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Guideline L-09-04 **Quality assurance of gamma cameras, SPECT and PET cameras**

The document will be translated in D/F/I in a later stage.
Note that the final layout will differ from the one of this document.

1. Objective and scope

This guideline is intended for suppliers and users of gamma cameras, single photon emission computed tomography (SPECT) scanners and positron emission tomography (PET) scanners. Its purpose is to indicate the quality assurance procedures to be followed during the commissioning, use and periodic revision of this equipment.

Quality assurance measures for organ-specific gamma cameras (e.g., those dedicated to cardiac or thyroid examinations) or PET cameras (e.g., for brain examinations), with direct application to human beings, shall in principle correspond to those defined in this guideline where relevant. The manufacturer's own methods are acceptable where necessary. In this case, the manufacturer must justify any deviations from the measures defined in this guideline.

Quality assurance measures for preclinical imaging devices, or more generally imaging devices without direct application to human beings (e.g., micro-PET or micro-PET-CT devices for small animal imaging, or devices developed for the assessment of excised tissues) are not defined in this guideline. However, manufacturers of such systems must define appropriate quality assurance measures for these cameras. For such devices, this guideline could serve as an example.

2. Legal provisions

Article 100 of the Radiological Protection Ordinance (RPO; SR 814.501; [1]) of 26 April 2017 specifies that nuclear medicine examination systems must be checked before being used for the first time and that they must also be regularly checked and revised. Article 62 of the Federal Department of Home Affairs' Ordinance on the Use of Radioactive Materials (UraM/OUMR/MMRa; RS 814.554; [2]) specifies that a reception test must be carried out by the supplier when these systems are put into service; that maintenance must be carried out every six months by an authorised qualified technician, with a status check at the end of this maintenance; and that regular checks must be carried out on the stability of these systems.

In principle, the system user will have the resources required to carry out the stability checks (phantoms, radioactive reference sources). These resources depend, among other things, on the type of construction of the equipment. They are generally determined by the supplier at the time of installation and during the acceptance test. Finally, in accordance with art. 60, para. 5 of the UraM/OUMR/MMRa, the company authorised under art. 9, let. g, RPO, to carry out quality assurance measures must notify the Federal Office of Public Health (FOPH) of the execution and results of acceptance tests and status checks.

In the case of hybrid PET-CT and SPECT-CT systems, the computed tomography (CT) scanner must also be checked in accordance with the FOPH directive on quality assurance for CT scanners [3].

If the parameters to be checked are established using a method other than that recommended in this guideline (for example in accordance with the standards [4,5,6,7]), the company responsible for the checks must prove and document the equivalent quality of the method, which has to be validated by



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FOPH upon application and before using the device. This may be the case in particular where there are manufacturing differences from the usual design.

For certain tests, instead of using unsealed radioactive sources whose activity is measured by dose calibrator, the use of sealed sources, the activity of which is officially calibrated, may be authorised, but must be approved by the FOPH. The application for approval must contain information on the manufacturing method, the activity of the source and its traceability.

The responsibility for carrying out the various quality checks and their frequency are summarised in the following Table:

	Reception tests	Status checks	Stability checks
Executing body	Licensed company	Licensed company	User
Periodicity	At reception and after major modifications ¹ , always before commissioning	Half-yearly	Regularly

Participation of the medical physicists in the execution of the quality assurance tests, in coordination with the radiographers and nuclear medicine physicians, is encouraged in the following way:

- Attending the reception tests and the half-yearly status checks performed by the manufacturer or licensed company, and involvement during the handing-over of the device from the manufacturer to the user.
- Direct involvement in the stability checks performed by the user.

The present guideline enters into force as of January 1st, 2025, with a one-year transition period, during which quality assurance checks may be carried out in accordance with the requirements of the previous version of this guideline, Revision 1, dated March 7th, 2018.¹

¹ Examples of major modifications: move of the device to another room, or addition of a ring of detectors.

3. Technical file and control records

During installation and reception testing carried out by the supplier, a technical file is drawn up for each gamma camera, SPECT camera and PET camera. This contains the documents and reports described in Annex 1 of this guideline. The results of reception tests (RT) and status checks (SC) must be recorded in a report and filed in the technical file. Due to their volume, the stability checks carried out by the user may be saved electronically or documented in a separate file. However, the technical file must contain at least the reference values established at the time of the last status check or reception test carried out by the competent personnel of the installation firms holding a FOPH authorisation and which serve as a reference for the stability checks.

