# File No: 11014/07/2021-QA Food Safety and Standards Authority of India

(A Statutory Authority established under the Food Safety and Standards Act, 2006) (Quality Assurance Division)

## FDA Bhawan, Kotla Road, New Delhi - 110002

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दिनांक: 16 अक्टूबर, 2024

# आदेश

Subject: Method for Residue Analysis of Ethylene oxide and 2-Chloroethanol in Food by Gas Chromatography Tandem Mass Spectrometry - reg.

The Method for Residue Analysis of Ethylene oxide and 2-Chloroethanol in Food by Gas Chromatography Tandem Mass Spectrometry has been approved by Food Authority in its  $44^{\rm th}$  meeting held on 19.06.2024 is enclosed herewith.

- 2. The approved method shall be implemented with immediate effect. The notified laboratories shall include the new method in their respective scope of accreditation within six months from the date of issue of this order.
- 3. Since the process of updation of test methods is dynamic, any changes happening from time to time will be notified separately. Queries/concerns, if any, may be forwarded to email: sp-sampling@fssai.gov.in.

Enclosure: As above

Dr. SATYEN

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RUMAR PANDA

Date: 2024.10.16
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(Dr. Satyen Kumar Panda) Advisor (QA)

To:

- 1. All FSSAI Notified Laboratories
- 2. All State Food Testing Laboratories
- 3. CEO, National Accreditation Board for Testing and Calibration Laboratories (NABL)

एफएसएसएआई	Method for Residue Analysis of Ethylene oxide and 2-Chloroethanol in	
Issai	Food by Gas Chromatography Tandem Mass Spectrometry	
भारतीय खाद्य सुरक्षाओर मानक प्रापिकरण Food Salely and Slandards Authorly of India स्वास्थ्य और परिवाद कल्याण मंत्रालय Ministry of Health and Family Welfare		
Method No.	FSSAI.OM.EO.001.2024 Revision No. & Date 0.0	
Scope	This is a selective and sensitive method for the residue analysis of ethy oxide (EO) in diverse matrices, namely oilseeds, cereals, tea, spices, he dehydrated fruits, food additives, and fruits & vegetables. When a	
	item is treated with EO (as a fumigant) to control spoilage and public health related microorganisms, it rapidly reacts with the matrix components, especially chloride, to form 2-chloroethanol (2-CE).	
	Here, a QuEChERS (EN 15662) technique is documented for the analysis of EO and 2-CE using (PTV or MMI)-GC-MS/MS or Headspace (HS)-GC-MS/MS or multi-step enrichment HS-trap GC-MS or GC-MS/MS. The method is validated in a wide variety of dry and fresh (high-moisture) food matrices, the measurement accuracies and precisions are noted in accordance with the analytical quality control criteria.	
	These methods have been validated in accordance with analytical quality control criteria and are documented here.	
Storage and Safety	Following storage and safety precautions shall be taken while handling:	
Precautions	• EO is highly volatile, and hence it was important to analyze the	
	samples at a temperature of < 10 °C.	
	• To prevent skin, eye, and inhalation contact, put on the proper protective eyewear and an apron with long sleeves.	
	Both EO and 2-CE are toxic and carcinogenic, thus require butyl rubber gloves for handling.	
Principle	EO is highly volatile, and hence it is important to prepare the samples at a temperature of < 10 °C. It is possible to estimate both compounds by liquid as well as headspace GC injections. It is also possible to analyze these compounds by automated headspace (HS)-trap GC-MS, in which, syringe-based HS is combined with cryogen-free trapping technology, exploiting the multi-step sample enrichment capability to increase the method sensitivity.	
	EO and 2-CE are analyzed within the same GC-MS/MS run. The recovery and precision, when checked through intra- and inter-laboratory validation studies, are highly satisfactory.	
Apparatus	A weighing balance with high precision is used to weigh the certified reference standards of EO and 2-CE. A heavy-duty grinder is used for crushing the samples. A vortex mixer (shaker), a highspeed refrigerated centrifuge, and a microcentrifuge are used at different stages of sample preparation.	

# Chemicals & Reference material

- (a) Chemicals EO (50 mg/mL, in methanol) and 2-CE (2000  $\mu$ g/mL, in methanol) having a purity >98%. HPLC grade water, anhydrous magnesium sulfate, sodium chloride, trisodium citrate dihydrate, and disodium citrate sesquihydrate. PSA: Bondesil, 40  $\mu$ m particle size) and octadecylsilane (C18, ODS). Other certified reference standard options may also be possible, e.g., EO (50 mg/mL in dichloromethane) and 2-CE (pure), diluted using a suitable solvent, e.g., LC grade hexane (>95% purity).
- (b) Materials— Polytetrafluoroethylene (PTFE) syringe filters (0.22  $\mu$ m) and Ultipor Nylon-6,6 membrane filters (0.2  $\mu$ m pore size and 13 mm diameter)

