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# Acoustic measurement of the triple point of neon $T_{Ne}$ and thermodynamic calibration of a transfer standard for accurate cryogenic thermometry

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#### Abstract

We used absolute primary acoustic gas thermometry (AGT) to calibrate a Pt–Co resistance thermometer on the thermodynamic temperature scale by measuring the speed of sound in helium at a temperature  $T^*$  chosen to be near the temperature of the triple point of neon,  $T_{\text{Ne}}$ . Prior to the present AGT, the Pt–Co thermometer was used with a neon triple-point cell as part of an interlaboratory comparison. Taken together, the results of the interlaboratory comparison and the present AGT redetermined the thermodynamic temperature  $T_{\text{Ne}} =$  $(24.55515 \pm 0.00024)$  K. This new value of  $T_{\text{Ne}}$  is consistent with other recent determinations obtained with various primary methods. After completing the AGT thermodynamic calibration, we used the Pt–Co thermometer to link  $T^*$  to the temperature ratios measured by single-pressure refractive-index gas thermometry (SPRIGT) in a different laboratory. (Gao *et al* 2020 *Metrologia* **57** 065006) Now, the  $T^*$ -linked SPRIGT system can calibrate other thermometers on the thermodynamic temperature scale T in the range  $5 \text{ K} \leq T \leq T_{\text{Ne}}$  without using the international temperature scale ITS-90. At most temperatures in this range, the uncertainties of the  $T^*$ -linked SPRIGT system are smaller than those of the ITS-90 systems used by National Metrology Institutes to calibrate resistance thermometers.

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Original content from this work may be used under the terms of the Creative Commons Attribution 4.0 licence. Any further distribution of this work must maintain attribution to the author(s) and the title of the work, journal citation and DOI. Keywords: primary thermometry, Pt Co thermometer, acoustic gas thermometry

Supplementary material for this article is available online

(Some figures may appear in colour only in the online journal)

#### 1. Introduction

Using the techniques of primary acoustic gas thermometry (AGT), [1, 2] we measured the thermodynamic temperature of the triple point of neon and obtained the result:  $T_{\text{Ne}} = (24.55515 \pm 0.00024)$  K. (All uncertainties reported in this work are one standard uncertainty corresponding to a 68% confidence level.) As shown in table 1 and figure 1, the present result for  $T_{\text{Ne}}$  is consistent, within combined uncertainties, with other recent determinations of  $T_{\text{Ne}}$  [3–6].

In figure 1, two literature values of  $T_{\rm Ne}$  are based on refractive-index gas thermometry (RIGT) [5, 6] and one is based on dielectric constant gas thermometry (DCGT) [4]. These values of  $T_{\rm Ne}$  were obtained during wide-range measurements of the differences  $T - T_{90}$  and their estimated uncertainties  $u(T - T_{90})$ . However, these references do not include values of  $u(T_{\text{Ne}})$ , We consulted the respective authors who graciously provided  $u(T_{\text{Ne}})$ . In 2006, two of the present authors (LP and MRM) obtained two values of T<sub>Ne</sub> using primary AGT [3]. (The two 2006 AGT values differ, in part, because they were traced to two different neon triple point cells.) The present AGT uses completely new hardware and software and a different neon triple-point cell. Furthermore, the present apparatus was not operated by LP and MRM. Thus, the 2006 AGT value of  $T_{\rm Ne}$  and the present AGT value are uncorrelated.

Our acoustic measurement of the thermodynamic temperature near  $T_{\text{Ne}}$  is preliminary and a key part of the program of Gao *et al* [7] to use single-pressure refractive-index gas thermometry (SPRIGT) to calibrate secondary thermometers (such as resistance thermometers) in the temperature range 5 K  $\leq T \leq T_{\text{Ne}}$ . Gao *et al* measure *ratios* of thermodynamic temperatures. By linking their ratio measurements to the present value of  $T_{\text{Ne}}$ , we have enabled their system to calibrate secondary thermometers with the thermodynamic temperature without using any input from the internationally accepted temperature scale ITS-90.

To link the SPRIGT to  $T_{\text{Ne}}$ , we determined  $T^*$  in two successive AGT runs in June and July 2019 at LCM-LNE-Cnam. After completing and analyzing the results of the AGT work, we obtained a thermodynamic calibration of one particular platinum–cobalt (Pt–Co) resistance thermometer, which was transferred to the TIPC-LNE Joint Laboratory of the Chinese Academy of Sciences, where—by September 2019—it was installed in the SPRIGT system to calibrate that system at  $T_{\text{Ne}}$ . As discussed in [7], the uncertainty of the  $T_{\text{Ne}}$ -linked SPRIGT system was 0.16 mK at  $T_{\text{Ne}}$  where it was dominated by imperfections in the AGT and resistance thermometry. (Thus, linking a more accurate value of  $T_{\text{Ne}}$  could further improve the SPRIGT system.) At 5.0 K, the uncertainty of the SPRIGT system was 63  $\mu$ K; it was dominated by the uncertainty of helium's 2nd and 3rd density virial coefficients *B* and *C*.

Figure 2 shows that, in the range 14 K  $\leq T \leq T_{\text{Ne}}$ , the uncertainties of T<sub>Ne</sub>-linked SPRIGT, primary AGT, and primary DCGT are all substantially smaller than those of  $T_{90}$  as disseminated, for example, by the National Metrology Institutes (NMIs) of France and Italy. [8] In this range, ITS-90 can be realized using two different instruments, the constant volume gas thermometer (CVGT) above 3 K, and the capsule standard platinum resistance thermometer (cSPRT) above 13.8 K. [9] The cSPRT must be calibrated at specific sets of defined temperature fixed points: the  $e-H_2$  triple point, two vapor pressure points of  $e-H_2$ , and  $T_{\text{Ne}}$ . During the last 20 years, no NMI in the world realized the  $e-H_2$  vapor pressure points, which gives the cSPRT an uncertainty on the order of 1 mK between these two points. Furthermore, only the NMI of Japan has operating CVGTs. Because operating these CVGTs is complex and time-consuming, these NMIs usually calibrate new thermometers by comparing them to previously calibrated thermometers. To summarize, the realizations of ITS-90 in the range 5 K  $\leq T \leq T_{Ne}$  are infrequent, complex, and have larger uncertainties than newer primary thermometers. In this range, newer primary thermometers are competitive with ITS-90.

Our realization of a primary acoustic gas thermometer was a helium-filled, copper-walled, quasi-spherical, cavity resonator. The theoretical principles and experimental techniques for determining  $T_{\text{Ne}}$  with this AGT are well-established [1, 2, 10]. Here, we briefly describe the AGT and our measurements while emphasizing the features that dominate our uncertainty budget.

