

Unparalleled Peptide Synthesis

cempeptides.com









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CEM Overview

Innovations in Microwave Peptide Synthesis

- **1978** CEM Corporation founded as a new company, based on microwave laboratory instrumentation
- **2001** CEM launches a single mode microwave system for chemical synthesis
- **2003** CEM develops the world's first automated microwave peptide synthesizer¹
- **2007** CEM publishes research for optimized methods for aspartimide formation and epimerization under microwave SPPS²
- 2013 Liberty Blue™ peptide synthesizer developed based on High Efficiency Solid Phase Peptide Synthesis (*HE*-SPPS)
- 2014 HE-SPPS methodology published³
- 2016 CEM launches new universal load resins eliminating the need for pre-loaded resins historically used
- **2016** CEM offers the world's first large-scale microwave peptide synthesis, with capabilities of up to 500 grams of a purified peptide, in a single batch
- 2016 CEM develops improved carbodiimide coupling methods for peptide synthesis at elevated temperature (CarboMAX[™])
- 2017 CEM develops a novel one-pot coupling/ deprotection process reducing SPPS cycle time and waste usage (Liberty PRIME[™])





Founding Fathers (circa 1980)

Chemist: Dr. Michael J. Collins (**Middle**) Electrical Engineer: Ron Goetchius (**Left**) Mechanical Engineer: Bill Cruse Jr. (**Right**)

- ¹ Collins J.M., Collins M.J., Steorts R.C. Biopolymers 71, 361 2003
- ² Palasek S., Cox Z., Collins J. J. Pept. Sci. 13, 143-148 2007
- ³ Collins, J., Porter K., Singh S., Vanier G. Org. Lett. 16, 940-943 2014



Corporate Legacy

Founded in 1978 by our current CEO, Dr. Michael J. Collins, CEM has pioneered the field of microwave chemistry. For nearly 40 years we have designed and developed laboratory instrumentation and scientific methods (both microwave-based and non-microwave technologies) that are used by major companies, prestigious research institutes, and universities around the world. The company's major products provide unique solutions for compositional analysis of food and chemical samples, acid digestion for elemental analysis, and chemical synthesis of peptides and small molecules. CEM is a private company with global headquarters outside Charlotte, North Carolina, along with offices in England, Germany, Japan, France, Italy, Singapore, and Ireland. The company's annual revenue is > 90M USD (2017) with more than 300 employees worldwide. Since 2003, CEM has pioneered the area of microwave peptide synthesis. The company sells an elite line of microwave-based peptide synthesizers, based on unique high efficiency solid phase peptide synthesis (*HE*-SPPS) which provides unmatched purity, ultra-fast cycle times, and up to a 90% reduction in total waste, compared to traditional technologies. More than 600 peptide synthesizers from CEM have been installed throughout the world.

HE-SPPS

High Efficiency Solid Phase Peptide Synthesis (*HE*-SPPS)

HE-SPPS is a significant advancement for solid phase peptide synthesis. It originates from our pioneering work in developing microwave assisted SPPS, introduced at the 2003 American Peptide Symposium.¹ At this time, we introduced a new process for making peptides, based on the use of microwave energy for both the deprotection and coupling steps in SPPS. This technology has demonstrated improvements for thousands of peptides with CEM's peptide synthesis instrumentation.² To support microwave SPPS, we have also utilized in-situ fiber optic temperature monitoring to provide true internal solution temperature control. This is essential for fast reaction heating with temperature control, as it is well known that the outside of a reaction vessel can be at a significantly different temperature than the inside.³

In 2013, we developed an improved methodology for microwave SPPS, based on the use of higher temperature carbodiimide based coupling at 90 °C, along with the elimination of all washing after each coupling step.⁴ These insights led to significant time and solvent savings, while providing peptides of incredibly high purity. The more acidic coupling environment with carbodiimide chemistry overcomes coupling issues for cysteine (epimerization) and arginine (y-lactam formation), which were previously an issue under more basic coupling conditions (ex. HCTU/DIEA). The instrumentation design used on CEM's Liberty Blue[™] peptide synthesizer is also a critical component of HE-SPPS to eliminate inefficient internal fluidic and reagent path cleaning that increases waste generated. HE-SPPS used on the Liberty Blue is now used in hundreds of laboratories worldwide and provides very fast, high purity peptides with incredibly low waste generated.

(1) Collins, J.M., Collins, M.J., and Steorts, R.C., "A Novel Method for Solid Phase Peptide Synthesis Using Microwave Energy" Biopolymers, 71, 361 2003.

