

The logo for Celerion features a stylized, wavy maroon line above the word "celerion" in a lowercase, sans-serif font. The background of the slide is white with a large, dark maroon wave shape at the bottom.

celerion

Quantitative LC-MS/MS Analysis of Glucagon

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June 21, 2011

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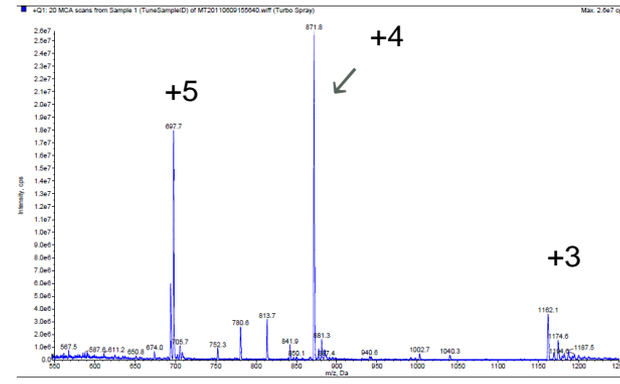
- Comparison with small molecule LC-MS/MS
- LC-MS/MS sensitivity of peptides detection
- Stability: neat vs. matrix solutions
- Method: extraction & LC-MS/MS
- Reference material
- Conclusions

Examples for Improvement of Peptides MS/MS Sensitivity

- Negative ESI MS/MS with loss of water
 - β -amyloid peptides, 100 pg/mL
- Sequence specific fragmentation
 - cleavage of peptide bonds involving Pro residues
- Chemical modification
 - Cys residues derivatization with iodoacetamide (terlipressin) increases MRM response by ~ 5-fold

Glucagon MS/MS

NH_2 -His-Ser-Gln-Gly-Thr-Phe-Thr-Ser-
Asp-Tyr-Ser-Lys-Tyr-Leu-Asp-Ser-Arg-
Arg-Ala-Gln-Asp-Phe-Val-Gln-Trp-Leu-
Met-Asn-Thr-COOH

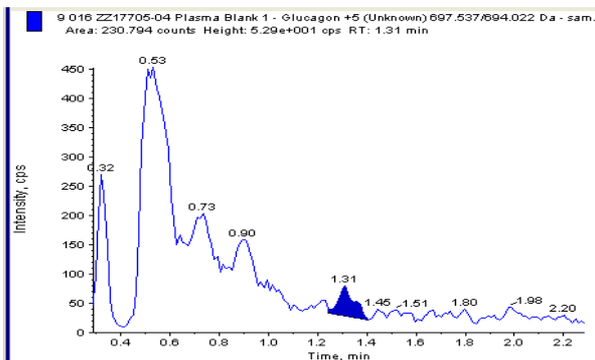


- Multiple-charged species in ESI mass spectra
- Unique highly efficient fragmentation of M^{+5} ions with loss of ammonia
 - Other peptides with N-terminal His share this fragmentation feature, including glucagon analogs missing Thr⁵ and Thr⁷

