# **Thermophysical Properties of Ethane**

Cite as: Journal of Physical and Chemical Reference Data **20**, 275 (1991); https://doi.org/10.1063/1.555881 Submitted: 28 November 1989 . Published Online: 15 October 2009

Daniel G. Friend, Hepburn Ingham, and James F. Fly



## **ARTICLES YOU MAY BE INTERESTED IN**

Thermophysical Properties of Fluids. II. Methane, Ethane, Propane, Isobutane, and Normal Butane

Journal of Physical and Chemical Reference Data 16, 577 (1987); https://doi.org/10.1063/1.555785

A Reference Equation of State for the Thermodynamic Properties of Ethane for Temperatures from the Melting Line to 675 K and Pressures up to 900 MPa Journal of Physical and Chemical Reference Data **35**, 205 (2006); https:// doi.org/10.1063/1.1859286

Ideal Gas Thermodynamic Properties of Ethane and Propane Journal of Physical and Chemical Reference Data 2, 427 (1973); https:// doi.org/10.1063/1.3253123



Where in the world is AIP Publishing? Find out where we are exhibiting next



## **Thermophysical Properties of Ethane**

## Daniel G. Friend, Hepburn Ingham, and James F. Ely

Thermophysics Division, National Institute of Standards and Technology, Boulder, CO 80303

#### Received November 28, 1989; revised manuscript received July 18, 1990

New correlations for the thermophysical properties of fluid ethane are presented. The correlations are based on a critical evaluation of the available experimental data and have been developed to represent these data over a broad range of the state variables. Estimates for the accuracy of the equations and comparisons with measured properties are given. The reasons for this new study of ethane include significant new and accurate data and improvements in the correlating functions which allow increased accuracy of the correlations—especially in the extended critical region. Short tables of the thermophysical properties of ethane are included. This study complements an earlier study of methane and uses the same correlating equations and format.

For the thermodynamic properties, a classical equation for the molar Helmholtz energy, which contains terms multiplied by the exponential of the quadratic and quartic powers of the system density, is used. The resulting equation of state is accurate from about 90 K to 625 K for pressures less than 70 MPa and was developed by considering PVT, second virial coefficient, heat capacity, and sound speed data. Tables of coefficients and equations are presented to allow the calculation of these and other thermodynamic quantities. Ancillary equations for properties along the liquid-vapor phase boundary, which are consistent with the equation of state and lowest order scaling theory, are also given.

For the viscosity of ethane, a contribution based on a theoretical fit of low-density data is combined with an empirical representation of the excess contribution. The approximate range of the resulting correlation is 90 K to 500 K for pressures less than 60 MPa. The correlation for the thermal conductivity includes a theoretically based expression for the critical enhancement; the range for the resulting correlation is about 90 K to 600 K for pressures below 70 MPa.

Key words: correlation; density; equation of state; ethane; heat capacity; phase boundary; pressure; speed of sound; thermal conductivity; thermophysical properties; transport properties; virial coefficients; viscosity.

## Contents

1.	Introduction	278
	1.1. Need for This Correlation	278
	1.2. Range of Correlations	279
2.	Correlating Equations	280
	2.1. Equation of State	280
	2. 2. Ideal Gas Reference State Equation	280
	2. 3. Liquid-Vapor Saturation Boundary	282
	2.4. Derived Property Equations	284
	2.5. Transport Property Correlations	286
	2.5.1. Dilute Gas Correlation	287
	2.5.2. Excess Property Correlation	290
	2.5.3. Critical Enhancement Correlation	290

Reprints available from ACS; see Reprints List at back of issue.

3.	Development of the Correlations	293
	3.1. Fundamental Constants, Fixed Points and	
	Ideal Gas Properties	293
	3.2. Ancillary Equations for the Two-Phase	
	Boundary	294
	3.3. Residual Helmholtz Energy	295
	3.4. Transport Property Correlations	298
	3.4.1. Viscosity	298
	3.4.2. Thermal Conductivity	299
4.	Comparisons of Derived and Experimental	
	Properties	299
	4.1. Two-Phase Boundary	300
	4.2. Ideal Gas Properties	304
	4.3. Thermodynamic Properties from the	
	SWEOS	306
	4.3.1. Second Viral Coefficient Data	306
	4.3.2. <i>PVT</i> Data	307
	4.3.3. Heat Capacities	314
	4.3.4. Sound Speed	320

<sup>©1991</sup> by the U.S. Secretary of Commerce on behalf of the United States. This copyright is assigned to the American Institute of Physics and the American Chemical Society.

	4.4. Transport Property Comparisons	324
	4.4.1. Viscosity	324
	4.4.2. Thermal Conductivity	328
5.	Conclusions	333
6.	Acknowledgments	333
7.	References	333
8.	Appendix	336

## List of Tables

1.	Fixed point constants and other parameters used in the correlations	280
2.	Exponents and coefficients for the residual	••••
3.	Coefficients for ideal gas Helmholtz energy.	288
	Eq. (3)	288
4.	Coefficients for liquid-vapor boundary correla-	200
~	tions	289
э. С	Ideal gas Helmholtz energy and its derivatives	-
6.	Residual Helmholtz energy and its derivatives.	289
7.	Thermodynamic property equations	291
8.	Coefficients for dilute gas transport properties.	291
9.	Coefficients for excess transport properties	292
10.	Constants for $\lambda_{cr}$ , Eq. (18) [using Eqs. (19) and	
	(20)]	292
11.	Statistics for thermodynamic property data ver-	
	sus SWEOS correlation	302
12.	Sources of <i>PVT</i> data	307
13.	Sources of heat capacity data	319
14.	Sources of sound speed data	322
15.	Sources of viscosity data at elevated pressures.	325
16.	Sources of thermal conductivity data at ele-	
	vated pressures	330
A1.	Properties of ideal gas at 0. 1 MPa and dilute	
	gas transport properties	336
A2.	Properties along saturation boundary	337
A3.	Properties of ethane in the single-phase	
	region	339
	6	

## List of Figures

Pressure-density plot with 2-phase dome and labelled isotherms	281
Pressure-temperature plot with saturation line	
and labelled isochores	282
Pressure-temperature data map for equation of	
state	283
Viscosity versus temperature and density	284
Pressure-temperature data map for primary	
viscosity data	285
Thermal conductivity versus temperature and	
density	286
Pressure-temperature data map for primary	
thermal conductivity data	287
	Pressure-density plot with 2-phase dome and labelled isotherms Pressure-temperature plot with saturation line and labelled isochores Pressure-temperature data map for equation of state Viscosity versus temperature and density Pressure-temperature data map for primary viscosity data Thermal conductivity versus temperature and density Pressure-temperature data map for primary thermal conductivity data

4.	Deviations of calculated saturation pressures	
	versus temperature	301
5.	Deviations of calculated saturated liquid densi-	
	ties versus temperature	303
6.	Deviations of calculated saturated vapor densi-	
	ties versus temperature	304
7.	Deviations of calculated ideal gas properties	
	versus temperature	305
8.	Deviations of calculated second virial coeffi-	
	cients versus temperature	306
<b>9</b> a	Deviations of calculated pressures versus pres-	200
<i>J</i> <b>u</b> .	sure $90-200$ K - primary data	300
9h	Deviations of calculated densities versus den-	507
<i>J</i> 0.	sity $00-200$ K - primary data	310
100	Deviations of colculated prossures vorsus pros	510
10a	cure 200 280 K primary data	311
105	Sure, $200-200 \text{ K} = \text{printary data}$	511
100	Deviations of calculated densities versus den-	212
11	sity, $200-280$ K – primary data	312
11a	Deviations of calculated pressures versus pres-	212
	sure, $280-315$ K – primary data	313
115	. Deviations of calculated densities versus den-	• • •
	sity, 280–315 K – primary data	314
12a	Deviations of calculated pressures versus pres-	
	sure, $315-625 \text{ K}$ – primary data	314
12b	Deviations of calculated densities versus den-	
	sity, 315–625 K – primary data	316
13.	Deviations of calculated pressures versus pres-	
	sure, secondary data	317
14.	Deviations of calculated densities versus den-	
	sity, secondary data	318
15.	Deviations of calculated molar isochoric heat	
	capacities versus density	319
16.	Deviations of calculated molar isobaric heat ca-	
	pacities versus pressure	320
17.	Deviations of calculated molar heat capacity of	
	the saturated liquid versus temperature	321
18	Deviations of calculated speeds of sound in sin-	
10.	ale-nhase region versus density	377
10	Deviations of calculated speeds of sound in sat-	
17.	urated liquid versus temperature	323
20	Deviations of calculated dilute gas viscosities	545
20.	Deviations of calculated diffute gas viscosities	225
	Versus temperature	323
21a	Deviations of calculated viscosities versus defi-	224
011	sity	320
216	Deviations of calculated viscosities versus den-	
	sity-data from [24,96]	327
22.	Deviations of calculated dilute gas thermal	
	conductivities versus temperature	329
23.	Critical enhancement of the thermal conductiv-	
	ity versus density	330
24a.	Deviations of calculated thermal conductivities	
	versus density-primary data	331
24b	Deviations of calculated thermal conductivities	
	versus density-secondary data	332

## THERMOPHYSICAL PROPERTIES OF ETHANE

# List of Symbols and Units

Symbol	Description	SI Units (used in text)	Reference
A	Molar Helmholtz energy	J·mol <sup>-1</sup> a	Ea. (1)
AAD	Average absolute deviation	_	Sec. 4
В	Second virial coefficient	dm³⋅mol <sup>-1</sup>	Table(7)
BIAS	Average deviation	_	Sec. 4
BWR	Benedict-Webb-Rubin EOS	-	Ref. [10]
С	Expansion coefficient in $\rho_{\sigma\nu}$	-	Eq. (6b)
Ci	Coefficients in $\Omega^{(2,2)*}$	-	Eq. (12), Table(8)
$C_P$	Molar isobaric heat capacity	J·mol <sup>-1</sup> ·K <sup>-1</sup>	Table(7)
$C_{\nu}$	Molar isochoric heat capacity	J·mol <sup>-1</sup> ·K <sup>-1</sup>	Table(7)
EOS	Equation of state		Sec. 1
F	Crossover function in $\lambda_{cr}$	-	Eqs. (18),(20)
$f_{\rm int}, f_{\rm i}$	Contribution from internal modes		Eqs. (13),(14), Table(8)
G	Molar Gibbs energy	J·mol <sup>-1</sup>	Table(7)
$G_i$	Coefficients in $\rho_{\sigma}$		Eq. $(5)$ , Table $(4)$
<b>g</b> i	Coefficients in $\eta_{ex}$	 I	Eq. $(15)$ , Table(9)
H	Molar enthalpy	J-mol	Tables(1),(/)
$H_i$	Coefficients in $P_{\sigma}$	-	Eq. $(4)$ , Table $(4)$
$J_i$	Coefficients in $\rho_{\sigma V}$		Eq. $(0)$ , Table $(4)$
Ji Ir	Roltzmann constant	-	Eq. $(17)$ , Table(9)
к M	Bolizinanii Constant Relative molecular mass	J'K	Table(1)
Nr N.	Avogadro constant	mol <sup>-1</sup>	Table(1)
n:	Coefficients in $\phi'$	-	$F_{a}(2)$ Table(2)
P	Pressure	MPa	-
P.*	Reduced saturation pressure, $P_{\sigma}/P_{c}$		Eq. (6a)
O <sub>i</sub>	Coefficients in $\phi^{id}$	-	Eq. $(3)$ , Table $(3)$
$\tilde{q}_{\rm D}$	Wavenumber cutoff	nm <sup>-1</sup>	Eq. (20). Table(10)
$\hat{R}$	Gas constant	J·mol <sup>-1</sup> ·K <sup>-1</sup>	Table(1)
RMS	Root mean square deviation		Sec. 4
r	Intermolecular separation	nm	Eq. (11)
r <sub>i</sub>	Exponent of δ	-	Eqs. (2),(15),(17),
			Tables(2),(9)
S	Molar entropy	$J \cdot mol^{-1} \cdot K^{-1}$	Eq. (24), Table(7)
s <sub>i</sub>	Exponent of $\tau$	_	Eqs. (2),(15),(17), Tables(2),(9)
t	Reduced temperature, $kT/\epsilon$		Eqs. (10),(12)
$T_{m+}$	Temperature, IPTS-68	K	_
<i>T</i> *	Reduced temperature, $(T_c - T)/T_c$	—	Eq. (4–6)
U	Molar internal energy	J·mol <sup>-1</sup>	Table(7)
u V	Unified atomic mass unit	 T	Table(1)
V	Intermolecular potential	J	Eq. $(11)$
W Z	Compressibility factor $B/BT_{-}$	In-S	Table(7)
L	Compressionity factor, $F/KT \rho$	_	Eq. (6)
Greek			
α	Scaling exponent	-	Ref. [15]
β	Scaling exponent in $\rho_{\sigma L}$ , $\rho_{\sigma V}$	-	Eqs. (5),(6), Table(4)
$\Gamma_0$	Critical amplitude	-	Eq. (19), Table(10)
γ	Potential parameter	-	Ref. [13]
Ŷ	Scaling exponent	-	Eq. (19), Table(10)
ŏ	Reduced density, $\rho/\rho_c$		

Symbol	Description	SI Units (used in text)	Reference
E	Scaling exponent in $P_{\sigma}$ , $2-\alpha$	_	Eq. (4),Table(4)
ε	Small temperature difference	K	Eq. (24)
E	Energy parameter in $V(r)$	J	See $\epsilon/k$
€/k	Energy parameter in $V(r)$	Κ	Eq. (11), Table(1)
η	Shear viscosity	µPa∙s	Eq. (8)
Λ	Coupling constant in $\lambda_{cr}$	_	Eq. (18), Table(10)
λ	Thermal conductivity	mW·m <sup>−1</sup> ·K <sup>−1</sup>	Eq. (9)
ν	Scaling exponent		Eq. (19), Table(10)
ξ	Correlation length	nm	Eqs. (18-20)
ξo	Correlation length amplitude	nm	Eq. (20), Table(10)
ρ	Molar density	mol·dm <sup>-3</sup>	_
σ	Distance parameter in $V(r)$	nm	Eqs. (10),(11), Table(1)
τ	Reduced inverse temperature, $T_{c}/T$	-	_
ф	Reduced Helmholtz energy, A/RT	-	Eq. (1)
$\Omega^{(2,2)*}$	Reduced collision integral	-	Eqs. (10),(12)
Superscripts			
id	Ideal gas contribution		Eq. (1), Table(1)
r	Residual contribution		Eq. (1)
Subscripts			
c	Value at critical point		Table(1)
cr	Critical contribution		Eqs. (9),(18)
ex	Excess contribution		Eqs. (8),(9),(15),(17)
exp	Value from experiment		Eq. (26)
t	Value at triple point		Table(1)
tL,tV	Value at triple point in liquid, vapor		Table(1)
σ	Value at saturation boundary		Eq. (4)
σL,σV	Value in saturated liquid, vapor		Eqs. (5),(6),(25)
δ	Partial derivative with respect to $\delta$		Tables(5),(6)
τ	Partial derivative with respect to $\tau$		Tables $(5), (6)$
0	Value at zero density		Eqs. (8)-(10),(13)

<sup>a</sup>Throughout this paper, extensive physical quantities are given on a molar basis. The elementary entities are the ethane (C<sub>2</sub>H<sub>6</sub>) molecules.

## 1. Introduction

## 1.1. Need for This Correlation

Because ethane is both an industrially important fluid and the second member of the vitally interesting alkane series, we have felt it necessary to examine the newly available thermophysical property data and to re-evaluate the older data to produce more useful and accurate correlations. We have studied the primary homologue, methane, in previous publications;<sup>1,2</sup> this work reports a parallel study for ethane, and the form of the correlations and presentation are essentially identical.

In this paper we present an empirical equation of state for ethane based on extensive multiproperty analysis, as well as correlations for ideal gas properties, the liquidvapor phase boundary, and for the viscosity and thermal conductivity of ethane. Tables of coefficients for these correlating equations and graphical representations of the functions for easy accessibility of estimated values of certain properties are included. Discussions of the accuracy of these correlations and their applicable ranges, and explicit comparisons with experimental data are also given. In an Appendix, we have compiled very brief tables of thermophysical properties in the ideal or dilute gas limit, along the liquid-vapor phase boundary, and in the one-phase region. Extensive tables of properties and comparisons with experimental data will be published separately.<sup>3,3a</sup>

Although there have been some experimental studies of ethane subsequent to the 1976 publication of the technical note by Goodwin *et al.*<sup>4</sup> and the more recent correlation of Younglove and Ely,<sup>10</sup> the primary reason for these correlations is to provide thermodynamic and transport property surfaces which are completely compatible with those developed for the methane fluid.<sup>1,2</sup> This compatibility will allow development and testing of mixture theories based on corresponding states models. The recently translated monograph by Sychev *et al.*<sup>5</sup> provides an extensive discussion of the available thermodynamic data through 1980 as well as an alternative wide-range correlation.

The present thermodynamic surface, based on the Schmidt-Wagner equation of state (SWEOS),<sup>6</sup> exhibits a greater conformity to known scaling behavior in the general region of the critical point than many established classical equations of state. Thus, while retaining the simplicity of classical algebraic representations, our correlation allows more accuracy in calculating thermodynamic properties around the critical point. We have not reformulated the equation of state based on implementation of an algorithm such as the evolutionary optimization method (EOM) or its variants.<sup>6a</sup> Our optimization is based on linear least-squares determination of the coefficients of the SWEOS with emphasis placed on data evaluation, selection, and weighting; non-linear routines are used in our treatment of the data. For modelling and theoretical work on mixture thermodynamics, it is useful to use identical formulations of the pure fluid equations of state. This is an important reason for retaining the SWEOS for our work on ethane. Other choices for the equation of state, such as those generated by an EOM or those with additional coefficients or other degrees of freedom, and different data selection would generate different statistical agreement between experimental data and the correlation; our SWEOS provides an optimum description of the thermodynamic surface within the constraints which we have outlined and with the estimated uncertainties as described in Sec. 4.

Among the extensive multiproperty data used in the algorithms for computing the coefficients in the SWEOS, we have incorporated both PVT and  $C_V$  points generated from a scaled EOS. As for any classical equation of state, the asymptotically critical region is not strictly correct.

For the transport properties in the dilute gas, we use the Chapman-Enskog theory<sup>7</sup> directly for viscosity and with an improved treatment of internal degrees of freedom for the thermal conductivity. The necessary collision integral, based on the 11–6–8,  $\gamma = 3$  interparticle potential and tabulated in Ref. 8, was represented by a simple function. The excess functions are represented by polynomial or rational polynomial approximations and have been scaled by corresponding states arguments. Finally, the critical enhancement term, vitally important for the thermal conductivity correlation over a broad region of the phase diagram, is based on a very recent mode-coupling theory of Olchowy and Sengers.<sup>9</sup> These transport property correlations differ from and supercede those recently published by Younglove and Ely.<sup>10</sup>

#### 1.2. Range of Correlations

For the equation of state correlation, we have examined *PVT* data in the range 90 K < T < 623 K, 0.1 MPa < P < 69 MPa, and 0.03 mol·dm<sup>-3</sup>  $< \rho < 22$  mol·dm<sup>-3</sup>.

as well as virial coefficient, heat capacity, and sound speed data. For the broad range of the phase diagram with pressures less than 70 MPa, we estimate the accuracy of our correlation to be about 0.2% (standard deviation) when evaluating the density given the temperature and pressure, and 1% when evaluating the pressure given the temperature and density. In the neighborhood of the critical point, the accuracy deteriorates to 0.5% for density calculations. Detailed comparisons with experimental data and estimates of the quality of the correlation in different regions of the phase diagram are given in Sec. 4. For the derived properties, the accuracy of the correlation is somewhat less, with details given in Sec. 4.3. In Figs. 1a and 1b, we give representations of the phase diagram for ethane, together with isochores and isotherms, from which the interested reader can determine a rough approximation to the scope of the PVT relation presented here. In Fig. 1c, we present a map of the primary data used to establish the SWEOS. Within each category of data shown in Fig. 1c, there may be several primary references. Detailed discussion of the data selection is given in Sec. 3.3.

To produce an accurate correlation for the equation of state and derived properties of a fluid, especially one that is to be useful for corresponding states calculations, it is necessary to have accurate values for the critical and triple-point parameters and a good correlation or an accurate and dense set of data for the two-phase boundaries and ideal gas properties. The selected values of the critical and triple point parameters, with uncertainties, are given in Table 1. Details of the selection are given in Sec. 3.1. The equations determined for the saturation pressures and the saturated liquid and vapor densities agree with lowest order scaling theory and are given in Sec. 2.3; they are illustrated in Figs. 1a and 1b. The estimated accuracies of these correlations are generally 0.1% for the pressure, 0.3% for the liquid density, and 0.3% for the vapor density, but the accuracies deteriorate near the triple point and critical point temperatures, as discussed in Sec. 4.1. The equation for the ideal gas properties follows that given in Ref.1 and reproduces spectroscopically derived ideal gas entropies to within 0.1% and isobaric heat capacities to within 0.2% in the temperature range 90-700 K.

The form of the SWEOS, together with values of certain reference point parameters (also given in Table 1) and the ideal gas properties, allows easy determination of many of the derived thermodynamic properties. In Table 7 below we have collected the explicit algebraic forms, in a manner quite similar to the original tables of Schmidt and Wagner,<sup>6</sup> which will allow one to evaluate several of these quantities. Comparisons with some experimental data and estimates of the accuracy of the derived property correlations are given in Sec. 4.3. The accuracy of the correlation decreases as the order of the derivatives increases and again, the accuracy generally is lower in the critical region than in other regions of the surface. One may use the SWEOS to determine any other thermodynamic properties, throughout the range of the

correlations, by using the usual relationships of thermodynamics; care must be taken in the critical region and when using higher order derivatives.

The available experimental data for the transport properties are not nearly as extensive as those for equilibrium properties. The temperature range for the viscosity is 95 to 500 K; the pressure range is 0.1 to 69 MPa; and the density range is 0.04 to 22 mol·dm<sup>-3</sup>. In this region of the phase diagram, the extrema of the viscosity are 9 and 1100 µPa·s. The viscosity correlation given in Sec. 2.5 has an associated uncertainty of about 2%. Figure 2a depicts the relationships among viscosity, temperature, and density, and again enables the reader to obtain a crude value for this property without evaluation of the algebraic expression given in Sec. 2. Figure 2b illustrates the pressure-temperature state points for the primary data used to determine the viscosity correlation. A comprehensive discussion of these primary data is given in Sec. 3.4.1.

The useful experimental data for thermal conductivity are between 112 and 600 K, 0.1 and 100 MPa, and 0.03 and 22 mol·dm<sup>-3</sup>, with resulting range in  $\lambda$  of 13 to 270  $mW \cdot m^{-1} \cdot K^{-1}$ . The thermal conductivity correlation has an uncertainty of about 2% and is shown in Fig. 3a. The primary data are shown in a pressure-temperature map in Fig. 3b. These data are discussed in Sec. 3.4.2.

TABLE 1. Fixed point constants and other parameters used in the correlations

Triple point: <sup>a</sup>	$T_{t} = 90.352 \pm 0.005 \text{ K}^{b}$ $P_{t} = 1.130 \pm 0.005 \text{ Pa}$ $\rho_{t} = 21.667 \pm 0.01 \text{ mol·dm}^{-3}$ $\rho_{tv} = 1.504 \pm 0.02 \text{ mol·dam}^{-3}$	Ref. [19] Eq. (4) Eq. (5) Eq. (6)
Critical point:	$T_{c} = 305.33 \pm 0.04 \text{ K}^{c}$ $P_{c} = 4.8718 \pm 0.005 \text{ MPa}$ $\rho_{c} = 6.87 \pm 0.1 \text{ mol·dm}^{-3}$ $Z_{c} = 0.279 34 \pm 0.005$	Ref. [22] Ref. [22] Ref. [22] Calculated
Intermolecular potential $\epsilon/k =$ parameters:	245.0 K $\sigma = 0.436 \ 82 \ nm$	Sec. 3.4.1
Ideal Gas reference point values:	(at 298.15 K and 0.101 325 MPa) $S^{id} = 229.12 J \cdot K^{-1} \cdot mol^{-1}$ $H^{id} = 11.874 kJ \cdot mol^{-1}$	Ref. [11] Ref. [11]
Miscellaneous: Relative molecular mass	$M_{\rm r} = 30.070$	Ref. [17]
Universal gas constant	$R = 8.314 \ 510 \ \mathrm{J} \cdot \mathrm{mol}^{-1} \cdot \mathrm{K}^{-1}$	Ref. [16]
Boltzmann constant	$k = 1.380 \ 658 \times 10^{-23} \ \mathrm{J}\cdot\mathrm{K}^{-1}$	Ref. [16]
Avogadro constant	$N_{\rm A} = 6.022 \ 136 \ 7 \ \times \ 10^{23} \ {\rm mol}^{-1}$	Ref. [16]
Unified atomic mass unit	$u = 1.660 \ 540 \ 2 \ \times \ 10^{-27} \ kg$	Ref. [16]

\*Uncertainties presented in this table are discussed in Sec. 3 and do not always agree with those given in the source references.

<sup>b</sup>This corresponds to 90.360 K on ITS-90 scale.

"This corresponds to 305.32 K on ITS-90 scale.

