Equations of state for the thermodynamic properties of binary mixtures for helium-4, neon, and argon

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(Dated: 12 March 2020)

Based upon the conceptual design reports for the Future Circular Collider cryogenic system, the need for more accurate thermodynamic property models of cryogenic mixtures of noble gases was identified. Both academic institutes and industries have identified the lack of a reliable equation of state for mixtures used at very low temperatures. Detailed cryogenic architecture modeling and design cannot be carried out without accurate fluid properties. Therefore, the helium-neon equation was the first goal of this work and it was further extended to other fluids beneficial for scientific and industrial applications beyond the particle physics needs. The properties of the noble gas mixtures of helium-neon, neon-argon, and helium-argon are accurately modeled with the equations of state explicit in the Helmholtz energy.

Key words: helium-4; neon; argon; equation of state; fundamental equation; thermodynamic properties.

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1. INTRODUCTION

The need for equations of state (EOS) capable of accurate representation of thermodynamic properties for cryogenic mixtures increases as more complex and more efficient cryogenic systems are being investigated and developed. Mixtures are presently considered as working fluids not only for large cryogenic systems but also for smaller, space-ready refrigerators. The potential applications of equations of state for noble gases include cooling cycles for fusion reactors or particle accelerators such as the Future Circular Collider,² (FCC) space refrigerators, and modeling of welding processes with inert gases.

No accurate model has been developed so far for binary mixtures of helium-4, neon, and argon.³ Until now, either linear or Lorentz-Berthelot^{4,5} classical rules for combining fluids have been used.^{6,7} Neither of them, however, represents quantum gas behavior correctly.⁸ Following the best practices from the domain of the equations of state, the quantum effects were included in the empirical model development with no special treatment.

The mixture models presented in this paper are explicit in the Helmholtz energy and use reducing parameters dependent on the mole fractions of mixture constituents and the critical parameters of the pure fluids. The Helmholtz energy a is defined as follows

$$a = u - Ts, \tag{1}$$

where u is internal energy, T is temperature and s is entropy. Since it is a fundamental value, all other thermodynamic properties can be calculated from analytical derivatives. The Helmholtz energy formulation takes advantage of the consistent description of all fluid states because temperature and density are the independent variables. The pure fluid contributions, and the contribution from mixing for each binary mixture use an empirical formulation, which is temperature, density, and molar composition-dependent. The pure fluid contributions take advantage of the highly accurate EOS of pure fluids. In this work, each mixture-specific departure function consists of three exponential terms and four to five Gaussian terms. Comparisons to available experimental data are given to validate the accuracy of thermodynamic properties calculated with each equation of state.

The accuracy of mixture models is dependent on the pure fluid models. The equation of state for the considered mixture constituents, helium-4, neon, and argon have state-of-theart accuracy. This important starting point allows for the development of mixture models that will become the industrial standard. The critical parameters of the pure fluids, used in Helmholtz energy equations of state (HEOS) are presented in Table 1. Additionally, the equations, in the form presented in this paper, can be implemented in fluid property modeling software, such as REFPROP,⁹ CoolProp,¹⁰ or TREND.¹¹

TABLE 1: Molar mass and critical parameters for pure fluids

	$M / (\text{kg mol}^{-1})$	<i>T</i> _c / K	$p_{\rm c}$ / MPa	$\rho_{\rm c} / ({\rm mol} \ {\rm dm}^{-3})$
⁴ He ¹²	4.0026×10^{-3}	5.195	0.22832	17.384
Ne ¹³	20.179×10^{-3}	44.400	2.66163	24.100
Ar ¹⁴	39.948×10^{-3}	150.687	4.86300	13.407

TABLE 2: Reducing temperature and density parameters

Yr	$Y_{c,i}$	$\beta_{Y,ij}$	Y_{ij}
T _r	$T_{\mathrm{c},i}$	$\beta_{T,ij}$	$\beta_{T,ij}\gamma_{T,ij} \left(T_{\mathrm{c},i}T_{\mathrm{c},j}\right)^{0.5}$
v _r	$rac{1}{ ho_{\mathrm{c},i}}$	$eta_{v,ij}$	$\beta_{\nu,ij}\gamma_{\nu,ij}\frac{1}{8}\left(\frac{1}{\rho_{\mathrm{c},i}^{1/3}}+\frac{1}{\rho_{\mathrm{c},j}^{1/3}}\right)^{3}$

2. FUNDAMENTAL EQUATION OF STATE

Among the available formulations of equations of state, the Helmholtz energy $a(T, \rho)$ has two important advantages. Unlike internal energy u(v,s) and enthalpy h(p,s), it is a function of measurable properties - temperature *T* and density ρ . Moreover, unlike the Gibbs energy g(T, p), it is continuous within the phase boundary, where *v* is specific volume, *s* is specific entropy, and *p* is pressure.

