Procedure for determining the specific activities of radionuclides in fish by gamma spectrometry

G-γ-SPEKT-FISCH-01

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1 Scope

The procedure described in the following is suitable for gamma spectrometrical analysis of samples of fish that have to be monitored according to the German Precautionary Radiation Protection Act (StrVG) in the IMIS-routine programme and the Guideline for the Monitoring of Emissions and Immissions of Nuclear Installations during Regular Operation.

The procedure is suitable for low-level measuring in the framework of radio-ecological studies due to its low limit of determination after dry-ashing of the sample.

2 Sampling

The sampling of fish – especially marine fish – relies to a wide extent on the cooperation of professionals in the respective industries (professional fishermen) and the product range available at fish markets.

It is in the responsibility of the person collecting the sample to ensure that a sample taken can be traced back to the sampled water body. The sampling position needs to be recorded as exact as possible. The name of the samples fish species is also an essential part of the sample label. In case of streams, the river kilometer has to be additionally reported, while for saltwater fish the FAO fishing areas or coordinates of the catch must be reported by the fishermen. According to an EU regulation, marketed fish must be labelled with its registered trade name and the fishing area (1).

The required sample size depends on the type of study to be carried out. For studies according to the Precautionary Radiation Protection Act and the Guideline for the Monitoring of Emissions and Immissions of Nuclear Installations, at least 1,5 kg of fish flesh is required. This equals a minimum of 3 kg non-filleted original sample. If the samples are to be used for additional analyses like strontium or isotopes of plutonium and americium, the sample size should be enlarged to 5 kg of fish flesh. The fish flesh will usually be preferred for analysis, but some species, e. g. sprat, must be examined as a whole fish because of their small size. The determining factor for sample preparation is the typical consumption of this species. In case of individual fishes, the person collecting the sample must ensure that only marketable fish is sampled.

The freshly collected fish are killed quickly at the point of sampling, sorted by species, packed in plastic bags, and transported to the laboratory in ice-filled containers. The fish samples are stored deep-frozen at ca. -20 °C until they can be processed further in the laboratory.

3 Analysis

3.1 Principle of the procedure

The whole fish, or fish flesh obtained after filleting, is dried at 100 °C to 110 °C and then ashed at a maximum temperature of 420 °C. The specific activity of radionuclides in the homogenised ash are analysed using a germanium semiconductor detector.

The specific activities of iodine isotopes cannot be determined quantitatively because of the low boiling point of iodine (184 °C). The quantitative determination of iodine isotopes in fish requires the use of the procedure $G-\gamma$ -SPEKT-FISCH-02.

3.2 Sample preparation

All items used for preparation of the ashes, e. g. stainless steel bowls, ceramic bowls, measuring beakers, etc. must be cleaned using cleaning solution prior each use.

During filleting of fresh or defrosted fish, attention needs to be paid to deboning them as completely as possible. The obtained fish flesh is roughly chopped up. Chopped fish flesh or whole fish samples are placed in suitably large stainless steel or ceramic bowls that are laid out with ash-free transparent paper. Frozen fillets are defrosted directly in prepared stainless steel bowls in order to include tissue fluids liberated during the defrosting process in the subsequent drying and ashing steps. The total fresh mass used is to be recorded.

Afterwards, fish samples are dried until constant mass is reached, i. e. about one day or two days at a temperature of about 110 °C. Thereafter, the dry mass (TM) is determined. The ratio between fresh mass and dry mass in fish meat is about 5.

The time to reach the maximum ashing temperature depends on the lipid content of the sample; for normal fish it takes 45 hours. If fish with high lipid content, e. g. eel, mackerel, herring or sprat are to be ashed, this time needs to be extended in order to avoid the ignition of the sample. In order to achieve a complete ashing of the sample, the maximum temperature is held for another 96 to 154 hours.

The following temperature program is commonly used for fish samples with average lipid content:

Step 1: Linear increase of the temperature to 230 °C within 3 hours;

Step 2: Linear increase to the maximum temperature 420 °C within

42 hours;

Step 3: Hold the maximum temperature for at least 96 hours, optimal

154 hours.

The final temperature of 420 $^{\circ}$ C is selected to avoid the volatilisation of caesium. Experiments to this effect, using the temperature program listed above, revealed ashing-induced losses of about 1 $^{\circ}$ C.

Note

The danger of ignition during the ashing process rises, when the sample consists of material high in lipid content, e. g. cod liver. Therefore, the bowls must contain only thin layer of these samples. The maximum temperature must be selected lower than 380 °C in these instances.

Once ashing has been completed and the sample has cooled down, the mass of the ash is recorded. For fish meat, a ratio of fresh mass to ash mass of about 80 (range 65 to 95) has usually to be expected; in whole fish, it may be lower by a factor of 2. Carbon residues still contained in the ash will not affect the results of the gamma spectrometric analysis. The ash is homogenised using a mortar, filled into a suitably large measuring beaker, and then carefully manually compressed with a stamp. The obtained filling height is recorded. The measurement geometry is calculated from the mass and filling height of the sample and the diameter of the measuring beaker. The compressed density is commonly around $0.5 \, \mathrm{g \cdot cm^{-3}}$.

3.3 Radiochemical separation

No radiochemical separation is required.

4 Measuring the activity

4.1 General

The basics of gamma spectrometry are discussed in the basic chapter γ -SPEKT/GRUNDL of this procedures manual and in the literature (2, 3, 4).

4.2 Calibration

4.2.1 Determination of the detection efficiency

For determining the energy-dependency of the detection efficiency, reference is made to the basic chapter γ -SPEKT/GRUNDL of this procedures manual.

A calibration using a calibration source is described here. A suitable calibration source is e. g. an aqueous solution of a traceable standard or a resin containing traceable amounts of radioactive compounds. Mathematical calibration of the detection efficiency may also be used if the required premises are fulfilled.

The relative standard uncertainties of the activities used should be less than 1 %. The relative standard uncertainty of Pb-210 may be larger due to conditions of manufacturing. The measurement duration of the calibration has to be selected to ensure standard uncertainties of the net count rates to be below 1%.

An example of a calibration using an aqueous solution with a density of 1,03 g·cm⁻³ (hydrochloric acid, 2 mol·l⁻¹) is described in the following.

Two alternatives are available which differ in the selection of radionuclides and depend on the consideration of coincidence summation.