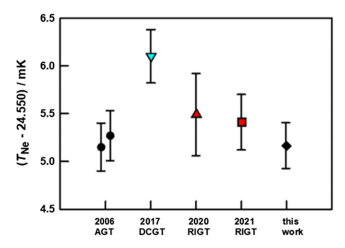
#### 2. Experiment setup

#### 2.1. Cryostat

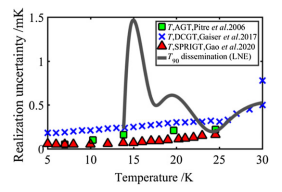
The AGT and its pressure vessel were contained within a cryogen-free cryostat cooled by a commercial 4 K pulse-tube cryocooler (Cryomech PT405, 0.5 W at 4.2 K cooling power)<sup>7</sup>. Compared to the cryostat used by Pitre *et al* in 2006 [3], the

<sup>&</sup>lt;sup>7</sup> In order to describe materials and procedures adequately, it is occasionally necessary to identify commercial products by manufacturer's name or label. In no instance does such identification imply endorsement by the institutions listed in the authors addresses, nor does it imply that the particular product or equipment is necessarily the best available for the purpose.

	Table 1.	Recent deterr	ninations of $T_{\text{Ne}}$ .	
Reference	Year	Method	$T_{\rm Ne}~{ m K}$	$u(T_{\rm Ne})  { m mK}$
			24.555 15 <sup>a</sup>	0.25
[3]	2006	AGT	$24.55527^{a}$	0.26
[4]	2017	DCGT	24.5561	0.28
[5]	2020	RIGT	24.555 49	0.43
[6]	2020	RIGT	24.555 41	0.29
This work	2020	AGT	24.555 15	0.24



**Figure 1.** Values of  $T_{\text{Ne}}$  from the literature and from this work. Literature values are identified, by technique, first author, year, and reference:  $\bigcirc$  AGT Pitre *et al* (2006) [3];  $\checkmark$  DCGT Gaiser *et al* (2017) [4];  $\blacktriangle$  RIGT Rourke (2020) [5];  $\blacksquare$  RIGT Madonna-Ripa *et al* 2021 [6];  $\blacklozenge$  AGT this work.



**Figure 2.** Uncertainties of realizations of T and  $T_{90}$ .

cryocooler is advantageous because it has long-term, continuous stability and it avoids the noise generated by boiling liquid helium. Figure 3 is a schematic diagram of the cryostat, which is the prototype of the cryostats used for several primary thermometers [6, 7]. To reduce the radiation thermal load on the pressure vessel surrounding the quasi-spherical resonator, the vessel was surrounded by two gold-plated heat shields attached to thermostatted flanges. When cooling began, a small amount of helium exchange gas (approximately 1000 Pa) was admitted into the sealed space between the shield designated 'heat switch' and the pressure vessel (figure 3). The lowest temperature reached by the pressure vessel was approximately 6 K. The methods for temperature control are the same ones that we used in previous work [7, 11-13]. Referring to figure 3, the finest temperature control of the quasisphere was achieved by reading cSPRT-1 with an F18 bridge and feeding the bridge's output through a proportional-integral feedback loop to heater-3. The temperature was monitored by cSPRT-2 using a second bridge. If the thermometry were ideal and if any temperature gradients were time-independent, the differences between the two thermometer readings  $\Delta T_{2-1} \equiv$  $(T_{cSPRT-2} - T_{cSPRT-1})$  would be time-independent. Averaged over each 12 days-long run,  $\langle \Delta T_{2-1} \rangle_{\text{Run } 2} - \langle \Delta T_{2-1} \rangle_{\text{Run } 1} =$ 3.5 µK. During all the measurements, imperfect pressure control generated very low-frequency fluctuations that were equivalent temperature changes as large as  $\pm 20 \ \mu$ K. These pressure fluctuations, together with the temperature fluctuations and possible temperature gradients, contributed 21 µK to the uncertainty of  $T_{\rm Ne}$ .

#### 2.2. Gas handling system

As shown in figure 4, the gas handling system was similar to that used by Pitre et al during their measurements of the Boltzmann constant [14, 15]. In [15], Pitre et al demonstrated that the cold trap and the getter in the fill line reduced the concentration of impurity gases to negligible levels. During most of the AGT measurements discussed in this work, the cold trap was not in use. A few, separate dedicated tests with the cold trap were conducted to check the possible presence and the effect of trace amounts of impurities, as discussed in section 4. The pressure control loop was installed in a temperature-controlled box that limited the maximum temperature excursion to 10 mK around a given set point. Two absolute pressure transducers (Paroscientific Digiquartz 745, full-scale pressure 0.7 MPa) monitored the pressure. These transducers were traceable to the LNE standard. The pressure in the resonator was regulated by one of the pressure transducers and a mass flow controller using a proportional-integral feedback loop written in the LabVIEW® software. A relative pressure stability of  $0.5 \times 10^{-6}$  was maintained throughout the operating range. As shown in figure 4, a 6 m-long capillary (ID 1.01 mm) connected the gas handling system to the quasi-spherical cavity. The electrical cables surrounded the capillary and were enclosed by a bellows. This arrangement avoided using hermetic feed throughs at low temperatures. On occasions when the cryostat was open at room temperature, valves 8 and 9 were used to flush the inside of the cavity with clean helium, thereby maintaining the cavity's cleanliness. During the measurement at low temperature, helium did not flow through the resonator to reduce the possible heat transfer from the room to the resonator and to minimize the cost of the helium.

#### 2.3. Quasi-spherical acoustic gas thermometer

The acoustic thermometer was the diamond-polished, copper, quasi-sphere known as 'TCU1'. This thermometer was described in detail and studied from a metrological point of view by Sutton *et al* 2010 [16]. Sutton *et al* used TCU1 to determine the Boltzmann constant k; their result was within 3 ppm (1 ppm = 1 part in 10<sup>6</sup>) of the final value of k adopted

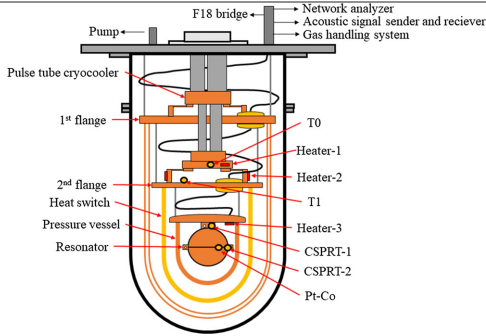


Figure 3. Schematic diagram of the cryogen-free cryostat used in this work.