## our correlations of the thermodynamic surface and transport properties. The data selection for establishing the coefficients and parameters is discussed in Sec. 3, and explicit and extensive comparisons between the correlations and experimental data are given in Sec. 4.

2. Correlating Equations

In this section, we describe all the equations used in

## 2.1. Equation of State

As indicated above, we have used the form of the residual equation of state presented by Schmidt and Wagner in Ref. 6. This SWEOS was introduced as an improvement over the Benedict-Webb-Rubin (BWR)<sup>10</sup> equation, especially to represent data in the extended critical region. The general form includes terms multiplied by  $\exp[-(\rho/\rho_c)^4]$  in addition to terms with  $\exp[-(\rho/\rho_c)^2]$  as occur in the BWR equation. The critical density  $\rho_c$  and critical temperature  $T_c$  are used as reduction parameters; the specific polynomial in reduced temperature and density (and the exponential terms) was determined by Schmidt and Wagner.<sup>6</sup> They used a selection algorithm with a set of 336 terms which were optimized to best fit the wide ranging data for oxygen. We have retained this choice of polynomial, because it seems also to improve the fit for methane in Ref. 1 and for ethane.

As in Ref. 6, we consider the molar Helmholtz energy A as the sum of ideal and residual terms and write

$$A(\rho,T) = A^{id} + A^r = RT\phi = RT(\phi^{id} + \phi^r).$$
(1)

With the definitions  $\delta = \rho/\rho_c$  and  $\tau = T_c/T$ , the dimensionless residual term becomes

$$\phi^{r} = \sum_{i=1}^{13} n_{i} \, \delta^{r_{i}} \, \tau^{s_{i}} \, + \, e^{-\delta^{2}} \, \sum_{i=14}^{24} n_{i} \, \delta^{r_{i}} \, \tau^{s_{i}} \, + \, e^{-\delta^{4}} \, \sum_{i=25}^{32} n_{i} \, \delta^{r_{i}} \, \tau^{s_{i}}.$$
(2)

The coefficients  $n_i$ , determined by fitting the data, and the specific exponents  $r_i$  and  $s_i$  are given in Table 2. Equation (2) and Table 2 are essentially the same as Eq. (11) in the paper by Schmidt and Wagner<sup>6</sup> and the corresponding Eq. (2) and Table 2 of Ref. 1.

#### 2.2. Ideal Gas Reference State Equation

Thermodynamic functions may be derived for the ideal gas by using statistical mechanical models with spectroscopic data for ethane. Because these functions are usually more accurate than the corresponding values obtained by direct evaluation using thermodynamic experiments, at least for simple molecules, it is customary to consider the results obtained from the spectroscopic data at the standard pressure of 0.101 325 MPa (1 atm) as a reference state with which the wide ranging correlations must agree. We have fitted the spectroscopically derived data of Chao et al.,11 and the corresponding values of the ideal gas entropy and enthalpy at standard conditions (298.15 K and 0.101 325 MPa) are given in Table 1. These values exclude any contributions from nuclear spin, so all



FIG. 1a. Ethane fluid isotherms from the SWEOS. Solid circle denotes the critical point, and two-phase boundary is from ancillary equations.

values for the entropy and for the thermodynamic potentials (internal energy, Helmholtz energy, Gibbs energy, and enthalpy) are relative to values assigned to two integration constants. The resulting ideal gas properties essentially agree with those adopted by Goodwin *et al.*<sup>4</sup> and by Sychev *et al.*<sup>5</sup> who used the same spectroscopic model and data.

We use an equation for the ideal gas Helmholtz energy which is identical to that presented in the methane manuscript<sup>1</sup> and which reflects the correlating equation developed by Goodwin for the methane fluid.<sup>12</sup> Thus we write, in dimensionless form,

$$\begin{split} \phi^{\rm id}(\delta,\tau) &= \mathcal{A}^{\rm id} / RT = Q_1 + \ln \delta + Q_2 \ln \tau + Q_3 \tau^{-1/3} \\ &+ Q_4 \tau^{-2/3} + Q_5 \tau^{-1} + Q_6 \ln (1 - e^{Q/\tau}) \end{split}$$
(3)

with the coefficients  $Q_i$  given in Table 3. Equation (3) must be evaluated at the experimental density and temperature, although for many thermodynamic properties there is no density (or pressure) dependence in the required derivatives. At 0.101 325 MPa (1 atm), the density of the ideal gas can be evaluated using the ideal gas equation of state and Eq. (3) becomes

$$\phi^{id}(\tau) = Q_0 + (Q_2 + 1) \ln \tau + Q_3 \tau^{-1/3} + Q_4 \tau^{-2/3} + Q_5 \tau^{-1} + Q_6 \ln (1 - e^{Q_7 \tau}).$$
(3a)



FIG. 1b. Ethane fluid isochores from the SWEOS. Solid circle denotes the critical point, and saturation boundary is from ancillary equation.

In Table 3, values of both  $Q_0$  and  $Q_1$  are given, although in practice  $Q_0$  is not needed for the evaluation of any property.

#### 2.3. Liquid-Vapor Saturation Boundary

Although the SWEOS allows calculation of saturation properties by a Maxwell construction technique within the two-phase region, it is useful to have separate correlations of the two-phase boundary. For this reason, we present new correlating equations for the saturation pressure  $P_{\sigma}$ , the density of the saturated vapor  $\rho_{\sigma V}$ , and the density of the saturated liquid  $\rho_{\sigma L}$ , all as functions of temperature. In fact, these saturation boundary correlations were also used as input in the development of the equation of state.

The three equations are identical in form to those presented in Ref. 1 for methane; the reference also provides motivation for these forms. For the density of saturated liquid methane, however, the coefficient  $G_5$  was 0; thus there is an extra term in the numerator of the ethane equation which was not indicated in Ref. 1. This is not suprising since the saturation boundary, extending from the triple point to the critical point, covers a much broader temperature range for ethane than for methane. We have written the vapor pressure equation in terms of  $T^* = (T_c - T)/T_c$  (which is positive along the saturation boundary) so that the critical behavior may be easily dis-



FIG. 1c. Primary data used to determine coefficients in the SWEOS. Several references may correspond to each type of data.

cerned by expansion of the exponential about  $T^* = 0$ . Thus we write

$$P_{\sigma}(T) = P_{c} \exp \left[H_{1}T^{*}/(1-T^{*}) + H_{2}T^{*} + H_{3}T^{*\epsilon} + H_{4}T^{*2} + H_{5}T^{*3}\right], \qquad (4)$$

where the dimensionless fitted coefficients  $H_i$  and the exponent  $\epsilon$  are given in Table 4. In Eq. (4), the saturation pressure  $P_{\sigma}$  precisely equals the critical pressure at the critical temperature, and the exponent  $\epsilon$  was forced to have its effective scaling theory value of 1.90 as in the methane manuscript.

For the density of the saturated liquid, we again have chosen an equation in  $T^*$  which reduces to the known (lowest order) scaling result in the limit  $T^* \rightarrow 0$ . The equation, which has the added flexibility available with a ratio of terms in  $T^*$ , is

$$\rho_{\sigma L}(T) = \rho_{c} \bigg[ 1 + \frac{G_{1}T^{*\beta} + G_{2}T^{*2} + G_{3}T^{*3} + G_{5}T^{*4}}{1 + G_{4}T^{*(1-\beta)}} , \quad (5)$$

where the coefficients  $G_i$  and exponent  $\beta$  appear in

Table 4. As in Ref. 1, the critical exponent  $\beta$  was assumed to have the effective value of 0.355.

For the density of the saturated vapor, it is desirable to use an equation which has the theoretically predicted behavior in both the low density (ideal gas) limit and in the neighborhood of the critical point. While these limits may not be simultaneously discernable at a glance, the equation

$$\rho_{\sigma V}(T) = \frac{P_{\sigma}(T)}{RT} \left\{ 1 + P_{\sigma}(T)\tau^{8} \frac{Z_{c} - 1}{P_{c}} \right\}$$

$$\left[ 1 + \frac{J_{0}T^{*\beta} + J_{1}T^{*2\beta} + J_{2}(T^{*} + T^{*4}) + J_{3}T^{*2}}{1 + J_{4}T^{*}} \right]^{-1}, (6)$$

with  $Z_c = P_c/(RT_c\rho_c)$ , the critical compressibility factor, has the ideal gas behavior at the lowest pressures and can be rewritten as

$$\rho_{\sigma V}(T) = \rho_{c} (1 - T^{*})^{7} \left\{ 1 - \frac{1}{Z_{c}} \left[ 1 - \frac{(1 - T^{*})^{8}}{P_{\sigma}^{*}} \right] + (1 - Z_{c}^{-1}) \frac{J_{0}T^{*\beta} + J_{1}T^{*2\beta} + J_{2}(T^{*} + T^{*4}) + J_{3}T^{*2}}{1 + J_{4}T^{*}} \right\}.$$
(6a)



FIG. 2a. Three-dimensional representation for the viscosity coefficient surface. The viscosity is in  $\mu$ Pa·s; the density is in mol·dm<sup>-3</sup>.

This last equation, with  $P_{\sigma}^* = P_{\sigma}(T)/P_c$ , reduces to the lowest order scaling result in the limit  $T^* \rightarrow 0$ :

$$\rho_{\sigma V} \propto \rho_c \ (1 - C \ T^{*\beta} + ...).$$
(6b)

The value of  $\beta$  for the vapor density correlations remains at the universal value of 0.355, and the amplitude  $C [= J_0(1-Z_c^{-1})]$  is forced to have the same value as  $G_1$ in the correlation for the saturated liquid density of Eq. (5). Because of the presence of the exponent  $2\beta$  in Eq. (6), higher order terms in the critical region expansion, Eq. (7), are not directly comparable. The equality of C and  $G_1$  reproduces the observed and theoretically predicted symmetry of the two-phase boundary around the critical point. Despite the presence of the  $T^{*2\beta}$  term in Eq. (6), the rectilinear diameter [defined as  $\frac{1}{2}(\rho_{oL} + \rho_{oV})$ ] determined by Eqs. (5) and (6) is very nearly linear in the critical region, as required by scaling theory. The coefficients for Eqs. (6) and (6a) are given in Table 4; Eqs. (6) and (6a) are completely equivalent. The primary data used in the development of these ancillay equations are discussed in Sec. 3.2; explicit comparisons with both primary and other data as well as estimates of the reliability for all three two-phase boundary correlations are given in Sec. 4.1.

## 2.4. Derived Property Equations

One can evaluate the thermodynamic properties of ethane by taking the appropriate derivatives of the Helmholtz energy as given in Eqs. (1), (2), and (3). In Tables 5 and 6, identical to tables in Ref. 1, we have col-



FIG. 2b. Pressure and temperature state points for primary data used to determine viscosity correlation.

lected the coefficients necessary to calculate the six lowest order derivatives for the ideal gas and residual contributions to the Helmholtz energy. These are the first two density derivatives along isotherms, the temperature derivatives along isochores, and the mixed derivatives. As in the paper by Schmidt and Wagner<sup>6</sup> and the methane manuscript,<sup>1</sup> we consider the reduced density and the inverse reduced temperature as independent variables, and we denote the derivatives as subscripts to the appropriate Helmholtz energy quantity. For example,

$$\phi_{\delta}' = \frac{\partial \phi'(\delta,\tau)}{\partial \delta} \bigg|_{\tau} = \frac{\rho_c}{RT} \left. \frac{\partial A'(\rho,T)}{\partial \rho} \right|_{T}$$
(7a)

is the isothermal (reduced) density derivative of the (reduced) residual Helmholtz energy. Second derivatives are analogously defined, but have two subscripts to indicate the parameters with respect to which the derivatives are taken.

To calculate derivatives of the ideal gas contribution to the Helmholtz energy, one is guided by the form of Eq. (3). The left-most column of Table 5 lists the terms necessary for the various derivatives. The remaining columns give the coefficients of these terms directly beneath the heading which indicates the quantity to be calculated. The resulting terms are to be added as in Eq. (3). For example, to calculate  $\tau \phi_{\tau}^{id}$  use the coefficients from the fourth column and the terms in the first column to write

$$\tau \phi_{\tau}^{id}(\delta,\tau) = Q_2 - \frac{Q_3}{3} \tau^{-1/3} - \frac{2Q_4}{3} \tau^{-2/3} - Q_5 \tau^{-1} - Q_6 Q_7 \tau (e^{-Q\tau} - 1)^{-1}.$$
(7b)

The values of the  $Q_i$  coefficients are given in Table 3. As indicated in Table 5, the density derivatives of the ideal gas Helmholtz energy are particularly simple:

$$\delta \phi_{\delta}^{id} = 1, \ \delta^2 \phi^{id}_{\delta \delta} = -1, \ \text{and} \ \delta \tau \phi^{id}_{\delta \tau} = 0.$$

For derivatives of the residual Helmholtz energy, Eq. (2) and Tables 2 and 6 can be used. As in Eq. (2), the derivatives are obtained by summing 32 terms of three general types. Each of the terms has factors consisting of powers of the reduced density and temperature with the explicit exponents  $r_i$  and  $s_i$  and coefficients  $n_i$  given for each value of i in Table 2. The additional exponential



FIG. 3a. Three-dimensional representation for the thermal conductivity coefficient surface. The thermal conductivity is in mW·m<sup>-1</sup>·K<sup>-1</sup> and the density is as in Fig. 2a. Note the divergence near the critical point.

factor, with its argument either the second or fourth power of the density, is indicated in the column heading of Table 6 where appropriate. The remaining coefficients relevant to the derivative being calculated are given in the appropriate row in that table.

In Table 7, the most common thermodynamic quantities of interest have been expressed in terms of the reduced derivatives of the molar Helmholtz energy. This table is similar to the first part of the Appendix in the paper by Schmidt and Wagner<sup>6</sup> and is identical to the table in Ref. 1. All nominally extensive quantities (that is, the various thermodynamic potentials and heat capacities) are given per mole. The density derivatives of the ideal gas contribution to the Helmholtz energy have been explicitly evaluated and included in the table where ap-

J. Phys. Chem. Ref. Data, Vol. 20, No. 2, 1991

propriate. It is straightforward to calculate, by using Tables 1-7, the most useful thermodynamic properties of ethane.

## 2.5. Transport Property Correlations

For both the viscosity and thermal conductivity, the present correlations reduce to the theoretically rigorous Chapman-Enskog theory<sup>7</sup> at the lowest densities. Added to these zero-density terms are functions which have been empirically determined to represent the excess portion of the transport coefficient. Because the critical enhancement of the viscosity is typically observed in only a small region around the citical point, and in the interest of simplicity of the correlating equations, we have chosen to



FIG. 3b. Pressure and temperature state points for primary data used to determine thermal conductivity correlation.

omit this term. For the ethane thermal conductivity, which clearly displays a critical enhancement in the experimental results, we also add a term which estimates the critical thermal conductivity enhancement. The additional term is based on the recent theoretical crossover description of the critical enhancement given by Olchowy and Sengers;<sup>9</sup> this differs from our treatment of methane in Ref. 1. The viscosity is given by

$$\eta(\rho,T) = \eta_0(T) + \eta_{ex}(\rho,T), \qquad (8)$$

and the expression for thermal conductivity is

$$\lambda(\rho,T) = \lambda_0(T) + \lambda_{ex}(\rho,T) + \lambda_{cr}(\rho,T).$$
(9)

These terms are explicitly described in this section.

#### 2.5.1. Dilute Gas Correlation

The Chapman-Enskog theory for the dilute gas viscosity gives, to lowest order in the Sonine polynomial expansion,

$$\eta_0(T) = \frac{5\sqrt{\pi \ u \ M_r \ k \ T}}{16 \ \pi \ \sigma^2 \ \Omega^{(2,2)*}(t)} \tag{10}$$

= 
$$12.0085 \sqrt{t} / \Omega^{(2,2)*}(t) \mu Pa \cdot s.$$
 (10a)

Equation (10) is identical to Eq. (8.2–10) in Ref. 7, whereas in Eq. (10a) the constants have been evaluated for ethane. The reduced collision integral  $\Omega^{(2,2)*}$  is a function only of the reduced temperature  $t = kT/\epsilon$  and the intermolecular potential function. For evaluation of Eq. (10), as for the methane fluid in Ref. 1, we have used the 11–6–8,  $\gamma = 3$  potential function,

$$V(r) = \epsilon \left[ \frac{12}{5} (r_m/r)^{11} - \frac{2}{5} (r_m/r)^6 - 3 (r_m/r)^8 \right]$$
(11)

where  $r_m = 1.1145\sigma$ . The parameters  $\epsilon$  and  $\sigma$ , which are defined by  $V(r_m) = -\epsilon$  and  $V(\sigma) = 0$ , were chosen to represent the low density transport data and are given in Table 1. The additional constants of Eq. (10), the relative molecular mass  $M_r$ , the unified atomic mass unit u, and the Boltzmann constant k, are also given in Table 1. A discussion of the utility of the 11–6–8,  $\gamma = 3$  potential in

	i	r <sub>i</sub>	Si			n <sub>i</sub>			
· · · · · · · · · · · · · · · · · · ·	1	1	0	0.462	154	305	60		· · · · · · · · · · · · · · · · · · ·
	2	1	1.5	- 0.192	369	363	87	х	10
	3	1	2.5	0.398	786	040	03		
	4	2	-0.5	0.160	545	323	72	×	10 <sup>-1</sup>
	5	2	1.5	0.128	952	422	19		
	6	2	2	0.354	583	204	91	x	10 <sup>-1</sup>
$\delta^{r_i} \tau^{r_i}$	7	3	0	0.349	278	445	40	×	10-1
	8	3	1	-0.113	061	833	80	×	10-1
	9	3	2.5	- 0.398	090	327	79	×	10-1
	10	6	0	0.830	319	368	34	×	10-3
	11	7	2	0.459	215	751	83	×	10-3
	12	7	5	0.175	302	879	17	×	10-6
	13	8	2	- 0.709	195	161	26	×	10-4
	14	1	5	- 0.234	361	622	49		
	15	1	6	0.845	746	976	45	×	10-1
	16	2	3.5	0.148	610	520	10		
	17	2	5.5	-0.100	168	578	67		
	18	3	3	- 0.592	648	243	88	×	10 <sup>-1</sup>
$e^{-\delta^2} \delta^{r_i} \tau^{s_i}$	19	3	7	-0.412	635	142	17	×	10-1
	20	5	6	0.218	551	618	69	×	10-1
	21	6	8.5	- 0.745	527	209	58	×	10-4
	22	7	4	- 0.988	590	855	72	×	$10^{-2}$
	23	8	6.5	0.102	084	164	99	×	10-2
	24	10	5.5	- 0.521	896	558	47	×	10-3
	25	2	22	0.985	921	620	30	×	10-4
	26	3	11	0.468	651	408	56	×	10 <sup>-1</sup>
	27	3	18	- 0.195	580	116	46	×	10-1
	28	4	11	- 0.465	571	616	51	×	10 <sup>-1</sup>
$e^{-\delta^4} \delta^{r_i} \tau^{r_i}$	29	4	23	0.328	779	053	76	×	10 <sup>-2</sup>
	30	5	17	0.135	720	901	85		
	31	5	18	-0.108	464	714	55		
	32	5	23	- 0.675	028	369	03	×	$10^{-2}$

TABLE 2. Exponents and coefficients for the residual Helmholtz energy  $\phi'$ , Eq. (2)

TABLE 3. Coefficients for ideal gas Helmholtz energy, Eq. (3)

	5
$Q_0 = -28.394 991 \qquad Q_4 = -3.307 373$	5
$Q_1 = -23.446$ 765 $Q_5 = -0.559$ 566	78
$Q_2 = 3.815  947  6 \qquad Q_6 = 5.072  226$	7
$O_3 = 8.602 \ 129 \ 9 \qquad O_7 = -5.507 \ 487$	4

## THERMOPHYSICAL PROPERTIES OF ETHANE

	Saturated vapor pressure Eq. (4)	Saturated liquid density Eq.(5)	Saturated vapor density Eq.(6) or (6a)
£	= 1.90	β = 0.355	$\beta = 0.355$
<i>I</i> 1	= -7.955 315	$G_1 = 1.930$ 740	$J_0 = -0.748 \ 371 \ 9$
$I_2$	= 1.532 027	$G_2 = -0.653$ 985 6	$J_1 = -1.372 895$
$\overline{I_3}$	= 14.780 68	$G_3 = 0.814  136  2$	$J_2 = -1.192597$
H.	= -13.43179	$G_4 = -0.339 743 0$	$J_3 = 1.861 505$
K	= 4.704 891	$G_5 = -0.383 814 1$	$J_4 = 1.313649$

TABLE 4. Coefficients for liquid-vapor boundary correlations

TABLE 5. Ideal gas Helmholtz energy and its derivatives

	φ <sup>id</sup> Eq.(3)	$\delta \phi_{\delta}^{id}$ ( -1 )	τφ <sub>τ</sub> <sup>id</sup>	$\delta^2 \phi_{\delta\delta}^{id}$ (1 )	τ²φ <sub>ττ</sub> <sup>id</sup>	δτφ <sub>δτ</sub> <sup>id</sup> ( -0 )
1	$Q_1$	1	$Q_2$	- 1	$-Q_{2}$	0
ln δ	1	0	0	0	0	0
ln τ	$Q_2$	0	0	0	0	0
$\tau^{-1/3}$	$Q_3$	0	$-Q_3/3$	0	4Q3/9	0
$\tau^{-2/3}$	Q₄	0	$-2Q_4/3$	0	10Q4/9	0
$\tau^{-1}$	Q5	0	$-Q_{5}$	0	2Q5	0
$\ln (1 - e^{Q_{T}})$	$Q_6$	0	0	0	0	0
$(e^{-Q_7\tau}-1)^{-1}$	0	0	$-Q_6Q_{77}$	0	0	0
$e^{Q_{7}\tau} (e^{Q_{7}\tau} - 1)^{-2}$	0	0	0	0	$-Q_6Q_7^2\tau^2$	0

TABLE 6. Residual Helmholtz energy and its derivatives

	$n_i  \delta^{r_i}  \tau^{r_i}$	$e^{-\delta^2} n_i \delta^{r_i} \tau^{s_i}$	$e^{-\delta^4} n_i \delta^{r_i} \tau^{s_i}$
	(i = 1  to  13)	(i = 14  to  24)	(i = 25  to  32)
φ <sup>r</sup>	1	1	1
δφ <sub>δ</sub> r	r <sub>i</sub>	$r_i - 2\delta^2$	$r_i - 4\delta^4$
τφ <sub>τ</sub> r	Si	Si	Si
$\delta^2 \varphi^r_{\delta \delta}$	$r_i(r_i-1)$	$[r_i(r_i-1)-2(2r_i+1)\delta^2+4\delta^4]$	$[r_i(r_i-1)-4(2r_i+3)\delta^4+16\delta^8]$
$\tau^2 \phi^r_{\tau\tau}$	$s_i(s_i-1)$	$s_i(s_i-1)$	$s_i(s_i - 1)$
δτφ <sup>ι</sup> δτ	T <sub>i</sub> S <sub>i</sub>	$s_i(r_i-2\delta^2)$	$s_i(r_i-4\delta^4)$

correlating transport property data has been given by Hanley and Klein.<sup>13</sup> This potential, with the present values of  $\epsilon$  and  $\sigma$ , has not been optimized for thermodynamic property evaluation. It represents an effective spherical approximation, applicable to the transport properties, to the true two-body interaction.

Rather than directly evaluate the collision integral, we fitted the tabulated results of Klein et al.<sup>8</sup> to the form

$$\Omega^{(2,2)*} = \left[\sum_{i=1}^{9} C_i t^{[(i-1)3 - 1]}\right]^{-1}.$$
 (12)

The coefficients  $C_i$  for Eq. (12) are listed in Table 8 and are identical to those published in Ref. 1. The fit agrees with the tabulated integration results<sup>8</sup> within about 0.1% in the (reduced) temperature range 0.5 < t < 200.

For the thermal conductivity  $\lambda_0$  of the dilute gas, a completely rigorous and calculable theory is not available for polyatomic molecules, due to the complexities of exchanging energy between internal and external (kinetic) degrees of freedom. As for methane, we have chosen a modified Eucken model<sup>14</sup> of the form

$$\lambda_0(T) = \frac{\eta_0(T)}{M_r} \left[ \frac{15R}{4} + f_{int}(C_p^{id} - 5R/2) \right]$$
(13)

$$= 0.276 \ 505 \ \eta_0(T) [ \ 3.75 \ - \ f_{int}(\tau^2 \Phi_{\tau\tau}^{id} + \ 1.5)]$$
$$m W \cdot m^{-1} \cdot K^{-1}, \qquad (13a)$$

where  $C_p^{id}$  is the (temperature dependent) ideal gas contribution to the molar isobaric heat capacity and  $f_{int}$  is a dimensionless function which describes the energy exchange mentioned above. When using Eq. (13a), the viscosity should be expressed in micropascal-seconds, as in Eq. (10a). The heat capacity contribution, having been written in terms of a derivative of the ideal gas Helmholtz energy, can be evaluated using Tables 3 and 5. The form for  $f_{int}$  is chosen empirically as

$$f_{\rm int} = f_1 + (f_2 / t) \tag{14}$$

with the coefficients f fitted to experimental data and also given in Table 8.