4.2.1.1 Calibration using single-line radionuclides

A calibration solution containing sufficient activity of a gamma emitting radionuclide with only one emission line in the energy range between 100 keV and 1200 keV, e. g. Sr-85, Cs-137, Mn-54 or Zn-65, respectively, is used. To extend this calibration curve to energies smaller than 100 keV, a second solution containing e. g. Pb-210, Am-241, Cd-109 und Co-57 is required. A curve is fitted to the values obtained by equation (1):

$$\varepsilon_{W}(E_{r,i}) = \frac{R_{n,r,i}}{A_{r} \cdot p_{\gamma}(E_{r,i})} \cdot f_{2}(E_{r,i}) \cdot f_{4,r}$$
(1)

The symbols in equation (1) mean:

 A_r activity of the radionuclide r used for the calibration at the reference date in Bq;

 $E_{r,i}$ energy of the gamma line *i* of radionuclide r in keV;

 $\varepsilon_W(E_{r,i})$ detection efficiency for water, depending on energy and fill level in Bq⁻¹·s⁻¹;

 $R_{n,r,i}$ net count rate of line *i* of radionuclide r in s⁻¹;

 $f_2(E_{r,i})$ self-absorption correction factor in water relative to hydrochloric acid (2 mol·l⁻¹):

$$f_2(E_{r,i}) = \frac{\varepsilon_{MC}(H_2O)}{\varepsilon_{MC}(HCI)};$$

 ε_{MC} detection efficiency, calculated by Monte Carlo (MC)-simulation in Bq⁻¹·s⁻¹;

 $f_{4,r}$ the decay correction factor of radionuclide r

$$f_{4,r} = \frac{\lambda_r \cdot t_m}{\left(1 - e^{-\lambda_r \cdot t_m}\right)} \cdot e^{-\lambda_r \cdot t_A};$$

 t_{A} time period between commencement of measuring and reference date of the standard in s;

 $t_{\rm m}$ duration of measurement in s;

 $p_{\gamma}(E_{r,i})$ probability of gamma emission of line *i* from radionuclide r.

K-40 is the only available single-line emitting radionuclide in the energy range greater than 1200 keV, but it requires large periods of measurement. Therefore, the multi-line gamma emitting nuclide Y-88 is used instead, but afterwards a correction for coincidence summing effects is needed to avoid a bias in the calibration curve (see section 4.2.1.2).

4.2.1.2 Calibration using multi-line radionuclides

The experimental effort may be reduced when corrections for coincidence summation can considered already during calibration, because then the energy range between 47 keV and 1836 keV may be covered by a single solution containing a mixture of radionuclides. This mixture consists of the radionuclides for the lower energy range mentioned in section 4.2.1.1 and additional multi-line radionuclides like Te-123m, Cr-51, Sn-113, Cs-134, Co-60 and Y-88. A calibration curve is fitted to values calculated by equation (2):

$$\varepsilon_{W}(E_{r,i}) = \frac{R_{n,r,i}}{A_{r} \cdot p_{\gamma}(E_{r,i})} \cdot f_{1}(E_{r,i}) \cdot f_{2}(E_{r,i}) \cdot f_{4,r}$$
(2)

The additional symbol in equation (2) means:

 $f_1(E_{r,i})$ correction factor for coincidence summation of line i emitted by the radio-nuclide r.

4.2.1.3 Filling height dependent calibration curves

The calibration must be carried out for different filling heights. The filling heights of the measuring preparations are selected according to the range of filling heights expected in samples. Four energy-dependent calibration curves for different filling heights are shown in Figure 1.

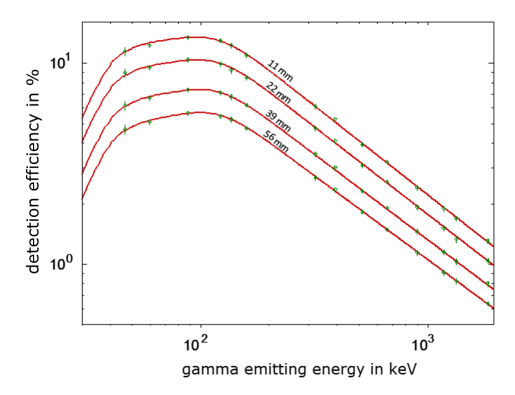


Fig. 1: Determined detection efficiency in % and fitted calibration curves for different filling heights between 11 mm and 56 mm, ordered from top to bottom; the calibration was carried out on an n-type detector with aluminum endcap.

4.2.2 Correction for self-attenuation

The self-attenuation of a sample depends on its density, its elemental composition and the energy-dependent mass attenuation coefficients of the single elements.

A self-attenuation correction factor $f_5(E)$ is used to correct the detection efficiency for water as obtained in section 4.2.1 to those of fish-ash. This correction factor depends on the gamma-ray energy, the density of the preparation, the elemental composition of the ash and the beaker and finally the filling height, respectively. Software-tools are essential for its calculation. Therein, an approximate knowledge of the detector's geometric dimensions is sufficient.

The elemental composition listed in Table 1 may be used to calculate the selfattenuation correction factor of ash from fish flesh as weighted mean of the massattenuation factor from single elements (5, 10).

Tab. 1: Typical composition of ash from fish flee	Гаb. 1:	i composition of asn	from fish fiesh
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Element resp. compound	relative mass fraction				
K	0,245				
PO ₄	0,460				
Cl	0,109				
Na	0,067				
ZnO	0,0576				
CaO	0,0308				
MgO	0,0305				

The following equation is used to calculate the energy depending self-attenuation factor $f_5(E)$ in the energy range larger than 45 keV:

$$f_{5}(E) = a_{1} \cdot E^{a_{2}} + a_{3} \cdot E^{a_{4}} = \frac{\varepsilon_{MC}(H_{2}O)}{\varepsilon_{MC}(Ash)}$$
(3)

with:

 $f_5(E)$ self-attenuation correction factor for ash relative to water;

E gamma-ray energy in keV;

 a_k parameter obtained through curve fitting (k = 1, ..., 4).

Figure 2 shows four curves fitted to calculated values of the energy dependent self-attenuation factor obtained for ash from fish flesh and hydrochloric acid (2 mol·l⁻¹). It is obvious that $f_5(E)$ strongly increases in the energy range below 45 keV. For energies significantly below 40 keV it might be necessary to include a third power function term in equation (3).

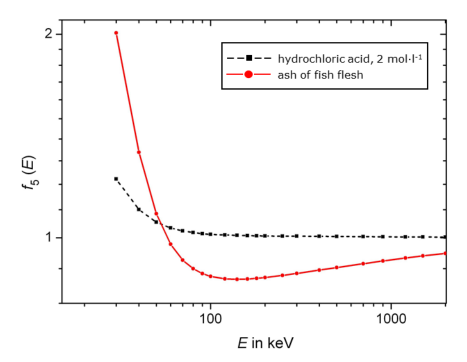


Fig. 2: Calculated self-attenuation correction factors as function of the gamma-ray energy after calibration using hydrochloric acid (2 mol·l⁻¹) and in ash from fish flesh; the filling-height are 40 mm, the density of the ash is 0,47 g·cm⁻³ and that of hydrochloric acid 1,03 g·cm⁻³.

Values of the relative standard uncertainty of $f_5(E)$ for photon energies above 100 keV have been determined to be around 1,5 %; the influence of measurement geometry and elemental composition of the ash increases for lower photon energies, which leads to a strong increase in relative standard uncertainty.

4.2.3 Detection efficiency in ash

The analysis for specific activities of gamma-ray emitting radionuclides in the sample requires to determine the detection efficiency in fish ash $\varepsilon_A(E)$. It is calculated as the quotient of the detection efficiency in water $\varepsilon_W(E)$ and the self-attenuation correction factor $f_5(E)$.

$$\varepsilon_{A}(E) = \frac{\varepsilon_{W}(E)}{f_{5}(E)}$$
(4)

In case that the standard uncertainty of the activities used for calibration is less than 1 %, values of the relative combined standard uncertainty of the detection efficiency in ash $\varepsilon_A(E)$ in the range of 2 % to 4 % for photon energies larger than 100 keV are achievable; they may be much larger for photon energies below 100 keV.

