholipid Technique



Interaction of Phospholipids with HybridSPE-Phospholipid



Featured Products

Description	Qty.	Cat. No.
HybridSPE-Plus Plate Essentials Kit		
Includes HybridSPE-Plus 96-well plate (575659-U), plate cap mat (as in 575680-U), sealing film (as in Z721581) and collection plate (as in Z717266)	1	52818-U
HybridSPE-Plus 96-Well Plates		
50 mg/well	1	575659-U
	20	575673-U
HybridSPE-Phospholipid Small Volume 96-Well Plates		
15 mg/well	1	52794-U
	20	52798-U
HybridSPE-Phospholipid Cartridges		
HybridSPE-Phospholipid Ultra Cartridge, 30 mg/1 mL	100	55269-U
HybridSPE-Phospholipid Cartridge, 500 mg/6 mL	30	55267-U
HybridSPE-Phospholipid Cartridge, 30 mg/1 mL	100	55261-U
HybridSPE-Phospholipid Cartridge, 30 mg/1 mL	200	55276-U

Protein Precipitation

96-Well Protein Precipitation Filter Plate

The 96-well protein precipitation filter plate is ideal for removing precipitated proteins from biological plasma/serum. The plate consists of a 0.2 µm hydrophobic graded filter/frit. Biological plasma is first added to the 96-well plate followed by a protein precipitating agent (e.g., acetonitrile). After a brief mixing step, vacuum is applied to the plate, and the filter/frit removes precipitated proteins from the sample. The resulting filtrate is ready for further processing and/or analysis.

Cat. No.	Description	Qty.
55263-U	2 mL	1 ea.

SupelMIP® Molecularly Imprinted Polymers

Key Features and Benefits

- Achieve lower detection limits through superior selectivity
- Reduce ion-suppression
- Save time and reduce cost via robust and rapid sample prep methodology
- Minimal to no method development required

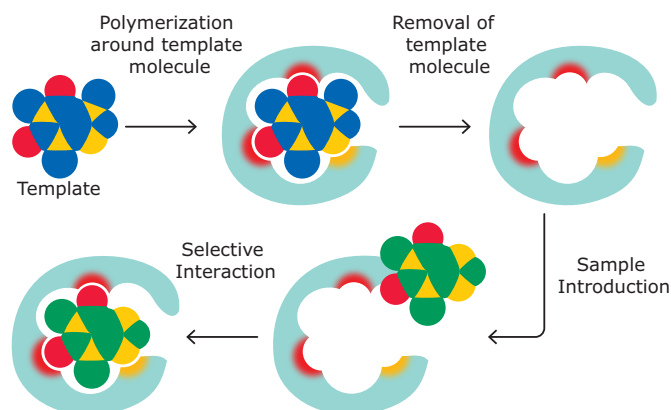
Highly Selective Extraction of Trace Analytes from Complex Matrices

Molecularly imprinted polymers (MIPs) are a class of highly cross-linked polymer-based molecular recognition elements engineered to bind one target compound or a class of structurally related target compounds with high selectivity. Selectivity is introduced during MIP synthesis in which a template molecule, designed to mimic the analyte, guides the formation of specific cavities or imprints that are sterically and chemically complementary to the target analyte.

SupelMIPs are available for these analyte and matrix combinations

Analytes	Matrix
Chloramphenicol	Milk, plasma, honey, urine, and shrimp/prawns
Clenbuterol	Urine
Fluoroquinolones	Bovine kidney, honey, and milk
PAHs	Edible oils
β-Agonists	Tissue, urine and wastewater
TSNAs (4 Different Tobacco-Specific Nitrosamines: NNK, NNN, NAB, NAT)	Urine and tobacco
NNAL (4-(Methylnitrosamino)-1-(3-pyridyl)-1-butanol)	Urine

Formation of MIPs



SupelMIP Molecularly Imprinted Polymer SPE Tubes

Cat. No.	Description	Pkg. Size
53225-U	SupelMIP SPE - β-agonists, bed wt. 25 mg, volume 3 mL	50
53202-U	SupelMIP SPE - β-agonists, bed wt. 25 mg, volume 10 mL	50
53209-U	SupelMIP SPE - Chloramphenicol, bed wt. 25 mg, volume 3 mL	50
53210-U	SupelMIP SPE - Chloramphenicol, bed wt. 25 mg, volume 10 mL	50
53201-U	SupelMIP SPE - Clenbuterol, bed wt. 25 mg, volume 10 mL	50
53269-U	SupelMIP SPE - Fluoroquinolones, bed wt. 25 mg, volume 3 mL	50
53222-U	SupelMIP SPE - TSNAs, bed wt. 50 mg, volume 3 mL	50
53221-U	SupelMIP SPE - TSNAs, bed wt. 50 mg, volume 10 mL	50
53203-U	SupelMIP SPE - NNAL, bed wt. 25 mg, volume 3 mL	50

Stable Isotope Labeled Bioactive Compounds

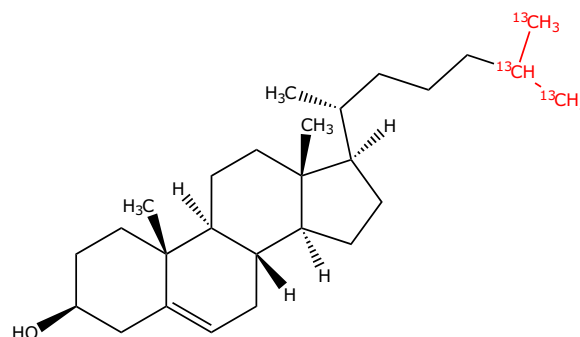
ISOTEC® Products for Use as Internal Standards

Stable isotope labeled compounds are used as internal standards for various MS techniques and within many applications. With chemical and ionization properties nearly identical to their unlabeled counterparts, stable isotope labeled compounds are often considered the top choice for an internal standard. Furthermore, the labeled standard and the analyte of interest can be easily differentiated by the mass shift between the two compounds, which is ideally three or more units.¹

ISOTEC Stable Isotopes offers a large selection of labeled products suitable for this purpose. Labeled standards have been utilized within numerous applications, including quantification of cholesterol in a clinical setting,² vitamin D within baby formula,³ and B vitamins in human milk.⁴ Labeled internal standards have also been employed in research on the diagnosis of Graves disease⁵ and hypertension,⁶ the study of fatty acid oxidation,⁷ and the analysis of androgenic steroids in wastewater.⁸

MS standards from ISOTEC have high chemical and isotopic purities with labeling patterns including ¹³C, ¹⁵N, and deuterium. The ¹³C and/or ¹⁵N labels do not exchange within the mass spectrometer source, providing further advantage.⁹

ISOTEC is also able to custom-synthesize labeled compounds upon request. Custom compounds can be designed with specific isotopes in specific locations. Whether a fully labeled or specifically labeled compound is of interest, let our expert team evaluate your needs.



References

1. Vogeser, M. and Seger, C., Pitfalls associated with the use of liquid chromatography-tandem mass spectrometry in the clinical laboratory. *Clin. Chem.*, **56**, 1234-1244 (2010).
2. Edwards, S.H., et al., Proposed serum cholesterol reference measurement procedure by gas chromatography-isotope dilution mass spectrometry. *Clin. Chem.*, **57**, 614-622 (2011).
3. Sullivan, D., Infant formula and adult/pediatric nutritional methods approved First Action using the AOAC voluntary consensus standards process. *J. AOAC Int.*, **95**, 1-4 (2012).
4. Hampel, D., et al., Ultra-performance liquid chromatography tandem mass-spectrometry (UPLC-MS/MS) for rapid, simultaneous analysis of thiamin, riboflavin, flavin adenine dinucleotide, nicotinamide and pyridoxal in human milk. *J. Chromatogr. B*, **903**, 7-13 (2012).
5. Higashi, T., et al., Stable isotope-dilution liquid chromatography/tandem mass spectrometry method for determination of thyroxine in saliva. *J. Chromatogr. B*, **879**, 1013-1017 (2011).
6. Taylor, P.J., et al., Measurement of aldosterone in human plasma by semiautomated HPLC-tandem mass spectrometry. *Clin. Chem.*, **55**, 1155-1162 (2009).
7. Mohammad, M.A., et al., Galactose promotes fat mobilization in obese lactating and nonlactating women. *Am. J. Clin. Nutr.*, **93**, 374-381 (2011).
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9. Ciccimaro, E. and Blair, I.A. Stable-isotope dilution LC-MS for quantitative biomarker analysis. *Bioanalysis*, **2(2)**, 311-341(2010).

