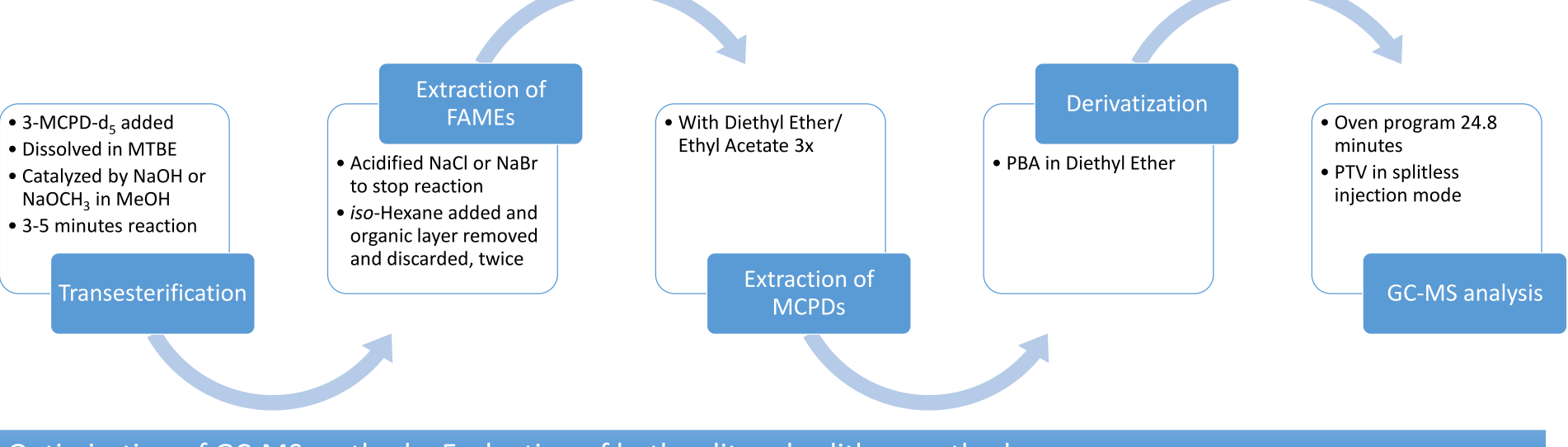
Optimizing GC-MS Analysis of 3-MCPD and Glycidyl Esters

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Abstract

Sample Preparation (according to AOCS Cd 13c-29)

3-MCPD and glycidyl esters in edible oils are contaminants that are formed through refining processes. Several of these substances have been classified as possible human carcinogens. Methods, which are similar to one another, have been developed by ISO, AOCS, and DGF for analyzing these contaminants. While these methods cover extraction and derivatization techniques in detail, very little attention is paid to the GC-MS methods. With emerging automated systems, it is important to simplify and speed up the instrument method by optimizing the parameters, to include evaluating split injection.



