

# Guide to the Realization of the ITS-90

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Partie 1

**Part 1**

## Platinum Resistance Thermometry

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Part 1: Platinum Resistance Thermometry**

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## Abstract

This paper is a part of guidelines, prepared on behalf of the Consultative Committee for Thermometry, on the methods how to realize the International Temperature Scale of 1990.

It discusses the major issues linked to platinum resistance thermometry for the realization of the International Temperature Scale of 1990 in the temperature range from 24.5561 K to 1234.93 K.

## 1. INTRODUCTION

In the temperature range from the triple point of equilibrium hydrogen (13.8033 K) to the freezing point of silver (1234.93 K), the ITS-90 is defined in terms of the temperature dependence of the electrical resistance of standard platinum resistance thermometers (SPRTs). This chapter provides mainly an overview of the practical realisation of the scale in the restricted range from the triple point of neon (24.5561 K) to the freezing point of silver, which does not require to realise the vapour-pressure points of equilibrium hydrogen in the vicinity of 17 K and 20.3 K or to apply an interpolating helium constant-volume gas thermometer. Furthermore, it includes an explanation of the scale definition, describes the different types of SPRTs used, gives advice on the use and calibration of SPRTs, and concludes with a brief summary of the sources of uncertainty. The chapter omits discussion of the fixed points used to calibrate SPRTs, as this is covered in [Chapter 2 Fixed Points](#) (<http://www.bipm.org/en/committees/cc/cct/guide-its90.html>) as well as in [Mangum *et al.* 1999 and 2000]. For more detail, the reader is referred to the references given at the end of the chapter and the [Technical Annex](#) of the *mise en pratique* of the definition of the kelvin (*MeP-K*) [Ripple *et al.* 2010, Fellmuth *et al.* 2016, [http://www.bipm.org/en/publications/mep\\\_kelvin/](http://www.bipm.org/en/publications/mep\_kelvin/)], which includes supplement-ary definitions and clarifications.

## 2. THE SPRT DEFINITIONS IN ITS-90

### 2.1. Overview of the scale definition

The ITS-90 specifies a set of fixed points (melting, freezing, triple, or boiling points of various pure substances, see *Guide* Chapter 2 *Fixed points*), which are used for the calibration of SPRTs in eleven temperature sub-ranges within the overall range from 13.8033 K to 1234.93 K. All the sub-ranges include the triple point of water (TPW) and extend to progressively higher or lower temperatures. The existence of overlapping sub-ranges implies the existence of a variety of temperature values  $T_{90}$  according to the ITS-90 for a given resistance value for a given SPRT (Type 1 non-uniqueness (sub-range inconsistency) [Mangum *et al.* 1997]), see Section 6.3.1. Mathematical procedures are specified for relating the measured resistances of the SPRTs to the corresponding ITS-90 temperatures.

SPRT resistances are not used directly but are first normalised by taking ratios to the resistance at the TPW (273.16 K). Thus, the ITS-90 resistance ratio,  $W(T_{90})$ , is defined as

$$W(T_{90}) = R(T_{90})/R(273.16 \text{ K}), \quad (1)$$

where  $R(T_{90})$  and  $R(273.16 \text{ K})$  are the resistances of the SPRT at the temperature  $T_{90}$  and at the TPW, respectively. Taking resistance ratios removes the need for traceability to absolute resistance standards in the calibration and use of an SPRT. The  $W$  values embody that SPRT characteristics are fundamentally similar, which allows the application of reference functions. Furthermore, the normalisation provides some compensation for instability, though this is not something the ITS-90 considers. The compensation is perfect for temperature-independent relative resistance changes caused for instance by dimensional changes in the platinum wire or by oxidation discussed in Section 5.1.1.

SPRTs must be made with strain-free wires of high purity platinum, so that the ratio  $W(T_{90})$ , which is essentially the ratio of the corresponding resistivities of the platinum wire, is very similar for all SPRTs. This is needed so that different SPRTs have similar interpolation properties; that is, they generate very similar realisations of the ITS-90. The purer and more strain-free the wire is, the higher is the temperature coefficient of resistance. Therefore, to ensure the suitability of an SPRT, the ITS-90 specifies criteria for the temperature coefficient, based on the resistance ratios at the melting point of gallium and at the triple point of mercury. The criteria are:

$$W(29.7646 \text{ }^{\circ}\text{C}) \geq 1.11807 \quad (2)$$

and

$$W(-38.8344 \text{ }^{\circ}\text{C}) \leq 0.844235. \quad (3)$$

These criteria are approximately equivalent and are used in sub-ranges above or below 273.16 K. They place lower limits on the temperature coefficient of the resistance of the platinum, and implicitly define the minimum purity requirements for the wire used in SPRTs.

In order to limit the effect of a possible breakdown of electrical insulation within High-Temperature SPRTs (HTSPRTs), the ITS-90 specifies an additional requirement for the resistance ratio at the freezing point of silver:

$$W(961.78 \text{ }^{\circ}\text{C}) \geq 4.2844. \quad (4)$$

Qualification criteria are discussed further in Section 4.4.2.

To interpolate between the values of  $W(T_{90})$  measured at a limited number of fixed points, the ITS-90 defines two reference functions (one above and one below the TPW), which relate the resistance ratio  $W_r(T_{90})$  of two chosen high-quality reference SPRTs, having high temperature coefficients of resistance, to  $T_{90}$ , see Section 2.2. The reference functions are continuous in their first and second derivatives throughout the whole range 13.8033 K to 1234.93 K.

The interpolation is done by comparing the values of  $W(T_{90})$  at fixed points with the corresponding values of  $W_r(T_{90})$  specified in the scale (Table 1 in the ITS-90 text). The differences (deviations)  $\Delta W = W(T_{90}) - W_r(T_{90})$  are mapped by low-order functions (mostly simple polynomials in  $(W-1)$  and/or  $\ln(W)$ , see Section 2.3.1, see p. 9), which then allow differences to be interpolated at any point in the sub-range. In each sub-range, different fixed points and interpolating equations are applied. For SPRTs using high-purity strain-free platinum, it follows that the  $W(T_{90})$  functions are very similar, and therefore that the deviations from the reference functions,  $W_r(T_{90})$ , are small. The deviation functions include the coefficients which are specific to the particular SPRT and whose values are determined from the calibration data  $\Delta W$  at the required fixed points.

Thus, the calibration of a particular SPRT has two components: the (relatively complicated) reference function,  $W_r(T_{90})$ , which is the same for all SPRTs, and the (relatively simple) deviation function,  $\Delta W$ , which contains the limited number of SPRT calibration coefficients derived from the fixed-point measurements.

As an example of a deviation equation, in the sub-range from the TPW to the freezing point of aluminium the specified equation is (omitting the qualifier  $T_{90}$ )

$$\Delta W \equiv (W - W_r) = a(W - 1) + b(W - 1)^2 + c(W - 1)^3. \quad (5)$$

The values of the coefficients  $a$ ,  $b$ , and  $c$  in Equation (5) are determined by requiring the equation to be satisfied at each of the defining fixed points for the sub-range, in this case the freezing points of tin, zinc, and aluminium. Hence:

$$W_{\text{Sn}} - W_{r,\text{Sn}} = a(W_{\text{Sn}} - 1) + b(W_{\text{Sn}} - 1)^2 + c(W_{\text{Sn}} - 1)^3, \quad (6)$$

$$W_{\text{Zn}} - W_{r,\text{Zn}} = a(W_{\text{Zn}} - 1) + b(W_{\text{Zn}} - 1)^2 + c(W_{\text{Zn}} - 1)^3,$$

$$W_{\text{Al}} - W_{r,\text{Al}} = a(W_{\text{Al}} - 1) + b(W_{\text{Al}} - 1)^2 + c(W_{\text{Al}} - 1)^3,$$

where  $W_{\text{Sn}}$ ,  $W_{\text{Zn}}$ , and  $W_{\text{Al}}$  are the measured values of resistance ratio at the indicated fixed points, and  $W_{r,\text{Sn}}$ ,  $W_{r,\text{Zn}}$  and  $W_{r,\text{Al}}$  are the values of reference resistance ratio assigned to the fixed points in the ITS-90. Since there are three equations, the three unknowns,  $a$ ,  $b$ , and  $c$ , are calculated exactly. The Equation (5) can now be used to calculate the deviation  $(W - W_r)$ , and hence  $W_r$ , for any input value of  $W$  within the sub-range.

The same procedure is followed in other sub-ranges, but an additional term is specified for an interpolation up to the silver point, and more complicated equations, involving terms in

$\ln(W)$ , are required for the sub-ranges below the TPW, see Section 2.3.1. Additionally, the sub-range from the triple point of mercury to the melting point of gallium extends above and below the TPW. Note that all of the ITS-90 interpolating equations are formulated so that at the TPW  $W_r = 1$ ,  $W = 1$  and  $\Delta W = 0$ .

Given the calibration coefficients for a particular SPRT in a particular sub-range, the calculation of  $T_{90}$  from a measured resistance can be seen as a three-step mapping,

$$R(T_{90}) \rightarrow W(T_{90}) \rightarrow W_r(T_{90}) \rightarrow T_{90}:$$

Step 1 *Calculate the resistance ratio  $W(T_{90}) = R(T_{90})/R(273.16 \text{ K})$ ,*

Step 2 *Use  $\Delta W(W)$  and the coefficients to calculate  $\Delta W$ , and hence  $W_r(T_{90}) = W(T_{90}) - \Delta W$ ,*

Step 3 *Use the reference function to calculate  $T_{90}$  from  $W_r(T_{90})$ .*

The first two steps are manipulations of the calibration data and interpolation involving  $W$ -values only; it is not until the Step 3 that temperature appears as a variable. In Step 3, calculating  $T_{90}$  from the reference function defining  $W_r(T_{90})$  requires iteration as it cannot be solved for  $T_{90}$  by direct substitution of values of  $W_r$ . However, as an alternative the ITS-90 specifies inverse functions  $T_{90}(W_r)$  which can be used with good accuracy, see Section 2.2.

## 2.2. Reference functions

Two separate reference functions are used in the SPRT sub-ranges of ITS-90, one for the range 13.8033 K to 273.16 K, and the other for 0 °C to 961.78 °C. In the range from 13.8033 K to 273.16 K the reference function is

$$W_r(T_{90}) = \exp \left( A_0 + \sum_{i=1}^{12} A_i \left[ \frac{\ln(T_{90}/273.16 \text{ K}) + 1.5}{1.5} \right]^i \right), \quad (7)$$

where the constants  $A_i$  are given in Table 1 below. This equation can be solved numerically to determine  $T_{90}$  from the value of  $W_r$  determined from the interpolation. Alternatively,  $T_{90}$  can be determined using the following inverse function, which is equivalent to Equation (7) within 0.1 mK:

$$T_{90} = 273.16 \text{ K} \left( B_0 + \sum_{i=1}^{15} B_i \left[ \frac{W_r(T_{90})^{1/6} - 0.65}{0.35} \right]^i \right). \quad (8)$$

The values of the  $B_i$  coefficients are given in Table 1 below. In the range from 273.16 K to 1234.93 K (0 °C to 961.78 °C), the reference function is defined by

$$W_r(T_{90}) = C_0 + \sum_{i=1}^9 C_i \left[ \frac{T_{90}/\text{K} - 754.15}{481} \right]^i. \quad (9)$$

An inverse function, equivalent to Equation (9) within 0.13 mK is

$$T_{90}/\text{K} = 273.15 + D_0 + \sum_{i=1}^9 D_i \left[ \frac{W_r(T_{90}) - 2.64}{1.64} \right]^i. \quad (10)$$

The coefficients  $A_i$ ,  $B_i$ ,  $C_i$ , and  $D_i$ , used in Equation (7) to Equation (10) are given below in Table 1.



**Table 1.** The constants  $A_i$ ,  $B_i$ ,  $C_i$  and  $D_i$  in the SPRT reference and inverse functions of ITS-90.

$i$	$A_i$	$B_i$	$C_i$	$D_i$
0	-2.135 347 29	0.183 324 722	2.781 572 54	439.932 854
1	3.183 247 20	0.240 975 303	1.646 509 16	472.418 020
2	-1.801 435 97	0.209 108 771	-0.137 143 90	37.684 494
3	0.717 272 04	0.190 439 972	-0.006 497 67	7.472 018
4	0.503 440 27	0.142 648 498	-0.002 344 44	2.920 828
5	-0.618 993 95	0.077 993 465	0.005 118 68	0.005 184
6	-0.053 323 22	0.012 475 611	0.001 879 82	-0.963 864
7	0.280 213 62	-0.032 267 127	-0.002 044 72	-0.188 732
8	0.107 152 24	-0.075 291 522	-0.000 461 22	0.191 203
9	-0.293 028 65	-0.056 470 670	0.000 457 24	0.049 025
10	0.044 598 72	0.076 201 285		
11	0.118 686 32	0.123 893 204		
12	-0.052 481 34	-0.029 201 193		
13		-0.091 173 542		
14		0.001 317 696		
15		0.026 025 526		

## 2.3. Interpolating equations

### 2.3.1. Interpolating equations prescribed in the ITS-90 text

For describing the temperature dependence of the resistance of an SPRT, it is necessary to interpolate between the deviations  $\Delta W = W_{T_{90}} - W_r(T_{90})$  obtained at the temperature fixed points. ITS-90 specifies eight different interpolating functions  $((W - W_r)$  in dependence of  $W$ ) to be used over the eleven SPRT sub-ranges. These are listed in Table 2, along with the fixed points used to calibrate the SPRT. Most are simple polynomials in  $(W - 1)$  and/or  $\ln(W)$ .

**Table 2.** The sub-ranges, deviation functions and calibration points for SPRTs used to define ITS-90.

SPRT Sub-range	Deviation function <sup>(a)</sup>	Fixed points
13.8033 K to 273.16 K	$a(W-1) + b(W-1)^2 + \sum_{i=1}^5 c_i [\ln(W)]^{2+i}$	e-H <sub>2</sub> , Ne, O <sub>2</sub> , Ar, Hg <sup>(b)</sup>
24.5561 K to 273.16 K	$a(W-1) + b(W-1)^2 + \sum_{i=1}^3 c_i [\ln(W)]^i$	e-H <sub>2</sub> , Ne, O <sub>2</sub> , Ar, Hg
54.3584 K to 273.16 K	$a(W-1) + b(W-1)^2 + d [\ln(W)]^2$	O <sub>2</sub> , Ar, Hg
83.8058 K to 273.16 K	$a(W-1) + b(W-1) \ln(W)$	Ar, Hg

−38.8344 °C to 29.7646 °C	$a(W-1) + b(W-1)^2$	Hg, Ga
0 °C to 29.7646 °C	$a(W-1)$	Ga
0 °C to 156.5985 °C	$a(W-1)$	In
0 °C to 231.928 °C	$a(W-1) + b(W-1)^2$	In, Sn
0 °C to 419.527 °C	$a(W-1) + b(W-1)^2$	Sn, Zn
0 °C to 660.323 °C	$a(W-1) + b(W-1)^2 + c(W-1)^3$	Sn, Zn, Al
0 °C to 961.78 °C	$a(W-1) + b(W-1)^2 + c(W-1)^3$ $+ d[W - W(660.323\text{ °C})^2]$	Sn, Zn, Al, Ag

- (a) the constants  $a, b$ , etc., take different numerical values in each sub-range, except that for the sub-range 0 °C to 961.78 °C the coefficients  $a, b, c$  are the same as used for the sub-range 0 °C to 660.323 °C. The coefficient  $d$  is determined from the measurement at the silver point.
- (b) For the sub-range 13.8033 K to 273.16 K, two of the calibration points are required to be in the vicinity of 17 K and 20.3 K and are determined using a hydrogen vapour-pressure thermometer or an interpolating helium constant-volume gas thermometer (see [Chapter 3 Vapour Pressure Scales and Pressure Measurements](#) and [Chapter 4 Gas Thermometry](#) for details).

### 2.3.2. Alternative interpolating equations for special applications

The ITS-90 interpolating equations (Table 2, see p. 9) can be written in many mathematically equivalent forms. Amongst the alternatives is a form, similar to Lagrange interpolation, that is useful for propagation-of-uncertainty calculations (Section 6.2, see p. 40), investigating the consistency of calibration data (Section 4.4.2, see p. 22), and understanding Type 1 non-uniqueness (Section 6.3.1, see p. 45) [White and Saunders 2007, White and Strouse 2009, White 2013]. It should be emphasised that the alternative equations are simply rearrangements of those prescribed in the text of the ITS-90; they give identical values in interpolations. Their solution is single-valued because the number of coefficients is equal to the number of equations. People being only interested in the calibration of SPRTs according to the ITS-90 can skip this alternative mathematics without any loss of information.

For the example treated in Section 2.1, the sub-range from the TPW to the freezing point of aluminium, the alternative interpolating equation is a sum of four functions each multiplied by one of the ITS-90 reference resistance ratios:

$$W_r(W) = W_{r,H_2O} f_{H_2O}(W) + W_{r,Sn} f_{Sn}(W) + W_{r,Zn} f_{Zn}(W) + W_{r,Al} f_{Al}(W), \quad (11)$$

where the four interpolating functions containing ratio differences both in the numerator and denominator are:

$$f_{H_2O}(W) = \frac{(W - W_{Sn})(W - W_{Zn})(W - W_{Al})}{(W_{H_2O} - W_{Sn})(W_{H_2O} - W_{Zn})(W_{H_2O} - W_{Al})}, \quad f_{Sn}(W) = \frac{(W - W_{H_2O})(W - W_{Zn})(W - W_{Al})}{(W_{Sn} - W_{H_2O})(W_{Sn} - W_{Zn})(W_{Sn} - W_{Al})}, \quad (12)$$

$$f_{Zn}(W) = \frac{(W - W_{H_2O})(W - W_{Sn})(W - W_{Al})}{(W_{Zn} - W_{H_2O})(W_{Zn} - W_{Sn})(W_{Zn} - W_{Al})}, \quad f_{Al}(W) = \frac{(W - W_{H_2O})(W - W_{Sn})(W - W_{Zn})}{(W_{Al} - W_{H_2O})(W_{Al} - W_{Sn})(W_{Al} - W_{Zn})},$$

Although these equations are not as compact as Equation (5), the interpolation is now expressed in terms of the interpolating functions Equation (12), which are actually the sensitivity coefficients for uncertainties in the fixed points (see Section 6.2, see p. 40). The functions have properties similar to a set of orthonormal basis functions, which simplify the manipulation of mathematical expressions for uncertainty. Note too that the interpolation Equation (11) now maps directly to  $W_r$  rather than the deviations  $W - W_r$ , and Equation (12) are rational functions of the various  $W$  values only (i.e., there are no  $W_r$  values). All

of the ITS-90 equations can be expressed in a similar form, see [Appendix 1](#) and for the most commonly used sub-ranges of the ITS-90, Table 4 in Section 6.2.