To find additional stable isotope labeled standards, visit SigmaAldrich.com/isotec

To inquire about Stable Isotopes pricing and availability, email us at isosales@sial.com

Amino Acids

Cat. No.	Description
485578	DL-Alanine-2,3- ¹³ C ₂
776408	L-Arginine-2,3,4,4,5,5-d ₇ hydrochloride
643440	L-Arginine- ¹³ C ₆ hydrochloride
492353	DL-aspartic acid-2- ¹³ C, ¹⁵ N
748935	L-Citrulline-5- ¹³ C,4,4,5,5-d ₄
578886	L-Citrulline-4,4,5,5-d ₄
741833	L-Citrulline-5,5-d ₂
900206	DL-Cysteine-3,3-d ₂
604984	DL-Glutamic acid- ¹³ C ₅
588644	DL-Histidine-1- ¹³ C
605239	L-Leucine- ¹³ C ₆
643459	L-Lysine- ¹³ C ₆ hydrochloride
749443	L-Ornithine-3,3,4,4,5,5-d ₆ hydrochloride
489107	DL-Serine-1- ¹³ C
592048	DL-Valine-2- ¹³ C amine

Additional products and labeling patterns are available.

Fatty Acids

Cat. No.	Description
735000	Arachidonic-5,6,8,9,11,12,14,15-d ₈ acid
616125	Decanoic-10,10,10-d ₃ acid
733326	<i>cis</i> -4,7,10,13,16,19-Docosahexaenoic-21,21,22,22,22-d ₅ acid
734322	<i>cis</i> -5,8,11,14,17-Eicosapentaenoic acid-19,19,20,20,20-d ₅
617040	Heptanoic-d ₁₃ acid
722774	<i>trans</i> -9-Hexadecenoic acid-1,2,3,7,8- ¹³ C ₅
451401	Lauric-d ₂₃ acid
605735	Linoleic acid- ¹³ C ₁₈
733148	Methyl heptadecanoate-d ₃₃
490865	Myristic acid-1,2- ¹³ C ₂
614165	Myristic acid-13,13,14,14,14-d ₅
722847	<i>trans</i> -6-Octadecenoic acid-1,2,3,4,5- ¹³ C ₅
722790	<i>trans</i> -9-Octadecenoic acid-1,2,3,7,8- ¹³ C ₅
722855	<i>trans</i> -11-Octadecenoic acid-1,2,3,9,10- ¹³ C ₅
605727	Octanoic acid- ¹³ C ₈
448214	Octanoic-d ₁₅ acid
490431	Oleic acid- ¹³ C ₁₈
605573	Palmitic acid- ¹³ C ₁₆
366897	Palmitic acid-d ₃₁

Additional products and labeling patterns are available.

Glycerides & Lipids

Cat. No.	Description
729663	Cholesteryl linoleate- ¹³ C ₁₈
729671	Cholesteryl-26,26,26,27,27,27-d ₆ oleate-1,2,3,7,8,9,10- ¹³ C ₇
616966	Glyceryl tri(palmitate-d ₃₁)
617121	Glyceryl tri(octanoate-d ₁₅)
605638	Glyceryl- ¹³ C ₃ trioleate
729523	Cholesteryl oleate- ¹³ C ₁₈
777862	Glyceryl tri(palmitate-1,2,3,4- ¹³ C ₄)
730041	2-Oleoyl-1-palmitoyl-rac-glycero-3-phosphocholine-(trimethyl-d ₉)
729507	Glyceryl-d ₅ trilinoleate
729515	Cholesteryl-26,26,26,27,27,27-d ₆ linoleate
723703	rac-Glyceryl-1,1,2,3,3-d ₅ -1,2-dioleate
646253	Glyceryl tri(oleate-1,2,3,7,8,9,10- ¹³ C ₇)
757438	1-Palmitoyl-rac-glycero-3-phosphocholine-(trimethyl-d ₉)
489514	Glyceryl tri(oleate-1- ¹³ C)
646245	Glyceryl tri(oleate-9,10- ¹³ C ₂)
757438	1-Palmitoyl-rac-glycero-3-phosphocholine-(trimethyl-d ₉)
492663	Glyceryl tri(stearate-1- ¹³ C)

For a full listing of labeled lipid and fatty acid products, visit [SigmaAldrich.com/lipid](https://www.sigmaaldrich.com/lipid)

To inquire about Stable Isotopes pricing and availability, email us at isosales@sial.com

Steroids and Hormones

Cat. No.	Description
706035	Aldosterone-2, 2, 4, 6, 6, 21, 21-d ₇
802883	Aldosterone-9,11,12,12-d ₄ solution
730645	4-Androstene-3, 17-dione-2, 3, 4- ¹³ C ₃ solution
749478	Cholesterol-2,3,4- ¹³ C ₃
488577	Cholesterol-2, 2, 3, 4, 4, 6-d ₆
707678	Cholesterol-25, 26, 27- ¹³ C ₃
802905	Corticosterone-9,11,12,12-d ₄
803146	Hydrocortisone-2,3,4- ¹³ C ₃ solution
803154	Cortisone-2, 3, 4- ¹³ C ₃ solution
709549	Dehydroepiandrosterone-2, 2, 3, 4, 4, 6-d ₆
723266	Dehydroepiandrosterone-2, 2, 3, 4, 4, 6-d ₆ sulfate sodium salt
710784	11-Deoxycortisol-2, 2, 4, 6, 6-d ₅
730637	Dihydrotestosterone-2, 3, 4- ¹³ C ₃ solution, 0.1 mg/mL
719552	17β-Estradiol-2, 3, 4- ¹³ C ₃
613967	17β-Estradiol-2, 4, 16, 16, 17-d ₅
731668	Estril-2, 3, 4- ¹³ C ₃
719544	Estrone-2, 3, 4- ¹³ C ₃
802921	Estrone-2,3,4- ¹³ C ₃ solution
710806	18-Hydroxycorticosterone
705594	Hydrocortisone-9, 11, 12, 12-d ₄
803081	17-α-Hydroxypregnenolone-20, 21- ¹³ C ₂ -16,16-d ₂
739545	Pregnenolone-20, 21- ¹³ C ₂ -16, 16-d ₂
740985	Pregnenolone-20, 21- ¹³ C ₂ -16, 16-d ₂ sulfate sodium salt
737143	Progesterone-2, 3, 4- ¹³ C ₃
803065	Progesterone-2, 3, 4- ¹³ C ₃ solution
730610	Testosterone-2, 3, 4- ¹³ C ₃ solution
750026	3α, 5β-Tetrahydroaldosterone
709719	3, 3', 5'-Triiodothyronine-(diiodophenyl- ¹³ C ₆) hydrochloride
709611	3, 3', 5'-Triiodothyronine-(tyrosine ring- ¹³ C ₆) hydrochloride
809667	Chenodeoxycholic-2,2,3,4,4,6,6,7,8-d ₉ acid
809837	Cholesterol-23,24,25,26,27- ¹³ C ₅
809845	Pregnenolone-2,2,4,4-d ₄
809853	Allopregnanolone-2,2,3,4,4-d ₅ solution
809861	Etiocholanolone-2,2,3,4,4-d ₅ solution
900079	Cortisone-2,3,4- ¹³ C ₃ 21-sulfate sodium salt solution
900080	Hydrocortisone-9,11,12,12-d ₄ 21-sulfate sodium salt
900067	L-Thyroxine-1",1",2,2",6-d ₅ hydrochloride solution
900170	Cortisone-2,2,4,6,6,9,12,12-d ₈
900182	Tetrahydrocortisol-2,2,3,4,4-d ₅
900183	Tetrahydrocortisone-2,2,3,4,4-d ₅

For a full listing of labeled vitamins, steroids and hormones, visit

[SigmaAldrich.com/sibio](https://www.sigmaaldrich.com/sibio)

To inquire about Stable Isotopes custom synthesis or pricing and availability, email us at isosales@sial.com

Vitamins

Cat. No.	Description
705268	Biotin-(ring-6, 6-d ₂)
802891	Coenzyme Q10-(ring-d ₉)
803162	Folic acid-(glutamic acid- ¹³ C ₅ , ¹⁵ N)
803049	Folic acid-(glutamic acid- ¹³ C ₅)
803030	25-Hydroxyvitamin D ₃ -(26,26,26,27,27,27-d ₆)
802913	(24R), 24,25-Dihydroxyvitamin D ₃ -26,26,26,27,27,27-d ₆ solution
803103	25-Hydroxyvitamin D ₃ -(23-24-25-26-27- ¹³ C ₅) solution
740217	25-Hydroxyvitamin D ₂ solution
740071	25-Hydroxyvitamin D ₂ (6, 19, 19-d ₃) solution
705497	25-Hydroxyvitamin D ₂ (6, 19, 19-d ₃)
739650	25-Hydroxyvitamin D ₃ solution
705888	25-Hydroxyvitamin D ₃ (6, 19, 19-d ₃)
762970	Nicotinamide-2, 4, 5, 6-d ₄
705187	Pyridoxal-(methyl-d ₃) hydrochloride
705322	Pyridoxamine-(methyl-d ₃) dihydrochloride
705292	Riboflavin-dioxypyrimidine- ¹³ C ₄ , ¹⁵ N ₂
731188	Thiamine-(4-methyl- ¹³ C-thiazol-5-yl- ¹³ C ₃) hydrochloride
731234	α-Tocopherol-(ring-5, 7-dimethyl-d ₆)
705837	Vitamin B ₅ (di-β-alanine- ¹³ C ₆ , ¹⁵ N ₂) calcium salt
803170	Vitamin B ₁₂ -(dimethylbenzimidazole- ¹³ C ₇) solution
705489	Vitamin D ₂ (6,19,19-d ₃)
739839	Vitamin D ₂ (6,19,19-d ₃) solution
740284	Vitamin D ₃ (6,19,19-d ₃) solution
731285	
615366	Vitamin E acetate-(trimethyl-d ₉)
705470	Vitamin K-d ₃ (5,6,7,8-d ₄ , 2-methyl-d ₃)
737836	Vitamin K ₃ -d ₈
809608	Biotin-2',2',3',3'-d ₄
809659	Pyridoxine-(methyl-d ₃) hydrochloride
809926	1α,25-Dihydroxyvitamin D ₃ -26,26,26,27,27,27-d ₆ solution
809748	(24R)-24,25-Dihydroxyvitamin D ₃ solution
809756	Vitamin D ₃ -25,26,27- ¹³ C ₃ solution
900234	Vitamin D ₃ -23,24,25,26,27- ¹³ C ₅ solution
809772	Vitamin D ₃ -23,24,25,26,27- ¹³ C ₅ solution
809799	Nicotinamide-2,6,7- ¹³ C ₃ -(pyridyl- ¹⁵ N)
809888	Vitamin K1-4a,5,6,7,8,8a- ¹³ C ₆
809896	Vitamin K2 (MK-4)-(5,6,7,8-d ₄ ,2-methyl-d ₃)
809918	Vitamin K2 (MK-4)-4',5,6,7,8,8'- ¹³ C ₆
900074	Vitamin K2 (MK-7)-(5,6,7,8-d ₄ ,2-methyl-d ₃)
900076	Vitamin K2 (MK-9)-(5,6,7,8-d ₄ ,2-methyl-d ₃)
900075	Vitamin K2 (MK-7)-4',5,6,7,8,8'- ¹³ C ₆
900077	Vitamin K2 (MK-9)-4',5,6,7,8,8'- ¹³ C ₆

