

Abstract

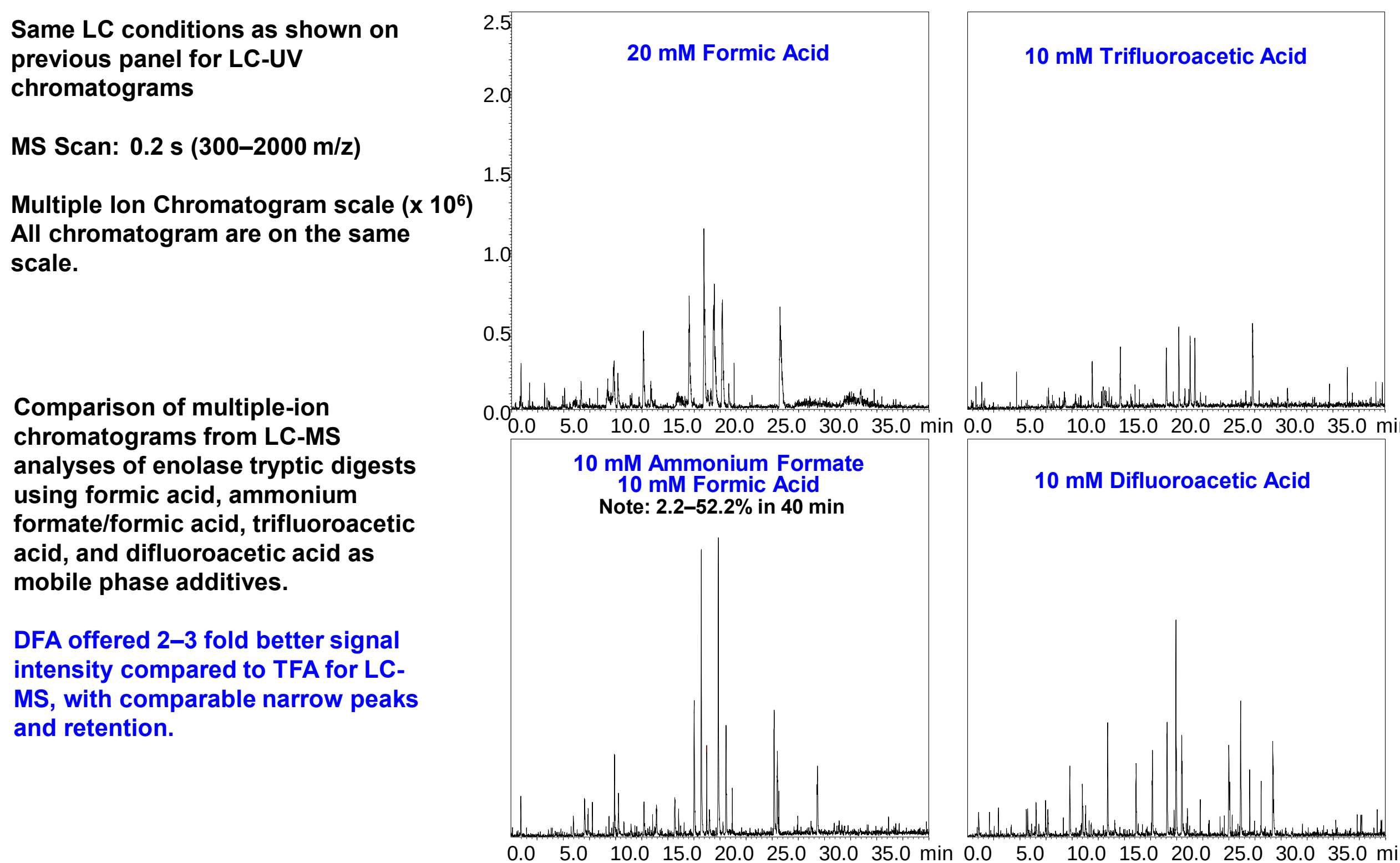
Peptide mapping is a critical competency and technique for both research and development and for quality assurance and product release for many new biopharmaceuticals including peptide drugs, monoclonal antibody drugs and antibody-drug conjugates (ADCs). New 2-µm superficially porous columns, capable of pressures up to 1000 bar, can deliver high resolution /peak capacity separations, and can be used at temperatures up to 60 °C. Moreover, a variety of mobile phase additives include trifluoroacetic acid (TFA), ammonium formate/formic acid buffers and a new additive, difluoroacetic acid (DFA), can be applied for effective development and optimization of rugged and robust separations of complex enzyme digests.

We will demonstrate the usefulness of these columns and additives in the development of peptide mapping separations, and will offer a stepwise method development and optimization strategy for such separations.