(2) US7393920; US7582728; US8058393; JP4773695

(3) M. Herrero, J. Kremsner and C.O. Kappe, "Nonthermal Microwave Effects Revisited: On the Importance of Internal Temperature Monitoring and Agitation in Microwave Chemistry," J. Org. Chem., vol. 73, pp. 36-47, 2008.

(4) J. Collins, K. Porter, S. Singh and G. Vanier, "High-Efficiency Solid Phase Peptide Synthesis (*HE*-SPPS)," Org. Lett., vol. 16, pp. 940-943, 2014.





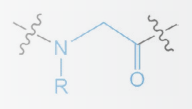
Selected Peptides Synthesized with HE-SPPS

Peptide	Sequence	UPLC Purity	Synthesis Time	Total Waste (mL)
65-74ACP	VQAAIDYING	93%	44 m	154 mL
JR-10mer	WFTTLISTIM-NH ₂	67%	49 m	170 mL
ABRF 1992	GVRGDKGNPGWPGAPY	82%	1 h 37 m	272 mL
ABC-20mer	VYWTSPFMKLIHEQCNRADG-NH ₂	73%	2 h 7 m	340 mL
Thymosin	SDAAVDTSSEITTKDLKEKKEVVEEAEN-NH ₂	61%	2 h 11 m	468 mL
¹⁻⁴² β-amyloid	DAEFRHDSGYEVHHQKLVFFAEDVGSNKGAIIGLMVGGVVIA-NH ₂	72%	3 h 49 m	1019 mL

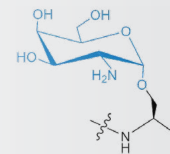
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- Branched Peptides
- Cyclized Peptides
- Disulfide Bonding
- Glycopeptides
- High Throughput Synthesis
- N-Methyl Peptides
- Peptide Thioesters
- Peptoids
- Phospho-peptides
- PNA

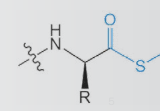


НО



Lvs

Lvs



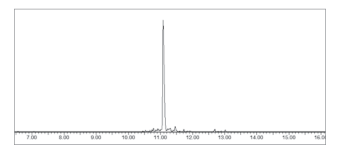


CarboMAX[™] (Enhanced Carbodiimide Chemistry)

Faster Coupling: Improved Purity with Less Epimerization

Coupling with carbodiimide chemistry has significant benefits over aminium/phosphonium salts (ex. HATU, HCTU, PyBOP) at elevated temperature. This includes major reductions of epimerization for cysteine and γ -lactam formation of arginine. However, activation by carbodiimides is relatively slow. We developed an improved coupling process which allows for faster formation of the key 0-acylisourea intermediate by increasing the amount of carbodiimide to 2 equivalents relative to the amino acid.¹

By forming more activated amino acid faster than standard carbodiimide chemistry the subsequent coupling will happen quicker. This provides not only a faster coupling time, but also less epimerization from less time as a sensitive activated amino acid. This methodology termed CarboMAX is patent pending and exclusively licensed for use on CEM's peptide synthesizers.



UPLC-MS Analysis of crude Liraglutide (CarboMAX)

Reduced Epimerization (ex. Liraglutide)

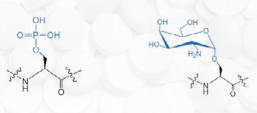
Epimer	DIC/Oxyma (%)	CarboMAX (%)
D-Asp	0.23	0.31
D-Ala	0.33	0.25
D-Arg	0.29	0.2
D-Glu	0.39	0.3
D-His	N/A	N/A
D-lle	< 0.10	< 0.10
L-allo lle	< 0.10	< 0.10
D-allo lle	< 0.10	< 0.10
D-Leu	0.17	0.13
D-Lys	< 0.10	0.1
D-Phe	0.2	0.16
D-Ser	0.16	0.12
D-Thr	< 0.10 < 0.10 < 0.10	< 0.10
D-Trp	0.24	< 0.10
D-Tyr	0.12	0.11
D-Val	< 0.10	< 0.10

¹ Patent Pending: US15686719; EP17188963.7

Improved Purity

Peptide	Sequence	% Crude Purity DIC/Oxyma	% Crude Purity CarboMAX
Thymosin	SDAAVDTSSEITTKDLKEKKEVVEEAEN	63	75
GRP	VPLPAGGGTVLTKMYPRGNHWAVGHLM	62	74
Bivalirudin	fPRPGGGGNGDFEEIPEEYL	80	82
¹⁻³⁴ PTH	SVSEIQLMHNLGKHLNSMERVEWLRKKLQDVHNF	67	85
³⁵⁻⁵⁵ MOG	MEVGWYRSPFSRVVHLYRNGK	77	91
Magainin 1	GIGKFLHSAGKFGKAFVGEIMKS	71	79
Dynorphin A	YGGFLRRIRPKLKWDNQ	74	82
Liraglutide	HAEGTFTSDVSSYLEGQAAK(γ-Glu-palmitoyl) EFIAWLVRGRG	74	88