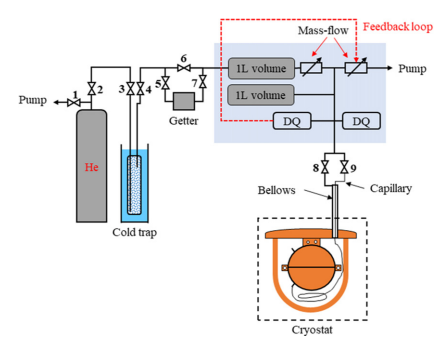


Figure 4. Gas handling system. The components in the gray rectangle were enclosed in a thermostatted box.

when the kelvin was redefined in 2019. TCU1 has an internal radius of 50 mm and a 5 mm thick wall. (A typical AGT with a 50 mm radius has a 10 mm thick wall.) TCU1's cavity is modeled as a triaxial ellipsoid with axes of length a,  $a(1 + e_1)$ , and  $a(1 + e_2)$ . In 2009, Sutton's microwave measurements determined  $e_1 = 0.001048$  and  $e_2 = 0.00079$  at the triple point of water. In this work, at the triple point of neon, using the TM11 mode, we determined  $e_1 = 0.001031$  and  $e_2 = 0.000792$ ; using the TM12 mode, we obtained  $e_1 = 0.001034$  and  $e_2 = 0.000792$  The small change in  $e_1$  might have resulted either from the movement of a microphone (or some other component of TCU1) or from anisotropic thermal expansion between  $T_{\text{TPW}}$  and  $T_{\text{Ne}}$ .

#### 2.4. Microwave measurements

Figure 5 is a block diagram of the microwave and acoustic measuring system. We used two straight antennas to measure the resonance frequencies and half-widths of the TM11 and TM12 microwave modes while TCU1 was thermostatted near  $T_{\rm Ne}$  and immersed in helium. For the TM11 mode, we used

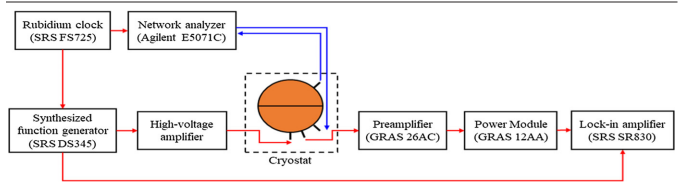


Figure 5. Block diagram of the microwave and acoustic measurement system.

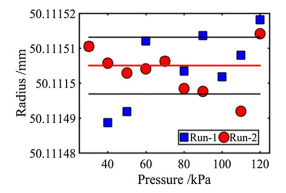
10 pressures ranging from 30 kPa to 110 kPa. For the TM12 mode, the data were much noisier and only two pressures were used: 30 kPa and 70 kPa. From the microwave data, we calculated  $a_{eq}$ , the radius of a sphere with the volume equivalent to that of the quasi-spherical cavity at zero pressure. To calculate  $a_{eq}$ , we corrected the measured frequencies for the microwave penetration depth, the refractive index of helium, the isothermal compressibility of copper (using the value  $-V^{-1}(\partial V/\partial P)_T = 19 \times 10^{-8} \text{ Pa}^{-1}$  from [7]), the effects of antennas and the fill duct [17], and the second-order shape correction [18, 19]. The statistical uncertainty of the extrapolated, zero-pressure radius is 0.14 ppm.

Figure 6 demonstrates that the values of  $a_{eq}$  from from two consecutive runs 1 and 2 were mutually consistent and that the repeatability of a single determination of  $a_{eq}$  was approximately 1 part in 10<sup>7</sup>. With this precision in mind, table 2 demonstrates two inconsistencies: (1) the values of  $a_{eq}$  measured using the TM11 and TM12 modes are inconsistent by up to 1.0 ppm and (2) the values of  $a_{eq,m1}$  and  $a_{eq,m2}$  differ by up to 2.1 ppm, depending upon the method used to correct the measured frequencies for the microwave penetration depth.

The first correction method determines the electrical conductivity  $\sigma$  of the copper resonator by fitting the half-widths of the TM11z triplet. The three components of the triplet were simultaneously fitted with the result  $\sigma = 9.69(3) \times 10^8 \Omega^{-1} \text{ m}^{-1}$ . This value of  $\sigma$  and the theory of the resonance half-widths leads to the values of  $a_{eq,m1}$  in table 2. Note: we did not use literature values of  $\sigma$  because impurities and/or mechanical stress in oxygen-free, high-conductivity copper can change its electrical conductivity near 24 K by more than one order of magnitude [20].

The second correction method allows each component of each triplet to have its own value of  $\sigma$  and leads to  $a_{eq,m2}$  in table 2. Method 2 adds the measured half-width of each component of the microwave triplets TM11 and TM12 to the measured frequency of that component. Compared with  $a_{eq,m1}$ , the values of  $a_{eq,m2}$  were closer to the values expected from the studies of Sutton *et al* [16]. Method 2 was used for the final value of  $a_{eq}$ . To account of the inconsistent values of  $a_{eq}$  between the two methods, we added 4.2 ppm to the total uncertainty in table 5.

At 24 K, the Q of the microwave resonances is three times larger than at 273 K; therefore, there is a higher chance that the microwave resonances were over-coupled to the external



**Figure 6.** Equivalent radius of quasi-sphere from the frequencies of the TM11 microwave mode. The central red line is the mean radius of runs 1 and 2. The outer black lines lie at  $\pm 1$  standard deviation from the mean radius.

**Table 2.** Scaled relative deviations  $a_{eq}$  from their mean 106  $(a_{eq}/\langle a_{eq} \rangle - 1)$  for two modes using two values of the electrical conductivity of copper  $\sigma_{Cu}$ . Here,  $\langle a_{eq} \rangle = 50.111506$  mm.

Mode	$a_{\mathrm{eq},m1}$	$a_{\rm eq,m2}$	$a_{\mathrm{eq},m1} - a_{\mathrm{eq},m2}$
TM11 TM12	-1.11 -0.09	1.00 0.20	-2.11 -0.29
TM11-TM12	-1.02	0.79	

instruments and/or that the perturbations of the microwave resonances by the microphones and antennas were relatively more important. We were unable to test these ideas; therefore, we added 4.2 ppm to the uncertainty of  $a^2_{eq}$  in table 5 and label this uncertainty 'electrical conductivity of shell'. The ratios  $g_n/f_n$ in ppm for the triplet of the TM11 (*x*, *y*, *z*) are (4.63, 4.71, 4.32) and for the TM12 (2.54, 2.64, 2.41). These values were independent of pressure.