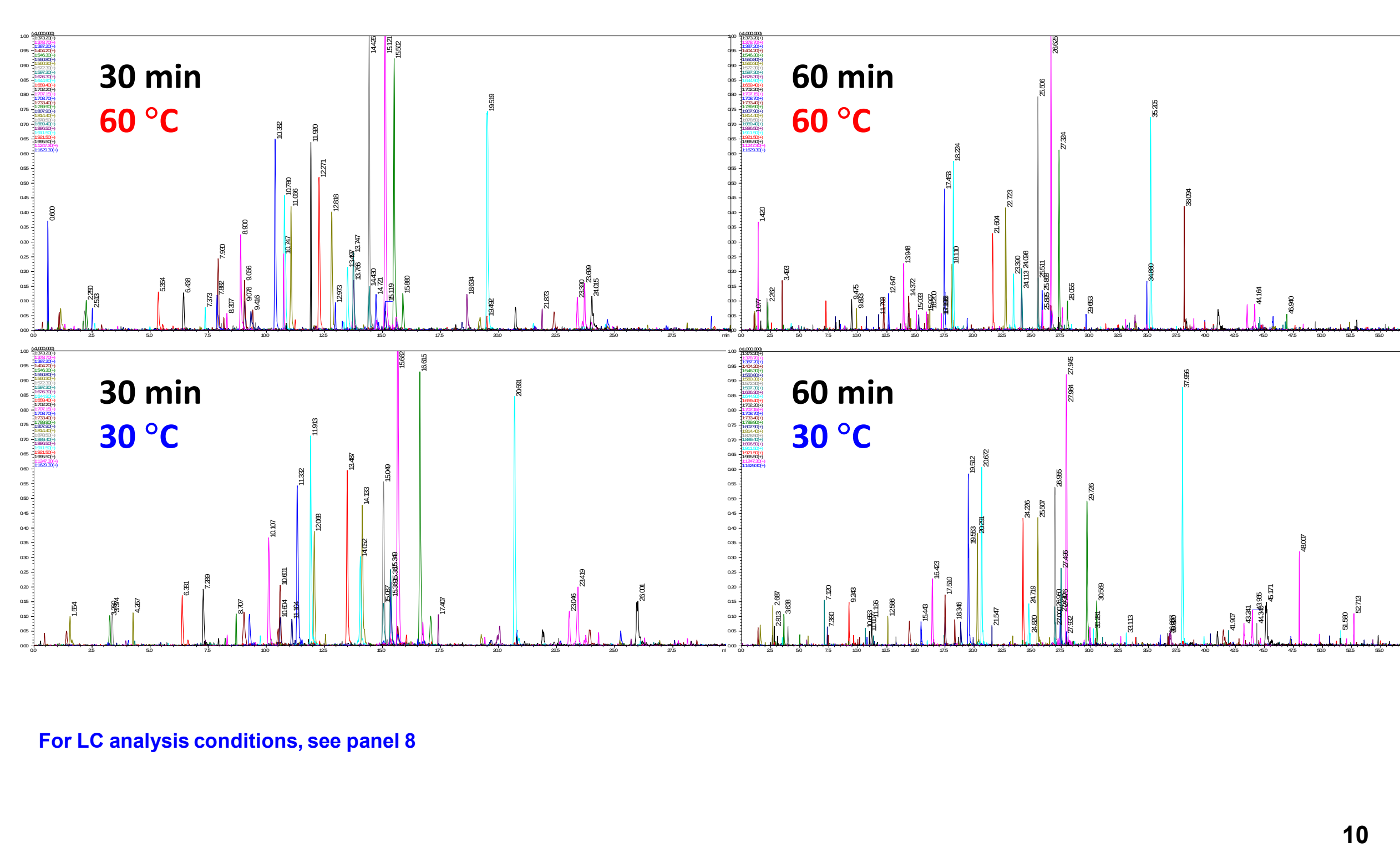
Parameters That Affect Peptide Reversed-Phase Gradient Separations

Parameter	Determined by	Impacts performance
Organic modifier	Most labs choose acetonitrile for viscosity, mass transfer, efficiency reasons, UV transparency, ionization efficiency in MS	- Peak width - Selectivity - Retention - Peak capacity
Mobile phase additive type and concentration	- UV transparency - LC-MS sensitivity (S/N) - Column stability at pH and temperature	- Peak shape - Peak width - Loading (dynamic range) - Selectivity - LC-MS signal
Gradient steepness	- User experience - Sample complexity - Experimentation and optimization software	- Selectivity - Peak capacity - Peak height
Column temperature	- Analyte requirements (recovery, stability) - Column stability requirements - Flow rate (thermal heating) - Instrument capability and pressure limit	- Selectivity - Recovery - Analysis time - Peak width - Peak capacity
Flow rate	- User choice - Column and instrument pressure limits - Flow rate - Column particle morphology and diameter	- Selectivity - Peak capacity
Pressure	- Packing particle size - Column length - Flow rate - Column temperature	- Retention - Selectivity