Metabolites

Cat. No.	Description
809616	5-Hydroxyindole-3a,4,5,6,7,7a- ¹³ C ₆ -3-acetic acid
809683	Sodium taurochenodeoxycholate-2,2,4,4-d ₄
809691	Sodium taurochenodeoxycholate-2,2,3,4,4,4,6,6,7,8-d ₉
900036	Sodium taurocholate-2,2,4,4-d ₄
900078	Sodium taurodeoxycholate-2,2,4,4,11,11-d ₆
900073	Sodium taurodeoxycholate-2,2,4,4-d ₄
809713	Sodium tauroolithocholate-2,2,4,4-d ₄
809721	Sodium tauroursodeoxycholate-2,2,4,4-d ₄
809780	Indoxyl-3a,4,5,6,7,7a- ¹³ C ₆ sulfate potassium salt

Other Bioactive Compounds

Cat. No.	Description
750913	L-Arbrine-(methyl-d ₃)
733865	Aldicarb-(N-methyl- ¹³ C,d ₃ , carbomoyl- ¹³ C)
733873	Aldicarb-(N-methyl- ¹³ C,d ₃ , carbomoyl- ¹³ C) sulfone
719579	(±)-Catechin-2,3,4- ¹³ C ₃
614122	Chenodeoxycholic acid-2,2,4,4-d ₄
614149	Cholic acid-2,2,4,4-d ₄
614130	Deoxycholic acid-2,2,4,4-d ₄
705349	Desethylamodiaquine-(ethyl-d ₅)
719528	3, 3'-Diiodo-L-thyronine-(phenoxy- ¹³ C ₆) (T2)
719536	3,3'-Diiodo-L-thyronine (T2)
749001	4,6-Dioxoheptanoic acid-3,4,5,6,7- ¹³ C ₅
722820	Ferulic acid-1,2,3- ¹³ C ₃
739723	Glycocholic-2,2,4,4-d ₄ acid
762962	Histamine-α, α,β, β-d ₄ dihydrochloride
793477	Kynurenic acid-3,5,6,7,8-d ₅
709891	Spermidine-(butane-d ₈) trihydrochloride
740780	Spermidine-(butane- ¹³ C ₄) trihydrochloride
705330	Spermine-(butane-d ₈) tetrahydrochloride
746274	Vinblastine- ¹³ C,d ₃
731242	Yohimbine-(methyl- ¹³ C,d ₃ ester)
900368	(±)-Epicatechin-2,3,4- ¹³ C ₃ gallate
900369	(±)-Epigallocatechin-2,3,4- ¹³ C ₃
900376	(±)-Epigallocatechin-2,3,4- ¹³ C ₃ gallate
900370	(±)-Catechin-2,3,4- ¹³ C ₃ gallate
900371	(±)-Gallocatechin-2,3,4- ¹³ C ₃
900372	(±)-Gallocatechin-2,3,4- ¹³ C ₃ gallate
900374	α-Tocopherol-(phenyl- ¹³ C ₆)
809594	11-Deoxycortisol-2,3,4- ¹³ C ₃ solution
809586	11-Deoxycorticosterone-2,3,4- ¹³ C ₃ solution
809640	Dehydroepiandrosterone-2,2,3,4,4-d ₅
809802	Exemestane-(3,4- ¹³ C ₂ -6-methylidene- ¹³ C)
809810	Clodinafop-propargyl-(phenoxy- ¹³ C ₆)
809829	Atrazine-(triazyl- ¹³ C ₃ , ¹⁵ N ₃)

Mass Spectrometry Metabolite Library of Standards

Supplied by IROA Technologies

Product Description

MSMLS™ (Mass Spectrometry Metabolite Library of Standards) is a collection of high quality small biochemical molecules that span a broad range of primary metabolism. These are high purity (>95%) compounds supplied in an economical, ready-to-use format. The library of standards are most commonly used to provide retention times and spectra for key metabolic compounds, help optimize mass spectrometry analytical protocols, and qualify and quantify mass spectrometry sensitivity and limit of detection. MSMLS comes with MSMLSDiscovery™, a software tool to support the extraction, manipulation, and storage of the data generated when using our MSMLS Library of authentic metabolomics standards.

Features and Benefits

Compounds

600 unique small molecule metabolites organized in a 96-well format.

Broad metabolite spectrum, key primary metabolites and intermediates covering key metabolic pathways, including the following classes of compounds:

- Carboxylic acids, amino acids
- Biogenic amines, polyamines
- Nucleotides, coenzymes and vitamins
- Mono- and disaccharides
- Fatty acids, lipids, steroids, and hormones



Convenient

- High purity, pre-weighed metabolites supplied dried. Added solubility instructions to ensure easy, efficient sample preparation
- The library is intended to be used for mass spectrometry metabolomics applications and provides a broad representation of primary metabolites

Formatted

MSMLS™ contains 600 small molecule metabolites:

- Arrayed in 96-well format
 - 7 polypropylene racks
 - Supplied as 5 µg dried weight
- Rack map provided upon purchase
 - Alphanumeric assigned position
 - Descriptors: Name, Parent CID, KEGG ID, molecular formula, molecular weight, CAS, ChEBI, HMDB/YMDB ID, PubChem Compound, Substance ID, Metlin ID
- Suitable for manual and automated work flow

Software

MSMLSDiscovery™ software package is distributed with and is tailored to work with MSMLS. A User Manual and video instructions are provided. For the second version additional data has been added for clarity. In addition a section has been added to outline the process for analyzing samples once the library has been created. The requirements of the program are that:

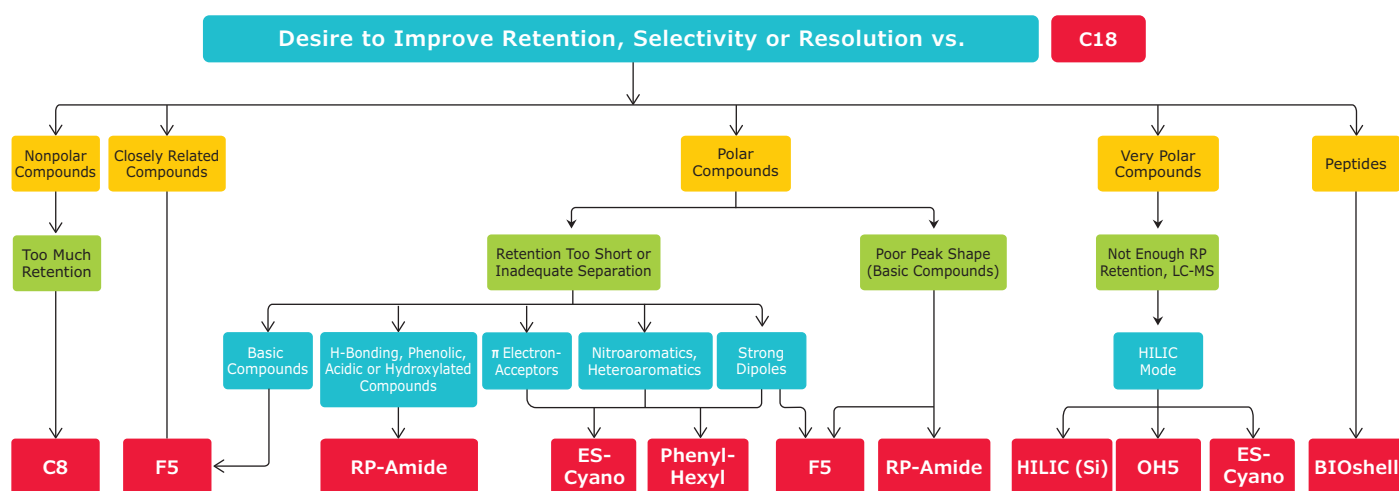
- The most recent version of Java 7 must be installed and callable
- The computer should have at least 8 GB of RAM
- You are running Windows 7 or higher

For more information, visit SigmaAldrich.com/msmls

Selecting The Right HPLC Phase Chemistry for Your Application

C18 column is the standard first choice when starting a new LC-MS method. You can consider selecting another stationary phase when C18 doesn't give the desired separation, or the sample contains compounds difficult to retain or resolve on C18. The Ascentis Express and BIOshell product lines offer a wide range of selectivities for making an effective choice.