Stabilizing acid-sensitive linkages



Phosphopeptides

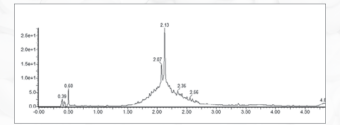
Glycopeptides

Many important side chain modifications are sensitive to acidic activators, such as Oxyma Pure and HOBt used under elevated temperature. With traditional carbodiimide chemistry, this can lead to undesirable cleavage of sensitive groups, such as phospho and O-linked carbohydrates.

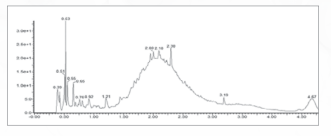
We developed a patented process of incorporation of < 1 equivalent base to stabilize these linkages while using carbodiimide chemistry at elevated temperature. This method which is part of CarboMAX chemistry is only available on CEM's peptide synthesizers and allows access to synthesis of these peptides at high temperatures with unmatched speed and purity.

³ Patent Pending: US20160176918; EP3037430; JP2016138090; CN105713066; AU2017204172

Multi-phosphorylated peptide -TpTGNGKPVECpSQPESSFKMpSFKR



CarboMAX Synthesis (Fmoc-AA/DIC/Oxyma Pure/DIEA) – 1/1/1/0.4



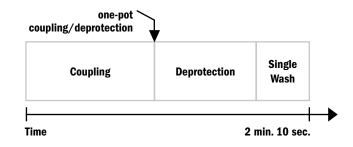
Standard Synthesis (Fmoc-AA/DIC/Oxyma Pure) – 1/1/1

One-Pot Coupling/Deprotection

Unparalleled speed and efficiency

Traditional solid phase peptide synthesis involves the use of iterative and separate deprotection and coupling steps with washing in-between. This is based on the assumption that undesirable amino acid insertions can occur without complete draining and washing between each step. In 2013, it was demonstrated that washing after the coupling step can be eliminated without effect on peptide purity.¹

The Liberty PRIME[™] takes this further by using a new onepot coupling and deprotection process.² This technique involves addition of the deprotection reagent (base) directly to the undrained post-coupling mixture. The ability to do this is based on the insight that faster reaction kinetics in the solution phase promote rapid hydrolysis or selfcondensation of the active ester, thereby avoiding potential side reactions at the resin bound amino functionality. The Fmoc removal then proceeds uninterrupted at elevated temperature. An optimized use of reagents results in an essentially neutral reaction mixture towards the end of deprotection step. This new procedure offers several advantages such as (a) approximately 90% reduction in solvent requirement for the deprotection step, (b) 75% reduction in solvent requirement for post-deprotection washings, (c) faster deprotection step since the microwave ramp time is not needed, and (d) shorter cycle time due to absence of the post-coupling drain step.



Utilization of the one-pot coupling/deprotection methodology requires the ability to consistently add precise small volumes of concentrated base. To achieve this, the Liberty PRIME incorporates a new dedicated pumping module with the ability to rapidly add the deprotection reagent precisely at the end of the coupling step in volumes as low as 0.25 mL. The pre-calibrated pump module does not require on-going calibration thereby avoiding drifting delivery amounts. Additionally, the main solvent and the activator (Oxyma Pure) are also delivered through similar individual pumps within the module for improved performance.

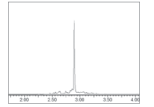
(1) J. Collins, K. Porter, S. Singh and G. Vanier, "High-Efficiency Solid Phase Peptide Synthesis (HE-SPPS)," Org. Lett., vol. 16, pp. 940-943, 2014.