#### 2.5.2. Excess Property Correlation

For the excess viscosity, we use a rational polynomial in the reduced density  $\delta$  and inverse reduced temperature  $\tau$ ; the viscosity is scaled by dimensional analysis and critical point variables. (Note that the parameter  $t = Tk/\epsilon$ , used above, is the temperature reduced by the interparticle potential energy scale, while  $\tau = T/T$  is the inverse temperature reduced by the critical temperature.) Thus, we write

$$\eta_{ex}(\rho,T) = \frac{P_c^{23} (M_r \mathbf{u})^{1/2}}{(T_c k)^{1/6}} \left[ \sum_{i=1}^{9} g_i \, \delta^{r_i} \, \tau^{s_i} \right] \left[ 1 + \sum_{i=10}^{11} g_i \, \delta^{r_i} \, \tau^{s_i} \right]^{-1}$$
(15)

$$= 15.977 \left[ \sum_{i=1}^{9} g_i \, \delta^{r_i} \, \tau^{s_i} \right] \left[ 1 + \sum_{i=10}^{11} g_i \, \delta^{r_i} \, \tau^{s_i} \right]^{-1} \, \mu \text{Pa·s, (15a)}$$

where the exponents  $r_i$  and  $s_i$  and the dimensionless fitted coefficients  $g_i$  are given in Table 9. The excess viscosity of fluids increases dramatically as the triple-point density of the liquid is approached; for conformality with other fluids a density reduced by  $\rho_{tL}$  could be used in the denominator of Eq. (15). However, we have incorporated the appropriate constants into  $g_{16}$  and  $g_{11}$  in order to simplify the equation. All of the terms in Eq. (15) were statistically significant and contributed to the reduction of the residuals for the primary data. The form was chosen to be consistent with the equation for methane,<sup>1</sup> and the temperature dependence in the excess function reflects the theoretical understanding of its presence.<sup>144</sup>

The excess thermal conductivity  $\lambda_{ex}$  has been correlated to a polynomial in  $\delta$  and  $\tau$ . As in Eq. (15), we include a prefactor with the appropriate dimensions and write

$$\lambda_{\rm ex}(\mathbf{p},T) = \frac{P_{\rm c}^{23} k^{56}}{T_{\rm c}^{16} (M_{\rm r} u)^{1/2}} \sum_{i=1}^{7} j_i \, \delta^{r_i} \, \tau^{s_i}$$
(16)

$$= 4.417 \quad 86 \left[ \sum_{i=1}^{7} j_i \, \delta^{r_i} \, \tau^{s_i} \right] \, \mathrm{mW} \cdot \mathrm{m}^{-1} \cdot \mathrm{K}^{-1}. \tag{17}$$

The exponents and dimensionless coefficients for Eq. (17) are given in Table 9; the form and exponents in this correlation differ slightly from those used for methane in Ref. 1. In both excess functions, Eqs.(15) and (17), there are strong density dependences and weak temperature dependences.

#### 2.5.3. Critical Enhancement Correlation

Both the viscosity and the thermal conductivity exhibit an increase near the liquid-vapor critical point, relative to values far from this singularity; theory indicates that the transport properties for pure fluids diverge to infinity at the critical point, although experimental difficulties limit the values found in the laboratory. The increase and theoretical divergence are presumably due to dynamic interactions among clusters formed because of large-scale density fluctuations.<sup>15</sup> It is only in the thermal conductivity that the divergence is strong and the enhancement is easily observed in a broad region around the critical point. For this reason, the present transport property correlations include an enhancement contribution only for the thermal conductivity.

Our treatment of the critical enhancement term is substantially different from that given in the methane manuscript;<sup>1</sup> the new mode coupling treatment of Olchowy and Sengers,<sup>9</sup> referred to briefly in Ref. 1, seems to make the older approach of Ref. 15, used in Ref. 1, obsolete. We have chosen to use a simplified version of the new theory, also presented by Olchowy and Sengers.<sup>9</sup> The simplified theory avoids the complications of implicitly defined functions and root-finding algorithms, and it

#### THERMOPHYSICAL PROPERTIES OF ETHANE

Pressure:	$P(\rho,T) = \rho RT (1 + \delta \phi_{\delta})$
Internal Energy:	$U(\rho,T) = RT \left( \tau \phi_{\tau}^{id} + \tau \phi_{\tau}^{\prime} \right)$
Enthalpy:	$H(\rho,T) = RT \left( 1 + \tau \phi_{\tau}^{id} + \tau \phi_{\tau}^{r} + \delta \phi_{\delta}^{r} \right)$
Gibbs free Energy:	$G(\rho,T) = RT \left( 1 + \phi^{id} + \phi^{r} + \delta \phi_{\delta}^{r} \right)$
Helmholtz Free Energy:	$A(\rho,T) = RT \left( \phi^{id} + \phi^{r} \right)$
Entropy:	$S(\rho,T) = -R \left( \phi^{id} + \phi^{r} - \tau \phi^{id}_{\tau} - \tau \phi^{r}_{\tau} \right)$
Isochoric Heat Capacity:	$C_{\nu}(\rho,T) = -R \left( \tau^2 \Phi_{\tau\tau}^{id} + \tau^2 \Phi_{\tau\tau}^{\prime} \right)$
Isobaric Heat Capacity:	$C_{\rho}(\rho,T) = C_{\nu}(\rho,T) + R \frac{(1+\delta\varphi_{\delta}^{\prime}-\delta\tau\varphi_{\delta\tau}^{\prime})^{2}}{1+2\delta\varphi_{\delta}^{\prime}+\delta^{2}\varphi_{\delta\delta}^{\prime}}$
Saturated Liquid	
Heat Capacity:	$C_{\sigma L}(T) = C_{\nu}(\rho_{\sigma L},T) - R (1 + \delta \phi_{\delta}^{r} - \delta \tau \phi_{\delta r}^{r}) \frac{T}{\rho_{\sigma L}} \frac{d\rho_{\sigma L}}{dT}$
Speed of Sound:	$w^{2}(\rho,T) = \frac{RT}{uN_{A}M_{r}} \frac{C_{\rho}(\rho,T)}{C_{v}(\rho,T)} \left(1 + 2\delta\phi_{\delta}^{\prime} + \delta^{2}\phi_{\delta\delta}^{\prime}\right)$
Second Virial Coefficient:	$B(T) = \frac{1}{\rho_c} \lim_{\delta \to 0} \phi_{\delta}^r$

TABLE 7. Thermodynamic property equations

seems appropriate within the present context of correlating equations developed for ease of use as well as accuracy.

Note added in final revision: We have recently presented expressions to evaluate the mode coupling integral in closed algebraic form.<sup>71</sup> Thus the full theory of Ref. [9] is now nearly as easy to use as the simplified theory and is to be preferred in future work.

In addition, we retain the completely classical description, as given above, of all thermodynamic quantities which must be calculated for the critical enhancement

TABLE 8. Coefficients for dilute gas transport properties

	Ω <sup>(2,2)*</sup> , E	Eq.(12)		f <sub>int</sub> , Eq. (14)			
<i>C</i> <sub>1</sub>	- 3.032	813	828 1				
$C_2$	16.918	880	086	$f_1$	1.710 414 7		
$C_3$	- 37.189	364	917				
$C_4$	41.288	861	858	$f_2$	-0.693 648 2		
<i>C</i> <sub>5</sub>	- 24.615	921	140				
$C_6$	8.948	843	095 9				
$C_7$	- 1.873	924	504 2				
$C_8$	0.209	661	013 90				
<i>C</i> 9	- 9.657	043	$707 4 \times 10^{-3}$				

term. While this again avoids certain complications inherent in the revised and extended scaling description of the thermodynamic surface, it causes substantial deviations from the expected behavior in the asymptotically critical region. The available data for ethane are not sufficiently close to the critical point to exhibit these deviations. Further details and appropriate ranges and uncertainties in the correlation are given in Sec. 4.

The expression for the enhancement from Ref. 9 is given by

$$\lambda_{cr}(\rho,T) = \frac{\Lambda k T \rho C_P}{6 \pi \eta(\rho,T) \xi} F(\rho,T)$$
(18)

$$= 1.55 \frac{\delta}{\tau} \frac{C_P}{\eta \xi} F(\delta, \tau) \text{ mW·m}^{-1} \cdot \text{K}^{-1}, \qquad (18a)$$

where the molar isobaric heat capacity is expressed in J·mol<sup>-1</sup>·K<sup>-1</sup>, the correlation length,  $\xi$  defined by Eq. (19), is in nm, and the viscosity is expressed in  $\mu$ Pa·s in Eq. (18a). The constant  $\Lambda$  has the value 1.01 and arises from dynamic mode-coupling theory, and the damping function F is also determined from theory and is defined in Eq. (20). The function  $\lambda_{cr}(\rho,T)$  of Eq. (18) is essentially identical to the expression in Ref. 1 in the asymptotically critical region; however, the function F differs significantly. When  $C_P$  in Eq. (18), as well as the viscosity and

η <sub>ex</sub> , Eq. (15)				λ <sub>ex</sub> , Eq. (17)					
i	r <sub>i</sub>	Si	g	'i	r <sub>i</sub>	Si		jı	
1	1	0	0.471 770	03	1	0	0.960	843	22
2	1	1	-0.239 503	3 11	2	0	2.750	023	5
3	2	0	0.398 083	3 01	3	0	- 0.026	609	289
4	2	1	-0.273 433	3 35	4	0	- 0.078	146	729
5	2	1.5	0.351 922	2 60	5	0	0.218	813	39
6	3	0	-0.211 013	3 08	1	1.5	2.384	956	3
7	3	2	-0.004 785	5 79	3	1	-0.751	139	71
8	4	0	0.073 781	29					
9	4	1	-0.030 425	5 255					
10	1	0	-0.304 352	2 86					
11	1	1	0.001 215	5 675					

=

TABLE 9. Coefficients for excess transport properties

derivatives in the definition of  $\xi$  given in Eq. (19), are evaluated from the analytical equation of state, the resultant  $\lambda_{cr}$  is not correct in the asymptotically critical region. In particular, the critical exponent describing  $C_P$  along the critical isochore has its classical value of 1 rather than the theoretical value near 1.24; considering all of the retained classical contributions, the critical exponent for  $\lambda_{cr}$ along the critical isochore is near 0.5 instead of having a value greater than 0.6 as is usually observed in the asymptotic region.

The correlation length used in Eq. (18) represents the quantity related to the critical region fluctuations; we subtract a background term and write

$$\xi = \xi_0 \Gamma_0^{-\nu/\gamma} \left( \frac{P_c}{\rho_c^2} \right)^{\nu/\gamma} \left[ \rho \left( \frac{\partial \rho}{\partial P} \right|_T - \frac{2T_c}{T} \left. \frac{\partial \rho}{\partial P} \right|_{T=2T_c} \right)^{\nu/\gamma} \quad (19)$$

$$= 0.428 \ (\delta\tau)^{0.507} \ [(1+2\delta\phi_{\delta}^{r} + \delta^{2}\phi_{\delta\delta}^{r})^{-1} \ - \ (1+2\delta\phi_{\delta}^{r}(1/2)$$

+ 
$$\delta^2 \phi_{\delta\delta}^r (1/2))^{-1} ]^{0.507}$$
 nm. (19a)

The critical amplitudes and exponents, obtained from Ref. 9, have been evaluated for ethane in Eq. (19a) and are given in Table 10; the choice  $2T_c$  as the cutoff temperature is arbitrary, but the resultant correlation is not very sensitive to the choice. In Eq. (19a), the arguments of the derivatives of the residual Helmholtz energy are the usual reduced density and temperature in the first terms. The final term in Eq. (19a) is evaluated at the reduced density of the state point for which  $\lambda_c$  is being evaluated and  $\tau = 0.5$  (as indicated) corresponding to a temperature of  $2T_c$ . When the term in square brackets becomes negative, very far from the critical point where the enhancement term is negligible, it should be set equal to 0.

The damping function in Eq. (18) is given by

$$F = \frac{2}{\pi} \left[ e^{-q_{\rm D} \xi \left[1 + (q_{\rm D} \xi)^3 / (3\delta^2)\right]^{-1}} + \frac{C_P - C_V}{C_P} \left( \tan^{-1}(q_{\rm D} \xi) + \frac{C_V}{C_P - C_V} q_{\rm D} \xi \right) \right]$$
(20)

and  $q_D^{-1} = 0.545$  nm is the only fitted parameter in the theory. The quantity  $q_D$  represents a cutoff wavenumber above which the contribution to the transport properties due to the coupling of hydrodynamic modes vanishes. Equivalently,  $q_D^{-1}$  represents a second length scale; fluctuations over lengths shorter than this scale do not contribute to the dynamical critical phenomena. The correlation length  $\xi$  is from Eq. (19), the reduced density is given by  $\delta = \rho/\rho_c$  as usual, and the heat capacities can be evaluated from the equation of state correlation. Again, we note that to the extent that the heat capacities do not have the correct critical behavior, the damping function will not extrapolate well in the asymptotically critical region.

TABLE 10. Constants for  $\lambda_{cr}$ , Eq. (18) [Using Eqs. (19) and (20)]

Fitted coefficient :	$q_{\rm D}^{-1} = 0.545 \ {\rm nm}$
Critical exponents:	$\begin{array}{l} \gamma \ = \ 1.242 \\ \nu \ = \ 0.63 \end{array}$
Universal constant:	$\Lambda = 1.01$
Fluid dependent amplitudes from Ref. 9:	$\xi_0 = 0.19 \text{ nm}$ $\Gamma_0 = 0.0563$

## 3. Development of the Correlations

## 3.1. Fundamental Constants, Fixed Points, and Ideal Gas Properties

The fundamental constants which were used are given in Table 1 and agree with the values recommended by the Committee on Data for Science and Technology,<sup>16</sup> CODATA in 1986. The relative molecular mass for ethane was derived from the atomic values of the recent IUPAC tables<sup>17</sup> and agrees with the value used by Goodwin *et al.*<sup>4</sup> The value 30.0694 adopted by Sychev *et al.*<sup>5</sup> is lower than ours by 0.002%, which is within the uncertainty of  $M_r$  computed from the uncertainties of the relative atomic masses given in Ref. [17]; the value from Ref. [5] was evidently derived from atomic masses promulgated by IUPAC<sup>18</sup> in 1980 or earlier. Uncertainties associated with these quantities can be found in the original references.

The values of the fixed point constants at the triple point of ethane were not needed for the present correlations. However, they determine the range of validity of many of the equations of Sec. 2 and may be of interest in their own right; therefore they are presented in Table 1. We have adopted the value of Pavese<sup>19</sup> for the triple point temperature of ethane; however we have increased the uncertainty in *T*, as as implied in the more recent study by Bedford *et al.*<sup>20</sup> The measurement in Ref. [20] was made by adiabatic calorimetry. Within uncertainties, our adopted value agrees with the derived result of Straty and Tsumura (90.348 ± 0.005 K)<sup>21</sup> which was used in Refs. 4 and 5. Additional historical values for *T<sub>t</sub>* are given in Refs. 5 and 19.

For the pressure at the triple point, we have calculated a value from our vapor pressure curve, Eq. (4), at the selected temperature. This extremely low pressure is difficult to measure experimentally. The uncertainty was established by estimating the accuracy of this equation, the uncertainty of the temperature, the value calculated using the Maxwell construction from the SWEOS at  $T_i$ , and comparisons with other determinations of  $P_i$ . Our value of  $1.130 \pm 0.005$  Pa agrees with that adopted by Sychev *et al.*<sup>5</sup> ( $1.13 \pm 0.01$  Pa) within stated uncertainties and essentially with that given by Goodwin *et al.*<sup>4</sup> (1.1308Pa). The SWEOS gives 1.131 Pa, a difference of less than 0.1% from our adopted value and well within our estimated uncertainty.

The fluid densities at the triple point were obtained from the ancillary Eqs. (5) and (6) using the stated value of  $T_i$ . Their uncertainties were estimated as above. The vapor density is identical to that obtained with the ideal gas equation of state at the given conditions. For comparison, we note that Goodwin *et al.*<sup>4</sup> adopted 21.680 mol·dm<sup>-3</sup> and 1.5154 mol·dam<sup>-3</sup>, without uncertainties, for the densities of the liquid and vapor, respectively. From the SWEOS, with the Maxwell construction, we obtain 21.665 mol·dm<sup>-3</sup> and 1.505 mol·dam<sup>-3</sup>.

The state variables at the critical point of ethane enter directly into many of the equations of Sec. 2. During the development of the present correlations, we attempted to improve the overall fit by allowing the critical parameters to vary. Since we did not find an alternative set of values which significantly improved the quality of the fits, we chose to use the critical values of Douslin and Harrison,<sup>22</sup> which they obtained from careful analysis of their nearcritical *PVT* data. Our choices conform to those adopted by Sengers and his collaborators<sup>23</sup> in their study of the asymptotically critical and crossover regions in the thermodynamic and transport properties of ethane. Sychev *et al.*<sup>5</sup> give an extensive table of various determinations of the critical point parameters.

Our selected critical temperature,  $305.33 \pm 0.04$  K, agrees within uncertainties with the temperatures used by Goodwin et al.<sup>4</sup> and by Sychev et al.<sup>5</sup> The value from Ref. 22, given without an estimate of the uncertainty, is based on the symmetry of the coexistence envelope in the temperature range 250 to 305.25 K. A visual determination of the critical point of ethane by Strumpf et al.24 gave the value  $305.368 \pm 0.005$  K; the uncertainty we present in Table 1 (0.04 K) is an estimate which does not rule out Strumpf's value. Burton and Balzarini<sup>24a</sup> estimate the critical temperature as  $305.229 \pm 0.03$  K, but, because they were mainly interested in temperature differences near the critical point, their thermometer was not calibrated according to IPTS-68. In a later fit of the coexistence data of Ref. 24a, based on a critical exponent  $\beta = 0.327$  and two Wegner correction terms<sup>24b</sup> with  $\Delta = 0.5$ , Pestak et al.<sup>24c</sup> obtained  $T_c = 305.2692$  K. Although no uncertainty was given with this determination and the calibration problem was not explicitly addressed, reasonable error bars would imply agreement with our value.

For the critical pressure, our selected value,  $4.8718 \pm 0.005$  MPa, is again from Douslin and Harrison,<sup>22</sup> who interpolated in density and extrapolated in temperature from their critical region *PVT* measurements. Goodwin *et et al.*<sup>4</sup> used 4.8714 MPa, 0.4 kPa below our value, based on their vapor pressure equation evaluated at the critical temperature. Sychev *et al.*<sup>5</sup> adopted the value from Ref. [4]. (Our vapor pressure equation, Eq. (4), is forced to the assigned critical pressure at the critical temperature.) Our assigned uncertainty, identical to that given in Ref. 5, is based on a study of our ancillary equations and the SWEOS when considering the uncertainty of the critical temperature; the value adopted in Refs. 4 and 5 falls within our estimated uncertainty.

Douslin and Harrison<sup>22</sup> computed the value of the critical density  $6.87 \text{ mol}\cdot\text{dm}^{-3}$ , which we have adopted, by using the linearity of the rectilinear diameter established from their saturated liquid and vapor measurements. Goodwin *et al.*<sup>4</sup> found  $6.80 \text{ mol}\cdot\text{dm}^{-3}$  (1% below our value) by computing rectilinear diameters from assorted experimental data; Ref. 5 used this value. Burton and Balzarini<sup>24a</sup> examined the behavior of the rectilinear diameter for their coexisting density data and obtained  $6.857 \pm 0.01 \text{ mol}\cdot\text{dm}^{-3}$ . Their densities were from an index-of-refraction experiment calibrated with a direct determination of the Lorentz-Lorenz coefficient. We have examined fits to the saturated liquid data from various sources (see Sec. 3.2) allowing the critical density to vary. In addition, we have used the liquid volume fraction method of Van Poolen *et al.*<sup>25</sup> which extrapolates from saturation densities far from the critical point (from 220 to 270 K, in our calculation). We obtained values for  $\rho_c$  between 6.870 and 6.893 mol·dm<sup>-3</sup>. Our selected value best fits the extensive critical region data of Ref. 22, and alternative values had little effect on the quality of the SWEOS. Our uncertainty of 0.1 mol·dm<sup>-3</sup> corresponds to a shift in the critical temperature of about 0.5 mK in our ancillary equations and seems realistic when considering the dispersion among different determinations and the difficulty of measuring critical point properties.

The intermolecular potential parameters  $\sigma$  and  $\epsilon$  in Table 1 were obtained by fitting low density transport data as discussed in Sec. 3.4.1. Our value for  $\epsilon$  is 2% above that reported by Hanley et al.<sup>26</sup> using earlier data, while that for  $\sigma$  differs by only 0.3%. The correlations of Ref. 26 formed the basis of the newer correlations in Ref. 10; in the present work, we have examined additional data. The same potential function, Eq. (11), was used in these studies. A very recent determination of the scaling parameters by Boushehri et al.,27 based on a corresponding states assumption, gives  $\epsilon/k = 241.9$  K (1.3% below our value) and  $\sigma = 0.4371$  nm (0.1% above the present value). This discrepancy is not surprising since the parameters depend on the form of the model potential which is assumed and on the particular properties being fitted. We emphasize that the intermolecular potential parameters of Table 1 are for the 11–6–8,  $\gamma = 3$  potential of Eq. (11) and are not appropriate when using the Lennard-Jones (12-6) potential; Mourits and Rummens<sup>28</sup> obtained 227.9 K and 0.4407 nm for the Lennard-Jones parameters. Good results cannot be expected when calculating equilibrium thermodynamic properties (such as the second virial coefficient) using Eq. (11) with the given parameters.

The remaining entries in Table 1, those concerning the values of the entropy and enthalpy at standard conditions, were from Chao *et al*.;<sup>11</sup> our correlation of the ideal gas Helmhotz energy, Eq. (3), gives these values, and they were also adopted in Ref. [10]. As noted in Sec. 2.2, these and all quantities evaluated using either the ideal gas Helmholtz energy [Eq. (3)] itself or its first temperature derivative are relative values. Contributions from the nuclear spin system have been excluded; the value of the ideal gas enthalpy is zero at zero temperature. The values in Table 1 are essentially equivalent to those found using the correlations of Refs. 4 and 5. A comparison with Wagman *et al.*,<sup>29</sup> based on a 1966 compilation, shows a discrepancy of 0.64% in the enthalpy and 0.16% in the entropy (adjusted to atmospheric pressure).

The development of the ideal gas correlations was discussed in Sec. 2.2. The tables of ideal gas properties of Chao *et al.*,<sup>11</sup> based on a rigid-rotor harmonic-oscillator model with contributions from the energy levels of the hindered internal rotation, formed the basis of our correlation. The isobaric heat capacities of the ideal gas<sup>11</sup> between the temperatures of 50 and 700 K were equally weighted in a linear least-squares algorithm to obtain the linear coefficients in  $\tau^2 \phi_{\tau\tau}^{id}$ , related to Eq. (3). The coefficient  $Q_7$  was found from a nonlinear algorithm, and an iterative procedure was used to obtain best values of all parameters. The linear fitter was constrained to the fixed value of the enthalpy at 298.15 K reported in Table 1, and the integration constant, related to  $Q_1$ , was established from S<sup>id</sup> at 298.15 K in Table 1. The ideal gas heat capacity and other ideal gas properties reported in Ref. 11 are compared to the resulting correlation in Sec. 4.2. The spectroscopically derived tables of Pamidimukkala et al.<sup>30</sup> use the same model as in the paper by Chao et al.<sup>11</sup> and give essentially identical results in the region of overlapping temperatures; they were not used as primary data because fewer points are available in the region of interest and the scatter relative to our correlating equation was somewhat higher. Comparisons for these data, other data, and other ideal gas correlations, and an assessment of the uncertainty associated with our ideal gas correlation, are given in Sec. 4.2.

## 3.2. Ancillary Equations for the Two-Phase Boundary

In the ancillary equations, Eqs. (4–6), for the saturated vapor pressure and the densities of the coexisting liquid and vapor, the critical constants were considered fixed at the values given in Table 1. In addition, the exponents  $\beta$  and  $\epsilon$  were given their effective scaling law values, as in Ref. 1. The forms and exponents in these equations are identical to those motivated in Ref. 1; only the nonzero value of  $G_5$  in Eq. (5) for the density of the saturated liquid, as mentioned in Sec. 2, and the values of all coefficients as given in Table 4, distinguish between the correlations for ethane. Unlike our study of methane<sup>1</sup> however, in this study we have used several experimental sources to provide primary data in establishing the coefficients in Eqs. (4–6).

For the pressure of the saturated vapor, 127 data from Pal et al.,<sup>31</sup> Pope,<sup>32</sup> Ziegler et al.,<sup>33</sup> Douslin and Harrison,<sup>22</sup> and Straty and Tsumura<sup>34</sup> in the range 100 to 305 K were used. All points were equally weighted in the fitting routine; however 7 points from the experimental papers, at temperatures near 214, 300.4, and 221 K in Ref. 31; 210, 255, and 305.4 K (above our critical temperature) in Ref. 32; and 160 K in Ref. [34], exhibited large deviations from our preliminary fits and were omitted from the final fit. The input data were essentially identical to those used in the correlation of Goodwin et al.<sup>4</sup> The monograph by Sychev et al.<sup>5</sup> lists additional (especially earlier) sources of data for the saturated vapor pressure, but uses the correlation of Ref. 4 to calculate vapor pressures. To obtain the coefficients  $H_i$ , Eq. (4) was linearized by taking the logarithm of both sides, and a linear least-squares fitting routine was used. Comparisons between the correlations and the experimental data are deferred to Sec. 4.1.