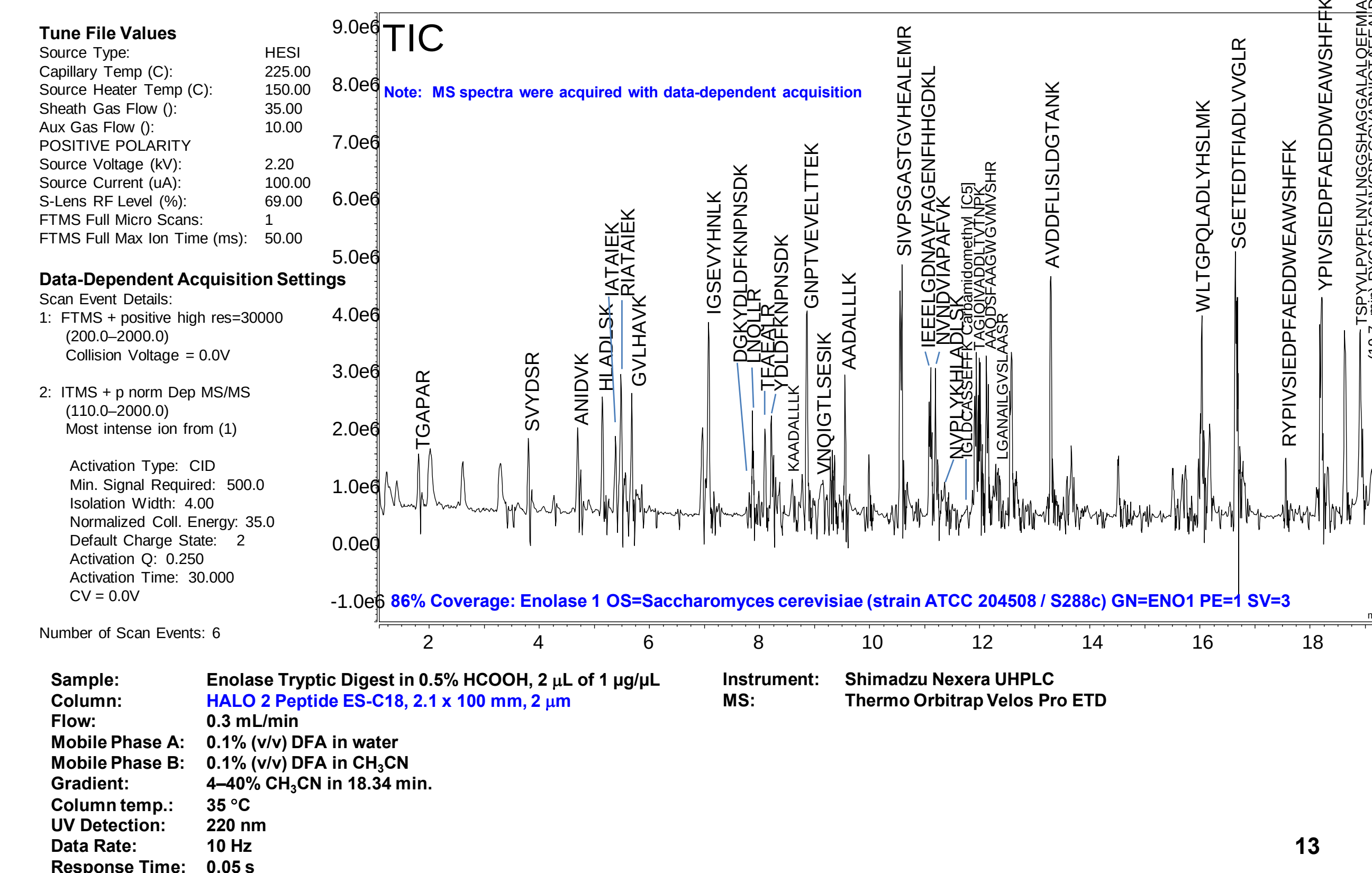
LC-MS Separations of Enolase Tryptic Digest Using Different Mobile Phase Additives



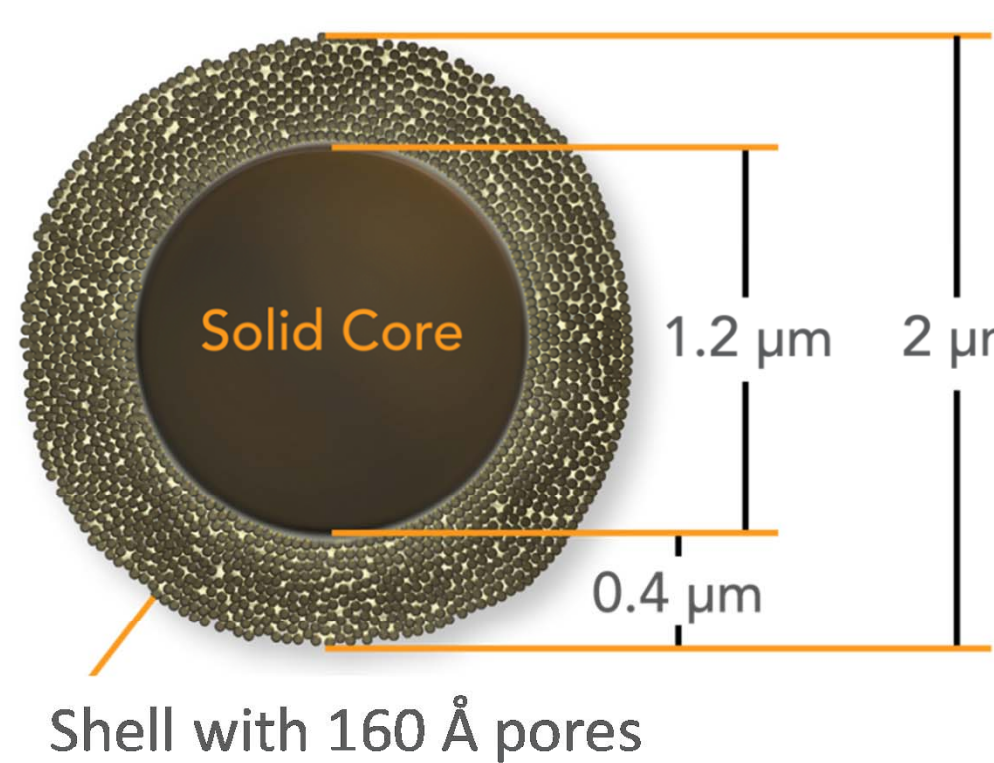
LC-MS Extracted-Ion Chromatograms for Gradient Input Runs at Two Temperatures