This decision tree will help you to select an alternative phase based on the particular compound type or separation challenge. All options displayed are relative to the C18 column that started your separation journey.



Key product features for LC/MS and (U)HPLC applications

Primary Application	Product Line	Particle Size (μm)	Pore Size (Å)	Surface Area (m^2/g)	Max Temperature	Pressure (bar)
Small molecules, metabolites and low molecular weight peptides	Titan	1.9	80	410	60	1,000
	Ascentis Express	2.0	90	120	60	1,000
		2.7	90	150	60	600
		5.0	90	100	60	600
Proteins, Peptides and large Bio-molecules	BIOshell	2.7	160	90	90	600
		3.4	400	15	90	600
		5.0	160	60	90	600

Available in a variety of analytical and capillary column dimensions

Column I.D	Column Length (cm)						
	2	3	5	7.5	10	15	25
75 μm			.			.	
100 μm			.			.	
200 μm			.			.	
2.1 mm
3 mm
4.6 mm

For Part Numbers, visit [SigmaAldrich.com/hplc](https://www.sigmaaldrich.com/hplc)

For a complete listing of LC/MS columns, visit [SigmaAldrich.com/hplc](https://www.sigmaaldrich.com/hplc)

LC/MS & (U)HPLC Columns

Ascentis Express & BIOshell™ Fused-Core® U/HPLC & LC/MS Columns

Key Features and Benefits*

- Maximize speed with sharp peaks even at ultra-high flow rates
- Stable low-bleed for LC-MS and LC-UV
- Suitable for any HPLC, UHPLC, and LC-MS instruments
- Achieve UHPLC performance on a traditional HPLC system
- Available in both 2.0, 2.7 and 5 µm particles
- Wide variety of pore sizes, ranging from 90 - 1000 Å, for small to large molecules

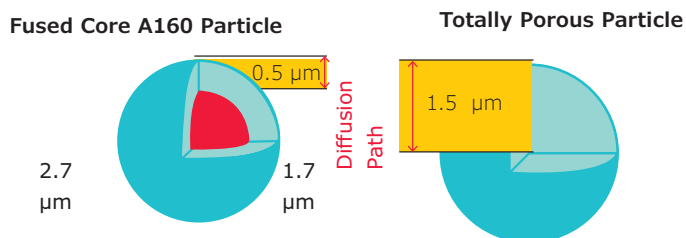
Ascentis® Express Fused-Core® Columns

Ascentis Express columns provide a breakthrough in (U)HPLC and LC/MS column performance. Based on Fused-Core particle technology, Ascentis Express columns provide the benefits of high speed and high efficiency. The Fused-Core particle consists of a solid core and a porous shell, allowing for a shorter diffusion path compared to conventional fully porous particles. Compared to totally porous particles typically used in HPLC, Ascentis Express Fused-Core particles generate approximately half the backpressure without loss of resolution. This permits for more resolving power, and faster flow rates, for higher throughput.

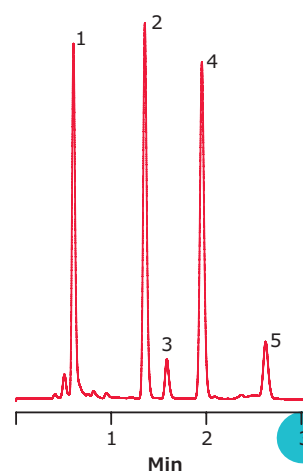
Ascentis Express Fused Core Columns are now available in 2.0, 2.7 and 5 µm particle sizes with 8 different phase chemistries. Available in pore size of 90 Å, Ascentis Express are ideal for LC/MS and (U)HPLC separations of small molecules, metabolites and low molecular weight peptides.

For more information, visit SigmaAldrich.com/express

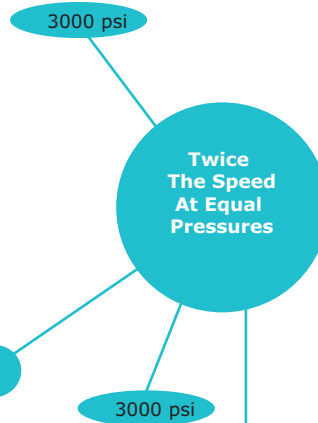
Comparison of Fused-Core and Standard HPLC Particle



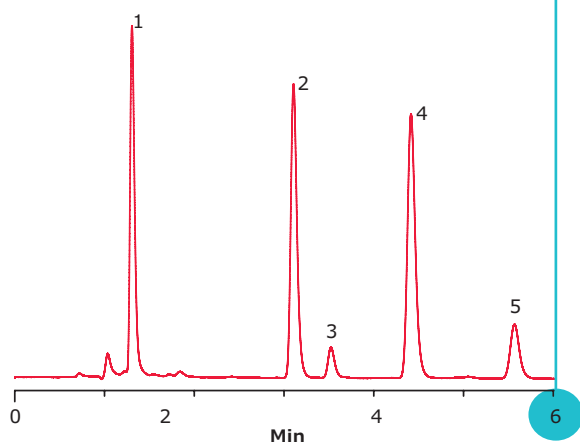
Ascentis Express C18
0.4 mL/min flow rate



1. Estriol (Cerilliant E-074)
2. 17β-Estradiol (Cerilliant E-061)
3. Unknown
4. Estrone (Cerilliant E-075)
5. Estrone degradant

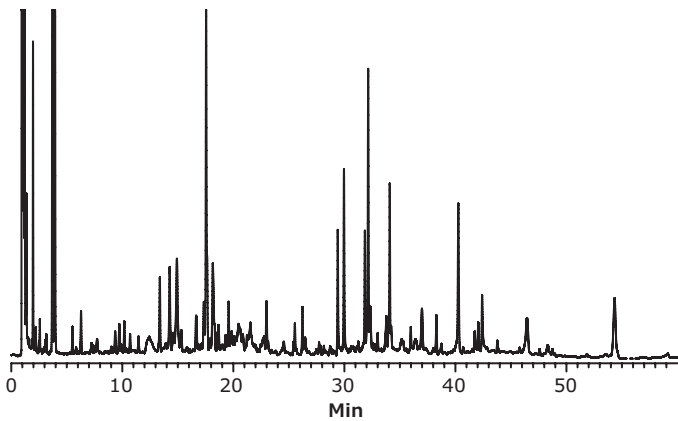


C18 Sub-2 µm
0.2 mL/min flow rate



Analysis of Tryptic Digests on BIOshell A160 Peptide ES-C18

Column	BIOshell A160 Peptide C18, 10 cm x 4.6 mm I.D. (66915-U)
Mobile phase A	0.1% (w/v) TFA in water
Mobile phase B	0.1% TFA (w/v) in 40:60
Water	acetonitrile
Gradient	initial = 3% B to 100% B in 53 min.
Flow rate	1.0 mL/min
Temp.	30 °C
Det.	UV at 215 nm
Injection	20 µL