The equation of state is used in a reduced form and is a function of non-dimensional quantities - reduced density and reciprocal reduced temperature^{15,16}

$$\tau = \frac{T_{\rm r}}{T} \quad \text{and} \quad \delta = \frac{\rho}{\rho_{\rm r}},$$
(2)

where T_r and ρ_r are the reducing temperature and density. The most recent formulation^{16,17} allows for the use of a common form for reducing parameters for the mixture (T_r and $v_r = 1/\rho_r$)

$$Y_{\rm r}(\bar{x}) = \sum_{i=1}^{N} x_i^2 Y_{{\rm c},i} + \sum_{i=1}^{N} \sum_{j=i+1}^{N} 2x_i x_j \frac{x_i + x_j}{\beta_{Y,ij}^2 x_i + x_j} Y_{ij}, \quad (3)$$

where *Y* is the parameter of interest, temperature *T* or molar specific volume *v* given in Table 2. $T_{c,i}$, $\rho_{c,i}$ are critical temperature and density of the pure constituents. $\beta_{T,ij}$, $\gamma_{T,ij}$, $\beta_{v,ij}$, and $\gamma_{v,ij}$ are the fitted parameters with both β parameters being asymmetric, that is $\beta_{T,ij} = \beta_{T,ji}^{-1}$, and $\beta_{v,ij} = \beta_{v,ji}^{-1}$.

The total reduced Helmholtz energy α can be divided into an ideal gas part α^{o} and a residual part α^{r}

$$\alpha(\delta,\tau,\bar{x}) = \frac{a(\rho,T,\bar{x})}{RT} = \alpha^{o}(\rho,T,\bar{x}) + \alpha^{r}(\delta,\tau,\bar{x}), \quad (4)$$

where \bar{x} is the mole fraction vector, and R = 8.314462618 J mol⁻¹ K⁻¹ is the gas constant.¹⁸ The ideal and residual contributions represent the Helmholtz energy for the mixture. Both are functions of the pure fluid Helmholtz energies

$$\boldsymbol{\alpha}^{\mathrm{o}}(\boldsymbol{\rho}, T, \bar{x}) = \sum_{i=1}^{N} x_i \left[\boldsymbol{\alpha}_{\mathrm{o}i}^{\mathrm{o}}(\boldsymbol{\rho}, T) + \ln x_i \right], \tag{5}$$

$$\alpha^{\rm r}(\delta,\tau,\bar{x}) = \sum_{i=1}^{N} x_i \alpha_{\rm oi}^{\rm r}(\delta,\tau) + \sum_{i=1}^{N-1} \sum_{j=i+1}^{N} x_i x_j F_{ij} \alpha_{ij}^{\rm r}(\delta,\tau), \quad (6)$$

where x_i is the molar fraction of component *i*, α_{oi}^{o} is the pure fluid ideal Helmholtz energy, and α_{oi}^{r} is the pure fluid residual Helmholtz energy, both taken from the pure fluid EOS. F_{ij} is the scaling parameter applied to the departure function, and here is set to unity because the departure functions have been fitted in this work for each binary mixture. $\alpha_{ij}^{r}(\delta, \tau)$ is a pairwise departure function, dependent only on reduced variables δ and τ .

The definition of the Helmholtz energy for mixtures allows to keep the pure fluid ideal α_{oi}^{o} and residual α_{oi}^{r} contributions unchanged as defined for the pure constituents. The mixture models presented in this paper are valid only with the pure fluid EOS used for their development. If new, and more accurate EOS for one of the constituents is developed, the equations for mixtures presented here may lose their validity and most probably must be refitted according to the updated pure fluid equations of state.

The last, previously undefined part of the Helmholtz energy for mixtures is the departure function, which depends on the reduced variables, and the adjustable parameters. Unlike in the GERG-2008 definition,¹⁶ the departure function used in this work is temperature dependent

$$\alpha_{ij}^{\mathrm{r}}\left(\delta,\tau\right) = \sum_{k=1}^{8} n_{ij,k} \delta^{d_{ij,k}} \tau^{t_{ij,k}} \\ \times \exp\left(-\eta_{ij,k} \left(\delta - \varepsilon_{ij,k}\right)^2 - \beta_{ij,k} \left(\tau - \gamma_{ij,k}\right)^2\right), \quad (7)$$

where $d_{ij,k}$, $n_{ij,k}$, $t_{ij,k}$, $\beta_{ij,k}$, $\gamma_{ij,k}$, $\varepsilon_{ij,k}$, and $\eta_{ij,k}$ are fitted parameters. Although they are empirical and arbitrary, constraints on their values were used during the fitting process in order to obtain physically correct EOS behavior (see Supplemental materials). The summation comprises the polynomial terms, and the Gaussian bell-shaped terms.