# Preparation of standards and reagents

## (a) Solutions—

- 1. Due to the high volatility of EO, its standard solutions were prepared at a low temperature < 10 °C) using a thermocol box containing ice bags.
- 2. As a diluting solvent, acetonitrile was placed in a freezer for at least 15 min before use.
- 3. The cold analytical standard solutions were pipetted into acetonitrile to generate the working standard solutions of EO (1 mg/mL) and 2-CE (1 mg/mL).
- 4. By serial dilution in acetonitrile, the calibration standards of 2.5, 5, 10, 25, and 50  $\mu$ g/L were prepared from this working standard.
- 5. The matrix-matched standards of the same concentrations were also separately prepared.
- 6. Prior to extraction, all stock solutions were preserved at a temperature of -20 °C to avoid degradation losses.

# Preparation of standards and reagents for multistep enrichment HStrap

#### (b) Solutions—

- 1. Due to the high volatility of EO, its standard solutions were prepared at a low temperature < 10 °C) using a thermocol box containing ice bags.
- 2. As a diluting solvent, hexane was placed in the freezer for at least 15 minutes before use.
- 3. All apparatus, including glassware, used for standard preparation were placed in the freezer for at least 15 minutes before use.
- 4. The cold analytical standard solutions were pipetted or weighed into hexane to generate the stock solutions, used for preparation of the working standard solutions, of EO (2 mg/mL) and 2-CE (2 mg/mL).
- 5. A combined stock solution was prepared containing EO and 2-CE each at a concentration of 1 mg/mL ( $1000 \text{ ng/}\mu\text{L}$ ).
- 6. By serial dilution in hexane, the working solutions of 10, 25, 50, 100, 200, 300, 400, 500 ng/µL were prepared from this stock solution.
- 7.  $1\mu$ L was introduced to 2 g of sample matrix, the resulting concentrations were 5, 10, 25, 50, 100, 150, 200, 250 ng/g ( $\mu$ g/kg).

- 8. Due to its structural similarity to 2-CE and its typical use as a surrogate standard during the analysis of volatile organic compounds, 1, 2-Dichloroethane- $d_4$  (pure) was used as the internal standard, diluted to 200 ng/ $\mu$ L in hexane.
- 9. Prior to extraction, all stock solutions were preserved at a temperature of -18 °C to avoid degradation losses. Additional measures, e.g., semi-transparent, flexible film (Parafilm®) to secure the vial lid, may be used to prevent potential evaporative loss during storage.

# Preparation of Test Samples

- (a) Sample type The samples of dry commodities [e.g., oil seeds (sesame seed), cereal (wheat), spices (cumin seed, turmeric powder, chilli powder, ginger powder), pulses (moong bean), dehydrated fruits/vegetables (kiwi, mango, onion flakes), medicinal herbal powder (e.g., ashwagandha, *Withania somnifera*), black tea powder], high-moisture foods (e.g., tomatoes, grapes), food additives (e.g., guar gum, locust bean gum), processed spices (e.g. coriander powder, curry powder mix (mutton, egg, and vegetable flavors) were evaluated for method performance. Before analysis, the samples were placed in separate sample collection bags, transferred into airtight containers, and maintained at -20 °C until further use.
- (b) Sample preparation With the modified QuEChERS (EN 15662) technique, the extraction of all dry and high-moisture matrices was performed at a temperature of ≤10 °C. The well-homogenized and precooled samples (4 g of powdered dry matrices and 10 g of high moisture matrices) were separately taken in 50 mL polypropylene centrifuge tubes. In the case of dry matrices, 5 mL of ice-cold water was added, and the sample was left standing for 15-20 min before vortexing for 2 min. To it, 10 mL of pre-cooled acetonitrile was added for extraction and vortexed for 15 min. With the exception of the addition of water, the same procedure was followed for the high moisture matrices, e.g., grape, tomato, etc. The mixture was vortexed for 2 min with MgSO<sub>4</sub> (4 g), NaCl (1 g), trisodium citrate dihydrate (1 g), and disodium citrate sesquihydrate (500 mg). Thereafter, the mixture was centrifuged for 5 min at 5000 rpm at  $\leq$ 10 °C. For the dry matrices, an aliquot of 1 mL of the cleaned supernatant was drawn and vortexed with 25 mg PSA + 25 mg C18 + 150 mg anhydrous MgSO<sub>4</sub>. But for certain other (relatively complex) dry matrices, such as coriander powder, curry powder mix (mutton, egg, and vegetable flavor), ashwagandha powder, guar gum, and locust bean gum, an aliquot of 1 mL of the cleaned supernatant was drawn and vortexed with 50 mg PSA + 50 mg C18 + 150 mg anhydrous MgSO<sub>4</sub>.