The correlation for the density of the liquid along the two-phase boundary, Eq. (5), was obtained in an analogous manner; 66 data from Douslin and Harrison,<sup>22</sup> Sliwinski,35 Canfield and collaborators,36 Rodosevich and Miller,<sup>37</sup> Haynes and Hiza,<sup>38</sup> McClune,<sup>39</sup> and Orrit and Laupretre<sup>40</sup> were used to determine the final coefficients. Data from other sources were also examined and are compared with the correlation in Sec. 4.1. Additional references to experimental work on  $\rho_{\sigma L}$  are given in Ref. 5. Points near 296 K and above 300 K in Ref. 35 and points from Ref. 40 not listed in the tables of Ref. 4 were exluded; from Ref. 22, only points near 302, 303, and 304 K were included in the determination of the coefficients listed in Table 4. These primary data, with temperatures ranging from 91 to 305 K, were equally weighted and a linear least-squares algorithm was used with a linearized form of Eq. (5).

Equation (6) for the density of the saturated vapor required expressions for both the vapor pressure and the density of the saturated liquid in order to ascertain values for the coefficients,  $J_i$ . As mentioned above, the value of  $J_0$  was completely determined from the fit for Eq. (5). The value 8 for the exponent of  $\tau$  was chosen to optimize the fit in Ref. 1. The expression was then linearized and the coefficients determined by a least-squares routine. The input data consisted of 32 equally weighted points from Goodwin et al.,<sup>4</sup> Douslin and Harrison,<sup>22</sup> and Sliwinski.<sup>35</sup> The points attributed to Ref. 4 comprised 18 points between 110 and 280 K obtained from the intersection of their vapor pressure and virial equations or other correlations; these are not experimental data but are needed to fill an important gap in the data. From Ref. 22, we used points below 290 K and near 303 and 304 K; from Ref. 35, data near 288 K and above 304 K were omitted. Other, especially earlier, sources of density data are cited in Ref. 5. All comparisons are deferred to Sec. 4.1.

## 3.3. Residual Helmholtz Energy

The coefficients associated with Eq. (2) were determined by multiproperty linear least-squares fitting, with the exponents used,  $r_i$  and  $s_i$ , identical to those determined by Schmidt and Wagner<sup>6</sup> and to those used for the methane correlation of Ref. 1. The critical parameters of Table 1 were introduced into the fitting routine as constraints on the pressure and its first two isothermal density derivatives at the critical point. Thus we required

and

$$P(\rho_c,T_c) = P_c \qquad (21)$$

$$\frac{\partial P(\rho_{\rm c},T_{\rm c})}{\partial \rho}\Big|_{T} = \frac{\partial^2 P(\rho_{\rm c},T_{\rm c})}{\partial \rho^2}\Big|_{T} = 0.$$
(22)

The resultant SWEOS gives the critical pressure exactly at the critical temperature and density; because of roundoff error, the first isothermal derivative is about  $10^{-11}$  MPa·dm<sup>3</sup>·mol<sup>-1</sup> and the second derivative is near  $10^{-12}$  MPa·dm<sup>6</sup>·mol<sup>-2</sup>. Because of this "accidental" deviation from 0, the isobaric heat capacity, strongly divergent in the critical region according to theory and experimental evidence, remains finite (but very large) at the critical point.

The densities and pressures along the two-phase boundary; second virial coefficient data; *PVT* data; molar heat capacities at isochoric and isobaric conditions and along the saturated liquid boundary; and speed of sound in the single phase and along the liquid phase boundary were used in the determination of coefficients. The sources of primary data and the details concerning the use of different types of thermodynamic data in our fitting scheme are discussed in this section. Figure 1c illustrates all the primary data in pressure-temperature coordinates. Comparisons between the data and the correlation, including figures and tables of statistical comparisons, are given in Sec 4.3.

The relative weights of data within the fitting routine were obtained from a determination of the type of thermodynamic data, the source of the experimental data, the region of the phase diagram, and the Gaussian error propagation formula. A complete discussion of the calculation of the weight for each of the approximately 1800 points used is not feasible in this paper, but some indication will be given in this section. The data and their weights, within the linear least-squares algorithm used to obtain the final values of the coefficients as given in Table 2, are tabulated in Ref. 3a.

The overall multiplier used to weight a particular property was determined by a nonlinear simplex algorithm.<sup>41</sup> At vertices of the generated simplex, the linear leastsquares problem was solved. This optimization technique provides a rational procedure for determining parameters which are not linearly related to the objective function. We used the algorithm in determining relative weights among data sets and data types. The assignment of accuracies in both dependent and independent variables by experimentalists is often overly optimistic, and the data are too often inconsistent. The linear fitting technique cannot easily compensate for these inconsistencies. Even with the simplex method, there is a difficulty in choosing the objective function. The simplex algorithm did provide a consistent method to exclude sets of data from the primary data set and to determine relative weights.

We sought to simultaneously obtain good agreement for all data types, with particular emphasis paid to reproducing the two-phase boundary defined by the ancillary equations. Relative weights ascribed to different data sources within a particular data type were also determined with a simplex nonlinear least-squares method.

It was impossible to fit all data within the stated experimental accuracy since data were frequently incompatible within the stated accuracies. Further it is sometimes impossible to fit certain highly accurate data to within experimental error even though they may be compatible with other data of lower accuracy. This is often the case when combining data such as speed of sound with other types of data. When data were incompatible in a particular region of the phase diagram, determination of emphasis and de-emphasis, reflected in the relative weight given to the data, was made by considering the intrinsic merits of the experimental methods used, as well as by examining the agreement with additional data both within and outside the region of interest.

The Gaussian error formula was invoked to establish the relative weights of data from a particular source or within a particular type. We assumed fixed relative or absolute accuracies for the various dependent and independent quantities involved in each type of data. Within each data source, the same accuracies were assumed throughout the range of the data. Thus, if z = z(x,y) with absolute accuracies  $\sigma_x, \sigma_y$ , and  $\sigma_z$  (or relative accuracies  $\sigma_z/z$ , etc.) the relative weight is given by

Wt 
$$\propto \left[\sigma_{z}^{2} + \left(\sigma_{x}\frac{\partial z}{\partial x}\Big|_{Y}\right)^{2} + \left(\sigma_{y}\frac{\partial z}{\partial y}\Big|_{x}\right)^{2}\right]^{-1/2}$$
. (23)

In a few instances individual data were omitted from the final determination of the fitted coefficients. These data were excluded when the deviations from a preliminary version of the correlation were well beyond deviations for similar points or differed greatly from trends determined from similar points. These points are enumerated in Ref. 3a where the assigned weight of 0 is shown and are indicated in this section.

The fitting of the saturation boundary PVT data was exceptional in that data generated from the ancillary equations were input to the least-squares routine. Thus, values of  $P_{\sigma}$ ,  $\rho_{\sigma V}$ , and  $\rho_{\sigma L}$  at 43 temperatures evenly spaced between 91 and 301 K were evaluated from the ancillary equations, Eqs. (4-6) of Sec. 2.3. This was done to ensure that the saturation properties determined from the equation of state are as accurate as possible; the twophase boundary determined from the Helmholtz energy correlation is completely compatible with the ancillary equations. Three distinct minimization conditions  $[P_{\sigma}(T)]$  $-P(\rho_{\sigma V},T), P_{\sigma}(T) - P(\rho_{\sigma L},T)$ , and the difference in the Gibbs energies of the two phases,  $A(\rho_{\sigma V}, T) + P_{\sigma}/\rho_{\sigma V} A(\rho_{\sigma L},T) - P_{\sigma}/\rho_{\sigma L}$ ] associated with the saturation boundary were incorporated within the fitting routine. In these expressions, the ideal gas contributions to the pressure and thermodynamic potential were subtracted, since only the residual molar Helmholtz energy was determined from the fitting procedure. The functions of density and temperature were evaluated as indicated in Table 7. All three of these conditions received heavy weighting, with extra emphasis placed on the density of the saturated liquid in the critical region. Below 180 K, the weights of both the vapor density and equality of the Gibbs energies were made somewhat lower than the weights obtained from the Gaussian formula and the simplex algorithm which gave an overall multiplier for this type of data. Figures 4-6 of Sec. 4.1 compare saturation properties calculated from the ancillary equations with those calculated from the SWEOS.

The data for the second virial coefficient which were used in this correlation were from Michels *et al.*,<sup>42</sup> Pope et al.,43 Douslin and Harrison,22 and Mansoorian et al.44 The 35 data spanned the range 210-623 K. Points at lower temperatures, based on corresponding states arguments by McGlashan and Potter<sup>45</sup> and adjusted by Goodwin et al.,<sup>4</sup> were used as primary data in Ref. 4 but have been excluded from our fit. The zero- density limit indicated in the expression for the second virial coefficient in Table 7 implies that only the parameters  $n_i$  with i =1,2,3,14, and 15 are directly affected by the virial coefficient data. The weighting for the virial data tended to be high. The simplex nonlinear fitting routine indicated that we could establish the greatest consistency with other thermodynamic data by heavily weighting the data from Refs. 22 and 44 and placing less emphasis on the points from Ref. 43. Other data were also considered, as discussed in the comparison section, Sec. 4.3.1. Experimentally determined third virial coefficient data were not used in the present correlation. These data are difficult to obtain and consequently suffer from inaccuracies.

The largest body of thermodynamic data for ethane consists of PVT measurements. We used, as primary data, 803 points from Pal et al.,<sup>31</sup> Straty and Tsumura,<sup>34</sup> Douslin and Harrison,<sup>22</sup> Michels *et al.*,<sup>42</sup> Parrish,<sup>46</sup> and Sengers.<sup>23</sup> From Ref. 31, only data at pressures greater than 35 MPa were weighted; these 45 data have a range of 36-69 MPa, 183-320 K, and 15-19 mol·dm<sup>-3</sup>. The 381 selected data from Ref. 34 span the range 96-320 K, 3-38 MPa, and 2-21.6 mol·dm<sup>-3</sup>. Some low density data along the 1.2 and 4.3 mol·dm<sup>-3</sup> isochores, near-critical data at densities near 6.5 mol·dm<sup>-3</sup>, and liquid data near the saturation boundary were not weighted in the correlation. Reference 22 provided 257 primary data with temperatures from 273 to 623 K, pressures from 1.4 to 40 MPa, and densities from 0.75 to 10.5 mol·dm<sup>-3</sup>; points outside this range and near the coexistence boundary and critical isotherm from Table 1 of Ref. 22 were not used to determine the final coefficients of Eq. (2). The primary data from Ref. 42 consisted of 81 points between 273 and 423 K, 1.6 and 10 MPa, and 0.85 and 3.6 mol·dm<sup>-3</sup>. The higher density data from Michels et al.<sup>42</sup> were not included in the fitting routine. The 9 PVT points from Ref. 46 were in the range 300-322 K, 5.5-9.7 MPa, and 4-12 mol·dm<sup>-3</sup>.

Finally, 30 single-phase PVT points in the critical region were generated from the (revised and extended) scaled equation of state provided by Sengers.<sup>23</sup> This equation of state was based largely on the PVT data of Ref. 22. These 30 data serve to ensure that the classical SWEOS formulation conforms closely to the known nonanalytic behavior of the fluid in the critical region. The data were within the region bounded by 304 and 325 K, 4.7 and 8.1 MPa, and 4.9 and 9 mol·dm<sup>-3</sup>. These and other data are compared with the SWEOS in Sec. 4.; Table 12 in Sec. 4.3.2 summarizes the data and statistical comparisons and Figs. 9–12 illustrate the deviations between the data and the SWEOS.

The determination of the relative weights of each source of PVT data was made using the simplex algorithm discussed above; other data sources, discussed in the

comparison section, Sec. 4.3.2, were excluded from the final fit on the basis of preliminary correlations. The data of Refs. 23 and 34 were assigned the largest overall weighting factors. The actual weight for each datum can be found in Ref. 3a. The *PVT* data entered the fitting routine through minimization of the expression  $P(\rho,T) - \rho RT = \rho RT \delta \phi'_{\delta}(\delta, \tau)$ .

For the molar heat capacity of ethane at constant volume  $C_V$ , we have selected 184 points from Roder<sup>47</sup> and 30 points from Sengers<sup>23</sup> as input into the SWEOS correlation. The data from Ref. 47 span the range 112-329 K, 1.6- 34 MPa, 1.6-21 mol·dm<sup>-3</sup>, and 44-63 J·mol<sup>-1</sup>·K<sup>-1</sup>. Some data near the critical point, near the saturated liquid boundary, and near 18.9 mol·dm<sup>-3</sup> have been excluded; data along the highest density isochore, 20.5 mol·dm<sup>-3</sup>, were given increased weighting in the fitting algorithm. The scaled equation of state from Ref. 23 was used to generate isochoric heat capacities at the same (near-critical) state points for which PVT data were generated; the resultant heat capacities range from 57 to 85 J·mol<sup>-1</sup>·K<sup>-1</sup>. A classical equation of state cannot incorporate the weak divergence of the isochoric heat capacity at the critical point, and our final SWEOS gives a local maximum in  $C_V$  of about 71.4 J·mol<sup>-1</sup>·K<sup>-1</sup> at  $T_c$  and  $\rho = 6.65$ mol·dm<sup>-3</sup>. For this reason, we did not generate points from the scaled equation<sup>23</sup> which were closer to the critical point and which might serve to distort the SWEOS in other regions; also, the input data from Ref. 23 were given less emphasis, by smaller relative weighting, than the wide ranging data of Ref. 47.

The experimental densities associated with the  $C_V$  measurements, as reported in Ref. 47, were based on an older equation of state<sup>4</sup> as well as absolute calibration of certain volumes. These densities were used as input in the development of the SWEOS, and the resultant uncertainty in the tabulated experimental densities seemed to have no significant effect on the present correlation. The ideal gas contribution to the heat capacity, which typically amounted to 70–90% of the total, was subtracted from the data. The second temperature derivative of the residual Helmholtz function, which is related to  $C_V$  as indicated in Table 7, was fitted to the resultant residual isochoric heat capacity. Comparisons between the  $C_V$  data and the SWEOS are summarized in Table 13 and illustrated in Fig. 15 of Sec. 4.3.3.

Two sources of molar isobaric heat capacities were used to obtain the final coefficients of Eq. (2): 241 points from Furtado<sup>48</sup> and 118 points from Bier *et al.*<sup>49</sup> The input data from Ref. 48 were in the range 100–378 K, 1.7–14 MPa, and 60–700 J·mol<sup>-1</sup>·K<sup>-1</sup>; several points with large deviations or within the general critical region were excluded from the fit. Bier *et al.* provided additional primary data from 283 to 473 K, 0.1 to 10 MPa, and 50 to 460 J·mol<sup>-1</sup>·K<sup>-1</sup>; again, points in the critical region, from 305 to 313 K and 4.6 to 4.9 MPa, were excluded. Because  $C_p$ is nonlinear in the residual Helmholtz energy and its derivatives, as seen in Table 7, and the experimental variables are pressure and temperature (whereas the independent variables in the Helmholtz energy are density and temperature), the experimental data could not be inputted directly into the linear least-squares fitting routine. The input density was calculated from the experimental *P*-*T* state point using a previous iteration of the SWEOS. The fit was linearized by subtracting two terms from the experimental heat capacity: the ideal gas isochoric heat capacity at the same temperature and a term equal to  $T(\partial P/\partial T)_{P}^{2}(\partial \rho/\partial P)_{T}/\rho^{2}$  evaluated from the previous SWEOS. The remainder is the residual isochoric heat capacity, linear in  $\phi'_{\pi}$ , and was fitted as such. In the final determination of the SWEOS coefficients, as indicated by the results of the simplex algorithm, the  $C_{P}$ data of Furtado were more heavily weighted. The data are summarized in Table 13, and deviations from the SWEOS are illustrated in Fig. 16.

The molar heat capacity while maintaining the liquid at saturation  $C_{\sigma L}$  was also measured by Roder<sup>47</sup> and used as primary input for the SWEOS correlation. The 106 points span the temperature range from 94 to 301 K and the heat capacities ranged from 68 to 212 J·mol<sup>-1</sup>·K<sup>-1</sup>. This thermodynamic quantity is not linearly related to the Helmholtz energy as seen in Table 7. The required orthobaric derivative (that is, the derivative along the saturation boundary) was approximated by differences according to

$$C_{\sigma L}'(T) = T \frac{\partial S'(\rho_{\sigma L}, T)}{\partial T} \bigg|_{\sigma} \approx T \frac{S[\rho_{\sigma L}(T + \epsilon)] - S[\rho_{\sigma L}(T - \epsilon)]}{2\epsilon},$$
(24)

where the function  $\rho_{\sigma L}(T)$  was evaluated using the ancillary Eq. (5), and  $\epsilon$  was chosen to be 0.01 K. The corresponding expression for the ideal gas contribution to  $C_{\sigma L}$ , which was subtracted from the experimental data, is

$$C_{\sigma L}^{id}(T) \approx T \frac{\Delta S^{id}}{\Delta T} - RT \frac{\Delta \ln(\rho_{\sigma L} T)}{\Delta T}.$$
 (25)

Here the deltas indicate differences as in Eq. (24), and the temperature difference was taken to be 0.01 K, consistent with Eq. (24). Alternative linearization schemes, such as that implied by the  $C_{\sigma L}$  entry in Table 7, could also be used. Such schemes could make use of the analytically known derivative properties of the ancillary equation [Eq. (5)] instead of calculating the difference as in Eq. (25). The saturated liquid heat capacities were thus entered into the linear fit for the Helmholtz energy; they were assigned relatively small weights, but with more emphasis on the higher density (lower temperature) data. The data are summarized and compared with the SWEOS in Table 13; Fig. 17 illustrates the deviations between the data and the correlations.

The final type of thermodynamic data which we considered was the speed of sound, both in the single phase region and in the liquid along the saturation boundary. For the single phase fluid, we selected 109 points from Tsumura and Straty,<sup>50</sup> (including 2 points published in Ref. 4 but not in Ref. 50), and 92 points from Terres *et al.*<sup>51</sup> The measurements from Tsumura and Straty ex-

tended from 100 to 323 K, 3.6 to 37 MPa, and 300 to 2000  $m s^{-1}$ ; points along the critical isotherm, as well as certain points at 300 K and 323 K, were excluded from the fit. The data of Ref. 51 were between 293 and 448 K, 0.1 and 12 MPa, and 225 and 400  $m s^{-1}$ . Reference 50 also provided 47 points for the saturated liquid at temperatures from 91 to 299 K and with sound speeds decreasing from 2000  $m \cdot s^{-1}$  near the triple point temperature to 290  $m \cdot s^{-1}$  at the highest temperature; values above 270 K were used as (slightly) adjusted in Ref. 4, and the 4 tabulated points above 299 K were not used as primary data. These data are summarized in Table 14 and the deviations between the data and correlation are illustrated in Figs. 18 and 19.

Again, as indicated in Table 7, these data are not linearly related to the Helmholtz energy. The mechanism by which these data entered the fit was iterative, in that a preliminary version of the SWEOS was used to calculate  $\frac{\partial P}{\partial \rho}_T$  from the sound speed data according to

$$\frac{\partial P}{\partial \rho}\Big|_{T} = u N_{A} M_{r} w^{2}_{\exp} \frac{C_{V}(\rho, T)}{C_{P}(\rho, T)}$$
(26)

where the heat capacities and the densities are calculated from the experimental P-T state point. For the data on the saturation boundary, the Maxwell construction, using the preliminary SWEOS, was used to find the density. The values of the density derivative, with the corresponding density and experimental temperature, were then used as input into the next iteration of the fitting process. The residual portion of this derivative, as seen in Table 7, can be expressed as a linear combination of  $\phi'_{\delta}$  and  $\phi'_{\delta\delta}$ and was used in the fitting routine.

The coefficients,  $n_i$  of Eq. (2), as listed in Table 2, represent the best least-squares fit of the residual Helmholtz energy for ethane within the constraints discussed above. In Sec. 4 we present a comparison between the correlation and the experimental data.

#### 3.4. Transport Property Correlations

#### 3.4.1. Viscosity

The dilute gas viscosity is completely specified by the Chapman-Enskog expression, Eq. (10), with only the form of the potential and its parameters unknown. The flexibility of the m-6-8 family of potentials, its theoretically-based justification, and the utility of the specific 11-6-8,  $\gamma = 3$  form have been established by Hanley and Klein.<sup>13</sup> Thus we have chosen this potential, Eq. (11), which was also used in the methane correlations.<sup>1,2</sup> A fit to the associated dimensionless collision integral  $\Omega^{(2,2)*}$ , according Eq. (12), was discussed in Ref. 1. The accurate low-density viscosity measurements of Kestin et al.52 in the range 300 to 480 K were used to establish the distance and energy parameters of the potential function; 2 additional points reported by Kestin et al. in Refs. 53 and 54 were also considered primary data. Values of  $\epsilon/k$  in the approximate range 230 to 245 K were considered, and the

corresponding values of  $\sigma$  were established by means of a least-squares fit to the data with the points at maximum and minimum temperatures receiving increased weights. The tabulated values (Table 1) minimized the root-meansquare (RMS) deviations for the data, although there are alternative pairs of  $\epsilon$  and  $\sigma$  whose fit is only slightly inferior; a much larger value for the well depth,  $\epsilon/k \approx 370$  K, decreased the systematic deviations between the primary data and the correlation, but is untenable because it differs greatly from other reported values of this quantity (for various potentials and corresponding states arguments)7,10,14,26-28 and would not extrapolate well at the lower temperatures. Comparisons with other low density viscosity data and correlations are given in Sec. 4.4.1; Fig. 20 illustrates the deviations between the data and the correlating equation.

The form of Eq. (15) for the excess viscosity and the exponents  $r_i$  and  $s_i$  of  $\delta$  and  $\tau$ , were optimized from a limited set of proposed terms by a study of a variety of pure fluids in addition to ethane; the equation and choice of exponents are identical to those used in the methane correlation.<sup>1,2</sup> The dimensionless coefficients  $g_i$  for ethane were established by using a least-squares algorithm with primary experimental data from Diller and Saber,55 Diller,<sup>56</sup> and Carmichael and Sage<sup>57</sup> after considering several other data sets as listed in Sec. 4.4.1 on comparisons between the correlations and experimental measurements. The data are summarized in Table 15, deviations are illustrated in Fig. 21, and a map in P-T coordinates was included as Fig. 2b. We chose the primary data by using the simplex algorithm to establish the relative weights of the data sets with an objective function based on residual deviations for all the data. The choice is thus based on consistency of the primary data sets relative to the form of correlating equation which was used.

The data from Ref. 55 comprise 144 points with temperatures from 95 to 320 K and pressures between 0.3 and 32 MPa measured with an oscillating quartz-crystal viscometer designed for cryogenic use. The 72 primary data from Ref. 56 were obtained at NIST with a new apparatus, also using an oscillating quartz crystal, built for high temperature measurements; these data were taken at temperatures from 295 to 500 K with pressures between 2 and 55 MPa. Finally, Ref. 57 provided 222 primary data with temperatures between 300 and 478 K with pressures from 1 to 36 MPa. These were obtained with a rotating cylinder viscometer and were also considered primary in the correlation of Hanley *et al.*<sup>26</sup> They served to increase our coverage of the high temperature region.

The primary data used to determine the excess viscosity were equally weighted; a few outlying points, as determined by a preliminary fit, were omitted from the final determination of the coefficients. In all cases, a value of the zero density viscosity obtained from Eq. (10) was subtracted from the experimental data to establish the "experimental" excess viscosity. The experimental pressure and temperature were used to calculate the density from the present equation of state for input into the leastsquares algorithm. Explicit comparisons between the total viscosity correlation and both the primary and other data are given in Sec. 4.4.1.

#### 3.4.2. Thermal Conductivity

Only the coefficients in Eq. (14) must be established in order to complete the correlation for the zero-density thermal conductivity. Data from the transient hot-wire measurements of Roder and Nieto de Castro<sup>58</sup> and of Prasad and Venart<sup>59</sup> were used for this purpose. Reference 58 provided 7 points with temperatures between 245 and 312 K; the point at 305 K was not given any weight. Roder and Nieto de Castro obtained these data by extrapolating the measurements, made along isotherms, to zero density. Their data at lower temperatures, where the vapor pressure was below 1 MPa so that vapor-phase measurements were difficult to obtain, were not used to establish our correlating coefficients. The additional 5 points from Ref. 59, in the range between 315 and 600 K, were reported as extrapolations of measured isotherms to a pressure of 0.1 MPa. We adjusted these data, using a preliminary version of Eq. (17) for the excess thermal conductivity, to zero density. These adjustments were less than 0.2% of the reported value. The data at higher temperatures were given increased weighting in our fitting routine. Equation (13) was solved for  $f_{int}$  using the above correlations for  $\eta_0$  and  $C_P^{id}$  and these experimentally derived values of  $\lambda_0$ . We used the value of  $\epsilon/k$  reported in Table 1 to obtain the reduced temperature, and a leastsquares algorithm was then used to calculate the two coefficients of Eq. (14). The resulting values of  $f_1$  and  $f_2$  are reported in Table 8. Comparisons with experimental data are given in Sec. 4.4.2, and deviations are illustrated in Fig. 22.