The mode-dependent inconsistencies [in table 2,  $|a_{eq,TM11} - a_{eq,TM12}| \gg u_{fit}(a_{eq})$ ] were up to 1 ppm near 24 K where  $u_{fit}(a_{eq})$  is the uncertainty due to the fit of the triplet. In contrast, Sutton *et al* [16], using the same quasi-sphere near 273 K, measured values of  $a_{eq}$  for the same modes that differed by only 0.12 ppm.

#### 2.5. Acoustic measurements

Two commercially-manufactured capacitive microphones (GRAS model 40BF) were installed in the cavity's wall so

that their diaphragms were flush with the cavity's internal surface. In order to reduce electrical crosstalk between the two microphones, the source microphone was excited with 50 V RMS so that it generated sound at twice the frequency of the sinusoidal driving signal. The driving signal originated in a function generator and was amplified to 50 V RMS by a TEGAM® model 2340 high-voltage amplifier (see figure 5). The second capacitive microphone detected the sound in the cavity resonator. The detecting microphone was connected via a coaxial cable to a preamplifier (a GRAS type 26AC) which was connected to a GRAS type 12AA power supply module located at room temperature. Finally, a lock-in amplifier (Stanford Research Systems SR830) was used to measure the in-phase and out-of-phase acoustic signal. At  $T_{\rm Ne}$ , this arrangement produced a satisfactory acoustic signal-to-noise ratio i.e. the fractional uncertainty of the resonance frequencies [1] was less than 10 ppm at the lowest pressure: 30 kPa.

We measured the acoustic resonance frequencies  $f_{(0,n)}$  and half-widths  $g_{(0,n)}$  of the radially-symmetric modes for n = 2, 3, 4, 5, 6, 8, and 9. The n = 7 mode was not included for two reasons. First, the frequencies of the (13, 2) and (0, 7) acoustic resonances overlap at low pressures where the modes broaden. (Ideally,  $f_{(13,2)}/f_{(0,7)} = 1.0017$ .) Secondly, the breathing mode of the empty copper shell of TCU1 strongly couples to the (0, 7) mode because both modes are resonant near 18.9 kHz at  $T_{\text{Ne}}$ .

We acquired data in two successive runs, each lasting approximately 12 days. Each run began at 120 kPa; then the pressure was decreased in steps of 10 kPa until the run ended at 30 kPa. After each pressure change, we waited 8 to 12 h for the temperature to stabilize. During the next 12 h, we repeatedly measured the resonance frequencies of the seven selected radial modes in approximately 70 cycles. Thus, each run consisted of approximately 4000 measurements of resonance frequencies.

The measurements were made under isothermal conditions at ten pressures between 30 kPa and 120 kPa. We applied all the corrections described by Moldover *et al* (2014) [1] to the measured values of  $f_{(0,n)}$  and  $g_{(0,n)}$ .

#### 3. Analyzing acoustic data

We corrected the measured values of  $f_{(0,n)}$  for the thermoacoustic boundary layer (using the theoretical value of the thermal conductivity of helium [21]) and we also applied small corrections for the mechanical compliance of the microphones [22], and the ducts that allow gas to enter and leave the cavity [16], and the ellipsoidal geometry of the cavity, as determined from the measurements of  $a_{eq}$ . However, the quasi-spherical copper shell is a complicated, bolted-together object; therefore, we and others [23–25] fit corrections to the measured acoustic frequencies to account for two shell-related phenomena: (1) the temperature jump in the helium gas where it contacts the copper shell and (2) the compliance of the copper shell in response to the radial oscillations of the helium within the shell.

**Table 3.** Relative importance of the terms in equation (1) in the present data (first two rows) in the measurement of k (last two rows) from reference [15]. The last column is proportional to the signal-to-noise ratio for measuring the resonance frequency  $f_{(0,5)}$ 

$\frac{T}{K}$	p MPa	$10^{6} \frac{A_{-1}}{pA_{0}}$	$10^6 \frac{A_0}{A_0}$	$10^6 \frac{A_1 p}{A_0}$	$10^6 \tfrac{A_2 p^2}{A_0}$	$\frac{10^{-6} p Q_{(0,5)}^2}{\rm MPa}$
24.5	0.03	7.5	1000 000	2108	15	1.4
24.5	0.12	1.9	1000 000	8433	240	24
273	0.15	9.3	1000 000	147	0.5	1.8
273	0.70	2.0	1000 000	685	9.9	45

On each isotherm, we represented the pressure dependence of the resonance frequencies (as corrected above) by the function:

$$\left(\frac{f_{(0,n)}}{z_{(0,n)}a_{eq}}\right)_{corr}^{2} = A_{-1}p^{-1} + A_{0} + A_{1}p + \Delta A_{1,(0,n)}p + A_{2}p^{2}.$$
 (1)

First, we discuss the terms in equation (1); then we discuss fitting the acoustic resonance frequencies with these terms.

In equation (1), z(0,n) is the eigenvalue appropriate for the (0,n) mode and  $A_0 \equiv w_0^2$  is the speed of sound in helium at zero pressure;  $w_0^2$  determines the thermodynamic temperature using the relation  $w_0^2 = 5 kT/(3m)$ . k is the Boltzmann constant and *m* the atomic mass of helium-4. The terms  $A_1p$ and  $A_2p^2$  account for the pressure dependence of the speed of sound. The parameters  $A_1$  and  $A_2$  are exact multiples of the temperature-dependent acoustic virial coefficients that have been calculated from the fundamental constants and the laws of quantum mechanics and statistical mechanics. For convenience, we fixed  $A_1(T)$  at the values that were accurately calculated by Czachorowski *et al* [26]. Fixing  $A_1$  affects neither our determination of  $T_{\text{Ne}}$  nor its uncertainty  $u(T_{\text{Ne}})$  because the coefficients of p in equation (1) are the sum  $[A_1 + \Delta A_{1,(0,n)}]$ and the uncertainties of  $\Delta A_{1,(0,n)}$  from the fitting are included in the uncertainty budget. We retained  $A_2(T_{\text{Ne}})$  as a fitted parameter because the uncertainty of the theoretical value of  $A_2(T_{\text{Ne}})$ from [27] appears to be larger than the uncertainty from fitting the present data. Within combined uncertainties, the fitted and theoretical values of  $A_2$  are mutually consistent. (See below.) Using the ab initio calculations of the fourth density virial coefficient of helium [28, 29], we crudely estimated  $A_3p^3/A_0$  $\sim 1 \times 10^{-6}$  at our highest pressure; therefore, we omitted  $A_3 p^3$ from equation (1).