BIOshell™ Fused-Core® Columns

Faster, Better Peptide and Protein Separations

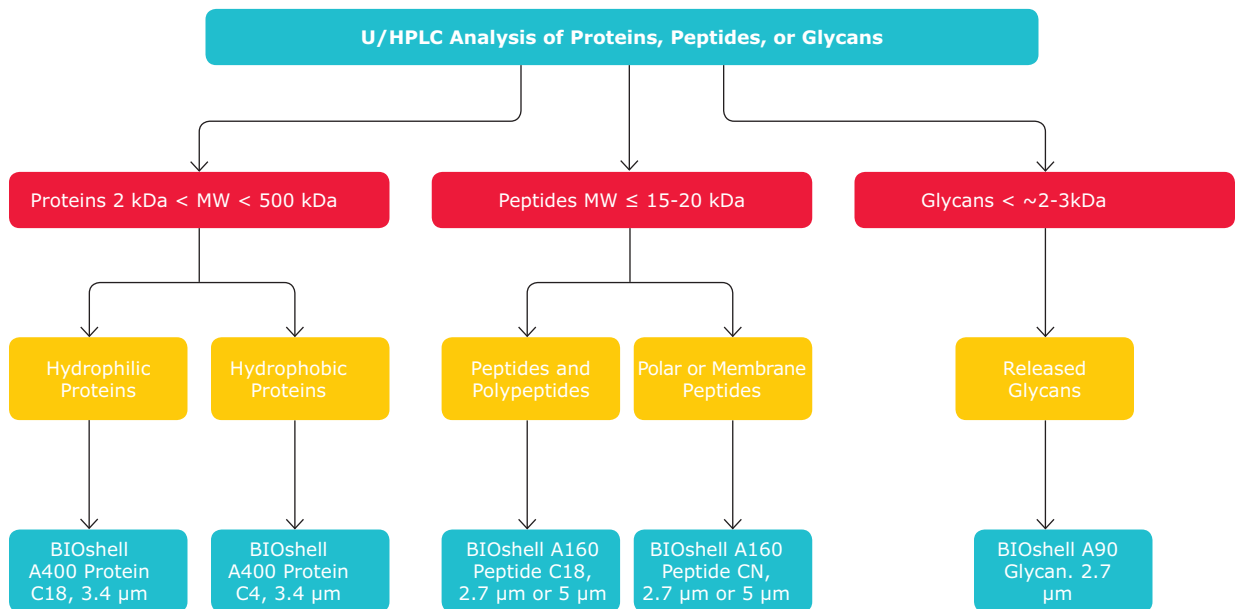
BIOshell columns are the most recent innovation in Fused-Core particle technology: high efficiency reversed-phase columns for protein and peptide separations. BIOshell columns can be operated in HPLC or UHPLC instrumentation equipped with a mass spectrometer or any other detector.

A complete BIOshell column lineup includes:

- BIOshell A160 Peptide C18 and CN
- BIOshell A400 Protein C4 and C18
- BIOshell A1000 IgG C4 (New)
- BIOshell A90 Glycan

For more information, visit SigmaAldrich.com/bioshell

Selecting the Right BIOshell Column



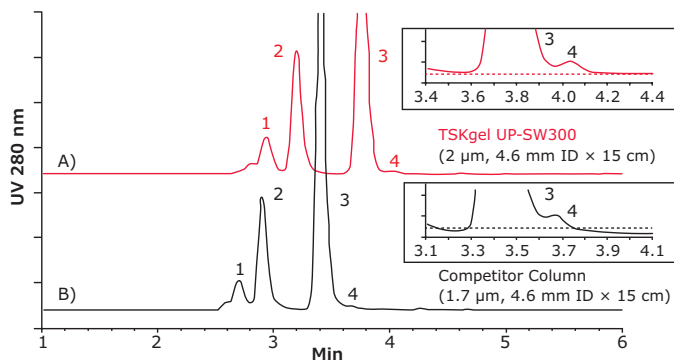
TSKgel® HPLC Columns for Protein Fractionation and Peptide Analysis

New 2 µm Particle TSKgel® UP-SW3000

SEC-UHPLC Column

Ideally Suited for Monoclonal Antibody (mAb) Aggregate and Fragment Analysis

- Speed up your HPLC -SEC analysis without giving up any resolution
- Gain more resolution within the same time frame as your current UHPLC -SEC analysis



Sample. Mouse-human chimeric mAb; 1: trimer; 2: dimer; 3: monomer ; 4: fragment.

Column	Rs (peak 1/2)	Rs (peak 2/3)
TSKgel UP-SW3000 2 µm	1.52	3.56
competitor UHPLC-SEC 1.7 µm	1.25	3.47

Quality, Consistency and Fast Delivery Time

Cat. No.	Description
80023448	TSKgel UP-SW3000 2 µm, 4.6 × 300 mm
80023449	TSKgel UP-SW3000 2 µm, 4.6 × 150 mm
80023450	TSKgel UP-SW Guardcolumn 2 µm, for P/N 0082348/9
80023451	TSKgel UP-SW DC Guardcolumn 2 µm, for 823448/9
69385	Protein Standard Mix 15-600Kda

TSKgel SW Product Line Overview

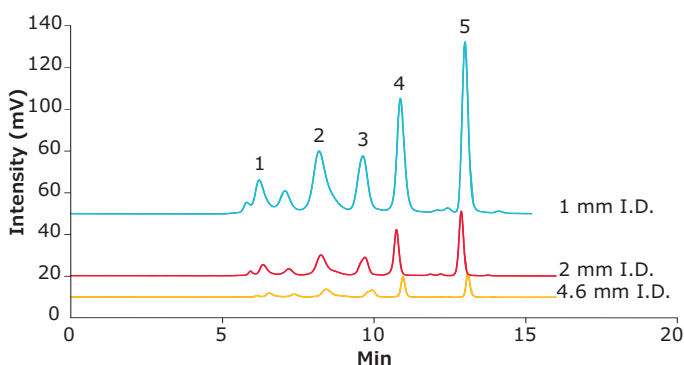
TSKgel Column Type	Particle Size [µm]	Pore Size [Å]	Molecular Weight (Globular Proteins)
SuperSW2000	4	125	5 – 150 kDa
GFC-200, G2000SW _{XL} , BioAssist 2SWXL	5	125	5 – 150 kDa
G2000SW	10, 13, 20	125	5 – 150 kDa
UP-SW3000	2	250	10-500 kDa

TSKgel Column Type	Particle Size [µm]	Pore Size [Å]	Molecular Weight (Globular Proteins)
SuperSW3000	4	250	10 – 500 kDa
SuperSW mAb (HTP, HR)	4	250	10 – 500 kDa
GFC-300, G3000SW _{XL} , BioAssist 3SWXL	5	250	10 – 500 kDa
G3000SW	10, 13, 20	250	10 – 500 kDa
UltraSW Aggregate	3	300	10 – 2.000 kDa
G4000SW _{XL} , BioAssist 4SWXL	8	450	20 – 7.000 kDa
G4000SW	13, 17	450	20 – 7.000 kDa

Cat. No.	Description	L (cm)	I.D. (mm)	Pore Diameter (Å)	Particle Size (µm)
818674	TSKgel® SuperSW2000 Size Exclusion HPLC Column	30	4.6	125	4
818675	TSKgel® SuperSW3000 Size Exclusion HPLC Column	30	4.6	250	4
821485	TSKgel® SuperSW3000 Size Exclusion HPLC Column	30	2.0	250	4
821845	TSKgel® SuperSW3000 Size Exclusion HPLC Column	30	1.0	250	4

Effect of Sample Mass on Detection Sensitivity

Columns	TSKgel SuperSW3000, 30 cm × 4.6 mm I.D. TSKgel SuperSW3000, 30 cm × 2 mm I.D. TSKgel SuperSW3000, 30 cm × 1 mm I.D.
Eluent	0.1 mol/L phosphate buffer + 0.1 mol/L Na ₂ SO ₄ + 0.05% Na ₃ (pH 6.7)
Flow rate	0.350 mL/min (4.6 mm I.D.) 0.650 mL/min (2 mm I.D.) 0.016 mL/min (1 mm I.D.)
Detection	UV at 280 nm
Detector cell volume	2 µL (4.6 and 2 mm I.D.) 35 nL (1 mm I.D.)
Temperature	25 °C
Injection volume	1 µL



For more information on TSKgel columns, visit SigmaAldrich.com/tsk

Chemical Derivatization Reagents for LC-MS

Modern mass spectrometry techniques such as APCI or ESI are highly successful in providing valuable structural information and allow the detection of very low analyte concentrations in various sample matrices. However, in today's advanced research and analytical areas, such as metabolomics, clinical and forensics analytics, such methods are sometimes insufficiently sensitive to deliver the solution to a particular analytical problem [1].

Therefore, derivatization is used in mass spectrometry to increase ionization efficiency, and thus enhance the sensitivity of the ionization used, to result in lower analyte detectability [2]. The derivatization reagents have functional groups with high proton (cation) affinity that stabilize positive charge. Of similar importance in derivatization is the improvement of qualitative analysis by modifying fragmentation behavior to form unique product ions, and shifting them to a specific, unique mass ("fingerprinting"), as well as precise quantitative analysis to profile comparatively small analyte molecules, particularly in metabolomics.

For more information, visit

SigmaAldrich.com/derivatization

References:

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9. (a) O.Y. Al-Dirbashi, et al., *J. Lipid Res.* 2008, 49, 1855-1862; b) Y. Tsukamoto, et al., *Biomed. Chromatogr.* 2005, 19, 802-808.
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19. C. O. Miles, et al., *Environ. Sci. Technol.* 2013, 47, 4080-4087.