Optimization of GC-MS method

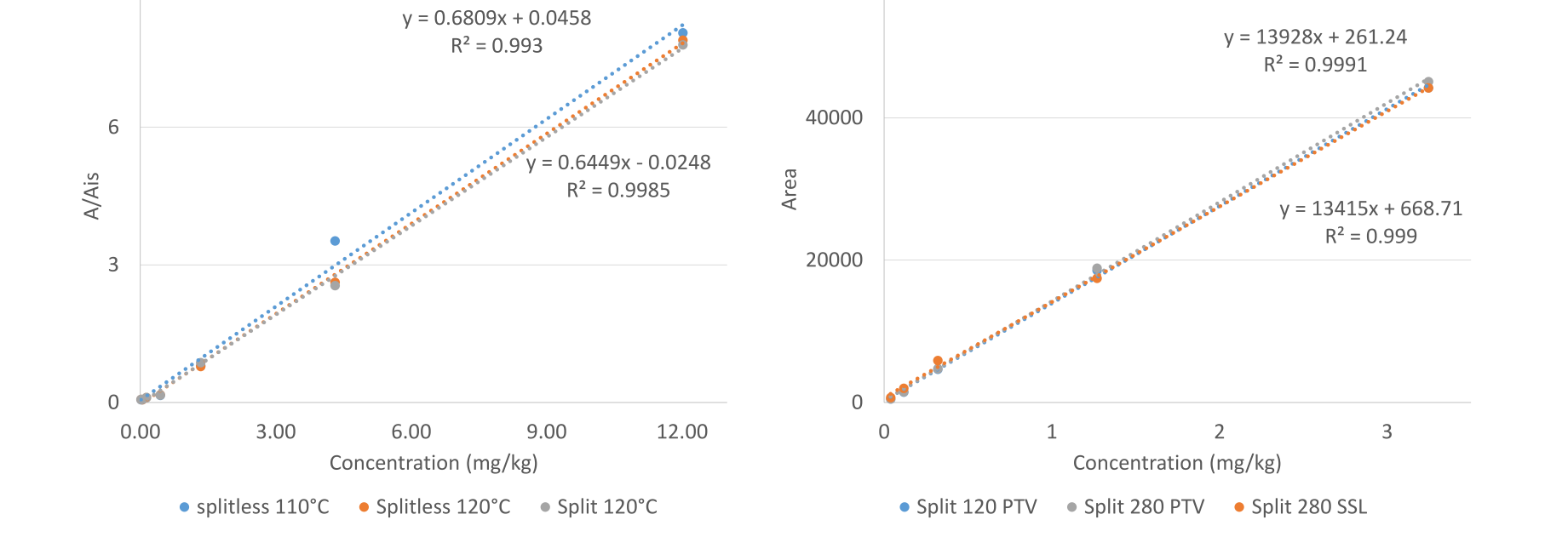
Initial temp. (°C)								
Splitless	RT1 (min)	RT2 (min)	Width1	Width2	Resolution			
95	8.03	8.07	0.030	0.035	0.762			
100	7.62	7.66	0.022	0.034	0.864			

Optimization of GC-MS method – Evaluation of both split and splitless methods

Comparison of calibration curves for both split and splitless injections

Comparison of calibration curves for split injection at different temperatures/inlets

105	7.22	7.26	0.023	0.021	1.126			
110	6.82	6.86	0.023	0.022	1.075			
115	6.42	6.46	0.025	0.025	0.944			
120	6.04	6.07	0.027	0.027	0.830			
Split								
95	8.03	8.07	0.020	0.028	1.033			
100	7.62	7.66	0.020	0.019	1.241			
105	7.22	7.26	0.021	0.019	1.209			
110	6.82	6.86	0.019	0.019	1.273			
115	6.43	6.47	0.019	0.019	1.242			
120	6.04	6.0	0.019	0.018	1.244			