Total-ion Chromatogram Showing Enolase Tryptic Digest Fragments Obtained Using Near-Optimum Conditions



HALO® 2 Peptide ES-C18



Bonded phase: di-isobutyl octadecylsilane
Endcapping: None
Carbon Load: 4%
Surface Area: 65 m²/gram
Max. Temp.: 60 °C at pH 1 to 3

Designed for:

- Peptide mapping (analysis of enzyme digests) for characterization and quality monitoring of synthetic protein and antibody drugs
- Analysis of therapeutic peptides and peptide biomarkers (protein surrogates)

Attributes

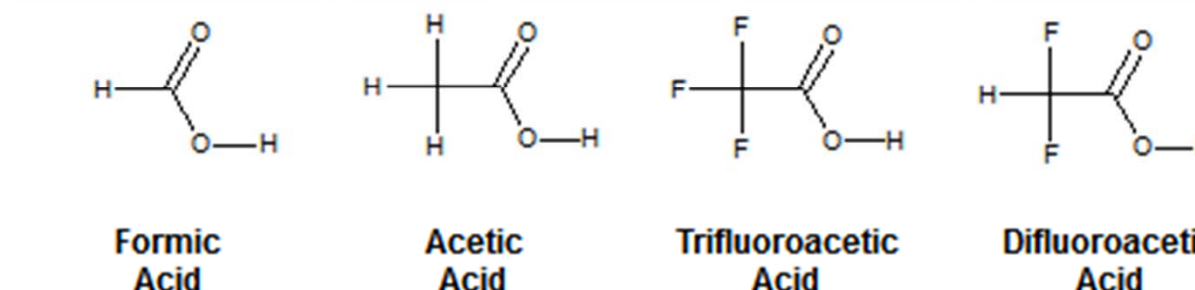
- Excellent for both fast and high resolution UHPLC separations
- High peak capacity
- Rugged, reliable performance
- Extremely stable up to 60 °C at low pH

Assessment of LC-UV and LC-MS Performance With Various Mobile Phase Additives

- Experiments were carried out using a 2.1 x 100 mm, HALO 2 Peptide ES-C18 column with the following mobile phase additives:
 - HCOOH (20 mM)
 - NH₄COOH/HCOOH (10 mM each)
 - TFA (10 mM)
 - DFA (10 mM)
- Mobile Phase A: water with additive
- Mobile Phase B: CH₃CN with additive
- Column Temperature: 60 °C
- Gradient: 2 to 47% CH₃CN/water
- Flow Rate: 0.3 mL/min
- Instrument: Shimadzu Nexera
- Instrument dispersion: 15.1 µL²

Traditional and New Mobile Phase Additives for Peptide and Protein RPLC Separations

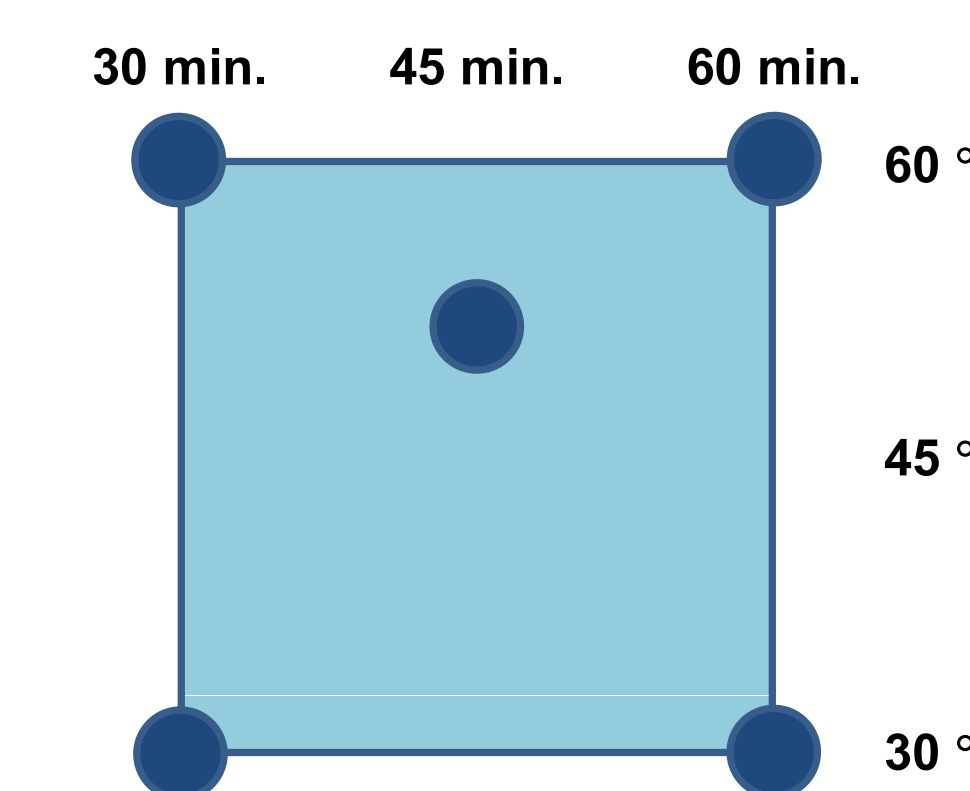
Additive	Property	Formic Acid	Acetic Acid	TFA	DFA
pK _a		3.75	4.75	0.95	2.0
Boiling Point		100.8 °C	118.1 °C	72.4 °C	133 °C
% Ionized at 10 mM		100%	4%	99%	65%
UV Cutoff (nm)		220 +	230 +	205	205



