Excellence in Pharmaceutical TOC Water Testing and Vaccine Quality Control multi N/C pharma Series





multi N/C pharma Series

The multi N/C pharma series offers tailored solutions for TOC cleaning validation, extractables testing from packaging materials, ultrapure water control and total protein analysis in pharmaceutical aqueous solutions.

Enhanced software features make sure for full data integrity and 21 CFR part 11 compliance. A complete service package including system qualification and software validation provides you with maximum comfort in operating your analyzer.

Your benefits:

 Compliance with international pharmacopoeia methods quaranteed

Exceptional long-term stable and cost-effective operation

High sensitivity

Individual systems tailored to your needs



multi N/C series - Features

- Focus Radiation NDIR Detector
 Highest radiation density for highest sensitivity and precision
- VITA Flow Management System
 Continues to work where classical TOC analyzers reach their limits
- Easy Cal Calibration has never been so easy
- High Power Long Life UV Reactor
 Convincing performance in wet chemical oxidation

multi N/C pharma Series

Excellence in pharmaceutical TOC water testing and vaccine quality control



multi N/C pharma UV

Highly sensitive TOC ultra trace analysis using wet chemical UV digestion

multi N/C pharma HT

Sensitive TOC ultra trace and TN analysis using catalytic high-temperature combustion

multi N/C 3100 pharma

High-throughput TOC trace and TN analysis in a wide concentration range, using catalytic high-temperature combustion

multi N/C 2100S pharma

Wide range total protein analysis by catalytic high-temperature combustion and CLD detection

multi N/C pharma – A Perfect Match for your Application

Tailored solutions for pharmaceutical industries

TOC ultrapure water (WFI) testing according to USP <643> (TOC monograph)

The TOC parameter is used for quality control in ultrapure water analyses, in particular for the analysis of water for injection purposes (WFI) and aqua purificata (purified water – AP). USP <643> represents the general method for TOC testing in pharmaceutical applications and provides guidance on how to qualify the analytical technique for use as well as guidance on how to interpret instrument results for use as a limit test, e.g. the 500 ppb limit for WFI – water for injection. The analysis can be performed using high-temperature combustion or wet-chemical/UV oxidation.

TOC cleaning validation

In cleaning validation TOC analysis plays an outstanding role since it provides a non-substance specific-method, which allows the measurement and quantitative determination of total organic residues of active ingredients, additives, reaction products and detergents in the trace range. Therefore it is used as a worst case screening parameter for cleaning validation testing of production equipment according to the established TOC monographs, e.g. USP <643>, PharmEur. 2.2.44 or JP 17, section 2.59. The instruments of the multi N/C series allow TOC cleaning validation analysis in the post-final rinse as well as swab test following elution or by direct high-temperature incineration. The particular advantage of aqueous rinse samples or swab eluates is that ultrapure water testing and cleaning validation of samples can be processed in a single TOC device using the same measuring method. In addition, the high-temperature combustion devices allow simultaneous total nitrogen (TN) cleaning validation when coupled to a chemiluminescence detector for NO detection.



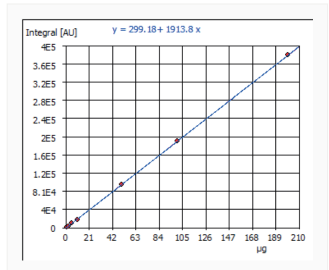






TOC testing for extractables from plastic packaging materials according to USP <661.1> and <661.2>

The characterization of the materials for pharmaceutical packaging systems is important to further improving the product safety of pharmaceuticals. TOC testing of water-based extracts is the method of choice for detecting organic leachables and is specified in the USP chapter <661>, addressing "Materials of Construction" USP <661.1> and "Plastic Packaging Systems for Pharmaceutical Use" USP <661.2>. For the analysis, purified water extractions from polymer packaging materials have to be tested for TOC according to USP <643>.