In equation (1) the terms  $\Delta A_{1,(0,n)}p$  account for the frequency-dependent compliance of the 5 mm-thick copperwalled shell. Equation (86) of reference [30] shows that the compliance of a perfectly-spherical, isotropic shell has the form  $\Delta A_1 = \text{constant} \times [1-(f_{(0,n)}/f_{\text{breathing}})^2]^{-1}$ , where the constant depends on known properties of the gas and the shell. (In this work  $f_{(0,7)} \approx f_{\text{breathing}} \approx 18.9 \text{ kHz.}$ ) This prediction for an idealized shell is approximately consistent with our data; however, it does not capture the full precision of the data. Therefore, we decided to fit the data for each (0,n) mode with its own value of  $\Delta A_1$  which we denote ' $\Delta A_{1,(0,n)}$ '.

**Table 4.** Sensitivity of results to the choice of *h*. We fit equation (1) to combined data from run 1 and run 2 twice; once fixing  $A_{-1}$  assuming h = 0.39 and once fixing  $A_{-1}$  assuming h = 0.34. The values of  $A_1$  were fixed by theory [26]. The last row of the table is the theoretical value of  $A_2$  [27].

h	Value 0.39	Value 0.34	Uncert <sup>a</sup> Fixed	Unit
$w^2$	85011.900	85011.707	0.017	$m^2 s^{-2}$
$\Delta A_{(0,2)}$	-2.77	0.32	0.51	$\mathrm{m}^2~\mathrm{s}^{-2}~\mathrm{MPa}^{-1}$
$\Delta A_{(0,3)}$	-3.75	-0.67	0.51	$m^2 s^{-2} MPa^{-1}$
$\Delta A_{(0,4)}$	-4.46	-1.39	0.51	$m^2 s^{-2} MPa^{-1}$
$\Delta A_{(0,5)}$	-7.37	-4.28	0.51	$m^2 s^{-2} MPa^{-1}$
$\Delta A_{(0,9)}$	19.63	22.72	0.51	$m^2 s^{-2} MPa^{-1}$
$A_2$	1414.4	1399.5	3.4	$\mathrm{m}^2~\mathrm{s}^{-2}~\mathrm{MPa}^{-2}$
$A_1$ (theory)	5973.89	5973.87	Fixed	$m^2 s^{-2} MPa^{-1}$
$A_{-1}$	0.0191	0.0226	Fixed	$m^2 s^{-2} MPa$
$T = M w_0^2 / g_0 R$	24.554 967	24.554 911	$5  imes 10^{-6}$	Κ
$A_2$ (theory)	1457 <sup>b</sup>		52 <sup>b</sup>	$m^2 s^{-2} MPa^{-2}$

<sup>a</sup>Statistical contribution to standard uncertainty only.

<sup>b</sup>Reference [27].

In equation (1), the term  $A_{-1}p^{-1}$  accounts for unknown details of the scattering of helium atoms by the wall of the quasispherical cavity. Indeed, we do not know if the wall is covered with oxides and/or adsorbed gases and/or molecules of, for example, pump oil. Therefore, we and other AGTs have used phenomenological theories to model gas-wall interactions and their consequences for measurements of acoustic resonance frequencies [31, 32]. These theories predict that the wall-gas heat transfer generated when a sound wave reflects off a wall alters the distribution of helium velocities within a thermal accommodation length  $l_a$  of the wall such that the usual hydrodynamic boundary condition  $T_{gas} = T_{wall}$  no longer applies. Instead, a temperature jump occurs at the boundary. The jump introduces into equation (1) the term  $A_{-1}p^{-1}$ , which for low-density helium in a quasi-spherical shell takes the form

$$\frac{A_{-1}}{pw_0^2} = \frac{4l_a}{3a_{\rm eq}} = \frac{2\kappa}{3a_{\rm eq}p} \left(\frac{\pi mT}{2k}\right)^{1/2} \frac{2-h}{h}.$$
 (2)

In equation (2),  $\kappa$  is the thermal conductivity of helium and the length  $l_a$  is on the order of the mean free path in the helium. In the AGT literature, the value of *h* in equation (2) has been obtained by fitting low-pressure speed-of-sound data.

Table 3 lists the relative sizes of the terms in equation (1) in the pressure range of this work and, for comparison, the corresponding values from a recent measurement of the Boltzmann constant [15]. The last column is proportional to the signal-to-noise ratio for measuring the  $f_{(0.5)}$ .

In table 3, the term  $A_2p^2$  is substantially larger at  $T_{\text{Ne}}$  than at  $T_{\text{TPW}}$  and, as calculated *ab initio*, the standard uncertainty of  $A_2$  is 94 times larger at  $T_{\text{Ne}}$  than at  $T_{\text{TPW}}$ . Therefore,  $A_2(T_{\text{Ne}})$ must be fitted to our data. Unfortunately, fitting the present data does not determine  $A_{-1}$ ,  $w_0^2$ ,  $\Delta A_{1,(0,n)}$  and  $A_2$  with satisfactory precision. (We discuss this problem in section 7: Conclusions.) Therefore, we relied on the AGT literature to estimate  $A_{-1}$  and its uncertainty. Reference [32] reviewed the literature prior to 2016 and noted that  $0.33 \leq h \leq 0.43$  for helium-4 at  $T_{\text{TPW}}$ interacting with walls made of stainless steel, oxygen-free high-conductivity copper, and electrolytic-tough-pitch (ETP) copper. More recently, Gavioso *et al* [24] reported  $0.38 \le h \le 0.42$  in the range 236 K  $\le T \le 430$  K for helium with ETP copper. With these observations in mind, we assumed  $h = (0.39 \pm 0.05)$  and fitted our acoustic data twice; once fixing h = 0.34 (equivalent to  $A_{-1} = 0.023$  m<sup>2</sup> s<sup>-2</sup> MPa) and a second time fixing h = 0.39 (equivalent to  $A_{-1} = 0.019$  m<sup>2</sup> s<sup>-2</sup> MPa). We assume that the  $w_0^2$  (h = 0.39) is the most accurate value of  $w_0^2$  (and  $T_{\text{Ne}}$ ) and that the uncertainty contribution from h to  $u(w_0^2)$  is  $\pm |[w_0^2(h = 0.39) - w_0^2(h = 0.34]|$ .

Now, we describe the iterative procedure for fitting the acoustic data.