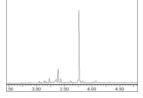
(2) Patent Pending: US20170226152; W02017070512

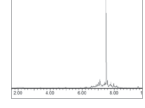


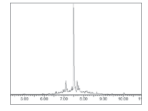
Peptide Synthesis on the Liberty PRIME³

Peptide	Sequence	Crude Purity (UPLC-MS)	Total Synthesis Time	Total Chemical Waste
⁶⁵⁻⁷⁴ ACP	VQAAIDYING-NH ₂	94%	25 m	92 mL
ABC-20 mer	VYWTSPFMKLIHEQCNRADG-NH ₂	83%	48 m	172 mL
JR-10 mer	WFTTLISTIM-NH ₂	70%	25 m	92 mL
Exenatide	$HGEGTFTSDLSKQMEEAVRIFIEWLKNGGPSSGAPPPS-NH_2$	57%	1 h 36 m	273 mL
⁷⁻³⁷ GLP1	HAEGTFTSDVSSYLEGQAAKEFIAWLVKGRG	47%	1 h 14 m	217 mL
PnIA(A10L)	GCCSLPPCALNNPDYC-NH ₂	77%	43 m	112 mL
Circulin A	GIPCGESCVWIPCISAALGCSCKNKVCYRN	79%	1 h 10 m	252 mL
Parigidin-br-1	GGSVPCGESCVFIPCITSLAGCSCKNKVCYYD	74%	1 h 14 m	264 mL









ABC-20 mer

JR-10 mer

Exenatide

7-37GLP1

(3) Refer to CEM Application Note, "Liberty PRIME – Ultrafast Peptide Synthesis at Elevated Temperature"

Minimal Epimerization

The potential for epimerization was then investigated on the elevated temperature coupling methods used on the Liberty PRIME. In particular, cysteine and histidine are known to be sensitive to epimerization during coupling. The epimerization level was therefore investigated through a well-known standard method involving hydrolysis, subsequent derivatization, and gas chromatography analysis (C.A.T. GmbH). Epimerization levels observed with HBTU/DIEA activation at room temperature were found to be higher than those from 90 °C standard or CarboMAX couplings, as well as from 105 °C CarboMAX coupling on the Liberty PRIME. Use of Fmoc-His(Boc)-OH instead of Fmoc-His(Trt)-OH allowed coupling temperatures of 90 °C or 105 °C without any increase in epimerization levels. These results further demonstrate that standard HE-SPPS or CarboMAX coupling methods are particularly wellsuited for peptide synthesis at elevated temperature.

Epimerization levels of cysteine and histidine in ABC 20mer

Epimer	Conventional RT - HBTU/DIEA	Liberty Blue 90 °C CarboMAX	Liberty PRIME 105 °C CarboMAX
D-His	1.79% ^{1,a}	1.12% ^b	1.05% ^b
D-Cys ^c	1.38% ¹	0.64%	0.68%

^aFmoc-His(Trt)-OH; ^bFmoc-His(Boc)-OH; ^cFmoc-Cys(Trt)-OH

Sequential vs Parallel

Unparalleled advantages

Sequential Peptide Synthesis Advantages

- \cdot Single peptides synthesized fast
- Faster Purification Workflow (purify as you go)
- Precise control at each step
- Higher purity peptides
- Simple to use & maintain instrumentation





Resin Loader Advantages

Optional modular resin loading options are available, which allows up to 24 peptides to be synthesized sequentially. The resin loader comes in a 12-position module with an additional 12-position module that can be coupled to it.

With rapid cycle times, and the ability to synthesize 12 - 24 peptides unattended, the Liberty Blue HT12TM and the Liberty PRIMETM provide exceptional throughput compared to conventional parallel systems, but with considerably better purity.

Synthesizer Comparison

Pick the synthesizer that's best for you.

	Liberty Lite [™]	Liberty Blue [™]	Liberty Blue HT12 [™]	Liberty PRIME [™]
Cycle Time	15 minutes	4 minutes	4 minutes	2 minutes
Waste/Cycle	40 mL (0.1 mmol)	16 mL (0.1 mmol)	16 mL (0.1 mmol)	8.5 mL (0.1 mmol)
Scale Range	0.005 - 5 mmol	0.005 - 5 mmol	0.005 - 5 mmol	0.005 - 5 mmol
Peptide Positions	1	1	12	24
Amino Acid Positions	20	27	27	27
Other Positions	4	4	4	4
Fluid Delivery	Flex-Add: Amino Acids & Activators Timed Delivery: Wash & Deprotection	Flex-Add: Amino Acids & Activators Timed Delivery: Wash & Deprotection	Flex-Add: Amino Acids & Activators Timed Delivery: Wash & Deprotection	Flex-Add: Amino Acids Pre-Calibrated Pumps: Wash, Deprotection, &
Dimensions	· 20" W x 18" D x 30" H · 51 cm x 46 cm x 76 cm	· 20" W x 18" D x 30" H · 51 cm x 46 cm x 76 cm	· 27" W x 18" D x 30" H · 69 cm x 46 cm x 76 cm	Oxyma Pure • 42" W x 18" D x 30" H • 107 cm x 46 cm x 76 cm
Upgrades	Upgrade to the Liberty Blue	· HT12 · HT24	· HT24	N/A
Accessories	Razor Cleavage System	 Integrated Camera, Razor Cleavage System Flex-Add Large Scale 	Integrated Camera Razor Cleavage System Flex-Add Large Scale	Integrated Camera Razor Cleavage System

