

Verification of the mycotoxin patulin from apple juice with isocratic HPLC

Kristin Folmert, Kate Monks; applications@knauer.net

KNAUER Wissenschaftliche Geräte GmbH, Hegauer Weg 38, 14163 Berlin; www.knauer.net

SUMMARY

The deleterious and mutagenic mould fungus product patulin can frame during wrong cidermaking process of fruits like apple. The permitted daily exposure for patulin, was declared as 0.4 µg/kg body weight, by the scientific committee for foodstuffs of the European commission. [1] The specified determination routines have to be performed with HPLC according to official guidelines. Here, a fast protocol for the determination of Patulin from apple juice with AZURA® HPLC plus and a Eurospher II column is described.

INTRODUCTION

The lactone Patulin belongs to the chemical group of polyketides. It is soluble in acidic water and many organic solvents like methanol. Different species of fungi, such as *Aspergillus*, *Byssoschlamys*, and *Penicillium*, growing on rotting parts of fruits including apples, cherries, plums, strawberries, blueberries and pears can produce the mycotoxin patulin. [2] Juices can be contaminated with patulin if affected fruits are used for the cidermaking process. As a result of its thermal stability, patulin cannot be destroyed by pasteurization or thermal denaturation. Next to its antibiotic qualities, patulin is implicated as a possible carcinogen but the toxicity of patulin is primarily through its affinity to sulphhydryl groups which results in inhibition of enzymes. [3] Major acute toxicity findings include gastrointestinal problems, neurotoxicity, pulmonary congestion, and edema. [3] To protect customers from patulin in juices like apple juice, the FDA and the European commission (EU) recommend a maximum daily ingestion concentration of 0.4 µg/kg body weight for humans according to 93/5/EWG. Furthermore, the EU has set a limit of 25 µg/kg in solid apple products and 10 µg/kg in baby food (EC 1881/2006). [1]

RESULTS

The very fast and robust method with AZURA HPLC plus for the identification of patulin in juices like apple juice enables the validation of the concentration according to the regulation of the EU and FDA and at lower concentrations. The minimum concentration, which was detected with a 50 µL injection was 0.1 µg/mL apple juice. The short retention times of hydroxymethylfurfural (HMF) and Patulin enable a high efficiency for the analysis. A good separation from all matrix peaks ($R_s > 1.5$) comfortably assures the quality of the measurements, as shown in Fig. 2 A for the concentration of 0.25 µg/mL patulin and HMF.