With their high oxidation power and advanced detection the instruments of the mLti N/C pharma series exceed the required specifications of the USP <661.1> and <661.2>:

- Linear dynamic range of 0.1–20 mg/L (required 0.2–20 mg/L)
- Detection limit <0.005 mg/L (USP max. 0.05 mg/L)

Total protein analysis in vaccines by total nitrogen measurement according USP <1057>, Method 7.2

The assay determination of protein is immanent for intermediate and finished products in pharmaceutical vaccine production. It is used to control the level of antigens by quantifying attenuated or devitalized viruses or bacteria via total protein contents. In allergenic vaccine production it is even possible to use one and the same total nitrogen test method (TN) for testing of raw materials like pollen lyophilisates as well as very different dosage forms of drug product like injectables for desensibilisation, prick test solutions or subligual applications. The application range of this TN method in the biopharmaceutical industry is as well very large.

Since total protein and TN correlate by well established conversion factors, the catalytic high temperature combustion with subsequent chemoluminescence (CLD) determination of the NO molecules is one of the methods recommended in pharmacopoeia and offers several advantages over other methods:

- Fast and precise results thanks to high sensitivity and selectivity
- No carry-over risk thanks to direct and septum-free sample injection by microliter syringe
- Wide linear working range of 0 up to 200 μg/ mL TN
- High degree of automation with up to 112 HPLC vials per sequence
- Low sample consumption requiring volumes of less than
 2 mL for multiple injection

The Perfect Solution for Your Requirements

No matter what your requirements and preferences are, the multi N/C pharma series provides a suitable device for all TOC/TN applications in the parmaceutical industry.

multi N/C pharma UV

The highly sensitive analyzer for uncompromised TOC determination in the ultra trace range using wet chemical UV digestion:

- TOC determination in direct NPOC mode free of blank interferences for highest sensitivity
- Injection volume up to 20.0 mL
- TOC detection <1 ppm requiring no oxidation reagent
- Lowest running costs no catalyst or combustion tube, no additional reagents
- Extension of working range up to 10,000 ppm using oxidation reagent which can easily be prepared by the user

Fields of application

- Optimal for WFI and Aqua Purificata applications
- Optimal for TOC cleaning validation for last rinse samples and swab extracts
- Optimal for testing of organic leachables from plastic packaging materials

multi N/C pharma HT

The flexible and sensitive instrument for TOC ultra trace analysis with best reproducibility using catalytic high-temperature combustion, and TN option:

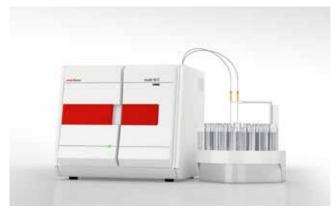
- TOC determination in direct NPOC mode for highest accuracy in the ultra trace range
- Injection volume up to 3.0 mL
- High-temperature combustion up to 950 °C
- Upgrade for direct swab combustion for TOC cleaning validation
- Optional TN cleaning validation

Fields of application

- Ideally suited for the analysis of WFI and AP
- TOC/TN cleaning validation for last rinse samples, as well as swab extracts
- TOC cleaning validation via direct swab combustion for water-insoluble or hardly soluble contaminants



multi N/C pharma UV



multi N/C pharma HT



multi N/C 3100 pharma

The high-throughput TOC analyzer for pharmaceutical applications requiring a wide concentration range, using catalytic high-temperature combustion, and TN option:

- TOC determination in direct NPOC mode for high accuracy in the ultra trace range
- Injection volume up to 1.0 mL
- Covering a TOC concentration range of 0–30,000 ppm
- Highest sample throughput with parallel purging and analyzing
- Highest degree of automation with automatic sample acidification and an effective backwash function preventing sample carryover in case of changing concentrations