4.3 Background

Background spectra need to be recorded at regular intervals to check the gamma spectrometer for contaminations. Measuring periods for this purpose should be at least 1,5 days to 3 days. It is recommended to obtain a mean value and standard deviation of the net count rate of each individual background gamma-ray energy from two or three background spectra. This will typically encompass lines relating to Ra-226 decay products which may exhibit some count rate fluctuation depending on the ventilation of the room. They may be reduced by e. g. the supply of gaseous nitrogen escaping from the dewar used for cooling of the detector to the measurement chamber.

For samples of fish, reliably determining the K-40 background count rate of the background effect is important in order to be able to use the K-40 activity determined in fish for verifying the correctness of the subsequent sample spectrum analysis.

4.4 Gamma spectrometrical measurement

The determination of the specific activity of gamma emitting radionuclides in fish ash does not necessarily require a low-level gamma spectrometer, because a significant input to the pulse height spectrum originates from the Compton spectrum of K-40. This input may increase the background of the sample compared to the background checks by a factor of three. Generally, the specific activity concentration of K-40 in fish flesh is around 110 Bq·kg⁻¹ FM and in whole fish around 80 Bq·kg⁻¹ FM.

Cylindrical polyvinylchloride (PVC) beakers with plane bottom are used for the analysis.

Note

Compared to other plastic materials, PVC has got the advantage that the detection efficiency for X-ray energies is reduced due to the higher absorption of smaller energies. This may influence the summing of coincidences (see section 4.2.1.2 and basic chapter γ -SPEKT/GRUNDL of this procedures manual).

The measurement is normally carried out over night. The measuring period may be enlarged, depending on the scope of the analysis or the required detection limit (see section 5.2).

A correction of coincidence summing effects according to section 4.2.1.2 in γ -SPEKT/GRUNDL has to be applied, if nuclides with the emission of multiple gamma-ray energies are to be determined; see also (6, 7, 8).

5 Calculation of the results

5.1 Equations

When determining specific activities of radionuclides in ash from fish flesh, interferences due to gamma lines of different radionuclides are scarcely occurring. Therefore, specific activities of radionuclides may be calculated either from a single gamma line or, in the case of multiple-line radionuclides, by using a weighted mean.

If interferences occur, reference is made to the basic chapter γ -SPEKT/INTERF of this procedures manual.

5.1.1 Equations for calculating the specific activity from a single gamma line

If a net count rate $R_{n,r}$ at the gamma-ray energy of the radionuclide r has been detected, the resulting specific activity a_r , relative to fresh mass (FM) and the date and time of sampling, is calculated according to equation (5):

$$a_{r} = \varphi \cdot R_{n,r} = \frac{f_{1} \cdot f_{3} \cdot e^{\lambda_{r} \cdot t_{A}}}{\varepsilon_{A} \cdot p_{\gamma} \cdot m_{A} \cdot q_{F}} \cdot R_{n,r}$$
(5)

with:
$$f_3 = \frac{\lambda_r \cdot t_m}{1 - e^{-\lambda_r \cdot t_m}}$$

Herein are:

 $R_{n,r}$ net count rate of the line of the radionuclide r in s⁻¹;

 φ procedural calibration factor in Bq·s·kg⁻¹;

 f_1 correction factor for coincidence summing;

 f_3 correction factor for the decay of the activity of the radionuclide r during the measurement;

 ε_A detection efficiency in ash in Bq⁻¹·s⁻¹ (see section 4.2.3);

 p_{γ} emission probability of the gamma line of the radionuclide r;

 m_A mass of the ash used for the measurement in kg;

 $q_{\rm F}$ ratio of fresh mass to ash mass;

 t_A time period between sampling and beginning of the measurement in s;

 $t_{\rm m}$ duration of measurement in s;

 λ_r decay constant of the radionuclide r in s⁻¹.

The net count rate of the gamma line of the radionuclide r is calculated according to equation (6):

$$R_{n,r} = R_{b,r} - R_{T,r} - R_{0,r}$$
 (6)

Herein are:

 $R_{\rm b,r}$ gross count rate of the line of the radionuclide r in s⁻¹;

 $R_{T,r}$ background count rate underneath the line of the radionuclide r, e. g. as count rate of a trapezoidal background, in s^{-1} ;

 $R_{0,r}$ net count rate at the line of the radionuclide r in a background spectrum in s⁻¹.

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The generalized expression to calculate the uncertainty of the net count rate is given in equation (7), with the coefficients μ_k are to be determined according to equation (8):

$$u^{2}(R_{n,r}) = \mu_{0} \cdot R_{n,r}^{2} + \mu_{1} \cdot R_{n,r} + \mu_{2}$$
(7)

with coefficients:

$$\mu_0 = 0$$
; $\mu_1 = \frac{1}{t_m}$; $\mu_2 = \frac{R_{T,r} + R_{0,r}}{t_m} + u^2(R_{T,r}) + u^2(R_{0,r})$ (8)

Using the trapezoidal method (assuming a linear background continuum) the coefficient μ_2 is given by equation (9):

$$\mu_{2} = \frac{R_{T,r}}{t_{m}} \cdot \left(1 + \frac{b}{2L}\right) + R_{0,r} \cdot \left(\frac{1}{t_{m}} + \frac{1}{t_{0}}\right) + \frac{R_{T,0,r}}{t_{0}} \cdot \left(1 + \frac{b_{0}}{2L_{0}}\right)$$
(9)

Herein are:

- t_0 duration of the background measurement in s;
- b, b_0 line widths of sample and background spectra at the respective peak baselines in channels;
- L, L_0 numbers of channels for the sample and for the background spectrum, respectively, over which the background continuum to the left and to the right of the peaks are estimated;
- $R_{T,0,r}$ background continuum count rate at the line of the radionuclide r within the background spectrum, e. g. as a trapezoidal background count rate in s^{-1} .

Equation (9) is also a sufficient approximation for a background estimated by an empirical background step function.

The standard uncertainty of the net count rate $u(R_{n,r})$ calculated by equations (7) to (9) is given by:

$$u^{2}(R_{r}) = \frac{R_{r}}{t_{m}} + \mu_{2} = \frac{R_{r}}{t_{m}} + \frac{R_{T,r}}{t_{m}} \cdot \left(1 + \frac{b}{2L}\right) + R_{0,r} \cdot \left(\frac{1}{t_{m}} + \frac{1}{t_{0}}\right) + \frac{R_{T,0,r}}{t_{0}} \cdot \left(1 + \frac{b_{0}}{2L_{0}}\right)$$
(10)

When no count rate occurs at corresponding gamma-ray energy of background spectrum equation (10) is reduced by the last two terms.