Fig. 1 Patulin contamination in apple juice

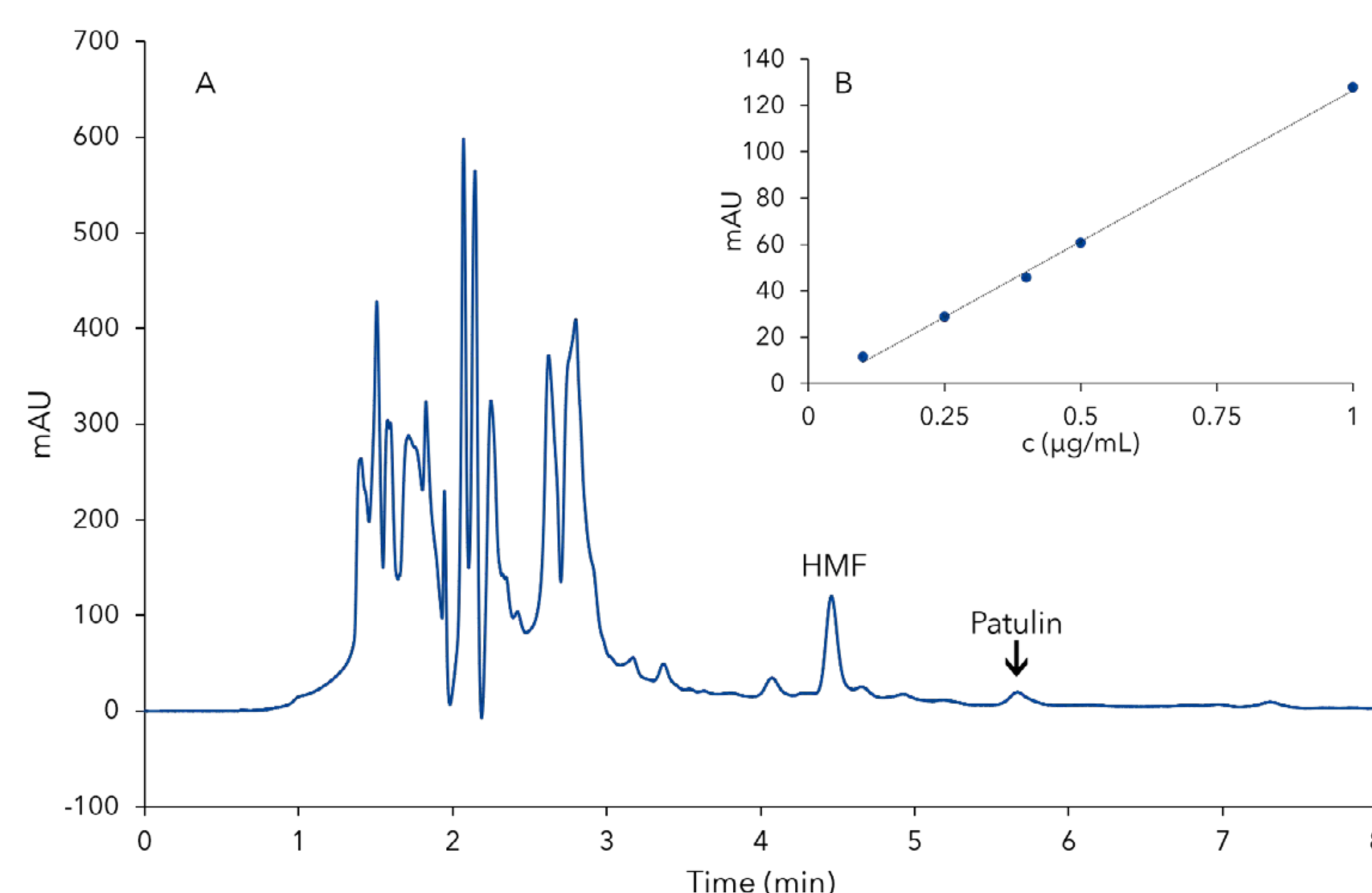


Fig. 2 Chromatogram of 50 µL injections from apple juice concentrate spiked with 0.25 µg/mL patulin and HMF (A) and an concentration curve of patulin with the same injection volume (B). Concentrations of 1.00 µg/mL, 0.50 µg/mL, 0.40 µg/mL, 0.25 µg/mL, 0.10 µg/mL and 0.05 µg/mL patulin (n=3) for the concentration curve

MATERIALS AND METHODS

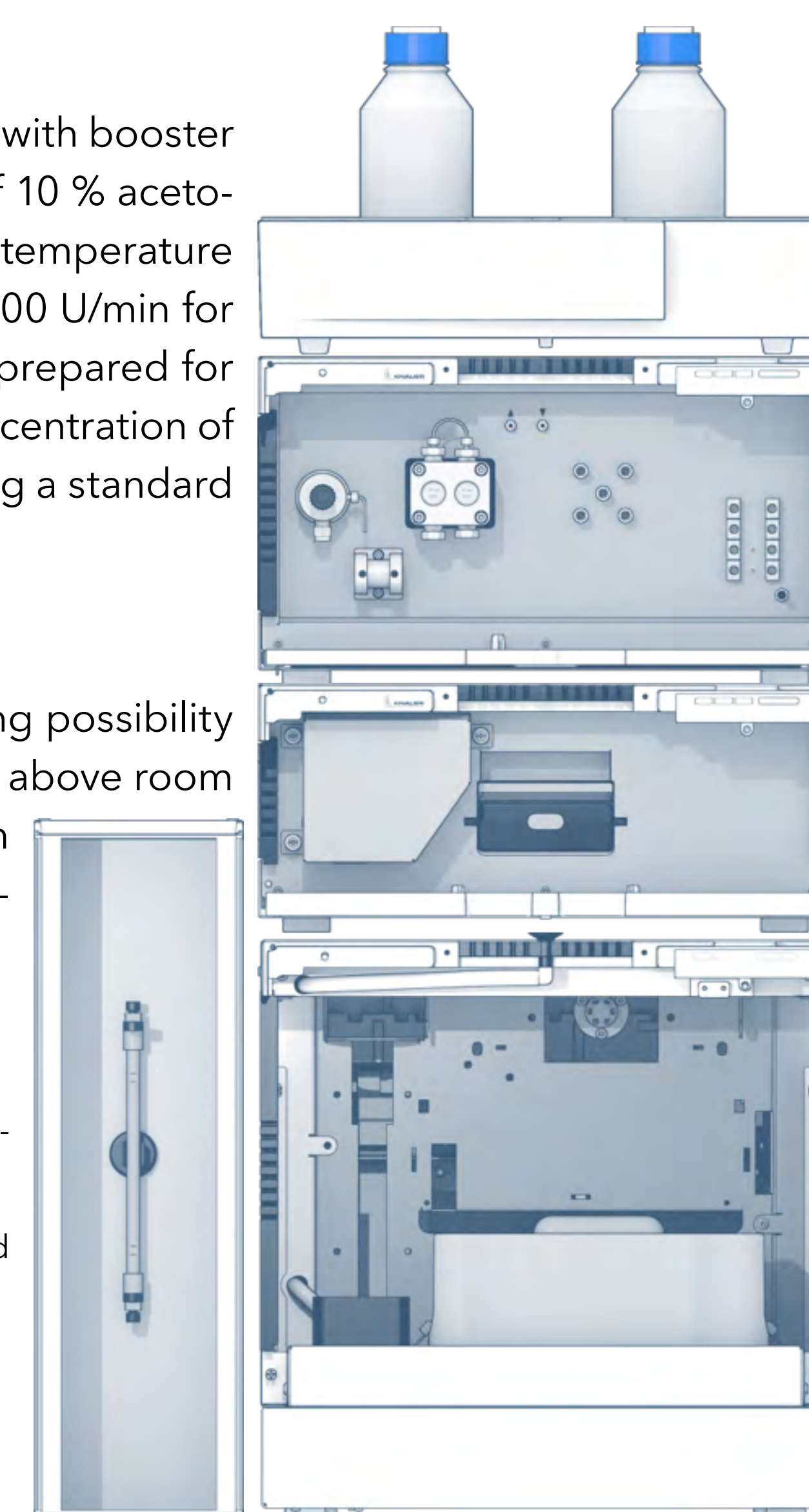
An AZURA HPLC plus system with the possibility to cool the autosampler rack to restore the quality of the food samples and a DAD detector with booster flow cell to monitor impurities at different wavelength was used to develop a simple and fast isocratic method. The eluent was a mixture of 10 % acetonitrile in water. The separation was realized within 7 minutes and with a phenyl endcapped Eurospher II 100-5 Phenyl column at an ambient temperature of 40°C. Therefore, 10 mL of gravy juice were mixed with 600 µL Pectinase enzyme and incubated at 37°C for 3 h before centrifuged at 3500 U/min for 10 min. After removing the centrifugate, the liquid was centrifuged again with the same conditions. The now concentrated apple juice is prepared for HPLC analysis. For the measurements in Fig. 2 A, the concentrated juice was mixed with a standard solution of HMF and Patulin to gain a concentration of 0.25 µg/mL. To evaluate the concentration in apple juice a concentration curve between 0.1 µg/mL to 1 µg/mL of patulin was measured using a standard of patulin in water (Fig. 2 B).

