



Order No.: MYS-05

Qualitative determination of the sum of **Fumonisins (FUM)**

Includes sample preparation with designed for use with **aokin FP analyzer**.

Package content:

Order No. MYS-05 **aokin**mycontrol**FUM**

Order No	<u>Material</u>	Content	Application	<u>Storage</u>
MYS-05-20 for 20 measurements	aokinmycontrol FUM Kit for qualitative determination of Fumonisin (FUM)	aokinExtractionSalt FUM ①, preparations 1 pouch for 20 preparations aokin Easy Extract Liquid FUM ②, 1 vial sticker for aokin EasyExtractBuffer FUM aokin ReactionBufferFUM 100x ④, 1 vial 1,8mL FUM Reagent ⑤, 1 unit FUM Tracer ⑥, 1 unit FUM positive control, Standard 1836.5ng/mL FUM B1 in Acetonitril/Water, 100µl ⑥; 1 unit	Measurement on aokin FP analyzer	Refrigerated Do not freeze
MYS-05-100 for 100 measurements	Aokinmycontrol FUM Kit for qualitative determination of Fumonisin (FUM)	aokinExtractionSalt FUM , 1 unit for 100 preparations aokin Easy Extract Liquid FUM (2), 1 unit 4x sticker for aokin EasyExtractBuffer FUM aokin ReactionBufferFUM 100x (4), 1vial 7mL FUM Reagent (5), 5 units FUM Tracer (5), 5 units FUM positive control, Standard 1836.5ng/mL FUM B1 in Acetonitril/Water, 100µl (5); 1 unit	Measurement on aokin FP analyzer	Refrigerated Do not freeze

Analytical-kit for rapid and quantitative determination of Fumonisin (FUM) in oats, wheat, corn and other grains.



Materials

All materials provided are precisely weighed and calibrated. Control of the volume and concentration of the individual solutions are essential for the precision of the analysis.

Quality control:

All materials and reagents are prepared according to strict quality control protocols. Exchanging reagents between kits having different Lot-numbers will lead to erroneous results and is not permitted.

Preparation of buffers:

MYS-05-20 for 20 measurements:

Mix 190 mL <u>destilled</u> or <u>deionized</u> water with 10 mL of **aokin** Easy Extract Liquid **FUM** ② and 800 mL of Methanol = **aokin** EasyExtractBuffer **FUM**.

Prepare the solution and label the container with the sticker **aokin** EasyExtractBuffer **FUM** included in the kit, on which you can note the preparation date.

Mix 1ml **aokin** ReactionBuffer**FUM 100x** ④ with 99ml <u>destilled</u> or <u>deionized</u> water = **aokin** ReactionBuffer**FUM** ④

The buffer has to be brought to room temperature (best 18-26°C) for usage.

MYS-05-100 for 100 measurements:

Mix 190 mL <u>destilled</u> or <u>deionized</u> water with 10 mL of **aokin** Easy Extract Liquid **FUM** ② and 800 mL of Methanol = **aokin** EasyExtractBuffer **FUM**.

Prepare 4x the solution and label the container with the sticker **aokin** EasyExtractBuffer **FUM** included in the kit, on which you can note the preparation date.

Mix 5ml **aokin** ReactionBuffer**FUM 100x** (4) with 495ml destilled or deionized water = **aokin** ReactionBuffer**FUM** (4)

The buffer has to be brought to room temperature (best 18-26°C) for usage.

Storage and stability:

The diluted **aokin** EasyExtractBuffer **FUM** ② is stable for 12 month without refrigeration (15° - 30°C).

aokin ReactionBuffer FUM 4) is stable for 12 months without refrigeration (15° - 30°C).

Kit reagents (**FUM** Reagent \odot , **FUM** Tracer \odot , positive control **FUM** \odot) must be stored in a refrigerator (2-10°C).

Any other components can be stored at ambient temperature.