The relative standard uncertainty of the procedural calibration factor is determined according to equation (11), where the uncertainty of the decay correction may be neglected:

$$u_{\text{rel}}(\varphi) = \sqrt{u_{\text{rel}}^{2}(f_{1}) + u_{\text{rel}}^{2}(f_{3}) + u_{\text{rel}}^{2}(\varepsilon_{A}) + u_{\text{rel}}^{2}(p_{\gamma}) + u_{\text{rel}}^{2}(m_{A}) + u_{\text{rel}}^{2}(q_{F})}$$
(11)

The combined standard uncertainty of the specific activity a_r using equation (5) is calculated as:

$$u(a_{\rm r}) = \varphi \cdot R_{\rm r} \cdot \sqrt{u_{\rm rel}^2(\varphi) + u_{\rm rel}^2(R_{\rm r})}$$
 (12)

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5.1.2 Equations for calculating the specific activity from multiple gamma lines

The specific activity a_r relative to fresh mass (FM) and the time of sampling of a number of j intensities (j greater than 1) at gamma-ray energies of radionuclide r sufficiently detected are calculated according to equation (13):

$$a_{r} = \varphi_{M} \cdot A_{r} = \frac{e^{\lambda_{r} \cdot t_{A}} \cdot f_{3}}{m_{A} \cdot q_{F}} \cdot A_{r}$$
(13)

Herein, A_r represents the activity (Bq) at the time of measurement, which is calculated as a weighted mean from the intensity of a number of single gamma-ray energies of nuclides,

$$A_{r} = u^{2}(A_{r}) \cdot \sum_{j} \frac{A_{j}}{u^{2}(A_{j})}$$
 (14)

wherein the standard uncertainty $u(A_r)$ is given by equation (15):

$$u^{2}(A_{r}) = \frac{1}{\sum_{j} \frac{1}{u^{2}(A_{j})}}$$
 (15)

The index j in these equations identifies single gamma-ray energies. The activities A_i of the individual gamma-ray energies are calculated according to equation (5):

$$A_{j} = \frac{R_{n,j} \cdot f_{1,j}}{\varepsilon_{A,j} \cdot p_{\gamma,j}} = R_{n,j} \cdot \varphi_{j}$$
(16)

Herein are:

 φ_{M} procedural calibration factor in kg⁻¹;

 φ_i procedural calibration factors in Bq·s;

 $R_{n,j}$ net count rate of the gamma-ray energy j in s⁻¹;

 $\varepsilon_{A,j}$ detection efficiency of the gamma-ray energy j in ash in Bq⁻¹·s⁻¹ (see section 4.2.3);

 $p_{\gamma,j}$ emission probability of the gamma-ray energy j;

 $f_{1,j}$ coincidence summing correction factor for the gamma-ray energy j;

 $f_{2,j}$ self-attenuation correction factor of the gamma-ray energy j, for ash relative to water solution:

$$f_2 = \frac{\varepsilon_{MC} (H_2 O)}{\varepsilon_{MC} (Asche)};$$

The variances of A_i are calculated according to the following equation:

$$u^{2}(A_{i}) = u^{2}(R_{n,i}) \cdot \varphi_{i}^{2} + R_{n,i}^{2} \cdot u^{2}(\varphi_{i})$$
(17)

When the net count rates are calculated using the trapezoidal background procedure, their standard uncertainties are calculated according to equation (10). If peak fitting is used instead, that factor (1+b/(2L)), representing the trapezoidal background procedure, changes into a factor $f_{\rm B}$, which depends on the method used for peak fitting:

$$u^{2}(R_{n,j}) = \frac{R_{n,j}}{t_{m}} + \mu_{2,j}$$
 (18)

$$\mu_{2,j} = \frac{R_{T,j}}{t_{m}} \cdot f_{B} + R_{0,r} \cdot \left(\frac{1}{t_{m}} + \frac{1}{t_{0}}\right) + \frac{R_{T,0,r}}{t_{0}} \cdot \left(1 + \frac{b_{0}}{2L_{0}}\right)$$
(19)

The factor f_B generally depends on the averaged background continuum per channel underneath the gamma-ray energy and on the ratio

$$\frac{R_{n,j} \cdot t_{m}}{\sqrt{R_{T,j} \cdot t_{m}}}$$

For net count rates close to the detection limit or the decision threshold, f_B may be approximated by a fixed value; see the calculation example in section 5.2.2.

Finally, the standard uncertainty of the specific activity is calculated as follows:

$$u(a_{\rm r}) = \varphi_{\rm M} \cdot A_{\rm r} \cdot \sqrt{u_{\rm rel}^2(\varphi_{\rm M}) + u_{\rm rel}^2(A_{\rm r})}$$
 (20)

with

$$u_{\text{rel}}(\varphi_{M}) = \sqrt{u_{\text{rel}}^{2}(m_{A}) + u_{\text{rel}}^{2}(q_{F})}$$
 (21)

Note

The calculations presented in this section are laborious if carried out manually; therefore, the software UncertRadio has been used to calculate the parameters of the example (see section 5.2.2). Such software is required for the calculation of the decision threshold and the detection limit by linear unfolding, which uses iteration, and therewith differs from the example shown in section 6.1.

5.2 Worked examples

5.2.1 Determining the specific Cs-137 activity

The following values are used for calculating the specific Cs-137 activity in 1,1 kg fish flesh (FM):

$$m_{A} = 0.01317 \text{ kg};$$
 $q_{F} = 83.54;$ $R_{n,Cs-137} = 0.00632 \text{ s}^{-1};$ $\varepsilon = 0.03056 \text{ Bq}^{-1} \cdot \text{s}^{-1};$ $R_{T,Cs-137} = 0.00271 \text{ s}^{-1};$ $b/(2L) = 0.523;$ $p_{\gamma} = 0.8499;$ $f_{1} = 1.000;$ $u_{rel}(m_{A}) = 0.004;$ $u_{rel}(q_{F}) = 0.02;$ $u(R_{n,Cs-137}) = 0.000381 \text{ s}^{-1};$ $u_{rel}(\varepsilon) = 0.029;$ $u_{rel}(p_{\gamma}) = 0.00235;$ $u_{rel}(f_{1}) = 0;$ $u_{rel}(f_{5}) = 0.008.$

The standard uncertainties of the following input quantities are negligible:

$$t_A = 1,9788 \cdot 10^7 \text{ s};$$
 $t_{Cs-137} = 9,483 \cdot 10^8 \text{ s};$ $t_m = 72000 \text{ s}.$

This leads to the correction factor f_3 for the decay during the measurement duration:

$$f_3 = \frac{7,30937 \cdot 10^{-10} \cdot 72000}{1 - e^{-7,30937 \cdot 10^{-10} \cdot 72000}} = 1,00003$$

The detection efficiency in ash ε_A is calculated according to equation (4):

$$\varepsilon_{A} = \frac{0,03056}{0,976} \text{ Bq}^{-1} \cdot \text{s}^{-1} = 0,03131 \text{ Bq}^{-1} \cdot \text{s}^{-1}$$