Cat. No.	Description	Analyte Functional Group	Typical Application	Reference
00721	4-(Dimethyl- <i>d</i> ₆ -amino)benzoyl chloride	Hydroxy	Deuterium mass shift	[3]
03334	Dansylhydrazine	Carbonyl	—	[2c]
03641	Dansyl chloride	Hydroxy	—	[2c]
05022	<i>N,N</i> -Dimethylglycine	Hydroxy	Cholesterol	[11]
05689	Diethyl ethoxymethylenemalonate	Amine	Amino acids	[12]
06696	3-Amino-9-ethylcarbazole	Hydroxy	Sugars	[13]
06963	4-(Diethylamino)benzhydrazide	Carbonyl	—	[3]
08843	2-Hydrazinopyridine	Carbonyl	Steroids	[14]
29208	(<i>N</i> -Succinimidylloxycarbonylmethyl) tris(2,4,6-trimethoxyphenyl) phosphonium bromide	Amine	Protein sequence analysis	[15]
42579	4-Phenyl-1,2,4-triazoline-3,5-dione	Diene	Vitamin D	[16]
59799	4-(Diethylaminomethyl) benzhydrazide	Carbonyl	—	[3]
61224	<i>N</i> -Succinimidyl 4-(dimethylamino) benzoate	Amine	Glycerophosphoethanolamine lipids	[4]
65562	2-Picolylamine	Carbonyl	Steroids	[14]
67954	4-(Dimethylamino) benzoyl chloride	Hydroxy	17β-Estradiol	[3]
69706	6-Bromo-3-pyridinylboronic acid	1,2-Dihydroxy	Brassinosteroids	[5]
72702	3,5-Dinitrobenzoyl chloride	Hydroxy	Tetrahydrocorticosterones	[6]
73177	1-Fluoro-2,4-dinitrobenzene	Amine	Prim./sec. aliphatic amines	[7]
74905	9-Anthracenemethanol	Carboxylic acid, amine, alcohol	—	[17]
75821	1,2-Benzo-3,4-dihydrocarbazole-9-ethyl- <i>p</i> -toluenesulfonate	Carboxylic acid	Fatty-/bile acids	[8]
79291	4-[2-(<i>N,N</i> -Dimethylamino)ethylaminosulfonyl]-7-(2-aminoethylamino)-2,1,3-benzoxadiazole	Carboxylic acid	Fatty acids	[9]
89397	Girard's reagent T	Carbonyl	Nucleosides	[18]
92989	4-(Dimethylamino)benzohydrazide	Carbonyl	—	[3]
93742	Pentafluorophenylhydrazine	Carbonyl	Oligosaccharides	[10]
94076	{1-[2-(Diethylamino)ethoxy]-2-isothiocyanatoethyl} benzene	Amine	—	[3]
97622	2-Mercaptoethanol	Double bond	Microcystins	[19]

Tools for Metabolite Analysis by GC-MS

Strategies to analyze small biological compounds in a metabolome range from analyzing a particular class of metabolites (targeted analysis) to separating and detecting as many metabolites as possible of a particular developmental stage (metabolite profiling or metabonomics). When gas chromatography (GC) is used as the separation technique, the analyst benefits from the high resolving power of capillary GC, but the task is complex, as not all compounds are volatile and therefore need to be derivatized before analysis.

This and other pages in this publication list selected product options for the analysis of volatile and semi-volatile metabolites, including metabolite standards, derivatization reagents, solid-phase microextraction (SPME), and selected GC columns and accessories. For detailed information, references 1 and 2 look at the role of GC and MS in metabolite analysis, while references 3 and 4 discuss compound identification and sample throughput, respectively.

References

1. D. Wishart, Chapter 10, "Metabolomics in Humans and Other Mammals", in *Metabolome Analysis: An Introduction*, SG Villas-Boas, J. Nielsen, J. Smedsgaard, M. Hansen, U. Roessner-Tunali, eds., John Wiley & Sons, 2007
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3. Applying In-Silico Retention Index and Mass Spectra Matching for Identification of Unknown Metabolites in Accurate Mass GC-TOF Mass Spectrometry, Kumari, S., et al., *Anal. Chem.* 2011, 83, 5895-5902
4. Fast, High Peak Capacity Separations in Gas Chromatography-Time-of-Flight Mass Spectrometry, Wilson, R.B., et al., *Anal. Chem.* 2012, 84, 4167-4173

SLB®-5ms, An MS-Grade Capillary GC Column for Metabolomics Research

The 5% phenyl equivalent phase provides a boiling point elution order with a slight increase in selectivity, especially for aromatic compounds. The low bleed characteristics, inertness, and durable nature make it the column of choice for the analysis of semivolatiles or, in general, any application that requires a low bleed non-polar column. Temp. Limits for ≤0.25 mm I.D. are -60 °C to 340 °C (isothermal) or 360 °C (programmed).

Cat. No.	I.D. (mm)	df (μm)	Length (m)	Beta Value	Qty.
28465-U	0.10	0.10	10	250	1 ea.
28466-U	—	0.10	15	250	1 ea.
28564-U	0.18	0.18	20	250	1 ea.
28566-U	—	0.30	12	150	1 ea.
28575-U	—	0.30	30	150	1 ea.
28576-U	—	0.36	20	125	1 ea.
28513-U	0.20	0.20	30	250	1 ea.
28467-U	0.25	0.10	30	625	1 ea.
28469-U	—	0.25	15	250	1 ea.
28471-U	—	0.25	30	250	1 ea.

Cat. No.	I.D. (mm)	df (μm)	Length (m)	Beta Value	Qty.
28472-U	—	0.25	60	250	1 ea.
28577-U	—	0.50	15	125	1 ea.
28473-U	—	0.50	30	125	1 ea.
28474-U	—	0.50	60	125	1 ea.
28476-U	—	1.00	30	63	1 ea.

Extend the Lifetime of Your Capillary Column

A guard column/retention gap is a short (1-5 m) piece of uncoated deactivated fused silica tubing which is placed in-line between the GC injection port and the capillary column. A guard column/retention gap consists of two parts: a short length of fused silica tubing and a connector. Match the deactivation of the fused silica tubing with the polarity of the injection solvent. In most cases, it is also recommended to match the I.D. of the capillary column.

For more information about guard column selection, visit SigmaAldrich.com/gc-guard

SPME, A Unique Sample Preparation Technique

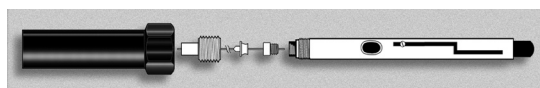
Solid Phase Microextraction is the sample preparation technique of choice for analyzing volatile and semi-volatile metabolites by GC-MS. SPME eliminates most drawbacks to extracting organics by more traditional methods. It requires no solvents or complicated apparatus, and can concentrate volatile and nonvolatile compounds, in both liquid and gaseous samples, for analysis by GC and GC-MS. SPME reduces sample preparation time by 70%, minimizes the use of solvents and their disposal, is cost-effective, can be used with any GC system, and can be automated.

An SPME fiber assembly consists of a length of fused silica fiber coated with a polymer material, in some cases mixed with a solid adsorbent. The fiber is attached to a stainless steel plunger sheathed by a protective needle.

Fiber holders are available for manual injection as well as for use with autosamplers. The holder protects the coated fiber, and controls exposure of the fiber during analyte adsorption and desorption. The holder is reusable indefinitely and accepts the replaceable fiber assembly. First time users must order both a holder and a fiber assembly. Fiber holders for use with an autosampler are also available.

Fiber Holder for Manual Sampling

An adjustable depth guide positions the fiber for sampling and for correct placement in the heated zone of the GC injection port. The fiber can be locked in the exposed position.



Cat. No.	Description	Qty.
57330-U	SPME Fiber Holder, for use with manual sampling	1 ea.

SPME Fiber Assemblies

SPME fiber assemblies can be reused for ≥ 100 analyses, depending on the application and the care they are given. For reuse, simply condition with heat before and after every analysis. Each assembly has a color-coded or notched hub indicating the type of coating on the fiber. Choose the appropriate assembly for the holder: manual or autosampler. The key to proper SPME performance is fiber selection.

For information on how to select a fiber, visit SigmaAldrich.com/spme

SPME Fiber Assortment Kit for Volatiles and Semivolatiles

Recommended starter kit for the extraction of volatile and semivolatile metabolites contains one fiber each of 85 μm polyacrylate coating, 100 μm polydimethylsiloxane coating, and 7 μm polydimethylsiloxane coating.

Cat. No.	For Use with	Needle	Qty.
57306	Manual holder	24 ga	1 kit
57307	Autosampler	24 ga	1 kit
57285-U	Autosampler	23 ga	1 kit

Achieve Sharper Peaks with SPME-GC Analyses Using Supelco® Inlet Liners

GC injection port liners are designed for optimal sample introduction for specific injection techniques. When using SPME, a 0.75 mm I.D. inlet liner increases linear velocity, compared to a conventional, larger volume 2 mm I.D. liner, and rapidly introduces analytes onto the column in a narrow band. To minimize sample loss or peak tailing, the inlet liner must be inert to minimize adsorption of active sample components. An inlet liner, in conjunction with efficient, solvent-free, SPME sample introduction, helps to achieve excellent chromatographic results. An inlet liner for several Agilent® GC systems is available.

For Agilent® (5890, 6890, and 7890)

Inlet Liner, Direct (SPME) Type, Straight Design (unpacked)

L × O.D. × I.D. _____ 785 mm × 65 mm × 0.75 mm

Cat. No.	Qty.
2637501	1 ea.

To select the appropriate inlet liner for your GC, visit SigmaAldrich.com/inletliners

GC Derivatization Reagents

A large number of reagents are used to prepare derivatives for gas chromatography. Derivatives are used for the following reasons:

- To improve resolution and reduce tailing of polar compounds (-OH, -COOH, =NH, -NH₂, -SH, and other functional groups)
- To analyze relatively nonvolatile compounds
- To improve analytical efficiency and increase detectability
- To improve stability of compounds

The following table lists the silylation reagents most commonly used together with acylation and alkylations.