Fields of application

- Optimal for TOC cleaning validation for last rinse samples and swab extracts
- Ideally suited for testing for organic leachables from plastic packaging materials
- Suited for the analysis of WFI and AP
- Optional TN cleaning validation

multi N/C 3100 pharma

multi N/C 2100S pharma

The dedicated total protein analyzer for aqueous pharmaceutical samples using high-temperature combustion in combination with septum-free direct injection via microliter syringe:

- Minimum sample consumption with a typical injection volume of 50–200 μL (max. 1.8 mL) saving costs for expensive products e.g. in vaccine quality control
- Wide working range of 0–200 ppm TN
- High degree of automation for large sample sequences (up to 112 sampling positions for HPLC vials)
- Unattended 24/7 operation thanks to self check system
- No sample carry-over due to direct injection principle by microliter syringe and effective rinse regime

Fields of application

- Total protein analysis in vaccine or biopharmaceutical production to monitor the level of antigens
- Optimal for TN cleaning validation



multi N/C 2100S pharma - the protein analyzer

Data Integrity and Reliable System Operation

Enhanced software features make sure for full data integrity and 21 CFR part 11 compliance. A complete service package, all offered from one hand provides you with maximum comfort in operating your analyzer.

Compliance with pharmacopoeia

The multiWin software fully complies with the requirements of the FDA 21 CFR part 11 and EudraLex Vol. 4 Annex 11. The user management of the software offers different user levels and personalized service and administrator logon, so that users can be authorized with different access rights. Individual passwords of adjustable complexity and expiration times guarantee that no unauthorized person has access to the system. All important events, such as logon/logoff, measurements, calibrations, electronic signatures as well as the messages generated by the Self Check System (SCS), are recorded in the audit trail.

All relevant information are protected, for example, the method and the calibration which have been used to generate the respective measurement results have to be checked and authorized by electronic signatures and can not be changed afterwards.

Individual electronic signatures are implemented into the multiWin software according to the 21 CFR part 11 requirements to sign methods, calibration reports and analysis reports of produced measurement data in a three stage process: created, examined and authorized. The principle of dual control is established in this process. The multiWin software enables LIMS import and export of data, as well as idividually customized manual or automatic PDF or CSV data export.





System suitability test (SST)

The system suitability test, which is mandatory by the pharmacopoeia regulations (e.g. USP <643>, EP 2.2.44 or JP 16, section 2.59), is an integrated function of the multiWin pharma software. Sucrose, p-benzoquinone and the water used to prepare these check solutions are measured by straight one-click selection of the SST function, which is integrated directly into the user interface. A specific SST result report is instantly generated, holding all relevant information like SST quotient and individual concentration results of the test solutions. SST tests are as well logged into the audit trail.

The Self Check System (SCS) provides valuable services in a pharmaceutical lab

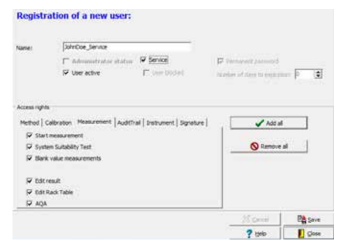
As an intelligent combination of hardware components and software functions, the SCS automatically ensures the trouble-free operation of the entire analytical system. Important parameters, such as gas flows, temperatures, pressures, system tightness, detector status, baseline stability, etc. are constantly checked, any deviations are recorded in the audit trail.

DQ, IQ, OQ, PQ and software validation – a completely reliable package!

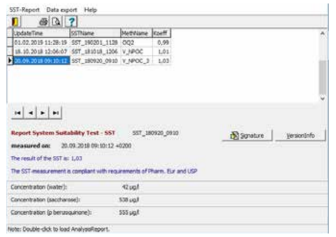
Analytik Jena provides assistance in design qualification, provides optimized and detailed installation qualification and operational qualification protocols for initial qualification or re-qualification after service or maintenance. Guidelines for performance qualifications are also offered as well as software validation services according our customized protocols.