Discover Bio[™]

Manual Microwave Peptide Synthesizer

Liberty Lite[™]

Microwave Peptide Synthesizer



Overview

The world's best selling microwave peptide synthesizer is also available in a research scale manual system that offers a costeffective alternative to purchasing peptides. Enhance your laboratory's capabilities with the benefits of microwave-assisted peptide synthesis.

Features

- Integrated module for washing and adding deprotection
- Easy access port for addition of activated amino acids
- True Internal fiber-optic temperature control
- 0.005 1 mmol scale range
- Ability to upgrade to Liberty Blue

Chemistry Technology

· CarboMAX



Overview

An entry-level option for the globally recognized Liberty Blue technology. The Liberty Lite provides advantages over existing peptide synthesizers.

Features

- Flex-Add[™] critical reagent delivery system (patented)
- True Internal fiber-optic temperature control
- · 20 amino acid positions
- 0.005 5 mmol scale range
- Ability to upgrade to Liberty Blue

Chemistry Technology

· CarboMAX

Liberty Blue[™] Microwave Peptide Synthesizer

Liberty Blue HT12[™]

High-Throughput Microwave Peptide Synthesizer



Overview

The Liberty Blue Automated Microwave Peptide Synthesizer is the gold standard for peptide synthesis. It features unmatched 4-minute cycle times along with a 90% solvent reduction based on High Efficiency Solid Phase Peptide synthesis (*HE*-SPPS), developed in 2013.

Features

- Flex-Add[™] critical reagent delivery system (patented)
- True Internal fiber-optic temperature control
- · 27 amino acid positions
- 0.005 5 mmol scale range
- Integrated Camera (optional)
- Ability to upgrade to Liberty Blue HT12

Chemistry Technology

- HE-SPPS
- CarboMAX



Overview

The Liberty Blue HT12 features all the advantages of the Liberty Blue with the added capability of the HT12 resin loader. This allows for the automated sequential synthesis of up to 12 different peptides.

Features

- Flex-Add[™] critical reagent delivery system (patented)
- True Internal fiber-optic temperature control
- · 27 amino acid positions
- 0.005 5 mmol scale range
- Integrated Camera (optional)

Chemistry Technology

- HE-SPPS
- CarboMAX



Liberty PRIME[™]

High-Throughput Microwave Peptide Synthesizer



Overview

The Liberty PRIME peptide synthesizer is the most advanced platform available for microwave peptide synthesis. It features a revolutionary one-pot deprotection and coupling process, allowing for a remarkable **2-minute cycle time**, with only **8.5 mL waste per cycle** (at 0.1 mmol).

- Individual 20-mers every 45 minutes
- Batches of 24 peptides (20-mers) every 20 hours, with only 4.5 liters total waste produced

Chemistry Technology

- One-Pot Coupling/Deprotection
- · CarboMAX

Performance

Scale	Cycle Time	AA Equiv.	Waste/Cycle
0.1 mmol	2 m 10 s	5	8.5 mL
0.3 mmol	3 m 40 s	5	20 mL
0.4 mmol	3 m 50 s	4	20 mL

Enhanced Hardware

The Liberty PRIME features enhanced delivery options of the main solvent and Deprotection solutions compared to the Liberty Blue system. These reagents are delivered by a pre-calibrated pumping system, not requiring calibration or affected by restriction changes in the delivery path. This reduces system maintenance and provides an ideal system for GMP environments that is free of calibration.

Accessories & Upgrades

Make your synthesizer even better.

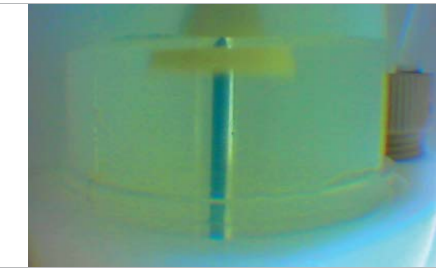
	68%
JR 10mer: WFTTLISTIM-NH ₂	
Scale: 1.0 mmol	
Cycle Time: 10 min	
man hand hand	hannen

Flex-Add (Large Scale)

Designed for routine syntheses at larger scales (≥ 0.5 mmol), this upgrade allows for optimized deliveries of larger volumes. At these scales, time savings $\geq 40\%$ are achieved.

