Volumetric one-component Karl Fischer Titration for aldehydes and ketones

1 to 100 mg of water in sample used

This document applies to volumetric Karl Fischer titrations determined with one-component based reagents. The used reagents are applicable for samples that contain aldehydes and ketones. The white cells in the table below show the applicable applications.

| | | One-component | | Two-component | |
|--------------------------------------|------------------------|------------------------------|----------------------------|----------------------------|----------------------------|
| Methanol- <u>based</u> titrations | Titrant in the syringe | Composite 5 | Composite 2 | Titrant 5 | Titrant 2 |
| | Solvent in the beaker | Methanol rapid | Methanol rapid | Solvent or Solvent CM | Solvent or Solvent CM |
| | Available application | 1-component methanol C5 | 1-component methanol C2 | 2-component methanol T5 | 2-component methanol T2 |
| | Application note | DOC316.52.93124 | | DOC316.52.93098 | |
| Methanol- <u>free</u> titrations | Titrant in the syringe | Composite 5K | | Titrant 5E | Titrant 2E |
| | Solvent in the beaker | Medium K or Working Medium K | | Solvent E | Solvent E |
| | Available application | 1-component C5K* | | 2-component ethanol T5E | 2-component ethanol T2E |
| | Application note | DOC316.52.93125 | | DOC316. | 52.93129 |

*used specially for samples containing aldehydes and ketones

Refer to the related application note to use other reagents.

Make sure not to mix titrant and solvent reagents of different systems.

1. Introduction

Water analysis is, without doubt, one of the most frequently performed laboratory analyzes in various industries such as the oil, pharmaceutical and food industries. For many products, water content needs to be determined at every stage of the manufacturing process from raw materials to finished goods. The quality and/or the preservation of the final product depend on the amount of water present.

Excess levels of moisture will, for example, allow bacterial growth in food, decrease the performance of oils and lubricants, modify the density and the viscosity of paints and inks, disturb the texture of powdered products by forming conglomerates and impede the combustion properties of fuels.

Titrations and settings described in this document have been chosen to cover sample types commonly examined for their water content. They should be used as a starting point for specific samples and, if needed, optimized to suit analysis requirements.

2. Principle

The Karl Fischer technique is the most commonly used for moisture determination because of its rapidity, accuracy and ease of use. The sample is generally injected into the titration cell where an appropriate solvent dissolves it. The cell content is then titrated to complete dryness with a solution containing iodine. Volumetric Karl Fischer determination covers the range from 1 to 100 mg of water in the sample taken.

The titrant

The determination of water content is based on the oxidation of sulfur dioxide by iodine in the presence of water:

$$I_2 + SO_2 + 2H_2O \rightarrow 2HI + H_2SO_4.$$

This chemical reaction is used in the Karl Fischer titration for quantitative water determination. The key components of the titrant are iodine and sulfur dioxide.

A base (noted RN, for example imidazole) and an alcohol (noted R-OH, for example methanol) are required for the reaction:

 $R-OH + SO_2 + RN \rightarrow (RNH) \bullet SO_3R$

 $(RNH) \bullet SO_3R + I_2 + H_2O \rightarrow (RNH) \bullet SO_4R + 2(RNH)I$

Recent studies have shown that oxidant molecules are not I_2 or $RN \bullet I_2$ but that I_2 partially splits up between I^{\dagger} and RNI^{\dagger} into polar solvents and that it is the RNI^{\dagger} which reacts.

The solvent

The solvent or working medium must ensure the stoichiometry of the Karl Fischer reaction. It must be able to dissolve the sample and the products of the reaction. It must also enable the detection of the end point.

The indicator electrode

A double platinum electrode is used, polarized with an alternating square wave current. The detection of the end point of the titration is based on the detection of a small excess of iodine. This occurs when water is no longer present in the KF cell. That may be indicated visually by a slight yellow coloring of the working media.

| 3. Electrode and reage | ents | | |
|------------------------|--|--|--|
| Electrode: | Combined Pt-Pt with Temperature sensor: Intellical MTC695 | | |
| Titrant: | Composite 5K reagent for volumetric one-component Karl Fischer Titrations | | |
| Working medium: | Medium K or Working Medium K reagent for volumetric one-component Karl Fischer Titrations Medium K is very recommended: toxic 2-chloroethanol is replaced with less hazardous alcohols When Working Medium K is used, solid potassium iodide may be necessary (refer to 4.3 Analysis steps) | | |
| Standard: | Standard 5.00 | | |

Ethanol

Note: For optimal results, the use of Hydranal[®] reagents is recommended.

Safety considerations: Reagents used for Karl Fischer titrations have to be handled carefully due to the chemical exposure hazard. Please read the MSDS document related to each chemical and adapt protective equipment accordingly (e.g. glasses, gloves, fume hood, white jacket, etc.)

4. Karl Fischer titration: Key points

4.1. Guideline for titrations

Step 1: Determination of the amount of sample. The quantity of sample taken is based on the expected amount of water. The manufacturer recommends the use of less than 20 g of sample. Also, the injected sample must add at least 1 mg of water into the cell.