2.1. Optimization algorithm

The algorithm used for the equations of state development is based on non-linear regression analysis and the Levenberg-Marquardt method.^{19,20} It was used to minimize the sum of squares of weighted data points, and constraints with applied limits on derivatives of the Helmholtz energy in Eq. (4). The expression for residual sum of squares is given as

$$SSQ = \sum_{i=1}^{N_{\rho}} W_{\rho,i} F_{\rho,i}^{2} + \sum_{i=1}^{N_{w}} W_{w,i} F_{w,i}^{2} + \sum_{i=1}^{N_{cv}} W_{cv,i} F_{cv,i}^{2} + \dots, \quad (8)$$

where W_x is the weight assigned to each data point of the thermodynamic property x, and F_x is the function used to minimize the relative deviation between the data points and the equation of state.²¹ Deviations do not have to be evaluated as the sum of squares. During the fitting process, the method for calculating the contribution to the total sum of deviations was changed periodically to use the sum of squares, cubes, fourthor sixth-powers to find the best possible solution.

Phase equilibrium data is first fitted with the algorithm, and later, when satisfactory agreement between the data and the model is reached at the phase boundary, the single-phase data points are used to improve the model (e.g., ρpT , speed of sound, etc.). The functional behavior is constantly constrained by applying weights on limits for the coefficients. All data points and applied constraints are provided in the supplemental materials.

The quality of the equation of state is determined through comparison of calculated deviations of data points and values from the equation of state. The errors for single-phase properties were evaluated numerically and deviations between the calculated values and available data points are discussed in section 3 3.1. The deviations at the phase equilibria, vaporliquid equilibrium (VLE) and gas-gas equilibrium (GGE) if it exists, were evaluated not only numerically but also visually by comparing the available data points with the isotherms calculated with the equation of state as shown in Fig. 1. A new approach, useful for the phase boundary uncertainty discussion, is presented in section 3 3.2.

2.2. Available data used for the equation of state development

The experimental data on thermophysical properties available in the literature were collected and are presented in Tables 3, 4, and 5, along with the number of data points in every data set and the number of points used in the EOS development process. When the experimental points were omitted (available vs. used data points for single entry from literature), it is because of either incomplete information needed for the EOS development or questionable accuracy.

Average absolute deviations in Tables 3, 4, and 5 were calculated as follows

$$AAD = \frac{100}{N} \sum_{i}^{N} \left| \frac{x_{d,i} - x_{EOS,i}}{x_{d,i}} \right|, \qquad (9)$$

where $x_{d,i}$ is a data point, $x_{EOS,i}$ is a respective value calculated from the equation of state, and *N* is the number of points in data set. If the data set was not used in the fitting process, the AAD is printed in bold.

	data po	ints					
authors	available	used	<i>T /</i> K	p / MPa	x _{He}	author claimed uncertainty	AAD ^a / %
					VLE data		
Heck, Barrick ²²	76	36	27 - 42	0.3 - 20	0.01 - 0.36	$\pm 3\%$ in p	15.0
22						$\int \pm 0.02 \text{ K in } T$	
Knorn ²³	22	22	25 - 27	0.6 - 5.1	0.002 - 0.03	$\{ \pm 0.02\% \text{ in } x \}$	4.3
						$\pm 0.1\%$ in y	
					ρpT data		
Holborn, Otto ²⁴	39	39	273 - 673	2.5 - 10	0.28	N/A ^b	0.26
Kestin et al. ²⁵	97	97	298	0.2 - 12.2	0.23, 0.80	$\pm 0.1\%$ in $ ho$	0.20
Kestin, Nagashima ²⁶	90	73	293 - 303	0.1 - 6.3	0.26 - 0.95	$\pm 0.1\%$ in $ ho$	0.24
Richardson, Leachman ²⁷	8	6	32 - 41	0.5 - 2.1	0.02 - 0.03	N/A ^b	2.0
Vogl, Hall ²⁸	51	51	233 - 313	0.1 - 3.6	0.49	$\pm 0.04\%$ in Z^{c}	0.081
-				Sp	eed of sound - w		
Pashkov et al. ²⁹	41	36	26 - 38	0.1 - 15	0.01 - 0.29	$\pm 0.1\%$ in w	9.6
				Second	l virial coefficient - B		
Berman et al. ³⁰	3	0	15 - 20			$\pm 5\%$ in B	55.3
Brewer, Vaughn ³¹	8	0	148 - 323			$\pm 5\%$ in B	3.9
Holborn, Otto ²⁴	5	0	273 - 673			N/A ^b	8.8
Iomtev et al. ³²	6	0	15 - 20			$\pm 6\%$ in B	57.6
Knobler et al. ³³	1	0	90			$\pm 3\%$ in B	5.9
Vogl, Hall ²⁸	3	0	233 - 313			$\pm 4\%$ in B	3.6

TABLE 3: Available and used data for the development of the equations of state for ${}^{4}\text{He} - \text{Ne}$

 $\frac{1}{a}$ average absolute deviation (AAD) of the data set from values calculated with EOS given in Eq. (9)