¹ Temporary footnote. As agreed in our last meeting, the one-year transition period will be introduced only if explicitly requested by the manufacturers and the users.



For reception tests and status checks, the target values and tolerance intervals, if not specified in the Annexes to this guideline, must correspond to the manufacturer's specifications and must be documented and passed on to the users.

4. Reception tests and status checks for gamma cameras, SPECT cameras and PET cameras

The authorized **supplier** performs the reception test and documents it in a report, after installation of the gamma camera, SPECT camera or PET camera and before its commissioning, or following major modifications (see the Table of §2 for examples of major modifications). The status checks are performed and documented in a report by the qualified personnel of the **companies possessing an FOPH license**, in the frame of the half-yearly maintenance. During the reception tests or the status checks, some reference values for the stability checks (see next §5) are established.

The parameters to be checked, their periodicity (at reception or during the half-yearly status checks), and the reference of the procedures and measurements to be followed (mainly the latest NEMA NU-1 2018 [8] and NEMA NU-2 2018 [9] standards) are given in Annex 2 for the gamma cameras and SPECT cameras and in Annex 4 for the PET cameras. The target values and tolerance intervals, if not specified in Annexes 2 and 4, are given by the manufacturer's specifications and must be documented and made available to the user.

The **companies** possessing a BAG license must perform preventive, manufacturer-specific maintenance checks in addition to the tests performed in the frame of the half-yearly status checks mentioned in Annexes 2 and 4.

The user's dose calibrator employed during the reception tests and the status checks must be the same as the one usually used for the applications of radiopharmaceuticals to the patients.

5. Stability checks for gamma cameras, SPECT cameras and PET cameras

The **user** is responsible for performing and documenting in reports the stability checks. In particular, in order to ensure the independence of these measurements, the user cannot mandate the supplier or the **authorised** company to perform the stability checks.

The parameters to be checked, their periodicity, and the procedures and measurements to be followed for the tests are given in Annex 3 for the gamma cameras and SPECT cameras, and in Annex 5 for the PET cameras. The target values and tolerance intervals, if not specified in Annexes 3 and 5, must be given by the manufacturer's specifications and must be documented and made available to the user.

If a stability check fails, the user has to contact the responsible medical physicist and corrections must be made before the machine can be used clinically.

The user's dose calibrator employed during the stability checks must be the same as the one usually used for the applications of radiopharmaceuticals to the patients.

6. References

[1] Radiological Protection Ordinance (RPO, SR 814.501) of 26 April 2017 (status as of 1 January 2022).

[2] Ordinance of the FDHA on the Use of Radioactive Materials (UraM/OUMR/MMRa, SR 814.554) of 26 April 2017 (status as of 30 January 2018).



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- [3] Guideline R-08-08: Quality assurance for computed tomography (CT) scanners, www.bag.admin.ch/rad-directives.
- [4] IEC 61675-1:2022, Radionuclide imaging devices – Characteristics and test conditions – Part 1: Positron emission tomographs.
- [5] IEC 61675-2:2015, Radionuclide imaging devices – Characteristics and test conditions – Part 2: Gamma cameras for planar, wholebody, and SPECT imaging.
- [6] IEC TR 61948-2:2019, Nuclear medicine instrumentation – Routine tests – Part 2: Scintillation cameras and single photon emission computed tomography imaging.
- [7] IEC TR 61948-3:2018, Nuclear medicine instrumentation – Routine tests – Part 3: Positron emission tomographs
- [8] NEMA Standards Publication NU 1-2018, Performance Measurements of Gamma Cameras.
- [9] NEMA Standards Publication NU 2-2018, Performance Measurements of Positron Emission Tomographs (PET).
- [10] Medical Devices Ordinance (MedDO, SR 812.213) of 1 July 2020 (status as of 26 May 2022).
- [11] IAEA Human Health Series No.6, Quality Assurance for SPECT Systems
- [12] American association of physicists in medicine, Report No.117, Research Needs of Radiation Protection



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Annex 1 Content of the technical file