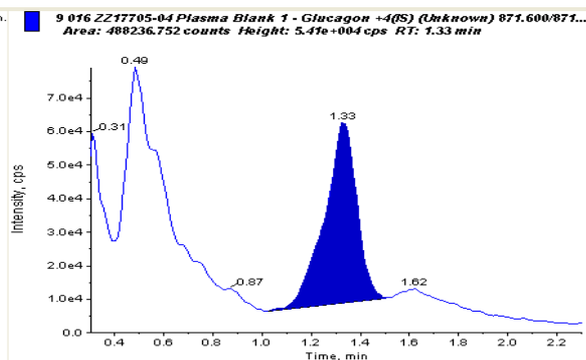
Selectivity/Efficiency of Glucagon MRM

Blank

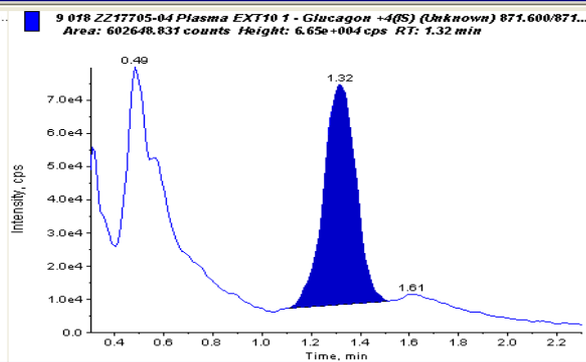
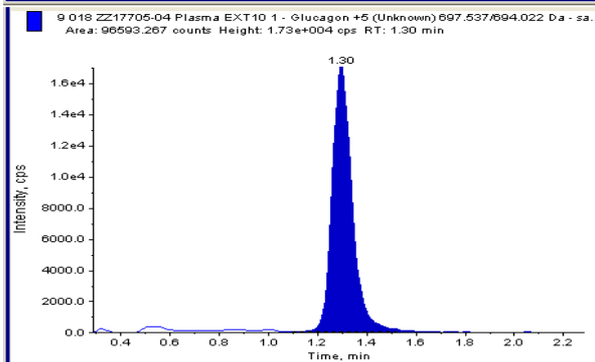
$M+5 \rightarrow (M-17)+5$



$M+4 \rightarrow M+4$



10 ng/mL



Selectivity: distinct difference between blank and spiked samples

MRM Efficiency: relative increase of the analyte response in spiked samples vs blanks for both transitions is approximately the same [peak area counts: 103,000 ($M+5$ MRM) vs. 114,000 ($M+4$ MRM)]

Glucagon Stability (Neat Solutions)

- Soluble in acidic (pH < 3) and basic (pH >9.5) solutions
- Chemically stable: decomposition of Trp, Met oxidation, deamidation of Asn/Gln, or peptide bonds hydrolysis only significant in relatively harsh conditions
- Prevention of adsorption: coating of polypropylene tubes with BSA
 - Glucagon solution below 50 µg/mL: addition of “keeper” peptides
 - Choice of “keeper” peptide/compound: lack of interferences, compatibility with method/analyte
- Extracted samples: no adsorptions (96 hours, 5°C)

Glucagon Stability (Plasma)

- Proteolysis rate is matrix lot-dependent
- In some lots of human plasma, aprotinin alone (250 KIU/mL) does not provide sufficient glucagon stability
- Cocktail of inhibitors was developed to enhance glucagon stability in human plasma and in whole blood
- Proteolysis rates of (des-Thr⁵)- and (des-Thr⁷)-glucagon variants are similar to glucagon degradation rates

Enhancement of Glucagon Stability in Plasma

- Human: Short-term stability (17 hours) of Test QC samples on an ice water bath

| Inhibitor | Aprotinin 250 KIU/mL | | Cocktail of inhibitors | |
|-----------------------------|-------------------------|--------|------------------------|--------|
| | Control QC | STS QC | Control QC | STS QC |
| | 5910 | 2180 | 6030 | 5830 |
| | 5930 | 2090 | 6050 | 5800 |
| | 5570 | 2160 | 6010 | 6050 |
| Mean | 5800 | 2140 | 6030 | 5890 |
| % CV | 3.5 | 2.2 | 0.3 | 2.3 |
| Stability (% of Control) | | 36.9 | | 97.7 |

- Rat: Acidification of plasma is also required to provide sample integrity along with addition of protease inhibitor cocktail

Glucagon Method: Extraction

- Ion-exchange 96-well plates
- Sample incubation with detergent & acetonitrile
 - Minimize protein binding
 - Improve accuracy of quantitation in matrix from multiple donors
- SPE washes with several organic solvents
 - Ensure consistency of the analyte/IS recovery
 - Lack of matrix effect
- Internal standard: (des-Thr⁷)-glucagon

Glucagon Method: LC-MS/MS

- Parallel-column system (Agilent Zorbax 300SB-C18, 3.5 μm , 50x2.1 mm)
 - Isocratic elution on column 1 (analysis)
 - Gradient regeneration on column 2
- Advantages:
 - Stability of LC-MS/MS system (ratio & response)
 - Lack of carry-over
 - Run time < 4 minutes

Glucagon (Human Plasma) Method Parameters

- Analytical range: 100-10,000 pg/mL
- Dilution integrity: up to 25,000 pg/mL
- Sample volume: 0.250 mL
- Sample collection and handling stability: 2 hours (5°C)
- Short-term stability in matrix: 14 hours (5°C)