Because it is difficult to unambiguously separate the excess thermal conductivity [Eq. (17)] from the critical enhancement contribution [Eq. (18)], the method of determining the relevant coefficients is more complicated than the analogous determination of coefficients relevant to the excess viscosity as discussed above. Preliminary determinations of the exponents and coefficients of Eq. (17) and of the cutoff wavenumber  $q_{\rm D}$  were made by separately fitting the two equations [Eqs. (17) and (18)] after restricting the data to lie within or outside the general critical region. This allowed an estimate of the size of the cuttoff parameter  $q_{\rm D}$  and assessment of the utility of various proposed terms in the excess function Eq. (17). However, the final estimation of the coefficients in Eq. (17) was made using all of the primary data. We simultaneously determined the cutoff parameter and the coefficients and temperature exponents of Eq. (17) by using the nonlinear simplex algorithm discussed in Sec. 3.3. Temperature dependencies were introduced into each power (through fifth order) in the density series, and the reported exponents in Table 9 represent an approximation to the best seven-coefficient fit. After approximating the parameter and exponents, as given in the tables, we refitted the coefficients with a linear least-squares routine. Because of recent theoretical progress,<sup>9</sup> we are using a more rigorous description of the critical enhancement contribution to the thermal conductivity than was available in the methane work.<sup>1</sup> The excess function also differs from that proposed in our study of methane; the older correlating equation was inadequate in describing the existing ethane thermal conductivity data with the current description of the enhancement.

The primary data for these fits comprised nearly 1100 points from Roder,<sup>60</sup> Prasad and Venart,<sup>59</sup> and Desmarest and Tufeu<sup>61</sup> as summarized in Table 16 of Sec. 4.4.2. Reference 60 provided 752 points in the range 112 to 328 K and 0.1 to 69 MPa. An additional 45 points reported in Ref. 60, including isolated data with large deviations and the portion of the 312 K isotherm denoted as the 78000 series in Ref. 60, were not used in the fit. The 235 primary data from Ref. 59 had temperatures between 293 and 600 K and pressures from 0.2 to 70 MPa; four points with large deviations were excluded from our primary data. Finally, Ref. 61 provided 111 points with temperatures from 308 to 365 K with pressures between 1 and 28 MPa. A pressure-temperature map of the primary thermal conductivity data is given as Fig. 3b.

The measurements of Roder<sup>60</sup> and Prasad and Venart<sup>59</sup> were made with a transient hot-wire method. The data from Roder at temperatures above the critical value were de-emphasized by decreased weighting in the fitting routine, and those from Prasad and Venart were de-emphasized below the critical temperature. Desmarest and Tufeu used a coaxial cylinder apparatus and emphasized the region around the critical point of ethane. An empirical weighting function was developed, and the resultant weights increased with an increase in density of the data and decreased with increasing value of the experimental excess thermal conductivity. Again, the sources of primary data were chosen by using the simplex algorithm to provide the best overall fit to the entire body of data (within constraints imposed by our estimates of the accuracy of the data). In each range of the independent variables, we chose to emphasize but one source of data in order to eliminate any spurious overfitting associated with inconsistent data. Comparisons of the correlation with both primary and other experimental data are considered in Sec. 4.4.2 and deviations are illustrated in Fig. 24.

## 4. Comparisons of Derived and Experimental Properties

In this section we compare the correlations discussed above with experimental data. These data consist of both primary data, which were explicitly used to develop the correlations, and other data, which were not used for any of several reasons including lack of adequate discussion of experimental details, unusually poor precision or accuracy, disagreement with better known results, or simply overly abundant data for a particular property within some region of the phase diagram. To avoid exceptionally long discussion, overly complicated figures, and too many figures, the comparisons given in this section are representative rather than exhaustive. References to other sources of experimental data can be found within the papers cited in our reference list.

In all the deviation plots, the zero line represents the appropriate correlation using the equations of Sec. 2 and the associated tables. The percentage deviations are computed as 100(cor - exp)/exp where "exp" represents the experimental value of a property and "cor" is the value computed from the correlation and the experimental value(s) of the independent variable(s).

The precision of a quantity is often measured by its reproducibility; for the correlations of this paper the precision concerns the round-off error and is not of primary interest. The accuracy is a measure of the deviation of a quantity from its true value; the estimates of the accuracies of our correlations represent our best assesment of the maximum difference between a quantity computed from the correlation given the independent variable(s) and the true physical quantity at the same value(s) of the independent variable(s). The accuracies of the correlations depend on the quantity to be calculated and the range of the independent variable(s) as discussed below. When an uncertainty band is quoted, we think that the true value lies within the band. Our error estimates represent the maximum deviation between a predicted value and the true value; these estimates are equivalent to the accuracy assessments. More than 99% of the primary data lie within our uncertainty estimates; thus, with conventional assumptions concerning the distribution of residuals, these are equivalent to 2.5 o accuracy estimates where  $\sigma$  is the standard deviation. Additional statistical quantities concerning the comparison of the correlations with experimental data are given below and in the tables.

## 4.1. Two-Phase Boundary

In Fig. 4, we show the deviation of experimental orthobaric pressure data from the correlation of Eq. (4). For the 127 primary data from Refs. 22 and 31-34, the average absolute percent deviation (AAD-%) is 0.04%. The consistency of these primary data, judged from the regions of overlap of the different experimental sources is also quite good, typically well within 0.1%. The data from Refs. 22, 31, 32, and 34 were obtained as part of PVT studies of ethane; expansion methods were employed by Pope<sup>32</sup> and Straty and Tsumura,<sup>34</sup> and an isochoric chamber technique and pycnometer were used in the work of Pal et al.<sup>31</sup> and Douslin and Harrison.<sup>22</sup> The excellent agreement with the lower temperature primary results of Ziegler et al.<sup>33</sup> is somewhat misleading, as these data were adjusted by Goodwin<sup>4,62</sup> to conform to his newer second virial coefficient data. The unadjusted data are also illustrated in Fig. 4 and have deviations to about 2.5% near the triple point temperature. Carruth and and Kobayashi63 report experimental determinations of these very low vapor pressures based on a steady-state gas-saturation technique with a reported accuracy of about 3%. These data, as well as the low temperature results of Regnier<sup>64</sup> and Djordjevich and Budenholzer,<sup>65</sup> are also indicated in Fig. 4. Data in the critical region, from Miniovich and Sorina,<sup>66</sup> agree to within 0.06% with our correlation and are included in the figure.

For the 220 data illustrated in Fig. 4, the AAD-% is 0.64%, with the major contribution to the deviations coming from the very small vapor pressures and hence large percentage deviations in the secondary data at low temperatures. A few data, reported for temperatures above our value of  $T_c$  or below our value of  $T_t$ , have been omitted from the figure and from the statistics. The figure also illustrates two earlier correlations, by Goodwin et al.4 and by Younglove and Ely,<sup>10</sup> which were based on substantially identical data. The difference between the ancillary correlation and vapor pressures computed from the SWEOS using the Maxwell construction technique is shown to be well under 0.1% from the triple point to the critical temperature. The ancillary equations were used to generate 43 points at 5 K increments as input for the SWEOS correlation. Statistics concerning a comparison of the Maxwell construction and the ancillary equation are given in Table 11. In addition to the AAD-%, the BIAS (average deviation) and root-mean-square deviation (RMS) are given. All statistics in Table 11 are presented on both a percentage and dimensional basis.

A compilation of sources of experimental vapor pressure data, including very early measurements, appears in the monograph by Sychev *et al.*,<sup>5</sup> that study adopted the correlation of Goodwin *et al.*<sup>4</sup> Additional discussions of the data are given by Goodwin,<sup>4,62</sup> Eubank,<sup>67</sup> and the experimental papers we have cited. Upon considering the uncertainties of both the triple point and critical point pressures (Table 1) and the quality of both primary and other experimental data, we make the subjective judgment that either the ancillary Eq. (4) or the SWEOS will provide values of the saturation pressures accurate within 0.1% above T - 165 K, within 1% between 130 and 165 K, and within 5% for temperatures from the triple point to 130 K where the vapor pressure is less than 0.003 MPa.

For the density of the saturated liquid, Fig. 5 illustrates the deviations between the correlation of Eq. (5) and experimental data and other correlations. The primary data, comprising 66 points from Refs. 22 and 35-40, give an AAD-% of 0.03% with a maximum deviation of less than 0.1%. These data were obtained by a variety of techniques. Buoyancy forces were measured by Haynes and Hiza<sup>38</sup> using a magnetic suspension densimeter with a barium ferrite buoy and by McClune<sup>39</sup> and Orrit et al.<sup>40</sup> using silica buoys and mechanical suspensions. Calibrated pycnometer volumes were used by extrapolation of single-phase isotherms to the measured saturation pressure (or near the critical point, to a break in the observed isotherm) by Douslin and Harrison.<sup>22</sup> Coexisting phases in a glass pycnometer were observed directly by Sliwinski35 and by Canfield and collaborators, 36 and an expansion technique was used by Rodosevich and Miller.<sup>37</sup>

Although the primary data were chosen to ensure consistency, Fig. 5 illustrates that there are more substantial differences among the other data. For instance, in the



Fig. 4. Saturation pressures compared to Eq(4). References are abbreviated with first authors only. Primary data from: Douslin<sup>22</sup>; Pal<sup>31</sup>; Pope<sup>32</sup>; Ziegler<sup>33</sup>; Straty<sup>34</sup>. Secondary data from: Straty<sup>34</sup>; Djordjevich<sup>65</sup>; Pal<sup>31</sup>; Carruth<sup>63</sup>; Miniovich<sup>66</sup>; Ziegler<sup>33</sup>; Regnier<sup>64</sup>; Pope<sup>32</sup>. Points exceeding the range include: 5 from Djordjevich; 10 from Carruth; 39 from Ziegler; 9 from Regnier; 1 from Pal; and 1 from Pope. Curves are from SWEOS, Goodwin<sup>4</sup>, and Younglove<sup>10</sup>.

temperature range from 253 to 298 K, the densities from Douslin and Harrison,<sup>22</sup> not considered primary, deviate nearly up to 0.4% from the correlation and are below the measurements of Haynes and Hiza,38 taken as primary in the region 100 to 270 K. Other values, such as those reported by Pal et al.,<sup>31</sup> using a method similar to that of Douslin and Harrison, and by Goodwin et al.<sup>4</sup> (as ID 5 in Table 2 of Ref. 4) derived from Pal's isochores and a different vapor pressure equation, are also below our correlation and much closer to the densities of Douslin and Harrison near 250 K. However, Havnes' data are supported at his lower temperatures (near 100 K) by other primary data and by data from Klosek and McKinley,68 as shown, by Sliwinski<sup>35</sup> near 280 K, somewhat above the highest temperature of Havnes and Hiza, and by the measurement of Kahre<sup>69</sup> at 267 K, considered secondary. Thus, the data of Haynes and Hiza, from their magnetic suspension densimeter, are to be preferred in this range, although the estimate of the uncertainty in our correlation must be increased.

Closer to the critical point, the primary data of Douslin and Harrison,<sup>22</sup> whose critical parameters we have adopted, agree fairly well with measurements of Sliwinski<sup>35</sup> and, to a lesser degree, with the secondary results of Miniovich and Sorina<sup>66</sup> and Tomlinson.<sup>70</sup> The saturated liquid densities reported by Pestak et al.<sup>24c</sup> were based on evaluation of the Lorentz-Lorenz experimental data of Burton and Balzarini.<sup>24a</sup> The points shown in Fig. 5 are from the tabulation in Ref. 24c using the critical temperature and density from that reference to convert the reduced quantities. The deviation of 1.6% at 305.17 K, their point closest to the critical temperature, is not shown in Fig. 5. This deviation is reduced to -0.3%, and general agreement with our correlation is greatly improved, if our critical temperature of 305.33 K is used to convert the reported reduced temperature. This amounts to a shift of about 60 mK for all their absolute temperatures. It is not clear whether this shift is due to lack of calibration of the thermometers used in the experiment of Ref. 24a, or is because our critical parameters reflect a classical, mean-

Туре	No. pts.	Source	AAD-%	BIAS – %	RMS-%	AAD	BIAS	RMS	
(percent)									
Ρσ	43	Eq.(4) <sup>a</sup>	0.020	0.015	0.023	0.32	0.32	0.62	kPa
ρσι	43	Eq.(5) <sup>a</sup>	0.030	0.017	0.052	0.004	0.002	0.006	mol·dm <sup>-3</sup>
ρσν	43	Eq.(6) <sup>a</sup>	0.120	0.001	0.137	0.89	0.42	1.98	mol·m <sup>-3</sup>
Bb	35	[22,42-44]	0.626	0.179	1.06	0.001	- 0.001	0.003	dm³•mol <sup>−1</sup>
B°	87	[22,42-44,74-79]	1.849	-0.649	2.843	0.005	0.002	0.011	dm <sup>3</sup> ·mol <sup>-1</sup>
PVT <sup>b,d</sup>	803	Table 12	0.064	-0.014	0.135	0.006	-0.002	0.011	mol·dm <sup>-3</sup>
PVT <sup>c,d</sup>	2112	Table 12	0.483	-0.113	1.509	0.030	-0.006	0.094	mol·dm <sup>-3</sup>
PVT <sup>b,€</sup>	803	Table 12	0.591	0.157	1.456	0.094	0.014	0.205	MPa
PVT <sup>c,c</sup>	2112	Table 12	2.844	2.301	11.574	0.193	0.081	0.552	MPa
$C_{V}^{b}$	214	Table 13	1.486	- 1.158	2.271	0.852	-0.683	1.760	J·mol <sup>-1</sup> ·K <sup>-1</sup>
$C_{V}^{c}$	239	Table 13	1.600	-1.255	2.247	0.904	- 0.729	1.699	J-mol <sup>-1</sup> ·K <sup>-1</sup>
$C_{P}^{b}$	359	Table 13	1.088	0.246	1.545	2.036	0.464	4.776	J·mol <sup>-1</sup> ·K <sup>-1</sup>
$C_r^{c}$	511	Table 13	2.353	- 0.078	6.201	9.55	- 0.952	43.46	J-mol <sup>-1</sup> -K <sup>-1</sup>
$C_{\sigma L}{}^{b}$	106	[47]	0.844	0.314	1.231	0.711	0.088	1.082	J·mol <sup>-1</sup> ·K <sup>-1</sup>
$C_{\sigma L}^{c}$	186	Table 13	1.014	0.599	1.308	0.828	0.393	1.185	J·mol <sup>-1</sup> ·K <sup>-1</sup>
$W^{b}$	201	Table 14	0.553	-0.269	0.738	3.400	- 1.426	4.257	m•s <sup>-1</sup>
W°	248	Table 14	0.486	-0.205	0.686	2.938	-1.123	3.922	m-s <sup>-1</sup>
$W_{\sigma L}{}^{b}$	47	[50]	0.493	0.366	0.634	3.424	1.434	4.265	m-s <sup>-1</sup>
Walc	120	Table 14	0.486	0.400	0.594	4.328	3.046	4.854	m·s <sup>-1</sup>

TABLE 11. Statistics for thermodynamic property data versus SWEOS correlation

\* These data were generated from the ancillary equations at equally spaced temperatures from 91 to 301 K.

<sup>b</sup> Primary data.

° Primary and secondary data.

<sup>d</sup> Statistics based on calculation of density from experimental temperature and pressure.

\* Statistics based on calculation of pressure from experimental temperature and density.

field bias [although we use an effective  $\beta$  of 0.355 in Eq. (5)] whereas the study of Ref. 24b, based on a more rigorous revised and extended scaling theory (with  $\beta = 0.327$  and two Wegner correction terms with  $\Delta = 0.5$  from the theoretically motivated mixing of the thermodynamic fields) does give the physical critical parameters.

Other comparisons among the available data are given in the cited experimental papers, and Sychev et al.<sup>5</sup> give an excellent bibliography of additional, especially earlier, sources of data on the density of saturated liquid ethane. For the 202 points considered, the AAD-% is 0.21%. Eight of these points have deviations greater than 1% as shown in Fig. 5; these comprise a single point from Miniovich and Sorina<sup>66</sup> at 305.26 K and near 1%, and 7 points from Pestak et al.<sup>24b</sup> in the range 305.08 to 305.17 K with deviations to 1.6%. We have also shown, in Fig. 5, earlier correlations from Goodwin et al.<sup>4</sup> and from Younglove and Ely,<sup>10</sup> as well as a comparison with saturated liquid densities calculated from the SWEOS. For the 43 points generated from the ancillary equation as input into the SWEOS fitting program, the deviations are all under 0.17%, and the AAD-% is 0.03%; additional statistics are in Table 11. The agreement between the SWEOS points from the Maxwell construction and the ancillary Eq. (5) is thus quite good as is agreement between the correlation of Goodwin et al., which used much the same primary data in its development, and the present ancillary equation. The larger deviations from the more recent correlation of Younglove and Ely arise partly from the different choice of critical parameters in that study.

At temperatures very close to the critical point, the oscillations of the correlations shown in Fig. 5 (and Fig. 6)

illustrate how difficult it is to produce the shape of the coexistence dome. The ancillary equation, as indicated in Sec. 2, expresses the theoretically described behavior, with a critical exponent of 0.355. The SWEOS, while approximating this flattened coexistence dome quite well, approaches the critical point with the mean field exponent of 0.5, and thus cannot accurately describe this region. Of course, measurements too are difficult and inaccurate in the asymptotically critical region. The SWEOS is constrained to give the reported critical density at the critical temperature. We conclude that either the ancillary equation or the value of the density from the SWEOS can be used with an estimated uncertainty of 0.1% from the triple point to 200 K, 0.3% from 200 K to 304 K, and deteriorating to about 1.5% to the critical point at 305.33 K.

Figure 6 shows the analogous deviation plot for the density of the saturated vapor. The primary data are from Refs. 4, 22, and 35; these 32 points have an AAD-% of 0.07% with a maximum deviation of 0.26% near the critical temperature. This excellent agreement is somewhat misleading; the accuracy of the data is not nearly this good. The primary data from Goodwin et al.4 were based on their virial equation of state and their vapor pressure equation, and thus, as indicated above, they are not direct measurements. These provide the sole source of primary data with temperatures from the triple point to about 250 K. Experimental data published in 1926 by Porter<sup>72</sup> and based on the intersection of measured isochores with an experimentally determined vapor pressure line, are also shown in the figure; there is a discrepancy of more than 1% at the lowest measured temperature, near



Fig. 5. Saturated liquid densities compared to Eq. (5). Primary data from: Douslin<sup>22</sup>; Sliwinski<sup>35</sup>; Chui<sup>36</sup>; Rodosevich<sup>37</sup>; Haynes<sup>38</sup>; McClune<sup>39</sup>; Orrit<sup>40</sup>. Secondary data from: Douslin<sup>22</sup>; Pal<sup>31</sup>; Sliwinski<sup>35</sup>; Orrit<sup>40</sup>; Kahre<sup>69</sup>; Miniovich<sup>66</sup>; Pestak<sup>24c</sup>; Goodwin<sup>4</sup>; Tomlinson<sup>87</sup>; Klosek<sup>68</sup>. Curves are as in Fig. 4.

184 K. The primary data of Douslin and Harrison<sup>22</sup> extend from 248 to 304 K. The data were obtained in the single phase in a pycnometer and were extrapolated to the intersection with the vapor pressure curve or to the break of the isochore when closer to the critical temperature. Some data from Sliwinski,<sup>35</sup> who also provided saturated liquid densities, were included in the primary data set and are shown to agree well with the correlation; the point at 305.15 K, his measurement closest to our critical temperature, however, deviates by 1% from our correlation and was not included in the primary data set.

The critical region results of Miniovich and Sorina,<sup>66</sup> considered secondary, show systematic deviations of up to 4.6% (at 305.325, not shown); their critical temperature of 305.35 differs slightly from ours, so the discrepancy in saturated vapor densities is not surprising. The densities of Pestak *et al.*<sup>24c</sup> were again reported versus reduced temperature; the illustrated deviations are based on the critical parameters in Ref. 24c and show a maximum deviation of -1.5%. Upon changing the critical temperature to our value, all deviations become positive and less than 1%, although the improvement is not

nearly as dramatic as for the saturated liquid densities. (Reference constants of 305.31 K and 6.86 mol·dm<sup>-3</sup> applied to the tabulated coexistence densities in Ref. 24c result in all deviations from the current correlation being well under 0.5%.)

The 102 data indicated in Fig. 6, including a single point from Ref. 66 which has a large deviation and which is not illustrated, have an AAD-% of 0.43%. Our correlation for the saturated vapor density agrees quite well with the ancillary equation from Goodwin et al.<sup>4</sup> with maximum deviations of about 0.25% from the triple point temperature to 303 K. The correlation from Younglove and Ely<sup>10</sup> systematically differs from our equation and exhibits a maximum deviation of 1.1%; although that correlation was based on substantially similar primary data, a shift in their selection of critical temperature and density induced the illustrated discrepancy. The saturated vapor densities calculated from the equation of state of Sychev et al.<sup>5</sup> (not shown in Fig. 6) show deviations of less than 0.5% below about 300 K, but these increase to more than 5% in the critical region; they used values from Refs. 4 and 22 as primary. Saturated vapor densities calculated



Fig. 6. Saturated vapor densities compared to Eq. (6). Data from: Goodwin<sup>4</sup> (Virial Equation); Douslin<sup>22</sup>; Sliwinski<sup>35</sup>; Miniovich<sup>66</sup>; Porter<sup>72</sup>; Pestak<sup>24b</sup>. One point from Miniovich is out of range. Curves are as in Fig. 4.

from the SWEOS agree with those calculated from Eq. (6) to within 0.25% for temperatures from the triple point to 300 K; closer to the critical temperature, a 1% deviation is seen. For the 43 equally spaced points generated from Eq. (6) as input into the SWEOS fitting program, the maximum deviation is 0.24% and the AAD-% is 0.12%.

At temperatures below 250 K, the density of the saturated vapor is less than 1 mol·dm<sup>-3</sup> and direct measurements are sparse. In this region, it is the uncertainty in the vapor pressure curve, and to a lesser degree, the uncertainty in the second virial coefficients, which can produce errors in a correlation for the saturated vapor density. Near the critical point, the flatness of the saturation envelope and the concomitant difficulty in making accurate measurements, can lead to uncertainty. As in the case of the saturated liquid, the SWEOS, and any analytic equation of state, incorporates an exponent of 0.5 to describe the curve near the critical point; the value should be closer to 0.3. Our analysis indicates that either the ancillary equation or the SWEOS will generate values for the density of the saturated vapor with uncertainties

as follows: from  $T_t$  to 130 K, 5%; from 130 to 240 K, 1.5%; from 240 to 303 K, 0.3%; and from 303 to  $T_c$ , 5%.

## 4.2. Ideal Gas Properties

The primary data for the correlation of ideal gas properties, Eq. (3), comprised the 11 points for  $C_P^{id}$  from Chao et al.<sup>11</sup> as well as the fixed points for the ideal gas entropy and enthalpy as given in Table 1 and discussed above: as seen in Table 7,  $C_P$  is related to  $\tau^2 \phi_{\tau\tau}$  and integration constants determined by S (proportional to  $\phi^{id}$  –  $\tau \phi_{\tau}$ ) and H are needed. The primary data were calculated from the spectroscopic model and spanned the temperature range 50 to 700 K; the AAD-% was 0.06% and the maximum deviation was 0.2% at 50 K (well below the triple point temperature). Figure 7 shows the deviations for the ideal gas properties. The deviations for the ideal gas enthalpy and entropy as calculated by Chao et al. in the same temperature range are also shown. The maximum deviation in the enthalpy is 0.17%, also at 50 K, and in the entropy, it is 0.02%.



Fig. 7. Ideal gas properties compared to Eq. (3). Data are from: Chao<sup>11</sup>; Pamidimukkala<sup>30</sup>; Bier<sup>49,49a</sup>; TRC Thermodynamic Tables<sup>30a</sup>. The curves are from Goodwin<sup>4</sup>; Buhner<sup>73</sup>; Younglove<sup>10</sup>; and Sychev<sup>5</sup>.

Pamidimukkala et al.<sup>30</sup> provide an alternative set of ideal gas properties. These were based on the same spectroscopic model, a rigid-rotor harmonic oscillator with contributions from the hindered internal rotation, using slightly different spectroscopic constants. As illustrated in Fig. 7, the results are quite similar to those of Chao et al.<sup>11</sup> in the region of overlap. The largest deviations are 0.3% for the isobaric heat capacity at 200 K, 0.3% for the enthalpy at 200 and 300 K, and 0.06% for the entropy at 100 K. Reference 30 also provides estimates of the uncertainties in their results. For  $C_P^{id}$ , the estimate is about 0.2%, for  $H^{id}$ , it is 0.4%, and for  $S^{id}$  it is 0.1%, all near 300 K. The TRC Thermodynamic Tables<sup>30a</sup> also provide a tabulation for ideal gas properties. Although these tables were revised 10/31/85, the latest source reference for ethane is dated 1974. As can be seen in Fig. 7, our correlation agrees with the TRC tabulation quite well. The maximum deviation in entropy in 0.02% (at 100 K), and deviations in the ideal gas isobaric heat capacity and enthalpy are less than about 0.2% at all temperatures between 50 and 700 K.