The fact that the alternative interpolation equations are in reality equal to those prescribed in the text of the ITS-90, i.e. that they contain the same terms, which are only arranged differently, can be illustrated by the following equations for the sub-range from the TPW to the melting point of gallium:

An alternative interpolation equation is

$$W_r(W) = f_{\text{H}_2\text{O}}(W) + W_{r,\text{Ga}} f_{\text{Ga}}(W), \quad (13)$$

with

$$f_{\text{H}_2\text{O}}(W) = \frac{(W - W_{\text{Ga}})}{(W_{\text{H}_2\text{O}} - W_{\text{Ga}})}, \quad f_{\text{Ga}}(W) = \frac{(W - W_{\text{H}_2\text{O}})}{(W_{\text{Ga}} - W_{\text{H}_2\text{O}})}. \quad (14)$$

Insertion of Equation (14) in Equation (13) yields with  $W_{\text{H}_2\text{O}} = 1$ :

$$W_r = \frac{(W_{r,\text{Ga}} - W_{\text{Ga}})}{(1 - W_{\text{Ga}})} + W \frac{(1 - W_{r,\text{Ga}})}{(1 - W_{\text{Ga}})}. \quad (15)$$

By comparing Equation (15) with the rearranged interpolation equation of the ITS-90,  $W_r = a + W(1 - a)$ , cf. Table 2, one obtains for the coefficient  $a$  the known expression

$$a = (W_{r,\text{Ga}} - W_{\text{Ga}})/(1 - W_{\text{Ga}}). \quad (16)$$

The general form for all of the alternative interpolation equations is similar to Equation (11) [White and Saunders 2007, White and Strouse 2009, White 2013]:

$$W_r(W) = \sum_{i=1}^N W_{r,i} f_i(W, W_2, W_3, \dots, W_N), \quad (17)$$

where  $N$  is the number of fixed points,  $W_i$  are the resistance ratios determined at the fixed points,  $W_{r,i}$  the corresponding reference resistance ratios, and  $W_1 = W_{r,1} = 1$  are the resistance-ratio values at the TPW. Alternative interpolating functions for each of the ITS-90 sub-ranges are listed in [Appendix 1](#), and for the most commonly used sub-ranges of the ITS-90 in Subsection 6.2, Table 4.

All of the interpolating functions satisfy the orthonormality property

$$f_i(W_j) = \begin{cases} 1, & j = i \\ 0, & j \neq i \end{cases}. \quad (18)$$

That is, they take the value one for the fixed point after which they are named and the value zero at all of the other fixed points in the interpolation. (This property is easily verified for the functions in Equations Equation (12).)

If any function  $g(W)$  can be interpolated exactly by the interpolating equations, then the function can be generated from samples  $g_{W_i}$  at the fixed points using

$$g(W) = \sum_{i=1}^N g(W_i) f_i(W), \quad (19)$$

and for all of the ITS-90 interpolations, this relation leads to the two identities

$$1 = \sum_{i=1}^N f_i(W), \quad (20)$$

and

$$W = \sum_{i=1}^N W_i f_i(W), \quad (21)$$

which are also useful for simplifying some expressions. The differences between the general ITS-90 interpolation Equation (17) and the two identities Equation (20) and Equation (21) generate two other useful forms for the ITS-90 interpolating equations:

$$W_r(W) - 1 = \sum_{i=1}^N (W_{r,i} - 1) f_i(W), \quad (22)$$

which leads to simpler expressions for Type 1 non-uniqueness [White and Strouse 2009], and

$$W_r(W) - W = \sum_{i=1}^N (W_{r,i} - W_i) f_i(W), \quad (23)$$

which is an alternative way of writing the ITS-90 interpolations in terms of the deviations.

## 2.4. Continuity of ITS-90

The ITS-90 was designed to be as close an approximation to thermodynamic temperature as was possible at the time it was formulated, so that experiments conducted into the thermal properties of systems and materials with the highest precision should not be affected by errors or inconsistencies in the scale. Discontinuities in value or derivatives between or within sub-ranges would, in particular, lead to spurious features in such data.

The reference and deviation functions specified for the various SPRT interpolation sub-ranges are all continuous as far as the second derivative, with the exception that there is in principle a small discontinuity, equal to  $2d$  ( $d$  is the coefficient in the deviation function, Table 2, see p. 9) in the second derivative of the sub-range to the silver point, at 660.323 °C (the aluminium point). However, the sub-ranges below and above the TPW are not forced to be continuous in first or second derivative at that point, and the (small) discontinuities in SPRT calibrations have been seen in the most precise determinations of thermodynamic temperature [Fischer *et al* 2011].

Insight into the continuity of ITS-90 can be gained by considering the first derivative of  $T_{90}$  with respect to the thermodynamic temperature,  $T$ ,

$$\frac{\partial T_{90}}{\partial T} = \left( \frac{\partial T_{90}}{\partial W_r} \right) \left( \frac{\partial W_r}{\partial W} \right) \left( \frac{\partial W}{\partial T} \right). \quad (24)$$

Note that ideally this should be 1, and a recent analysis [Fischer *et al* 2011] suggests that it is, in practice, always within about  $10^{-4}$  of 1.

The three terms identified in parentheses in Equation (24) relate to the derivatives of the three mathematical transformations used in the definition of ITS-90. The first term of Equation (24) is the derivative of the ITS-90 reference function, which, by design, has continuous first and second derivatives [Kemp 1991]. The third term is proportional to the derivative of the SPRT

resistance with thermodynamic temperature, which is believed to have continuous first and second derivatives, there being no structural or other transformations in platinum.

Any discontinuities therefore arise in the second term, which is the derivative of the interpolating equations, and specifically in the first derivatives of the various sub-ranges which terminate at the TPW. These are indicated by the differences between the  $a$ -coefficients in the interpolation functions below and above the TPW (but exceptionally,  $a + c_1$  for the sub-range to the triple point of neon). The differences have been reported to be in the range from 0 to  $-6 \times 10^{-5}$  [Rusby 2010]. The bias to negative values is the result of the well-documented inconsistency between the ITS-90 reference resistance ratios at the mercury and gallium points [Rusby 1993, Singh *et al.* 1994, Hill 1995]. The range of magnitudes results partly from the experimental uncertainties in the  $a$ -values, but mainly from inconsistencies between the various sub-ranges and the different behaviour of individual SPRTs (Types 1 and 3 non-uniqueness).

Types 1 and 3 non-uniqueness (Section 6.3, see p. 44) will also lead to small differences between the first and second derivatives of  $T_{90}$  in the various sub-ranges, which could affect precise measurements of thermal properties, such as heat capacities. The magnitude of these effects can be estimated from the slopes of non-uniqueness plots as functions of temperature, and for the most part is  $< 0.01\% (0.1 \text{ mKK}^{-1})$ , except perhaps at low temperatures, approaching 14 K.

## 3. DESIGN AND OPERATION OF SPRTS

### 3.1. Operating principles and overview

All pure metals exhibit an almost linear temperature dependence of electrical resistance at sufficiently high temperatures. Amongst the metals that can be used for resistance thermometers, platinum is preferred because of the very wide temperature range over which it exhibits good immunity against chemical and physical effects that influence the resistance-temperature characteristic of the thermometers.

Under the influence of an electric field, the conduction electrons in a metal (i.e., those that are not bound to a particular atom) are free to move through the crystal lattice and so to conduct electricity. In an ideally pure metal at the absolute zero of temperature, there is no resistance to the current because no lower-energy states are available for the electrons to scatter into. At non-zero temperatures the electrons are scattered by thermal vibrations in the lattice and by other electrons, and this gives rise to the temperature-dependent ‘ideal’ resistivity of the metal. In a real metal the electrons are also scattered by impurities and by imperfections in the lattice, such as interstitial atoms, dislocations, vacancies and grain boundaries. According to Matthiessen’s rule, this additional resistivity is, to a first approximation, temperature-independent. The loss of energy due to scattering of the electrons is the origin of the Joule heating in the metal.

In the manufacture of a thermometer, the aim is to ensure that scattering due to impurities, etc., is minimised, leaving only the nearly ideal resistance of the pure platinum. Other influence effects originating outside the platinum metal contribute measurement errors, such as electrical leakage in the insulating components, which shunts the platinum resistor, and thermal resistance between the platinum resistor and the surroundings, which restricts the dissipation of the Joule heating. These should also be minimised, as far as possible, in the design of the thermometer.

### 3.2. Typical designs of SPRTs

The platinum wire for the sensing elements in all SPRTs is obtained “hard drawn”, as it is easier to handle in this condition, but it is thoroughly annealed during the manufacture of the thermometer. The sensing-element winding is usually bifilar, but occasionally other low inductance configurations are used. A low inductance is important if the thermometer is to be used in ac measurement circuits, and reduces the SPRT sensitivity to electromagnetic interference. In all SPRTs, the sensing element must be supported in a strain-free manner on a structure, usually made of clean high-purity polished silica, with no rough or sharp edges. As a result, a well designed and manufactured SPRT does not show hysteresis during thermal cycling. The sensing element is sealed in a suitable atmosphere and connected within the sheath to four platinum leads, two for the passage of the current and two for sensing the voltage. This is to enable a truly four-wire resistance measurement, in which lead resistance effects are eliminated.

Mechanical, electrical and thermal constraints dictate that no SPRT can be used over the whole temperature range from 13.8033 K to 1234.93 K, and in practice three distinct types are used. ‘Capsule-type’ SPRTs are designed for operation in the temperature range from 13.8 K to

about 430 K. ‘Long-stem’ SPRTs are used in the temperature range from  $-189.3^{\circ}\text{C}$  to  $660^{\circ}\text{C}$ . Both of these types typically have a resistance of about  $25\ \Omega$  at the TPW, giving a nominal sensitivity of  $0.1\ \Omega/\text{K}$ . At a measuring current of 1 mA, the self-heating effect is usually in the range from 0.2 mK to 4 mK. ‘High-temperature’ long-stem SPRTs (HTSPRTs) are used at temperatures up to  $962^{\circ}\text{C}$ . Their resistance at the TPW is typically between  $0.2\ \Omega$  and  $2.5\ \Omega$  and higher measuring currents are used (see below).

### 3.2.1. Capsule-type standard platinum resistance thermometer

Capsule-type standard platinum resistance thermometers (CSPRTs) are typically used between 13.8 K and  $30^{\circ}\text{C}$ , but sometimes as high as  $156^{\circ}\text{C}$ , and very occasionally to  $232^{\circ}\text{C}$ . A schematic diagram of a typical design of a  $25\ \Omega$  CSPRT is shown in Figure 1. The platinum sensor is mounted on an insulating former and inserted into the sheath, which is a platinum or glass tube about 5 mm in diameter with a closed end. The four short (30 mm to 50 mm) platinum lead wires emerge through a glass seal at the open end of the sheath. Electrical connections can be made to the leads with ordinary soldering techniques; however, care must be taken to avoid straining the leads where they emerge from the glass seal as they are prone to breaking there. A common solution is to tie the (insulated) leads together above the top of the capsule and, if need be, ply them back from there. The capsule is filled with helium gas, usually at a pressure of about 30 kPa at room temperature, to ensure good thermal coupling between the wire and the sheath.

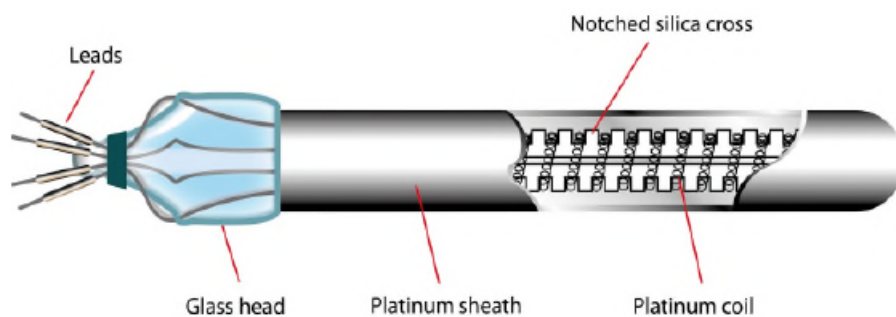
For calibration at the fixed points of mercury, water and gallium, the capsule may be housed in an adaptor made of a small copper sleeve to fill the gap between the capsule and the inner wall of the cell. The sleeve is attached to a stainless-steel capillary which leads up to room temperature. Two thermal shunts between the capillary and the cell, one close to the capsule and one near the top of the cell, reduce heat flow along the leads. Such an adaptor eliminates the problems associated with simply suspending the capsule from its wires.

The low temperature limit for the use of CSPRTs has been set at 13.8 K because at lower temperatures their resistance and sensitivity become inconveniently small. Also, the low-temperature characteristics become increasingly dependent on non-thermal resistance effects due to strain, lattice defects, and impurities. To compensate for the loss of sensitivity, measuring currents below 24.5 K are generally increased from the 1 mA usually used above that temperature, to about 5 mA. At this level, because of the low values of resistance in this range (down to about  $30\ \text{m}\Omega$ ), the self-heating should not exceed 0.2 mK, and the minimum sensitivity is  $\sim 30\ \mu\text{VK}^{-1}$ .

The use of CSPRTs above  $30^{\circ}\text{C}$  is limited by electrical leakage in the glass seal, particularly if the surface is contaminated, e.g. by solder flux, and the diffusion of helium through the glass. Calibrations up to 505 K (the freezing point of tin) are possible, though the results at this temperature are not likely to be of the best quality and the calibration at low temperatures may be adversely affected. Because of these limitations, CSPRTs cannot be annealed, so that any change in resistance due to mechanical shock or other influences effectively causes a permanent shift in the CSPRT resistance. Regular checks should be carried out to look for such shifts, either by measuring the ‘residual resistance’ in liquid helium, where the temperature coefficient of the CSPRT is very small, or the resistance at the triple point of water, which can be measured easily and accurately. The residual resistance measurement is often more effective for detecting small changes, however, as part of a cryogenic run, but care must be taken because the temperature dependence of the resistivity near 4 K is not zero [Tew *et al.* 2013]. If a significant change is found in the residual resistance it cannot be reversed. It may,

as a first approximation, be satisfactory to correct for it by subtracting this resistance change from all measured values, but more usually a new calibration is indicated.

CSPRTs are usually used “totally immersed”, meaning that the entire capsule is immersed in the medium of interest; preferably inserted in a well in a copper block in the cryostat. Because thermal conductivity and electrical conductivity are closely related phenomena, the thermal conductivity of thermometer leads is relatively high at low temperatures. To avoid heat leaks through the leads, connections are usually made via long fine copper wires thermally anchored to the block, or another object at a similar temperature, to reduce or eliminate heat flow into or out of the thermometer. This is particularly important where the wires are in the cryostat vacuum space, with little heat exchange along their lengths. When the anchoring is done correctly, the effect of heat leaks can be reduced to well below 0.1 mK [Hust 1970, Kemp *et al.* 1976, Gaiser and Fellmuth 2013a]. Heat leaks also influence measurements in higher-temperature applications where CSPRTs are used at the fixed points of mercury, water and gallium. Immersion in air within fixed-point cells provides insufficient thermal coupling so close-fitting sleeves, a suitable oil, or an adaptor as discussed earlier, are needed.



**Figure 1 — Schematic diagram of a typical 25 Ω capsule-type SPRT. The sensing element is formed from fine, ~0.07 mm diameter, coiled platinum wire supported on a notched high-purity silica cross. The four lead wires are welded to the sensing element, two at each end, and pass through the glass seal. The platinum sheath is about 5 mm in diameter and 50 mm long. (Illustration published on courtesy from FLUKE®)**

### 3.2.2. Long-stem standard platinum resistance thermometer

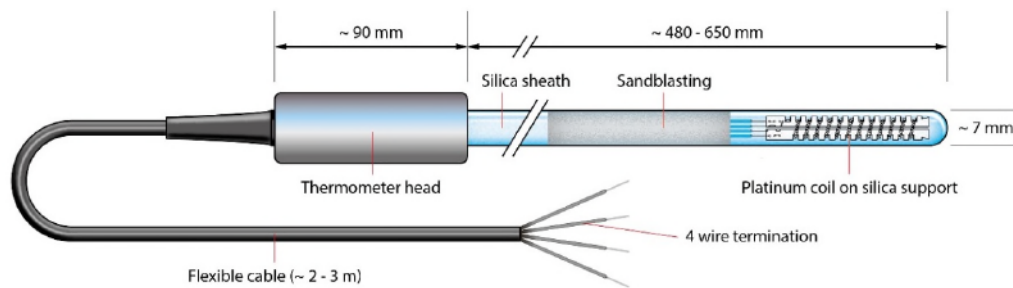
Long-stem standard platinum resistance thermometers (LSPRTs) are applied in the temperature range between the triple point of argon ( $-189.3442\text{ °C}$ ) and the freezing point of aluminium ( $660.323\text{ °C}$ ). A typical design is shown in Figure 2. The sensor element is platinum wire of 0.05 mm to 0.1 mm diameter, which may be wound onto the former in a variety of ways. The wound element is placed into a fused-silica tube with a length of 480 mm to 650 mm and outside diameter up to about 8 mm. The tube is sandblasted or blackened along part of the lower length to reduce radiative heat transfer (light-piping) between the element and the head at room temperature. Some metal-sheathed SPRTs are also made (see comments in Sec 5.1.1). The former for the sensing element is usually made of fused silica, occasionally Vycor or ceramic, and in older thermometers, mica. The length of the element is typically 35 mm to 50 mm, and it has a nominal resistance at the temperature of the TPW of 25 Ω, giving the SPRT a sensitivity of  $0.1\text{ Ω/°C}$ .