Method Development Approaches for Peptide Mapping Separations

Systematic Approach using Computer Simulation (DryLab® 4)

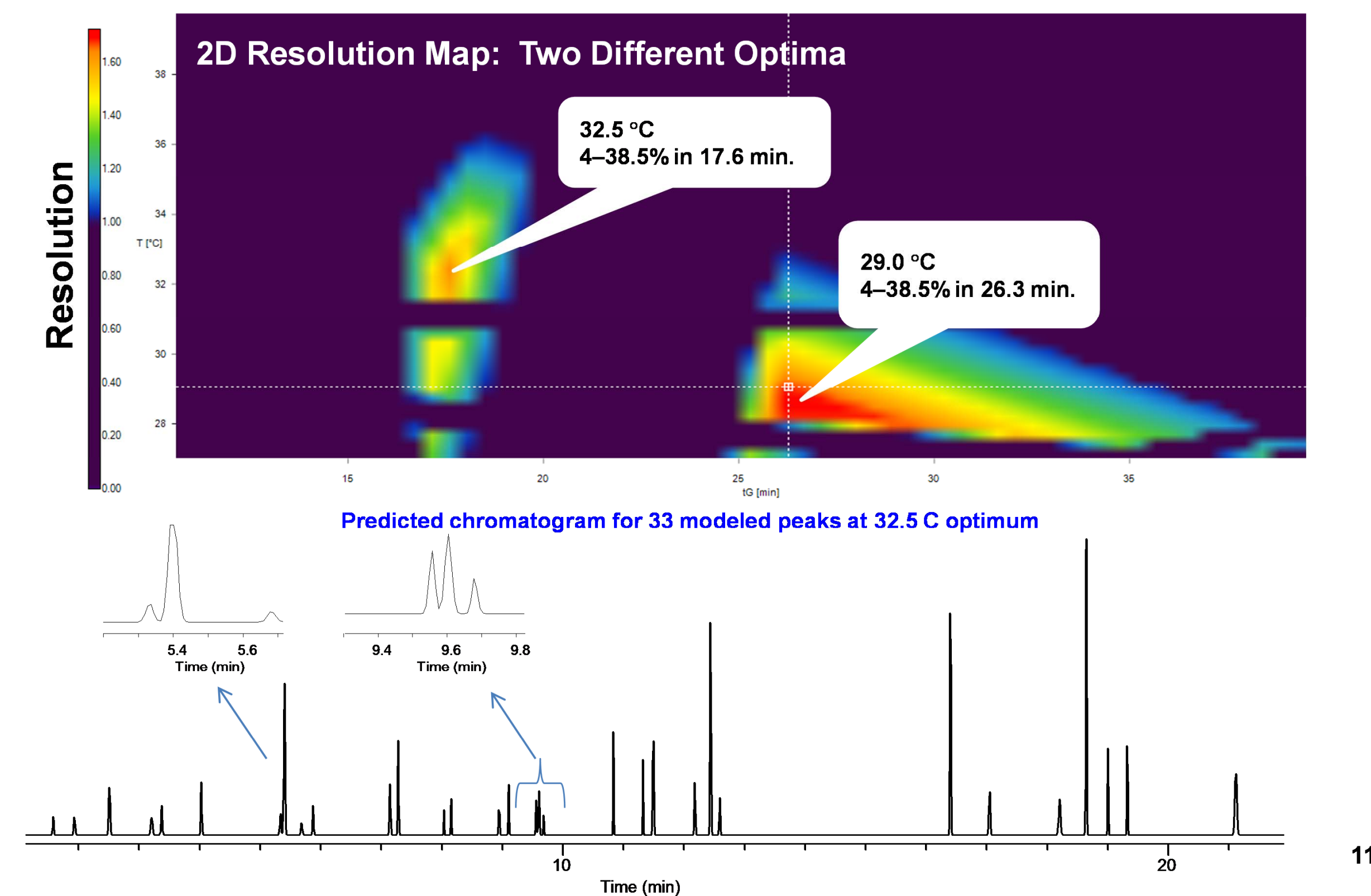
- Two-dimensional Design
 - t₀ x T
 - 2–3 gradient steepnesses
 - 2–3 column temperatures



Optimization was carried out to identify a robust gradient time and temperature combination for a run time < 40 minutes