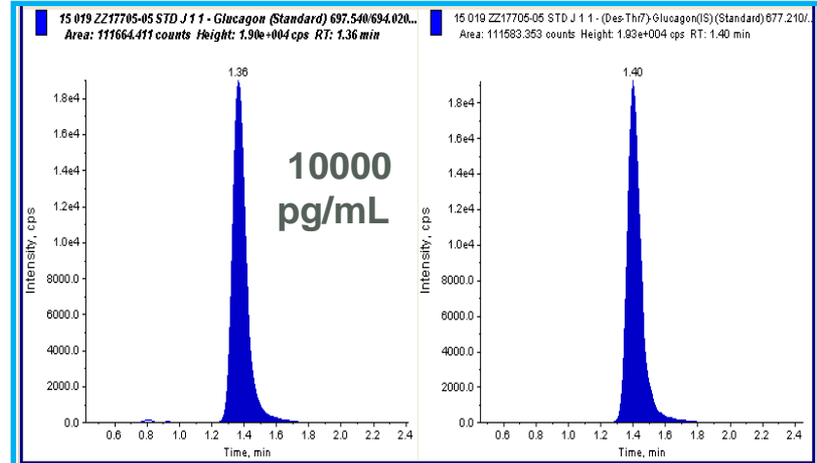
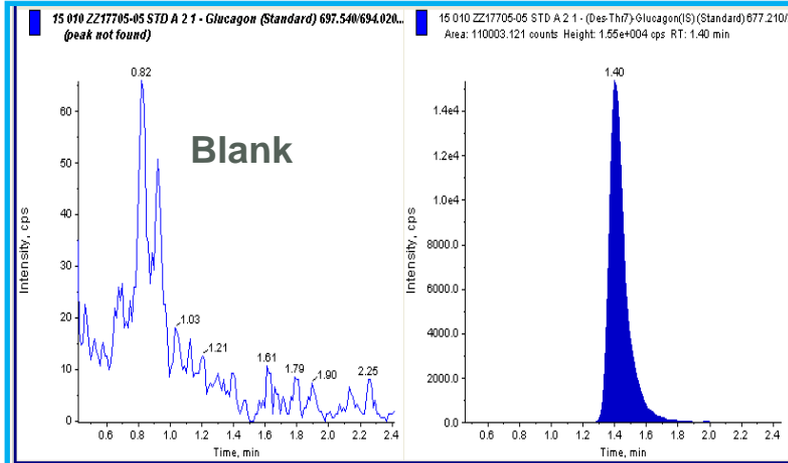
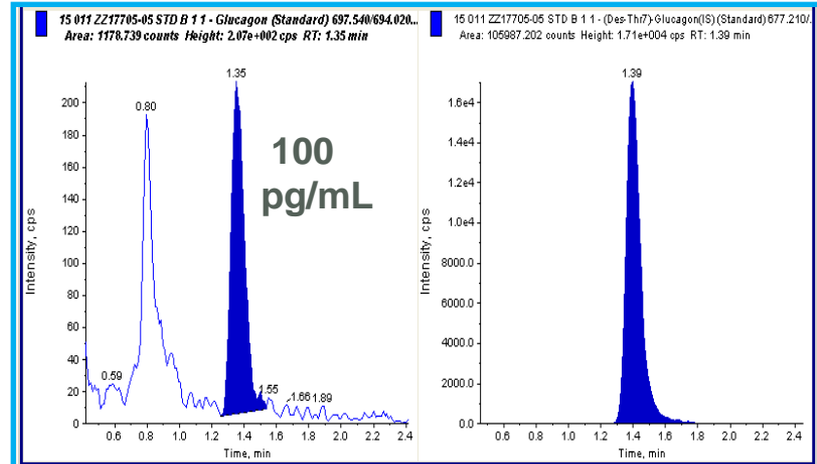
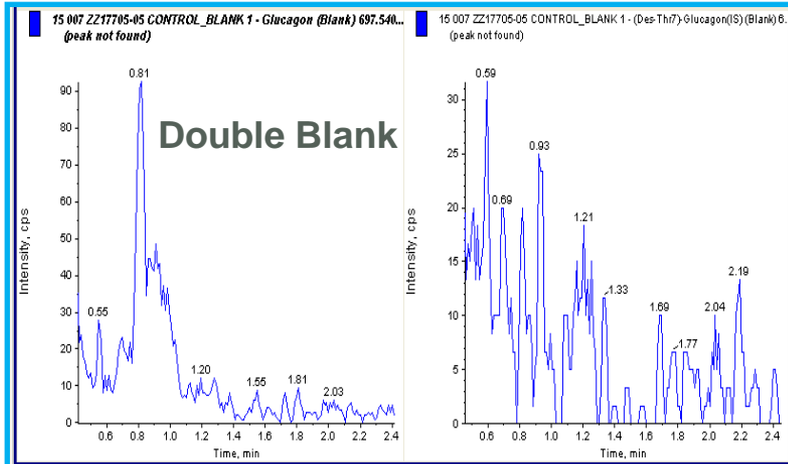
Glucagon Extracted Samples