Do not use expired or contaminated components, or components from other kits. Do not mix components from different manufactured lots. A certificate of analysis (COA) is available upon request at info@aokin.com.

Materials and instrumentation required but not provided:

aokin FP analyzer, 10x75 mm borosilicate glass test tubes for **aokin FP analyzer**, Micro-pipette: 10 – 100 μ L, 100-1000 μ L and related pipette tips, 2 mL collection tubes, tube vortex mixer, centrifuge, timer, deionized water for buffer reconstitution, methanol.

Alternative to the **aokin FP analyzer** other FP single tube readers or FP microplate readers can be used. Depending on instrumentation the applied volumes need adjustment. Black flat bottom microtiter plates are recommended for use with FP microplate readers.



Introduction

aokinmycontrol **FUM** is a rapid screening method for Fumonisin (FUM). It has been specifically designed and calibrated for the analysis of food and feed and includes a sample preparation with matrix removal columns (**aokin**QuickClean **FUM**). Samples in the μ g/kg range (μ g/kg = ppb) can be analysed for FUM in under 40 minutes.

aokinmycontrol **FUM** is available with a calibration, which has been validated for grains and other food and feed products. Daily checks of negative control and positive control (either liquid standard or reference matrix extract) are required to ensure reliable results.

FPA technology works on clear homogeneous liquids without particles or emulsions, therefore avoid practices that may contaminate the test controls, test buffer, or test reagents. **aokin** will gladly assist you customizing the test for your specific sample type and application. Please do not hesitate to contact us.

Polarization readings are affected by temperature. Polarization readings decrease by around 3 mP (millipolarization units) for a temperature increase of 4°C.

The method is best used at constant temperatures between controls and sample measures. If a temperature change of >10°C is observed, a new calibration is recommended.

Health:

All materials in this kit should be treated as any other laboratory chemicals. Avoid ingestion, eye contact and other potential detrimental exposure. A material safety data sheet (MSDS) is available upon request.

Negative controls:

Negative control should be run in triplicate at regular intervals with the single tube assay or on every microtiter plate, respectively. Avoid temperature variations of more than 1°C during one determination. Mix well with a vortexer and wait 60 s before taking the value. Be careful to avoid bubbles when pipetting into microtiter plates.

Negative control FUM = **aokin** ReactionBuffer **FUM** ④

Positive controls:

Positive controls are recommended as an additional measure to adjust the calibration for small temperature changes. Reference matrix materials are best used to determine the recovery rate and to do a recovery rate correction if needed.

Positive controls FUM (5) = Standards diluted in **aokin** ReactionBuffer **FUM** (4)

Dilute included **Fumonisin B1 1836.5** ng/ml in Acetonitrile/Water Lot:00511837240720279 \odot with Negative control FUM IAC (Dilutionfactor **1:70** -> example: add 10 µL of **Fumonisin B1 1836.35** ng/mL in Acetonitrile/Water to 690 µL Negative control FUM (= *aokin ReactionBuffer* FUM 4) [26.2ng/mL]. Use 5 to 90 µL within the assay, fill up with Negative control FUM for overall 100 µL sample volume.

(Example: 50 μ L of **1:70 diluted Fumonisin B1 1836,5 ng/mL in Acetonitrile/Water** [26.2ng/mL] + 50 μ L Negative control FUM = 13.1 ng/mL Fumonin B1 within the assay)

Fumonisin

Fumonisin is a mycotoxin. It naturally occur in molds by *Aspergillus* and *Penicillium* species and are toxic to humans and animals. As a consequence, it is strongly recommended to monitor the content in grain and corn food and feed raw materials and products.