- Column
 - HALO 2 Peptide ES-C18, 2.1 x 100 mm, 2 µm
- Mobile Phase
 - A Solvent: 0.1% DFA in water
 - B Solvent: 0.1% DFA in CH₃CN
 - Φ_{initial}: 2% B
 - Φ_{final}: 50% B
- Column Temperatures
 - 30 and 60 °C
- Gradient Times
 - 30 and 60 min.
 - 45-min x 50 °C run used for peak matching

Input Data for DryLab Software and 2-D Resolution Map With Optima



Recommended Method Development Strategies for RPLC-UV and RPLC-MS of Protein Enzymatic Digests

Systematic Approach (DryLab)

- Select desired column geometry.
- Choose initial flow rate based on column ID.
- Select two different gradient times varying by 2- or 3-fold.
- Select two column temperatures varying by 20–40 °C.
- Set gradient range of 0–5 to 60% CH₃CN or 0–5 to 50% CH₃CN.
- Carry out gradient runs at T1 and T2 with two different gradient times.
 - For example, 30 and 60 min. at 30 and 60 °C.
- Carry out an additional gradient run at intermediate temperature and gradient time.
 - For example, 45 min. at 45 °C.
- Enter RTs and areas into DryLab software.
 - Note: Peak matching can be tricky and difficult
- Compare the predicted intermediate gradient run vs. actual intermediate run to assess peak matches.
- Determine whether there are any optimum regions of robust resolution in the 2-D resolution map.
- Optimize linear gradient in gradient editor or choose a two or three segment gradient to find optimum conditions.

Wang, Carr et al. Approach*

- Choose desired column geometry.
- Set flow rate for column max. pressure or 75% of max. pressure.
- Set Φ_{final} so that last peak of interest elutes at very end of gradient.
- Choose temperature based on column stability and "fluidity" 60°C?
- For example, use 2–50% CH₃CN for initial run.
- Set gradient time for maximum allowable time.
- Calculate %B at elution of last peak of interest (using RT, %B/min, t₀ and t₉).
- Rerun sample with the 2–X %B limit for gradient time corresponding to that final %B.

* Xiaoli Wang, Dwight Stoll, Adam Schellinger, and Peter Carr
Peak Capacity Optimization of Peptide Separations in Reversed-Phase Gradient Elution Chromatography: Fixed Column Format
Anal. Chem. 2006, 78, 3406–3416

Typical Peptide Mapping Conditions and Desirable Mobile Phase Additive Properties

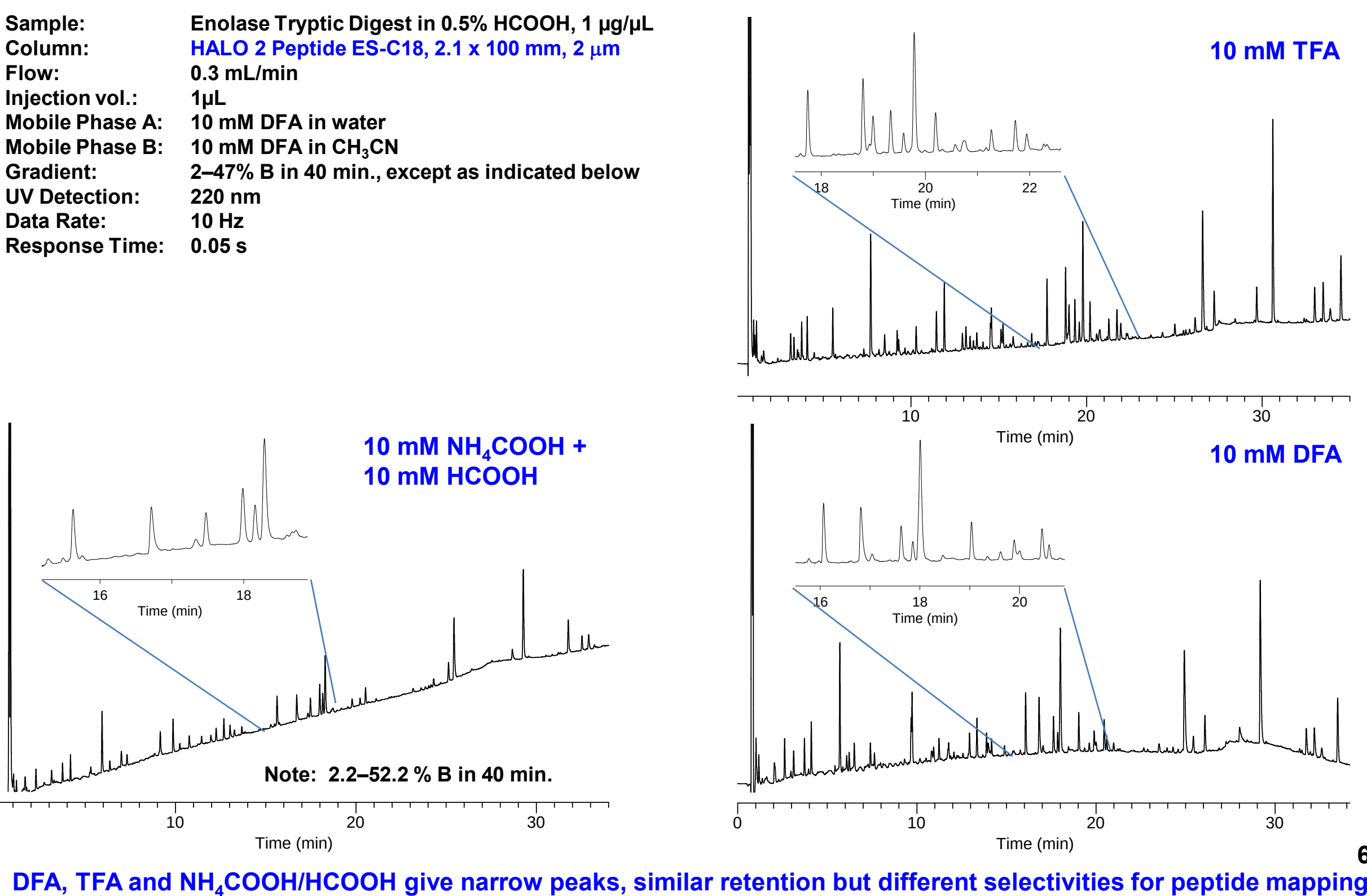
Typical Conditions for RPLC Separations of Peptides

