



Determination of Total Chlorine in Palm Oil with the Elemental Analyzer multi EA 5100

Introduction

Palm oil and palm kernel oil production has grown drastically in the last 50 years. With more than 30%, it is the largest contributor to global oil and fat production. Traditionally the food and cosmetics industries are amongst the main consumers with a share of 70% of the world palm oil production, followed by usage for biofuel production. Palm oil is used in products like margarine, oil, spreads, chocolate, cleaning agents, cosmetics, candles and many more. To improve taste characteristics and shelf life palm oil needs to be refined. But refined palm oil can pose a health risk to consumers.

Various studies carried out during the last years have proven the presence of 3-monochloropropane-1,2-diol (3-MCPD) fatty acid ester in refined palm oil. 3-MCPD is formed when fats and oils are exposed to high temperatures, in the presence of organic and inorganic bound chlorine, which often happens during the refining process. 3-MCPD is suspected to raise the risk of cancer. Animal experiments have shown that a higher uptake of 3-MCPD triggers kidney and liver damage and benign tumors. That's why, the World Health Organization has set a threshold level for the tolerable daily intake of 2 micrograms 3-MCPD per kilogram of body weight.

Used as a food ingredient, palm oil is subject to strict quality and food safety controls. Besides the determination of antioxidants, water content or trace metals (Pb, As, Cd and Hg) also the determination of total chlorine plays an important role. Perfectly suited for this purpose is the organic elemental analysis, coupling oxidative combustion and micro coulometric detection.

Challenge

Fully automated, fast and accurate determination of total chlorine content in different types of palm oil

Solution

Organic Elemental Analysis - Matrix- and time optimized combustion with flame sensor technology using the multi EA 5100

Materials and Methods

Samples and Reagents

Different aliphatic and aromatic hydrocarbons and their mixes (e.g., toluene, isooctane, etc.) have been analyzed.

- Isooctane (C₈H₁₈), Suprasolv®, GR for gas chromatography (Merck Art.-No.: 1.15440.1000)
- Dibenzothiophene (C₁₂H₈S), GR for synthesis (Merck Art.-No.: 8.20409.0025)
- Kit calibration solutions 0.1 - 10 mg/L Chlorine (Analytik Jena, Art.-No.: 402-889.071)

Sample Preparation

Two different palm oil samples were analyzed. Red palm oil, which is a viscous red-orange liquid used as high quality cold-pressed cooking oil. And 100% natural palm oil from a cosmetic production, a white-ivory paste-like solid substance. Depending on the quality of raw material, production method and level of treatment, the homogeneity of palm oil samples differs, which has an effect on the reproducibility of results.

Due to the fact that the proportion of the chlorine-containing compounds in such matrices is also inhomogeneous, a sufficient pre-treatment strategy is crucial to ensure reliable measurement results.

That's why two different strategies were applied to prepare homogeneous sample aliquots. First, the samples were diluted 1:3 (w/w) with *o*-xylene. Second, they were carefully melted while gently shaking in a water bath, shortly before sampling. For this purpose, closed vials were used to avoid evaporation losses of the Cl-containing components.

Instrumentation Settings

A multi EA 5100 in horizontal operation mode, including the flame sensor technology, was used for the analysis.

For sample feeding the automatic boat drive in combination with the multi matrix sampler – MMS was used. In solids mode the undiluted samples were introduced directly by means of sample boats. In liquids mode 100 µL of the diluted samples resp. the liquid standards were injected by means of a µL-syringe.

The sample digestion is carried out by efficient catalyst-free high temperature combustion in a quartz reactor. This process is controlled and adopted to the special needs of every matrix component fully automatically thanks to the flame sensor technology. This ensures matrix-independent, optimal results in the shortest possible time. The process is split into two phases. In the first phase evaporation of light components and pyrolysis of the heavier ones takes place within an inert argon atmosphere. The resulting gaseous products are converted in the pure oxygen atmosphere of the combustion zone. In the second phase the system switches completely to oxygen and the remaining components are combusted quantitatively. The implemented Auto-Protection System guarantees highest operational safety (particle and aerosol trap) and a complete transfer (no condensation loss) of the formed HCl into the "high sensitive" cell. Afterwards the determination of the chlorine content is carried out by means of a micro-coulometric titration. The multi EA 5100 enables a detection limit of 50 µg/L Cl. Depending on the cell type used, Cl contents up to 10 wt% can be analyzed directly.

Method Parameters

Standard method settings for the horizontal operation mode are applied. The combustion process parameters are shown in Table 1 and the TCl detection parameters in Table 2.

