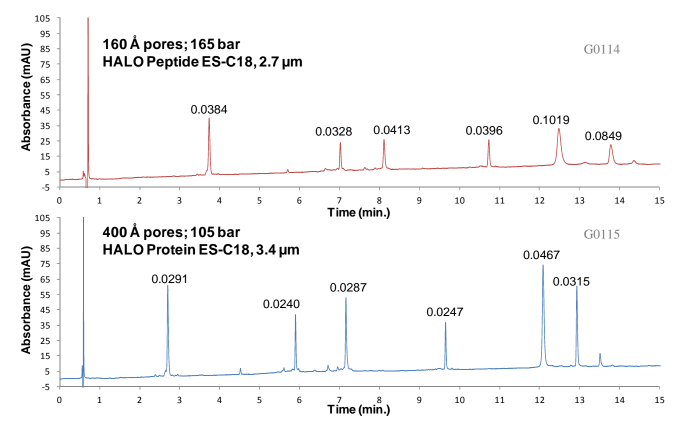
HALO: | Fused-Core® Particle Technology

Application Note: 130-PR

Effect of Silica Pore Size on Protein Separations



TEST CONDITIONS: Columns:

4.6 x100 mm, HALO Peptide ES-C18, 2.7 µm Part Number: 92124-602 4.6 x 100 mm, HALO Protein ES-C18, 3.4 µm Part Number: 93414-602 Mobile Phase: A= 0.1% Trifluoroacetic acid in water B= 0.1% Trifluoroacetic acid in acetonitrile Flow Rate: 1.5 mL/min. Gradient: 23% B to 50% B in 15 minutes Starting pressure: As indicated on chart Temperature: 60℃ Detection: UV 215 nm, VWD Injection Volume: 5 µL Sample Solvent: mobile phase A Response Time: 0.12 sec. Data Rate: 14 Hz

LC System: Agilent 1100 Quaternary Flow Cell: 5 µL semi-micro

PEAK IDENTITIES:

2.	Ribonuclease A Cytochrome <i>c</i> Lysozyme	13.7 kDa 12.4 kDa 14.3 kDa
4. 5.	α-Lactalbumin Catalase Enolase	14.2 kDa tetramer of ~ 60 kDa each 46.7 kDa

Sharper, taller peaks are observed using the HALO 400 Å Protein ES-C18 column because the larger pore size allows unrestricted diffusion for these biomolecules into and out of the porous shell. The half height peak widths above each protein peak are significantly smaller with the HALO Protein column despite the larger particle size of the packing material, emphasizing the importance of larger pores when separating proteins.



FOR MORE INFORMATION OR TO PLACE AN ORDER, CONTACT:

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