- 150 or 250 mm C18 column
- Mobile phase A: water with 0.02–0.2% (v/v) additive
- Mobile phase B: CH₃CN with same % additive or 10–30% lower than in A solvent
- Temperature: 30 to 60 °C
- Flow rate set so that linear velocity is ~2 mm/sec
- Initial % CH₃CN: ≤ 5%
- Final % CH₃CN: 40–60%
- Gradient slope: ranges from 0.1% to 2% CH₃CN per min.
- Gradient time: can vary from 30 to 240 min.

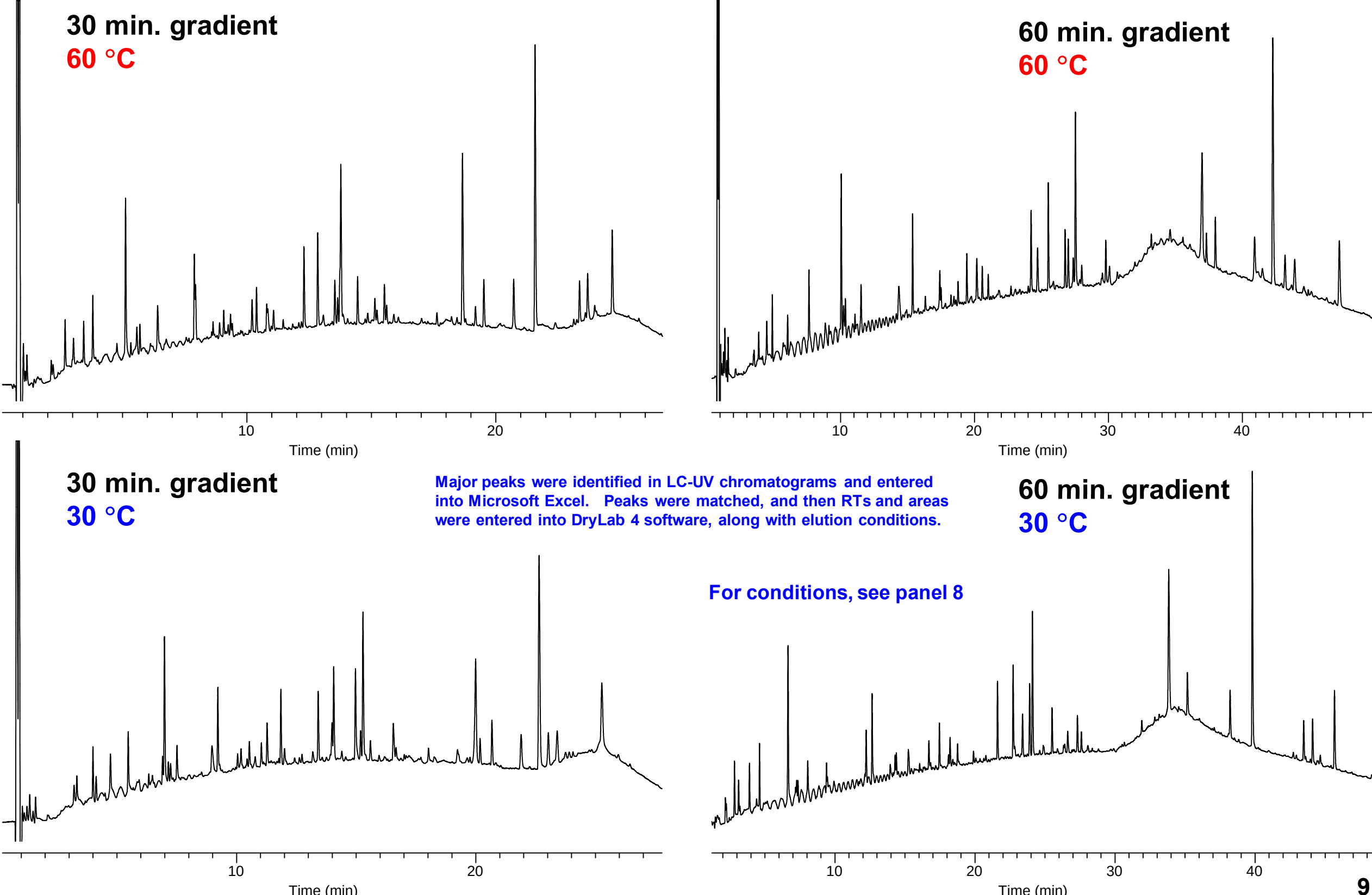
Desirable properties for additives

- Very high purity, quality, stability and reproducibility
- Low UV absorbance with acceptable baselines and blank runs
- Ability to provide good retention and symmetrical peak shapes for acidic, basic and zwitterionic analytes
- Volatile
- Acceptably low or no suppression of LC-MS signal
- Miscible with water and organic modifiers