The upper temperature limit for LSPRTs having mica formers or mica insulation is  $500\text{ °C}$  due to the release of water of crystallisation from the mica at higher temperatures. Once driven from the mica, the water condenses at temperatures near  $0\text{ °C}$  causing electrical leakage and

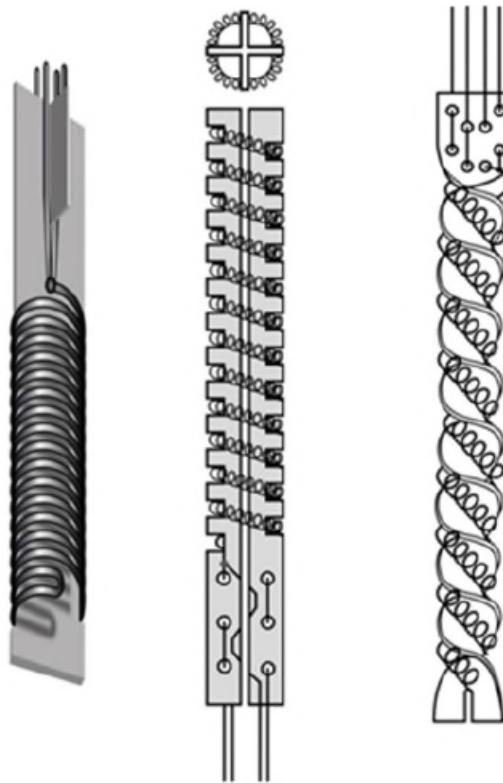


measurement errors. Above 660 °C, LSPRTs with silica insulators exhibit an exponentially increasing electrical breakdown of the insulation that can lead to errors of a few tens of millikelvin at 960 °C (see Section 3.2.3, see p. 19). Mechanical problems can also arise due to the large difference between the thermal-expansion coefficients of platinum and fused silica, leading to mechanical deformation and short circuits either in the sensor element or between lead wires.

Figure 3 shows two of the most frequently used designs of the resistance element of LSPRTs. Because the plasticity of platinum increases sharply at temperatures above 200 °C, the sensor element should have a design that prevents the coil from sliding along the former and causing short-circuiting of the wire turns. Four platinum leads with a diameter of 0.2 mm to 0.4 mm connect the resistance element to the external copper cable and connections. The lead wires are usually kept apart by silica (or mica) discs. In some designs, the leads may be placed into separate fused silica capillaries to prevent short-circuiting. The silica or mica discs also reduce convection within the thermometer.



**Figure 2 — Schematic diagram of a typical 25 Ω LSPRT. The sensing element is a platinum coil with a typical length of 35 mm to 50 mm. (Illustration published on courtesy from FLUKE®)**



**Figure 3 — Schematic designs of sensitive elements of SPRTs: HTSPRT with a simple bifilarplatinum coil supported on a notched high-purity silica plate,  $R(0.01\text{ }^{\circ}\text{C}) \sim 0.25\text{ }\Omega$  (left); LSPRT with the platinum coil laid in notches in a silica cross,  $R(0.01\text{ }^{\circ}\text{C}) \sim 25\text{ }\Omega$  (centre and top-view); LSPRT with the platinum coil laid in the helical turns of a twisted silica plate,  $R(0.01\text{ }^{\circ}\text{C}) \sim 25\text{ }\Omega$  (right). In all cases two platinum wires are welded to the coils at each end to create four-lead connections ("Y" joints). (Illustration published on courtesy from FLUKE®)**

To ensure the thermometer sheath is hermetically sealed at the head where the platinum wires exit the assembly, ‘graded’ seals are often used. The seals are designed to bond to the platinum wires to form an air tight seal, and are formed from layers of glass with graded linear coefficients of thermal expansion to match that of platinum where the seal joins the wire and match the fused silica where it joins the sheath. The platinum leads are connected in the head of the thermometer to 4-wire multi-strand copper cable to allow connection to the measuring instrument. The leads may terminate as plain wires, spades, banana plugs or BNC connectors. Sometimes a cable screen is used to connect the sensor or the thermometer head to a ground at the resistance bridge.

The thermometer sheath in an LSPRT is usually filled with a mixture of inert gas plus oxygen at a partial pressure of about 2 kPa. The total pressure is equivalent to one atmosphere at the maximum temperature of the thermometer’s application range. The low level of oxygen is sufficient to oxidise metallic contaminants and hold them at grain boundaries: they might otherwise migrate into the platinum lattice and affect the temperature dependence of the resistivity. At the same time, it limits the formation of an oxide film on the surface of the platinum wire (see Section 5.1.1, see p. 24).

### 3.2.3. High-temperature standard platinum resistance thermometer

High-temperature standard platinum resistance thermometers (HTSPRTs) are designed for use in the temperature range up to the freezing point of silver (961.78 °C). Their design is very similar to that of LSPRTs, except that they use lower resistance sensors (with larger gauge platinum wire) in order to reduce the effects of electrical leakage at temperatures above the freezing point of aluminium. Figure 3 shows also one design of HTSPRT sensing elements.

At temperatures approaching the silver point, the electrical conductivity of the fused silica formers and discs, which make up the insulating components of thermometer, becomes appreciable. The effects shunt the sensing element leading to errors of several tens of millikelvin. By reducing the resistance  $R(0.01\text{ °C})$  from the  $25\ \Omega$  value used in LSPRTs to  $0.25\ \Omega$  the influence of the unwanted shunting resistance is reduced by a factor of 100. In practice  $R(0.01\text{ °C})$  values ranging from  $0.2\ \Omega$  to  $2.5\ \Omega$  are used. A measuring current in the range 5 mA to 10 mA is typical for HTSPRTs, which partially compensates for the low resistance and loss of voltage sensitivity.

The shorter length of larger diameter wire (0.3 mm to 0.5 mm) used in the low-resistance sensors simplifies the construction and has other beneficial effects. Firstly, it improves structural stability of the element against the large difference between the thermal expansion of silica and platinum. The differential thermal expansion also means that about 1 cm of space must be left at the bottom of the sheath, when the thermometer is cold, to allow the platinum lead wires to expand. A second advantage of the thicker wire is that it reduces the surface/volume ratio of the wire, which slows the impact of contaminants, making the thermometer more stable, and reduces the effects of oxidation at the surface of the wire.

At the higher temperatures, metals and other potential contaminants of platinum become increasingly mobile and volatile, with some contaminants able to diffuse through the silica sheath. Cleanliness of the sheath and preventing metallic contamination of the sensor element is critical for long-term reliability of HTSPRTs.

## 4. STANDARD PLATINUM RESISTANCE THERMOMETER USE AND CARE

### 4.1. Mechanical Treatment and Shipping Precautions

SPRTs are delicate instruments. Shock, vibration, or any other form of acceleration may cause the wire to bend between and around its supports, producing strains and mechanical damage. In the worst case, careless day-to-day handling of a thermometer over a year has been observed to increase its resistance at the TPW by an amount equivalent to as much as 0.1 K, and on rare occasions single incidents have caused similar changes or complete failure. Changes may be caused over long periods by an apparatus that transmits vibrations to the thermometer, or by shipping the thermometer in an unsuitable container.

It is strongly advisable to hand-carry SPRTs to maintain the integrity of calibration. If a thermometer must be shipped, it should first be placed in a rigid and moderately massive container that has been lined with soft material which conforms to the thermometer shape and protects it from mechanical shocks or vibrations. This container should then be packed in an appreciably larger box with room on all sides for soft packing material that will substantially attenuate any shocks that might occur during shipment.

### 4.2. Thermal Treatment and Annealing

Generally, the greatest mechanical damage to thermometers occurs during manufacture and shipping. While thermometers are annealed in the process of their production, this annealing is not always sufficient. Therefore, with new SPRTs and SPRTs recently shipped, it is advisable to measure first the resistance at the TPW and then beginning annealing.

LSPRTs (used up to 660 °C) should be placed in an auxiliary furnace at 480 °C to 500 °C before the temperature is increased to about 675 °C over a period of about 45 minutes to 60 minutes. The LSPRT is then annealed at this temperature for four hours. Afterwards, the temperature is reduced again to about 480 °C over a 4 hour period, after which the PRT is removed directly from the furnace to the room-temperature environment and, as soon as practical, measured at the TPW. If, after annealing, the resistance of the sensitive element has not changed by more than 0.5 mK temperature equivalent, it is ready for use. If the change of resistance is significant, annealing should be repeated. If the resistance increases after each annealing cycle, the thermometer is probably contaminated or affected by three-dimensional oxidation and should not be used.

HTSPRTs (used up to 962 °C) are annealed in the same way, but the annealing temperature is 975 °C and they are annealed for four to six hours. After annealing they should be cooled slowly to 480 °C at no more than 50 °C per hour. At high temperatures, the SPRTs become particularly sensitive to thermal and mechanical shock and should be handled very carefully.

To remove the defects and strain caused by normal handling, annealing should be performed before LSPRTs are employed at the aluminium freezing point or HTSPRTs at the silver freezing point. After an annealing period of about 30 minutes the SPRT should be transferred quickly, but gently, into the fixed-point cell. After completion of the measurements at these fixed points, the SPRTs should be again transferred to an annealing furnace at a temperature of about 675 °C (LSPRTs) or 975 °C (HTSPRTs), maintained at that temperature for at least

30 minutes, and then cooled slowly to near 480 °C to restore the low-temperature vacancy concentration. From this temperature, the SPRT must be removed relatively quickly (< 10 minutes) from the furnace into the room-temperature environment to prevent oxidation and measured at the TPW as soon as possible to ensure that the two measurements performed for the determination of the resistance ratio  $W$  are obtained in the same oxidation state of the platinum sensor.

With measurements above the aluminium freezing point, great care must be taken to avoid contamination of SPRTs by metallic impurities, and some means of protecting the thermometer may be required. Sometimes a platinum foil layer is inserted or sapphire tubes are used.

No preliminary treatment of CSPRTs is necessary. The stability of the thermometer is checked by monitoring its resistance at the triple point of water, and also its resistance at a low temperature, such as 4.2 K or at the triple point of hydrogen (see Section 3.2.1, see p. 15).

### 4.3. Devitrification

The sheaths of LSPRTs and HTSPRTs are usually made from fused silica; silica in a glassy state that is largely impervious to most contaminants. However, there are some contaminants, notably sodium chloride from perspiration, that cause the silica to return to its crystalline state (much like quartz, the natural crystalline form of silica), which is an irreversible phase transition. In its devitrified state, silica is milky white, very brittle, and permeable to gases. Devitrification tends to occur at high temperatures and is catalytically enhanced by alkali compounds.

Before using silica-sheathed platinum resistance thermometers above 100 °C, they should be carefully cleaned with pure ethanol and dried with clean paper or cloth. The cleaning with diluted nitric acid followed by washing with clear water is also suitable. This is especially important for using HTSPRTs above 660 °C. It serves to remove all traces of fingerprints that would otherwise trigger devitrification at high temperatures, and may cause patterns of devitrification to become visible. The first traces of devitrification of the outer surface of the sheath should be removed by sandblasting with alumina powder in order to stop the process. Protection from contamination of the element and from devitrification of the sheath becomes increasingly difficult at higher temperatures.

### 4.4. Calibration

#### 4.4.1. Calibration Procedures

For the lowest uncertainties, SPRTs should always be calibrated in their most reproducible state; that is, the resistance should correspond to the zero-current resistance when the SPRT is in an unoxidised state, and with an equilibrium concentration of vacancies. While this state is generally not entirely accessible to CSPRTs, it is the ideal operating state of LSPRTs and HTSPRTs.

Before the calibration begins, LSPRTs and HTSPRTs should be annealed until the  $R(0.01\text{ °C})$  value is stable; typically within 0.2 mK for a LSPRT, and 0.8 mK for a HTSPRT. Once the SPRT is stable, fixed-point measurements should be made progressing from the highest

temperatures to the lowest temperatures. For LSPRTs and HTSPRTs and fixed points temperatures below and including the zinc point, the  $W$  values for the fixed points should be calculated using TPW measurements made in the same oxidation state as during the fixed-point measurement. This means doing a TPW measurement within a few hours after the fixed-point measurement. If the slow cool-down from a high temperature is run overnight and continues much below 480 °C, the SPRT should be re-heated to 480 °C for 1 hour, and then removed to room temperature before the TPW measurement is made.

Where corrections to fixed point resistance measurements are required, such as for self-heating, hydrostatic pressure, gas pressure, impurity, or isotope effects, they should be made to the measured resistance values, before the ITS-90 interpolating equations are applied.

#### 4.4.2. Consistency checks

SPRT calibrations involve a large number of measurements and corrections with considerable opportunity for operator mistakes, either in the signs of corrections, or operating conditions of the fixed points, or in the values of standard resistors. Therefore, it is helpful to include a number of checks in a procedure to build confidence and reduce the chances of making mistakes.

As much as is possible, validated software should be used for making the corrections and calculating the various  $W$  values from the bridge ratio measurements and standard resistor values. The software should be validated using either synthetic data (including pressure corrections, etc.), or older data from a real SPRT that has been checked using independent software or a calculator.

Where possible, a sequence of fixed-point measurements should include repeat measurements or additional fixed points as a redundancy check. Some laboratories use the indium point as the redundant point for LSPRT and HTSPRTs, to check that the value of  $W_{r,In}$  deduced from the measured  $W_{In}$  is close to the ITS-90 reference value. Plots of the calculated deviation function  $\Delta W$  versus  $W$  should also be done. For temperatures above 50 K, the curve should have weak quadratic shape and be smooth.

When calibrating a group of SPRTs, at least one SPRT should be included for which the calibration data are already known. Many laboratories employ check thermometers for specific fixed points. This helps to ensure that there are no major changes in the furnace operating conditions, and ensure the fixed-point cells have not developed leaks to atmosphere and have not become contaminated.

It is also helpful to calculate the parameter

$$S_i = \frac{W_i - 1}{W_{r,i} - 1} \quad (25)$$

for each fixed point. These  $S_i$  values should be very similar for all fixed points above 50 K, although there is a step down in crossing to temperatures above 273.16 K due to a small inconsistency in the ITS-90 definitions. (The use of the  $S_i$  values for checking purposes has been proposed in [White and Strouse 2009] applying the alternative interpolation Equation (22). The near constancy of the  $S_i$  values for each SPRT is a consequence of the resistance closely following Matthiessen's rule. This rule states that the additional resistivity caused by lattice imperfections is nearly independent of temperature.)

The measurements of  $W$  should be checked to ensure that the SPRTs meet the ITS-90 quality criteria Equation (2) and Equation (3); most SPRTs meet these criteria comfortably. Note that

the two relations defined by ITS-90 are not quite consistent: an alternative but very similar requirement is that  $S_i$  values from Equation (25), calculated for all fixed points, should be greater than 0.9994 [White and Strouse 2009]. This requirement has the advantage of being applicable to fixed points other than mercury and gallium. Additionally, a large drop in the  $S_i$  value for the silver-point measurements is a more sensitive indicator of insulation breakdown than the ITS-90 quality criterion Equation (4).

#### 4.4.3. Reporting

The simplest and minimal option for the presentation of SPRT calibration data is to report the measured  $W$  values for all of the fixed points, and their uncertainties. The uncertainties should be presented in terms of resistance ratio, though it is helpful to give the equivalent temperature uncertainty. It is also essential to include the coefficients for the interpolating equations realisable using those fixed points as additional information.

It is also necessary to report the resistance value at the TPW for the SPRT. This value is required to track the stability of the SPRT with shipping to and from the calibration laboratory. This value of  $R(0.01\text{ °C})$  should not be used to calculate  $W$  values: instead the user should measure the  $R(0.01\text{ °C})$  on their own equipment, as this minimises the propagation of uncertainties associated with the bridges and standard resistors, and real changes in the SPRT resistance.

[Appendix 2](#) provides a table of typical resistance ratios and sensitivity coefficients at the fixed points, which are useful for calculating the temperature equivalents of measurement uncertainties.

## 5. EXPERIMENTAL SOURCES OF UNCERTAINTY

This subsection provides a brief summary of the known sources of uncertainty affecting platinum resistance thermometry, see also the overview of factors influencing the resistance given in [Pokhodun 2002, Meyer and Ripple 2006]. The subsection explicitly excludes effects relating to fixed-point realisations (impurities, isotope effects, hydrostatic corrections, pressure effects, etc), since sources of uncertainty associated with fixed points are covered in [Chapter 2 Fixed Points](#) (<http://www.bipm.org/en/committees/cc/cct/guide-its90.html>), and effects associated with the triple point of water are covered in [Section 2.2 Triple Point of Water](#) of this chapter. The budget for estimating the uncertainty of the ITS-90 realisation with SPRTs calibrated at the fixed points is presented in Subsection 6.

### 5.1. Factors affecting SPRT resistance

#### 5.1.1. Oxidation

SPRT sheaths are filled with a gas for improving the thermal coupling between the sensing element and the sheath. In the past, dry air was widely used, but nowadays the gas consists mainly of a mixture of an inert gas, often argon or helium, with a partial pressure of oxygen between about 2 kPa to 10 kPa. An oxygen content of at least 1 kPa is necessary to prevent contamination of the sensing element by metallic impurities reduced from their oxides. On the other hand, effects associated with the oxidation of the platinum cause hysteretic changes in the SPRTs resistance. This effect is complicated further by its dependence on the partial pressure of oxygen, the operating temperature, the presence of impurities, and crystal size and orientation [Wang and Yeh 1998]. To reduce oxidation effects, the oxygen content should be as small as practical. A partial pressure of oxygen of about 2 kPa is a good compromise between the need for oxygen and its deleterious effects.

The chemical interactions between platinum and oxygen are complex, with as many as a dozen possible oxides and allotropes. The oxides most relevant to platinum thermometry are PtO, PtO<sub>2</sub> and Pt<sub>3</sub>O<sub>4</sub> [Berry 1978, 1980, 1982a, 1982b, Seriani *et al.* 2006, Sakurai and Tamura 2011, Jursic and Rudtsch 2014]. Platinum oxidation has been investigated by variously applying calorimetry,

thermogravimetry, mass spectrometry, electron diffraction, acid solubility, and thermodynamic modelling [Seriani *et al.* 2006, Sakurai *et al.* 2008, Sakurai and Tamura 2011].

For partial pressures of oxygen of a few kPa, the formation and dissociation of platinum oxides takes place with temperature-dependent rates as follows. Up to about 350 °C, which is dependent on the partial pressure of oxygen, less than one monolayer of oxide is formed, so the oxide is described as two-dimensional. The rate of formation is slow, being just detectable after one hour at 200 °C, and is still progressing after many tens of hours at 300 °C. In saturation, its temperature equivalent of the effect amounts to about 1 mK to 2 mK in 25 Ω thermometers.

At higher temperatures, the two-dimensional oxides dissociate. But at temperature up to about 550 °C, a three-dimensional surface layer may slowly propagate deeper into the wire, apparently limited only by diffusion and without signs of saturation. The temperature equivalent of the resistance change may reach 10 mK or more in 25 Ω thermometers, and it depends on the heat-treating period. This effect also depends on the partial pressure of oxygen in the exchange gas; both the rate of oxidation and the dissociation temperature decrease as the



partial pressure is decreased [Ancsin 2003]. Studies performed by Berry (1980, 1982a, 1982b) and Sakurai and Tamura (2011) show that the formation of the three-dimensional oxide can usually be suppressed by applying partial oxygen pressures of only a few kPa. Heat treating the SPRTs at 600 °C or higher will cause the three-dimensional oxide to dissociate and should largely restore the SPRT. However, extended heat treatment for very long periods ( $> 10$  hours) may be required, and in some SPRTs, the effect can be difficult to suppress. Sakurai and Tamura (2011) have also observed anomalous phenomena that indicate a kind of irreversibility associated with oxidation.