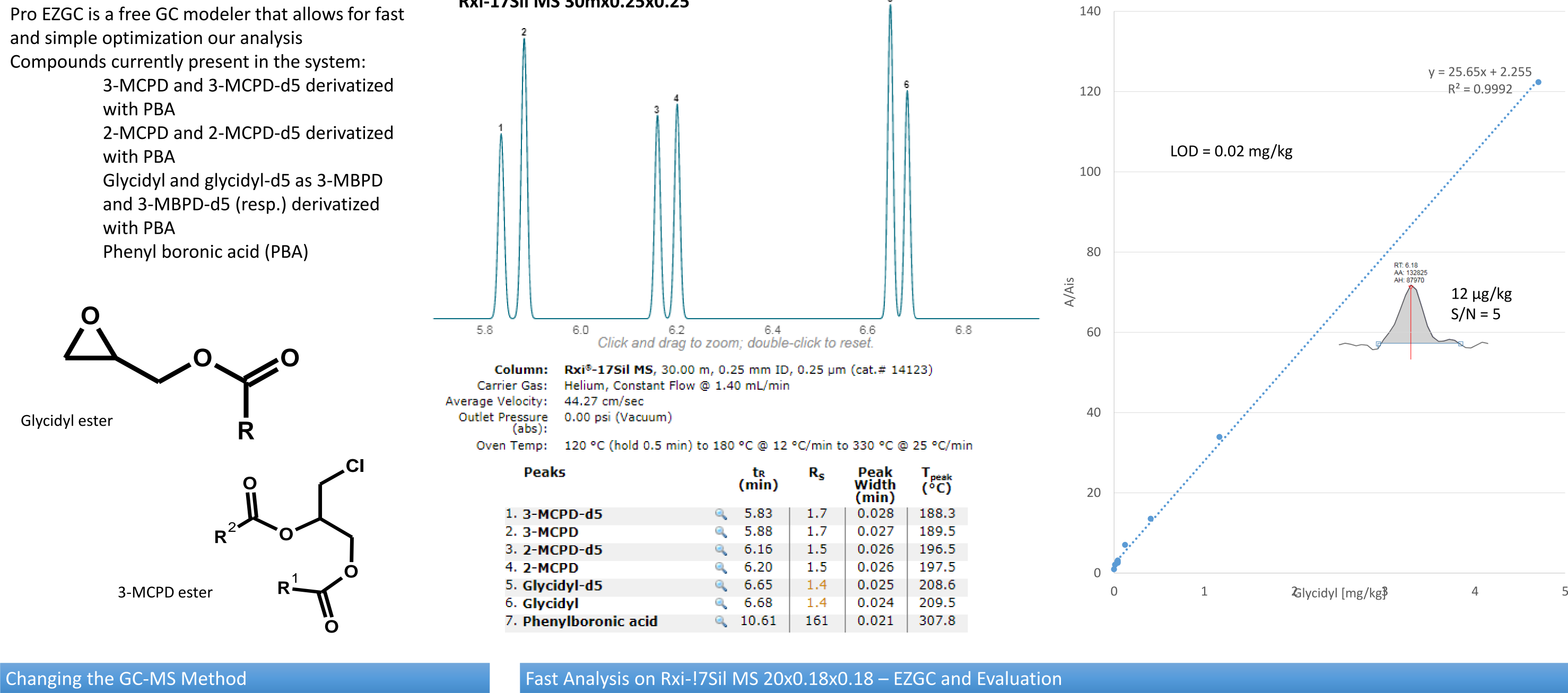


60000

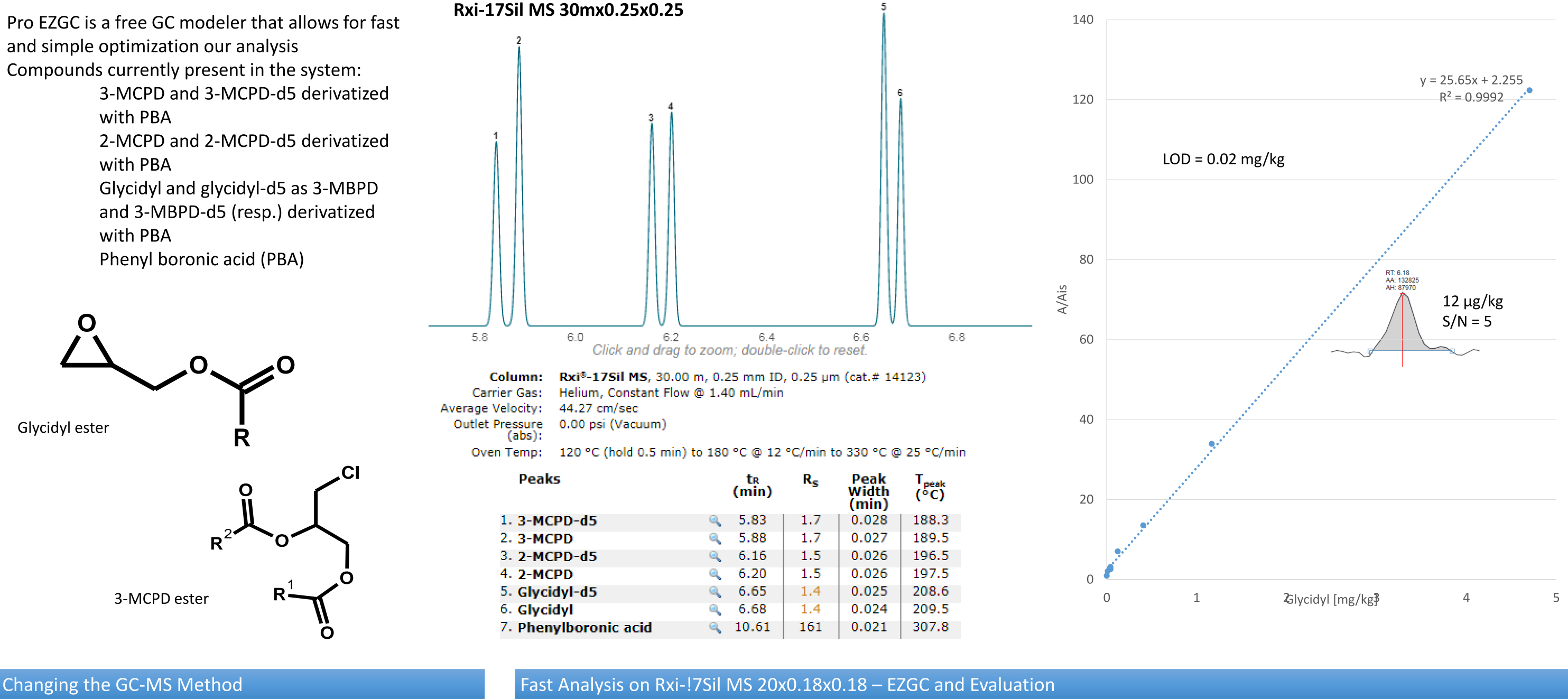
ProEZGC Model on Rxi-17Sil MS 30x0.25x0.25