Content	Remarques
FOPH licence	Licence to operate a radiology system for medical use for SPECT-CT and PET-CT. Licence to use unsealed radioactive sources for independent gamma cameras (without CT).
Reception test reports	Includes records of reception tests and records of handover of the installation to the user. Target values and tolerance intervals, if not specified in the Annexes to this guideline, must correspond to the manufacturer's specifications and must be documented and passed on to users.
Status checks reports	Certain reference values for stability checks are redefined at the time of each status check. The target values and tolerance intervals, if not specified in the Annexes to this guideline, must correspond to the manufacturer's specifications and must be documented and passed on to users.
Revision checklists (maintenance)	According to the manufacturer's data.
Reference values for the stability checks	Filing of reference values and procedures to be followed for the stability checks; the reports of the stability checks are filed in electronic form or on paper.
Indication of the version and storage location of the instructions for use	The instructions for use must be available at all times in the language used in the unit.
CE Declaration of Conformity	In accordance with the Medical Devices Ordinance (MedDO; SR 812.213; [10]).
Indication of ambient dose rates in the room where the camera is installed and in adjacent rooms	The maximum permitted ambient dose rates must be respected in all rooms. For PET equipment, the radiation protection plans and calculation tables in accordance with the guideline "Radiation protection measures relating to the construction of PET installations" [11] must be available.



Annex 2 Reception tests and status checks for gamma cameras and SPECT cameras: parameters to be checked, periodicities, references and methodologies
Executing body: manufacturer or licensed company

RT = reception test

SC = status check (half-yearly if not specified otherwise)

N°	Parameter	Periodicity	Reference and method	Devices concerned	Nuclids, activities and phantoms
G1	Intrinsic flood field uniformity	RT and SC	NEMA NU 1-2018, part 2.4 The intrinsic uniformity of the system is measured, i.e., the response of the system without a collimator to a uniform flux of radiation from a point source.	All, except those ones for which the collimators cannot be removed	Contrarily to what is requested by NEMA, it is sufficient to measure this parameter for two nuclides, Tc-99m and another one with photons of higher energy (if Tc-99m is the only nuclide used at the center, the test is executed with Tc-99m only).
G2	System image homogeneity & collimator check	RT and SC	Both the status/condition of the collimators and the images produced with the collimators (homogeneity) shall be checked visually.	All	RT: check of all the collimators. SC: check of the mostly used collimator.
G3	Intrinsic energy resolution	RT and SC	NEMA NU 1-2018, part 2.3 The ratio of the photopeak FWHM ² to the photopeak center energy shall be calculated.	All	Use of Tc-99m and/or Co-57 if the latter is used for daily quality assurance.
G4	Intrinsic spatial resolution	RT, and SC if the G5 (system)	NEMA NU 1-2018, part 2.1 The FWHM and FWTM ³ shall be determined.	All, except direct conversion	Use of Tc-99m.

² Full width at half maximum.

³ Full width at tenth maximum.



		spatial resolution without scatter) SC is out of tolerance		detectors, such as CZT ⁴ detectors	
G5	System spatial resolution without scatter	RT and yearly SC	NEMA NU 1-2018, part 3.1 The FWHM and FWTM of the line spread function shall be measured.	All	The test has to be performed for all collimators. Use of two capillary tubes ⁵ .
G6	System planar sensitivity and collimator penetration and scatter	RT and SC	NEMA NU 1-2018, part 3.3 The ratio of collimated counts to the known activity of a planar source shall be determined.	All	RT: all the collimators. SC: with the mostly used one only. The activity has to be injected into Petri dishes (as requested by NEMA), but the use of traceable sources is allowed.
	System volume sensitivity	RT and yearly SC	NEMA NU 1-2018, part 4.4 Measurement of the total system sensitivity to a uniform concentration of activity, and determination of the average volume sensitivity per axial centimeter.	RT: all SPECT devices SC: only for SPECT devices using absolute	Use of Tc-99m only. Use of a fillable cylindrical phantom.

⁴ Cadmium zinc telluride (CdZnTe).