Comprehensive Service – IQ, OQ, software validation and more

Analytik Jena offers various services including maintenance, software update and validation as well as system calibration all from one hand.



multiWin user management permits the granting of individual access rights, e.g personalized service login



Clear presentation of the measurement results in the SST report

Highest Precision Meets Outstanding Stability and Robustness

Highest accuracy in the TOC trace range

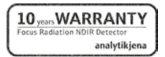
Advanced detector technology – Focus Radiation NDIR Detector

A combination of high-quality optics and the latest detector technology provide a detection system of unchallenged performance. High-energy IR radiation is focused on the micro detector by means of integrated optics. The obtained radiation density exceeds classical detectors and guarantees higher sensitivity and precision over a measuring range from 0 to 30,000 mg/L TOC without sample dilution.

Using corrosion-free materials and an electronically pulsed radiation source instead of classical, mechanically moving components, the detector is extremely stable, even when working with aggressive samples.

Focus Radiation NDIR Detector - Your benefits

- Highest measurement sensitivity and precision
- Large range detector: undiluted measurements from 0-30.000 mg/L TOC
- 10 years long-term warranty*



Efficient gas management – VITA Flow Management System

The VITA Flow Management System continues to work where classical TOC analyzers reach their limits. It enables the quick injection of large sample volumes in high-temperature TOC devices and to obtain flow-independent measurement curves. This significantly improves both the precision of measurement results and sensitivity in the trace range.

VITA also guarantees highest operating safety and reliable analysis results by continuous internal tests in combination with the self check system (SCS), like the automatic leak check.

* according to our warranty conditions: www.analytik-jena.com

VITA Flow Management System - Your benefits

- Quick injection of large sample volumes: Increase of sensitivity
- Compensation of carrier gas fluctuations for maximum precision
- Permanent leak test
- Enables Easy Cal and thus minimum calibration effort with maximum long-term stability

Calibration made easy - Easy Cal

Calibrations with VITA can be made on the basis of a single standard using different injection volumes. The obtained calibration curves are flow-independent: the calibration remains stable! This technique is ideally suited for calibration in the trace range.

Reliable sample digestion using durable technology

Both digestion techniques used in the multi N/C pharma series provide efficient digestion of the samples. The proven TOC furnace technology of the high-temperature combustion devices permits the complete oxidation of particulate samples and makes simultaneous TN determination possible.

The High Power Long Life UV Reactor of the multi N/C pharma HT uses the particularly energy-rich UV radiation of two wavelengths, 254 nm and 185 nm, for UV supported wet chemical digestion. This allows fast and complete oxidation of even the most stable organic compounds.

For both techniques, we offer a long-term warranty that guarantees stable performance and reliable analyses over many years.





multi N/C pharma Series – Specifications and Recommended Fields of Application

multi N/C pharma series	multi N/C pharma UV	multi N/C pharma HT	multi N/C 3100 pharma	multi N/C2100S pharma
			mema	
Sample digestion principle	UV/Persulfate	нтс	НТС	НТС
Injection principle	flow injection	flow injection	flow injection	direct injection
Measuring range [mg/l] TC/TOC/NPOC/TIC	0-10,000	0-30,000	0-30,000	-
Measuring range TN [mg/I] (CLD)	-	0-200	0-200	0-200
Highest injection volume [µL]	20,000 μL	3000 μL	1000 μL	500 μL
Best precision in the TOC trace range	х	X	(x) ¹⁾	-
Highest automation comfort for NPOC (automatic acidification)	X	-	X	-
Extra rinse function (best carry-over prevention)	х	-	х	х
Self Check System / VITA / Easy Cal	x	×	x	х
Best suited for following application				
WFI, aqua purificata	x	x	(x) ¹⁾	-
Extractables (TOC) from packaging materials (USP 661)	Х	X	X	-
TOC cleaning validation (last rinse / swab extracts)	X	X	X	-
TOC cleaning val. (direct swab combustion)	-	X	-	-
TN cleaning validation	-	×	×	x
Total nitrogen in vaccines / aqueous protein solutions	-	-	-	х

¹⁾ Suited for side application

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