When oxidation effects occur in SPRTs, the relative change of the resistance ratio  $W(T_{90})$  is much smaller than that of  $R(T_{90})$ . This is caused by the fact that oxidation and dissociation change the cross-sectional area of the platinum core, while the resistivity of the core remains unchanged. Berry (1980, 1982a) has developed a two-zone model with a relatively poorly conducting oxide film at the surface for describing the two-dimensional effect. The moderately good stability of  $W$  with respect to oxidation reduces the problem to a manageable level in most cases. The main difficulty for high-accuracy measurements is that the measurements of  $R(T_{90})$  and  $R(273.16\text{ K})$  used for the  $W(T_{90})$  calculation must correspond to the same oxidation state, and hence  $R(273.16\text{ K})$  must be measured frequently if the oxidation is causing substantial drifts. Particular care may be necessary when the SPRT passes from an oxidising to a dissociating temperature range. If a frequent measurement of  $R(273.16\text{ K})$  is not feasible, dedicated experiments are required to investigate the magnitude of the effects and make a reliable estimate of the uncertainty.

### 5.1.2. Impurities

In an ideal metal conductor, electrical resistance is caused mainly by the scattering of electrons due to the thermal motion of the metal atoms. Additional scattering of electrons caused by impurities gives rise to an additional contribution to the electrical resistance, which is approximately independent of temperature [Berry 1963]. Impurities cause irreversible changes in the resistance-temperature dependence of SPRTs, and are a main cause of long-term drift. The impurities may originate from the production of the wire, including insufficient purification of platinum, contamination during preparation of the sensing element or production of the thermometer. Impurities may also originate in the sheaths after the thermometer is assembled, by diffusion especially from metal sheaths, at temperatures above 450 °C. Some impurities can diffuse through fused-silica sheaths at temperatures above about 900 °C [Marcarino *et al.* 1989]. For thermometers used at temperatures below 450 °C, the long-term drift is usually very small, no more than 1 mK with many hundreds of hours of use [Berry 1962]. Above 450 °C the drift increases with increasing temperature and length of exposure and may reach 5 mK/100 h at the silver point [Berry 1966, Fellmuth *et al.* 2005]. The form and speed of drift depend on quantitative and qualitative composition of impurities in the platinum wire [Pokhodun *et al.* 2005].

### 5.1.3. Strain and hysteresis

Elastic strain on the platinum wire, which causes a temporary distortion of the atomic lattice, modifies the resistance. From research on platinum strain gauges it is known that electrical resistance of platinum increases when the wire is under tension and decreases when under compression. Strain typically arises from differential thermal expansion between the platinum wires and its insulating supports, or from mechanical movement of the wire, or a mixture of both. Because it is a purely elastic effect, the deformation disappears on cessation of the

mechanical forces. However, where mechanical and thermal causes combine, the effect gives rise to hysteresis, which becomes apparent with cyclic excursions in temperature of more than several tens of degrees.

The magnitude of the hysteresis is strongly influenced by the design of the SPRT, the materials used in its construction, and the degree to which the platinum is allowed to expand and contract relative to support structures. The effects are typically no more than 0.3 mK peak-to-peak, but can be as low as 0.1 mK peak-to-peak and range up to 1.8 mK peak-to-peak [Berry 1983].

#### 5.1.4. Vacancies and defects

In a manner, very much like the effects of impurities, defects in the crystal structure of the platinum lattice also increase the resistance of the wire. The defects may be induced thermally or mechanically.

##### 5.1.4.1. Thermally induced defects

At temperatures above about 450 °C, the thermal motion of the atoms occasionally causes some atoms to jump out of position in the lattice, creating vacancies (absence of an atom), interstitials (extra atom where one shouldn't be) or other crystal dislocations. The effect increases exponentially with rising temperature, and at higher temperatures, is responsible for the formation of more complex, multi-atom (higher energy) defects, corresponding to many tens of millikelvin temperature equivalent of the resistance change at 962 °C.

Given sufficient time at any temperature, the concentration of thermally induced vacancies reaches an equilibrium concentration, causing an increased resistance of approximately [Berry 1966]

$$\delta R \approx 1200R(273.16 \text{ K}) \exp(-E_a/kT), \quad (26)$$

where  $E_a$  is the activation energy for the formation of the vacancy or defect in the lattice ( $\approx 1.5$  eV for simple vacancies in platinum). The additional electrical resistance caused by the vacancies is considered to be an integral part of the  $R(T_{90})$  characteristic of the thermometer, but it becomes detectable after quenching. Because the equilibrium concentration is determined by the rate at which vacancies are created or destroyed, the rate at which the equilibrium is reached is also temperature dependent. This means that defects anneal out according to an exponential law with a half-life,  $\tau$ , dependent on the activation energy for the defect:

$$\tau = \tau_0 \exp(E_a/kT), \quad (27)$$

where  $\tau_0$  is a time constant characteristic of the diffusion and equilibration process. For simple low-energy vacancies, the half-life ranges from milliseconds at 960 °C to hours at temperatures below 400 °C. Thus, equilibration occurs very quickly when moving to higher temperatures but very slowly when moving to low temperatures. If an SPRT is cooled too quickly, the vacancies can become quenched-in and cause errors in the  $R(T_{90})$  characteristic at lower temperatures. Recommended cooling rates vary but should be no more than 50 °C per hour when cooling from temperatures above 500 °C [Mangum *et al.* 1990]. The annealing of defects with higher activation energies requires higher temperatures and longer annealing times. Once the SPRT has cooled to 500 °C from higher temperatures, the residual effects of vacancies are negligible, and the thermometers should be cooled quickly (a few minutes) to room temperature to avoid oxidation.

#### 5.1.4.2. Mechanically induced defects

The most troublesome defects are those arising from mechanical damage to the wire. Typically, a large number of defects, including high-energy defects, are introduced during the cold drawing of the platinum wire prior to the manufacture of the SPRT. Although the SPRT is thoroughly annealed at the time of manufacture, many of the higher-energy defects will persist and be a permanent part of the thermometer behaviour. Mechanical shock or vibration during use is a major cause of defects and, hence, drift in SPRTs. Mechanical damage, to some degree, occurs during all use of the thermometers with effects of the order of microkelvin accumulating each time an SPRT is knocked [Berry 1962]. Berry (1983) suggested that the mechanical damage could be classified as inelastic or plastic according to the degree of damage it caused.

Inelastic deformation occurs when forces below the yield point of the wire are applied, but the deformation does not disappear with cessation of the mechanical force. The SPRT can be restored by annealing, so the effect also gives rise to hysteresis. Berry (1983) describes this effect as similar to internal friction, and is probably due to the creation of low-energy defects.

Plastic deformation arises when the mechanical forces exceed the yield point of the platinum wire, and is usually caused by mechanical shock (i.e., it is rarely due to thermal effects). It leads to strong deformations of the crystal lattice accompanied by the generation of many defects, only some of which will be removed by annealing.

In principle, all crystal defects can be removed with sufficient annealing [Berry 1966, 1972, Berry and Lamarche 1970]. However, for the highest-energy defects, the annealing temperatures may be beyond the material limits for the SPRT sheaths and support structures. In these cases, the effects of such defects are practically irreversible. Long-term drifts in SPRTs, whether caused by impurities, highest-energy defects or dimensional changes, are evident from changes in the  $R(273.16\text{ K})$  values that cannot be removed by annealing. Note that long-term downward drifts in resistance are unusual, but may be due to insufficient annealing during manufacture.

Some of the causes of long-term drift effects can be distinguished. In particular, dimensional changes arising from plastic deformation, volatilisation of platinum, and changes incurred during an episode of three-dimensional oxidation, lead to a change in the resistance  $R(273.16\text{ K})$  but not the  $W(T_{90})$  value [Berry 1966] (“ $\delta R/R \approx \text{const. instability}$ ”). On the other hand, defects and impurities tend to cause an increase in resistance that is more or less independent of temperature (“ $\delta R \approx \text{const. instability}$ ”), so that both  $R(273.16\text{ K})$  and  $W(T_{90})$  change. The resulting change of the resistance ratio is given by the relation  $\delta W = (1 - W)\delta R(273.16\text{ K})/R(273.16\text{ K})$ .

For capsule-type thermometers, the upper temperature limit (typically  $156^\circ\text{C}$ ) means defects cannot be removed by annealing and are, therefore, practically indistinguishable from impurity effects. A valuable (high resolution) indicator of resistance shifts in capsule-type SPRTs is the residual resistance at liquid-helium temperature. It is therefore good practice to check it at regular time intervals. But a careful check requires consideration of the small temperature dependence of the resistivity at these temperatures [Tew *et al.* 2013].

#### 5.1.5. Moisture

Berry (1966) and Zhang and Berry (1985), established that the electrical insulation resistance of mica-insulated SPRTs deteriorates once the thermometer has been used much above  $500^\circ\text{C}$  for an extended period. Marcarino *et al.* (1999) subsequently demonstrated an effect as large

as 1 mK or more. The problem is caused by the release of water of crystallisation from the mica when the SPRT is exposed to high temperatures. Quartz insulated SPRTs may also exhibit a moisture effect due to the small quantities of water trapped within the sheath during manufacture, although the effect is usually much smaller than for mica. The effect of the water is greatest near 0 °C, where it condenses or freezes on the internal surfaces of the SPRT. The effect decreases exponentially as the temperature moves away from 0 °C [Berry 1966]. As the temperature decreases below 0 °C, the conductivity of the ice falls, while above 0 °C an increasing fraction of the water is vapour. At –39 °C, the effect is very much reduced and by 200 °C the effect is negligible.

SPRTs affected by moisture typically exhibit long (> 10 minutes) settling times, and sometimes instability at the TPW. They will also exhibit hysteresis due to the migration of moisture within the sheath, which may be confused with oxidation effects. The effect may be sensitive to the operating frequency of the resistance bridge, so it is sometimes detectable by changing the frequency (current-reversal times for dc bridges) [Marcarino *et al.* 1999].

The presence of moisture can be assessed by using dry ice to cool the upper end of the SPRT sheath with the SPRT in a water triple point cell. The dry ice condenses the moisture away from the electrically conducting elements of the SPRT. The observed change in the triple-point resistance typically ranges from undetectable for good quartz SPRTs to 100 µK or more for older mica-insulated SPRTs.

#### 5.1.6. High-temperature insulation breakdown and contamination

At sufficiently high temperatures, all of the insulating materials used in SPRTs show the thermistor-like decrease in electrical resistivity characteristic of large-band-gap semiconductors. The resistance decreases exponentially with temperature, causing significant effects in 25 Ω SPRTs at temperatures above 700 °C and inducing errors as large as 400 mK at 960 °C. Berry (1995) and earlier Zhang and Berry (1985) investigated the effect and demonstrated a complex dependence on a variety of influence variables, including:

- The electrical operating conditions, including any ground and screen configuration of the measurement circuit, grounding or screening of components in the fixed-point furnace, any dc polarising voltage, and the time that the SPRT is subjected to the polarising voltage.
- The structure of the thermometer, including the insulator material, the geometry of the insulator material and the platinum winding, and contact between the platinum winding and the insulator.
- Thermal operating conditions including the temperature distribution along the thermometer sheath and the thermal history.

There are also peculiar effects, similar to the electrical charge and discharge of batteries, associated with the insulation [Berry 1995, Moiseeva 2005]. White *et al.* (2007) and Yamazawa *et al.* (2007) showed that most of the observed complexity is explained by the influence of metal-semiconductor diodes (also known as Schottky-barrier or point-contact diodes) formed at the points of contact between the platinum and the silica insulators in the SPRT. It also seems likely that the conduction in fused silica is ionic rather than electronic, being due to impurities, which explains the battery-like effects. In practice, the leakage effect is far more complicated than a single conductance in parallel with the SPRT resistance. There are conductances distributed between each of the leads along the full length of the thermometer, each subject to different voltages and temperature profiles. Additionally, there are shunt resistances and voltage differences between the SPRT and the furnace.

Evans (1984) estimated the magnitude of both internal and external (through the sheath) leakage, and showed that guarding helps suppress some leakage resistance effects for several major models of thermometer. Berry (1995) showed that a dc offset of the correct polarity (approximately +6.4 V between thermometer wires and ground), applied to the measurement circuit or to conducting screens, allows to reduce significantly the conductance of the shunt (the resistance is increased). Some resistance bridges have the facility for introducing the offset. Recent experiments [Widiatmo *et al.* 2013] suggest this may be a way of switching the platinum-silica diodes off so that the leakage resistance effect can be measured.

Measurements of the shunt resistance by many workers, e.g. [Pokhodun *et al.* 1990, Berry 1995, Yamazawa and Arai 2003, 2005, Moiseeva 2005], suggest that typical values for a lumped shunt resistance, in the absence of a dc bias, for thermometers with quartz insulation are in the range 0.5 M $\Omega$  to 20 M $\Omega$  at 960 °C, which produce leakage effects of 3 mK to 0.08 mK for a sensor with  $R(273.16\text{ K}) = 0.25\ \Omega$ . Note that this effect may increase or decrease following calibration, when the thermometer is in use, depending on the temperature profiles along the thermometer sheath. These observations suggest a standard uncertainty of the order of 1 mK at 960 °C for 0.25  $\Omega$  thermometers. For thermometers employing glass and/or mica insulation, the leakage conductance is much greater and becomes apparent at lower temperatures with errors of about 1 mK at 600 °C.

During the calibration of HTSPRTs at the silver freezing point, the diffusion of silver through quartz may lead to contamination of the platinum sensing element. Different attempts to utilize passive barriers (glassy carbon crucible, sapphire thermowell, platinum sheath) and dc electric fields to protect the HTSPRTs from silver have been described [Hill 2015].

## 5.2. Thermal effects arising in use

### 5.2.1. Conduction and immersion effects

Static temperature-measurement errors are caused by the sensing element of the SPRT not being in direct thermal contact with the object of interest. The temperature indication of the thermometer is in practice affected by the many thermal conductances between the sensor and neighbouring objects. These include the axial thermal conductances of the measuring leads and the sheath, which extend to room temperature for LSPRTs and HTSPRTs, and the radial thermal conductance, which is influenced by the thermal conductivity of the exchange gas and the sheath and the heat-transfer coefficient between the sheath and its environment. A low radial thermal conductance also means a high self-heating effect, see Section 5.3.3.

The thermometer immersion problem has been investigated theoretically in [White and Jongenelen 2010, Gaiser and Fellmuth 2013a and 2013b] and experimentally, for instance, in [White and Jongenelen 2010, Riddle *et al.* 1973]. It has been shown that after the whole sensing element is immersed in the temperature environment to be measured, the static temperature-measurement error decreases exponentially with further immersion. The immersion characteristic can be used experimentally to estimate the residual errors. A thermometer is sufficiently immersed when there is no detectable change in the indicated temperature with additional immersion in a constant-temperature environment.

For fixed-point measurements, the hydrostatic pressure effect must be considered (see Chapter 2 *Fixed Points*), the magnitude of which is given for the different fixed-point substances in Table 2 of the text of the ITS-90. Usually the results of measurements of a fixed-point

temperature are considered as acceptable, if the SPRT immersion curve corresponds to the linear relation, describing the effect of the hydrostatic pressure on the temperature of the solid-liquid interface, over a distance of at least (3–5) cm from the cell bottom. The uncertainty due to the immersion effect can be evaluated with the following equation [Strouse 2005]:

$$u^2(\delta T_{\text{SPRT}}) = \frac{1}{N-1} \sum_{i=1}^N \left[ T_{90}(h_i) - T_{90}(h_0) - \frac{dT_{90}}{dh}(h_i - h_0) \right]^2, \quad (28)$$

where  $N$  is the number of measurements of the immersion characteristic,  $h$  is the distance along the axis of the cell, and  $dT_{90}/dh$  is the hydrostatic-pressure coefficient given in the text of the ITS-90. The determination of an immersion curve requires high-accuracy measurements of very small temperature differences. It is preferable to make measurements at two measuring currents with subsequent extrapolation of the results to the zero power in order to eliminate the influence of changes in the heat-exchange conditions when moving the SPRT in the re-entrant well of the cell. It may be preferable to make measurements on insertion of the SPRT, rather than on its withdrawal, because on withdrawing the SPRT the heat transfer fluid (air, water, oil) flows down into the measurement zone, with potentially harmful thermal effect, whereas expelling the fluid on insertion does less harm (though any vertical movement will create some disturbance to the temperature profile along the stem, which must be allowed to subside).

The influence of poor thermometer immersion on fixed point measurements can also be assessed by applying to the furnace a small,  $\sim 1$  K, temperature oscillation with a period of at least 30 min while observing the plateau [Fahr and Rudtsch 2008]. The amplitude of the observed fluctuations in the plateau temperature provides a direct measure of the coupling between the SPRT and the furnace. Note that the value of the thermal coupling constant varies throughout the melt or freeze of the cell [Fahr and Rudtsch 2008]. The difference between the furnace temperature and the fixed-point temperature can be determined from the monitoring SPRT once the cell has stabilised after the freeze.

The difference between the mean temperature of the sensing element and that of the solid-liquid interface depends on several factors: the depth of the thermometer immersion into the fixed-point cell, the presence of a continuous interface along the height of the fixed-point substance in the cell, the SPRT design, as well as heat exchange in the thermometer well and outside the cell. The influence of each of these factors can be evaluated by analysing the change of the thermometer immersion curve after a change in particular conditions during the fixed-point realisation. The depth of thermometer immersion into the cell is generally considered to be the main factor influencing the static temperature-measurement error. Above room temperature, the necessary depth increases with increasing fixed-point temperature up to about 450 °C. Then it decreases slightly as heat exchange by radiation improves. At higher temperature especially, longitudinal radiation baffles and measures to inhibit radiation piping through the sheath are essential, see Section 5.2.2. For different SPRT designs, the necessary immersion depth may differ by as much as 10 cm. Typically, about 20 cm to 25 cm of immersion (in addition to the length of the sensing element) in an isothermal zone is required to reduce the uncertainty component to 0.1 mK at 400 °C.