⁵ As an alternative to the method described in the NEMA standard, see the IAEA Human Health Series No.6 [11] for the methodology to create a system spatial resolution phantom with four capillary tubes.



				quantification in clinical examinations	
G7	System count rate performance with scatter	RT	<p>NEMA NU 1-2018, part 3.5 or AAPM report 117</p> <p>The maximum observed count rate and the observed count rate for a 20% count loss shall be measured, and a curve of observed count rate with scatter versus input count rate shall be made.</p> <p>As an alternative to the NEMA methodology, the use of one of the methods mentioned in the AAPM report 117 [12] is allowed.</p>	All	Use of Tc-99m and of a cylindrical phantom.
G8	Intrinsic spatial linearity	RT	<p>NEMA NU 1-2018, part 2.2</p> <p>The goal of the test is to identify potential distortions in the images. The intrinsic spatial differential and absolute linearity has to be reported.</p>	All, except direct conversion detectors, such as CZT detectors	Use of Tc-99m and a lead mask with parallel slits.
	System linearity	RT and yearly SC	In addition, a visual check for linearity and absence of distortions shall be performed.	All	Use of the data from G5.
G9	Whole-body system spatial resolution	RT	<p>NEMA NU 1-2018, part 5.1</p> <p>The system spatial resolution without scatter shall be measured parallel and perpendicular to</p>	All, except step-and-shoot whole-body planar acquisition devices	Use of two capillary tubes ⁶ of Tc-99m. The camera shall be equipped with a collimator.

⁶ As an alternative to the method described in the NEMA standard, see the IAEA Human Health Series No.6 [11] for the methodology to create a system spatial resolution phantom with four capillary tubes.



	without scatter		the direction of continuous motion and expressed as FWHM or FWTM of the line spread function.		
G10	System alignment	RT and SC	NEMA NU 1-2018, part 4.1, or according to the manufacturer The transaxial alignment of acquired images with the system's mechanical center of rotation shall be measured. Likewise, for multi-head SPECT imaging systems, the axial alignment of images from the individual heads shall be measured.	SPECT devices only	If the NEMA method is used, three Tc-99m or Co-57 point sources shall be used.
G11	Detector-detector sensitivity variation	RT and yearly SC	NEMA NU 1-2018, part 4.5 Assessment of the relative difference in sensitivity of the individual detectors. All the projection images collected using each detector shall be summed up. The difference between the largest and the smallest of these quantities shall be computed.	RT: all multi-detector camera systems, when applicable Yearly SC: only if the yearly SC G6 (system volume sensitivity), if applicable, fails.	Use of the data from the second part of G6. The use of one collimator is enough.
G12	Tomographic contrast and absolute	RT and SC	NEMA NU 1-2018, part 6 Images simulating those obtained in a total body imaging study with hot and cold lesions of	SPECT devices only	Use of the NEMA/IEC image quality phantom (same phantom as for the PET reception test P5) ⁷ . Use of Tc-99m. Background activity shall be 20 kBq/ml. The sphere-to-background activity concentration shall be 8:1.

⁷ Or an equivalent phantom, such as the PET ACR phantom described the AAPM report n° 126.



	quantificatio n accuracy		<p>different diameters are produced. Measurement of the contrast of cold and hot spheres in a warm background, of the variability of the background regions, and of the deviation of the large, cold reconstructed lung region without activity.</p> <p>For centers using absolute quantification (image output in Bq/ml), the absolute activity concentration shall also be measured and compared to the injected activity concentration.</p> <p>Reference values from the second part of the reception test (absolute quantification accuracy) shall be determined for the later stability checks KG4.</p> <p>The half-yearly SC concerns the tomographic contrast part of the test only.</p>		<p>The user shall be involved in the half-yearly SC (tomographic contrast part of the test).</p> <p>The user must take note of the values of the second part of the reception test (absolute quantification accuracy) for the later stability checks KG4.</p>
G13	Image homogeneity and precision of quantificatio n	RT	<p>Verification of the background activity. The mean coefficient of variation (image homogeneity) and mean background activity concentration (precision of quantification) shall be computed according to the paragraph "Analysis" of Annex 6.</p> <p>The corresponding stability check is KG5.</p>	<p>Image homogeneity: all devices.</p> <p>Precision of quantification: only for SPECT devices using absolute quantification in clinical examinations.</p>	<p>Use of the data from the second part of G6.</p> <p>Use of Tc-99m only.</p> <p>Tolerance interval: see paragraph "Tolerance" of Annex 6.</p>



G14	SPECT-CT coregistration accuracy	RT and SC	<p>Preferably NEMA NU-1 2018, part 7, or according to the manufacturer</p> <p>NEMA NU-1 2018 method: The alignment accuracy between SPECT and CT image volumes shall be measured, by using data acquired with SPECT and CT fiducial markers at six locations.</p>	SPECT-CT only	<p>Radionuclide for the SPECT portion of the fiducial markers: Tc-99m.</p> <p>It is allowed not to use the weights prescribed by NEMA NU-1 2018.</p>
G15	Pixel size	RT and SC	<p>According to the manufacturer.</p> <p>The goal of the test is to make sure that the system determines distances correctly, by comparing a known length (typically 30 cm) with the length measured by the system.</p>	All	According to the manufacturer.