For high moisture matrices, e.g., grapes and tomatoes, 50 mg of PSA and 150 mg of anhydrous MgSO<sub>4</sub> were added and vortexed for 30 s, followed by centrifugation at 10000 rpm for 5 min (≤10 °C).

In the case of the sesame seed matrix,  $100~\mu L$  of the extract was directly analyzed by HS-GC-MS/MS. For chilli powder, turmeric powder, and guar gum, an aliquot of 1 mL of the supernatant was cleaned by d-SPE with 50 mg of PSA. After centrifugation at 10,000~rpm for 5 min, an aliquot of  $100~\mu L$  was taken in a 20~mL HS vial and carefully sealed before the final analysis.

The extracts were analyzed by GC-MS/MS after filtration. A PTFE syringe filter (0.22  $\mu$ m) was used for filtration of dry matrices, and a Nylon-6,6 membrane filter (0.2  $\mu$ m) was used for the filtration of wet matrices.

In the dynamic headspace (HS), an aliquot (100  $\mu$ L) of the sample extract was incubated at 110 °C for 15 min. In a subsequent enrichment process, the vapor from the 20 mL capacity HS vial was adsorbed on the trap, which was electrically cooled to -10 °C. Finally, the trap was desorbed at 280 °C to transfer the analytes into GC-MS/MS.

In multi-step enrichment HS-trap, the test samples (whole, uncontaminated sesame seeds and powdered guar gum) were weighed (2 g  $\pm$  0.05 g) directly into the vial (20 mL). The working solutions (1  $\mu L$ ) were introduced rapidly, including the internal standard (1  $\mu L$ ) at a fixed concentration (equating to 100  $\mu g/kg$ ). The vials were quickly capped and placed onto the auto sampler for immediate analysis. Vials were incubated at 70 °C for 10 min with agitation at 300 rpm. To concentrate the analytes, three headspace volumes (5 mL each) were collected from each sample and injected to a focusing trap, which is electrically cooled to -30 °C throughout the enrichment process. An incubation period of 3 min in between each extraction re-establishes the headspace equilibrium. Finally, the trap is desorbed at 250 °C to transfer the analytes to the GC–MS system for separation and detection.

# **Chromatography** conditions

### GC- MS/MS analysis by liquid injection—

- 1. A triple quadrupole GC-MS/MS with auto sampler.
- 2. The GC separation can be achieved on a Wax column, or fused silica cross-bonded capillary column (e.g. dimension: 30 m, 0.20-0.25 mm).
- 3. Ultrapure helium (99.9999%) was used as the carrier gas, with a flow rate of 1.2 mL/min.
- 4. The oven temperature program to be set as follows: an initial temperature of 45 °C (2 min hold) was ramped to 230 °C at 50 °C/min (5.3 min hold), which resulted in a total run time of 11 min.
- 5. The transfer line and ion source temperatures were maintained at 250 and 230 °C, respectively. The injector program started at 90 °C (0.8 min hold) and increased to 250 °C (10 min hold) by rapid heating at 12 °C/s.
- 6. Split injection was employed with a baffled PTV injector linear

- (dimension, e.g.:  $2\times2.75\times120$  mm). The injection volume was 2  $\mu$ L.
- 7. The MRM transitions may be optimized for EO and 2-CE by using the auto-MRM feature of the software.
- 8. The MRM method was automatically optimized in terms of the precursor ions, product ions, and collision energies by adjusting the dwell time for each transition to achieve the highest sensitivity (S/N).

GC-MS/MS analysis by headspace injection— A gas chromatograph with auto-injector and headspace sampler is used with a triple quadrupole mass spectrometer. The software is used for data analysis and quantitation. The hardware system allows the sample vial to enter the oven from the bottom, reducing heat loss during the process. Through an advanced flow control system, the accuracy of flow rate is maintained. The pressure in the vial is kept constant at 192 kPa.

# Dynamic headspace GC-MS/MS parameters for EO and 2-CE:

Column	Rtx-VMS (example name) GC column: $60 \text{ m} \times 0.45$
	mm, 2.55 µm or equivalent
Flow rate	Helium, 3.0 mL/min
Injection	Split, split ratio: 20:1
mode	
Headspace	Incubation temperature: 110 °C, trap cooling temp: -
program	10 °C, Trap desorb temp: 280 °C, pressurizing gas
	pressure: 192 kPa, Equilibrium time: 15 min
Oven	35 °C (5 min hold); ramped at 20 °C/min to 235 °C (5
temperature	min hold)
program	