The procedural calibration factor according to equation (4) is obtained from:

$$\varphi = \frac{1,0 \cdot 1,00003}{0,03131 \cdot 0,8499 \cdot 0,01317 \cdot 83,54} \cdot e^{7,30937 \cdot 10^{-10} \cdot 1,9788 \cdot 10^{-7}} \text{ Bq} \cdot \text{s} \cdot \text{kg}^{-1} = 34,655 \text{ Bq} \cdot \text{s} \cdot \text{kg}^{-1}$$

The specific activity in the fish flesh relative to the fresh mass (FM) results from equation (5):

$$a_{\text{Cs-}137} = 34,655 \cdot 0,00632 \;\; \text{Bq\cdot kg}^{-1} = 0,219 \;\; \text{Bq\cdot kg}^{-1}$$

The relative standard uncertainty of the procedural calibration factor is calculated by equation (11):

$$u_{\rm rel}(\varphi) = \sqrt{0.008^2 + 0.029^2 + 0.00235^2 + 0.004^2 + 0.02^2} = 0.03642$$

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The standard uncertainty of the net count rate is obtained by equation (10) with neglecting the last two terms:

$$u(R_{\text{Cs-}137}) = \sqrt{\frac{0,00632}{72000} + \frac{0,00271}{72000} \cdot (1 + 0,523)} \cdot s^{-1} = 3,809 \cdot 10^{-4} \text{ s}^{-1}$$

The relative standard uncertainty of the specific activity is calculated according to equation (12):

$$u_{\text{rel}}(a_{\text{Cs-}137}) = \sqrt{0.03642^{-2} + \left(\frac{0.0003809}{0.00632}\right)^2} = 0.0704$$

Finally, the specific Cs-137 activity is obtained as:

$$a_{\text{Cs-}137} = (0.219 \pm 0.015) \text{ Bq} \cdot \text{kg}^{-1} \text{ (FM)}$$

5.2.2 Determining the specific Cs-134 activity

For calculating the specific Cs-134 activity, the two main gamma energies at 604,7 keV and 795,9 keV were evaluated in a sample 3,7 kg (FM) of flesh from North Atlantic cod off Greenland, whose ash was measured over a period of about 15 days on a Germanium detector with relative efficiency of 35 %. The numerical values used for the calculations are the following:

$$m_A = 0,05121 \text{ kg};$$
 $q_F = 72,4;$ $u_{rel}(m_A) = 0,004;$ $u_{rel}(q_F) = 0,02.$

The values for calculating the activities of the two Cs-134 gamma lines are:

Gamma line at 604,7 keV:

$$R_{\text{n,1}} = 4,7955 \cdot 10^{-4} \text{ s}^{-1};$$
 $\varepsilon_1 = 0,02071 \text{ Bq}^{-1} \text{ s}^{-1};$ $R_{\text{T,1}} = 4,9727 \cdot 10^{-3} \text{ s}^{-1};$ $p_{\gamma,1} = 0,9763;$ $f_{1,1} = 1,1332;$ $f_{5,1} = 0,9243;$ $u_{\text{rel}}(\varepsilon_1) = 0,01950;$ $u_{\text{rel}}(f_{1,1}) = 0,01209;$ $u_{\text{rel}}(f_{5,1}) = 0,01.$

Gamma line at 795,9 keV:

$$R_{\text{n,2}} = 3,835 \cdot 10^{-4} \text{ s}^{-1};$$
 $\varepsilon_2 = 0,01638 \text{ Bq}^{-1} \text{ s}^{-1};$ $R_{\text{T,2}} = 4,9608 \cdot 10^{-3} \text{ s}^{-1};$ $p_{\gamma,2} = 0,854;$ $f_{1,2} = 1,1368;$ $f_{5,2} = 0,932;$ $u_{\text{rel}}(\varepsilon_2) = 0,01632;$ $u_{\text{rel}}(f_{\gamma,2}) = 0,00105;$ $u_{\text{rel}}(f_{5,2}) = 0,01415;$ $u_{\text{rel}}(f_{5,2}) = 0,01.$

The standard uncertainties of the following input quantities are negligible:

$$t_{A} = 1,365 \cdot 10^{7} \text{ s};$$
 $t_{Cs-134} = 6,517 \cdot 10^{7} \text{ s};$ $t_{m} = 1,314 \cdot 10^{6} \text{ s}.$

In this example, a value of 1,08 is attributed to the factor f_B . For the factor φ_M from equation (13) a value $\varphi_M = 0,31406 \text{ kg}^{-1}$ is obtained with $e^{\lambda_r \cdot t_A} \cdot f_3 = 1,1644$; its relative standard uncertainty according to equation (21) is obtained as 0,020396 kg⁻¹.

Values $\varphi_1 = 51,7973$ Bq s (604,7 keV) and $\varphi_2 = 75,6845$ Bq s (795,9 keV) are obtained for the φ_j in equation (16). By multiplying values of $R_{n,i}$ and φ_j the activities associated with the two lines are calculated as $A_1 = 0,02484$ Bq and $A_2 = 0,02903$ Bq.

The uncertainties of the two net count rates calculated by equations (18, 19) are $u(R_{n,1}) = 6,6712 \cdot 10^{-5} \text{ s}^{-1}$ and $u(R_{n,2}) = 6,6077 \cdot 10^{-5} \text{ s}^{-1}$, respectively. The squared relative uncertainties of the φ_i values become:

$$u_{\text{rel}}^{2}(\varphi_{1}) = (0.01209^{-2} + 0.01^{-2} + 0.01950^{-2} + 0.00082^{-2}) \text{Bq}^{2} \cdot \text{s}^{2} =$$

$$= 6.2709 \cdot 10^{-4} \text{ Bq}^{2} \cdot \text{s}^{2}$$

$$u_{\text{rel}}^{2}(\varphi_{2}) = (0.01415^{-2} + 0.01^{-2} + 0.01632^{-2} + 0.00105^{-2}) \text{Bq}^{2} \cdot \text{s}^{2} =$$

$$= 5.6767 \cdot 10^{-4} \text{ Bq}^{2} \cdot \text{s}^{2}$$

The variances of the activity values of the two lines are calculated according to equation (16):