Step 2: Solvent determination. Specific working medium is necessary for samples that contain aldehydes and ketones. These classes of compounds form acetals and ketals in conventional reagents at the same time of the release of water, which can give erroneous high water content results. With aldehydes, a second side reaction (bisulfite addition) can also occur. This reaction consumes water, so the determined water content can be too low. Replace the methanol used in conventional reagents by other alcohols to prevent the formation of acetals and ketals.

Step 3: Frequency of solvent renewal. This parameter is based on the nature of the sample. Experimentally determine the number of possible titrations with the same solvent. Successive titrations with the same solvent are possible until the drift of the measurements caused by the pollution of the cell.

4.2. Sampling, storage and sample introduction

Sampling depends on the nature of the sample but for moisture determination, handling has to exclude moisture absorption or desorption. Atmospheric moisture can affect the accuracy of the result.

Moisture determination of the sample has to be done as soon as possible after sampling. In the case of storage, keep the sample in a tightly sealed bottle.

The optimal accuracy on the measurement is obtained using the back weighing technique. The sample and the vessel (e.g. plastic or glass syringe for liquid samples or weighing boat for solid samples) are weighed before the introduction of the sample into the cell and the empty vessel is weighed again after sample introduction. The weight obtained by the difference of the two will be used for further calculations.

For liquid samples: Adapted sampling steps to prevent atmospheric moisture absorption have to be considered, especially for low water content sample.

For solid samples: They can be directly introduced into the cell quickly after being weighed, to limit atmospheric exposure.

The table that follows gives sample weights based on the expected water content and the selected titrants:

| Expected water content (%) | Sample weight (g) with titrant concentration:5 mg/mL | |
|----------------------------|---|--|
| 100 | < 0.02 | |
| 10 | 0.05 – 0.20 | |
| 1 | 0.5 – 2.0 | |
| 0.1 | 5 – 20 | |
| 0.01 | - | |

4.3. Analysis steps

Refer to the user manual for additional information.

1. Fill the cell with the appropriate solvent in such a way that both the bottom of the electrode and the anti-diffusion tip are properly immersed in the solvent. An indication mark on the cell helps you to add the minimum volume required.

The relative positions of the electrode and the anti-diffusion tip in the cell are crucial to achieve optimum results. The height between the bottom of the electrode and the bottom of the cell is defined by the supplied Karl Fischer white Teflon® conical adapter. The correct positioning of the anti-diffusion tip is defined by the conical adapter pre-assembled on the tube. It is highly recommended not to modify these settings. Refer to the user manual for additional information about the installation.

- 2. Launch the application. Cell conditioning begins and any moisture present in the cell is removed. **Note:** When Working Medium K is used, PtPt electrode saturation can occur after a solvent renewal, which can block the cell conditioning. To prevent this, add a few pinches of solid potassium iodide KI in the cell until the cell conditioning starts.
- 3. Prepare the sample and weigh both the sample and the sample container.
- 4. When required, introduce the sample into the cell and start the titration.

Tips:

- Avoid spills onto the cell walls.
- When using a syringe equipped with a needle for introducing the sample, put the needle directly into the working medium to inject the sample. This makes sure that the entire sample is injected into the working medium and none is lost on the cell walls or electrode. Do not rinse the syringe by pumping back some solvent.
- 5. Weigh the empty sampling container and, by difference, determine the exact weight of the sample injected into the KF cell.
- 6. At the end of the titration and when prompted, enter the exact weight of the sample calculated in Step 5.

4.4. Recommendations for settings modifications

If needed, adapt the following parameters to optimize the titration to make it suit the analysis requirements.

- Name of the sample: With "?" the sample will be automatically incremented with a number for each analysis
- **Stirring:** An efficient stirring of the titration cell is crucial to have a homogeneous solution and avoid over-titration. Increase the speed until a vortex occurs without creating any bubbles. Make sure that both the titrant addition anti-diffusion tip and the PtPt electrode wires are kept immersed.
- **Delay before titration:** The time of stirring before the beginning of the titration helps to extract the water from the sample and has to be adjusted for each sample to increase the repeatability and accuracy of the measurement.
- **Drift threshold:** This value is set at 50 µg/min by default, but can be adjusted with regards to the level of humidity surrounding the titrator.
- Minimum and Maximum titration times: The minimum titration time setting avoids under estimation of the water content. The maximum titration time setting avoids taking into account any side reactions after the true determination of water content.

5. Hardware configuration

The KF1121 titrator is dedicated to the determination of moisture in different samples. It integrates automatic moisture drift compensation to avoid moisture drift errors that might cause a deviation from the true measurement. This instrument is equipped with a specific water tight cell, an MTC695 electrode installed onto the probe holder with a specific electrode adaptor, an addition tip equipped with an anti-diffusion tip, a tube to remove the cell content and a removable watertight cap.

To reinforce the repeatability and the accuracy of the measurements, the position of each element (electrode and addition tip) has been carefully defined and no modifications are required.