Bier *et al*.<sup>49,49a</sup> used a flow calorimeter to determine  $C_P$  at low pressures (from about 0.1 to 1 MPa) and have ex-

trapolated to zero pressure to obtain ideal gas values. Their data range in temperature from 283 to 473 K and they estimate their uncertainty in the ideal gas heat capacity as about 0.2%. The discrepancy between these thermodynamic values and the spectroscopically derived values had been noted by Bier *et al.*<sup>49a</sup> and, as seen in Fig. 7, reach 1.4% at 373 K.

While this disagreement has not been satisfactorily resolved, we have chosen to retain the spectroscopically derived data to develop our ideal gas correlation and maintain consistency with other correlations. As shown in the figure, correlations from Goodwin *et al.*,<sup>4</sup> Younglove and Ely,<sup>10</sup> and Sychev *et al.*<sup>5</sup> agree quite well with our own. They were based on the same primary data. Buhner *et al.*<sup>73</sup> used primary data from both Ref. 11 and 49. We estimate the uncertainty in the ideal gas correlation as follows: for the enthalpy, 1.5%; for the isobaric heat capacity, 1.5%; and for the entropy, 1.0%; all from the triple point temperature to 700 K. If the measurements of Ref. 49 are replicated or new spectroscopic models or data become available, development of a new ideal gas correlation would be warranted.

## 4.3. Thermodynamic Properties from the SWEOS

In this section we discuss the comparisons between experimental measurements and the correlations for several of the thermodynamic properties calculable from the Helmholtz energy using the algebraic expressions in Table 7. In the first subsection, 4.3.1, we discuss the comparisons for the second virial coefficient. In Sec. 4.3.2, comparisons for the extensive PVT data are given, and following sections discuss the heat capacities and sound speed data.

#### 4.3.1. Second virial coefficient data

The deviation plot for second virial coefficient data is shown in Fig. 8. The 35 primary data from Refs. 22 and 42-44 have an AAD-% of 0.6% and an AAD of 0.001  $dm^3 \cdot mol^{-1}$ ; additional statistics are included in Table 11. Douslin and Harrison<sup>22</sup> obtained second virial coefficients from fits to a truncated virial equation of state using their unsmoothed low density data. The *PVT* data were obtained with a thin-walled pycnometer inside a compressibility bomb. The largest deviation from our SWEOS is about 1% at 623 K, the highest temperature, and this corresponds to a deviation of 0.0002 dm<sup>3</sup>·mol<sup>-1</sup>; deviations of less than 0.2% are found in the range 273 to 500 K. Michels et al.<sup>42</sup> used a glass piezometer, and their data for densities less than 2.25 dm<sup>3</sup>·mol<sup>-1</sup> were fitted by a virial series truncated at the third virial coefficient; their temperatures were adjusted to conform to IPTS-68, but these adjustments from 8 to 20 mK correspond to changes of less than 0.01% in the second virial coefficients. The largest deviation for these data is about 0.5% at 273 K. The data of Pope et al.43 are based on an isochorically coupled Burnett method and extend downward from 306 to 210 K, our lowest temperature for primary second virial coefficients. At 210 K, a deviation of nearly 4% is seen; this is contrasted to the accuracy of about 2.4% for this point reported in Ref. 43. Mansoorian et al.<sup>44</sup> also use an isochoric-Burnett method and their data in the range 323-473 K show a maximum deviation of 2.4% (compared to their estimated error of 1.5%). Better agreement among alternative data in this region serve to discount the importance of the outliers from Ref. 44.



Fig. 8. Deviations for second virial coefficients versus temperature. Data are from: Douslin<sup>22</sup>; Pope<sup>43</sup>; Mansoorian<sup>44</sup>; Michels<sup>42</sup>; Eucken<sup>74</sup>; Hoover<sup>76</sup>; Gunn<sup>77</sup>; Dymond<sup>75</sup>; Strein<sup>79</sup>. Curves are from Boushehri<sup>27</sup> and Goodwin<sup>4</sup>. Two low temperature points from Eucken are out of range.

Among the secondary data, we have illustrated the 1933 results of Eucken and Parts,<sup>74</sup> which, though often discredited,75 represent the most extensive set of low temperature measurements and include the point at the lowest reported temperature, 192 K with a deviation of 8%. The data from Hoover et al.76 extend down to 215 K and are seen to lie considerably above our correlation and the primary data of Pope et al.<sup>43</sup> Data from Gunn,<sup>77</sup> reported by Huff and Reed,<sup>78</sup> and from Strein et al.<sup>79</sup> again indicate a spread in the measured values. Finally, we have illustrated the values selected by Dymond and Smith<sup>75</sup> from their analysis of these and other experimental data as recorded in Ref. 75. The deviation of 2.6% for their lowest temperature can be compared to their estimate of 2.4% of the uncertainty in that point. References 75 and 5 provide additional citations to experimental studies of the virial coefficients of ethane.

There are many calculations of the virial coefficients based on either experimental measurements or direct calculation from a model intermolecular potential whose parameters may be based on experiment. We have illustrated the correlation of Goodwin et al.<sup>4</sup> who used much of the same data as primary for an ancillary correlation of the second virial coefficient. At low temperatures, however, Goodwin et al. based their correlation on data attributed to McGlashan and Potter<sup>45</sup> after an adjustment for better agreement at higher temperatures. Reference 45, in turn, had constructed an equation for the second virial coefficients of the alkanes based on two- parameter corresponding states and measurements above room temperature on the higher alkanes; the critical density used for ethane in the correlation differs by 2% from the value which we have selected. Thus, we do not consider the difference of more than 6% between our correlation and the actual data from Ref. 45 at 150 K (not shown) either surprising or disturbing. Further, at 150 K the vapor pressure is about 0.01 MPa and the density of the saturated vapor is only about 0.008 mol·dm<sup>-3</sup>: the ideal gas equation alone will give this density within about 0.4%. The correlation of Boushehri et al.,<sup>27</sup> also shown in Fig. 8, is based on a corresponding states model including corrections and contributions from the nonsphericity of the ethane molecule; the minimum temperature considered in Ref. 27 is 250 K.

For the 87 data indicated in Fig. 8, the AAD-% is 1.85%. Additional quantification of the comparison is given in Table 11. Our assessment of the data and correlation leads to an estimate of the uncertainty in second virial coefficients calculated from the SWEOS as follows. At temperatures from 150 to 200 K, there is an uncertainty of 10%; from 200 to 250 K, this decreases to 3%; from 250–500 K, 1%; and from 500 to 600 K, the maximum temperature considered, the estimated uncertainty is 2%. The SWEOS correlation cannot be used to estimate values of the second virial coefficient at temperatures below 150 K.

#### 4.3.2. PVT Data

For the *PVT* data, there are two types of evaluations which can be informative. One can calculate the pressure from the SWEOS directly from the experimental temperature and density, or one can use a root-finding technique to calculate the density using the experimental temperature and pressure. To implement the second type of comparison, we used a standard Newton-Raphson algorithm.

			Temperature	Pressure	range	Pressure	Density
First author	Ref.	No. pts.	range, K	range, MPa	mol·dm <sup>-3</sup>	AAD-%*	AAD-%⁵
Beattie	82	96	298-523	1.1-20	0.5–5	0.43	0.49
Besserer	82a	68	311-394	0.7-10	0.2-12	0.67	0.83
Douslin <sup>e</sup>	22	257	273-623	1.4-40	1. – 11	0.07	0.06
Douslin	22	449	243-623	0.1-41	1. – 16	1.10	0.24
Golovskiy	83	111	90270	1.2-60	16-22	8.88	0.10
Khazanova	85	86	299-318	0.5-7	0.2-11	2.23	3.02
Law	84	56	240-350	1.1-34	2-17	2.03	0.49
Michels <sup>e</sup>	42	81	273-423	1.6-10	0.8-3.6	0.03	0.04
Michels	42	101	273-423	1.6-22	0.8-8.2	0.03	0.05
Miniovich	86	63	303-307	4.6-4.9	4.5-9.3	0.03	1.41
Pal <sup>c</sup>	31	45	183-320	36-69	15-19	0.96	0.09
Pal	31	309	157-344	0.3-69	1-19	8.38	0.78
Parrish <sup>c</sup>	46	9	300-322	5.5-9.7	4-12	0.32	0.23
Reamer	86a	176	311-511	1.4-55	0.3-16	1.13	0.50
Sengers <sup>c,d</sup>	23	30	304-325	4.7-8.1	4.9–9	0.03	0.30
Straty <sup>c</sup>	34	381	96-320	3-38	1.9-22	1.07	0.04
Straty	34	477	93-320	0.2-38	1.2-22	2.73	0.10
Tomlinson	87	61	280-325	4.4-14	4-14	0.48	0.51
Wallace	88	20	248-348	0.1-0.2	0.03-0.09	0.06	0.06

TABLE 12. Sources of PVT data

<sup>a</sup>Average absolute percent deviation when pressure is calculated from experimental temperature and density.

<sup>b</sup>Average absolute percent deviation when density is calculated from experimental temperature and pressure.

<sup>c</sup>Only primary data are included in these statistics.

<sup>d</sup>These data were obtained from a scaled equation of state; they are not experimental data.

A third possibility, using pressure and density as independent variables, is not often required and is not considered here. For each of 15 references, Table 12 summarizes the data and certain statistics regarding the fit to the present SWEOS correlation. The pressure AAD-% gives the average absolute percent deviation when pressure is calculated from the SWEOS. The last column in Table 12, the density AAD-%, gives the analogous statistic when the density is calculated. The primary data, used to develop the equation of state, are indicated in the table. References 22, 31, 34, and 42 are each listed twice in the table; the first line refers to data selected as primary and the second line includes the entirety of the data as given in the references.

In Table 11, we present several overall statistics for the 803 primary data summarized in Table 12 and for the entire set of 2112 points. In addition to the average absolute deviation, we present the average deviation (or BIAS), and the root-mean-square deviation (RMS). Each of these three statistics is given on both a pecentage and dimensional basis. Of course, the quality of the fit and the quality of the data depend strongly on the region of the phase diagram being considered. For this reason deviation plots are useful and we have several comments on Figs. 9–12 which cover different temperature ranges for the primary data; deviations for other data are plotted in Figs. 13–14.

The primary data were taken from Refs. 22, 23, 31, 34, 42 and 46 and included temperatures from 96 to 628 K with pressures to 69 MPa. For these data the AAD-% is 0.59% when calculating pressures and is 0.06% when calculating densities. Douslin and Harrison<sup>22</sup> used their thinwalled pycnometer in a compressibility bomb to measure the PVT surface of a highly pure sample with an uncertainty in the compressibility factor ranging from 0.03% at low temperatures to 0.3% at the highest temperatures; this source also provided saturation data and virial coefficients, as discussed above. Data from Ref. 22 with densities above 10.5 mol·dm<sup>-3</sup> were excluded from the primary data set in favor of the measurements from Refs. 31 and 34. Also, points tabulated only in Table 1 of Ref. 22, in the coexistence region, were not considered primary; thus we excluded the data very near the critical point, including subcritical isotherms at 305.15 and 305.25 K and supercritical isotherms at 305.35, 305.37, and 305.39 K. From the deviation plots shown and the statistics given in Table 12, the primary data from Ref. 22 are seen to conform quite well with our SWEOS. The maximum deviation is 0.4% when calculating the pressure near 323 K and 10.5 mol·dm<sup>-3</sup>; the primary data from Ref. 34 show a smaller deviation in this region. When calculating densities, the deviations are all under 0.3%. The isotherm at 623 K shows the largest deviations and the maximum devation is at the highest measured density  $(7 \text{ mol} \cdot \text{dm}^{-3})$  at 623 K. Reference 22 provided all of the primary PVT data with temperatures above 425 K.

For the secondary data from Ref. 22, the deviations are in general much larger. When calculating the pressure for the liquid near the coexistence line at low temperatures,

we see deviations above 60%. The steepness of the isotherms is of primary importance when calculating the system pressure from the temperature and density. In the compressed liquid at low temperatures the slope,  $\partial P/\partial \rho|_T$ , is extremely large, so that small (experimental) uncertainties in the density can lead to very large errors in the calculated value of the pressure. For instance, at 100 K, the slope of about 77 MPa·mol<sup>-1</sup>·dm<sup>3</sup> near the two-phase boundary means that an error of 0.1% in the density, typical of many experimental measurements, leads to an error of almost 500% in the calculated value of pressure. At 243 K, the lowest temperature measured by Douslin and Harrison,<sup>22</sup> the isotherms are not this steep; here the 60%discrepancy in pressure corresponds to less than 0.5% in density. Thus, the large percentage deviations are not suprising. In addition, the SWEOS shows better agreement with secondary data from Ref. 34 in this region, and the absolute deviations in pressure are less than 1 MPa.

When calculating densities from the experimental pressure and temperature, the maximum deviation for the data of Ref. 22 is 4.2% and occurs near the critical point. Here the difficulty is caused by the flattness of the isotherms; we calculate a slope  $(\partial P/\partial \rho|_T)^{-1}$  of about 1900 mol·MPa<sup>-1</sup>·dm<sup>-3</sup> at the critical density and 305.37 K. The 4.2% deviation in density corresponds to a deviation of 0.004% in pressure. The experimental difficulties at nearcritical conditions, for example, the importance of gravitationally induced density gradients, the long time necessary for equilibration of an experimental cell, and the difficulty of accurate temperature measurement, are also well known. We have used the scaled equation of state<sup>23</sup> to calculate a PVT point corresponding to the experimental temperature and pressure from Ref. 22 with the largest deviation from the SWEOS. This calculated datum has a deviation of 0.4% from the density calculated from the SWEOS.

Pal et al.<sup>31</sup> used a stainless steel isochoric chamber, calibrated at low pressures by Burnett expansions, and used differential weighing to determine the amount of fluid in the cell. The accuracy was estimated to be 0.2% except near the critical point; this source also provided saturated fluid properties and was mentioned above. All of our primary PVT data with pressures greater than 40 MPa were taken from Ref. 31. Among the primary data, the maximum deviation in the calculated pressure is near 3% and occurs near 252 K and 42 MPa (corresponding to 17  $mol \cdot dm^{-3}$ ). The same point exhibits a large density deviation, but this is less than 0.3%. This again illustrates how the steep isotherms are associated with large uncertainties in the determination of pressure. For these primary data, an average deviation of less than 0.1% in density, less than the claimed accuracy, corresponds to 1% average deviation in pressure. The overlapping primary data from Ref. 34 show even better agreement with the SWEOS calculations.

Points from Ref. 31 with pressures below 35 MPa were not considered primary data and some show extreme deviations from the SWEOS. The problems are the same as discussed above. A deviation of more than 200% (or 0.8



Fig. 9a. Pressure deviations versus experimental pressures of primary data for temperatures at or below 200 K. Data are from Straty<sup>34</sup> and Pal<sup>31</sup>.

MPa) seen in the calculation of pressure in the compressed liquid at 177 K corresponds to a difference of 0.13% in the density; this is within the experimental uncertainty quoted by Pal *et al*.<sup>31</sup> Also along some supercritical isotherms, density predictions near the critical point differ greatly from experimental values. Further, many of these points showing extreme deviations were not considered in Refs. 4 and 5, and some of the data which appeared in the thesis<sup>32</sup> did not appear in the smoothed tables of Ref. [31]. For these reasons, the large deviations shown in our figures are not a cause of much concern.

Straty and Tsumura<sup>34</sup> used a cell of nearly constant (and calibrated) volume to measure *P*-*T* points along quasi-isochores. The amount of sample was determined by expansion at low pressure into large calibrated volumes. Their estimated experimental uncertainty in density ranged from 0.1% at low temperatures, to 0.2% at higher temperatures and lower densities, to 1% in the critical region. This source provided all of the primary *PVT* data below 183 K and most of the low pressure data below about 275 K. The primary data from Ref. 34 are again reproduced quite well by the SWEOS as indicated by the statistics in Table 12. Again, the compressed liquid at low temperatures and pressures (near the saturation boundary with densities greater than about 20 mol·dm<sup>-3</sup>) show large deviations in the calculated value of pressure. The largest deviation, nearly 18%, corresponds to a density variation of 0.02%; the correlation of Ref. 4 indicates a deviation of about 15% for this point. For the density calculation, a point at 304 K and 9 mol·dm<sup>-3</sup>, that is, quite near the critical point, exhibits a 0.34% deviation. Data from Ref. 34 at densities below 11 mol·dm<sup>-3</sup> were not considered in the correlation of Goodwin *et al.*<sup>4</sup>

Some points from Ref. 34 near the saturated liquid boundary, usually the last two points along the measured isochore, were not included in the primary data. In addition to the correlational problems described above and caused by the steepness of the isotherms, there are experimental problems in this region. Although the dead volumes in the experimental procedure of Straty and Tsumura<sup>34</sup> were made very small, measured, and accounted for, when measurements are made close to the saturation boundary, a meniscus can be formed in these experimental volumes, and the correction will become less accurate. However, for the data with densities above 20 mol·dm<sup>-3</sup>, all density deviations are within 0.1%; the pressure deviations range up to 40% (1.5 MPa) for these data. The measured isochore near 6.5 mol·dm<sup>-3</sup> and low


Fig. 9b. Density deviations versus experimental density. References and temperature range are as in Fig. 9a.

densities near 1.2 and 4.3 mol·dm<sup>-3</sup> (restricted to temperatures above 270 K) were also considered secondary. The critical region suffers the same problem in the density calculation as discussed above, and for the low densities we have preferred the data from Douslin and Harrison.<sup>22</sup>

The measurements from Michels et al.<sup>42</sup> were published in 1954 and are the oldest set of primary data which we have used. The experiment was performed at the van der Waals Laboratory using a calibrated glass piezometer and expansions to determine the amount of material. Temperatures were adjusted to IPTS-68 according to the table of Powell et al.,80 but the adjustments were very small in this temperature range. The data substantially overlap those of Ref. 22 and were given slightly less relative weight in the fit; agreement with the SWEOS is excellent. The isotherm at 273.15 K provides the most extreme deviations in both pressure and density calculations; the largest deviations are less than 0.2% in pressure and 0.3% in density. The corresponding isotherm from Ref. 22 agrees even better with the SWEOS; the deviation in pressure is about 0.03% and in density is about 0.05%. Secondary data from Ref. 42, that is those points with densities above 4 mol $\cdot$ dm<sup>-3</sup>, also agree quite well with the equation.

Parrish<sup>46</sup> recently used a vibrating tube densimeter in a study of mixtures and also published some data on pure ethane. The apparatus was calibrated with argon and propane using modified BWR equations of state. An uncertainty in ethane density of 0.2%, based on comparison with a modified BWR equation, is implied; however, an outlier with deviation of about 1.25% is noted in Ref. 46. These data overlap those of Ref. 22 and were given slightly less weight. The deviations from our SWEOS are somewhat larger than those for other primary data in this region. The point referred to above shows a deviation in density of slightly less than 1% from our SWEOS; a deviation in pressure of 0.5% is seen for the point at highest temperature and pressure.

The final source of primary *PVT* data was the scaled equation of state due to Sengers.<sup>23</sup> The parametric equation (with critical exponents  $\beta = 0.325$ ,  $\gamma = 1.24$ ,  $\nu = 0.63$ ,  $\Delta = 0.5$ ,  $\xi_0 = 19$  nm, and the same critical point as that adopted here) has been described by Sengers and Levelt Sengers<sup>81</sup> for other fluids. The formalism incorporates corrections to asymptotic scaling (Wagner corrections) and accounts for the mixing of the thermodynamic fields within the linear model for the asymptotic behavior of the potential; many of the fluid-dependent parameters



Fig. 10a. Pressure deviations versus experimental pressures of primary data for temperatures between 200 and 280 K. Data are from: Pal<sup>31</sup>; Straty<sup>34</sup>; Douslin<sup>22</sup>; Michels<sup>42</sup>.

in the equation were derived from the experimental data of Douslin and Harrison.<sup>22</sup> The range of application is considered 300-350 K with densities in the range 4.8-9.2 mol·dm<sup>-3</sup>; however, the accuracy of thermodynamic quantities calculated from the equation is difficult to assess. Because the classical SWEOS can reasonably mimic a scaled equation of state in the general critical region<sup>6</sup> (although not in the asymptotically critical region), at least to the extent of reproducing the shape of the coexistence dome, we have used data generated from the scaled equation to exploit this capability. The SWEOS is capable of reproducing the pressures evaluated from the scaled equation; the deviations are generally less than 0.05% with a single outlier at 0.13% at the edge of the generated data. When using the SWEOS to calculate densities, we see deviations of 1 and 2% for primary data closest to the critical point. The flatness of the supercritical isochores near the critical point is reflected in these results. We have studied secondary data, not illustrated in the figures or included in the tables, and find that even at temperatures as close to critical as 305.331 K, density deviations of calculated values from the SWEOS and the scaled equation of state are less than 2.5%.

There are many other measurements of the PVT surface of fluid ethane. We have illustrated the deviations for some of these data in Figs. 13-14 and have included statistics in Tables 11 and 12. Our selection of data is not meant to be exhaustive: citations to earlier data are given in Ref. 5 and some of these data are included in Ref. 3. Data from Refs. 22, 31, 34, and 42 which were not included among the primary data are shown in these figures and are included in the statistics for the secondary data. From all these sources of data, only points reported at temperatures below our triple point temperature, inside the two-phase boundary constructed with our ancillary equations, or otherwise incompatible with our selection of critical parameters and phase boundaries have been excluded from consideration. In a few cases, obvious misprints, including transposition of digits, have been corrected. Outliers which are inconsistent with the scale of the figures are indicated in the sub-plots and are nonetheless included in the statistical summaries. We will only briefly summarize our observations concerning the secondary data.

Generally, the problems seen when comparing the SWEOS with the secondary data are similar to those dis-



Fig. 10b. Density deviations versus experimental density. References and temperature range are as in Fig. 10a.

cussed above: pressure calculations for the liquid at low temperatures and density calculations near the critical point. The extreme deviation in pressure for the data of Golovskiy,<sup>83</sup> for example exceeds 100%. This occurs near 92 K at a density near 22 mol·dm<sup>-3</sup>; for almost all densities above 19 mol·dm<sup>-3</sup>, the pressure deviations from Ref. 83 exceed several percent. Generally the lowest pressure data point for each isochore of Ref. 83 (the point closest to the liquid saturation boundary) shows large deviations. As is typical of the primary liquid data, however, the density deviations from Ref. 83 are quite good. Besserer and Robinson,<sup>82a</sup> who used a refractive index technique and calculated densities from the refractivity virials of Sliwinski,<sup>35</sup> have pressure deviations approaching 3% in the supercritical fluid; the corresponding deviations for the primary data of Douslin and Harrison<sup>22</sup> are much less. Data from Khazanova and Sominskaya,85 who used a piezometer of height 2 cm for critical region measurments and observed a critical point at 305.34 K, 4.88  $\pm$ 0.01 MPa, and 6.78  $\pm$  0.03 mol·dm<sup>-3</sup>, also show anomalously large pressure deviations throughout the measurements. The large pressure deviations in the data from Law<sup>84</sup> are greatest in his 240 K and 260 K isotherms.

Deviations of secondary experimental densities from

J. Phys. Chem. Ref. Data, Vol. 20, No. 2, 1991

those calculated from the SWEOS are dominated by the critical region measurements of Khazanova and Sominskaya<sup>85</sup> and of Miniovich and Sorina.<sup>86</sup> Reference 86 indicated a critical point at 305.35 K,  $4.8749 \pm 0.005$  MPa, and 6.793  $\pm$  0.01 mol·dm<sup>-3</sup>. Of the 80 points reported in Ref. 86 to lie in the single-phase region, 17 were in the two-phase region as defined by our ancillary equations and were excluded from consideration. The 104 points from Ref. 86 which were reported as on or within the saturation boundary were also not included even though some of these fall in the single-phase region as defined by our correlations and other experimental data. The large density deviations for Ref. 86, between 6 and 16%, occur for supercritical temperatures between 305.345 and 305.38 K at pressures within about 0.1% of our value of the critical pressure. Subcritical data, in both the liquid and vapor phases, have much smaller deviations, typically less than 0.3%. The density deviations from the data of Khazanova and Sominskaya<sup>85</sup> are suprisingly large even well away from the critical point. In particular, many of the measurements along the isochores near 0.2, 0.5, and 0.9 mol·dm<sup>-3</sup> have deviations between 8 and 10%; the pressure deviations for these data are also large. Much better agreement with the primary data of Douslin and



Fig. 11a. Pressure deviations versus experimental pressures of primary data for temperatures between 280 and 315 K. Data are from: Pal<sup>31</sup>; Straty<sup>34</sup>; Douslin<sup>22</sup>; Michels<sup>42</sup>; Parrish<sup>46</sup>; Sengers<sup>23</sup>.

Harrison<sup>22</sup> supports our SWEOS in this region. Sychev et al.<sup>5</sup> report a better statistical agreement with data from Khazanova and Sominskaya;<sup>85</sup> however, only 66 points of the 87 tabulated in Ref. 85 are indicated in their table. (We found that one point from Ref. 85 lies inside the two-phase region.) Further, the equation of state from Ref. 5 agrees very well with our SWEOS in this region. Data from Besserer and Robinson<sup>82a</sup> agree well with the correlation at the lowest pressures; along the 311 K and 372 K isotherms these data deviate from the SWEOS by nearly 5%. Other data in this range agree with the SWEOS.