^b undefined or poor uncertainty analysis ^c compressibility factor

TADLE 4	A		(. C 1	11	. C (1		. 6	c 4	TT.	
IABLE 4:	Available a	na usea aa	ta for the	e development	of the	equations	of state	lor -	He –	Ar

	data po	oints					
authors	available	used	<i>T /</i> K	<i>p</i> / MPa	x _{He}	author claimed uncertainty	AAD ^a / %
				V	LE / GGE data	(
Mullins ³⁴	354	0	68 - 108	2.0 - 12	0.001 - 0.02	$\begin{cases} \pm 0.03 \text{ K m } T \\ \pm 0.5\% \text{ in } p \\ \pm 2\% \text{ in } x \\ \pm 3\% \text{ in } y \end{cases}$	24.0
Mullins ³⁵	187	187	85 - 140	0.5 - 12	0.00006 - 0.02	$\begin{cases} \pm 0.03 \text{ K in } T \\ \pm 0.5\% \text{ in } p \\ \pm 2\% \text{ in } x \\ \pm 3\% \text{ in } y \end{cases}$	3.4
Mullins, Ziegler ³⁶	50	0	86 - 108	2.0 - 12	0.002 - 0.02	$\begin{cases} \pm 0.03 \text{ K in } T \\ \pm 0.5\% \text{ in } p \\ \pm 1 - 2\% \text{ in } x, y \end{cases}$	3.6
Sinor, Kurata ³⁷	29	0	93 - 148	1.7 - 14		$\begin{cases} \pm 0.5\% \text{ in } p \\ \pm 1\% \text{ in } x, y \end{cases}$	12.0
Skripka, Lobanova ³⁸	77	64	90 - 115	2.9 - 22	0.0005 - 0.002	± 0.02 vol. % in x, y	3.5
Streett ³⁹	56	48	91 - 148	1.4 - 69	0.002 - 0.44	$\begin{cases} \pm 0.02 \text{ K in } T \\ \pm 2 \text{ mol } \% \text{ in } x, y \\ \leftarrow 0.1\% \text{ in } n \end{cases}$	15.0
Streett, Hill ⁴⁰	202	176	91 - 160	0.6 - 422	0.02 - 0.60	$\begin{cases} \pm 0.1 \text{ mm } p \\ \pm 0.01 \text{ K in } T \\ \pm 0.1 \text{ mol } \% \text{ in } r \text{ y} \end{cases}$	15.0
Streett, Erickson ⁴¹	81	74	150 - 199	386 - 1048	0.27 - 0.72	N/A ^b	7.7
					ho pT data		
Blancett et al. ⁴²	288	288	223 - 323	0.3 - 73	0.21 - 0.80	$\begin{cases} \pm 0.03 \text{ K in } T \\ \pm 0.06\% \text{ in } Z \end{cases}$	0.72
Kanishev, Pitaevskaya43	41	41	373, 473	50 - 180	0.66, 0.95	N/A ^b	1.8
Kosov, Brovanov ⁴⁴	59	59	293 - 353	29 - 59	0.24 - 0.79	$\begin{cases} \pm 0.05 \text{ K in } T \\ \pm 0.3\% \text{ in } \rho \end{cases}$	2.2
Maslennikova <i>et al.</i> ⁴⁵	116	116	293 - 423	180 - 800	0.31, 0.74	$\pm 0.5\%$ in ρ	2.4

^a average absolute deviation (AAD) of the data set from values calculated with EOS given in Eq. (9)

Continued on next page...

^b undefined or poor uncertainty analysis

Popov <i>et al.</i> ⁴⁶	45	45	293 - 673	5.0 - 40	0.11 - 0.98	$\begin{cases} \pm 0.05 \text{ K in } T \\ \pm 0.05\% \text{ in } p \\ \pm 0.02\% \text{ in } x_{\text{He}} \end{cases}$	1.3
Provine, Canfield ⁴⁷	212	212	143 - 183	0.2 - 68	0.22 - 0.80	N/A ^b	1.2
Vidal <i>et al</i> . ⁴⁸	31	31	298	0.1 - 800	0.50	$\pm 0.2\%$ in $ ho$	1.5
Vidal <i>et al</i> . ⁴⁹	13	13	298	80 - 800	0.50	$\pm 0.2\%$ in $ ho$	1.9
				Spe	eed of sound - w		
Hanayama ⁵⁰	51	27	298	109 - 1972	0.50 - 0.90	$\pm 0.5\%$ in w	5.1
Kachanov et al. ⁵¹	22	20	298 - 422	100 - 700	0.21	$\pm 0.4\% - 0.63\%$ in w	1.3
Nishitake, Hanayama ⁵²	47	47	298	201 - 1696	0.50 - 0.90	$\pm 0.5\%$ in w	11.0
Vidal <i>et al.</i> ⁴⁹	13	13	298	80 - 800	0.50	$\pm 0.2\%$ in w	2.4
				Second	virial coefficient - B		
Blancett et al. ⁴²	15	0	223 - 323			$\begin{cases} \pm 0.03 \text{ K in } T \\ \pm 0.3\% \text{ in } B \end{cases}$	21.0
Brewer, Vaughn ³¹	8	0	148 - 323			$\pm 5\%$ in B	11.0
Kalfoglou ⁵³	60	0	303 - 773			$\pm 0.86\% - 1.31\%$ in <i>B</i>	16.9
Knobler et al. ³³	1	0	90			$\pm 2 - 3\%$ in <i>B</i>	6.1
Martin ⁵⁴	3	0	290 - 320			greater than 1.2%	5.9
Tanner ⁵⁵	14	0	298			N/A ^b	18.9