Cat. No.	Description
43340	1,1,3,3-Tetramethyl-1,3-diphenyldisilazane
69649	4-(Trimethylsiloxy)-3-penten-2-one
14755	Bis(dimethylamino)dimethylsilane
15256	BSA+TMCS
76750	Chlorodimethyl(pentafluorophenyl)silane
90383	Chlorotriethylsilane
89595	Chlorotrimethylsilane
52619	Hexamethyldisilazane
01565	Hexamethyldisiloxane
91566	<i>N</i> -(Trimethylsilyl)acetamide
15235	<i>N,N</i> -Bis(trimethylsilyl)methylamine
89539	<i>N,O</i> -Bis(<i>tert</i> -butyldimethylsilyl)trifluoroacetamide
15269	<i>N,O</i> -Bis(trimethylsilyl)acetamide
15222	<i>N,O</i> -Bis(trimethylsilyl)trifluoroacetamide
15209,	<i>N,O</i> -Bis(trimethylsilyl)trifluoroacetamide with
15238	trimethylchlorosilane
69479	<i>N</i> -Methyl- <i>N</i> -(trimethylsilyl)trifluoroacetamide
69478	<i>N</i> -Methyl- <i>N</i> -(trimethylsilyl)trifluoroacetamide with 1% trimethylchlorosilane
00942	<i>N-tert</i> -Butyldimethylsilyl- <i>N</i> -methyltrifluoroacetamide with 1% <i>tert</i> -Butyldimethylchlorosilane
33036,	BSA Derivatization Grade
33035-U,	
33037-U	
33030,	BSA+TMCS+TMSI
33151,	
33031-U	
33149-U,	BSTFA + TMCS
33154-U,	
33155-U,	
33148	
33024,	BSTFA, Derivatization Grade
33027,	
33084	
33014	Chlorotrimethylsilane
33350-U	HMDS, Derivatization Grade
33038,	HMDS+TMCS+Pyridine
33039	
33175	Silica Column Regeneration Solution
505846	Silylation Sampler Kit
33065-U	Sylon CT
33092-U	<i>tert</i> -Butyldimethylsilylimidazole solution
33068-U	TMSI, Derivatization Grade
33156-U,	TMSI+Pyridine
33159-U	

To learn more, view the Derivatization Reagents for Selective Responses Guide at SigmaAldrich.com/derivatization

Solvents and Blends for LC-MS

LC-MS has become an important tool in today's analytical labs. In order to obtain accurate and reproducible results, high demands are made on the purity of chemicals. We offer high purity solvents specifically to meet the stringent requirements of LC-MS applications, ensuring high UV transmittance, baseline stability and lowest impurity levels.

We have developed and introduced high purity solvents pre-blended with acetic acid (HA), formic acid (FA) or trifluoroacetic acid (TFA) to provide ready-to-use mobile phases for LC-MS. With this comprehensive portfolio, we set the standard for accurate, reproducible and high-resolution analytical separations.

Features:

- Ready-to-use
- LC/MS suitability
- Minimal metal adduct formation
- Minimal ionization suppression
- Batch to batch consistency
- Filtered through 0.2µm

Benefits:

- Time- & cost-saving
- Reliable LC/MS application
- Less laborious mixing procedure
- Reduced contamination danger
- Safer - less exposure to hazardous chemicals
- No glassware cleaning
- Reduced solvent/acid excess
- Less storage room needed

Cat. No	Name	Description	Pkg Size
159004.2500	Acetonitrile + 0.1% Acetic acid (v/v)	Hypergrade for LC-MS LiChrosolv®	2.5 L
159002.2500	Acetonitrile + 0.1% Formic acid (v/v)	hypergrade for LC-MS LiChrosolv®	2.5 L
159014.2500	Acetonitrile + 0.1% Trifluoroacetic acid (v/v)	hypergrade for LC-MS LiChrosolv®	2.5 L 4 L
159007.2500	Water + 0.1% Acetic acid (v/v)	hypergrade for LC-MS LiChrosolv®	2.5 L
159013.2500	Water + 0.1% Formic acid (v/v)	hypergrade for LC-MS LiChrosolv®	2.5 L
480112.2500 480112.4000	Water + 0.1% Trifluoroacetic acid (v/v)	hypergrade for LC-MS LiChrosolv®	2.5 L 4 L
100029.1000 100029.2500 100029.9010 100029.9030	Acetonitrile	hypergrade for LC-MS LiChrosolv®	1 L GL* 2.5 L GL* 10 L ST 30 L ST
106035.1000 106035.2500	Methanol	hypergrade for LC-MS LiChrosolv®	1 L GL* 2.5 L GL*
115333.1000 115333.2500 115333.4000 115333.9010 115333.9030	Water	hypergrade for LC-MS LiChrosolv®	1 L GL* 2.5 L GL* 4 L GL* 10 L ST 30 L ST
103649.1000 103649.2500	Ethyl acetate	hypergrade for LC-MS LiChrosolv®	1 L 2.5 L
103701.1000 103701.2500	Hexane	hypergrade for LC-MS LiChrosolv®	1 L 2.5 L
103654.1000 103654.2500	Heptane	hypergrade for LC-MS LiChrosolv®	1 L 2.5 L
102781.1000 102781.2500	2-Propanol	hypergrade for LC-MS LiChrosolv®	1 L 2.5 L

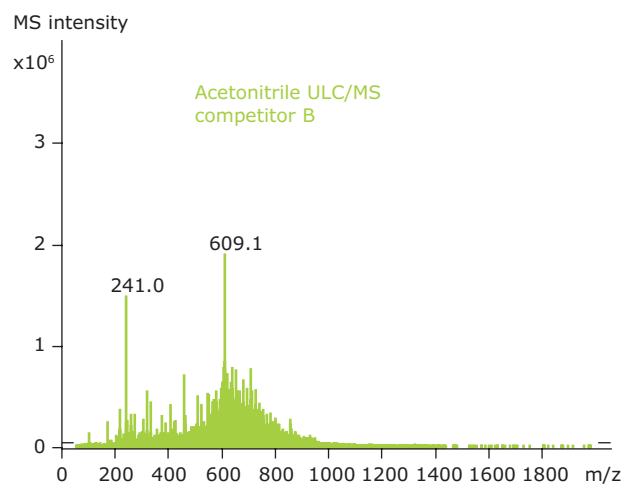
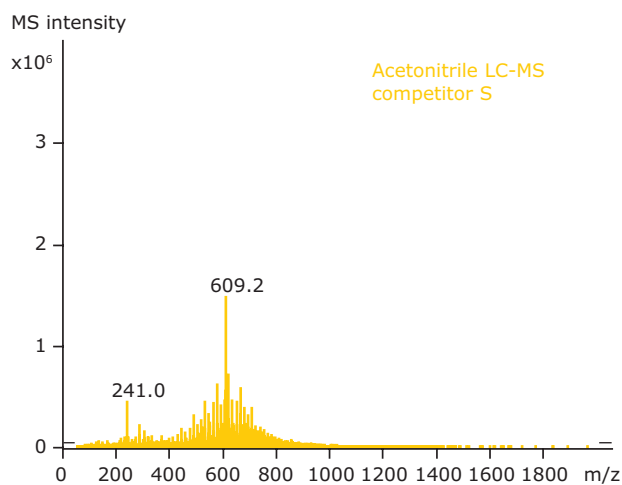
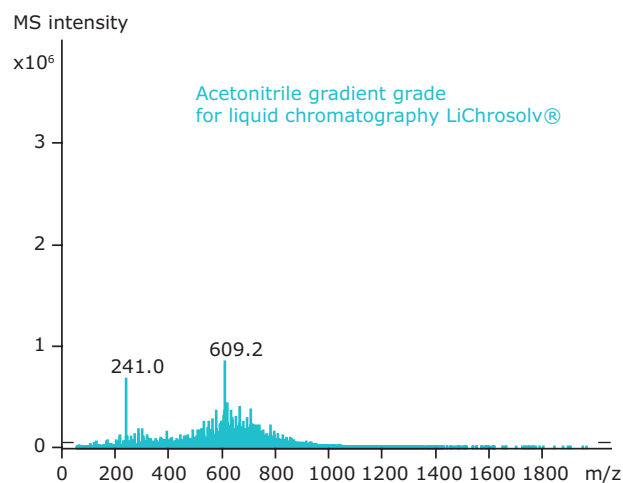
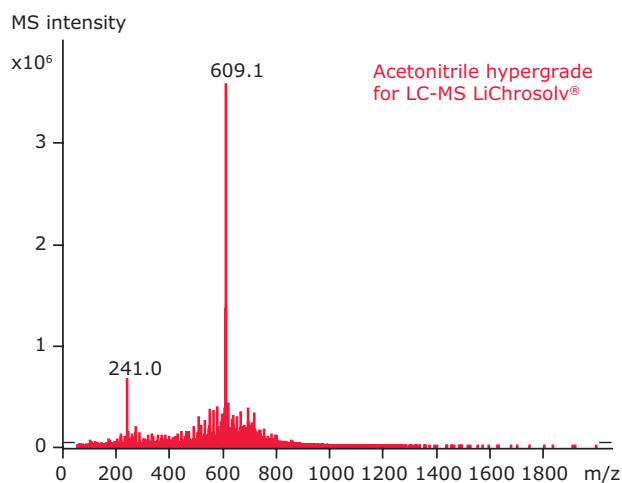
* special treated amber glass bottle

All solvents are filtered through 0.2 µm. | GL = glass bottle | ST = stainless steel returnable barrel

MS conditions

System	Bruker Esquire 3000+ ion trap MS
Detection	Pos. ESI-MS, m/z range 50 – 2000
Flow rate	0.2 mL/min via syringe pump
Temperature	25°C
Sample	Reserpine (m/z 609.1), internal standard (m/z 241.0)

The mass spectra of four different acetonitrile grades clearly show the variation in the intensity of the reserpine signal ($[M+H]^+ = 609$) as well as the extent of the background signals. The differences in the intensity of the reserpine signal are caused by ion suppression. This effect occurs due to interfering trace contaminants that can be present in acetonitrile.



