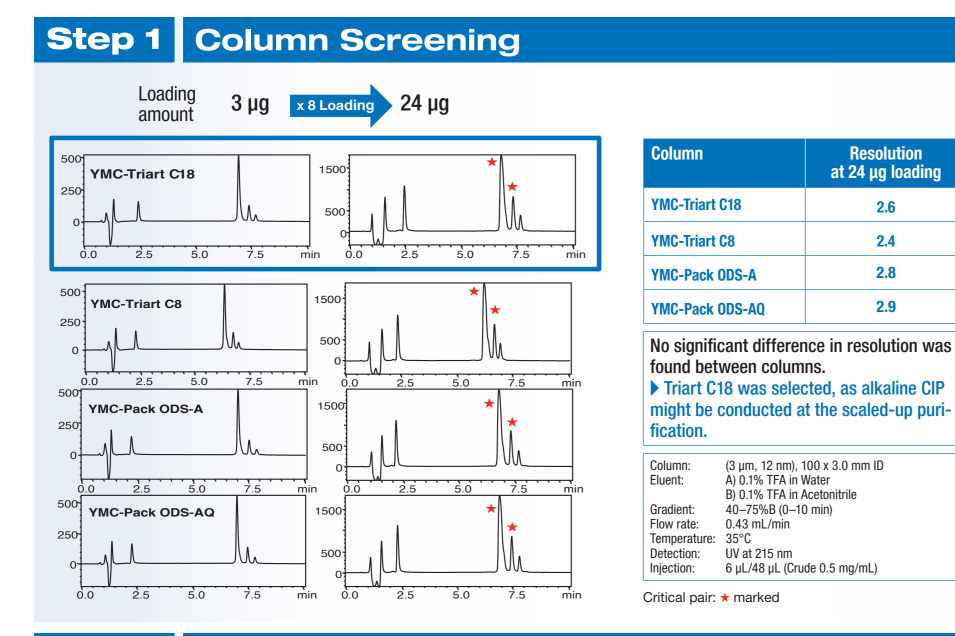
Purification method development for Liraglutide

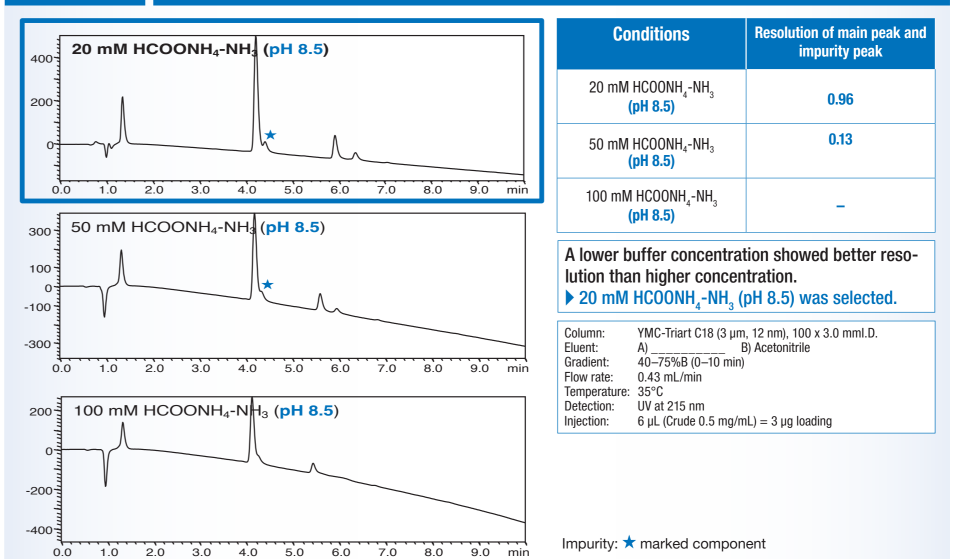


urification is the most critical step in the manufacturing process of peptide therapeutics. The right choice of chromatography media is crucial for cost-effective production. With its wide pH range (pH 2-10), YMC-Triart Prep C18-S provides you with full flexibility in the method development of peptide purification. Simple scale-up procedures ensure the