Glucagon

Internal Standard

Glucagon

Internal Standard

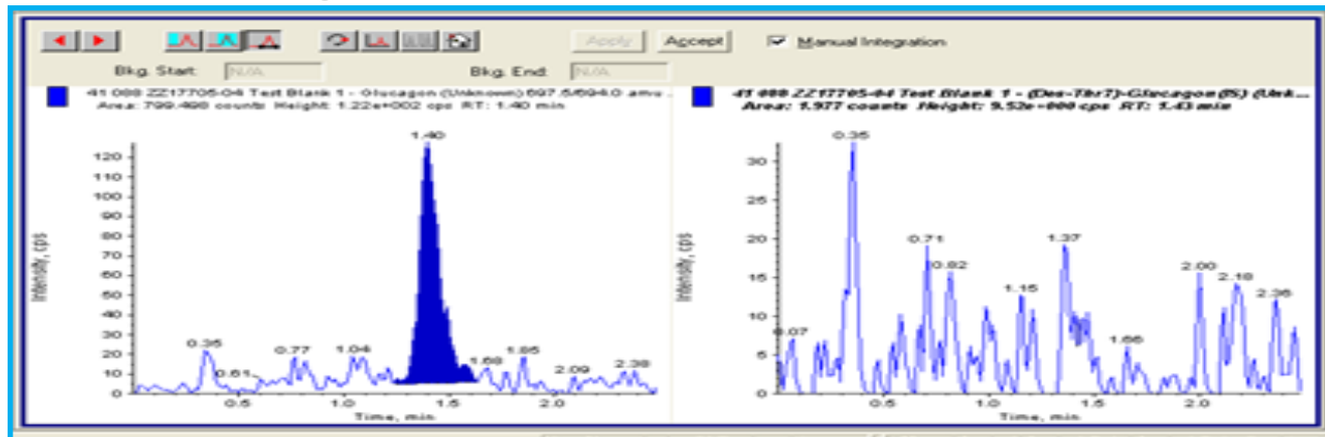


Evaluation of Glucagon Endogenous Level

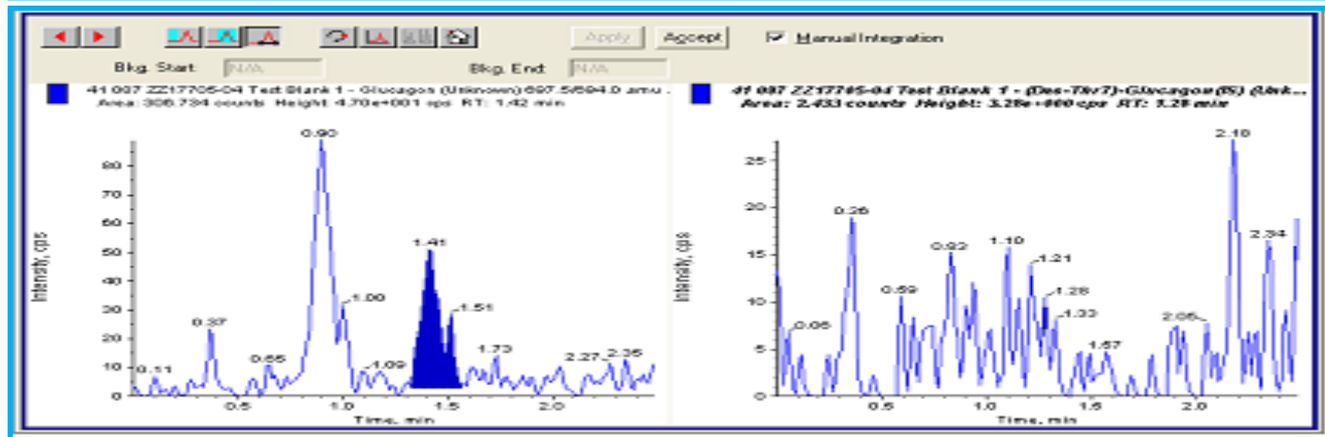
Glucagon

Internal Standard

Lot 1
(~55 pg/mL)