Mass spectra displaying the results of the reserpine test of different acetonitrile qualities from Merck and two alternative competitors.

LiChropur® LC-MS Reagents

It is common practice in LC-MS to add certain chemicals to the mobile phase or introduce them post-column prior to the interface to influence analyte ionization. Most often, an improvement in the analyte signal is the goal. However, some additives may be used to suppress unwanted signals or selectively enhance the signal of particular compounds in a mixture, for example glycosidic species in a mixture of peptides.

We offer a wide range of high purity mobile phase additives for LC-MS applications. Our offer includes the most commonly used acids, bases and volatile salts. All are of high purity and are rigorously tested for LC-MS application suitability, offering many advantages for both small and large molecule analysis.

Impurities, such as alkali ions, plasticizers and surfactants, found in lower-grade reagents are

particularly problematic as they interfere strongly with LC-MS, resulting in higher background noise and formation of adducts. Only highly pure reagents allow high signal-to-noise ratios.

Features:

- LC-MS application tested for consistent quality
- Improves ionization and resolution
- Extremely low levels of inorganic and organic impurities
- Manufactured specifically for accurate and fast LC-MS
- Highest quality acids, bases & salts

For more information, visit

SigmaAldrich.com/lcms-reagents

Cat No.	Product Name	Description	Package Size
5.33001.0050	Acetic acid	100% for LC-MS LiChropur®	50 ml
5.33002.0050	Formic acid	98-100% for LC-MS LiChropur®	50 ml
5.33003.0050	Ammonia solution	25% for LC-MS LiChropur®	50 ml
5.33004.0050	Ammonium acetate	for LC-MS LiChropur®	50 ml
5.33005.0050	Ammonium hydrogen carbonate	for LC-MS LiChropur®	50 ml

LC-MS Certified Spiking Solutions and Reference Materials

Single and multi-component solution standards (both stable-labeled and unlabeled) designed, manufactured and tested specifically for use as reference standards for laboratories performing bioanalysis, therapeutic drug monitoring, diagnostic and toxicology testing. Cerilliant's products address the stringent and complex requirements of forensic toxicology, clinical toxicology, clinical chemistry/ immunoassay, therapeutic drug monitoring, pain management and pharmaceutical analysis. Products manufactured at Cerilliant® are fully documented through the use of batch records to provide traceability of materials used, traceability of equipment utilized, calibration records, and detail of all work performed and staff utilized, all backed by a comprehensive Certificate of Analysis. Cerilliant's quality credentials include accreditations to ISO Guide 34, ISO/IEC 17025 and certification to ISO 13485 and ISO 9001. Cerilliant's quality system incorporates cGMP and GLP requirements.

Cerilliant's portfolio includes:

Catalog and Custom

- Metabolites including P450 and Glucuronides
- Impurities / Degradants
- Internal Standards

- Many analyte classes, including
 - Drugs / Drugs of Abuse
 - TDM Drugs / Immunosuppressants / Catecholamines
 - Hormones including Thyroid / Steroids – Alcohol / Ethanol
 - Vitamins (A, B, D and E)
 - Natural Products / Phytochemicals



For a complete listing of Cerilliant certified standards, visit

SigmaAldrich.com/cerilliant

ZipTip Pipette Tips: Proteomics Sample Prep in Seconds

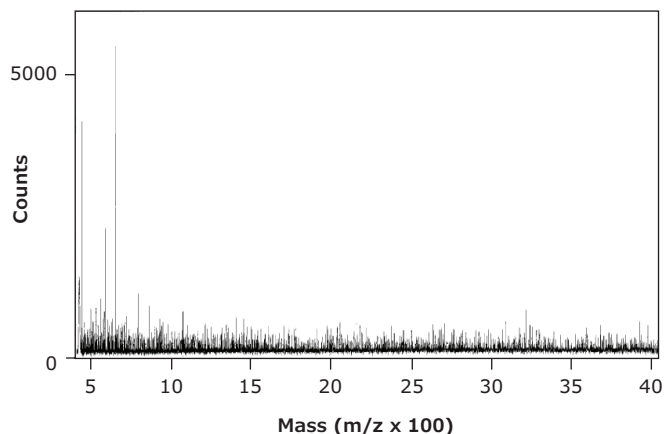


A staple of every mass spectrometry lab, ZipTip® is a 10 µL pipette tip with a 0.6 or 0.2 µL bed of chromatography media fixed at its end with no dead volume. It is ideal for concentrating and purifying peptides or proteins in seconds prior to mass spectrometry, HPLC, and capillary electrophoresis. The ZipTip® pipette tip provides a reproducible, high recovery method for concentrating, purifying or even fractionating femtomoles to picomoles of peptides, proteins and oligonucleotides for improved data quality.

ZipTip® Advantages:

- Single-step desalting, concentration, and purification
- Fractionate complex samples for more meaningful data
- Ideal for peptides, proteins, nucleic acids, and more
- No dead volume for maximum recovery
- Eliminates time-consuming chromatography

A. Direct Spotting



ZipTips® increase sensitivity of mass spectrometric analysis. MALDI MS spectra of a tryptic peptide digest from an in-gel 2D digest. The top spectrum represents a contaminated sample prior to sample clean-up. The lower spectrum represents the sample after treatment with a ZipTip®C18 prior to MALDI-ToF MS analysis.

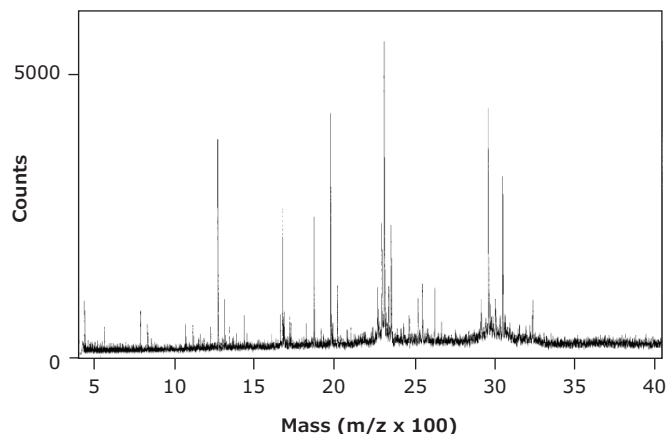
The ZipTip pipette tip is simple and easy to use.

- Place the tip on a single- or multi-channel pipettor, standard 22-gauge blunt-end HPLC needle, or compatible automated liquid handling/sample preparation station.
- For sample binding, aspirate and dispense through the media several times.
- Similarly, aspirate and dispense appropriate wash solvent to remove contaminants.
- The concentrated, purified sample is eluted in 1–4 µL of compatible solvent with direct transfer to a mass spectrometer target, vial, or other analysis step.

For applications requiring smaller elution volumes (e.g., <1 µL), a micro-bed format containing 0.2 µL of media is available.

Cat. No.	Description	Qty/Pk
ZipTip® Pipette Tips		
ZTSCXS008	ZipTip® with 0.6 mL strong cation resin	8
ZTSCXS096	ZipTip® with 0.6 mL strong cation resin	96
ZTC04S008	ZipTip® with 0.6 mL C4 resin	8
ZTC04S096	ZipTip® with 0.6 mL C4 resin	96
ZTC04S960	ZipTip® with 0.6 mL C4 resin	960
ZTC18S008	ZipTip® with 0.6 mL C18 resin	8
ZTC18S096	ZipTip® with 0.6 mL C18 resin	96
ZTC18S960	ZipTip® with 0.6 mL C18 resin	960
ZTC18M008	ZipTip® with 0.2 mL C18 resin	8
ZTC18M096	ZipTip® with 0.2 mL C18 resin	96
ZTC18M960	ZipTip® with 0.2 mL C18 resin	960

B. After ZipTip®µ-C18



To place an order or receive technical assistance, please visit:

SigmaAldrich.com/order

For all locations across Europe and the world, please visit:

SigmaAldrich.com/offices

For technical service email: ets@sial.com

merckgroup.com/life-science