TABLE 4 - continued from previous page

^a average absolute deviation (AAD) of the data set from values calculated with EOS given in Eq. (9)

^b undefined or poor uncertainty analysis

	data noi	nts					
authors	available	used	<i>T </i> K	p / MPa	x _{Ne}	author claimed uncertainty	AAD ^a / %
					VLE data		
Skripka, Lobanova ³⁸	84	54	91 - 120	2.9 - 20	0.03 - 0.31	± 0.02 vol. % in x, y	8.0
Streett ⁵⁶	54	54	84 - 130	0.4 - 7.3	0.05 - 0.73	$\begin{cases} \pm 0.01 \text{ K in } T \\ \pm 0.01 \text{ MPa in } p \end{cases}$	11.0
Streett ⁵⁷	37	36	95 - 130	7.5 - 62	0.09 - 0.57	N/A ^b	12.0
Streett, Hill ⁵⁸	58	58	87 - 93	6.4 - 103	0.002 - 0.09	$\begin{cases} \pm 0.1\% \text{ in } p \\ \pm 0.02 \text{ K in } T \\ \pm 0.1 \text{ mol } \% \text{ in } x, y \end{cases}$	6.7
Trappeniers, Schouten ⁵⁹	67	59	93 - 138	0.5 - 101	0.002 - 0.57	$\pm 0.1 \text{ mol } \% \text{ in } x, y$	13.0
					ho pT data		
Kestin et al. ²⁵	21	21	298	0.4 - 13.7	0.23, 0.73	$\pm 0.1\%$ in $ ho$	0.43
Streett ⁵⁷	109	107	102 - 121	3.0 - 55.2	0.04 - 0.50	$\pm 1\%$ in ρ	2.5
Vidal <i>et al</i> . ⁴⁸	31	31	298	0.1 - 800	0.50	$\pm 0.2\%$ in $ ho$	1.6
Vidal <i>et al.</i> ⁴⁹	13	13	298	80 - 800	0.50	$\pm 0.2\%$ in $ ho$	2.8
				S	peed of sound - w		
Konovodchenko et al.60	10	0	102, 121	1.3 - 15	0.22	N/A ^b	60.0
Vidal <i>et al.</i> ⁴⁹	13	13	298	80 - 800	0.50	$\pm 0.2\%$ in w	1.2
				Secon	d virial coefficient - B		
Brewer, Vaughn ³¹	9	0	123 - 323			$\pm 5\%$ in <i>B</i>	9.7
Knobler et al. ³³	1	0	90			$\pm 2 - 3\%$ in <i>B</i>	5.9

TABLE 5: Available and used data for the development of the equations of state for Ne - Ar

^a average absolute deviation (AAD) of the data set from values calculated with EOS given in Eq. (9)

^b undefined or poor uncertainty analysis

Even though the virial coefficients were collected, and are presented in Tables 3, 4, and 5, they were not fitted. Physics dictates that B_{12} should only be a function of temperature, but our mixing rule also produces a dependence on composition in the multi-fluid models.

When compared to the pure fluids, the region of existing state points is always broader for mixtures, since it is also composition-dependent. It is therefore more difficult to cover a comparable number of states for mixtures as for pure fluids. The equilibrium data available for the mixture of helium-neon are limited in pressure up to 20 MPa. Therefore, the shape of the phase envelope at higher pressures is unknown. However, a pressure of 20 MPa is higher than most engineering applications, and the model shows satisfactory accuracy at lower pressures with the extrapolation behavior being acceptable. Both the equilibrium data, and ρpT data for helium-argon and neon-argon mixtures reach higher pressures up to 1 GPa for

 TABLE 6: Binary specific parameters obtained in this work

i-j	$\beta_{T,ij}$	γ T,ij	$\beta_{v,ij}$	γ _{v,ij}	Fij
4 He $-$ Ne	0.793	0.728	1.142	0.750	1.0
⁴ He – Ar	1.031	1.113	1.048	0.862	1.0
Ne – Ar	1.033	0.967	0.919	1.035	1.0

helium-argon, and 100 MPa for neon-argon.