Collections of experimentally determined immersion curves are available within reports on Key Comparisons performed with LSPRTs at the TPW (CCT-K7 [Stock and Solve 2005, Stock *et al.* 2006]) and at the other fixed points in the range 84 K to 933 K (CCT-K3 [Mangum *et al.* 2002]). In several cases, the curves do not follow the linear behaviour corresponding to the hydrostatic-pressure coefficient, even near to the maximum possible immersion. These

departures increase the uncertainty estimate from Equation (28). The analysis of possible causes for such deviations is somewhat easier for the TPW because cells can be checked visually to confirm the thermometer well is completely surrounded by a solid-liquid interface and the temperature field around the cell is homogeneous along its height. The immersion curves of an SPRT were obtained for 23 different TPW cells with the immersion depth varying from 205 mm to 289 mm [Stock and Solve 2005], where the completeness of the interface was visually noticed. The analysis of immersion curves is more complicated for the other fixed points because the position of the interface is invisible and it is difficult to control the homogeneity of the temperature field along the cell and around the thermometer sheath. The following tests could be helpful:

- Improvement of the thermal contact between the thermometer sheath and the re-entrant well by introducing sleeves or contact liquids in the gap.
- Check of the influence of the temperature around the cell, e.g. by adjusting the temperature periodically, which may allow corrections for disturbing influences [Fahr and Rudtsch 2008]. Large influences may be caused by thermal links between the thermometer well and the outer wall of the cell. In the beginning of a freeze such links may consist of connecting spaces filled with liquid fixed-point substance and at the end of the freeze through spaces filled with solid substance [White and Mason 2011].
- Application of different methods for initiating an inner solid-liquid interface around the thermometer well, on which the completeness of the interface depends [Ivanova *et al.* 2013, White and Mason 2011]. In particular, it depends on the starting point of the initiation after observation of supercooling, on the duration of the cooling with a rod inserted in the thermometer well, and on the thermal conductivity and heat capacity of the rod that is used. A complete outer interface at the wall of the cell forms gradually. The time necessary for this process is dependent on the temperature around the cell.
- Check if thermometers that have transparent (glass, quartz or sapphire) sheaths are protected from intrusive, or escaping, radiation. (Metal sheathed thermometers need no further radiation shielding, but the heat conduction down the sheath is usually such as to require deeper immersion.)

The experimental immersion curve is in general not a reliable estimate of the temperature profile along the axis of the thermometer well because a displacement of the thermometer leads to a change of the profile due to its influence [Ilin 2003]. However, the slope of the immersion curve is an indicator of the closeness of the measurement conditions to the thermal equilibrium between the solid-liquid interface and the thermometer. During the measurements, not only the thermometer immersion depth must be taken into account, but also the design of the furnace or bath and the chosen modes of its operation.

A computer simulation of the heat exchange in a zinc cell during the phase transition has shown that lifting the LSPRT by 10 cm may lead to a change in the temperature in the lower end of the thermometer well by 0.7 mK [Batagelj 2004, Batagelj *et al.* 2005]. Another simulation yielded a relationship between the change in the temperature of the upper heater of the furnace and the temperature profile along the axis of the thermometer well at the freezing points of tin and zinc [Ivanova and Ilin 2005]. Furthermore, the influence of the position and the dimensions of the sheath area, over which the surface is roughened to decrease radiation effects, see Section 5.2.2, on the heat removal along the thermometer was estimated. This allowed assessment of the difference between the interface temperature and the temperature detected by the sensing element of an LSPRT [Batagelj 2004, Batagelj *et al.* 2005].

In capsule-type SPRTs, there is often an additional effect due to background heat fluxes, and static temperature-measurement errors are caused by the heat flow  $\phi$  via the sensing

electrical leads and the thermometer [Gaiser and Fellmuth 2013a]. To evaluate the errors, two thermometer parameters are needed: the thermal resistance  $R_T$  of the CSPRT between the glass seal at the thermometer head and the sheath, and the so-called reduction factor  $V = (T_T - T_S)/(T_H - T_S)$  ( $T_T$  mean temperature of the sensor element,  $T_S$  sheath and  $T_H$  head temperature). These parameters have to be determined in dedicated experiments. The error caused by the heat flow is given by  $T_T - T_M \approx V\phi R_T$  since the difference between the temperature of the measuring object  $T_M$  and  $T_S$  can usually be made vanishingly small.

### 5.2.2. Radiation effects

A significant source of error, especially at high temperatures, is heat flux to and from the platinum coil, via thermal radiation. If the sensor is in line of sight of a surface that is appreciably hotter or colder than that being measured, the power gained or lost by thermal radiation will result in a change of its temperature. In TPW measurements, thermal radiation from incandescent lights in the room, which may be incident upon the emerging thermometer sheath or the inner wall of the triple point cell, can produce an error of up to 0.2 mK [McLaren and Murdock 1966, Riddle *et al.* 1973]. The TPW cell should, therefore, be immersed in a bath in which no extraneous radiation from sources above room temperature can reach the sensor of the thermometer. If this precaution is omitted, even with radiation entry restricted to the top of the cell, a thermometer may require an immersion below the vapour-liquid level of 35 cm (which is seldom available), much more than the immersion of 15 cm for a thermometer with excellent immersion characteristics, in order for it to achieve an accuracy approaching 0.05 mK. For high-resolution measurements, such as TPW cell comparisons, opaque radiation shields can be used to cover the SPRTs and the top of the cells to confirm the absence of any effect.

At higher temperatures, radiation escaping via the SPRT sheath will cool the sensor element and can lead to errors as large as a few tenths of a millikelvin at 420 °C (zinc point) rising to several tens of millikelvin at 962 °C (silver point). This heat loss is due to “light piping” through the sheath due to total internal reflections within the wall of the thermometer sheath. This error can be substantially reduced by roughening the lower part of thermometer sheath from just above the sensor for some 15 cm to 20 cm by sand blasting, or by coating this part of the sheath with graphite paint [Widiatmo *et al.* 2006, Žužek *et al.* 2014]. The same heat-loss mechanism can affect the apparent temperatures of fixed point cells that have fused silica thermometer wells. These are less easily dealt with by sand blasting or painting but it is a fairly simple matter to arrange that they ‘see’ only opaque surfaces substantially at their own temperature, and not suffering net loss or gain of heat.

## 5.3. Resistance measurement

Resistance measurements in platinum thermometry are usually made using automatic low-frequency resistance bridges. The reading of the bridges is typically a dimensionless ratio  $X = R(T_{90})/R_S$ , where  $R(T_{90})$  is the resistance of the SPRT under the current conditions of measurement, and  $R_S$  is the resistance of a calibrated reference or standard resistor also connected to the bridge. The bridges all employ a 4-terminal-coaxial or guarded 4-terminal definition of resistance [Awan *et al.* 2011] to reduce the effect of lead resistances and stray impedances. The major sources of uncertainty associated with the resistance measurements include the reference resistor value and its stability, and self heating of the SPRT due to the



sensing power. There are additional minor effects associated with the connecting cables and the resistance bridge itself.

### 5.3.1. Resistance bridges

There are two basic types of thermometry resistance bridges in common use: ac bridges using a sinusoidal sensing current in the frequency range 10 Hz to 90 Hz, and so-called dc bridges, which use a low-frequency square-wave (ac) sensing current, in the range 0.01 Hz to 10 Hz. The bridges typically have a resolution of six to nine digits and a specified uncertainty in resistance ratio between  $2 \times 10^{-8}$  and  $4 \times 10^{-6}$ , corresponding to equivalent uncertainties in temperature measurements near the TPW ranging from about 5  $\mu$ K to 1 mK. Some of the sources of error within thermometry bridges and bridge calibration methods are summarized in [White 2003]. Rudtsch *et al.* (2005), Strouse and Hill (2003), and White (1997) give additional information on typical performance of some resistance bridges. The errors in bridges can be classified as noise, large scale or ‘integral’ non-linearities (INL), and short-range or ‘differential’ non-linearities (DNL).

The resolution of a bridge is determined by a combination of the thermal noise generated in all resistances and amplifiers, and quantisation error due to the conversion of the signal to a digital reading [White and Edgar 2013]. The resolution is readily measured as the standard deviation when measuring a single stable resistance. Uncertainty due to resolution can be reduced by decreasing the operating bandwidth of the bridge or, equivalently, increasing the measurement time or averaging many results.

Noise in the resistance measurement may also arise from electromagnetic interference (EMI), particularly electrical noise coupled from ac furnace controllers, electric motors, and other ac resistance bridges operating nearby on the same carrier frequency. In a bridge with resolution limited by thermal noise, the standard deviation of measurements will change as the square root of the bandwidth or measurement time. If EMI due to ac machinery is present then often the standard deviation may change more rapidly or not at all. Digital interface connections between the bridge and a computer for data acquisition can also cause interference which arises from current flow caused by differing ground potentials between the two devices; this can be overcome by using optical interface connections.

INL are those non-linearities that change slowly and smoothly over a wide range of bridge readings. These effects are caused by finite input and output impedances, power coefficients of resistors, finite gains of amplifiers, and stray capacitance and resistance of sub-circuits of the bridge. They tend to be well modelled by quadratic or cubic functions of bridge reading. With many bridges, the INL effects are below the manufacturers’ accuracy specification and, therefore, less significant than many other sources of uncertainty in the temperature measurement. INL errors larger than the bridge manufacturer’s specification are usually indicative of a faulty sub-circuit within the bridge or poor adjustment. Additionally, because the ITS-90 is defined in terms of resistance ratio, only the non-linear bridge errors (offsets, quadratic, and cubic terms) propagate and affect the temperature measurement.

Further, if the same bridge is used for the calibration and use of an SPRT, then the SPRT interpolating equations largely correct the INL errors of the bridge. In such cases the INL correction and uncertainty can be omitted from the uncertainty analysis.

DNL are commonly caused by errors in the transformer windings or analogue-to-digital converters. They may occur at every change in reading and are generally too complex to measure, model and correct. The effects tend to be below the manufacturers’ specifications,

but can increase with time and mechanical vibration as transformer cores lose magnetic permeability, and ADC circuits drift out of adjustment. DNL are the dominant source of uncertainty in resistance measurements for high resolution or differential measurements (e.g., fixed-point comparisons and self-heating measurements). The magnitude of the resulting uncertainty can be inferred from the standard deviation of residual error in the calibration curves for bridges.

### 5.3.2. Reference resistor

The two main contributions to the uncertainty in the value of the standard resistance,  $R_S$ , are caused by the calibration uncertainty (the value of the resistance) and the temperature dependence of  $R_S$ . The effects can be evaluated using a model of the temperature dependence of the resistance:

$$R_S(T_{\text{bath}}) = R_S(T_{\text{cal}})[1 + \beta(T_{\text{bath}} - T_{\text{cal}})] \quad (29)$$

where  $T_{\text{bath}}$  is the temperature of the oil or air bath used to maintain the resistor during use,  $T_{\text{cal}}$  is the temperature of the bath used to maintain the resistor when it was calibrated, and  $\beta$  is the temperature coefficient of the resistor at  $T_{\text{cal}}$ .  $\beta$  is normally expressed as the fractional change per degree; i.e.,

$$\beta = \frac{1}{R_S} \frac{dR_S}{dT_{90}}, \quad (30)$$

with typical values for good-quality resistors within the range  $\pm 5 \times 10^{-6}/^\circ\text{C}$ . If the resistor is not maintained at its calibration temperature, then a correction can be applied using Equation (29). The cumulative uncertainty in the resistance value due to these terms is

$$u^2(R_S) = u^2(R_{S,\text{cal}}) + R_S^2[(T_{\text{bath}} - T_{\text{cal}})^2 u^2(\beta) + \beta^2 u^2(T_{\text{bath}})] \quad (31)$$

where  $u(R_{S,\text{cal}})$  is the calibration uncertainty as supplied by the electrical calibration laboratory,  $u(T_{\text{bath}})$  is the uncertainty arising from fluctuations in the resistor-bath temperature, and  $u(\beta)$  is the uncertainty in the temperature coefficient of the resistor. Where  $T_{\text{bath}}$  and  $T_{\text{cal}}$  are nominally equal, the uncertainty in the temperature coefficient can usually be ignored, as can any quadratic temperature dependence of  $R_S$  in Equation (29). Also, where the same resistance bridge is used for the two resistance measurements used to calculate the  $W(T_{90})$  value (Equation (1), see p. 6), uncertainty in the value of  $R_S$  can be ignored and only the stability of  $R_S$  needs be considered. The uncertainty in the  $R_S$  does need to be considered where the  $R(0.01^\circ\text{C})$  value of a thermometer is reported.

The calibration value for the standard resistor,  $R_{S,\text{cal}}$ , is nearly always the dc value. If the resistor is to be used with an ac bridge, standard resistors with a low ac-dc difference should be used [Wilkins and Swan 1970]. For frequencies below 100 Hz, ac-dc differences arise from stray inductance and capacitance, dielectric absorption, and eddy current effects. Errors in dc and low frequency measurements can arise from thermoelectric EMFs produced as a result of Peltier heating at the connections [Kirby and Laubitz 1973], but these are usually small. Some standard resistors exhibit relative ac-dc differences as large as  $10^{-5}$ . With resistors of the Wilkins' design, the relative uncertainty due to ac-dc difference,  $u(\delta R_{S,\text{ac-dc}}/R_S)$ , see Table 3 in Section 6.1.1, is below  $10^{-7}$ .

In bridges based on dc current comparators (see for example [McMartin and Kusters 1966]), the current  $i_0$  through the standard resistor changes with bridge reading. This causes variable

power dissipation in the standard resistor and an error with a cubic dependence on the measured resistance ratio,  $X = R(T_{90})/R_S$ :

$$\delta X_{\text{power}} = -i_0^2 R_S \left( \frac{R(T_{90})}{R_S} \right)^3 \left( \frac{1}{R_S} \frac{\partial R_S}{\partial P} \right) = -i_0^2 R_S X^3 \left( \frac{1}{R_S} \frac{\partial R_S}{\partial P} \right), \quad (32)$$

where  $P$  is the power dissipated in the standard resistor, and the term involving the derivative is the power coefficient. Electrical calibrations of standard resistors are normally carried out at a specified power. The power coefficient is the product of the temperature coefficient and the thermal resistance between the resistor windings and the environment, so good quality standard resistors are generally quite bulky devices. The Wilkins' type resistors [Wilkins and Swan 1970] have a power coefficient of less than  $4 \times 10^{-6}/\text{W}$ . The error is then much less than  $1 \mu\text{K}$ . But this error can be considerably larger if a purpose-built standard resistor is not used. If sufficient time is allowed for the resistor to stabilise after current changes (tens of minutes), the effect is compensated by the self-heating correction for the SPRT.

Considering also the uncertainty components due to ac-dc differences and the power dissipation, from Equation (31) it follows:

$$u^2(R_S) = u^2(R_{S,\text{cal}}) + R_S^2 (T_{\text{bath}} - T_{\text{cal}})^2 u^2(\beta) + \beta^2 u^2(T_{\text{bath}}) + u^2(\delta R_{S,\text{ac-dc}}) + u^2(\delta R_{S,\text{power}}) \quad (33)$$

### 5.3.3. Self heating

Measurements with resistance thermometers necessarily involve passing a current through the thermometer sensing element, which gives rise to Joule heating. As a result, the sensing element warms to a slightly higher temperature than that of the object or environment whose temperature is to be measured. The warming continues until the temperature difference is sufficient to dissipate the heat and the steady state is reached. Considering SPRTs, most of the heat flows by conduction through the filling gas to the sheath, though there is some convection and also conduction through solid components. This internal so-called “self-heating” is fixed by the design of the thermometer, but there is also an external effect between the sheath and the heat sink (for example, the liquid-solid interface in a fixed-point cell) [Riddle *et al.* 1973], which in a well-designed experiment can be made relatively small.

The self-heating effect depends on the thermal conductance between the sensor and the measurement environment, and is therefore a useful indicator of whether good thermal contact has been achieved, and subsequently for routine checks. Influencing factors are the design of the sensor element, the thermal conductivity of the exchange gas, and the heat transfer to the heat sink, i.e. the solid-liquid interface of the fixed-point material. Thus, the magnitude of the effect has to be checked individually for each experimental setup. It is not alone determined by the parameters of the SPRT. The measurement current must be chosen so as to achieve good resolution without excessive self heating. For  $25 \Omega$  SPRTs the current is often  $1 \text{ mA}$  (dissipation of  $25 \mu\text{W}$  at  $0^\circ\text{C}$ ) and the self heating effect in a TPW cell is typically between  $0.2 \text{ mK}$  and  $4 \text{ mK}$ . With modern resistance bridges, a temperature resolution of  $< 0.1 \text{ mK}$  is easily achieved. Self heating has usually a typical variation with temperature, showing a maximum slightly above  $200^\circ\text{C}$ .

The self-heating effect is determined by making measurements at two currents and calculating the temperature rise from the change in measured resistance. The user then has a choice of whether or not to apply a correction for the effect. If a thermometer is measured using the same current as was used in its calibration, and if the external conditions for the SPRT are also similar, then the effect will be similar and the thermometer can be used without making

corrections. This is convenient and is common practice in secondary-level calibrations and measurements.

For precise work, self-heating corrections must be applied [Batagelj *et al.* 2003a, 2003b]. Since the power dissipated increases as the square of the current, the resistance at zero current,  $R_0$ , can be calculated from resistance measurements  $R_1$ , and  $R_2$  made using measuring currents  $i_1$  and  $i_2$  using the following relation:

$$R_0 = \frac{R_2 i_1^2 - R_1 i_2^2}{i_1^2 - i_2^2}. \quad (34)$$

Commonly, the current ratio  $i_2/i_1$  is chosen to be  $\sqrt{2}$ . In that case a simple measurement sequence  $i_1: i_2: i_1$  allows the zero-current resistance to be determined from the corresponding measurements  $R_1: R_2: R_3$  as  $R_0 = R_1 + R_3 - R_2$ . Pearce *et al.* (2012, 2013) have shown that the uncertainty in the extrapolation to zero current is minimised if the current ratio is 2 and the measurement time is apportioned in the ratio 4:1:4.

The zero-current bridge ratio  $X_{\text{meas}}^0$  is calculated corresponding to Equation (34). Its uncertainty can be estimated using the equation:

$$\begin{aligned} u^2(X_{\text{meas}}^0) = & \frac{i_2^4 + i_1^4}{(i_2^2 - i_1^2)^2} [u^2(\delta X_{\text{DNL}}) + 2u^2(\delta X_{\text{noise}}) + u^2(R_S)] \\ & + \frac{2i_1^4 i_2^4}{(i_2^2 - i_1^2)^4} [X_{\text{meas}}^2 - X_{\text{meas}}^1]^2 \left[ \frac{u^2(i_1)}{i_1^2} + 2\frac{u(i_1, i_2)}{i_1 i_2} + \frac{u^2(i_2)}{i_2^2} \right] \end{aligned} \quad (35)$$

In Equation (35),  $u(\delta X_{\text{DNL}})$  is the uncertainty component due to the DNL error between the measurements of  $X_{\text{meas}}^1$  and  $X_{\text{meas}}^2$  with the two currents  $i_1$  and  $i_2$ ,  $u(\delta X_{\text{noise}})$  is the component due to electrical noise, which is assumed to have the same order of magnitude for both currents, and  $u(i_1, i_2)$  is the estimated covariance associated with  $i_1$  and  $i_2$ . If the relative errors of the two currents are equal, i.e. if they are fully correlated, the second term vanishes.