Annex 3 Stability checks for gamma cameras and SPECT cameras: parameters to be checked, periodicities, references and methodologies
Executing body: user

N°	Parameter	Periodicity	Reference and method	Devices concerned	Nuclids, activities and phantoms	Tolerances
KG1	Background count rate	At the beginning of each working day	Perform a blank scan of 2.5 to 3 minutes. A given number of counts (manufacturer-specific) should not be exceeded to make sure that no radioactive contamination is present in the system. In case of contamination, measures have to be taken.	All	--	According to the manufacturer
KG2	Check of the energy window	Qualitatively: ideally before each exam, but at least once a day Quantitatively: once a week	Check of the correct position of the Tc-99m peak. A visual check shall be performed before each exam, with the injected patient himself (no additional source needed). Once a week, a quantitative measure shall be performed and documented, in order to follow any shift in the energy window.	All	Qualitative tests: use of Tc-99m. Quantitative tests: use of Tc-99m or of Co-57 pencils.	According to the manufacturer
KG3	Homogeneity	Weekly	Methodology (with or without collimator, maximal number of counts, point source or flat source) according to the manufacturer.	All	See column "Reference and method".	According to the manufacturer
KG4	Absolute quantification accuracy	Yearly	The same method shall be used as for G12, but only for the part of the test related to absolute quantification accuracy. The results shall be compared with the reference values from the reception test G12.	Only for devices using absolute quantification	See G12.	According to the manufacturer.



				in clinical examinations		
KG5	Image homogeneity and precision of quantification	Half-yearly	<p>See Annex 6 for detailed instructions and tolerance intervals.</p> <p>Verification of the background activity. The coefficient of variation and mean background activity are measured and compared with given tolerance intervals.</p>	<p>Only for SPECT devices using absolute quantification in clinical examinations</p>	<p>Use of Tc-99m only.</p> <p>Radionuclide: use of Tc-99m only with 10 - 20 MBq/kg. Use of a cylindrical fillable phantom (same as for P7 and KP6).</p>	<p>See paragraph "Tolerance" of Annex 6.</p>





Annex 4 Reception tests and status checks for PET cameras: parameters to be checked, periodicities, references and methodologies
Executing body: manufacturer or licensed company

RT = reception test

SC = half-yearly status check

N°	Parameter	Periodicity	Reference and method	Devices concerned	Nuclids, activities and phantoms
P1	Spatial resolution	RT	NEMA NU 2-2018, part 3 Imaging point sources in air and then reconstructing images by a linear reconstruction method, such as filtered back projection, with no smoothing or apodization. The FWHM and FWTM of the image point spread function are measured.	All	Use of F-18 or Na-22.
P2	Scatter fraction, count losses, and randoms	RT	NEMA NU 2-2018, part 4 The relative system sensitivity to scattered radiation, the effects of system dead time and the generation of random events are measured.	All	Use of a dedicated phantom (solid right circular cylinder with a hole drilled parallel to the central axis of the cylinder) with approximately 500 MBq of F-18.
P3	Sensitivity	RT	NEMA NU 2-2018, part 5 The sensitivity of the system (rate in counts per second that true coincidence events are detected for a given source strength) is measured.	All	Use of F-18 with approximately 10 MBq activity. Use of the NEMA sensitivity phantom (five sleeves of different diameters). The use of a cylinder with Ge-68 sources (according to the former NEMA NU 2-1994) is not allowed ⁸ .
P4	Accuracy: corrections for	RT	NEMA NU 2-2018, part 6	All	Test performed with the measurement data from P2.