Figure 1: Chemical formula for Fumonisin B1, C₃₄H₅₉NO₁₅, Molecular weight: 721.83 g/mol



Standard sample preparation for wheat and corn (Page 7/7)

1. Preparation

1.1. Prepare buffer:

Mix 5ml **aokin** ReactionBuffer**FUM 100x** ④ with 495ml <u>destilled</u> or <u>deionized</u> water = **aokin** ReactionBuffer**FUM** ④

Allow aokinReactionBuffer to reach room temperature (15°C to 30°C) before use.

aokin EasyExtractBuffer **FUM**: Mix 190 mL <u>destilled</u> or <u>deionized</u> water with 10 mL of **aokin** Easy Extract Liquid **FUM** and 800 mL of Methanol.

2. Instrument adjustment

Instruments must be set up and calibrated according to manufacturer's specifications. For detailed information, please consult instrument manual and contact technical support.

3. Sample collection, grinding and mixing

The analysis sample is collected, ground, and mixed/homogenised according to an approved procedure. For convenience volumes may be ground and mixed by using a knife-blender.

4. Weighing and extraction

Weigh 15 g of your sample, add one spoon (1.5 g) of **aokin**ExtractionSalt **FUM** and 35 ml **aokin** EasyExtractBuffer **FUM** directly into an extraction beaker. Preferentially the exact volume is applied using a dispensette.

Extract sample by blending for 3.5 minutes at high speed. The recommended protocol has blending times alternating with resting time to avoid heating of the sample and is as follows: mix for 30 seconds, pause for 1 minute, mix for 30 seconds and so on (until 3.5 minutes of blending time). Preferably use a preprogrammed timer (*aokin*watchbox) to conveniently and automatically complete this extraction protocol.

5. Filtration or centrifugation

Filter the raw extract through a fluted filter. Discard the filter cake and use the filtrate. Alternatively centrifuge the raw extract and use the supernatant.

6. Precipitation

Add 400 μ l of the filtrate to 400 μ L of the **aokin** ReactionBuffer **FUM** ④, mix it well and centrifuge with maximum g-force (> 10.000 x g) for 20 minutes. Transfer 100 μ l of the supernatant into 400 μ L of the **aokin** ReactionBuffer **FUM** ④, mix it well and centrifuge with maximum g-force (> 10.000 x g) for 10 minutes. Transfer the supernatant into a clean tube. Your sample is now ready for analysis.

7. Introduction of general testing procedure

- 7.1. Pipette 965 µl of **aokin**ReactionBuffer **FUM** (4) into the test tube.
- 7.2. Add 100 µl of sample extract <u>or</u> a control into the test tube. (if necessery fill up with **aokin** ReactionBuffer **FUM** 4) if sample eluat is too concentrated, example: 50 µl sample + 50 µl **aokin** ReactionBuffer **FUM** 4) for overall 100 µl sample volume) <u>or</u> a control into the test tube.

Note: Run negative controls in triplicate, and positive control and samples as duplicates for best performance.

- 7.3. Add 15 µL of **FUM** Reagent (**VIOLET** cap) into the test tube.
- 7.4. Mix well/vortex (without spilling).
- 7.5. Obtain background measurement readings of all samples and controls. Wait a minimum of 1 minute and a maximum of 15 minutes between mixing and reading.
- 7.6. Add 20 µL of **FUM** tracer (**YELLOW** cap) into the first test tube.
- 7.7. Mix well.
- 7.8. Incubate for 60 seconds.
- 7.9. Obtain tracer measurement of the sample.
- 7.10. Repeat steps 8.6 to 8.9 for all test tubes.

Note: Include one set of control after each hour of testing, or on each microtiter plate. Performing measurements of negative controls regurarly will ensure the accuracy of your determinations.



8. Data acquisition and analysis

- 8.1. Mix and Read Process
- 8.1.1. Mix buffer, sample and reagent in exact volumes and at exactly specified time intervals. Read background fluorescence in horizontal and vertical direction.
- 8.1.2. Add tracer, incubate for exactly the specified time. Read emission values in horizontal and vertical direction.