### 5.3.4. Connecting cables and lead resistances

Figure 4 shows a simplified equivalent circuit for the measurement of a 4-terminal resistance, which includes the finite shunt resistance of the connecting cables. The model can also be used to characterise the input and output impedances of the bridge, hence the following nomenclature:  $R_{\text{out}}$  is the resistance shunting the current source  $i_0$ , and  $R_{\text{in}}$  is the resistance shunting the voltage measurement  $V_m$ . The equivalent circuit also includes the lead resistances  $R_L$  in each of the four leads to the measured resistance. Analysis of Figure 4 shows that the error in the measured resistance ratio (the bridge reading),  $X$ , is well approximated by

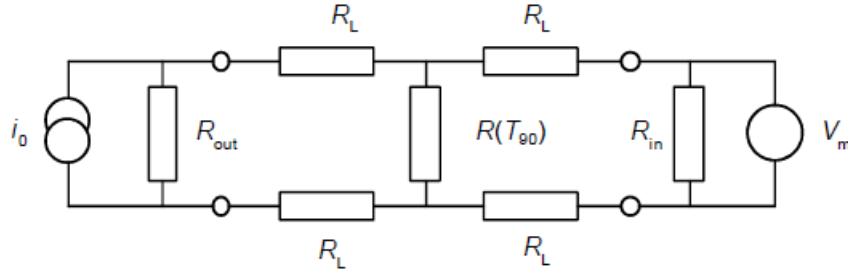
$$\delta X_{\text{lead}} = \frac{1}{R_S} \left( \frac{V_m}{i_0} - R(T_{90}) \right) \approx - \left( \frac{R(T_{90})}{R_S} \right)^2 \frac{R_S}{R_{\text{eff}}} - \left( \frac{R(T_{90})}{R_S} \right) \frac{R_L}{R_{\text{eff}}} = -X^2 \frac{R_S}{R_{\text{eff}}} - X \frac{R_L}{R_{\text{eff}}}, \quad (36)$$

where

$$R_{\text{eff}} = \frac{R_{\text{out}} R_{\text{in}}}{R_{\text{out}} + R_{\text{in}}}.$$

Thus, the finite resistances,  $R_{\text{out}}$  and  $R_{\text{in}}$ , give rise to an error proportional to the square of the resistance ratio. With care, cable shunt resistances can normally be maintained at greater than  $10^{10} \Omega$ , ensuring the effects are negligible. Significant effects can be observed in very long cables, or unscreened cables immersed in oil (the standard resistor bath), and with some

coloured PVC cables. With ac bridges, dielectric loss in the cables gives rise to a similar error that increases as the square of the resistance and increases linearly with frequency. Dielectric loss can be a major problem in low-temperature measurements where many meters of lead wire is used for thermal anchoring. Operating an ac bridge at lower frequencies or using a dc bridge can significantly reduce the effects. The effects of lead resistances are usually negligible in comparison to the effects of the finite cable shunt resistance since  $R_L \leq R_S$  in most resistance measurements.



**Figure 4 — The equivalent circuit for a 4-terminal resistance measurement.**

In ac bridges, the measurements can also be affected by the stray capacitance of the connecting cables. Any capacitance in parallel with  $R(T_{90})$  (Figure 4, see p. 37) causes the impedance ‘seen’ by the resistance bridge to be

$$Z = \frac{R(T_{90})}{1 + \omega^2 R(T_{90})^2 C^2} - \frac{j\omega R(T_{90})C}{1 + \omega^2 R(T_{90})^2 C^2}, \quad (37)$$

where  $\omega = 2\pi f$  is the angular frequency,  $C$  is the cable capacitance (about 100 pF/m for coaxial and twisted pair cable), and  $j = \sqrt{-1}$  represents a 90° phase shift. The real part of Equation (37) is the resistance seen by an ideal ac thermometry bridge; the imaginary part contributes to the quadrature component of the bridge signal, which must be balanced or rejected by the bridge. The real part of Equation (37) contributes a resistance ratio error with a cubic dependence on the measured resistance ratio:

$$\delta X_{\text{cap}} = \frac{\text{Re}(Z(T_{90}) - R(T_{90}))}{R_S} \approx -\omega^2 C^2 R_S^2 \left( \frac{R(T_{90})}{R_S} \right)^3 = -\omega^2 C^2 R_S^2 X^3. \quad (38)$$

Normally, the effect of stray capacitance is negligible. However, because of the square-law dependence on capacitance, in measurement systems with multiplexers and long leads, the capacitance may overload the quadrature-balance capability of the bridge or lead to relative errors in resistance ratio as large as  $10^{-4}$  (many millikelvin). AC bridges used at low temperatures may also be affected by the self-inductance of any long high-resistance lead wires used for thermal anchoring. In principle, the measured resistance is independent of the inductance (unlike the case for capacitance). However, the large quadrature signal can overload the bridge leading to excessive noise.

For dc resistance bridges the effect of stray capacitance is complicated by the many harmonics in the ‘square-wave’ sensing current, but normally below the limit of detection. Where detectable, it is apparent with changes in the current-reversal times and settling times [Zhang and Berry 1985].

In systems with short leads, most cable effects can be reduced to negligible levels with care. If the effects are significant they are often indicative of fault conditions.

## 6. UNCERTAINTY IN SPRT CALIBRATIONS

The estimation of the total uncertainty of a temperature value  $T_{90}$  realised with an SPRT (Section 6.4, see p. 50) starts with the estimation of the uncertainty of the calibration at the fixed points (Section 6.1, see p. 38). But for a realisation at any temperature, it is necessary to propagate the calibration uncertainty (Section 6.2, see p. 40) and consider the non-uniqueness of the scale realisation due to interpolation errors (Section 6.3, see p. 44). The estimation for practical measurements starts with the calibration uncertainty as basic component. It is complex due to the variety of sub-ranges with different interpolation equations, the option to use either single or multiple TPW measurements, possibly with different TPW cells, and the options for the use of different resistance bridges or standard resistors for different resistance measurements. It requires, furthermore, to consider the SPRT effects discussed in Subsection 5 because usually a separate TPW measurement cannot be performed for each single temperature determination. Since the practical application of SPRTs is outside the scope of this guide, here only the most important cases routinely met during calibration of SPRTs are covered. For cases not covered here, and for detailed explanations and derivations, see [White *et al.* 2018]. In all cases, the necessary corrections and measuring errors are sufficiently small for estimating the uncertainty using a linear mathematical model.

Some subsections below provide graphs giving numerical values for some of the uncertainty contributions and the total uncertainty. The values in these graphs are purely indicative, and are given only to illustrate the approximate magnitude and temperature dependence of the various contributions. Users are expected to determine uncertainties appropriate to their measurements and use these in the equations.

[Appendix 3](#) gives values for the range of some of the contributions to the uncertainty in fixed-point measurements. These values are also indicative. Most of the ranges encompass the uncertainties reported from recent measurement comparisons, as well as the expected extremes determined from the application of the models given in previous subsections. (For comparisons involving the TPW, see Stock and Solve (2005), Renaot *et al.* (2005). For comparisons involving low temperature fixed points ( $< 0^\circ\text{C}$ ) and SPRTs, see Steele *et al.* (2002), Fellmuth *et al.* (2012), Head *et al.* (1997), Pavese *et al.* (1984). For comparisons involving high-temperature fixed points ( $> 0^\circ\text{C}$ ) and SPRTs, see Mangum *et al.* (2002), Nubbemeyer and Fischer (2002).)

### 6.1. Uncertainty of SPRT calibration at the fixed points

#### 6.1.1. Uncertainty in SPRT resistance at the triple point of water

The main purpose of the TPW resistance value  $R_{\text{TPW}}$  reported on a calibration certificate is to enable users of the SPRT to assess the stability of the SPRT and detect major changes that may have occurred through shipping, or to detect damage that may have occurred during use. For this purpose, it is best if the SPRT is in its most reproducible state, i.e. the SPRT is well annealed and heat-treated so that it is strain-, vacancy-, and oxide-free. The  $R_{\text{TPW}}$  value in this state may be different from that used for the calculation of resistance ratios or temperature. The mathematical model for estimating the uncertainty of  $R_{\text{TPW}}$  is given by:

$$R_{\text{TPW}} = R_{\text{meas}} \left( 1 + \frac{dW_{\text{r}}}{dT_{90}} \delta T_{\text{FP}} \right) \quad (39)$$

with the measured resistance value

$$R_{\text{meas}} = R_S (T_{\text{cal}}) [1 + \beta (T_{\text{bath}} - T_{\text{cal}})] [X_{\text{meas}}^0 + \delta X_{\text{INL}}] \quad (40)$$

and the necessary correction of the realised fixed-point temperature

$$\delta T_{\text{FP}} = \sum_{k=1}^8 \delta T_{\text{FP}}^k \quad (41)$$

The individual corrections  $\delta T_{\text{FP}}^k$  are explained in Table 3. Most of them were already discussed in [Fellmuth *et al.* 2001, 2005, Bonnier *et al.* 2005].

The total Type B uncertainty in the zero-current TPW resistance, using sensing currents  $i_1$  and  $i_2$ , is

$$u^2 (R_{\text{TPW}}) = R_S^2 u^2 (X_{\text{meas}}^0) + R_S^2 u^2 (\delta X_{\text{INL}}) + (X_{\text{meas}}^0)^2 u^2 (R_S) + \left( R_{\text{meas}} \frac{dW_r}{dT_{90}} \right)^2 \sum_{k=1}^8 u^2 (\delta T_{\text{FP}}^k), \quad (42)$$

where the uncertainty of the zero-current bridge ratio  $X_{\text{meas}}^0$  is given by Equation (35) in Section 5.3.3 and that of the standard resistor  $R_S$  by Equation (33) in Section 5.3.2.

**Table 3.** Summary of Type B uncertainty components (Note: “No.  $k$ ” indexes the terms of Equation (41) and [Appendix 3](#)).

No. $k$	Symbol	Description of the correction	Reference
1	$\delta T_{\text{hyd}}$	Hydrostatic head correction	Section 5.2.1
2	$\delta T_p$	Gas pressure correction	ITS-90 text, Table 2
3	$\delta T_{\text{imp}}$	Influence of impurities	Chapter 2, <a href="#">Section 2.1</a>
4	$\delta T_{\text{iso}}$	Influence of the isotopic composition	<i>MeP-K</i> , Technical Annex
5	$\delta T_{\text{cry}}$	Influence of crystal defects and strain	Chapter 2
6	$\delta T_{\text{stat}}$	Static thermal effects	Section 5.2
7	$\delta T_{\text{dyn}}$	Dynamic thermal effects	Section 5.2
8	$\delta T_{\text{liq}}$	Extrapolation to the liquidus point	Chapter 2, Section 2.1
	$\delta X_{\text{INL}}$	Integral resistance bridge non-linearities	Section 5.3.1
	$\delta X_{\text{DNL}}$	Differential resistance bridge non-linearities	Section 5.3.1
	$\delta R_{\text{Sac-dc}}$	ac-dc difference of the reference-resistor value	Section 5.3.2
	$\delta R_{\text{Spower}}$	Power dependence of the reference-resistor value	Section 5.3.2

### 6.1.2. Uncertainty in SPRT resistance ratios at fixed points

In general, when measuring the resistance at the gallium, indium, tin, or zinc points, it is not possible to keep the SPRT in an oxide-free state, see Section 5.1.1. If the SPRT is in the same oxidation state for the measurements at fixed point  $i$  and the TPW ( $i = 1$ ), the uncertainty of the resistance ratio  $W_i$  is given by

$$u^2 (W_i) = \frac{1}{R_{\text{TPW}}^2} [u^2 (R_i) + W_{r,i}^2 u^2 (R_{\text{TPW},i}^2)] \quad (43)$$

where  $R_{\text{TPW},i}$  is the corresponding TPW value. If furthermore both measurements are performed with the same reference resistor under stable conditions (negligible short-term

instability), the Type B uncertainty in the SPRT resistance for each fixed point can be estimated applying the following equation, which considers correlations:

$$u^2(R_i) = R_S^2 u^2(X_{\text{meas},i}^0) + R_S^2 u^2(\delta X_{\text{INL}}) + \left(R_{\text{TPW}} \frac{dW_{r,i}}{dT_{90}}\right) \sum_{k=1}^8 u^2(\delta T_{\text{FP},i}^k) \quad (44)$$

Under these conditions, the uncertainty of  $R_{\text{TPW},i}$  is also given by Equation (44) ( $i = 1$ ).

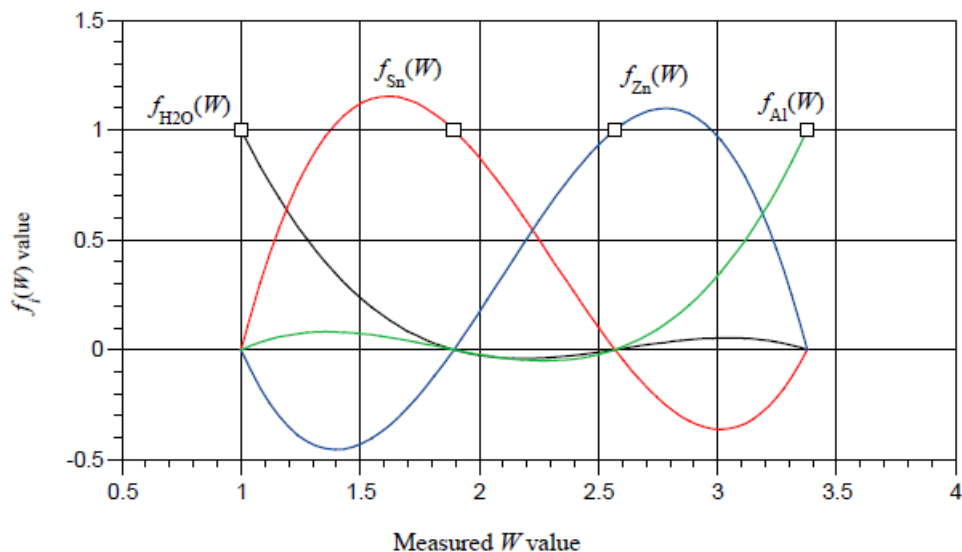
## 6.2. Propagation of calibration uncertainty

The many interpolating equations of the ITS-90 greatly complicate the propagation of uncertainty for temperatures measured using SPRTs. Complications also arise from the option to use either single or multiple TPW measurements, possibly with different TPW cells, and the options for the use of different resistance bridges or standard resistors for different resistance measurements. In principle, each combination of these possibilities results in a different equation for total uncertainty. Further, the complexity of ITS-90 is such that it is generally impractical to obtain correct expressions for total uncertainty without some supporting mathematics. This subsection covers the two cases routinely met during calibration of SPRTs: LSPRTs/HTSPRTs and CSPRTs. For cases not covered here, and for detailed explanations and derivations, see [White *et al.* 2018].

### 6.2.1. Fixed-point sensitivity coefficients and propagation equation without correlations

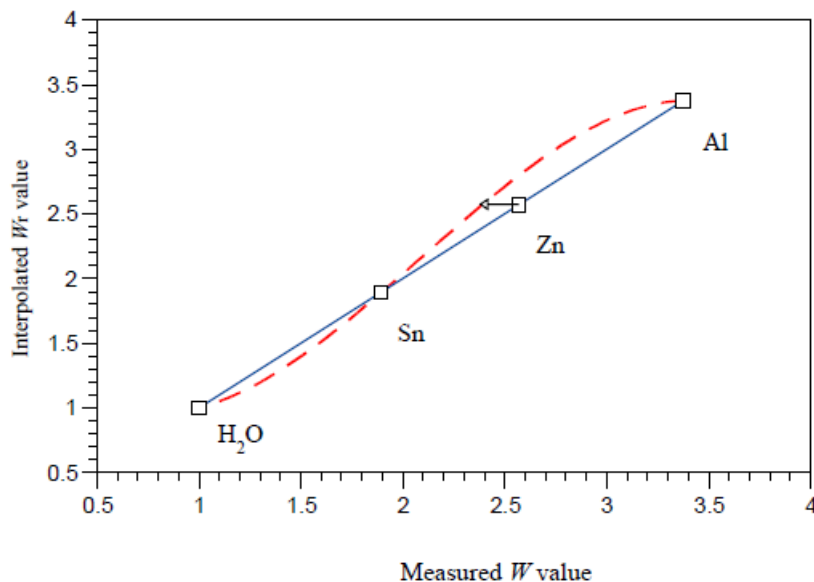
In order to estimate the uncertainty in SPRT realisation of the ITS-90 between the fixed points, it is necessary to calculate how errors in the fixed-point measurements, made during the SPRT calibration, propagate to other temperatures in the sub-range. The derivation of the propagation functions can be demonstrated for the example from Subsection 2 applying the alternative interpolation Equation (11) for the water-aluminium sub-range. Figure 5 plots the four functions,  $f_i(W)$ , of Equation (12a-d). Each function takes the value 1.0 exactly at the fixed point, for which it is named, and is zero at the other fixed points. Note that each of the interpolating functions is a cubic function of  $W$ , and any linear combination of them, such as Equation (11), is therefore also cubic.





**Figure 5 — The four ITS-90 interpolating functions for the water-aluminium sub-range.**

We can illustrate the effect of the fixed-point sensitivity coefficients by comparing two SPRT calibration curves. Figure 6 shows a graph of two water-aluminium interpolating equations. The solid curve passes through all four marked points. In the second, dashed curve, an error has been introduced into the measured zinc-point value,  $W_{\text{Zn}}$ . The dashed curve passes through the new point but continues to pass through the other original points. The difference between the two curves is one of the four sensitivity coefficients we seek, and is equal to the  $f_{\text{Zn}}(W)$  function of Figure 5 multiplied by the error in the zinc-point measurement.



**Figure 6 — The influence of an error in a fixed-point measurement. The solid line shows an interpolation between the measured  $W$  values and the  $W_r$  values. The dashed line shows the effect of a small error in the measured  $W_{\text{Zn}}$  value. The difference between the two curves is proportional to the  $f_{\text{Zn}}(W)$  function.**

All of the ITS-90 interpolating equations can be expressed in a form similar to Equation (11): the sum Equation (17) of interpolating functions multiplied by the corresponding reference resistance ratios, where the numerical index  $i$  refers to the sequence of fixed points used to calibrate the SPRT (e.g. H<sub>2</sub>O, Sn, Zn, etc.). The index  $i = 1$  is reserved for the TPW. The  $f_i(W)$  are a different set of functions for each ITS-90 sub-range. Table 4 below itemises the functions for the most commonly used sub-ranges of the ITS-90. Algebraic expressions for the functions for each sub-range are described in detail in [Appendix 1](#) and [White \_et al.\_ 2018].