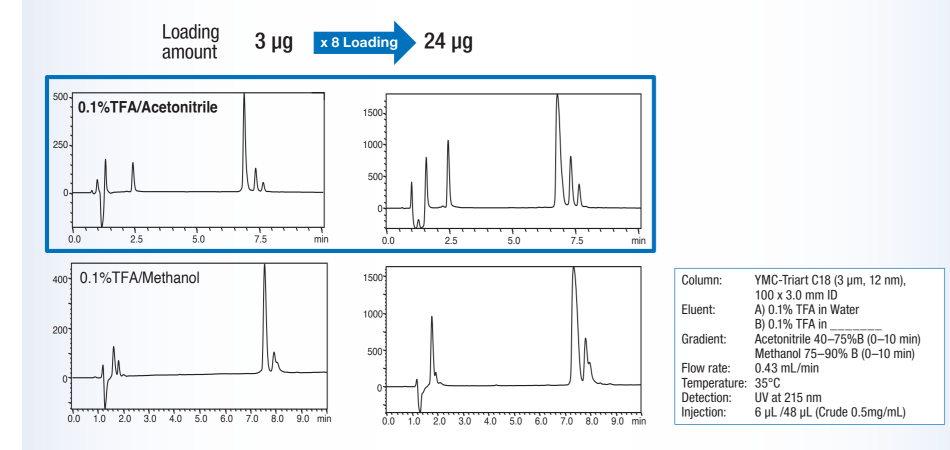
reproducible result at manufacture-scale. A method for the purification of liraglutide with high resolution (antidiabetic peptide therapeutic, marketed by Novo Nordisk as Victoza®) was successfully developed with YMC-Triart Prep C18-S under alkaline condition.

The purity obtained for the target compound was 99.5%.



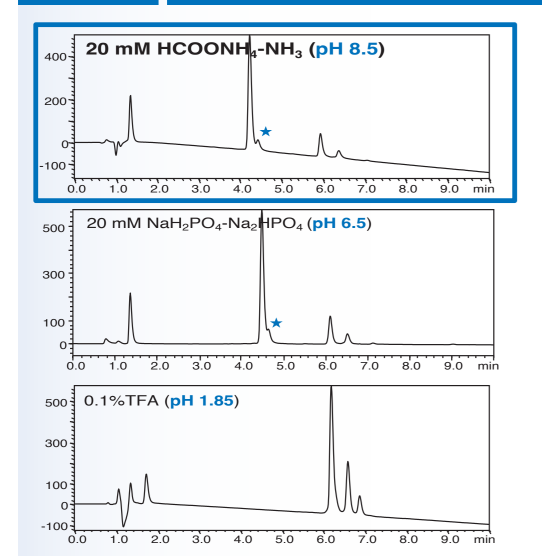
Step 5 Influence of Buffer Concentration

Step 2 Influence of Organic Solvent



Acetonitrile gave better resolution of the critical pair than methanol. Furthermore, longer retention when using methanol is not good for prep purpose from the stand point of post-chromatography step. Acetonitrile was selected.

Step 3 Influence of pH

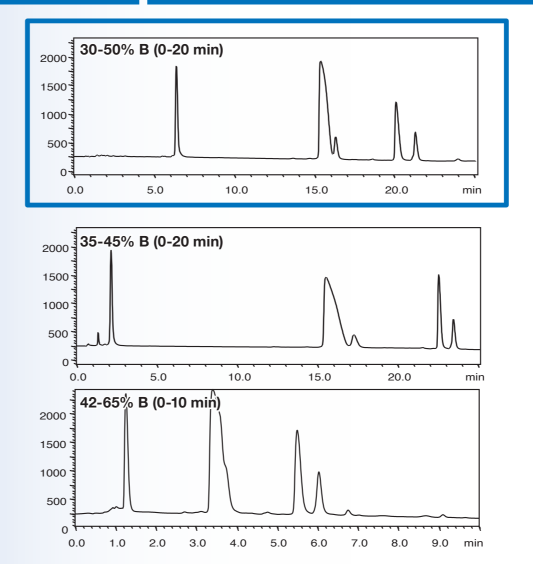


Conditions	Resolution of main peak and impurity peak			
20 mM HCOONH ₄ -NH ₃ (pH 8.5)	0.96			
20 mM NaH ₂ PO ₄ -Na ₂ HPO ₄ (pH 6.5)	0.50			
0.1% TFA (pH 1.85)	_			
Higher pH gave better resolution, and a new impu- rity peak appeared and separated at higher pH.				

▶ pH 8.5 was selected, by considering chemical durability of the packing material and resolution.

Column:	YMC-Triart C18 (3 µm, 12 nm), 100 x 3.0 mm ID
Eluent:	A) B) Acetonitrile
Gradient:	40–75%B (0–10 min)
Flow rate:	0.43 mL/min
Temperature:	35°C
Detection:	UV at 215 nm
Injection:	$6 \ \mu L$ (Crude 0.5 mg/mL) = 3 μg loading