Our examination of both primary and other experimental data, including the experimental techinque, reported uncertainty of the data, purity of the fluid ethane, and deviations from the SWEOS and other equations of state, have led us to these guidelines concerning the uncertainty of a *PVT* surface generated from the SWEOS. For the vapor at temperatures below 250 K, where the vapor pressure is about 1.3 MPa and the vapor density is about 0.8 mol·dm<sup>-3</sup>, calculations of both density and pressure from the SWEOS have uncertainties of about 0.1%. Liquid densities in this temperature range also have uncertainties of 0.1%; the calculation of pressure from a liquid density at these low temperatures is much more difficult. For pressures less than 10 MPa, the percent deviation in pressure may be very large, and it increases near the saturation boundary. Typically, the absolute deviation is within about 1 MPa. For higher pressures, the uncertainty ranges from 5% at temperatures below 200 K down to 1% for temperatures from 200 to 250 K.

Temperatures from 250 K to 300 K are still far enough from the critical point that accurate density calculations from the SWEOS are possible. For the vapor in this range, we estimate an uncertainty of 0.2% for both density and pressure calculations. For the liquid state near the saturation boundary at the lower temperatures of this range, to 265 K, large percentage deviations are possible. For pressures greater than 5 MPa or temperatures from 265 to 300 K, we estimate the uncertainty in the pressure calculation as 3%. Critical phenomena evidently do not lead to problems for the SWEOS for calculations at subcritical temperatures below 305.3 K: at 305.3 K, the reduced temperature,  $(T_c - T)/T_c$ , is about 10<sup>-4</sup> but the two-phase region excludes densities between about 6.4



Fig. 11b. Density deviations versus experimental density. References and temperature range are as in Fig. 11a.

and 7.4 mol·dm<sup>-3</sup>, that is within about 7% of the critical density. For the vapor at temperatures between 300 and 305.3 K, we estimate the uncertainty in both density and pressure calculations as less than 0.5%. For the liquid in this temperature range, uncertainties in both density and pressure calculations are estimated as 0.5% at pressures to 10 MPa, but the density uncertainty drops to 0.1% above 10 MPa.

When conditions are very close to the critical point, that is, with temperatures from 305.3 to 307 K with pressures between 4.6 and 5.2 MPa, the uncertainty in calculated densities may be 5%; the uncertainty in pressure in this region is 0.2%. Outside this critical region, from 305.3 to 315 K, the uncertainty in density is estimated at 0.3% and the uncertainty in pressure is 1% for pressures below 40 MPa and 3% for pressures above 40 MPa. For the higher supercritical temperatures, we estimate the uncertainty in density as 0.2% for pressures below 40 MPa and 0.5% for higher pressures. The pressure uncertainties are 0.5% for pressures below 40 MPa and 3% for pressures.

#### 4.3.3. Heat Capacities

Deviation plots for the isochoric heat capacity, isobaric heat capacity, and heat capacity of the saturated liquid are presented in Figs. 15-17. The sources of the data, with ranges and summary statistics, are listed in Table 13; Table 11 gives additional overall statistics for the molar heat capacity data. For the 214 primary  $C_V$  data of Roder<sup>47</sup> and Sengers,<sup>23</sup> the AAD-% is 1.5%; the tabulated densities and temperatures were used as the independent variables for this comparison. Roder used a constant-volume adiabatic calorimeter and reported experimental uncertainties ranging from 2% to about 5% as the critical point is approached. Figure 15 shows that the SWEOS correlates the primary data from Ref. 47 essentially to within the experimental uncertainty. The largest deviation among these primary data is 3.8% and occurs at 189 K and 18 mol·dm<sup>-3</sup>; this point had a 2.1% deviation from the equation of state of Ref. 4.

Secondary  $C_{\nu}$  data from Ref. 47 were chosen because they seemed to be outliers, were not included in the



Fig. 12a. Pressure deviations versus experimental pressures of primary data for temperatures above 315 K. Data are from: Straty<sup>34</sup>; Douslin<sup>22</sup>; Michels<sup>42</sup>; Parrish<sup>46</sup>; Sengers<sup>23</sup>; Pal<sup>31</sup>.

tables of Ref. 4, or were near the saturation boundary or critical point; these are also shown in Fig. 15. Even for these data, the deviations are less than 6%; the largest deviation occurs at 110 K and 21 mol·dm<sup>3</sup>. We noted in Sec. 3.3 that the densities tabulated in Ref. 47, were based on the equation of state of Goodwin *et al.*<sup>4</sup> and an absolute calibration of certain volumes; for the 209 experimental  $C_V$  points in Roder's table,<sup>47</sup> the AAD-% deviation between densities listed in Ref. 47 and those calculated from his tabulated pressures using the SWEOS was 0.4%. We conclude that the uncertainty in density to be associated with the  $C_V$  data of Ref. 47 is not important in our study.

The scaled equation of state of Sengers<sup>23</sup> provided additional  $C_{\nu}$  data near the critical point of ethane. This equation was described in Sec. 4.3.2, as it also provided *PVT* data in the critical region. The large deviations seen in Fig. 15 (the maximum is about 17% at 306 K and 6.9 mol·dm<sup>-3</sup>) reflect the inability of the classical SWEOS to describe the fluctuation-induced weak divergence of the isochoric heat capacity near the critical point. Theoretical and observational evidence indicate a divergence described by  $C_{\nu} \sim (T - T_c)^{-\alpha}$  along the critical isochore;  $\alpha \approx 0.11$ . This critical exponent is incorporated in the scaled equation of state,<sup>23</sup> but it is not approximated by the SWEOS.

Calculation of the isochoric heat capacity using the SWEOS is not recommended for temperatures from 305 to 306 K with densities between 5.5 and 8.0 mol·dm<sup>-3</sup>. Uncertainties of 10%, in  $C_{\nu}$  calculations, are estimated for temperatures from 305 to 307 K with densities between 5 and 8.5 mol·dm<sup>-3</sup>. Outside this general critical region, we estimate the uncertainty in calculation of the isochoric heat capacity as 2.5%, except below 150 K, where the uncertainty rises to 5%.

Furtado<sup>48</sup> and Bier *et al.*<sup>49</sup> provided the primary data for the isobaric heat capacity. For these 359 points, the AAD-% is 1.1%. Bier *et al.*<sup>49</sup> used a flow calorimeter and estimated accuracies of about 0.1% at low pressures, 0.2% at higher pressures, and between 0.5 and 2.5% near the critical point. Their extrapolations to zero pressure<sup>49×49a</sup> have been discussed in Sec. 4.2. As seen in Fig. 16, as well as in the statistical summary of Table 13, deviations from the SWEOS of about 1.5% are typical throughout the entire range of the data. A large part of the discrepancy can be attributed to the contribution



Fig. 12b. Density deviations versus experimental density. References and temperature range are as in Fig. 12a.

ascribed to the ideal gas state; as a fraction of the total measured isobaric heat capacity, this contribution amounts to more than 99% for data at 0.1 MPa and close to 40% for data at 10 MPa. (In the region of the critical anomaly, the proportion attributable to the ideal gas state drops; for the maximum measured  $C_P$  of Ref. 49, the ideal gas contributes somewhat more than 10%.) The deviations for  $C_P^{id}$  from our ideal gas correlation, shown in Fig. 7, propagate strongly in the pressure-dependent  $C_P$  comparisons.

The largest deviation for the primary data of Ref. 49 is 6% and occurs near 317 K and 6 MPa. This point is in the general critical region, and a change of 0.5% in the pressure will reduce the deviation in  $C_P$  to about 0.2%; we also note that the scaled equation of state<sup>23</sup> gives a value of  $C_P$  which differs from that calculated from the SWEOS by about 1.2%. We have excluded 3 points of Ref. 49 from our primary data set; these are also in the critical region and are indicated in Fig. 16. For the point with the largest deviation, 16% at 305.35 K and 4.6 MPa, the deviation from the equation of state of Goodwin *et al.*<sup>4</sup> is 17%; the point is outside the stated range of Sengers' scaled equation.<sup>23</sup>

Furtado<sup>48</sup> also used an isobaric flow calorimeter. He reported a general accuracy of 0.7% for  $C_P$  measurements in the single phase. The 299 points we have indicated in Table 13 are identical to those reported in Ref. 4, except that points extrapolated to zero pressure have not been included in comparisons for the SWEOS. Also, points from Ref. 48 reported on the two-phase boundary and duplicate points are excluded; 3 replicate state points with (slightly) different experimental values of  $C_P$  are included in our data. Among the primary data, there are several regions in which the deviations from the SWEOS are large. All of the data below 133 K (10 points) have deviations between 1.9 and 3.1%; these comprise almost all of the data with densities above 20 mol·dm<sup>-3</sup>. In addition, some of the data in the general region of the critical point were retained as primary data and exhibit large deviations. Some of the measurements along the 5.6 MPa isobar, between 310 amd 314 K, have deviations between 4 and 6%; in the correlation of Goodwin et al.<sup>4</sup> deviations of up to 35% are seen for data along this isobar. There are a few other outliers among the primary data, and the point with the maximum deviation of 8.6% is near 300 K and 6.9 MPa; this point has a deviation of almost 10%



Fig. 13. Pressure deviations versus experimental pressures of secondary data. Data are from: Pal<sup>31</sup>; Tomlinson<sup>87</sup>; Straty<sup>34</sup>; Reamer<sup>86a</sup>; Douslin<sup>22</sup>; Michels<sup>42</sup>; Beattie<sup>82</sup>; Law<sup>84</sup>; Golovskiy<sup>83</sup>; Besserer<sup>82a</sup>; Khazanova<sup>85</sup>; Miniovich<sup>86</sup>; Wallace<sup>88</sup>.

from the correlation of Ref. 4, and upon reflection, it should have been dropped from the primary data set.

Secondary data from Furtado,48 including many points in the critical region and points identified as outliers from preliminary fits, show much larger deviations. Along the isobars near 4.7 and 4.9 MPa, large deviations of 20 to 40% arc sccn in Fig. 16. Along these isobars, the correlation of Goodwin et al.<sup>4</sup> has deviations from the experimental data of up to 36%. Most of the large deviations occur at subcritical temperatures, near the saturated liquid boundary; absolute determination of mass flowrates from volumetric flowrates is difficult in this region. Among the supercritical data, only a point near 306 K and 4.9 MPa displays a large (31%) deviation. For this point, a 0.5% change in the pressure assigned to this nearly critical equilibrium point reduces the deviation to less than 1%. Further, the scaled equation<sup>23</sup> gives a value of  $C_P$  which differs by only 1% from the value calculated using the SWEOS at the reported state point.

We have included critical region  $C_P$  data of Miyazaki et al.<sup>89</sup> in Fig. 16 and in the statistical summary. These again were obtained with a flow calorimeter and an ethane purity of 99.7% was specified; pressure gradients

driving the flow ranged from 0 to 0.001 MPa and the temperature differences upon which the primary enthalpy measurements were made ranged from 1.5 to 28 K. The assignment of a state temperature to the value of a mean  $C_P$  across a finite temperature difference was based on a smoothing technique which may be subject to error near the critical point. Figure 16 shows that the agreement of these data with our SWEOS is generally poor, with deviations up to 71%. For the point with the largest deviation, at 309.15 K and 5.167 MPa, the associated experimental measurement has a temperature change of nearly 17 K; a change of less than 1 K in the assigned temperature brings the value of  $C_P$  into complete agreement with the SWEOS calculation. For the point as tabulated in Ref. 89, the scaled equation<sup>23</sup> and the SWEOS yield values of  $C_P$  which differ by less than 2%.

We estimate the uncertainty in calculations of the isobaric heat capacity from the SWEOS as 1.5% throughout the liquid and vapor states at temperatures from 150 K to 290 K and in the supercritical region above 320 K. An exception is made for the compressed or saturated liquid with densities above 20 mol·dm<sup>-3</sup>, where the uncertainty increases to 4%. Below 150 K, the vapor isobaric heat



Fig. 14. Density deviations versus experimental densities of secondary data. References are as in Fig. 13.

capacity has an uncertainty near 5%. At points closer to the critical point, but outside a region bounded by 305 and 307 K, and by 5 and 8.5 mol·dm<sup>-3</sup>, the uncertainty is 6%. At points within this boundary, we note that  $C_P$  as calculated by the classical SWEOS diverges along the critical isochore with the mean field exponent of 1 rather than the observed exponent of about 1.2; in addition, roundoff error causes the finite value of  $C_P \approx 4 \times 10^{12}$ J·mol<sup>-1</sup>·K<sup>-1</sup> at the critical point. Thus, percent errors can be very large in this region, but physical measurements of these extremely large heat capacities are exceedingly difficult.

Roder<sup>47</sup> also provided primary data for the molar heat capacity while maintaining the liquid at saturation; deviations for these data are shown in Fig. 17. Roder used the same constant-volume adiabatic calorimeter for  $C_{\sigma L}$  as discussed previously in the context of  $C_{\nu}$  measurements. The uncertainty in the experimental measurements was reported to range from 0.5% to 5% within a few kelvins of the critical point. Systematic trends in the deviations are clear from Fig. 17; we were not able to fit these data well at low temperatures without distorting the thermo-

dynamic surface in other regions. The maximum deviation of 3.3% occurs at a temperature near 107 K; in Ref. 4, explicit comparisons with these data were not made, and in Ref. 5, deviations between these data and their equation of state were 7.5% near the triple point temperature and 5.4% near the critical point.

At low temperatures, older data for  $C_{\sigma L}$  of Wiebe *et al.*<sup>90</sup> and of Witt and Kemp<sup>91</sup> agree surprisingly well with the more recent work of Roder.<sup>47</sup> These secondary data are also shown in Fig. 17. Above the normal boiling point of ethane, about 184.6 K, scatter and systematic deviations of the data from Ref. 90 are evident; these are most likely due to the quality of the data rather than being a reflection of the SWEOS correlation. The correlation from Ref. 47 shown in Fig. 17 was a fit to only experimental  $C_{\sigma L}$  data from that source; the correlation from Ref. 4 indicates the problems of correlating  $C_{\sigma L}$  data with a wide ranging equation of state.

We estimate the uncertainty in calculations of  $C_{\sigma L}$  from our SWEOS as 3.5% from the triple point to 145 K and 0.6% from 145 to 280 K. Closer to the critical point, the usual problems with the classical equation of state occur;

First author	Ref.	Туре	No. pts.	Temperature range, K	Pressure range,MPa	Density range mol·dm <sup>-3</sup>	AAD – %
Bier <sup>a</sup>	49	C <sub>P</sub>	118	283-473	0.1-10	0.03-14	1.1
Bier	49	$C_P$	130	283-473	0.1-10	0.03-14	1.2
Furtado <sup>a</sup>	48	CP	241	100-378	1.7-14	0.6-21	1.1
Furtado	48	C <sub>P</sub>	299	100-378	1.7-14	0.6-21	2.2
Miyazaki	89	$C_P$	45	298-323	4.5-13	2.5-13	7.9
Roder <sup>a</sup>	47	$C_{V}$	184	112-329	1.6-34	1.6-21	1.2
Roder	47	$C_{\nu}$	209	112-329	1.6-34	1.6-21	1.3
Roder <sup>a</sup>	47	$C_{\sigma L}$	106	94-301	10-6-4.5	9.7-22	0.84
Sengers <sup>a,b</sup>	23	$C_{V}$	30	304-325	4.7–6	4.9-9	3.5
Wiebe	90	$C_{\sigma L}$	41	97-295	10 <sup>-5</sup> -3.9	11-21	0.95
Witt	91	$C_{\sigma L}$	39	95-181	10-6-0.1	18-22	1.5

TABLE 13. Sources of heat capacity data

\* Only primary data are included in these statistics.

<sup>b</sup> These data were obtained from a scaled equation of state; they are not experimental data.



Fig. 15. Deviations for molar isochoric heat capacity versus density. Data are from Roder<sup>47</sup> and Sengers<sup>23</sup>.



Fig. 16. Deviations for molar isobaric heat capacity versus pressure. Data are from: Furtado<sup>48</sup>; Bier<sup>49</sup>; Miyazaki<sup>89</sup>.

the uncertainty increases to 5% for temperatures from 280 to 303 K. For temperatures between 303 and  $T_c$ , very large errors are possible.

#### 4.3.4. Sound Speed

Data for the speed of sound both in the single-phase region and in the saturated liquid are compared with the SWEOS in Figs. 18 and 19 and are summarized in Table 14. Tsumura and Straty<sup>50</sup> used a pulse-echo method at 1 and 10 MHz and a pulse-electric method at 10 MHz for points where critical region attenuation was important; quartz crystals were used to generate and detect the acoustic wave. The estimated experimental uncertainty in the sound speed ranged from 0.05 to 0.1%.

For the 109 primary single-phase data from this source, the AAD-% is 0.36% and systematic trends are seen in the deviations of Fig. 18. For instance, at the extreme right of the figure, the 100 K isotherm corresponds to densities of about 21.4 to 21.7 mol·dm<sup>-3</sup>; the corresponding deviations seem to be linear in density and range from 0.1 to 0.8%. The difficulty in fitting these data arises from the large magnitude of the slope  $\partial P/\partial \rho|_T$ , as also discussed in Sec. 4.3.2 above. As indicated by Eq. (26), the primary speed-of-sound data were used to minimize the residuals associated with this slope (in conjunction with all the other data in the multiproperty fit); however, deviations for this quantity range to 1.4% along the 100 K isotherm. The equation-of-state correlation of Goodwin *et al.*<sup>4</sup> generates deviations from 1.5 to 2% for these sound speed data at 100 K; Sychev *et al.*<sup>5</sup> indicate deviations to 30% at 100 K and show deviations from about -4 to 4% for the data from Ref. 50 along the 120 K isotherm. The largest deviation between the SWEOS and the primary single-phase sound speed data of Tsumura and Straty<sup>50</sup> is 0.94% and occurs at 280 K near 5 MPa.

Data from Ref. 50 along the critical isotherm and some data at 300 and 323 K were considered secondary and are indicated in Fig. 18; these data were not included in the tabulation of Ref. 4. The deviations are seen to be comparable to those of the primary data, with the maximum at nearly 0.9% for the point closest to critical conditions at  $T_c$  and 4.93 MPa. The SWEOS correlation does not produce a vanishing sound speed at the critical point as is indicated by theory and a substantial body of experimental data for other fluids. The finite value of about 181

320



Fig. 17. Deviations for molar heat capacity along saturated liquid boundary versus temperature. Data are from: Roder<sup>47</sup>; Wiebe<sup>90</sup>; Witt<sup>91</sup>. Curves are from Goodwin<sup>4</sup> and Roder<sup>47</sup>.

m·s<sup>-1</sup> is caused by the finite, rather than weakly divergent, value of  $C_{\nu}$  and the nonzero value of  $\partial P/\partial \rho|_T$  (which is about  $10^{-11}$  MPa·dm<sup>3</sup>·mol<sup>-1</sup> due to round-off error) at the critical point.

Terres et al.<sup>51</sup> provided primary sound speed data in the single phase region at higher temperatures and in the compressed vapor; their tabulated extrapolations to zero pressure and the single point on the phase boundary were not considered. The interferometric method used in Ref. 51 operated near 500 kHz, and the estimated accuracy was 0.3%. As seen in Fig. 18, some of the deviations between these data and the SWEOS are rather large. The 4% deviation occurs near 323 K and 8.8 MPa; this point is quite close to a measurement from Ref. 50 at 323 K and 9 MPa which has a deviation of 0.05%. The discrepancy between measurements from Refs. 50 and 51 in the region of overlap was noted by Tsumura and Straty<sup>50</sup> but could not be explained; we prefer the more recent work of Ref. 50. Deviations for the 323 K isotherm range from -2.7% to 4% compared to the SWEOS and from -2.5to 5.8% when compared to the correlation of Ref. 4. The isotherm at 293 K, extending to -2.2% near 2.3

mol·dm<sup>-3</sup>, and a single outlier near 373 K and 3 mol·dm<sup>-3</sup> are also apparent in Fig. 18.

Primary sound speed data measured along the saturated liquid boundary by Tsumura and Straty<sup>50</sup> are illustrated in Fig. 19. Systematic deviations between the data and the SWEOS are again apparent. Near the triple point, corresponding to the highest densities for these data, the deviations reach only 0.5% but follow a clear trend. For these points, below 100 K, the equation of state of Ref. 4 shows deviations of about 1.5% and Ref. 5 indicates deviations from 20 to 40% between 91 and 110 K. At temperatures above 245 K, the systematic deviations are even more pronounced; the maximum deviation for the primary  $W_{\sigma L}$  data is 1.8% near 280 K. The equation of state from Goodwin *et al.*<sup>4</sup> gives deviations in the critical region extending to 9% at 299 K, and Sychev *et al.*<sup>5</sup> indicate deviations exceeding 10%.

The secondary data for  $W_{\sigma L}$  include the 4 points from Tsumura and Straty<sup>50</sup> between 300 and 303 K as well as data from Vangeel<sup>92</sup> and from Poole and Aziz.<sup>93</sup> A single point from Ref. 93 was reported at a temperature below the triple point as given in Table 1 and was not consid-

First author	Ref.	Туре	No. pts.	Temperature range, K	Pressure range, MPa	Density range mol·dm <sup>-3</sup>	AAD - %
Poole	93	W <sub>oL</sub>	25	93-199	10-6-0.2	17.5–22	0.40
Terres <sup>a</sup>	51	W	92	293-448	0.1-12	0.03-10.5	0.79
Tsumura <sup>a</sup>	4,50	W	109	100-323	3.6-37	4.8-22	0.36
Tsumura	4,50	W	156	100-323	3.6-37	4.8-22	0.31
Tsumura <sup>a</sup>	4,50	$W_{\sigma L}$	47	91299	10-6-4.3	10-22	0.49
Tsumura	4,50	$W_{\sigma L}$	51	91-303	10-6-4.6	9-22	0.56
Vangeel	92	$W_{\sigma L}$	44	96-289	10-6-3.4	12-21	0.44

TABLE 14. Sources of sound speed data

\*Only primary data are included in these statistics.



Fig. 18. Deviations for speed of sound in single-phase region versus density. Data are from Tsumura<sup>50</sup> and Terres<sup>51</sup>.



Fig. 19. Deviations for speed of sound of saturated liquid versus temperature. Data are from: Tsumura<sup>50</sup>; Poole<sup>93</sup>; Vangeel<sup>92</sup>. Curves are from Goodwin<sup>4</sup> and Tsumura<sup>50</sup>.

ered. The critical region data from Tsumura and Straty show deviations to 2.5% and again demonstrate the difficulty with the classical SWEOS in this region where the sound speed should approach zero. The measurements of Vangeel are seen to agree quite well with those of Tsumura and Straty except below about 125 K. Tsumura and Straty<sup>50</sup> attribute this difference to dispersion associated with the dimensions of the experimental cell. The data from Poole and Aziz<sup>93</sup> differ from the other sources especially below 175 K. The reasons for this discrepancy are not clear, but the more recent data of Ref. 50 are to be preferred. The maximum deviation of the data of Ref. 93 from the SWEOS correlation is 0.8% near 100 and 115 K.

The difficulty in correlating the sound speed to within the reported experimental accuracies while obtaining good agreement with other data and using the classical SWEOS formulation is clear. The problems are most serious at high densities and near the critical point. We can summarize our observations by the following guidelines for the use of the SWEOS in predicting the speed of sound in ethane. Below 100 K, in the saturated or compressed liquid, we estimate the uncertainty as 1%. Between 100 and 225 K, the liquid phase sound speed has an uncertainty of 0.6%. Between 225 K and 290 K and above 310 K, the uncertainty increases to about 1%. Between 290 and 310 K, for densities outside the range 5 to 10 mol·dm<sup>-3</sup>, a 2% uncertainty is estimated. Closer to the critical point, the uncertainty ranges to 10%; at points between 305 and 306 K, with densities from 5.5 to 8 mol·dm<sup>-3</sup>, the classical SWEOS cannot be used. For the gas below 290 K, we estimate an uncertainty of 0.6%; below 150 K, this rises to 1%. For pressures above 35 MPa at subcritical temperatures and above 12 MPa for supercritical temperatures, there are no data and we estimate an uncertainty of 2%.

These comparisons with experimental data (and the process of determining the coefficients of the SWEOS, as described in Sec. 3), only probe four derivatives of the dimensionless, residual Helmholtz energy, namely  $\phi_{\delta_1} \phi_{\delta_2} \phi_{\delta_3}$ ,  $\phi_{\tau\tau}'$ , and  $\phi_{\delta\tau}'$ . While this should describe the actual surface quite well, additional uncertainties will enter any calculation which requires other derivatives of  $\phi'$  or integrals of the Helmholtz function. We hesitate to make any quantitative predictions of the errors involved in calculating any thermodynamic quantities not discussed in

this section, but we conjecture that these errors will be comparable to those found with any other precise equation of state for ethane.

### 4.4. Transport Property Comparisons

#### 4.4.1. Viscosity

The viscosity of the dilute ethane gas, as described by Eq. (10), is compared with experimental data in the deviation plot of Fig. 20. The 6 primary data of Kestin et al.<sup>52-54</sup> were obtained using an absolute method in an oscillating disk instrument and had a reported accuracy of 0.2%; the comparison between these data and our correlation of Eq. (10) gives an AAD-% of 0.85%. Secondary data from Refs. 52, 53, 57, and 94-100 are also shown in the figure, and additional, especially earlier, data sources are discussed in these references and in Refs. 26, 27, and 101. Where appropriate, the data have been adjusted to zero density by subtracting the (small) value of the excess viscosity calculated from the correlation of Eq. (15) at the experimental pressure, which was usually near 0.1 MPa. The adjustment is less than 0.25% for all temperatures. Secondary data in the gas phase at pressures less than 0.12 MPa were included in dilute gas comparisons and not below with the total viscosity. For the 76 points illustrated, in the range from 250 K to 523 K, the comparison of the data to Eq. (10) gives an AAD-% of 0.89%, a BIAS-% of 0.18%, and an RMS-% of 0.98%.