8.2. **Background information**

The fluorophore is excited with polarized light, depending on the rotational velocity of the fluorophore (or the fluorophore antibody complex) the emission will also be polarized. The emission intensities using horizontally directed and vertically directed polarization filters are detected for the background and for the reaction. Followed by an automatic calculation of polarization from the value changes in vertical and horizontal direction. $P = (\Delta H - \Delta V)/(\Delta H + \Delta V)$.

8.3. Software: direct use of excel worksheet - aokin mycontrol version of software - alternatively use user interface

Advice: Use a copy of the empty original excel worksheet to do all measurements of one series. When you start with a new series including negative controls, positive controls and sample analysis measurements, use a new copy of the empty original worksheet saved previously with the name of your series and the date.

Note:

- Connect computer with aokin FP analyser
- Install the **aokin** software
- One excel row = one experiment
- Green cells can be written into
- Yellow cells are for measuring: highlight a cell in the correct line and press F3
- Brown cells are for information only
- Light green cells contain evaluations = calculations and end results

10. Daily Check I:

- a. Use negative control (= aokin ReactionBuffer FUM 4) in Step 7.2
- b. Software: Choose Daily Check I in the user interface or alternatively Offset at given $T \circ C$ Sheet in particular excel version of software
- 10.1. Pipette 965 μl of **aokin**ReactionBuffer **FUM** (4) into a test tube.
- 10.2. Add 100 µl of a negative control into the test tube. The negative control for FUM is the **aokin** EasyExtractBuffer **FUM** as prepared above.
- 10.3. Add 15 μL of **FUM** Reagent (**VIOLET** cap) into the test tube.
- 10.4. Mix well/vortex (without spilling).
- 10.5. Obtain background measurement reading. Wait a minimum of 1 minute and a maximum of 15 minutes between mixing and reading.
- 10.6. Add 20 μ L of **FUM** Tracer (YELLOW cap) into the first test tube.
- 10.7. Mix well.
- 10.8. Incubate for 60 seconds.
- 10.9. Obtain tracer measurement

Note: Run negative controls in triplicate.

- 11. Daily Check II-A:
- a. Use liquid positive control in Step 7.2
- b. Software: Choose Daily Check II, Liquid standard in the user interface or Recovery rate at given T°C Sheet in particular excel version of software
- 11.1. Pipette 965 μ l of **aokin**ReactionBuffer **FUM** 4 into a test tube.
- 11.2. Add 100 µl of a positive control into the test tube. The positive control is a liquid standard is 26.2ng/mL Fumonisin B1 Standard diluted in **aokin** ReactionBuffer **FUM** ④, use 5 to 90µl within the assay); fill up with **aokin** ReactionBuffer **FUM** ④ for overall 100µl sample volume (example: 50µl positive control Fumonisin B1 + 50µl **aokin** ReactionBuffer **FUM** ④ = 13.1ng/mL Fumonisin B1 within the Assay)
- 11.3. Add 15 µL of **FUM** Reagent (**VIOLET** cap) into the test tube.
- 11.4. Mix well/vortex (without spilling).
- 11.5. Obtain background measurement reading. Wait a minimum of 1 minute and a maximum of 15 minutes between mixing and reading.



- 11.6. Add 20 µL of **FUM** Tracer (YELLOW cap) into the first test tube.
- 11.7. Mix well.
- 11.8. Incubate for 60 seconds.
- 11.9. Obtain tracer measurement

Note: Run positive controls in duplicate.

12. Alternative Daily Check II-B:

- a. Use of final extract of a reference matrix material as positive control in Step 7.2
- b. Software: Choose Daily Check II, Reference Matrix in the user interface or Recovery rate at given T°C Sheet in particular excel version of software
- 12.1. Choose a reference matrix sample most similar to your samples. Example: In case you analyse wheat use a wheat reference matrix sample containing a known amount of mycotoxin.
- 12.2. Prepare the reference matrix sample according to Steps 4, 5, 6, 7.
- 12.3. Pipette 965 µl of **aokin**ReactionBuffer **FUM** ④ into a test tube.
- 12.4. Add 100 µl of the Reference Matrix extract into the test tube.
- 12.5. Add 15 μL of **FUM** Reagent (**VIOLET** cap) into the test tube.
- 12.6. Mix well/vortex (without spilling).
- 12.7. Obtain background measurement reading. Wait a minimum of 1 minute and a maximum of 15 minutes between mixing and reading.
- 12.8. Add 20 µL of **DON** Tracer (YELLOW cap) into the first test tube.
- 12.9. Mix well.
- 12.10. Incubate for 60 seconds.
- 12.11. Obtain tracer measurement