$$u^{2}(A_{1}) = (6,6712 \cdot 51,7873)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (6,2709 \cdot 10^{-4} \cdot 51,7873)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (6,2709 \cdot 10^{-4} \cdot 51,7873)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (6,2709 \cdot 10^{-4} \cdot 51,7873)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (6,2709 \cdot 10^{-4} \cdot 51,7873)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (6,2709 \cdot 10^{-4} \cdot 51,7873)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (6,2709 \cdot 10^{-4} \cdot 51,7873)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (6,2709 \cdot 10^{-4} \cdot 51,7873)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (6,2709 \cdot 10^{-4} \cdot 51,7873)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (6,2709 \cdot 10^{-4} \cdot 51,7873)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (6,2709 \cdot 10^{-4} \cdot 51,7873)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (6,2709 \cdot 10^{-4} \cdot 51,7873)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (5,6767 \cdot 10^{-4} \cdot 75,6845)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (5,6767 \cdot 10^{-4} \cdot 75,6845)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (5,6767 \cdot 10^{-4} \cdot 75,6845)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (5,6767 \cdot 10^{-4} \cdot 75,6845)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (5,6767 \cdot 10^{-4} \cdot 75,6845)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (5,6767 \cdot 10^{-4} \cdot 75,6845)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (5,6767 \cdot 10^{-4} \cdot 75,6845)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (5,6767 \cdot 10^{-4} \cdot 75,6845)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (5,6767 \cdot 10^{-4} \cdot 75,6845)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (5,6767 \cdot 10^{-4} \cdot 75,6845)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (5,6767 \cdot 10^{-4} \cdot 75,6845)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (5,6767 \cdot 10^{-4} \cdot 75,6845)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (5,6767 \cdot 10^{-4} \cdot 75,6845)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (5,6767 \cdot 10^{-4} \cdot 75,6845)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (5,6767 \cdot 10^{-4} \cdot 75,6845)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (5,6767 \cdot 10^{-4} \cdot 75,6845)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (5,6767 \cdot 10^{-4} \cdot 75,6845)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (5,6767 \cdot 10^{-4} \cdot 75,6845)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (5,6767 \cdot 10^{-4} \cdot 75,6845)^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (5,6767 \cdot 10^{-4} \cdot 10^{-4})^{2} + (4,7955 \cdot 10^{-4})^{2} \cdot (5,6767 \cdot 10^{-4})^{2} + (4,7955 \cdot 10^{-4})^{2} + (4,7955 \cdot 10^{-4})^{2} + (4,7955 \cdot 10^{-4})^{2} + (4,7955 \cdot 10^{-4})^$$

First, the variance of the Cs-134 activity is calculated according to equation (14):

$$u^{2}(A_{Cs-134}) = \frac{1}{\frac{1}{1,23226 \cdot 10^{-5}} + \frac{1}{2,54883 \cdot 10^{-5}}} Bq^{2} = \frac{1}{81151,7 + 39233,7} Bq^{2} = \frac{1}{81151,7$$

Afterwards, the activity of Cs-134 is obtained using equation (13):

$$A_{\text{Cs-}134} = u^{2} (A_{\text{Cs-}134}) \cdot \sum_{j} \frac{A_{j}}{u^{2} (A_{j})}$$

$$A_{\text{Cs-}134} = 8,30666 \cdot 10^{-6} \cdot \left(\frac{0,02484}{1,23226 \cdot 10^{-5}} + \frac{0,02903}{2,54883 \cdot 10^{-5}} \right) \text{Bq} =$$

$$= 8,30666 \cdot 10^{-6} \cdot (2015,808 + 1138,95) \text{Bq} = 0,0262 \text{ Bq}$$

Finally, the value and the associated standard uncertainty of the specific Cs-134 activity are determined according to equations (12, 20):

$$a_{\text{Cs-}134} = 0.31404 \cdot 0.0262 \text{ Bq} \cdot \text{kg}^{-1} = 8.2278 \cdot 10^{-3} \text{ Bq} \cdot \text{kg}^{-1}$$

$$u(a_{\text{Cs-}134}) = 8.2278 \cdot 10^{-3} \cdot \sqrt{0.020396}^{2} + \left(\frac{2.882 \cdot 10^{-3}}{0.0262}\right)^{2} \text{ Bq} \cdot \text{kg}^{-1} = 9.205 \cdot 10^{-4} \text{ Bq} \cdot \text{kg}^{-1}$$

5.3 Consideration of uncertainties

The combined standard uncertainty of the specific activity of Cs-137 amounts to be less than 10 % in the IMIS routine measurement program, where the largest contribution originates from the net count rate. If the self-attenuation correction for ash from fish flesh is neglected, another systematic error of up to 5 % will be the result.

For multi-line radionuclides such as Cs-134, the uncertainty contribution of the coincidence summing correction factor (see basic chapters γ -SPEKT/GRUNDL and γ -SPEKT/SUMESC of this procedures manual) has to be taken into account, which may lead to additional few per cents in case of significant corrections.

6 Characteristic limits of the procedure

The calculations of decision threshold and detection limit follow the standard ISO 11929 (9). For single-line emitters (see section 5.1.1) explicit equations for calculating decision threshold and detection limit can be applied. For this purpose an Excel file (see section 7.3.1) as well as a project file for the program UncertRadio (see section 7.3.2) are available on the website of this procedures manual.

For multi-line emitters (see section 5.1.2) these equations can only be calculated with software support. At present, a corresponding project file for the program UncertRadio (see section 7.3.2) is available on the website of this procedures manual.

6.1 Equations

6.1.1 Equations for calculatation of the detection limit of the specific Cs-137 activity

Before calculating the detection limit $a_r^{\#}$ the decision threshold of the radionuclide's specific activity a_r^{*} is determined using equation (22):

$$a_{r}^{*} = \varphi \cdot k_{1-\alpha} \cdot \sqrt{\mu_{2}} = \varphi \cdot k_{1-\alpha} \cdot \sqrt{\frac{1}{t_{m}} \cdot (R_{T,r} + R_{0,r}) + u^{2}(R_{T,r}) + u^{2}(R_{0,r})}$$
 (22)

This allows the detection limit $a_r^{\#}$ to be calculated by equation (23):

$$a_{r}^{\#} = \frac{a_{r}^{*} \cdot \psi}{\theta} \left\{ 1 + \sqrt{1 - \frac{\theta}{\psi^{2}} \left(1 - \frac{k_{1-\beta}^{2}}{k_{1-\alpha}^{2}} \right)} \right\}$$
 (23)

with auxiliary quantities:

$$\theta = 1 - k_{1-\beta}^2 \cdot \left(u_{\text{rel}}^2(\varphi) \right)$$

$$\psi = 1 + \frac{k_{1-\beta}^2}{2 \cdot a_r^*} \cdot \left(\frac{\varphi}{t_m}\right)$$

Herein are:

 $k_{1-\alpha}$ quantile of the normal distribution associated with the error of first kind;

 $k_{1-\beta}$ quantile of the normal distribution associated with the error of second kind.

6.1.2 Equations of calculation of the detection limit of the specific Cs-134 activity

While the decision threshold for multi-line emitters may be calculated directly, the subsequent derivation of the detection limit requires iteration. Details are given in the basic chapter EKG/NWG-ISO-01 of this procedures manual.