Depending on the environmental relative humidity conditions, it is recommended to replace the molecular sieve in the desiccant tubes with a fresh/dry one as frequently as required. This will prevent any moisture contamination of the measuring setup (titrant bottle, solvent bottle and KF cell). It is advised to use silica gel crystals which act as a visual indicator.

Titrations and calibrations are performed using a 5 mL syringe.

6. Titrant calibration with Standard 5.00

To increase the accuracy of the measurement of moisture determination of the sample, it is recommended to perform a calibration of the titrant every day. When titrations are performed in very wet conditions, a calibration can be performed twice a day. Always perform a calibration for a new titrant installation.

6.1. Procedure

Standard 5.0 is used as a standard for volumetric Karl Fischer titration: nominal concentration of \approx 5.00 mg/ml water corresponds to \approx 5.8 mg/g at 20 °C (refer to analysis report from supplier for accurate value).

Standard 5.0 is added using a 2.5 mL glass syringe.

The amount of standard should be estimated so that the total volume of the syringe is added.

6.2. Settings

The detection of the end point of the titration is based on the detection of an excess of iodine based on the following settings: End Point at 200 mV and current at 80 μ A.

At the end of the titrant calibration the concentration of the titrant is calculated and expressed in mg/mL. The saved value (real concentration) will be used for moisture determination.

6.3. Calibration of Composite 5K

6.3.1. Standard and procedure

Standard: When a 5 mg H₂O/mL titrant is used in a 5 mL-syringe, add exactly 2.5 mL Standard 5.00.

6.3.2. Settings

| Name | Default parameters | Units |
|---------------------------------------|--------------------|----------|
| Application | | |
| Application name | 1-component C5K | |
| Advisable syringe | 5 mL (Hamilton) | |
| Electrode | | |
| Туре | mV | |
| Recommended electrode | MTC695 | |
| Current (titrant calibration)* | 80 | [µA] |
| Titrant: KF C5K 5 mg/mL | | |
| Name | KF C5K | |
| Real concentration | 5.000 | [mg/L] |
| KF C5 5 mg/mL method : Titrant calibr | ation | |
| Active | Yes | |
| Calibration frequency | 1 | [Day(s)] |
| Stirring speed | 25 | [%] |
| Injection autodetect | No | |
| Proportionnal band* | 600 | mV |
| Drift threshold | 50 | [µg/min] |
| Min. titration duration | 30 | [s] |
| Max. titration duration | 300 | [s] |
| Min. titrant conc. | 4.5 | [mg/L] |
| Max. titrant conc. | 5.5 | [mg/L] |
| Standard name | Standard 5.00 | |
| Standard amount | 2 | [g] |
| Min. amount | 0 | [g] |
| Max. amount | 5 | [g] |
| Concentration** | 0.58 | [%] |

* Do not change this parameter in Expert mode. The parameter was specially adapted for this application.

** Edit and modify the value of the standard concentration based on the value in the analysis report from the supplier.

7. Moisture determination with 1-component C5K

7.1. Reagent

| Titrant: | Composite 5K |
|-----------------|--------------|
| Working medium: | Medium K |

7.2. Procedure

The sample is injected using a plastic syringe with a needle. The actual weight is determined from the difference between the full and empty vessel.

7.3. Settings

| Name | Default parameters | Units |
|-------------------|--------------------------|-------|
| Application | | |
| Application name | 1-component methanol C5K | |
| Advisable syringe | 5 mL (Hamilton) | |
| Sample | | |
| Name | Sample? [*] | |
| Amount | 2.0000 | [g] |
| QC | | |

| Name | QC Sample | | |
|-----------------------------|---------------|----------|--|
| Electrode | | | |
| Туре | mV | | |
| Recommended electrode | MTC695 | | |
| Current (sample analysis)** | 80 | [µA] | |
| Titrant: KF C5K 5 mg/mL | | | |
| Name | KF C5K | | |
| Real concentration | 5.000 | [mg/L] | |
| Method : KF titration | | | |
| Stirring speed | 25 | [%] | |
| Delay | 30 | [s] | |
| Injection autodetect | No | | |
| Proportionnal band** | 600 | mV | |
| Drift threshold | 50 | [µg/min] | |
| Min. titration duration | 30 | [s] | |
| Max. titration duration | 300 | [s] | |
| Result 1 (R1) name | Water content | | |
| R1 hide | No | | |
| R1 min. | 0 | [%] | |
| R1 max. | 100 | [%] | |
| R1 QC min. | 0 | [%] | |
| R1 QC max. | 100 | [%] | |

* "?" in the name, indicates that the instrument automatically increments the sample name with a number for each analysis.
** Do not change this parameter in Expert mode. The parameter is specially adapted for this application.

7.4. Example of water determination in acetylacetone

This is an example of moisture determination in acetylacetone with the default parameters.

Sampling: 2 g of acetylacetone. Eight tests have been done.

| | | Units |
|----------------------|-------|-------|
| Number of replicates | 8 | |
| Average | 0.494 | [%] |
| Standard deviation | 0.003 | [%] |
| RSD | 0.5 | [%] |