Note: Run positive controls in duplicate. Repeat with a different dilution (7.4), if needed.

- 13. Testing procedure Sample:
- a. Use sample extract in Step 7.2
- b. Software: Choose *Analysis* in the user interface or *Analysis at given T°C* Sheet in particular excel version of software
- 13.1. Pipette 965 μl of **aokin**ReactionBuffer **FUM** ④ into the test tube.
- 13.2. Add 100 μ l of clear sample extract or a control into the test tube.
- 13.3. Add 15 µL of **FUM** Reagent (**VIOLET** cap) into the test tube.
- 13.4. Mix well/vortex (without spilling).
- 13.5. Obtain background measurement readings of all samples and controls. Wait a minimum of 1 minute and a maximum of 15 minutes between mixing and reading.
- 13.6. Add 20 μL of **FUM** Tracer (**YELLOW** cap) into the first test tube.
- 13.7. Mix well.
- 13.8. Incubate for 60 seconds.
- 13.9. Obtain tracer measurement of the first sample.
- 13.10. Repeat steps 13.6 to 13.9 for all test tubes.

Note: Run samples as duplicates. Repeat with a different dilution (8.2), if needed.

14. Test validation

- 14.1. The mean negative control must read between 160 and 230 mP.
- 14.2. The positive control must always read lower than the negative control. Expected values of the positive control are 10 mP to 50 mP lower than the negative control.
- 14.3. If the negative or positive control produce values outside of the expected values, the instrument should be recalibrated using the procedures described in the instrument manual.
- 14.4. Check the accuracy of your measurements by regularly analysing reference materials (e.g. **aokin**ReferenceMatrixMaterials), or other verified materials with known amounts of contaminants. Participation in proficiency tests is recommended as good laboratory practice.

Recommendation: If you notice erroneous measurement values, change the test tube and repeat the measurement after performing measurement of controls. Check that the liquid in the tube is not cloudy and that there are no bubbles after mixing.

Note: Repeat the daily checks after each hour of testing, or on each microtiter plate. Performing measurements of controls regularly will ensure the accuracy of your determinations. The results of the daily checks are used to ensure the analytical performance.



Be aware: The general composition of controls and samples should always be identical in salt concentration, pH and solvent ratio.

StandardSample preparation for **wheat, corn**

		T		
		Weighing:		
		15 g	sample	
		1.5 g	aokinExtractionSalt FUM	
Ē		35 mL	aokin EasyExtractBuffer FUM	
Extraction		Extraction:		
Ext		3.5 min	mixing with aokin watchbox	
		Filtration:		
			collect filtrate	
			(discard filter cake)	
Precipitation		Precipitation:		
		400 µl	•	
	\Diamond		ReactionBuffer FUM	
		20 min	centrifuge at $> 10.000 \times g$	
		100 ul	supernatant into 400 µl of aokin	
		200 μ.	ReactionBuffer FUM	
		20 min	centrifuge at $> 10.000 \times g$	
			transfer supernatant into clean 2	
			mL reaction tube for measurement	
	Quantification			
Measurement			add into tube:	
		965 µl	aokin ReactionBuffer FUM	
	Mix and read		sample	
		15 µl	FUM Reagent VIOLET	
			mix well	
		20 1		
		20 μΙ		
			measure tracer	
		20 μΙ	measure background add FUM Tracer YELLOW mix well measure tracer	