The decision threshold of the specific Cs-134 activity is obtained by equation (24):

$$a_{r}^{*} = k_{1-\alpha} \cdot \varphi_{M} \cdot u(A_{r} = 0) = k_{1-\alpha} \cdot \varphi_{M} \cdot \sqrt{\frac{1}{\sum_{j} \frac{1}{\varphi_{j}^{2} \cdot \frac{R_{T,j}}{t_{m}}} f_{B}}}$$
(24)

The detection limit of the specific Cs-134 activity is estimated according to equation (25).

$$a_{r}^{\#} \approx a_{r}^{*} + k_{1-\beta} \cdot u(a_{r}^{\#'})$$
 (25)

with an iterated standard deviation $u(a^{\#}_{r})$:

$$u\left(a_{r}^{\#}\right) = \sqrt{\left(\frac{a_{r}^{*}}{a_{r}^{*}}\right)^{2} + \left[u^{2}(a_{r}) - \left(\frac{a_{r}^{*}}{k_{1-\alpha}}\right)^{2}\right] \cdot \frac{a_{r}^{\#}}{a_{r}}}$$

6.2 Worked examples

6.2.1 Decision threshold and detection limit of the specific Cs-137 activity

Using the values of the input quantities from section 5.2.1 the coefficient μ_2 is derived according to equation (9):

$$\mu_2 = \frac{0,00271}{72000} \cdot (1 + 0,523) \text{ s}^{-2} = 5,732 \cdot 10^{-8} \text{ s}^{-2}$$

Using this value, the decision threshold of the specific activity $a_{\text{Cs-137}}^*$ is obtained from the procedural calibration factor φ and the normal distribution quantile $k_{1-\alpha}=3$:

$$a_{\text{Cs}-137}^* = 34,65 \cdot 3,0 \cdot \sqrt{5,732 \cdot 10^{-8}} \text{ Bq } \cdot \text{kg}^{-1} = 0,0249 \text{ Bq } \cdot \text{kg}^{-1}$$

According to equation (14) the detection limit of the specific activity $a_{Cs-137}^{\#}$ is:

$$a_{\text{Cs-137}}^{\#} = \frac{0,0249 \cdot 1,0262}{0,9962} \cdot \left\{ 1 + \sqrt{1 - \frac{0,9962}{1,0262}} \cdot \left(1 - \frac{1,645^{2}}{3,0^{2}} \right) \right\} \text{ Bq } \cdot \text{kg}^{-1} = 0,02565 \cdot \left\{ 1 + \sqrt{1 - 0,9460 \cdot (1 - 0,30067)} \right\} \text{Bq } \cdot \text{kg}^{-1} = 0,0406 \text{ Bq } \cdot \text{kg}^{-1}$$

with values of the auxiliary quantities

$$\theta = 1 - 1,645^{2} \cdot (0,0374^{2}) = 0,9962$$

$$\psi = 1 + \frac{1,645^2}{2 \cdot 0.0249} \cdot \left(\frac{34,65}{72000}\right) = 1,0262$$

6.2.2 Decision threshold and detection limit of the specific Cs-134 activity

First, the following auxiliary quantity from equation (24) is calculated using the values of quantities from section 5.2.2 and the normal distribution quantile, $k_{1-\alpha} = 3$:

$$\sum_{j} \frac{1}{\varphi_{j}^{2} \cdot \frac{R_{T, j}}{t_{m}}} = \dots = 1,3401 \cdot 10^{5} \text{ Bq}^{-2}$$

Therewith, the decision threshold of the specific Cs-134 activity amounts to:

$$a_{\text{Cs-}134}^{*} = 3.0 \cdot 0.31404 \cdot \sqrt{\frac{1}{1.3401 \cdot 10^{5}}} \text{ Bq } \cdot \text{kg}^{-1} = 2.57 \cdot 10^{-3} \text{ Bq } \cdot \text{kg}^{-1}$$

This decision threshold value agrees quite well with that value obtained from using UncertRadio (see section 7.3.2). It is smaller compared to the two single-line decision threshold values, which are $3,12 \text{ mBq} \cdot \text{kg}^{-1}$ and $4,55 \text{ mBq} \cdot \text{kg}^{-1}$, respectively.

The detection limit of the specific Cs-134 activity is estimated with a first step of iteration according to equation (25) as follows:

$$u(a_{r}^{\#}) = \sqrt{\left(\frac{2,5736 \cdot 10^{-3}}{3,0}\right)^{2} + \left[\left(9,21 \cdot 10^{-4}\right)^{2} - \left(\frac{2,5736 \cdot 10^{-3}}{3,0}\right)^{2}\right] \cdot \left(\frac{\frac{3+1,645}{3} \cdot 2,5736 \cdot 10^{-3}}{\frac{3}{8,2278 \cdot 10^{-3}}}\right)} \operatorname{Bq} \cdot \operatorname{kg}^{-1} = 8,888 \cdot 10^{-4} \operatorname{Bq} \cdot \operatorname{kg}^{-1}$$

$$= 8,888 \cdot 10^{-4} \operatorname{Bq} \cdot \operatorname{kg}^{-1}$$

$$a_{Cs-134}^{\#} \approx 2,5736 \cdot 10^{-3} + 1,645 \cdot 8,888 \cdot 10^{-4} \operatorname{Bq} \cdot \operatorname{kg}^{-1} = 4,036 \cdot 10^{-3} \operatorname{Bq} \cdot \operatorname{kg}^{-1}$$

This iteratively estimated detection limit is in good agreement with the result from UncertRadio, $4,031\cdot10^{-3}$ Bq·kg⁻¹ (see section 7.3.2.2).

Version January 2016

7 Catalogue of chemicals and equipment

7.1 Chemicals

Cleaning agent:
 e. g. RBS-50-Super-Flüssigkonzentrat 2 %.

7.2 Equipment

Ordinary equipment of a radiochemical laboratory.

7.2.1 Sampling

- ice container / cooling boxes;
- plastic bags;
- deep freezer (ca. -18 °C), if the samples have to be stored.

7.2.2 Sample preparation

- filleting board made of plastic;
- sharpened filleting knifes;
- cut-resistant gloves.

7.2.3 Drying/ashing

- incineration ovens with catalytical treatment of exhaust gases; organic exhaust gases shall largely be burned to CO₂ und H₂O;
- stainless steel bowls (V4A) or ceramic bowls adapted in size to the internal volume of the oven,
- transparent paper with an areal density of about 90 g·m⁻² as backing material for the stainless steel containers;

7.2.4 Calibration and measurment

- gamma mitting radionuclides, preferably single-line emitters; if applicable, within a single solution covering the whole energy range to be calibrated;
- cylindrical plastic containers, e. g. made from PVC, with an inner diameter of about 7 cm and a capacity of ca. 220 ml, preferably with plane bottom;
- High-purity Germanium-semiconductor detector with a relative efficiency preferably between 20 % and 60 % relative to a 3" x 3" NaI(Tl)-crystal and a full-width-at-half-maximum less than 2,0 keV for the 1,33 MeV gamma-line of Co-60;
- measuring electronics;
- lead shielding with ca. 10 cm wall thickness.

7.2.5 Evaluation

- software for accumulating and evaluating gamma-ray spectra;
- software tools for calculating the peak and total efficiencies and correction factors for coincidence summing and self-attenuation.