(Iter 1) Start by guessing the temperature  $T_{\text{trial}}$  as an initial estimate of  $T_{\text{Ne}}$ . (The guess  $T_{\text{trial}} = 20$  K is close enough!) With this guess, define the data set:

Data 
$$(p, T_{\text{trial}}, n, h) = \left(\frac{2\pi a_{\text{eq}} f_{(0,n)}}{z_{(0,n)}}\right)_{\text{corr}}^2 - A_{-1} (T_{\text{trial}}, h) p^{-1} - A_1 (T_{\text{trial}}) p.$$
 (3)

and the model that will be fitted to the data:

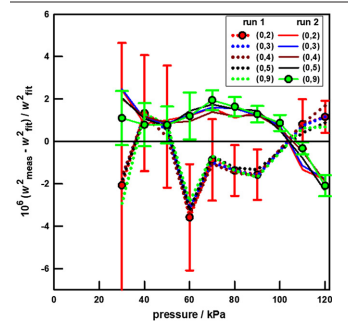
Model
$$(p, T, n) = w_0^2(T) + \Delta A_{1,(0,n)}p + A_2p^2.$$
 (4)

(Iter 2) Using  $T_{\text{trial}}$ , correct the microwave frequencies (for the microwave, penetration depth, the refractive index of helium, etc) and correct the acoustic frequencies (for the thermal boundary layer, ducts, etc). Using the corrected frequencies, minimize the sum

$$\min_{p, T \text{trial, } A2} \left\{ \sum \frac{\left[ \text{Data}(p, T_{\text{trial}}, n, h) - \text{Model}(p, T, n) \right]^2}{\sigma^2(p, T, n)} \right\}.$$
(5)

by varying  $A_{-1}$ ,  $w_0$ , and  $\Delta A_{1,(0,n)}$  for each acoustic mode at each pressure.  $\sigma^2$  is the squared standard deviation of the measured squared speed of sound.

(Iter 3) Replace  $T_{\text{trial}}$  with an improved  $T_{\text{trial}}$  obtained from  $T_{\text{trial}} = M w_0^2 / (\gamma_0 R)$ , where  $w_0$  resulted from the minimization, R is the gas constant, M is the molar mass of helium-4 and  $\gamma_0 \equiv 5/3$  is the heat capacity ratio at zero density.



**Figure 7.** Residuals after iteratively fitting  $T_{\text{Ne}}$ ,  $\Delta A_{1(0,n)}$ ,  $A_{-1}$  in equation (1) to the data for both runs while fixing h = 0.39. The legend identifies each radial mode. The outputs from the iteration are in table 4. Each plotted point represents the average of approximately 70 measurements; the uncertainty bars are the standard deviation of the 70 measurements from their mean. (The standard deviations of the means are  $(70)^{1/2} \approx 8,37$  times smaller.)

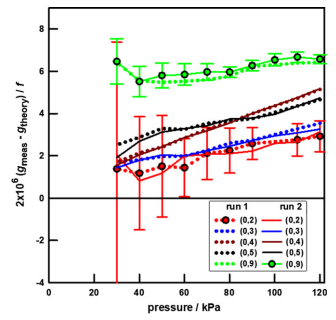
(Iter 4) Repeat steps (Iter 1) to (Iter 3) until the difference between successive values of  $T_{\text{trial}}$  is less than  $10^{-6} T_{\text{Ne}}$ .

The iteration converged rapidly; after three iterations, the differences between successive values of  $T_{\text{trial}}$  was  $10^{-4}$  mK. For run 2, the numerical results of the iterated fit for  $w_0^2$ ,  $A_{-1}$ ,  $\Delta A_{1(0,n)}$ , and  $A_2$  are listed in table 4. We emphasize that neither the iteration nor the theoretical values of  $A_1$  and  $A_2$  have any relationship to ITS-90.

After a preliminary analysis, we removed the (0,6) and (0,8) modes from our analysis because these modes had large, smooth deviations from the five remaining modes. We speculate that the frequencies of the (0,6) and (0,8) modes lay too close to  $f_{\text{breathing}}$  to represent the shell's compliance correction as the linear function of the pressure  $\Delta A_{1(0,n)}p$ . The remaining data set comprised approximately 6000 measurements.

Figure 7 displays the deviations of the data from equation (1) for 5 modes measured during runs 1 and 2 using the parameters in table 4 that resulted from the iteration with h = 0.39. Each plotted point represents the average of approximately 70 measurements for a single mode at nearly-identical temperatures and pressures. Each uncertainty bar is the standard deviation of the 70 measurements from their mean.

In figure 7, the relative standard deviation of the averaged data from their mean is  $1.51 \times 10^{-6}$ , which corresponds to 37  $\mu$ K. The differences between the two runs is a quantitative indicator of the long-term (12 days) stability of the complete measurement system, including any possible impurities in the helium. At each pressure, the relative standard deviation of the data for the five modes from their average value was  $0.23 \times 10^{-6}$ , which corresponds to 5.7  $\mu$ K. This is a measure



**Figure 8.** The excess half-widths of the resonances scaled by the factor  $2 \times 10^6/f$  as a function of pressure. Each plotted point represents the average of approximately 70 measurements; the uncertainty bars are the standard deviation of the 70 measurements from their mean.

of the stability of the measurement system during intervals of 12 h.

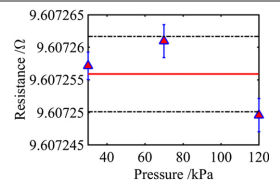
Inspection of table 4 reveals that the uncertainties of  $w^2$ ,  $\Delta A_{(0,n)}$ , and  $A_2$ , as determined by fitting, are much smaller than the differences between the best-fit values of these parameters with h = 0.39 and with h = 0.34. Thus, our assumption that  $h = (0.39 \pm 0.05)$  is crucial for determining the uncertainty  $T_{\text{Ne}}$ .

It is satisfying that the best-fit value of  $A_2$  is consistent with the *ab initio* calculation by Garberoglio *et al* [27] Under the assumption  $h = (0.39 \pm 0.05)$ , the uncertainty  $u(A_2)_{\text{measured}}$  is 38% of  $u(A_2)_{\text{theory}}$ .

