

Single Quad LC/MS Analysis of Organic Acids Using an Agilent Hi-Plex Column

Application Note

Food Testing

Abstract

Organic acids are highly hydrophilic and difficult to retain in reversed-phase mode. Conversely, ion exclusion mode is excellent for separating organic acids. Direct detection by UV or by postcolumn addition of an indicator compound however, does not provide the necessary level of sensitivity for many applications. This application note examines the separation of organic acids by ion exclusion mode, followed by direct detection on a single quadrupole MS.

Authors

Hayashi Keiko, Hiroki Kumagai, Kuniaki Matsushita, Kyoko Yasuda, Hirokazu Sawada, and Adam Bivens Agilent Technologies, Inc.



Introduction

Organic acid analysis by LC/MS is often performed using a reversed-phase mode. However, since many organic acids are highly hydrophilic, and have little retention in a reversed-phase column, coelution of interfering compounds in samples containing complex matrices is a substantial problem.

For organic acid analysis using an ion exclusion column, both detection by direct UV or by the postcolumn addition of a pH indicator such as BTB have been used. However, it is difficult to detect low concentrations. In this application note, organic acids were separated by ion exclusion mode, and examined by single quadrupole MS detection.

Experimental Conditions

Table 1. LC/MS Analysis Conditions

Parameter	Value
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Acquisition parameters

Agilent 1290 Infinity II LC			
Column	Agilent Hi-Plex H, 250 × 4.6 mm, 8 µm (p/n PL1570–6830)		
Mobile phase	0.01 % Aqueous formic acid/ACN = 80/20		
Flow rate	0.2 mL/min		
Postcolumn additive	0.01 % Aqueous $NH_4OH/ACN = 80/20$		
Additive flow rate	0.1 mL/min		
Temperature	50 °C		
Injection volume	10 µL		
Agilent 6120 Single Quadrupole LC/MS			
lon source	ESI		
Drying gas	N ₂ , 12 L/min		
Dry gas temperature	200 °C		
Nebulizer pressure	50 psi		
Capillary voltage	3,000 V		
Mode	Negative		

SIM

Table 2.	SIM	SIM Ion List		
		М	M-H	
Acatio	aid	60	E0	

	М	M-H	Fragmentor (V)
Acetic acid	60	59	80
Glyoxylic acid	74	73	80
Propionic acid	74	73	80
Glycolic acid	76	75	80
Butyric acid	88	87	80
Pyruvic acid	88	87	80
Lactic acid	90	89	80
Oxalic acid	90	89	80
Valeric acid	102	101	80
Malonic acid	104	103	60
Fumaric acid	116	115	80
Levulinic acid	116	115	80
Maleic acid	116	115	80
Succinic acid	118	117	80
Pyroglutamic acid	129	128	60
Itaconic acid	130	129	60
Malic acid	134	133	60
Adipic acid	146	145	80
Ketoglutaric acid	146	145	80
Tartaric acid	150	149	100
Ascorbic acid	176	175	100
Citric acid	192	191	100
Gluconic acid	196	195	100

Post column ammonium hydroxide addition was made using a Micro T-connector (p/n 5042-8519).

Results and Discussion

Figure 1 shows the SIM chromatogram of a 1 ppm standard solution of each organic acid. A peak was detected for all organic acids.

The S/N value (signal/[peak-to-peak noise] × 2) is shown in Table 3.

Analysis of commercially available yogurt was conducted. Yogurt whey was sampled and filtered through a 0.22 μ m filter, and diluted 100 times to prepare a sample. Figure 2 shows the chromatogram of the detected organic acids in this yogurt sample. In Figure 3, a 0.1 ppm malic acid standard was added to this sample, and a recovery test was conducted. The recovery rate was 93 %, with good overall results. In Figure 4, excellent linearity is demonstrated using malic acid over a range of 10 ppb to 0.25 ppm.



Table 3. Retention Time of Each Organic Acid and S/N at 1 mg/L

5.897

6.048

6.237

6.49

6.791

6.964

7.051

7.789

8.005

8.088

8.154

8.463

8.921

9.059

9.745

10.029

10.123

10.714

11.016

12.049

12.764

13.117

13.964

RT (min)

S/N

5.8

829.3

36.3

593.8

527.5

18.2

392.9

84.0

52.1

1,254.1

2,740.3

429.4

226.8

555.0

1,709.1

1,339.1

2,682.3

687.1

652.6

1,711.9

1,288.4

2,873.6

58.6

[M-H]-

89

115

145

87

191

103

149

133

73

115

195

175

129

117

145

75

89

128

115

59

73

87

101

Oxalic acid

Maleic acid

Pyruvic acid

Malonic acid

Tartaric acid

Malic acid

Glyoxylic acid

Fumaric acid

Gluconic acid

Ascorbic acid

Itaconic acid

Succinic acid

Adipic acid

Lactic acid

Glycolic acid

Levulinic acid

Propionic acid

Butyric acid

Valeric acid

Acetic acid

Pyroglutamic acid

Citric acid

Ketoglutaric acid

Ketoglutaric acid Citric acid Malic acid Succinic acid Lactic acid Pyroglutamic acid

Figure 2. Organic acids detected from commercially available yogurt.

Figure 1. Chromatogram of organic acid standard solution (1 mg/L).



Figure 3. Chromatogram of the malic acid addition recovery test.

Table 4. Area Value in the Malic Acid Addition Recovery Test

Sample	Area
Sample	1,746.8
Sample + 0.1 mg/L spike	10,117.8
0.1 mg/L	7,690.7

Conclusion

A panel of organic acid standards was separated by ion exclusion chromatography, and detected with single quadrupole MS. Good separations were obtained, and peaks were detected at 1 ppm for all organic acids tested.

When applied to a yogurt sample, key organic acids were detected with good sensitivity. Further tests on this real-world sample showed excellent recovery and linear detection down to 10 ppb.

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Figure 4. Linearity of malic acid 0.01–0.25 ppm (A) and SIM chromatogram (B).

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