The interpolation Equation (17) appear simple, but using them directly for propagating uncertainty would be troublesome because the  $f_i(W)$  depend on all  $W_j$ . White and Saunders (2007) have deduced simple expressions for the sums of terms containing the many partial derivatives. Their propagation-of-uncertainty equation resulting from Equation (17) is given by:

$$\delta W_r \approx - \sum_{i=2}^N f_i(W) \delta W_{\hat{p}} \quad (45)$$

which shows again that the  $f_i(W)$  functions are the sensitivity coefficients for errors in the fixed-point resistance ratios. The sum omits the term with the index  $i = 1$ , which corresponds to the TPW. Because the TPW measurements are used to compute all resistance ratios, we need to break Equation (45) down so that it separates the resistance measurements. This is done by repeatedly applying the identity

$$\delta W = \frac{1}{R_{\text{H}_2\text{O}}} (\delta R - W \delta R_{\text{H}_2\text{O}}), \quad (46)$$

which follows from the definition of  $W$ . This leads to the expression

**Table 4. The interpolating equations and fixed-point sensitivity functions for commonly used sub-ranges.**

Temperature sub-range	Interpolating equation for $W_r$ and interpolating functions
83.8058 K to 273.16 K	$W_r = f_{\text{H}_2\text{O}} + W_{r,\text{Hg}} f_{\text{Hg}} + W_{r,\text{Ar}} f_{\text{Ar}}$ $f_{\text{H}_2\text{O}} = \frac{1 - f_{\text{Hg}} - f_{\text{Ar}}}{(W-1)(\ln W - \ln W_{\text{Hg}})}, f_{\text{Hg}} = \frac{(W-1)(\ln W - \ln W_{\text{Ar}})}{(W_{\text{Hg}}-1)(\ln W_{\text{Hg}} - \ln W_{\text{Ar}})},$ $f_{\text{Ar}} = \frac{(W_{\text{Ar}}-1)(\ln W_{\text{Ar}} - \ln W_{\text{Hg}})}{(W-1)(W-W_{\text{Ga}})}$
-38.8344 °C to 29.7646 °C	$W_r = f_{\text{Hg}} W_{r,\text{Hg}} + f_{\text{H}_2\text{O}} + f_{\text{Ga}} W_{r,\text{Ga}}$ $f_{\text{Hg}} = \frac{(W_{\text{Hg}}-1)(W_{\text{Hg}}-W_{\text{Ga}})}{(1-W_{\text{Hg}})(1-W_{\text{Ga}})}, f_{\text{H}_2\text{O}} = \frac{(W-W_{\text{Ga}})(W-W_{\text{Ga}})}{(1-W_{\text{Hg}})(1-W_{\text{Ga}})}, f_{\text{Ga}} = \frac{(W-W_{\text{Hg}})(W-1)}{(W_{\text{Ga}}-W_{\text{Hg}})(W_{\text{Ga}}-1)}$
0.01 °C to 29.7646 °C	$W_r = f_{\text{H}_2\text{O}} + f_{\text{Ga}} W_{r,\text{Ga}}$ $f_{\text{H}_2\text{O}} = \frac{(W-W_{\text{Ga}})}{(1-W_{\text{Ga}})}, f_{\text{Ga}} = \frac{(W-1)}{(W_{\text{Ga}}-1)}$
0.01 °C to 156.5985 °C	$W_r = f_{\text{H}_2\text{O}} + f_{\text{In}} W_{r,\text{In}}$ $f_{\text{H}_2\text{O}} = \frac{(W-W_{\text{In}})}{(1-W_{\text{In}})}, f_{\text{In}} = \frac{(W-1)}{(W_{\text{In}}-1)}$
0.01 °C to 231.928 °C	$W_r = f_{\text{H}_2\text{O}} + f_{\text{In}} W_{r,\text{In}} + f_{\text{Sn}} W_{r,\text{Sn}}$ $f_{\text{H}_2\text{O}} = \frac{(W-W_{\text{In}})(W-W_{\text{Sn}})}{(W-1)(W_{\text{In}}-1)(W_{\text{In}}-W_{\text{Sn}})}, f_{\text{In}} = \frac{(W-1)(W-W_{\text{Sn}})}{(W_{\text{In}}-1)(W_{\text{In}}-W_{\text{Sn}})},$ $f_{\text{Sn}} = \frac{(W_{\text{Sn}}-1)(W_{\text{Sn}}-W_{\text{In}})}{(W-1)(W_{\text{In}}-W_{\text{Sn}})}$
0.01 °C to 419.527 °C	$W_r = f_{\text{H}_2\text{O}} + f_{\text{Sn}} W_{r,\text{Sn}} + f_{\text{Zn}} W_{r,\text{Zn}}$ $f_{\text{H}_2\text{O}} = \frac{(W-W_{\text{Sn}})(W-W_{\text{Zn}})}{(W-1)(W_{\text{Sn}}-1)(W_{\text{Sn}}-W_{\text{Zn}})}, f_{\text{Sn}} = \frac{(W-1)(W-W_{\text{Zn}})}{(W_{\text{Sn}}-1)(W_{\text{Sn}}-W_{\text{Zn}})},$ $f_{\text{Zn}} = \frac{(W_{\text{Zn}}-1)(W_{\text{Zn}}-W_{\text{Sn}})}{(W-1)(W_{\text{Sn}}-W_{\text{Zn}})}$
0.01 °C to 660.323 °C	$W_r = f_{\text{H}_2\text{O}} + f_{\text{Sn}} W_{r,\text{Sn}} + f_{\text{Zn}} W_{r,\text{Zn}} + f_{\text{Al}} W_{r,\text{Al}}$ $f_{\text{H}_2\text{O}} = \frac{(W-W_{\text{Sn}})(W-W_{\text{Zn}})(W-W_{\text{Al}})}{(1-W_{\text{Sn}})(1-W_{\text{Zn}})(1-W_{\text{Al}})}, f_{\text{Sn}} = \frac{(W_{\text{Sn}}-1)(W_{\text{Sn}}-W_{\text{Zn}})(W_{\text{Sn}}-W_{\text{Al}})}{(W_{\text{Sn}}-1)(W_{\text{Sn}}-W_{\text{Zn}})(W_{\text{Sn}}-W_{\text{Al}})},$

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Temperature sub-range      Interpolating equation for  $W_r$  and interpolating functions

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$$f_{Zn} = \frac{(w-1)(w-w_{Sn})(w-w_{Al})}{(w_{Zn}-1)(w_{Zn}-w_{Sn})(w_{Zn}-w_{Al})}, f_{Al} = \frac{(w-1)(w-w_{Sn})(w-w_{Zn})}{(w_{Al}-1)(w_{Al}-w_{Sn})(w_{Al}-w_{Zn})},$$


---

$$\delta W_r = -\frac{1}{R_{H_2O}} \sum_{i=2}^N f_i(W) (\delta R_{H_2O,i}), \quad (47)$$

where  $R_{H_2O,i}$  are the TPW resistance values used to calculate each  $W_i$ . The  $R_{H_2O,i}$  value may be different for each  $W_i$  value. From here the calculations diverge depending on how an SPRT is calibrated, in particular, it depends on how the  $R_{H_2O,i}$  values are used, and this depends largely on the type of SPRT.

The propagation of uncertainty based on the approximate Equation (45) neglects the correlation of the different  $W_i$  values. But the possibly resulting overestimation of the uncertainty seems to be acceptable because primarily non-dominant uncertainty components are correlated. In most cases, the uncertainty estimates  $u(W_i)$  are dominated by components caused by the influence of impurities on the fixed-point temperatures, see *Guide* Section 2.1 *Influence of impurities* in Chapter 2 *Fixed Points* (<http://www.bipm.org/en/committees/cc/cct/guide-its90.html>). This influence is not correlated between the different fixed-point materials. A complicated program for considering correlations completely applying the interpolation equations of the ITS-90 has been developed by Gaiță *et al.* [Gaiță and Iliescu 2001]. But it is questionable if such a huge effort is necessary for decreasing the overall uncertainty estimate only slightly. For most applications, Equation (47) will yield reliable uncertainty estimates. Uncertainty propagation using the ITS-90 mathematics is also discussed in [Meyer and Ripple 2006].

### 6.2.2. Propagated uncertainty

For LSPRTs and HTSPRTs, a separate TPW measurement is used to calculate each  $W_i$  value. This is done to minimise the effects of the oxidation state of the SPRT. If we assume the uncertainties associated with each resistance measurement in Equation (47) are not correlated, we can immediately write down the corresponding expression for the uncertainty in the interpolated  $W_r$  value:

$$u^2(W_r) = f_{H_2O}^2(W) \frac{u_{\text{mean}}^2(R_{H_2O})}{R_{H_2O}^2} + \sum_{i=2}^N f_i^2(W) \frac{(u^2(R_i) + W_i^2 u^2(R_{H_2O,i}))}{R_{H_2O}^2} \quad (48)$$

The set of terms under the summation propagates the uncertainties resulting from the calibration at the fixed points. The first term considers the mean uncertainty of the measurements at the TPW.

Equation (48) is well suited to use for the ITS-90 sub-ranges using LSPRTs and HTSPRTs, where separate TPW measurements are used to eliminate oxidation effects. A useful feature of the equation is that we can identify a single term that can be associated with the uncertainty in the  $W_i$  value for each fixed point:

$$u^2(W_i) = \frac{u^2(R_i) + W_i^2 u^2(R_{H_2O,i})}{R_{H_2O}^2}. \quad (49)$$

The calibration of CSPRTs is a little different from above because the same TPW value  $R_{\text{H}_2\text{O,cal}}$  is used to calculate all of the  $W_i$  values for the fixed points. In that case, Equation (47) simplifies together with Equation (21) to

$$u^2(W_i) = \frac{1}{R_{\text{H}_2\text{O}}^2} \left[ f_{\text{H}_2\text{O}}^2(W) u^2(R_{\text{H}_2\text{O,cal}}) + (W - f_{\text{H}_2\text{O}}(W))^2 u^2(R_{\text{H}_2\text{O,cal}}) + \sum_{i=2}^N f_i^2(W) u^2(R_i) \right] \quad (50)$$

With this simplification, the total propagated uncertainty due to uncertainty in the TPW value has been significantly reduced and the uncertainty associated with each  $W_i$  value has the form

$$u(W_i) = \frac{u(R_i)}{R_{\text{H}_2\text{O}}}. \quad (51)$$

### 6.3. Non-uniqueness due to interpolation errors

Ideally, the interpolating equations of ITS-90 should accommodate the real variations in the  $W(T_{90})$  relationship for different SPRTs. In practice, the functional form of the equations is insufficient to characterise the many complex physical effects that occur, and hence the SPRT calibration equations are subject to interpolation error. Interpolation error is responsible for non-uniqueness [Mangum *et al.* 1997]: ambiguity in the scale that arises because there is more than one equation used (Type 1 non-uniqueness), more than one type of interpolating instrument used (Type 2 non-uniqueness), or different instances of the designated interpolation instrument applied (Type 3 non-uniqueness). Over most of the SPRT sub-ranges of ITS-90, non-uniqueness is a major source of uncertainty.

In this subsection, we consider only Type 1 and Type 3 non-uniqueness in the SPRT sub-ranges. (Type 2 non-uniqueness contributes only between the triple point of equilibrium hydrogen (13.8033 K) and the triple point of neon (24.5561 K), where the SPRT and the interpolating gas thermometer sub-ranges overlap.) Type 1 and Type 3 non-uniqueness contribute up to 0.5 mK or more to uncertainty in the temperatures interpolated between fixed points.

The physical cause of non-uniqueness is presumed to be due to different crystal orientations, different grain sizes, and different impurities in different concentrations in the platinum wire, as well as variations in surface and vacancy effects. The general mathematical form of both types of non-uniqueness is known since they are forms of interpolation error [White and Saunders 2007]:

$$\delta W_r = (W - W_1)(W - W_2)(\dots)(W - W_N)g(W), \quad (52)$$

where  $W_1 \dots W_N$  are the measured resistance ratios at the fixed points and  $g(W)$  is function of resistance ratio that may or may not be known depending on the type of non-uniqueness. Although Types 1 and 3 non-uniqueness are not independent (because this would imply that Type 3 non-uniqueness between two SPRTs would be the same in different sub-ranges), the correlations between them have not yet been fully established [Rusby *et al.* 2017], and for the present they are treated separately. The definition of the uncertainty for non-uniqueness is problematic because of the multiple sub-ranges and definitions, some of which are infrequently used, each of which gives rise to very different, typically non-normal distributions of temperatures. There is also a paucity of data in some sub-ranges.

### 6.3.1. Type 1 non-uniqueness (sub-range inconsistency)

Sub-range inconsistency (SRI), for a given SPRT, is calculated from the difference between the interpolating equations for two different sub-ranges. For example, it has been shown that the SRI between the water-zinc and water-aluminium sub-ranges, expressed in terms of the difference in calculated reference resistance ratio, is

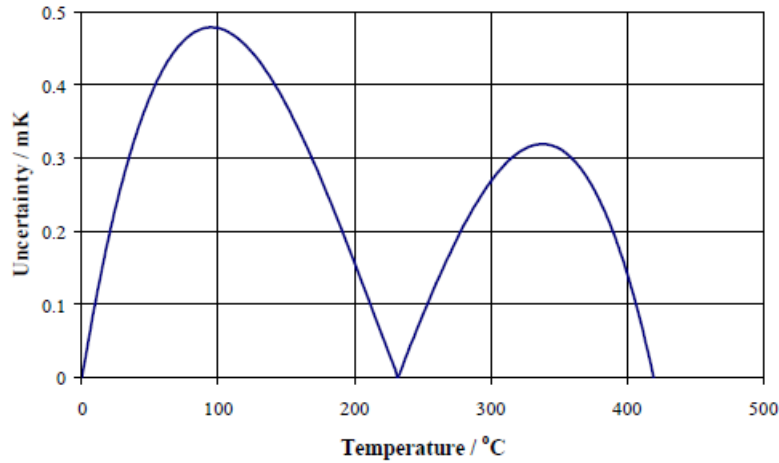
$${}^{\text{Al}}W_r(W) - {}^{\text{Zn}}W_r(W) = (W - 1)(W - W_{\text{Sn}})(W - W_{\text{Zn}})c, \quad (53)$$

where the superscript indicates the sub-range and  $c$  is the coefficient of  $(W - 1)^3$  in the ITS-90 water-aluminium interpolating equation for the SPRT [Zhiru *et al.* 2002, White and Strouse 2009]. SRI has two key features, both illustrated by Equation (53). Firstly, it has a well-defined mathematical form determinable from the different interpolating equations and measured fixed-point resistance ratios. Secondly, it has zeros at the fixed points shared by the two sub-range equations. The proportionality with  $c$  is to be expected for dimensional reasons but also because if  $W_{\text{Al}}$  lies on the extrapolation of  ${}^{\text{Zn}}W_r(W)$ , then it follows that the two interpolations are identical: the same values of  $a$  and  $b$  apply,  $c$  is zero, and there is no sub-range inconsistency. The SRI has a minor “scale intrinsic” component arising from inconsistencies in the  $W_{r,i}$  values specified at the fixed points.

For temperatures between 420 °C (zinc point) and 962 °C (silver point), there are no overlapping equations and, therefore, there is no SRI. Between 0.01 °C (TPW) and 420 °C (zinc point), the SRI is described by 15 different equations arising from the overlap of the water-gallium, mercury-gallium, water-indium, water-tin, water-zinc, and water-aluminium sub-ranges. Over most of the 0.01 °C to 420 °C range, the SRI is dominated by the difference between the water-zinc and water-aluminium sub-ranges. There is a lot of good data for this sub-range with studies by Mangum *et al.* (1990), Strouse (1992a), Moiseeva and Pokhodun (1992), Ahmed (2005), and Ancsin (1996). The two most extensive studies are by Zhiru *et al.* (2002), and White and Strouse (2009), who determined the SRI for 65 and 50 SPRTs, respectively. The maximum values for the SRI occur near 93.15 °C, where the two studies found mean SRI values of 0.05 mK and 0.12 mK, suggesting that the ITS-90 definitions are broadly consistent. However, the standard deviations for the two groups of SPRTs were 0.33 mK and 0.48 mK, respectively, the latter result probably reflecting a wider range of SPRT models and manufacturers in the sample. Figure 7 gives the SRI uncertainty versus temperature, calculated according to

$$u(W_{r,\text{SRI}}) \approx 8.0 \times 10^{-6} (W - 1)(W - W_{\text{Sn}})(W - W_{\text{Zn}}), \quad (54)$$

which is derived from the measurements of White and Strouse (2009). Note that the equation gives the uncertainty in the interpolated value of the reference resistance ratio as a function of measured resistance ratio, while Figure 7 has been scaled so that both axes are in terms of temperature.



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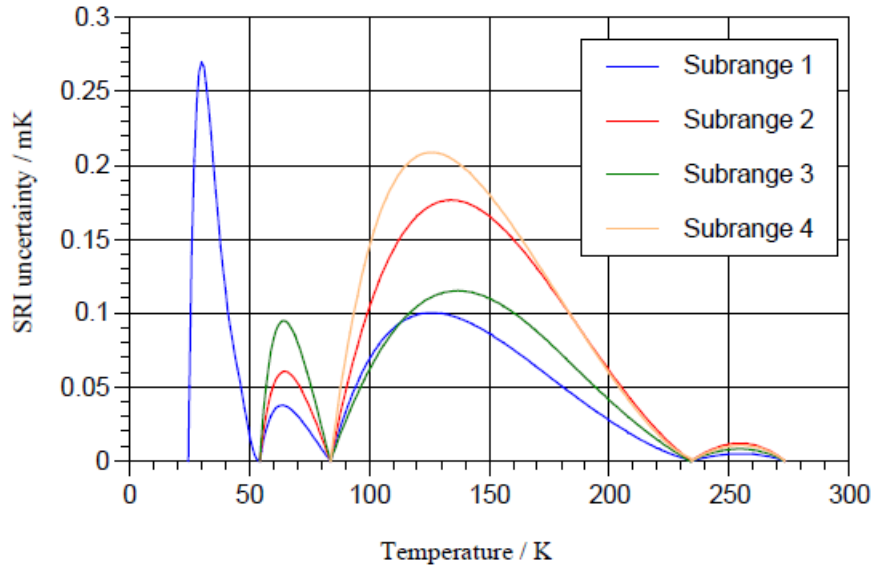
**Figure 7 — Standard deviation of the sub-range inconsistency over the range 0 °C to 420 °C for a group of 50 SPRTs, from White and Strouse (2009). The maxima occur at 93.15 °C ( $W = 1.366\,227$ ) and 336.57 °C ( $W = 2.274\,97$ ).<sup>(1)</sup>**

For temperatures below 0.01 °C, the SRI arises from the overlap of the different sub-ranges between the TPW and each of the triple points of equilibrium hydrogen, neon, oxygen, argon and mercury as well as the sub-range between the triple point of mercury and the melting point of gallium. (The latter range is omitted below.) Recent data for these sub-ranges comes from Meyer and Tew (2006), and Steele (2005), with very similar results. With the data from 18 thermometers, Meyer and Tew determined the standard deviations of the SRI shown in Figure 8. Each curve in Figure 8 is described by a polynomial of the form

$$u(T_{90,\text{SRL}i}) = \sum_{j=1}^5 A_{ij}(T_{90} - T_0)^j, \quad (55)$$

where the coefficients  $A_{ij}$  and the constants  $T_0$  are given in Table 5.