Pro EZGC° Chromatogram Modeler Pro EZGC is a free GC modeler that allows for fast and simple optimization our analysis Compounds currently present in the system: 3-MCPD and 3-MCPD-d5 derivatized with PBA 2-MCPD and 2-MCPD-d5 derivatized with PBA

MCPD and GE analysis



Evaluation on GC-MS/MS



Original method:

Temp program: 85°C (0.5), 6°C/min to 150°C, 12°C/min to 180°C, 25°C/min to 280°C (7); total time: 24.8 min

New, optimized method: Temp program: 120°C (0.5), 12°C/min to 180°C, 25°C/min to 330°C (5); **total time: 16.5** min **Split 10:1**

Fast MCPD and GE analysis

Rxi-17Sil MS 20x0.18x0.18

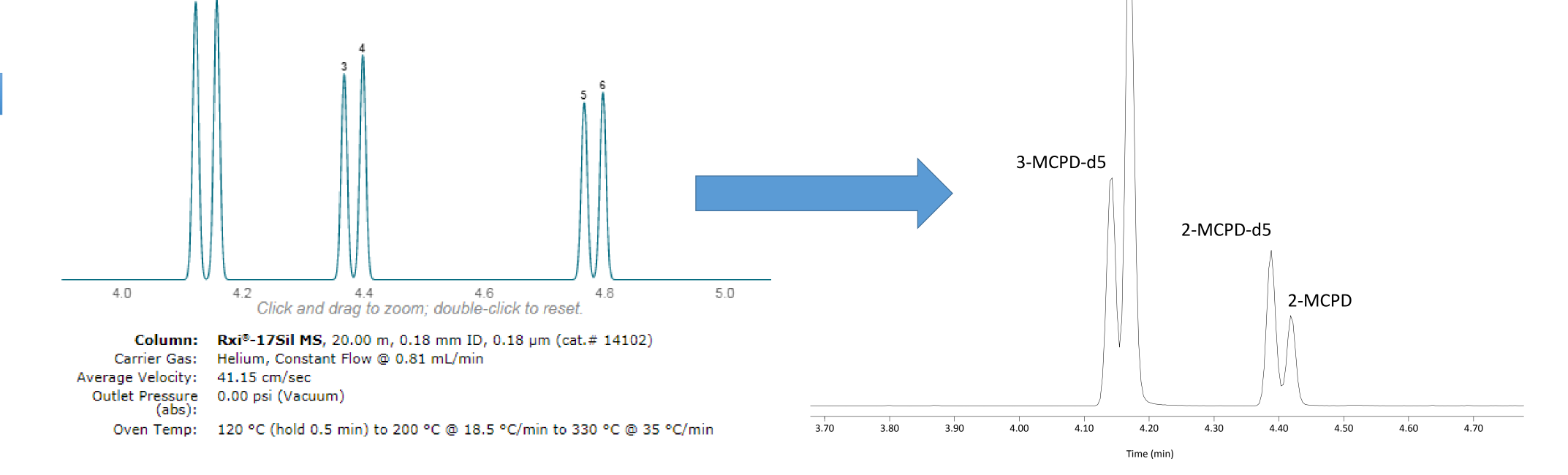
3-MCPD

Splitless time 0.5-1 min

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Conclusions

- An optimized GC-MS method led to improved peak shapes without detrimental effect on resolution. Manually optimized temperature program saved 8 minutes per analysis. **EZGC model can save up to 20** minutes.
- Switching to split injection had no negative effect on limits of detection.
- Using regular split/splitless injector had no effect on the performance. However, using guard column is recommended.





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