CONCLUSION

A fast and efficient HPLC method was realized with AZURA HPLC plus and an isocratic method in only 7 minutes retention time. The cooling possibility of the autosampler elongates the life time of sensitive food samples and a column thermostat enables a continuous separation atmosphere above room temperature, which is crucial for the high robustness of the method. With these advantages the identification of patulin from apple juice can be realized down to the very low concentration of 0.1 µg/mL, which is four times less than the recommended maximum daily ingestion concentration for humans according to 93/5/EWG of the EU.

REFERENCES

- [1] Protokoll of the 120. meeting of the scientific foodstuff committee 8th-9th march 2000 in Brüssel: https://ec.europa.eu/food/sites/food/files/safety/docs/cs_contaminants_catalogue_patulin_out55_en.pdf
- [2] G. C. Llewellyn, J. A. McCay, R. D. Brown, D. L. Musgrove, L. F. Butterworth, A. E. Munson, Jr. K. L. White, Immunological evaluation of the mycotoxin patulin in female B6C3F1 mice. Food Chem. Toxicol. 1998, 36, 1107-1111.
- [3] O. Puel, P. Galtier, I. P. Oswald, Biosynthesis and Toxicological Effects of Patulin, Toxins, 2010, 2 (4): 613-631.



Verification of the mycotoxin patulin from apple juice with isocratic HPLC

ADDITIONAL MATERIALS AND METHODS

Tab. A1 Method parameters

Eluent	ACN: _{dd} H ₂ O/1:9 (v:v)		
Flow rate	1 mL/min		
Column temperature	40°C	Injection volume	50 µL
Autosampler temperature	4°C	Injection mode	Full loop
Detection	275 nm	Data rate	10 Hz
		Time constant	0.1 s

Tab. A2 Instrument set up

Instrument	Description	Article No.
Pump	AZURA P6.1L, LPG	APH34EA
Autosampler	AZURA AS 6.1L	AAA01AA
Detector	AZURA DAD 6.1L	ADC11
Flow cell	LuightGuide 50 mm	AMD59XA
Column thermostat	AZURA CT 2.1	A05852
Column	KNAUER Vertex Plus column, Eurospher II 100-5 Phenyl, 150 x 4.6 mm	15VE050E2J
Software	ClarityChrom 7.4.2 - Workstation, autosampler control included	A1670
	ClarityChrom 7.4.2 - PDA extension	A1676

RELATED KNAUER APPLICATIONS

[VFD0042J](#) - Analysis of Flavonoids in Fruit Juice

[VFD0002J](#) - Determination of Naringine and Hesperidine in Fruit Juices

[VFD0152](#) - Determination of Aflatoxin M1 in Milk