Table 1: Process parameters

Parameter	Specification
Furnace temperature	1050°C
2nd combustion	60 s
Ar flow (1st phase)	200 mL/min
O ₂ main flow	200 mL/min
O ₂ flow (2nd phase)	200 mL/min
Purge*	100 s
Draw up**	1 µL/s
Injection**	3 µL/s

Table 2: Detection parameters

Parameter	Specification
Max. integration time	1200 s
Threshold value	300 cts
Max. drift	100 cts/s
Threshold	25 cts
Cell temperature	23 °C
Titration delay	30 s

* for the solids method only, ** for the liquids method only

Calibration

Liquid calibration standards based on 2,4,6-trichlorophenol in isooctane were used to calibrate the analysis system in the concentration range from 0.1 to 5 mg/L TCI. The calibration curve is shown in Figure 1. The calibration was checked with a certified reference standard.

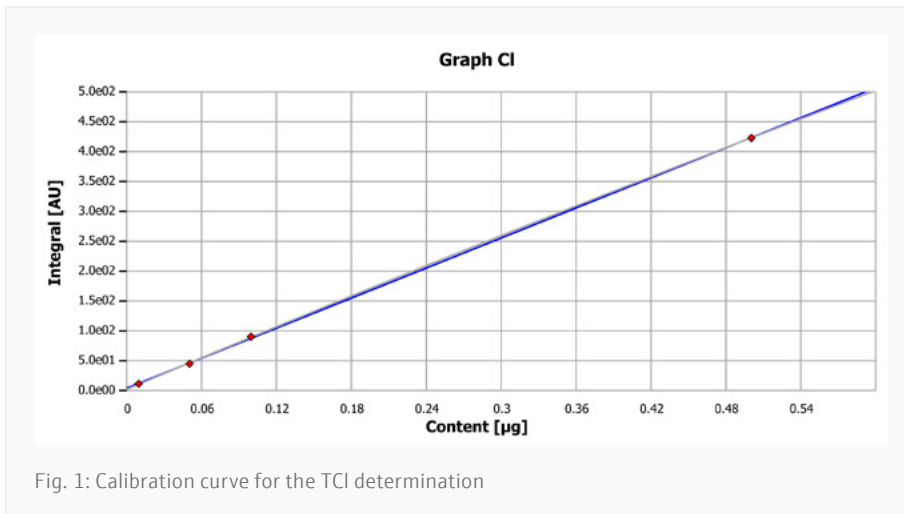


Fig. 1: Calibration curve for the TCI determination

Results and Discussion

The two different palm oil samples were analyzed two times – first directly, using the solids mode, and second in liquids mode, diluted with o-xylene. The results and the measurements of two selected CI standards are summarized in Tables 3 (undiluted) and 4 (diluted). The results are averages of three replicate analyses. Typical measuring curves are shown in Figures 2 (undiluted) and 3 (diluted).

Table 3: Results of TCI determination in undiluted palm oil using the solid method

Measurement	TCI	RSD	Sample quantity
red palm oil	1.27 mg/kg	± 5.95 %	~ 55 mg
white palm oil	3.03 mg/kg	± 25.7 %	~ 25 mg
TCI standard 1.45 mg/kg	1.49 mg/kg	± 0.29%	100 µL