Integrated Camera

The newly introduced camera for the Liberty Blue system enables researchers to monitor microwave peptide synthesis as never before. This fully integrated accessory provides complete visibility of the reaction vessel at any time with 5 mega-pixel quality (2560 x 1920) and 720p video (1280 x 720) capability. This option is beneficial for optimization of synthetic methods and system troubleshooting.





Pressure Rated Bottles

Pressure rated, stainless steel bottles incorporating visual liquid level detection. Available in 2 L, 5 L, 10 L, and 20 L sizes (GL45 cap size).

Razor®

Rapid Parallel Peptide Cleavage At Elevated Temperature

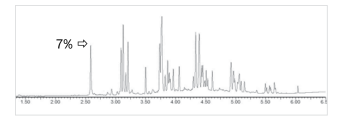


Overview

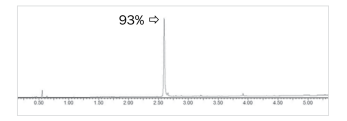
Cleave up to 12 peptides in only 30 minutes. The Razor features a compact design that easily fits in a fume hood and allows for temperature at +/- 1 °C control for up to 12 different vessels. Cleavage is typically complete in 30 minutes for standard peptides and allows for draining into a centrifuge tube for subsequent centrifugation. This system is ideal for both single peptides and large batches.

- Elevated Temperature Cleavage Block With +/- 1 °C Control
- Valve Control For Independently Draining Each Vessel
- Convenient Tray For Holding & Transporting Each Collection Tube
- Compact Design That Easily Fits In Standard Fume Hoods

Peptide: Fmoc-YGRKKRRQRRR Conditions: TFA/TIS/H20/DODT (92.5/2.5/2.5/2.5)



30 minutes, room temperature

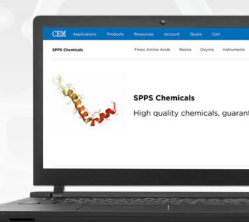


30 minutes, Razor

SPPS Chemicals

High quality chemicals, available online.

CEM offers a complete suite of peptide synthesis reagents for optimized SPPS, using microwave irradiation. This includes a complete library of standard and unique, high-quality Fmoc amino acids, PEG and polystyrene resins, and the powerful Oxyma Pure activator. Using CEM's unique high-quality reagents provides the highest purity peptides, with CEM's innovative methodology and instrumentation.



Fmoc N CO₂H

Fmoc Amino Acids

Extremely high quality at an affordable price.

Overview

Using Fmoc amino acids of lower quality can have a significant impact on peptide purity and yield, resulting in hard to separate impurities and even total synthesis failures. CEM's Fmoc amino acids are the highest quality available on the market and provide the best purities and yields possible for peptide synthesis.

Standard Specifications

- HPLC purity \geq 99.0%
- Enantiomeric purity \geq 99.8%
- 100% fully synthetic amino acids
- Continuously used and tested in CEM's peptide synthesis laboratory



Pre-weighed

Eliminate your weighing step by using amino acids that have been pre-weighed specifically for your Liberty system.



Full Library

A catalogue of Fmoc amino acids is available for synthesizing standard and modified peptides, for use with any peptide synthesizer.



Oxyma Pure

The perfect activator for elevated temperature.

Overview

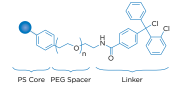
Oxyma Pure used with DIC produces peptides with increased yield and decreased epimerization, when used as an alternative to HOBt.¹ This safe, non-explosive auxiliary nucleophile works with carbodiimide coupling strategies to provide the best results for a peptide synthesis. Additionally, the use of DIC/Oxyma avoids side reactions associated with high levels of base (\geq 1 equiv. DIEA), using onium salt methods such as HBTU/DIEA.

¹R. Subirós-Funosas, et al. (2009) Chem. Eur. J., 15, 9394.



Overview

A full library of PEG and polystyrene resins for SPPS. CEM's SPPS resins are of the highest quality and optimized for the synthesis of standard and difficult peptides, with a variety of linkers.



ProTide™ Resins

Based on a PEG-PS core with optimal swelling, ProTide is recommended for synthesis of very long and difficult peptides.

