

Quantitation and Identification of the Pesticide Malathion in Fruit Samples using MRM³ Workflow

On the QTRAP[®] 5500 System

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LC-MS/MS instruments operating in Multiple Reaction Monitoring (MRM) are widely used for targeted quantitation on triple quadrupole and hybrid triple quadrupole linear ion trap (QTRAP[®]) systems because of their well-known selectivity and sensitivity. Although this double mass filtering greatly reduces noise there is always a chance that elevated background levels or matrix signals interfere with the targeted analyte.

One possibility of improving quantitative results is to use a more selective detection mode, such as MRM³ workflow¹. In comparison to MRM mode, MRM³ often provides higher selectivity due to one additional fragmentation step, effectively reducing interferences and improving quantitation.

Improved Selectivity for Quantitation

Here, a homogenized apple was spiked with the organophosphorus pesticide Malathion (10 ppb), then extracted



using a QuEChERS procedure, diluted 50 times to minimize matrix effects, and analyzed by LC-MS/MS. The resulting chromatograms using two MRM transitions and two MRM³ experiments are shown in Figure 1. The MRM transition 331/127 showed expected selectivity, while the second transition 331/99 had an elevated background level and also matrix interference. In contrast both MRM³ experiments showed superior selectivity for reliable quantitation. Table 1 shows signal-to-noise ratios (S/N) of the chromatograms described above.

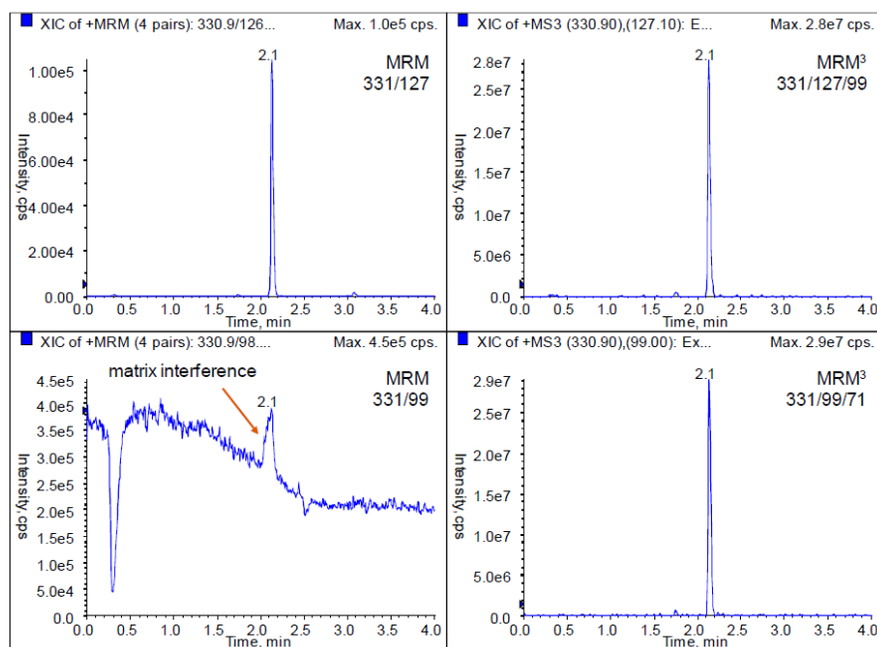


Figure 1: Comparison of MRM vs MRM³ for Detecting Malathion (10ppb) in Apple Extract. (Left) While the MRM transition 331/127 showed expected selectivity, the second transition 331/99 had an elevated background level and also matrix interference. This results in incorrect ion ratios and prevents confident identification. (Right) In contrast, both MRM³ experiments showed minimal interferences and higher selectivity.

Improved Confirmation

The quantifier MRM and the quantifier MRM³ had very similar sensitivity. Both experiments allowed quantifying Malathion at sub ppb levels. However, pesticide testing requires identification. Thus, a second MRM or MRM³ signal has to be recorded to allow ratio calculation (qualifier/quantifier). In this example, the second MRM showed a dramatic loss in S/N because of the elevated background. In this case the quantitation and identification of Malathion in fruit matrix using MRM³ was much more sensitive than in MRM mode.