⁸ Among other arguments because in 3D mode acquisition, the scatter is not negligible and the test is more precise with the sleeves foreseen by the NEMA norm.



	count losses and randoms		The accuracy of corrections for dead time losses and random event counts is measured.		
P5	Image quality, accuracy of corrections	RT	<p>NEMA NU 2-2018, part 7</p> <p>Images simulating those obtained in a total body imaging study with hot lesions of different diameters are produced, with activity also be present outside the region of interest of the scanner to reproduce the clinical routine. Image contrast and background variability ratios for hot spheres are used as measures of image quality. The accuracy of corrections is determined from the uniform background and cold lung insert regions.</p>	All	Use of the NEMA/IEC image quality phantom ⁹ (body phantom). Radionuclide to be used: F-18 with an activity concentration¹⁰ of 5.3 MBq/kg . Use as well of the solid right circular cylinder (scatter phantom, same as for P2) with same activity concentration. Fill the spheres with a unique activity ratio of 4:1.
P6	Qualitative contrast test	RT	<p>Qualitative determination of the contrast by counting and reporting the number of visible lesions in images simulating those obtained in a total body imaging study with hot lesions of different diameters.</p> <p>The size of the smallest visible lesion detected during RT will be the reference value for the later stability checks KP5.</p>	All	<p>Use of the NEMA/IEC image quality phantom¹¹ prepared for P5 one hour after P5 has been performed¹². The test has to be performed <u>without</u> the scatter phantom and with the clinically recommended algorithm.</p> <p>For the centers undergoing accreditation tests for quantitative measurements (e.g., EARL), if an additional concentration ratio is used at reception (in addition to the 4:1 ratio specified by NEMA for P5), this additional concentration ratio can be used for P6 and KP5.</p>

⁹ Or an equivalent phantom, such as the PET ACR phantom described in the AAPM report n° 126.

¹⁰ Corresponding to 370 MBq for a mass of 70 kg.

¹¹ Or an equivalent phantom, such as the PET ACR phantom described in the AAPM report n° 126.

¹² To have an activity concentration of 3.5 MBq/kg of F-18, corresponding to the diagnostic reference level (DRW/NRD).



P7	Image homogeneity and precision of quantification	RT	<p>See Annex 6 for detailed instructions and tolerance intervals.</p> <p>Verification of the background activity. The coefficient of variation and mean background activity are measured and compared with given tolerance intervals.</p> <p>The corresponding stability check is KP6.</p>	All	Radionuclide: F-18 with 30-100 MBq. Use of a cylindrical fillable phantom covering the axial field of view (diameter at least 20 cm).
P8	PET-CT coregistration accuracy	RT and SC	<p>PET-CT devices: preferably NEMA NU-2 2018, part 9, or according to the manufacturer. PET-MR devices: according to the manufacturer.</p> <p>NEMA NU-2 2018: The coregistration error between PET and CT data is determined by using data acquired with PET and CT fiducial markers at six locations within the PET and CT field of view.</p>	For PET-CT and PET-MR devices only	<p>Radionuclide for the PET portion of the fiducial markers: F-18 or Na-22 as prescribed by NEMA NU-2 2018, or another radionuclide according to the manufacturer.</p> <p>It is allowed not to use the weights prescribed by NEMA NU-2 2018.</p>
P9	Time-of-flight resolution	RT	<p>NEMA NU-2 2018, part 8, has to be followed for devices sold after this Guideline enters into force and for devices that underwent a major upgrade. For the older devices, the test has to be performed with the datasheet of the system provided by the manufacturer.</p> <p>The time-of-flight resolution (uncertainty in detecting the arrival time-difference of two photons in a coincidence event) is assessed by measuring the FWHM of the detector response.</p>	For devices with time-of-flight mode	If NEMA NU-2 2018 is followed: use of the measurement data from P2.



Annex 5 Stability checks for PET cameras: parameters to be checked, periodicities, references and methodologies

Executing body: user

N°	Parameter	Periodicity	Reference and method	Devices concerned	Nuclids, activities and phantoms	Tolerances
KP1	PM check	Every working day	Check of the amplification factor (gain), offset PM, and homogeneity.	All	Use of an external positron source, except for those devices equipped with an appropriate internal radioactive source (for example lutetium sources).	According to the manufacturer
KP2	Control of the energy window	Every working day	Check of the setting and FWHM resolution.	All	Use of an external positron source, except for those devices equipped with an appropriate internal radioactive source (for example lutetium sources).	According to the manufacturer
KP3	Coincidence timing	Weekly check for the semiconductor devices, on every working day for the other (PM) devices.	The parameter to be checked is as defined by the manufacturer.	All	Use of an external positron source.	According to the manufacturer
KP4	Visual verification of the system	Every working day	Comparison of the sinograms.	All	Use of an external positron source, except for those devices equipped with an appropriate internal radioactive source (for example lutetium sources).	According to the manufacturer
KP5	Qualitative contrast test	Yearly	Qualitative determination of the contrast by counting and reporting the number of visible lesions in images simulating those obtained in a total body imaging study with hot lesions of different diameters.	All	Prepare the NEMA/IEC image quality phantom ¹³ with an activity concentration of 3.5 MBq/kg of F-18 and fill the spheres with a single activity ratio of 4:1. The test has to be performed <u>without</u> the scatter phantom and with the clinically recommended algorithm.	Minimal sphere size has to be the same (at least) as the one observed during reception test P6.