LC-UV Separations of Enolase Tryptic Digest Using Different Mobile Phase Additives



LC-UV Chromatograms for Gradient Input Runs at Two Temperatures For DryLab Input



Peptide Identification Using Proteome Discoverer Software

Confidence	Single Letter Amino Acid Sequence	RT [min] Squest HT	Modifications	# PSiMS	# Missed Cleavages	Theo. MH+ [Da]	XCorr Squest HT	ΔM [ppm] Squest HT
High	LTGAPAR	1.99	1	0	572.31509	1.85	-1.32	
High	ANIVK	5.22	1	0	689.37227	1.99	-2.06	
High	HLADLQK	5.59	2	0	753.42593	2.87	-0.89	
High	IATAIEK	6.32	1	0	745.44543	2.26	-1.14	
High	ISSEVYHLK	7.76	2	0	1159.6106	2.98	0.04	
High	DSKVDLPNPNDSK	8.46	3	2	1755.81842	5.11	-0.17	
High	LNQLLR	8.75	2	0	756.47265	2.01	-1.38	
High	THAEALR	8.94	1	0	807.43588	2.86	-1.23	
High	YDLDPNPNDSK	9.13	1	1	1455.67505	3.81	-0.89	
High	DSKVDLPK	9.22	2	1	1100.52387	2.52	-1.5	
High	KADALLLK	9.97	2	1	947.59674	2.61	-0.38	
High	YDLDFK	10.10	1	0	806.3925	1.74	-1.28	
High	GNPTVEVLTTEK	10.29	1	0	1418.72187	3.57	-0.73	
High	YNDLTSESK	10.68	2	0	1288.70717	2.87	-1.3	
High	SVSPSASTGVHEALEMR	10.76	1	0	1858.91709	4.23	-0.29	
High	IEEELGDAVFASENFHRSK	11.36	2	0	2328.05273	4.73	-2.91	
High	AQDSFAAGWGMVSHR	11.71	1	0	1865.80978	4.72	-1.41	
High	SVSPSASTGVHEALEMR	12.17	2	0	1849.92217	4.22	-0.27	
High	IEEELGDAVFASENFHRSK	12.70	1	1	2441.13679	5.34	-0.91	
High	YNDLSESK	12.82	2	0	1286.71031	2.87	-1.51	
High	YLDLASSPEPK	13.60	1	0	1373.64058	3.79	-0.46	
High	AQDSFAAGWGMVSHR	13.63	1	0	1789.84386	5.28	-0.45	
High	TAGIQVADLTVMRK	14.31	2	0	1755.9487	4.97	0.18	
High	LDANALGVSLASR	15.02	1	0	1412.82199	4.57	-1.23	
High	AVDRLSLDTGANK	15.40	3	0	1578.80998	5.83	-0.48	
High	WLTGPQLADLYSLRK	16.89	1	0	1888.96258	4.8	-1.3	
High	WLTGPQLADLYSLRK	16.95	2	0	1877.96167	5.32	-0.18	
High	SEETEDFAVSLVSLR	19.42	1	0	1821.82288	6.42	-0.33	
High	RYVPSIEDPFAEDDWEAWSHFFK	20.64	1	1	2984.38808	7.39	-1.18	
High	RYVPSIEDPFAEDDWEAWSHFFK	21.51	2	0	2828.29187	5.48	-0.78	
High	YKASGWDGEGGVNWKAEALDLYDAIK	24.08	1	0	3257.81721	4.97	-0.88	

Data Set was generated using the 30-min, 60 °C Xcalibur software raw file

PSM = Peptide Spectrum Match

XCorr = Cross Correlation Factor

Summary and Conclusions

- A sample of enolase was digested with trypsin, and was analyzed using gradient UHPLC using a 100-mm HALO 2 Peptide ES-C18 column.
- Various mobile phase additives were compared in terms of LC-UV and LC-MS signal, peak shape, peak width and retention.
- Difluoroacetic acid (DFA) offers peak shape, peak width and retention comparable to TFA and ammonium formate/formic acid, but with much better LC-MS ionization efficiency (signal-to-noise).
- Systematic method development for optimizing peptide separations allows one to find temperature and gradient time combinations for robust, effective analyses.
- HALO 2 Peptide ES-C18 columns facilitate high resolution peptide mapping of complex tryptic digests with short analysis times (20–30 minutes).

Acknowledgements: Supported in part by NIH Grant GM116224 (BEB). Opinions expressed are solely those of the authors. HALO and Fused-Core are registered trademarks of Advanced Materials Technology, Inc.