#### 4. Quality tests

Figure 8 displays a powerful test of our understanding of the acoustic resonator. The theory for the half-widths  $g_{(0,n)}$ of the acoustic resonances contains no adjustable parameters. If the theory and the half-width measurements were perfect, the excess half-widths  $\Delta g_{(0,n)} \equiv (g_{\text{measured}} - g_{\text{theory}})$  would be zero at all pressures, within combined uncertainties. Any phenomenon that removes energy from an acoustic resonance will increase  $g_{(0,n)}$ ; therefore, values of  $\Delta g_{(0,n)} > 0$  are measures of imperfect understanding of the resonator. Figure 8 plots averaged values of  $\Delta g_{(0,n)}$  measured during runs 1 and 2 after scaling by the factor  $2 \times 10^6 / f_{(0,n)}$ . With this scale factor, non-zero values of  $2\Delta g_{(0,n)}/f_{(0,n)}$  are comparable to fractional errors in  $T_{\text{Ne}}$ , expressed in parts per million. (Similarly, fractional errors in  $2 \times 10^6 \Delta f_{(0,n)}/f_{(0,n)}$  correspond to ppm errors in  $T_{\text{Ne}}$ .)

At corresponding pressures, the averaged values of  $\Delta g_{(0,n)}$ in runs 1 and 2 differed from each other by only a few tenths of a part per million. This reproducibility is comparable to that



**Figure 9.** Pressure dependence of the resistance of the Pt–Co thermometer at  $T_{\text{Ne}}$ . The central horizontal red line is the mean resistance; the outer dashed lines are separated from the mean resistance by  $\pm 1$  standard deviation corresponding to  $\pm 0.036$  mK.

**Table 5.** Standard uncertainties (in mK) for calibrating a Pt–Co thermometer with the AGT near  $T_{Ne}$ .

Source	mK
Microwave determination of $a_{eq}^2$	
Repeatability of runs 1 and 2	0.002
TM11 and TM12 inconsistency	0.049
Electrical conductivity of shell	0.10
Fitting $a_{eq}(p)$	0.007
Acoustic determination of $w_0^2/a_{eq}^2$	
Compare $h = 0.39$ with $h = 0.34$	0.055
Repeatability between runs 1 and 2	0.010
Pressure fluctuation	0.002
Impurity	0.007
Fitting $w_0^2$	0.005
Thermal	
Repeatability between runs 1 and 2	0.004
Temperature stability, gradient, fluctuations	0.026
Sensor	
Bridge and standard resistor	0.004
Reproducibility between 2018 and 2019	0.08
Repeatability and self-heating of the Pt-Co	0.049
Upper bound to pressure dependence	0.036
Combined standard uncertainty $(k = 1)$	0.16

of  $w^2$  in figure 7. (Figures 7 and 8 have equivalent scales.) We fitted the values of  $2\Delta g_{(0,n)}/f_{(0,n)}$  to linear functions of the pressure. The zero-pressure intercepts for the modes (0,2), (0,3), (0,4), and (0,5) range from 0.4 ppm to 1.8 ppm. (1.8 ppm corresponds to 0.044 mK, which is much smaller than  $u(T_{\text{Ne}}) = 0.24$  mK.) Apparently, an unknown, pressure-independent energy loss mechanism significantly increases  $\Delta g_{(0,9)}$ ; however it does not affect  $f_{(0,9)}$  as it is show in figure 7.

In prior realizations of AGT, including the use of this resonator filled with argon at  $T_{\text{TPW}}$ , it was found that  $\Delta g_{(0,n)}$  for many modes increases linearly with pressure [16]. Usually, this increase is attributed to energy losses associated with oscillations of the resonator's shell driven by the radial acoustic oscillations of the gas within the shell. **Table 6.** Standard uncertainties (in mK) for measuring  $T_{\text{Ne}}$ , including transfer of the Pt–Co thermometer from the neon triple-point cells to AGT.

Source	mK	
Microwave determination of $a_{eq}^2$ (table 5)	0.11	
Acoustic determination of $w_0^2/a_{eq}^2$ (table 5)	0.057	
Thermal (table 5)	0.026	
Sensor (table 5)	0.11	
$T_{\rm Ne}$ from ITS-90 realization at LNE		
Triple point value	0.10	
Isotopic composition	0.02	
Temperature stability, gradient, fluctuations	0.08	
Interpretation of the plateau	0.03	
Electrical measurements	0.10	
Combined standard uncertainty $(k = 1)$		

In common with other realizations of AGT, problems in measuring  $g_{(0,n)}$  develop as the pressure is decreased. The values of  $g_{(0,n)}$  increase (in accord with theory); this increases the overlap of the (0, n) modes with nearby non-radial modes. The increasing overlap with decreasing pressure increases the uncertainty of fitting the resonance frequency and half-widths and introduces bias. The bias has been documented when the (0, 7) mode begins to overlap with the (13, 2) mode [33].

To test for the presence of undetected impurities in the helium, we compared the acoustic resonance frequencies with and without operating the liquid-helium cooled trap shown in figure 4. This comparison is sensitive to impurities at the subppm level and essentially insensitive to the corrections to the acoustic frequencies, the geometry of the shell, etc. First, with helium that passed through the cold trap at ambient temperature, we measured the acoustic resonance frequencies of the cavity near  $T_{\rm Ne}$  at pressures ranging from 30 kPa to 120 kPa. From this isotherm, we deduced a value of  $T_{\text{Ne,mix}}$  and we measured the resistance of a cSPRT that had a well-known history and sensitivity. To clean the system, we evacuated the pressure vessel and then flushed it three times with helium that had passed through the helium-filled cold trap. Using this 'purified' helium, we remeasured the cSPRT and the acoustic resonance frequencies at two pressures: 50 kPa and 70 kPa. (To save time and helium, we did not use other pressures). We found that the difference between our acoustic determinations of  $T_{\rm Ne}$ , by alternatively flowing in the apparatus a presumably slightly impure mixture  $T_{mix}$ , or by using the cold-trap  $T_{pure}$ , was  $\Delta T = T_{
m mix} - T_{
m pure} = -0.028$  mK at 50 kPa and  $\Delta T =$ -0.024 mK at 70 kPa with an uncertainty of 0.007 mK in both cases. These observations would be consistent with the presence in the mixture of traces of neon with a mole fraction  $2.6 \times 10^{-7}$ . In order to account for this estimate, a correction of 0.026 mK was added to obtain the thermodynamic temperature of  $T_{\rm Ne}$  previously listed in table 1

#### 5. Pt-Co thermometer

To transfer  $T_{\text{Ne}}$  to the SPRIGT apparatus, we calibrated the standard platinum–cobalt resistance thermometer, serial number RS144.08, manufactured by Chino Corporation in 2016.