Method Details

Chromatography: Separation was performed using a Shimadzu UFLC_{XR} system with a Phenomenex Synergi Fusion-RP (2.5 μ m) column and a fast gradient of water and methanol with ammonium formate buffer.

Mass Spectrometry: A looped MS experiment was performed using the SCIEX QTRAP 5500 System with Turbo VTM Source and ESI probe. The first experiment consisted of two MRM transitions of 100 msec dwell time each. The second experiment was two MRM³ scans using a scan speed of 20000 Da/s with 20 msec fill time and 25 msec excitation time. Total cycle time of only 0.33 sec which allowed ~15 data points across the UHPLC peaks that had a base-to-base peak width of only 5 sec. The detected masses in MRM and MRM³ mode with compound dependent parameters are shown in Figure 2.

Data Processing: Data was processed with MultiQuantTM Software.

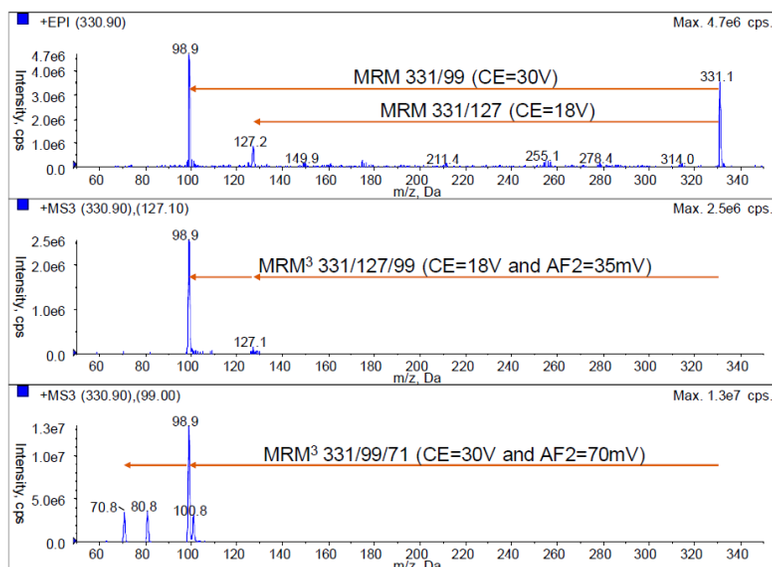


Figure 2. Developing the MRM and MRM³ Workflow Methods. (Top) Full scan MS/MS of Malathion shows two fragments at m/z 127 and 99 for use in the MRM experiment. (Middle) MS/MS/MS of 331/127 shows a fragment at m/z 99 for use in the first MRM³ experiment. (Bottom) MS/MS/MS of 331/99 shows a fragment at m/z 71 for use in the second MRM³ experiment.

Improved Reproducibility

Several fruit samples were fortified with 10 ppb Malathion and analyzed in replicates. While MRM detection suffered from matrix interference, MRM³ resulted in more accurate and reproducible data. The %CV values in apricot, apple, pear, and orange were <5% with accuracies ranging from 90% to 110%. In addition, the MRM³ ratio calculation clearly identified the presence of Malathion in the analyzed fruits.

Table 1. Improvements in Detection Using MRM³ Workflow. Signal-to-noise ratios (S/N) of MRM and MRM³ of 10 ppb Malathion in a 50-times diluted QuEChERS extract of apple.

MRM	S/N	MRM ³	S/N
331 / 127	922	331 / 127 / 99	497
331 / 99	8*	331 / 127 / 71	147

* High background – see Figure 2

Summary

LC-MS/MS methods were developed to quantify and identify the organophosphorus pesticide Malathion in fruit samples using the QTRAP[®] 5500 System. The traditional MRM and MRM³ modes were compared regarding selectivity, sensitivity, accuracy and reproducibility. The results show that the higher selectivity of MRM³ eliminates background and matrix interference, resulting in better data quality. MRM³ gave comparable data versus MRM for the quantifier signal but much better sensitivity, accuracy, and reproducibility for the qualifier signal in fruit matrix.

References

1. MRM³ Quantitation for Highest Selectivity in Complex Matrices. SCIEX Technical Note RUO-MKT-02-2739-A.

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