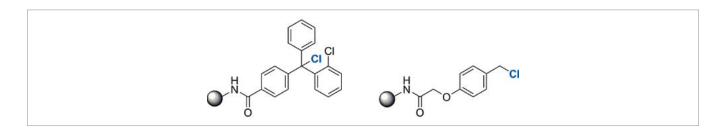


PS (1% DVB)

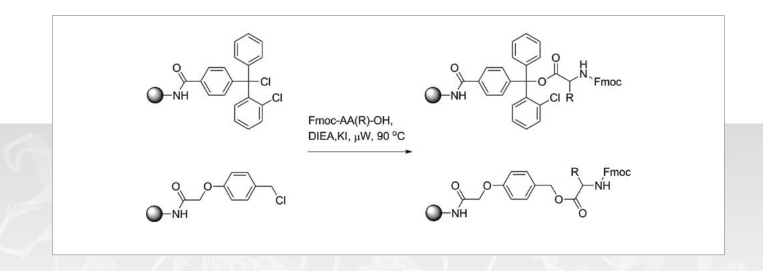
Polystyrene Resins

High quality, preloaded, polystyrene resins are great for synthesis of standard and difficult peptides.

Optimized PEG Resin Core with CI-TCP(CI) and CI-MPA Universal Linkers



ProTide resins contain an ideal PEG and polystyrene core, leading to an optimized environment for the synthesis of difficult peptides, with excellent swelling properties. New CI-TCP(CI) and CI-MPA linkers incorporated onto ProTide, eliminate the necessity for purchasing resins with pre-loaded C-terminal amino acids. The C-terminal amino acid reacts with the linker-chloride, in the presence of potassium iodide (KI)¹, N,N-diisopropylethylamine (DIEA), and microwave irradiation. The process is automatically carried out on CEM's microwave peptide synthesizers, using pre-programmed methods in the software. The result, any amino acid can be loaded on the resins in 10 minutes.

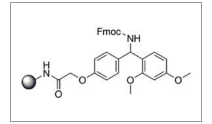


Key Advantages:

- Automated, high-temperature loading procedure complete in 10 min, whereas room temperature takes up to 24 hours
- Avoids coupling reagents; therefore, eliminating epimerization and dipeptide formation that can occur during loading
- No longer need to buy/store > 20 different, pre-loaded acid-linked resins

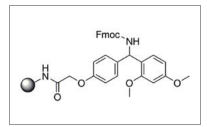
- Exhibit strong stability towards hydrolysis during storage and handling
- TCP(CI) is hyperacid sensitive and will produce protected peptides with 1% TFA/DCM and further minimizes diketopiperazine and 3-(1-piperidinyl) alanine formation²

¹Sandhya K., Ravindranath B. Tetrahedron Lett. 49, 2435 2008
 ²Heinlein C., Silva D., Tröster A., Schmidt J., Gross A., Unverzagt C. Angew. Chem. 50, 6406 2011



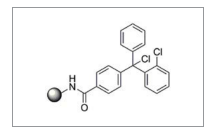
Rink Amide ProTide Resin (LL)

The ultimate resin recommended for longer and more difficult sequences of peptide amides. Based on ideal swelling properties from a TentaGel[®] core, incorporating PEG PS with a loading of 0.15 - 0.25 mmol/g. This resin is unmatched for the routine synthesis of difficult peptides, even > 75 amino acids.



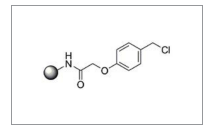
Rink Amide ProTide Resin

A powerful resin recommended for the synthesis of peptides with amide linkages < 30 amino acids. Based on ideal swelling properties from incorporating a PEG PS core with a loading of 0.55 - 0.8 mmol/g.



CI-TCP(CI) ProTide Resin

A powerful resin recommended for the synthesis of peptides with acid linkages < 30 amino acids. Based on ideal swelling properties from a TentaGel^{*} core, incorporating PEG PS with a loading of 0.4 - 0.6 mmol/g. This resin features an activated chloride linker, allowing for attachment of the first amino acid in an unactivated form. This resin is recommended for protection of C-terminal cysteine and proline residues, due to its steric protection against diketopiperazine formation and 3-(1-Piperidinyl) alanine formation.