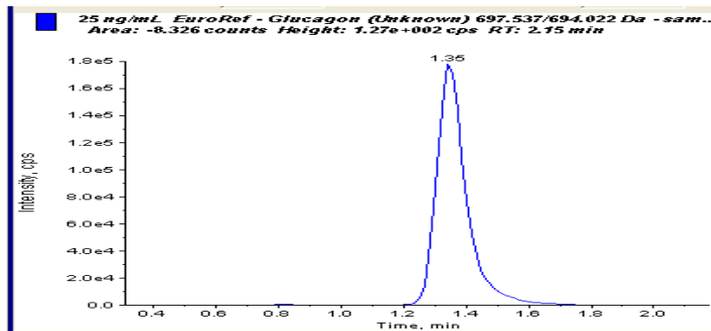
Lot 2
(~25 pg/mL)



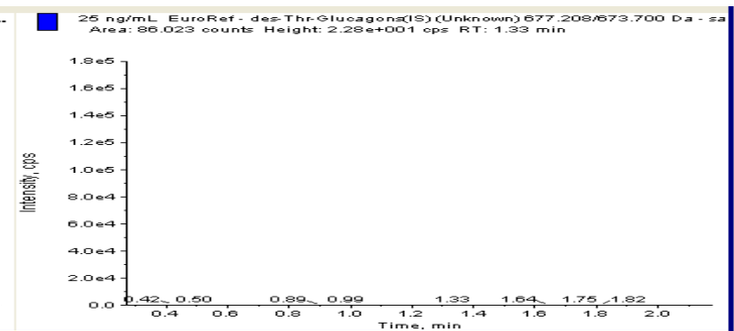
Glucagon Reference Standards

Ph. Eur.
Glucagon

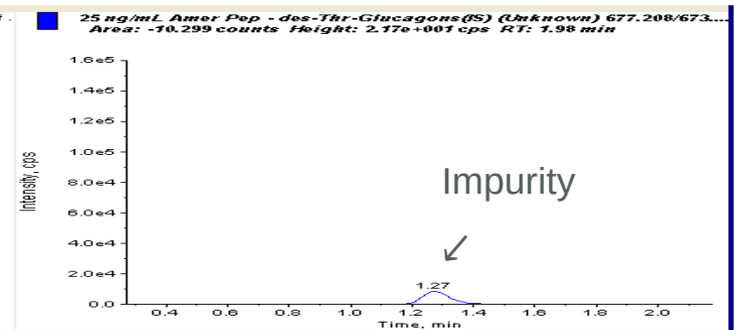
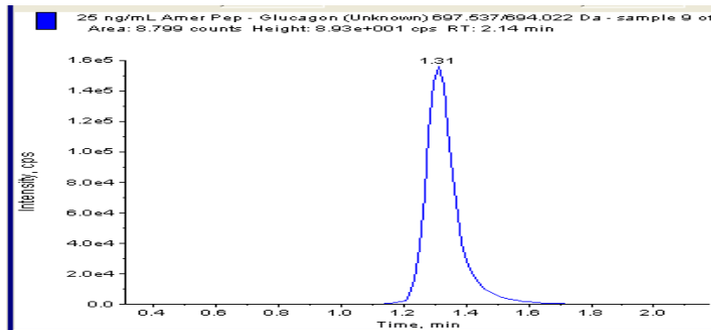
Glucagon



Internal Standard



Synthetic
Glucagon



- Concentrations of glucagon as European Pharmacopeia Reference Standard and glucagon from commercially available Eli Lilly Glucagon Emergency Kit matched well
- Some synthetic preparations may contain a significant amount of peptide impurities not shown in Certificate of Analysis

LC-MS/MS vs Immunochemical Methods (Selectivity)

- Glucagon (RIA vs LC-MS/MS)
 - Low concentration quality control samples (140 pg/mL, RIA method) had no detectable intact glucagon in LC-MS/MS method
- 13,14-dihydro-15-keto Prostaglandin F_{2α} (ELISA vs LC-MS/MS)
 - Analytical samples: up to 200-fold difference in concentrations between methods

Conclusions

- Some challenges in LC-MS/MS with peptides similar to those with small molecules
- Glucagon MRM transition with loss of ammonia provides an easy LC-MS/MS solution
- Several inhibitors are required to ensure glucagon stability in human plasma
- LC-MS/MS glucagon method advantages
 - More selective than immunochemical
 - Lack of matrix effect
 - Large linear range

Acknowledgements

- Paul Brown
- Chris Kafonek
- Alan Dzerk
- Brendon Retke
- Corey Ohnmacht
- Ridha Nachi
- Sarah Roby
- Patrick Miller
- Curtis Sheldon