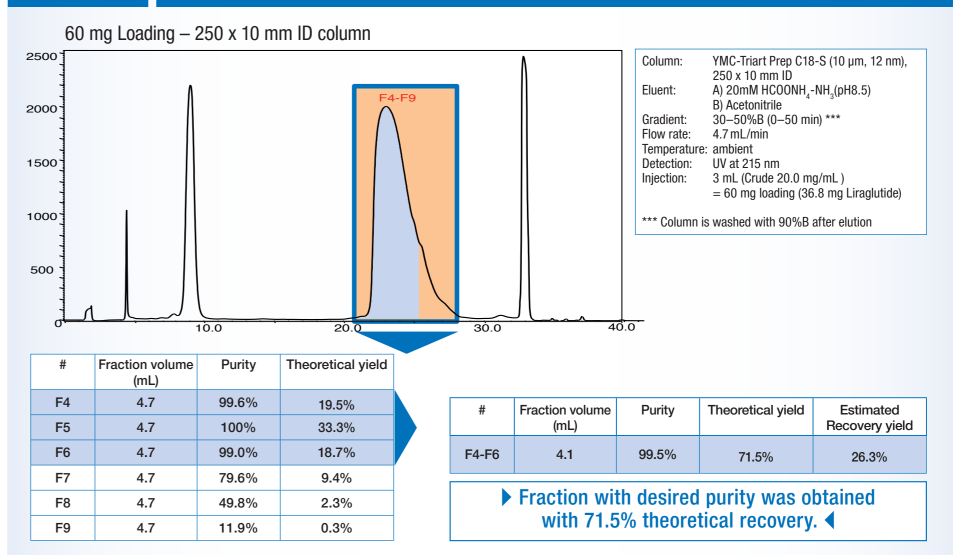
Step 6 Gradient Optimization



At increased loading condition, influence of gradient curve was evaluated. ▶ 30-50%B (0-20 min) condition was selected when considering balance of resolution of critical pair and fraction volume.

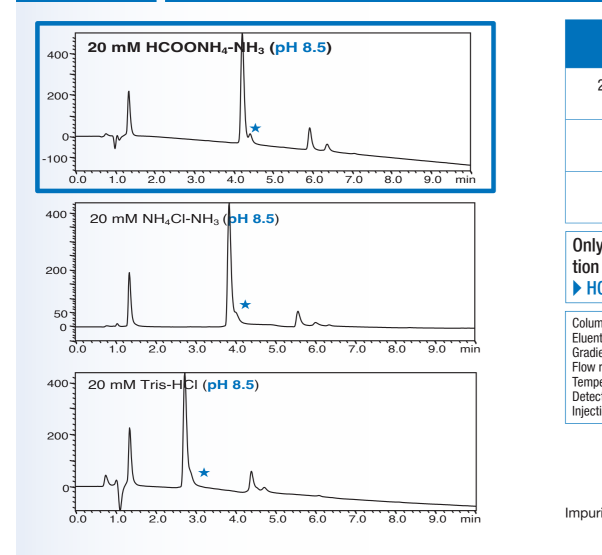
YMC-Triart C18 (3 µm, 12 nm), 100 x 3.0 mm ID Column: A) 20mM HCOONH₄-NH₃(pH8.5) Eluent: B) Acetonitrile Flow rate: 0.43 mL/min Temperature: 35°C Detection: UV at 215 nm 3 μL (Crude 20.5 mg/mL) = 61.5 μg loading Injection:

Result Purification Run



Impurity: **★** marked component

Influence of Buffer Type Step 4



Conditions	Resolution of main peak and impurity peak		Column	
M HCOONH ₄ -NH ₃ (pH 8.5)	0.96		Eluent	
mM NH ₄ CI-NH ₃ (pH 8.5)	-		Detection	
) mM Tris-HCl (pH 8.5)	-		Temperature	
γ HCOONH ₄ -NH ₃ system showed a fair separa-			Cycle time	
of the main peak and impurity peak. $COONH_4$ -NH ₃ was selected.			Column dimension	
nn: YMC-Triart C18 (3 μm, 12 nm), 100 x 3.0 mm ID t: A) B) Acetonitrile ent: 40–75% B (0–10 min) rate: 0.43 mL/min erature: 35°C tion: UV at 215 nm ion: 6 μL (Crude 0.5 mg/mL) = 3 μg loading			Flow rate	
			Loading/run	
			Fraction volume / run	
			Liraglutide recovery / run	
ity: ★ marked component			Liraglutide recovery /day	

Theoretical Scaling Up Calculation

Column	YMC-Triart Prep C18-S (10 µm, 12 nm)					
Eluent	A) 20 mM HCOONH ₄ -NH ₃ (pH 8.5) B) Acetonitrile 30-50% B (0-50 min)					
Detection	UV at 215 nm					
Temperature	Ambient					
Cycle time	60 min/run – 8 cycles/day					
Column dimension	250 x 100 mm ID	250 x 450 mm ID	250 x 600 mm ID			
Flow rate	0.47 L/min	9.52 L/min	16.92 L/min			
Loading/run6.0 gFraction volume / run1.4 L		121.5 g	216.0 g			
		28.6 L	50.8 L			
Liraglutide recovery / run	2.6 g	53.4 g	94.9 g			
Liraglutide recovery /day	20.8 g	427.2 g	759.2 g			

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