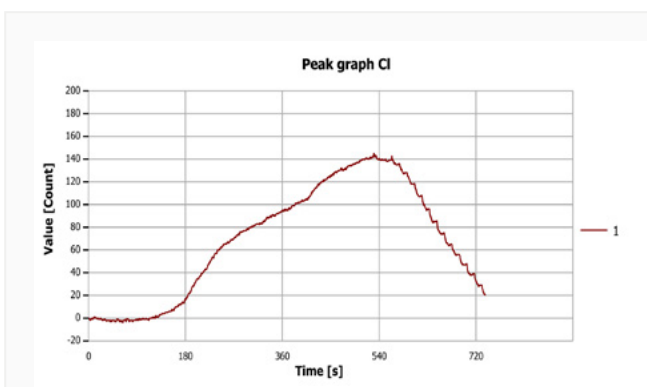


Fig. 2a: TCI measuring curve for the undiluted red palm oil

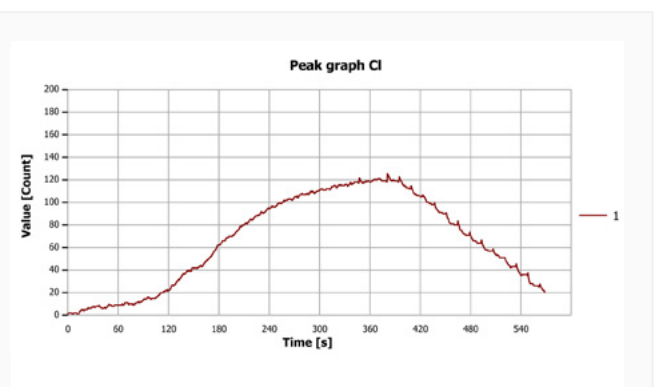


Fig. 2b: TCI measuring curve for the undiluted white palm oil

Table 4: Results of TCI determination in diluted palm oil samples

Measurement	TCI	RSD	Dilution [g in g]
red palm oil	1.30 mg/kg	± 1.99 %	3.5559 in 7.3622
white palm oil	3.16 mg/kg	± 2.91 %	3.4382 in 8.1002
TCI standard 0.72 mg/kg	0.71 mg/kg	± 0.54 %	-

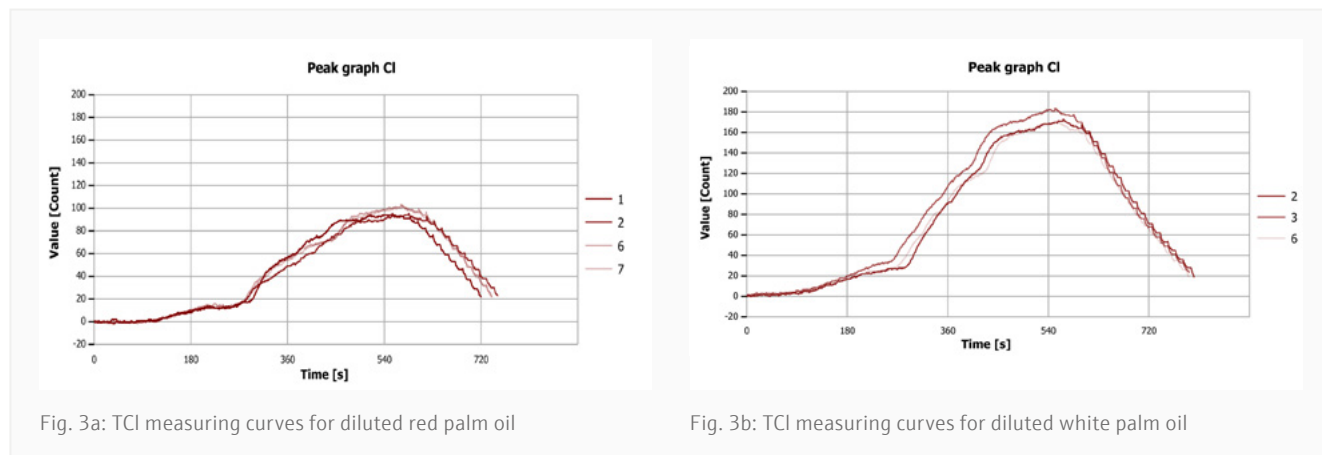


Fig. 3a: TCI measuring curves for diluted red palm oil

Fig. 3b: TCI measuring curves for diluted white palm oil

The different pre-treatment strategies deliver comparable results with deviations of less than 5%, thus demonstrating the general suitability of both principles for analysis of diverse palm oil samples. Nonetheless using dilution leads to smaller deviations of the single analyses. This as well as the easier sample handling, and faster processing, make the dilution strategy the superior one.

Conclusion

This work has successfully demonstrated that the multi EA 5100 with ABD and flame sensor technology provides a fast, safe and reliable solution for the analysis of Cl traces in palm oil and related matrices. Due to the time- and matrix-optimized digestion process by means of the flame sensor, the quantitative combustion of any sample component is ensured. No formation of soot or other undesired pyrolysis products is observed. This and the efficient Auto-Protection system enable superior reproducibility even for the smallest chlorine quantities. A high sample throughput is easily achieved using the multi-matrix sampler MMS that is able to introduce solid as well as liquid sample matrices fully automatically either by quartz boats or by direct syringe injection.

Thanks to the modular design of the multi EA 5100 the user has any flexibility for later upgrades to determine other parameters like Nitrogen, Sulfur or Carbon in different matrices like LPG and gases.

References

- [1] EFSA CONTAM Panel (EFSA Panel on Contaminants in the Food Chain); SCIENTIFIC OPINION ON THE RISKS FOR HUMAN HEALTH RELATED TO THE PRESENCE OF 3- AND 2-MONOCHLOROPROPANE-DIOL (MCPD), AND THEIR FATTY ACID ESTERS, AND GLYCIDYL FATTY ACID ESTERS IN FOOD. EFSA Journal. 2016, 14/5, 4426, 159