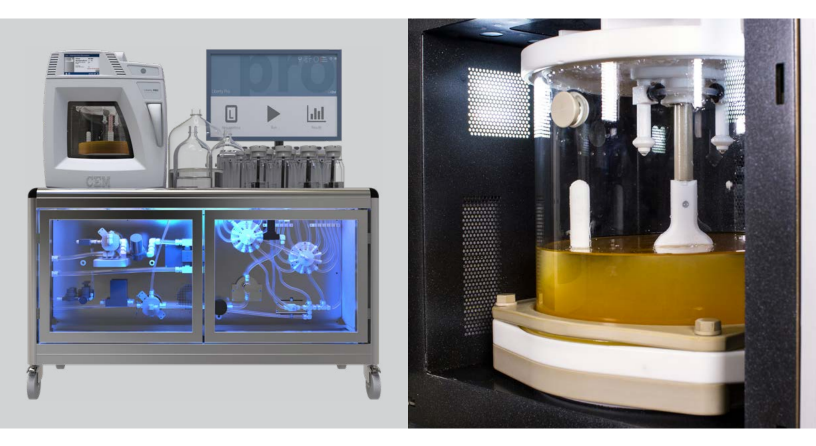


CI-MPA ProTide Resin (LL)

The ultimate resin recommended for longer and more difficult sequences of peptide acids. Based on ideal swelling properties from a TentaGel[®] core, incorporating PEG PS with a loading of 0.15 - 0.25 mmol/g. This resin is unmatched for the routine synthesis of difficult peptides even, > 75 amino acids. This resin features an activated chloride linker, allowing for attachment of the first amino acid in an unactivated form.

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- Improved synthesis efficiency with microwave irradiation
- Batch sizes up to 500 grams of purified peptide (15 L reaction vessel)
- Match synthesis profiles of peptides made on the Liberty Blue[™]

Available Options

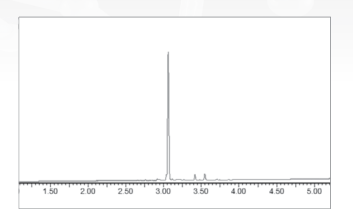
- Instrumentation (Liberty PRO)
- Large-scale custom peptide synthesis services

Contact Us To Get Started

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- peptide.support@cem.com

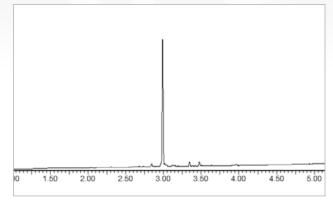
Synthesis Examples

Peptide Sequence: 9 mer Resin: Rink Amide AM PS (0.75 mmol/g) Excess Reagents: 2.0 fold



Liberty Blue™

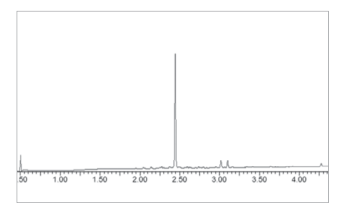
Scale: 0.1 mmol

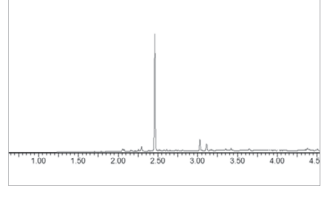


Liberty PRO™

Scale: 700 mmol Synthesis Time: 9 h

Peptide Sequence: 7 mer Resin: Rink Amide AM PS (0.97 mmol/g) Excess Reagents: 2.5 fold





Liberty Blue

Scale: 0.1 mmol

Liberty PRO

Scale: 360 mmol Synthesis Time: 6 h



"The Liberty Blue is fast, reliable, and makes difficult peptides in high purity. We are very satisfied with the Liberty Blue and would highly recommend it for both protein synthesis and methodological development."

Prof. Fernando Albericio Group Leader Chemistry & Molecular Pharmacology Institute for Research in Biomedicine (IRB) University of Barcelona

"The Liberty Blue system is unquestionably the best peptide synthesizer available today and represents the major workhorse for PeptiDream (12 systems). It is highly recommended to any company looking to synthesize peptides chemically, specifically those containing nonstandard amino acids."

Dr. Patrick C. Reid President and Director PeptiDream Inc.





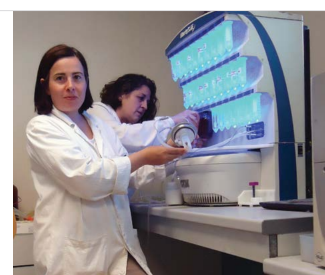
"We are very satisfied with the Liberty Blue system. The system is one of the best peptide synthesizers available today for research and medicinal chemists...*HE*-SPPS using Liberty Blue, features overwhelming speed."

Dr. Hajime Hibino Research Chemist Peptide Institute

"The Liberty Blue is the best peptide synthesizer on the market. It's synthesis speed and purity are unmatched. Using the Liberty Blue has also made our subsequent purifications easier, which is a major benefit."

Prof. Anna Maria Papini

Coordinator of Interdepartmental Laboratory PeptLab



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