Applying the alternative interpolation Equation (22), White and Strouse (2009) could show that all terms in the equations describing the SRI are proportional to differences of inverse  $S_i$  values defined by Equation (25). This means the SRI is zero if the  $S_i$  values for a single SPRT are the same for all fixed points.



**Figure 8** — Uncertainty due to Type 1 non-uniqueness over the range 24 K to 273 K, with respect to the mean of the four overlapping sub-ranges terminating at the triple points of hydrogen, neon, oxygen and argon (Meyer and Tew 2006, see p. 56). The curves of sub-ranges 1 and 2 are identical from 25 K to 54 K.<sup>(1)</sup>

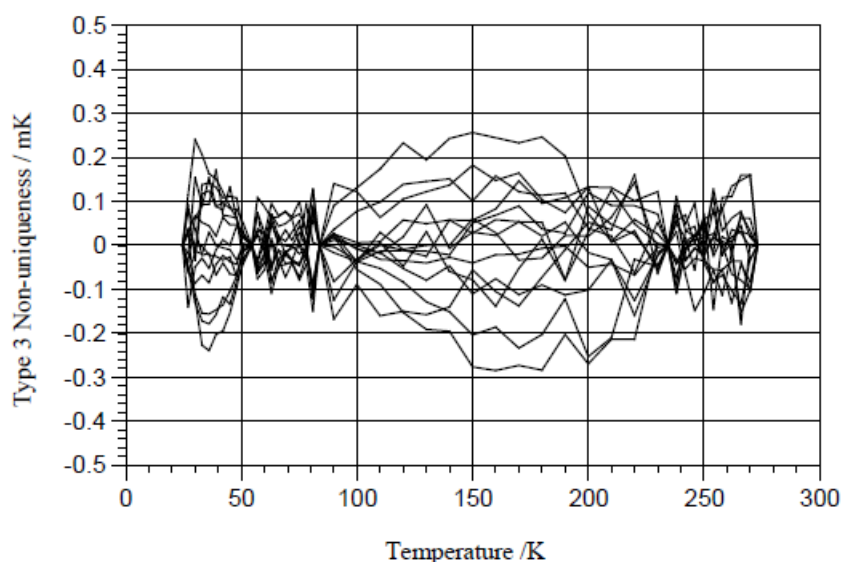
**Table 5.** Table of coefficients  $A_{ij}$  and constants  $T_0$  for the Equations Equation (55) giving the uncertainty due to sub-range inconsistency for temperatures below 273.16 K [Meyer and Tew 2006].

Interval		24 K to 54 K	54 K to 84 K	84 K to 234 K	234 K to 273 K
$T_0$		24.5561 K	54.3584 K	83.8058 K	234.3156 K
Sub-range 1	$A_{11}$	$1.226\,72 \times 10^{-1}$	$1.035\,03 \times 10^{-2}$	$5.937\,67 \times 10^{-3}$	$4.292\,53 \times 10^{-4}$
	$A_{12}$	$-1.882\,93 \times 10^{-2}$	$-9.836\,57 \times 10^{-4}$	$-1.190\,04 \times 10^{-4}$	$-8.511\,44 \times 10^{-6}$
	$A_{13}$	$1.162\,74 \times 10^{-3}$	$4.041\,73 \times 10^{-5}$	$9.798\,45 \times 10^{-7}$	$-6.165\,75 \times 10^{-8}$
	$A_{14}$	$-3.320\,79 \times 10^{-5}$	$-8.844\,29 \times 10^{-7}$	$-4.117\,01 \times 10^{-9}$	0
	$A_{15}$	$3.612\,10 \times 10^{-7}$	$8.185\,25 \times 10^{-9}$	$7.437\,45 \times 10^{-12}$	0
Sub-range 2	$A_{21}$	$1.226\,72 \times 10^{-1}$	$1.469\,65 \times 10^{-2}$	$8.179\,35 \times 10^{-3}$	$1.121\,23 \times 10^{-3}$
	$A_{22}$	$-1.882\,93 \times 10^{-2}$	$-1.225\,28 \times 10^{-3}$	$-1.174\,48 \times 10^{-4}$	$-2.418\,84 \times 10^{-5}$
	$A_{23}$	$1.162\,74 \times 10^{-3}$	$4.421\,59 \times 10^{-5}$	$5.216\,56 \times 10^{-7}$	$-1.187\,54 \times 10^{-7}$
	$A_{24}$	$-3.320\,79 \times 10^{-5}$	$-9.021\,57 \times 10^{-7}$	$-6.277\,18 \times 10^{-10}$	0
	$A_{25}$	$3.612\,10 \times 10^{-7}$	$8.088\,77 \times 10^{-9}$	$-3.524\,29 \times 10^{-13}$	0
Sub-range 3	$A_{31}$		$2.446\,61 \times 10^{-2}$	$4.516\,32 \times 10^{-3}$	$7.571\,90 \times 10^{-4}$
	$A_{32}$		$-2.176\,72 \times 10^{-3}$	$-4.246\,06 \times 10^{-5}$	$-1.631\,14 \times 10^{-5}$
	$A_{33}$		$8.410\,30 \times 10^{-5}$	$-1.541\,05 \times 10^{-7}$	$-8.058\,81 \times 10^{-8}$
	$A_{34}$		$-1.782\,94 \times 10^{-6}$	$2.652\,34 \times 10^{-9}$	0
	$A_{35}$		$1.627\,20 \times 10^{-8}$	$-7.178\,17 \times 10^{-12}$	0
Sub-range 4	$A_{41}$			$1.262\,90 \times 10^{-2}$	$9.616\,22 \times 10^{-4}$
	$A_{42}$			$-2.625\,39 \times 10^{-4}$	$-1.952\,92 \times 10^{-5}$

$A_{43}$	$2.322\,29 \times 10^{-6}$	$-1.276\,28 \times 10^{-7}$
$A_{44}$	$-1.071\,72 \times 10^{-8}$	0
$A_{45}$	$2.110\,43 \times 10^{-11}$	0

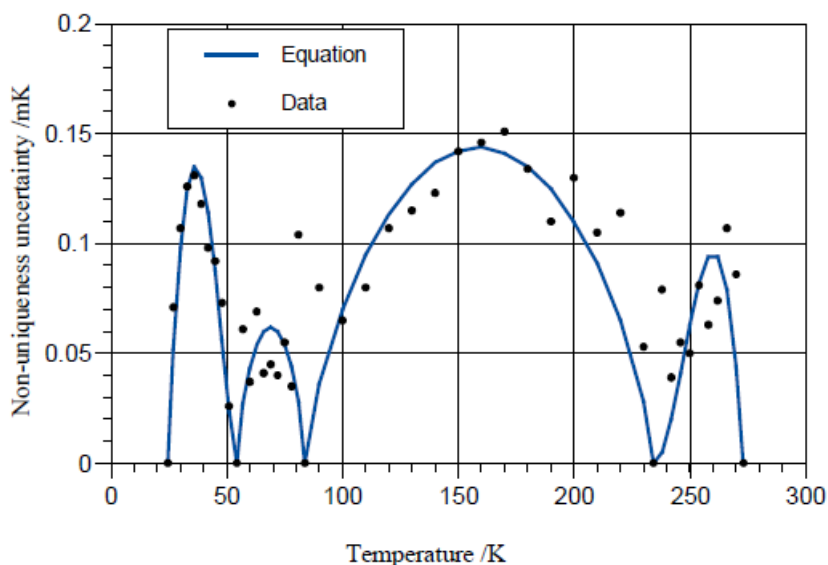
### 6.3.2. Type 3 non-uniqueness

Conventionally described simply as non-uniqueness, Type 3 non-uniqueness arises from the differences in the interpolated  $W_r$  values due to the use of different SPRTs over the same sub-range. It has the form of Equation (52) in that it exhibits zeros at all of the fixed points, but the  $g(W)$  function is unknown. It is known that the non-uniqueness is made worse in the sub-ranges where the fixed-point sensitivity coefficients amplify uncertainties [White and Saunders 2007]. There have been several investigations of non-uniqueness at temperatures between 14 K and 273 K, e.g. [Ward and Compton 1979, Head 1997, and Hill and Steele 2003]. Figure 9 shows the non-uniqueness for the temperature range between 24 K and 273.16 K based on Hill and Steele (2003). Figure 10 shows the standard deviations versus temperature for the data of Figure 9 as well as functions fitted to the measured standard deviations. The functions are given in Table 6.



**Figure 9 — The Type 3 non-uniqueness of ITS-90 between 24.5561 K and 273.16 K (from the data of Hill and Steele (2003), see p. 54)**





**Figure 10** — The measured uncertainty (standard deviation) due to Type 3 non-uniqueness in the range 24.5561 K to 273.16 K (from the data of Hill and Steele (2003), see p. 54) and the functional approximation of Table 6.

**Table 6.** Uncertainty due to Type 3 non-uniqueness for intervals in the range 24.5561 K to 273.16 K.

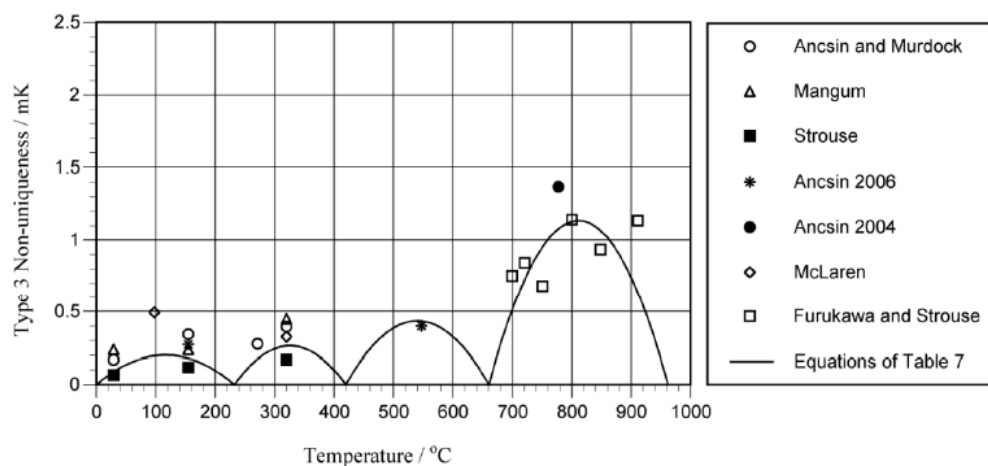
Temperature range / K	Uncertainty / mK ( $T$ in K)
24.5561 to 54.3584	$1.5 \times 10^{-4} (T - 24.5561)(54.3584 - T)^{1.5}$
54.3584 to 83.8058	$1.1 \times 10^{-3} (T - 54.3584)^{0.75} (83.8058 - T)^{0.75}$
83.8058 to 234.3156	$2.2 \times 10^{-4} (T - 83.8058)^{0.75} (234.3156 - T)^{0.75}$
234.3156 to 273.16	$1.1 \times 10^{-5} (T - 234.3156)^2 (273.16 - T)$

In the temperature range between 0 °C and 962 °C there is limited data on Type 3 non-uniqueness because of the problems with oxidation and drift of SPRTs as well as the difficulty of making accurate comparison measurements. Most data is based on calibrations at redundant fixed points, e.g. [Ancsin and Murdock 1990, Strouse 1992b, Furukawa and Strouse 2002, Ancsin 2004, 2006], which are summarised in Figure 11. Higher values above 660 °C are given by Marcarino *et al.* (1997).

The paucity of data makes the development of an equation for the uncertainty due to non-uniqueness problematic. Fellmuth *et al.* (2005) applied a constant value of 0.6 mK to simply represent the uncertainty for this temperature range; however, that neglects the knowledge that the uncertainty should be zero at the fixed points. Figure 11 shows an improved approximation based on quadratic functions that pass near the lowest points in the interpolation intervals. The equations for the curves are given in Table 7.

**Table 7.** Approximate expressions for uncertainty due to Type 3 non-uniqueness above 0 °C.

Temperature range	Uncertainty / mK ( $t$ in °C)
0.01 °C to 231.928 °C	$1.5 \times 10^{-5} t (231.928 - t)$
231.928 °C to 419.527 °C	$3 \times 10^{-5} (419.527 - t) (t - 231.928)$
419.527 °C to 660.323 °C	$3 \times 10^{-5} (660.323 - t) (t - 419.527)$
660.323 °C to 961.78 °C	$5 \times 10^{-5} (961.78 - t) (t - 660.323)$

**Figure 11** — Summary of available data on Type 3 non-uniqueness of ITS-90 above 0 °C. The data of Mangum and McLaren are from Hill and Woods (1992). The solid line shows the curves defined by the polynomials of Table 7.

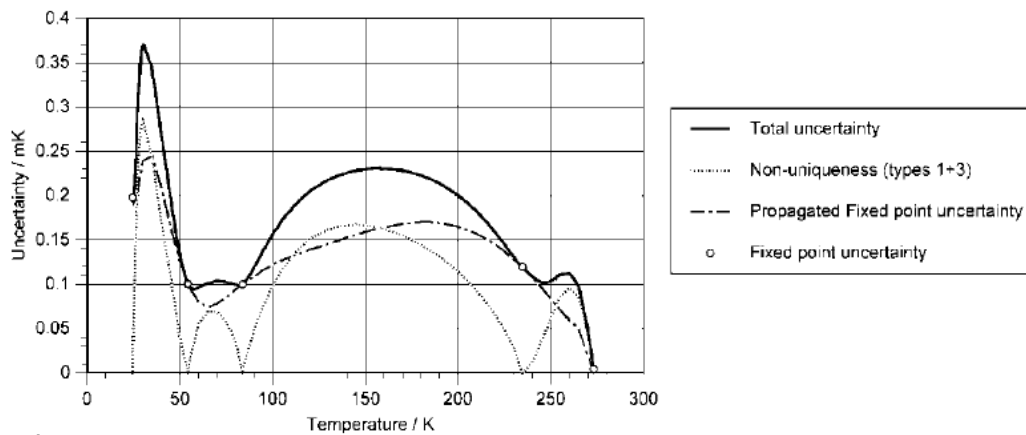
## 6.4. Total uncertainty in realised temperature

The application of Equation (48) and Equation (50) requires estimation of the uncertainty of the determination of the SPRT resistance values at the defining fixed points including the TPW ( $u(R_i)$ ,  $u(R_{H_2O})$ ). Relevant uncertainty components result from fixed-point effects and the resistance measurement. [Appendix 3](#) provides a complete tabular summary of the typical range of uncertainties at each of the fixed points that is purely indicative. It is assumed that all resistance measurements are performed applying the same equipment. In most cases, the influence of impurities is the largest fixed-point effect. Its estimation is treated in *Guide Section 2.1 Influence of impurities* of Chapter 2 *Fixed points*. Once the uncertainty in the interpolated resistance ratio  $u(W_r)$  is calculated, the total uncertainty in the realised temperature is then given by

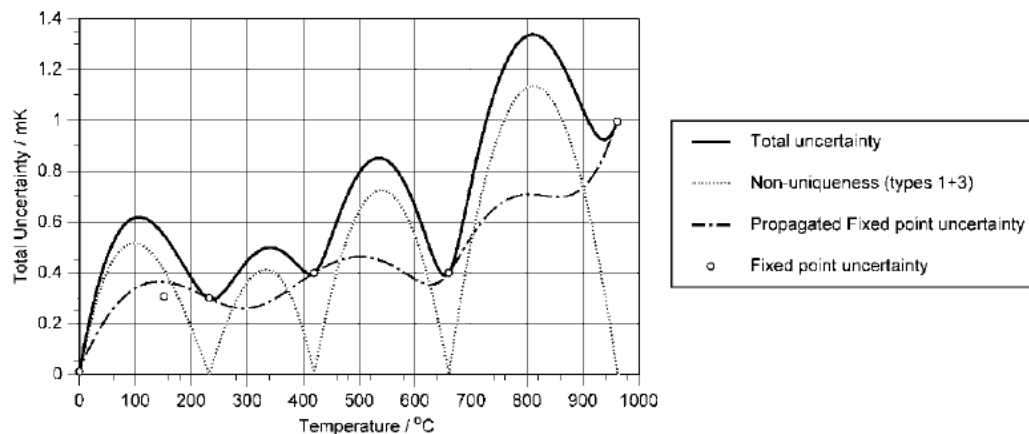
$$u_{\text{total}}^2(T_{90}) = \left( \frac{dT_{90}}{dW_r} \right)^2 [u^2(W_r) + u^2(\delta W_{\text{SRI}}) + u^2(\delta W_{\text{NU3}})] \quad (56)$$

where  $u(\delta W_{\text{SRI}})$  and  $u(\delta W_{\text{NU3}})$  are the uncertainties due to Type 1 (sub-range inconsistency) and Type 3 non-uniqueness (NU3). Figure 12 and Figure 13 present examples of evaluations of Equation (56) and showing the contributions of the uncertainties associated with the fixed

points and the two types of non-uniqueness, but not those associated with the SPRT effects discussed in Section 5.1.



**Figure 12** — Example of the total calibration uncertainty for a SPRT over the neon-water sub-range, showing the contribution of uncertainty due to fixed points and the two forms of non-uniqueness. The values shown are standard uncertainties and are indicative of a high-quality ITS-90 realisation.



**Figure 13** — Example of the total calibration uncertainty for a SPRT over the water-silver sub-range, showing the contribution of uncertainty due to fixed points and the two forms of non-uniqueness. (In the range between the freezing points of zinc and aluminium, the contribution of the Type 3 non-uniqueness is slightly increased compared with Figure 11 because of the lack of sufficient data.) The values shown are standard uncertainties and are indicative of a high-quality ITS-90 realisation.

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