¹³ Or an equivalent phantom, such as the PET ACR phantom described in the AAPM report n° 126.



					For the centers undergoing accreditation tests for quantitative measurements (e.g., EARL), if an additional concentration ratio is used at reception (in addition to the 4:1 ratio specified by NEMA for P5), this additional concentration ratio can be used for the reception test P6 and KP5.	
KP6	Image homogeneity and precision of quantification	Half-yearly	See Annex 6 for detailed instructions. The coefficient of variation and mean background activity are measured and compared with given tolerance intervals.	All	Radionuclide: F-18 with 30-100 MBq. Use of a cylindrical fillable phantom (same as for P7). For the centers undergoing accreditation tests for quantitative measurements (e.g., EARL), this test can be skipped. The results of the accreditation tests shall be filed in the stability checks file.	Tolerance intervals are given in Annex 6.



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Annex 6 Detailed instructions for reception tests G13 and P7 and stability checks KG5 and KP6: image homogeneity and precision of quantification

Purpose: To verify the accuracy of image quantification and image homogeneity.

Materials: Cylindrical phantom which should cover¹⁴ the axial FOV filled with ¹⁸F or Tc-99m. Diameter of the phantom should be at least 20 cm.

Procedure: Prepare a syringe with about¹⁵ 50 - 100 MBq of ¹⁸F, or with a Tc-99m activity corresponding to an activity concentration of 10 – 20 MBq/kg, by accurately measuring it in the same dose calibrator used for clinical routine, writing value of activity and time of assay. The clocks used for recording the time assay should be checked against the scanner time. Put the activity in the phantom, mixing with water solution thoroughly to get a uniform radioactivity distribution. Measure the residual activity in the syringe and evaluate the net activity put into the phantom. Add water until the phantom is completely filled. Place the phantom on the phantom holder provided by the manufacturer and move the phantom at the beginning of the CT FOV, making sure that the phantom is centered in the FOV also with respect to the height. In case of no phantom holder, simply place it on the patient's table.

The acquisition should be performed by using the protocol provided by the manufacturer (same protocol for the reception tests P7 and the later stability checks KP6). In case no acquisition protocol is provided, the standard protocol used for clinical routine should be used (body, head, ...). Make sure that a low dose CT for attenuation and scatter correction purposes is included in the procedure. Attention should be paid when inserting data regarding ¹⁸F activity assayed by the dose calibrator, time of assay and weight of the volume of the ¹⁸F solution used to fill the phantom. Acquisition stop condition should be set for 100 million counts.

Analysis: For the acquired images, draw one circular ROI with a diameter greater than 5 cm, at least 2 cm from the phantom edge on the reconstructed central slice and on ± 5-6 adjacent slices (total of 12 ROIs). Measure the mean and standard deviation activity concentration for each ROI and calculate the coefficient of variation (COV) for each slice:

$$\text{coefficient of variation (\%)} = \frac{\text{standard deviation}}{\text{mean}} \times 100$$

The mean COV and mean background activity concentration can then be calculated among the different ROIs.

Tolerance: The mean background activity concentration shall be within ±10% of the true activity concentration¹⁶ and the mean coefficient of variation should be < 15%.

¹⁴ Phantoms covering the whole FOV might not be commercially available for long-axial FOV PET-CTs. In such a case, the user can either use several small phantoms, or move the same phantom to different bed positions. The adopted method shall allow checking the whole FOV.

¹⁵ With this activity range, the user can choose to comply with the requirements of the EANM Research GmbH (EARL) initiative.

¹⁶ A value of ±5% should be achievable for most of the devices installed in Switzerland.