2.3. Fluid properties scaling

Depending on the mixture, the phase envelope varies in shape; it can be either closed or open, it can either have the GGE locus or just the VLE dome. The name gas-gas equilibrium implies that a phase separation occurs beyond the critical temperature of the heavier component.⁶¹ For the heliumargon mixture, the gas-gas coexistence region detaches from the vapor-liquid region with increasing temperature. When temperature further increases, the regions separate completely (see two 148 K isotherms in Fig. 1d - VLE closing at 27 MPa, and GGE opening at 57 MPa). The vapor-liquid coexistence terminates at 150 K and for higher temperatures only the gasgas equilibria exist. Helium often shows this type of behavior when mixed with other substances.^{61,62} It is therefore probable that two other mixtures considered in this paper behave similarly but this question is not further investigated due to the lack of high-pressure data for 4 He – Ne and Ne – Ar. Scaling the properties of helium-4 and neon with any physical theory in the regions of missing data can be challenging because of the quantum phenomena influencing the fluid's behavior. Classical or quantum effects will dominate depending on the length scale of interactions vs. the thermal de Broglie wavelength

$$\lambda_{\rm th} = \sqrt{\frac{h^2}{2\pi m k_{\rm B} T}},\tag{10}$$

where *T* is the temperature at which $\lambda_{\rm th}$ is calculated, $h = 6.626\ 070\ 15 \times 10^{-34}$ J s is the Planck constant,¹⁸ $m = M/N_{\rm A}$ is the particle mass with molar mass *M* given in Table 1, $N_{\rm A} = 6.022\ 140\ 76 \times 10^{23}\ {\rm mol}^{-1}$ is the Avogadro constant,¹⁸ and $k_{\rm B} = 1.380\ 649 \times 10^{-23}$ J K⁻¹ is the Boltzmann constant.¹⁸

Quantum effects are important in determining the thermodynamics of helium, and to a lesser extent neon; argon can be considered classical for most practical purposes. Despite dealing with the fluids that show high and moderate quantum effects, the discussion on the quantum physics is secondary, since the presented equations are empirical and the question about the origins of the intermolecular forces is irrelevant.

3. RESULTS

In addition to the four reducing parameters from Table 6, the model optimization and development included deriving the departure functions according to Eq. (7). The temperature and density dependent parts of the departure functions for three binary fluid mixtures are presented in Table 7. The phase behavior of the EOS is presented in Fig. 1 and is analyzed further in this section. To illustrate the accuracy of the equations, deviations of thermodynamic properties calculated with the EOS are compared against the experimental values.

Along with the errors discussed below, each data set has a measurement uncertainty. The uncertainties claimed by the authors should be viewed skeptically, especially in light of reporting standards for uncertainty that have evolved over time. With this in mind, the uncertainties specified by the authors in their publications are listed in Tables 3, 4, and 5. The largest uncertainty for the equilibria measurements is attributed to composition and varies from 0.1% to 3% in mole fraction for bubble points *x* and dew points *y*. Regarding the density measurements, the uncertainty is often claimed to be on the order of 0.1%, except for 1% for Ne-Ar densities.

3.1. Uncertainty discussion for single-phase

Fig. 4 presents the EOS deviations from the data used in the development of each equation. The density deviations for the helium-neon equation are smaller than for the other two pairs: 0.5% dropping the lower accuracy data at 30 - 40 K. However, similarly as with the equilibrium data, the pressure range for this binary pair is also more narrow for ρpT when compared to the other two mixtures. For neon-argon, the density measurements reach 55 MPa and for helium-argon, they go up to 1 GPa. When comparing the equation error for similar low-pressure data (0 – 10 MPa), the helium-argon EOS has errors of 0.5% – 2.5% for 95% of data points used in the EOS development; and neon-argon has errors of 3% for 90% of density data.

The deviations in density at lower pressures for each EOS are comparable with the measurement uncertainties presented in Tables 3, 4, and 5. However, for pressures above 20 MPa, the equations deviate from data more than the specified uncertainties.

At the same time, the temperature regions of data availability has to be taken into consideration. For the helium-argon mixture the data set is broad, not only in pressure but also in temperature (starts close to the two-phase region and reaches 573 K). It is not the case for helium-neon, which is missing accurate low-temperature data, nor for neon-argon, where density measurements are narrow in temperature range. The exact pressure and temperature ranges are presented in Tables 3, 4, and 5 and Fig. 4.

3.2. Uncertainty discussion at phase boundary

When applying the isothermal error calculations based on the comparison of the bubble or dew pressures for a given mixture composition, very large calculated deviations can be obtained when the isotherm in the $\log p - x$ plane is vertical or very steep, although the data point may be still very close to



(a) Helium-neon closed phase envelope



(b) VLE isotherms for helium-neon with experimental data plotted in respective temperature color code



(c) Class III⁶¹ phase envelope of helium-argon with GGE opening



experimental data



(f) VLE isotherms for neon-argon and experimental data

FIG. 1: The phase envelopes for three binary mixtures shown in three-dimensional space of p - T - x with isobars in red, and isotherms in black (a, c, e). Their projection on 2D plane of p - x (b, d, f) constructed with the algorithm for equilibria of binary mixtures¹.