# **Mass Spectrometric parameters:**

MS	Ionization mode: Electron Ionization			
parameters	Transfer line temperature: 230 °C			
	Ion source temperature: 230 °C			
MRM	Retention	Precursor	Product ion	CE
transitions	time	ion (m/z)	(m/z)	
	(min)			
		44	14	20
EO		44	28	5
	After the	44	29	5
	void	80	31	5
2-CE	volume	80	43	5
		80	44	5

# Multi-step enrichment headspace-trap example GC parameters for EO and 2-CE:

Column	VF-624 (example name) GC column: 60 m $\times$ 0.25 mm, 1.4 $\mu$ m or equivalent
Flow rate	Helium, 2.0 mL/min
Headspace program	Incubation temperature: 70 °C (10 min, 300 rpm). Extraction volume: 5 mL. Number of extractions (enrichment): Three- 5 mL, 3 min enrichment delay between each. Trap cooling temp: -30 °C. Trap desorb temp: 250 °C. Split flow (trap desorb): 10 mL/min (6:1 split ratio)
Oven temperature program	35 °C (5 min), to 100 °C at 10 °C/min (1 min) reaching 230 °C at 20 °C/min (5 min)

Mass Spectrometric parameters – *see example parameters for Dynamic headspace GC-MS/MS* 

When using multi-step enrichment HS-trap, due to the concentration factor achieved, it may be possible to use a single quadrupole mass spectrometer in Single-Ion-Monitoring (SIM) mode. Example MS parameters for EO and 2-CE are as follows:

MS parameters	Ionization mode: Electron Ionization		
	Transfer line temperature: 230 °C		
	Ion source temperature: 250 °C		
SIM ions	Target ion (m/z)	Qualifier ion (m/z)	
EO	29	44	
2-CE	31	44	

#### **Results**

In this study, a simple and rapid temperature-controlled extraction method was established for the analysis of EO in a wide variety of dry and high-moisture food matrices. The optimized method provided satisfactory homogeneity, sensitivity, accuracy, and precision for both the target compounds in compliance with the method performance criteria of the SANTE/11312/2021v2 guideline. The LOQ of both compounds was  $\leq 0.01$  mg/kg.

Furthermore, a satisfactory performance in the intra- and inter laboratory validation studies indicates its ruggedness and reproducibility. Owing to its satisfactory performance, this method is recommended for the determination of EO and its reaction product 2-CE in governmental and commercial food testing laboratories. The method is expected to facilitate EU–India trade in a variety of fresh and processed commodities due to its high reproducibility.

Calculation	The detected residues were quantified through matrix-matched calibration, and EO (sum) was calculated using the following formula:		
	EO (sum) [mg/kg] = EO + 2-CE $\times$ 0.55 (the conversion factor)		
	In the case of multi-step enrichment HS-trap-GC-MS, EO and 2-CE were		
	individually quantified using matrix-matched calibrations for both target analytes.		
LOQ	The LOQ was set as the lowest concentration at which the results met the		
	method performance evaluation criteria and were estimated using matrix-		
	matched standards. At LOQ, the S/N of the quantifier MRM was >10:1.		
	The recoveries at LOQ were above 70%.		
	In the case of multi-step enrichment HS-trap-GC-MS, recoveries for EO		
D -f	and 2-CE at 3 levels (5, 50 and 250 µg/kg) were between 80-115%.		
Reference	1. Analytical quality control and method validation procedures for pesticide residues analysis in food and feed. SANTE/11312/2021 v2.		
	2. EN 15662:2018. Foods of plant origin. Multimethod for the determination of pesticide residues using GC- and LC-based analysis following acetonitrile extraction/partitioning and cleanup by dispersive SPE. Modular QuEChERS-method.		
	3. Nerpagar A., Banerjee Kaushik (2023). Dynamic headspace GC-MS/MS analysis of ethylene oxide and 2-chloroethanol in dry food commodities: a novel approach <i>Journal of Environmental Science and Health</i> , <i>Part B</i> https://doi.org/10.1080/03601234.2023.2264740.		
	4. Patil R., Langade N., Banerjee Kaushik (2023). Development and validation of a residue analysis method for ethylene oxide and 2-chloroethanol in foods by gas chromatography tandem mass spectrometry. <i>ACS Agricultural Science and Technology</i> https://doi.org/10.1021/acsagscitech.2c00319.		
	5. Hearn L, Szafnauer R., Banerjee Kaushik, Amin P. (2024). Automated, cryogen-free headspace-trap with gas chromatography-mass spectrometry analysis of ethylene oxide and 2-chloroethanol as residual fumigants in foods. <i>Journal of Environmental Science and Health, Part B</i> https://doi.org/10.1080/03601234.2023.2298169.		
Approved by	Scientific Panel on Methods of Sampling and Analysis		

**Note:** The brand / model of equipments/ accessories/ column / Chemicals & reagents given in the methods are for reference purpose only. The end user may use equivalent specifications of equipments/ accessories/ column / Chemicals & reagents.

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