7.3 Program supported evaluation

7.3.1 View of the Excel sheet

7.3.1.1 Calculating the specific Cs-137 activity

	Procedure for the gamma spectromet							
	G-γ-SPEKT-FISCH-01						Version Ja	nuary 201
	Procedures manual for monitoring of	radioacti	ve substanc	es in the envir	onment and of	external radia	ation (ISSN 18	65-8725)
	Sample identification:				Variant 1: sing	le-line radion	uclide Cs-137	
	#Number of parameters p:	13		Colors:	values from VI	basic		
	k_alpha: quantile for (1-α)	3			Excel formulae (user)			
	k_beta: quantile for (1-β)	1,645			Manual input of values			
	gamma: probability for conf.limits	0,05			Def. Excel-Varia	ble (user)		
					#Keywords			
	Data input:	I India.	variable nar			Uncertainty b		la al ac a A
	#Values of parameters p:	Unit:		Input values	StdDov"	partial	uncertainty	budget in %
-1	#Niveshou of gross accounts Nive		NI-	CEO 40	StdDev:	derivatives	budget:	
01	#Number of gross counts Ng:		Ng	650,16		0,000481292		
02	trapezoidal BG count rate	1/s	RT	2,71000E-03		,	0,004861951	
03	b/2L		b2L	5,23000E-01	•			-,
o4 -	detection efficiency		eps	3,05600E-02	*		0,006351203	
05	emission probability, 661,7 keV		p_gamm	8,49900E-01			0,000514598	
o6	ash mass	kg	mA	0,01317000	•		0,000876028	
o7	ratio fresh mass / ash mass	_	qF	83,54000000		*		
08 -0	Time elapsed Sampling>StartMeasum	S	tA	1,97880E+07		1,6008E-10		0,00
o9	half-live of Cs-137	S	tCs137	9,48300E+08		-3,3466E-12	0,001752058	0,00
010	self-attenuation correction factor	_	_f2	0,97600000 72000,00		*	1	
	counting duration	s 1/s	tm RnNE	0,000E+00	•	-		0,00 0,00
o12	net count rate of the BG peak counting duration of the BG spectrum		tNE	72000,00			0	0,00
313	counting duration of the bo spectrum	3	LIVE	72000,00	0,000002100	O O		0,00
	(the list may be continued here)							
	Model section		c = Factor *	Rn				
	auxiliary equations:			(formulae)				
	#Gross counting rate Rg:	1/s	Rg	9,03000E-03				
h1	decay correction measuring period		_f3	9,99974E-01				
ո2	decay correction Sampling>StartMe	S	_f4	1,01457E+00				
	(the list may be continued here)							
	((formulae)				
	#Net counting rate Rn:	1/s	Rn	6,32000E-03				
	#Calibration factor, proc.dep.:	Bq*s/kg		34,65304141				
	#Value output quantity:	Bq/kg	Result	2,19007E-01	0,040542041	< output valu	e modifiable by	VBA
	#Combined standard uncertainty:	Bq/kg	uResult	1,54230E-02				
	#Desision threshold:	Da/Ira		0,024890348				
	#Decision threshold:	Bq/kg			Cale	culate!		
	#Detection limit:	Bq/kg		0,040542029				
	further derived quantities (ISO 11929): auxiliary quantity omega		Omogo	1,00000000				
	Best estimate	Bq/kg	Omega RostWort	2,19007E-01				
	Uncertainty of best estimate	Bq/kg Bq/kg	BestWert	2,19007E-01 1,54230E-02	•			
	lower confidence limit	Bq/kg Bq/kg		1,54230E-02 1,88779E-01				
	upper confidence limit	Bq/kg Bq/kg		2,49236E-01				

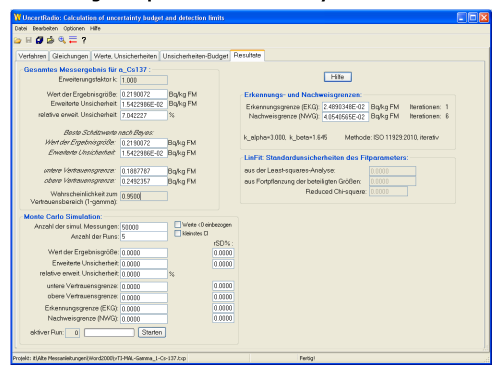
The Excel file can be found on the website of this procedures manual.

7.3.1.2 Calculating the specific Cs-134 activity

Because of the complexity of the calculations an Excel file is not available.

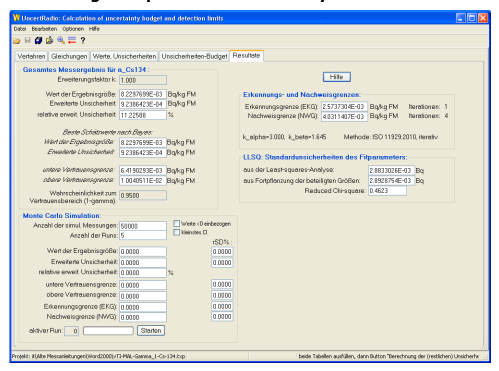
7.3.2 View of the result page of UncertRadio

7.3.2.1 Calculating the specific Cs-137 activity



The UncertRadio project file can be found on the website of this procedures manual.

7.3.2.2 Calculating the specific Cs-134 activity



The UncertRadio project file can be found on the website of this procedures manual.

Literature

- (1) Verordnung (EG) Nr. 2065/2001 der Kommission vom 22. Oktober 2001 mit Durchführungsbestimmungen zur Verordnung (EG) Nr. 104/2000 des Rates hinsichtlich der Verbraucherinformation bei Erzeugnissen der Fischerei und der Aquakultur. Amtsblatt der Europäischen Gemeinschaften 23.10.2001, L 278/6
- (2) Gilmore, G.: *Practical Gamma-Ray Spectrometry*. 2nd Edition. Chichester: John Wiley & Sons Ltd., 2008. ISBN 978-0-470-86196-7.
- (3) Knoll, G. F.: *Radiation detection and measurement*. 4th Edition. New York: John Wiley & Sons Inc., 2010. ISBN 978-0-470-13148-0.
- (4) Debertin, K., Helmer, R. G.: *Gamma- and X-Ray Spectrometry with Semicon-ductor Detectors*. Amsterdam: North Holland, 1988. ISBN 978-0-444-87107-7.
- (5) Berger, M. J., et al.: *XCOM: Photon Cross Sections Database*. NIST Standard Reference Database 8 (XGAM) [Online-Datenbank]. Letztes Update: November 2010 [Zugriff am: 03.03.2016]. Verfügbar unter: http://www.nist.gov/pml/data/xcom/index.cfm
- (6) Debertin, K., Schötzig, U.: Bedeutung von Summationskorrektionen bei der Gammastrahlen-Spektrometrie mit Germaniumdektoren. PTB-Ra-24. Braunschweig: Physikalisch-Technische Bundesanstalt, 1990. ISBN 3-89429-010-9.
- (7) Kanisch, G., Vidmar, T., Sima, O.: *Testing the equivalence of several algorithms for calculation of coincidence summing corrections*. Appl. Radiat. Isot., 2009, Vol. 67, S. 1952–1956.
- (8) Vidmar, T., Kanisch, G.: *Intrinsic error of the point-source and the LS-curve approximations in treatment of true coincidence summing*. Appl. Radiat. Isot., 2010, Vol. 68, S. 1785–1788.
- (9) ISO 11929:2010, Determination of the characteristic limits (decision threshold, detection limit and limits of the confidence interval) for measurements of ionizing radiation Fundamentals and application.
- (10) Souci, S. W., Fachmann, W., Kraut, H.: *Lebensmitteltabelle für die Praxis*. 5. Auflage. Stuttgart: WVG, 2011. ISBN 978-3-8047-2679-6.