The characteristics and stability of Pt-Co thermometers have been studied by the National Metrology Institute of Japan [16, 34, 35]. They showed the performance of Pt-Co thermometers is comparable to that of rhodium-iron (Rh-Fe) resistance thermometers. Before the present calibration, we checked the stability of RS144.08 at the triple point of neon at LNE-Cnam in 2018 and 2019 in a cryostat describe in [8]. The difference between the two calibrations was 0.14(26) mK, which is indistinguishable from zero. We used this difference as an estimate of possible drift of the Pt-Co sensor during one year. Assuming a rectangular distribution, we added 0.080 mK to the uncertainties in table 5. During our two AGT runs, we compared the Pt-Co thermometer to other thermometers mounted on the quasi-spherical resonator. The comparisons included a Rh-Fe thermometer (serial number 229 080) calibrated by NPL and a cSPRT (serial number B398) calibrated by LNE-Cnam. These diverse thermometers gave mutually consistent temperatures, thereby providing evidence that the drift of the Pt–Co thermometer at  $T_{\rm Ne}$  did not make an important contribution to the uncertainty of  $T_{\rm Ne}$ . After the Pt-Co was installed in the cryostat described in section 2.1, we measured its self-heating at  $T^*$  while it was immersed in helium at 30 kPa 70 kPa, and 120 kPa. When extrapolated to zero current and zero pressure, the resistance of the Pt-Co thermometer was 9.607 2558  $\Omega$  with a standard deviation of  $6.3 \times 10^{-6} \Omega$  (figure 9), which corresponds to 0.036 mK. We included this value in the uncertainties in table 5.

## 6. Uncertainty of $T_{Ne}$ and of calibrating a Pt–Co thermometer with AGT

Table 5 lists the uncertainty contributions greater than 1  $\mu$ K for calibrating Pt–Co thermometer RS144.08 in the present AGT. The uncertainty is dominated by the drift of the thermometer's resistance during an interval of one year. Less important (but non-negligible) uncertainties resulted from the inconsistent microwave results, the uncertain value of the accommodation coefficient *h*, and the thermometer's self-heating in helium at the pressures 30 kPa, 70 kPa and 120 kPa. The sum in quadrature of all the uncertainties in table 5 is 0.16 mK. This is the uncertainty of calibrating the Pt–Co thermometer using the present AGT at a thermodynamic temperature near  $T_{\text{Ne}}$ . In [7], we reported a value of  $T^*$  for the calibration of the same Pt–Co that differs from the present value by only 26% of its standard uncertainty in k = 1. We consider this difference to be insignificant.

At LNE-Cnam in 2018 and in 2019, we used the cryostat and triple point cell described in [8] to study the stability of the Pt–Co thermometer RS144.08 at  $T_{\rm Ne}$ . Table 6 adds the uncertainties from this interlaboratory comparison of a neon cell to the uncertainties of the AGT calibration of the Pt–Co thermometer in table 5. The two largest additions are 'triple point value' and 'electrical measurements'. 'Triple point value' is the documented reproducibility of neon triple points during 2012 interlaboratory comparisons [36]. It is likely that future work can reduce this uncertainty to 0.015 mK (reproducibility of the fix point has been substantially improved and LNE-Cnam has recently started a series of bilateral intercomparisons [8]). The item 'electrical measurements' in table 6 includes the table 5 items 'bridge and standard resistor' and 'repeatability and self-heating of the Pt–Co'. However, 'electrical measurements' refers to the bridge, standard resistor, and cryostat that were parts of the inter-comparison apparatus, not the present AGT.

The sum in quadrature of all the uncertainties listed in table 6 is 0.24 mK. This is the uncertainty of the present determination of the thermodynamic temperature of the triple point of neon  $T_{\text{Ne}}$ .

#### 7. Conclusion

We used a quasi-spherical AGT to calibrate the Pt–Co RS144.08 resistance thermometer with the result: (9.607255  $8 \pm 0.0000063$ )  $\Omega$  at the thermodynamic temperature  $T^* = (24.55499 \pm 0.00016)$  K. Prior to this calibration, the same Pt–Co thermometer had been compared with other thermometers that had been calibrated at the triple point of neon. By relying on the comparisons, the present AGT redetermines the thermodynamic temperature of the triple point of neon with the result  $T_{\text{Ne}} = (24.55515 \pm 0.00024)$  K. (Because of the uncertainty of the comparisons, the uncertainty of  $T_{\text{Ne}}$  is larger than the uncertainty of  $T^*$ .) After calibration, we sent the Pt–Co thermometer to the Technical Institute of Physics and Chemistry of the Chinese Academy of Sciences in Langfang to be used as a reference for single-pressure refractive-index gas thermometry.

In this work, the uncertainty u(h) is the largest contributor to the uncertainty of the acoustic determination of  $w_0^2/a_{eq}^2$  but only a minor contributor to  $u(T_{Ne})$  and to the uncertainty of the calibration of the Pt–Co thermometer RS144.08 at  $T_{\rm Ne}$ . Based on AGT results for helium-4 at temperatures from 236 K to 430 K, we assumed  $h = 0.39 \pm 0.05$  at  $T_{\rm Ne}$ . If new measurements surprise us by showing h = 1, our value of  $T_{\text{Ne}}$ will increase by 0.23 mK, which is 0.95 times the estimated standard uncertainty of  $T_{\rm Ne}$  in table 6. We had to make *some* assumption concerning h because, in the expansion of  $w^2(p)$ , the ratio of the terms  $(A_2p^2)/(A_{-1}/p)$  is approximately 50 times larger at  $T_{\text{Ne}}$  than at  $T_{\text{TPW}}$ . (See table 3.) Much of this factor of 50 resulted from our decision to use gas densities at  $T_{\rm Ne}$ that were approximately twice those used at  $T_{\text{TPW}}$ . If the densities ( $\rho$ ) had been the same at both temperatures, the quality factors (Qs) of the resonances would have been approximately the same; however, the signal-to-noise ratio, which scales as  $pQ^2 = \rho RTQ^2$  for our capacitive transducers [30], would have been reduced by the factor  $T_{\rm Ne}/T_{\rm TPW} \approx 0.090$ . Simply stated: we went to higher densities to recover some of the signalto-noise ratio that we lost by going to lower temperatures. Future acoustic measurements will be possible at lower pressures, either by increasing integration times and/or by using transducers systems optimized for lower pressures.

#### **Electronic supplement**

Supporting information for this article comprises the original acoustic data for run 1 and 2. We tabulate the corrected resonance frequencies for 7 separate acoustic modes as a function of pressure on isotherm  $T^*$ . These are available online at https://stacks.iop.org/MET/58/045006/mmedia.

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