			⁴ H	e – Ne							
k	N_k	t_k	d_k	η_k	eta_k	γ_k	ϵ_k				
1	-4.346 849	1.195	1	0	0	0	0				
2	-0.884 378	1.587	2	0	0	0	0				
3	0.258 416	1.434	3	0	0	0	0				
4	3.502 188	1.341	1	0.157	0.173	1.310	1.032				
5	0.831 330	1.189	2	0.931	1.070	1.356	1.978				
6	2.740 495	1.169	3	0.882	0.695	1.596	1.966				
7	-1.582 230	0.944	4	0.868	0.862	1.632	1.709				
8	-0.304 897	1.874	4	0.543	0.971	0.766	0.583				
-	⁴ He – Ar										
k	N_k	t_k	d_k	η_k	eta_k	γ_k	ϵ_k				
1	-2.643 654	1.030	1	0	0	0	0				
2	-0.347 501	0.288	2	0	0	0	0				
3	0.201 207	0.572	3	0	0	0	0				
4	1.171 326	1.425	1	0.371	0.320	1.409	0.378				
5	0.216 379	1.987	1	0.081	1.247	1.709	0.741				
6	0.561 370	0.024	2	0.375	1.152	0.705	0.322				
7	0.182 570	1.434	3	0.978	0.245	1.162	1.427				
8	0.017 879	0.270	4	0.971	1.030	0.869	2.088				
-			Ne	e-Ar							
k	N_k	t_k	d_k	η_k	eta_k	γk	ϵ_k				
1	-1.039 686	0.723	1	0	0	0	0				
2	0.593 776	1.689	2	0	0	0	0				
3	-0.186 531	1.365	3	0	0	0	0				
4	-0.223 315	0.201	1	1.018	0.360	1.119	2.490				
5	0.160 847	0.164	2	0.556	0.373	1.395	1.202				
6	0.405 228	0.939	2	0.221	0.582	1.010	2.468				
7	-0.264 563	1.690	3	0.862	0.319	1.227	0.837				
8	-0.033 569	1.545	4	0.809	0.560	1.321	2.144				

TABLE 7: Departure function coefficients



FIG. 2: Schematic representation of the orthogonal error calculations for the phase envelopes. Two experimental compositions for liquid and vapor from one measurement are marked with circles (the same pressure and temperature) and the respective isotherm evaluated with the EOS is plotted in two-dimensional space of reduced pressure $\hat{p} = p/p_r$ and mole fraction in liquid *x* and vapor *y*.

the phase envelope. This is simply a degeneracy in the means of error quantification.

In order to evaluate the phase equilibria errors, the orthogonal length scale for data versus the EOS was defined. The error is calculated by finding the smallest distance between the experimental data point $((x_d, \hat{p}_d), T_d)$ and the respective isotherm evaluated with given EOS ($(x_c, \hat{p}_c), T_c$), where $T_d = T_c$, and $\hat{p} = p/p_r$. The orthogonal error itself should be dimensionless, therefore its evaluation in the pressure-mole fraction plane requires the pressure to also be dimensionless, therefore reduced. After comparing the impact of the reducing scales on the error representation, the logarithmic reducing scale has been selected among the linear, and density-temperature reducing scale ($p_r = p(\rho_r, T_r)$)

$$\begin{cases} p_{\rm r,x} = \exp(x_1 \ln(p_{\rm c1}) + x_2 \ln(p_{\rm c2})) \\ p_{\rm r,y} = \exp(y_1 \ln(p_{\rm c1}) + y_2 \ln(p_{\rm c2})) \end{cases}, \tag{11}$$

where $p_{r,x}$ reduces the pressure for the bubble points and $p_{r,y}$ reduces the pressure for the dew points.

The error itself is defined as a sum of distances between the data point x_d and the closest point from the respective isotherm in the earlier defined $\hat{p} - x$ plane

$$\begin{cases} \operatorname{err}_{x}^{vle} = \pm \sqrt{(x_{d} - x_{c}^{\min})^{2} + \left(\ln\left(\hat{p}_{d}\right) - \ln\left(\hat{p}_{c,x}^{\min}\right)\right)^{2}} \\ \operatorname{err}_{y}^{vle} = \pm \sqrt{(y_{d} - y_{c}^{\min})^{2} + \left(\ln\left(\hat{p}_{d}\right) - \ln\left(\hat{p}_{c,y}^{\min}\right)\right)^{2}} \end{cases}, \quad (12)$$

where x_d and \hat{p}_d are molar composition and reduced pressure for the experimental data point, and x_c^{\min} and $p_{c,x}^{\min}$ are molar composition and reduced pressure evaluated with the HEOS at



(a) Deviations in density as a function of pressure presented as percentage differences between calculated and measured density



FIG. 3: Relative deviations of the ${}^{4}\text{He}$ – Ne equation to data

(a) Deviations in density calculations for pressures up to 100 MPa in two left columns and up to 1 GPa in the right column





(a) Deviations in density calculations

FIG. 5: Relative deviations of the Ne – Ar equation to data



orthogonal error of 1.89

maximal orthogonal error of 39.99

FIG. 6: Three mixture phase envelopes with color scales representing the absolute orthogonal error values plotted for 99.99% of data points for ${}^{4}\text{He} - \text{Ne}$, 91.36% of data points for ${}^{4}\text{He} - \text{Ar}$, and 99.23% of data points for Ne - Ar. The dotted line in each figure represents the reducing pressure scale from Eq. (11)



FIG. 7: Vapor-liquid and gas-gas equilibria orthogonal error values for three binary mixtures. Color intensity increases with increasing number of points in the hexagonal unitary area. The marginal histograms represent the number of data points in different data bins in whole pressure range (x-axis) and orthogonal error range (y-axis)

the same temperature as the data point. This error definition is represented schematically in Fig. 2. The sign convention is chosen so the positive error represents the EOS calculating too large a pressure when compared to data, and negative when calculating too small a pressure. It is important to highlight that the compositions at the minimum point and experimental point can be different. Results for described error calculations are presented in Fig. 6 and 7. To our knowledge, this is the first time that an error metric like this has been used to quantify error from mixture equations of state.

While the composition for the phase equilibrium calculations is always in the range of [0, 1], the logarithm of reduced pressure $\ln(\hat{p})$ can reach values up to 7 for the absolute pressure of 1 GPa and defined pressure metric. Therefore, the error definition from Eq. (12), and the chosen pressure metric emphasize the pressure error more than the composition



FIG. 8: Cumulative percentage of points with error less than the specified value of the VLE orthogonal error for each EOS. ⁴He – Ar errors are considered for two pressure ranges:

complete data availability pressure range and up to 100 MPa with smaller orthogonal error

error. Even though all the data were fitted to the ${}^{4}\text{He} - \text{Ar}$ EOS, it largely deviates from the experimental points at very high pressures. Those high deviations are visible in Fig. 6c as the dark blue circles and in Fig. 7b as the faint hexagons with large orthogonal error. Additional error comparison for helium-argon at pressures below 100 MPa is given in Fig. 8, where it shows very satisfactory agreement between the data and the EOS in limited pressure range, and worse EOS behavior for high pressure.

3.3. Ternary mixture

With the three binary fluid mixture equations developed, the model for ternary mixture of all constituents is valid from the theoretical point of view for the Helmholtz energy EOS. Even though it is mathematically valid, no experimental data for the ternary mixture were found to compare against the equation. The accuracy of the EOS for the ternary mixture of heliumneon-argon remains unknown.

3.4. Validation data

When implementing new equations of state in any software, the EOS should be validated against the data points provided to verify if their implementation was successful. If the software uses the same pure fluid equations as given in this work (helium-4,¹² neon,¹³ and argon¹⁴), the implementation of mixture equations should return the same results in pressure calculations up to the last significant digit. The difference between the current gas constant¹⁸ and that used in the argon EOS has an effect on the calculated pressure values even at the 3rd digit. A script in the Python language is provided in the supporting information demonstrating that calculations with CoolProp (version 6.3.1) reproduce exactly the values in Table 8.

 TABLE 8: Data points at equimolar (0.5/0.5) composition for EOS validation

	Т / К	$\rho / (\text{mol dm}^{-3})$	р / Ра
4 He $-$ Ne	200.0	10.0	18 430 775.292 601
⁴ He – Ar	200.0	10.0	17 128 034.388 363
Ne – Ar	200.0	10.0	15 905 875.375 781

4. CONCLUSIONS

Three equations of state explicit in the Helmholtz energy for mixtures of noble gases with high potential of application at cryogenic temperatures were developed. The equations are not limited to the cryogenic temperatures and can be successfully used at ambient and above ambient temperatures calculations at the phase equilibrium, as well as in the single phase. The EOS performances vary for each equation depending on the pressure. The ⁴He – Ne EOS shows errors in range of uncertainties of the experimental data points for both single phase and the phase envelope. ⁴He – Ar shows satisfactory performance at pressures up to 100 MPa, and increasing errors above. The Ne – Ar EOS errors are low at the equilibrium conditions and degenerate in the single phase. Small artifacts might occur immediately above the critical points of the heavier components for given equations.

The necessary improvements introduced to the fitter let us think that development of the Helmholtz energy equations of state for further challenging mixtures should be successful.

5. SUPPLEMENTARY MATERIALS

The supplementary material contains files with data, constraints, and limits applied on the fitted function for every binary mixture. It also contains the hmx.bnc text file with developed equations, which can be implemented in REFPROP in the given form. The included python 3 script allows to validate the equations with CoolProp.

ACKNOWLEDGMENTS

This research was partially supported by EASITrain – European Advanced Superconductivity Innovation and Training. This Marie Skłodowska-Curie Action (MSCA) Innovative Training Network (ITN) has received funding from the European Union's H2020 Framework Program under Grant Agreement no. 764879.

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