The statistics above indicate considerable scatter among the data; this is also illustrated by Fig. 20. The two correlations shown, by Boushehri et al.27 and by Younglove and Ely,<sup>10</sup> also indicate some systematic differences. Boushehri *et al.*<sup>27</sup> did not consider any data below about 300 K (they do not include Ref. 97 among their secondary data) and indicate an accuracy of 0.5% in their entire range from 250 to 500 K; extrapolation of the curve to lower temperatures is not warranted. At 300 K, near the most reliable data, the difference between our correlation and that from Ref. [27] is about 0.3%. Both correlations differ from the primary data in this region: a deviation of about 1-1.2% is seen for the present correlation, and a deviation of about 0.8% is seen for the correlation of Boushehri et al. The original data papers specified an accuracy of 0.1-0.2%. At higher temperatures, the spread of even the most accurate data is apparent. At 420 K, for instance, the point from Ref. [52] differs by about 0.8% from the more recent data by Abe et al.95 Although the true value of the dilute gas viscosity may indeed lie between these two values, as fit by Boushehri et al., our correlation gives a value about 0.4% lower than the datum from Ref. [95] which we did not consider primary. Although the differences between our correlation and that from Ref. [27] are systematic, the maximum deviations of about 1.5% (near 250 K, the lower limit in Ref. [27]) are within combined uncertainties.

Younglove and Ely<sup>10</sup>, whose correlation for  $\eta_o$  and selection of primary data are quite similar to ours, give a value at 100 K which differs by less than 3% from that calculated using Eq. (10). All of the correlations considered here, from our Eq. (10), from Ref. [10], and from Ref. [27], are based on two adjustable parameters and have difficulty fitting the best experimental data over a wide range in temperature. We estimate the uncertainty of our correlation as 1% in the range 300 to 375 K; 1.5% from 250 to 300 K and above 375 K; and 5% from the triple point temperature to 250 K.

There are also several sources for the viscosity of ethane at elevated pressures. The total viscosity is correlated by the sum of contributions from Eqs. (10) and (15). In addition to comparisons to the 438 primary data of Diller and Saber,<sup>55</sup> Diller,<sup>56</sup> and Carmichael and Sage,<sup>57</sup> we have included comparisons with data from Refs. 24, 96, 100, 102, and 103. The general experimental accuracies were specified as about 2% for Refs. 55 and 56 and about 0.5% for Ref. 57. Table 15 summarizes these data and comparisons for the entire set of 1225 points. Additional, especially older, data sources were cited by Hanley *et al.*<sup>26</sup> and in the monograph by Stephan and Lucas;<sup>104</sup> some of these are included in the comparisons of Ref. 3.

Figure 21a illustrates the deviations for the primary data and much of the secondary data. Data of Iwasaki and Takahashi<sup>96</sup> and of Strumpf et al.<sup>24</sup> emphasized the region of the critical point and indicate a critical enhancement. Because of the small size of the critical region relevant to the viscosity enhancement and in the interest of simplicity of the correlating equations, we have not studied the theoretically predicted enhancement of the viscosity, nor have we critically evaluated the data of Refs. 24 and 96. Our correlation does not account for any critical enhancement in the viscosity; therefore, the critical region, 305–306 K and 6–7.5 mol·dm<sup>-3</sup>, should be excluded when making calculations with this viscosity correlation. Because of the larger deviations between the correlation and data from Refs. 24 and 96, which may be of interest to some readers, these data are separately illustrated in Fig. 21b.

The AAD-% is 0.75% for the 438 selected primary viscosity data. Including the few secondary data from Diller and Saber,<sup>55</sup> Diller,<sup>56</sup> and Carmichael and Sage,<sup>57</sup> as well as data from Baron *et al.*,<sup>102</sup> Eakin *et al.*,<sup>100</sup> and Swift *et al.*,<sup>103</sup> we have 718 points with an AAD-% of 1.05%. When the data compared include the critical region measurements of Iwasaki and Takahashi<sup>96</sup> and Strumpf *et al.*,<sup>24</sup> the AAD-% for the resulting 1225 points increases to 3.07% if the pressures of Ref. 96 are used and 2.01% if the densities of Ref. 96 are used. These statistical comparisons and the appearance of Figs. 21a and 21b warrant some discussion.

We are aware of no accurate data for the vapor phase below 290 K. From the subcritical data at 290 K and above, as well as supercritical data in the low density region, and our study of the dilute gas correlation, we can assess the uncertainty of the viscosity correlation in the vapor phase. From the triple point temperature to 250 K,



Fig. 20. Deviations for dilute gas viscosity coefficient versus temperature. Primary data are from: Kestin<sup>52</sup>; Kestin<sup>53</sup>; Fleeter<sup>54</sup>. Secondary data are from: Kestin<sup>52</sup>; Kestin<sup>53</sup>; Iwasaki<sup>96</sup>; Trautz<sup>94</sup>; Craven<sup>98</sup>; De Rocco<sup>97</sup>; Lambert<sup>99</sup>; Carmichael<sup>57</sup>; Eakin<sup>100</sup>; Abe<sup>95</sup>. Curves are from Boushehri<sup>27</sup> and Younglove<sup>10</sup>.

First author	Ref.	No. pts.	Temperature range, K	Pressure range, MPa	Density range mol·dm <sup>-3</sup>	AAD-%
Baron	102	40	325-408	0.7-55	0.2–15	1.85
Carmichael <sup>a</sup>	57	222	300-478	0.1-36	0.04-15	0.76
Carmichael	57	226	300-478	0.1-36	0.04-15	0.81
Diller <sup>a</sup>	55	144	95-320	0-32	0.3-22	0.69
Diller	55	164	95-320	0-32	0.3-22	0.87
Diller <sup>a</sup>	56	72	295-500	1.7-55	0.7-16	0.83
Diller	56	76	295-500	1.7-55	0.7–16	0.96
Eakin	100	198	294-511	0.7-69	0.2-17	1.33
Iwasaki <sup>b</sup>	96	402	298-348	0.2-13	0.1-10	5.6
Iwasaki <sup>b,c</sup>	96	402	298-348	0.2-13	0.1-10	2.4
Strumpf	24	105	302-322	4.5-8.2	6.7-10	7.4
Swift	103	14	193–305	0.2–5	9–18	1.82

TABLE 15. Sources of viscosity data at elevated pressures

<sup>a</sup>Only primary data are included in these statistics.

<sup>b</sup>These measurements emphasized the critical region.

This comparison is based on experimentally determined density; all other comparisons are based on experimental pressure.



Fig. 21a. Deviations for viscosity versus density for primary and secondary data. Data are from: Carmichael<sup>57</sup>; Diller<sup>55</sup>; Diller<sup>56</sup>; Baron<sup>102</sup>; Swift<sup>103</sup>; Eakin<sup>100</sup>. Curves from correlation of Younglove<sup>10</sup> at 325 and 600 K.

we estimate the uncertainty in the correlation for the vapor phase as 5%; in the saturated vapor at 250 K, the excess portion of the viscosity contributes less than 6% to the total viscosity. There are a few primary data from Diller and Saber,<sup>55</sup> and secondary data from Eakin *et al.*<sup>100</sup> and from Iwasaki and Takahashi;<sup>96</sup> these extend from 290 K to  $T_c$  in the vapor phase; typical deviations for these data are within 2% except for an outlier in Ref. 100 near 9% and near 294 K and 2.3 mol·dm<sup>-3</sup>. We estimate a 2.5% uncertainty for the vapor at temperatures from 250 K to the critical point.

Data for the saturated liquid are available<sup>55</sup> down to 95 K; for the compressed liquid, the 100 K isotherm from Diller and Saber<sup>55</sup> is the lowest temperature. We have fitted these primary liquid data to well within the experimental uncertainty of about 2%; except for a single point from Ref. 55 (considered secondary) at 110 K and 32 MPa with a 2.6% deviation, the largest deviation is 1.7% and occurs in the saturated liquid at 225 K. Additional liquid data from Refs. 24, 57, 100, and 103 are also fitted well by the correlation; exceptions include a 4.3% devia-

tion for a point of Swift *et al.*<sup>103</sup> near 193 K and 4.1 MPa and some larger deviations (to about 5%) for the liquid data of Eakin *et al.*<sup>100</sup> at 294 K and pressures above 50 MPa. The highest pressure points reported by Eakin *et al.*<sup>100</sup> were extrapolations of their experimental measurements. For the liquid from the triple point to the critical point for pressures from the saturation boundary to 30 MPa, we estimate the uncertainty of the correlation as 2%. For higher pressures, the uncertainty increases to about 5%.

Many of the data illustrated in Figs. 21a and 21b are at supercritical temperatures, and most of these primary data have been fitted to better than 2%. Among the outliers, 21 points from Carmichael and Sage,<sup>57</sup> especially near 311 K, have deviations exceeding 2%. A point along the 311 K isotherm with a pressure of 4.96 MPa has a deviation of -2.8%; a replicate point listed in Ref. 57 and included in our data set has a deviation of less than 0.1%. The largest deviations among the supercritical primary data from Ref. 57 are near 3% and occur at the highest measured temperature (478 K) near 14 MPa. The lowest



Fig. 21b. Deviations for viscosity versus density for secondary data from Iwasaki<sup>96</sup> and Strumpf<sup>24</sup>.

density data along this 478 K isotherm also have deviations of more than 2%. In addition to the subcritical data discussed above, Diller and Saber<sup>55</sup> measured a single supercritical isotherm at 320 K; a single primary point from this set has a deviation of 2% and the 7 points with pressure less than 4 MPa have deviations to 3.4% and have been excluded from the primary data. The high temperature data of Diller<sup>56</sup> include a few isolated points with deviations of near 2% and a point near 400 K and 6.5 MPa with a deviation of -2.6%; 4 points from Ref. 56 had deviations between 2.9 and 3.8% and were not considered primary data.

With the exception of the data in the critical region as mentioned above, even the secondary supercritical data are reproduced fairly well by our correlation. The data from Ref. 102 along the 325 K isotherm have deviations which reach 6.7% at the highest pressure (55 MPa). The secondary data of Carmichael and Sage<sup>57</sup> include a few replicate points in the general critical region with deviations to about 5%; these have temperatures of 305.372 K and pressures of 4.86 MPa. The deviations for the supercritical data of Eakin<sup>100</sup> are all less than about 4%.

The deviations in the critical region are large and systematic. Iwasaki and Takahashi<sup>96</sup> measured both viscosity and density as functions of temperature and pressure in the critical region. Their PVT data differ considerably from our thermodynamic correlation, and thus a comparison with the viscosity correlation depends dramatically upon whether the experimental pressure or density is considered an independent variable. For instance, if the experimental pressure is taken as the independent variable, the maximum deviation from the viscosity data of Ref. 96 is about 27% and occurs along the isotherm at 305.65 K ( $T^* = 10^{-3}$ ) and 4.8934 MPa. For this state point, we calculate a density of 5.615 mol $\cdot$ dm<sup>-3</sup>; the reported density is 7.855 mol·dm<sup>-3</sup>. There is only a 0.4%deviation when the viscosity is calculated at the experimental density. The maximum deviation between the supercritical viscosity data of Ref. 96 and our correlation when the experimental density is used as the independent variable is 5.7% at 305.65 K and 4.5357 MPa; at this state point, the calculated and experimental densities are much closer, and the viscosity deviation is 5.5% when the experimental pressure is used.

The source of the discrepancy between the PVT surfaces near the critical point is not clear; we have given comparisons for both types of calculation in Table 15 but have illustrated only the larger deviations when using

the experimental pressures. Strumpf *et al.*<sup>24</sup> also measured densities for their isochoric data, but they did not measure pressure. Their viscosity data in the critical region differ considerably from those reported in Ref. 96 and large deviations from our correlation occur even at 322 K and 9.6 mol·dm<sup>-3</sup>; here the deviation is 7%. After considering the agreement among the alternate data, we have concluded that the data from Ref. 96 may suffer some systematic error. We estimate the uncertainty of the viscosity correlation in the supercritical region as generally 2%. At pressures greater than 50 MPa, the uncertainty increases to 5%, and in the critical region, from 305 to 307 K and 5 to 8.5 mol·dm<sup>-3</sup> as indicated above), the uncertainty is also 5%.

#### 4.4.2. Thermal Conductivity

For the dilute ethane gas, the correlation of Eq. (13), using Eq. (14) for the factor  $f_{int}$ , describes the thermal conductivity, and Fig. 22 illustrates the deviations of experimental data from this correlation. In addition to the primary data of Roder and Nieto de Castro<sup>58</sup> and of Prasad and Venart,<sup>59</sup> the figure includes points from Refs. 54, 98, 104a-106, 110 and 111. Other experimental work is cited in these references, and some data which incidentally include data at low pressures are included in the comparisons for the density-dependent thermal conductivity below. In all cases, we have used either the tabulated zero-density extrapolations of the authors, or we have subtracted a small value for the excess thermal conductivity as calculated from the correlation of Eqs. (17-18) at the experimental conditions. For the primary data, comprising 12 points spanning the temperature range 245-600 K, the AAD-% is 0.67%. Overall, for the 45 experimental points included in the figure, the AAD-% is 1.96%; the BIAS-% is -0.48%; and the RMS-% is 2.38%.

The agreement among the sources of data as well as the quality of the present dilute gas correlation are worse than those seen in the dilute gas viscosity correlation. This last point is perhaps not surprising, since the theory for  $f_{int}$  is not completely rigorous and the form we have chosen for this factor may be overly simplified.

Note added in final revision: A theoretical approach based on an approximate solution to the Wang Chang-Uhlenbeck equation has very recently been studied in analyzing new ethane thermal conductivity data.<sup>111</sup> This approach requires empirical evaluation of several collision integrals in addition to estimates of vibrational and rotational collision numbers. The new dilute gas extrapolations are shown in Fig. 22. Deviations from the correlation and from both primary and secondary data are substantial and systematic. However these new data agree with the correlation within combined uncertainties.

The correlations shown also represent fits of limited data. The curve from Roder and Nieto de Castro<sup>58</sup> is their polynomial description of their zero-density extrapola-

tions, and the curve from Hanley *et al.*<sup>26</sup> is based on earlier data. Among the primary data, the largest deviation is 1.4% and occurs at 245 K for a point from Ref. 58. The statistical uncertainty which Roder and Nieto de Castro associate with the extrapolation needed to obtain this zero-density value is 2.7% and the general accuracy of the data is specified as 1.6%.

The  $\lambda_0$  data from Ref. 58 at 225 K and 235 K were not considered primary; at 225 K, 8 measurements with various temperature rises at only 2 distinct densities were extrapolated. Although the statistical uncertainty given in Ref. 58 is 1.1%, a systematic change in these 8 data by only 0.5% will change the extrapolated value of  $\lambda_0$  from a deviation of 2.8% to a deviation of about 0.2% with respect to our correlation. The experimental uncertainty increases when extrapolations of limited measurements at low temperatures and pressures are made. The dispersion among the alternate data throughout the temperature range is unfortunate. Fleeter et al.,54 who also used a transient hot-wire instrument, report a 0.3% uncertainty from their regression to obtain  $\lambda_0$  at 301 K; this point has a deviation of 3.2% from our correlation. The low density results from Ref. 58 seem to show a slightly larger value of  $\partial \lambda / \partial \rho |_T$  at 295 and 305 K and extend to lower densities than the isotherm at 301 K from Ref. 54. The experimental accuracy of the measurements from Yakush et al.<sup>105</sup> is not specified in that paper, and Le Neindre et al.,<sup>106</sup> who used a coaxial cylinder method in the range 130-625 K, indicate a general experimental accuracy of 1.5%.

It is difficult to assess the uncertainty of our correlation for temperatures below that for which data are available. At the lowest temperatures, between the triple point and 200 K, where the transfer of energy between kinetic and internal degrees of freedom is expected to be most inhibited, the correlation seems to exaggerate this effect, and the calculated values of  $f_{int}$  and hence the thermal conductivity may be too small. Thus, at temperatures below 200 K, where the vapor pressure is about 0.2 MPa, the correlation could give errors in excess of 10%. In the range 200 to 350 K, the uncertainty in the correlation is about 3%, and between 350 K and 600 K we estimate the uncertainty as 4%.

Additional measurements, especially at low temperatures and with sufficient isothermal data to permit extrapolation to zero density, could help improve the correlation and might justify a more elaborate expression for  $f_{int}$  than the 2-parameter expression given in Eq. (14).

For higher pressures, the thermal conductivity has contributions from its dilute gas limit, Eq. (13), and the critical enhancement, Eq. (18), as well as from the excess function of Eq. (17). The critical enhancement is shown along isotherms in Fig. 23; the experimental points in the figure were obtained from the original data by subtracting the dilute gas and excess contributions and adjusting the data to lie along true isotherms as indicated. Our correlation gives a reasonable description of the enhancement throughout the range of the data. As indicated above, the distinction between the excess and enhance-



Fig. 22. Deviations for dilute gas thermal conductivity coefficient versus temperature. Primary data are from Prasad<sup>59</sup> and Roder<sup>58</sup>. Secondary data are from: Craven<sup>98</sup>; Yakush<sup>105</sup>; Le Neindre<sup>106</sup>; Fleeter<sup>54</sup>; Prasad<sup>59</sup>; Roder<sup>58</sup>; Clifford<sup>104a</sup>; Tufeu<sup>110</sup>; Millat<sup>111</sup>. Curves are from Hanley<sup>26</sup> and Roder<sup>58</sup>.

ment contributions is operationally ambiguous; our iterative method ensures a good fit for the total thermal conductivity, but may not give a definitive division among the contributions to this total. Olchowy and Sengers<sup>o</sup>, who used a different thermodynamic surface in the critical region, different dilute gas correlation, different (and temperature independent) excess correlation, and a different choice of primary experimental data, obtained a value of  $q_D^{-1} = 0.29$  nm for the single kinetic parameter in the simplified mode coupling theory; their agreement with experimental data is comparable to the agreement of the present correlation.

The deviations of experimental measurements from the total conductivity correlation are shown in Figs. 24a and b, and Table 16 gives additional information about the experimental data from Refs. 54, 59–61, and 106–110. There are additional, especially earlier, sources of thermal conductivity data for ethane listed within these references; Prasad and Venart<sup>59</sup> give an excellent bibliography and Hanley *et al.*<sup>26</sup> cite other literature. The AAD- % for the 1098 primary data of Roder,<sup>60</sup> Prasad and Venart,<sup>59</sup> and Desmarest and Tufeu,<sup>61</sup> which were used to establish the total thermal conductivity correlation, is 1.21%; deviations for these data are shown in Fig. 24a. For the 1357 points listed in the table, the AAD-% is 1.47%. The deviations indicated by these statistics and illustrated in the figures again warrant some discussion.

The major deviations between the correlation and data occur in the general region of the critical point, despite the improved method of correlating the crossover region expressed in Eqs. (18–20). For the primary data of Roder,<sup>60</sup> the estimated experimental accuracy<sup>58</sup> is 1.6%, but it is undoubtedly worse near the critical point. Our representation of the thermal conductivity in the gas phase at low temperatures gives some systematic deviations as indicated in the discussion of  $\lambda_0$ ; near the 225 K isotherm the deviations range to 2.5% at the lowest pressure and systematic deviations to 1.4% are seen near 245 K. There is a single point with 2% deviation near 303 K and 0.24 MPa, but the major disagreements be-



Fig. 23. Critical enhancement of the thermal conductivity. Background term has been subtracted from data and data have been adjusted to 308 K, 312 K, and 325 K isotherms. Data are from: Roder<sup>58</sup>; Prasad<sup>59</sup>; Desmarest<sup>61</sup>. Curves are from present correlation (Eq. [18]).

First author	Ref.	No. pts.	Temperature range, K	Pressure range, MPa	Density range mol·dm <sup>-3</sup>	AAD-%
Carmichael	107	31	278-444	0.1-37	0.03-16	3.81
Desmarest <sup>a</sup>	61	111	308-365	1-28	0.3-12	1.85
Fleeter	54	12	300-301	0.6-3.4	0.3-2	1.60
Gilmore <sup>b</sup>	108	18	348	0.1-304	0.04-20	1.80
Le Neindre	106	113	307-801	0.1-119	0.02-17	1.63
Leng	109	12	341	0.1-26	0.04-13	1.71
Prasada	59	235	294-600	0.2-70	0.07-17	1.58
Prasad	59	239	294600	0.2-70	0.07-17	1.67
Roder <sup>a</sup>	60	752	112-327	0.169	0.04-22	1.01
Roder	60	797	112-327	0.1-69	0.04-22	1.19
Tufeu	110	24	307-500	0.5-30	0.1-7	2.58

TABLE	16.	Sources	of	thermal	conductivity	data
-------	-----	---------	----	---------	--------------	------

\*Only primary data are included in these statistics.

<sup>b</sup>Statistics based on tabulated smoothed data.

330



Fig. 24a. Deviations for thermal conductivity versus density for primary data. Data are from: Roder<sup>58,60</sup>; Prasad<sup>59</sup>; Desmarest<sup>61</sup>.

tween the correlation and these data occur at densities closer to the critical point and along the supercritical isotherms. Along the isotherm nominally denoted 305 K, but spanning the range 302.7 to 307.4 K, the deviation reaches 3.5% at 3.8 MPa or 2.2 mol·dm<sup>-3</sup>; there are no data above 2.4 mol·dm<sup>-3</sup> along this isotherm. Along the 312 K isotherm, there is a deviation of 4.3% near 3 mol·dm<sup>-3</sup>, the largest deviation at subcritical densities for this temperature, and large systematic deviations between 2 and 5.6% at supercritical densities. A few data close to the critical density (between 5.5 and 6.5 mol·dm<sup>-3</sup>) were considered secondary data and have deviations to 6.2%; other points close to the critical density were fitted quite well by our equations. An additional set of measurements along this isotherm was made earlier by Roder and was included in Ref. 60, but these data were considered secondary for our correlation; deviations for these data range to 6.9% at about  $4.9 \text{ mol}\cdot\text{dm}^{-3}$ . The next isotherm, near 325 K and the upper limit of the apparatus, also shows some large deviations. There is a 3% deviation at 5.2 mol·dm<sup>-3</sup>, below the critical density, and large systematic negative deviations to 6% between 6.4 and 11.2 mol·dm<sup>-3</sup>. A few data near 9.7 mol·dm<sup>-3</sup> were not included in the fit and have deviations to 10.4%.

Prasad and Venart<sup>59</sup> indicate experimental uncertainties between 0.7 and 3%, increasing in the critical region. Their data near 295 K include both gas and liquid states and seem to indicate an enhancement near the phase boundary which is not described well by the correlation; the deviations in the liquid phase reach 5.1% and in the gas phase 2 points with deviations to 8.8% were excluded from the primary data set. The isotherm near 318 K has a deviation of 6.8% near 9.7 mol·dm<sup>-3</sup>, and a point with a deviation of 8.3% was excluded from our primary data. The isotherm near 350 K has deviations of up to 5.5% (near 7.2 mol·dm<sup>-3</sup>), and for higher temperatures, the maximum deviation is 2.6% near 600 K and 4.1 MPa.

The primary data of Desmarest and Tufeu<sup>61</sup> emphasized the critical region and show a maximum deviation of 5.1% from our correlation. This point is at 308.75 K, 5.23 MPa, and about 6.9 mol·dm<sup>-3</sup>, and is the primary datum closest to the critical point of ethane. For this point, the experimental value of  $\lambda$  is 83.77 mW·m<sup>-1</sup>·K<sup>-1</sup>; our calculated value is 79.49 mW·m<sup>-1</sup>·K<sup>-1</sup> which comprises contributions of 22.27 mW·m<sup>-1</sup>·K<sup>-1</sup> from the dilute gas term, 24.12 mW·m<sup>-1</sup>·K<sup>-1</sup> from the excess conductivity, and 33.10 mW·m<sup>-1</sup>·K<sup>-1</sup> from the critical enhancement. The isotherms from Ref. 61 at higher temperatures show



Fig. 24b. Deviations for thermal conductivity versus density for secondary data. Data are from: Roder<sup>58,60</sup>; Fleeter<sup>54</sup>; Le Neindre<sup>106</sup>; Prasad<sup>59</sup>; Carmichael<sup>107</sup>; Tufeu<sup>110</sup>; Leng<sup>109</sup>; Gilmore<sup>108</sup>;

smaller deviations, with a maximum of 4% at 311.5 K and 3.3 mol·dm<sup>-3</sup>. As indicated by the statistical summary, the typical deviations for all the primary data are under 2%. The secondary data which we have included in Fig. 24b and Table 16 show deviations from our correlation which appear similar to those of the primary data, although the scales of the two plots are different. The data of Le Neindre et al.<sup>106</sup> extend to 800 K, well above the upper limit of the primary data. Deviations for these high temperature points range from 2% to 5.6% at the most extreme point, 800 K and 114 MPa. The data of Carmichael et al.<sup>107</sup> were obtained with an apparatus using concentric spheres and exhibit the largest deviations from our correlation; the maximum deviation is 6.6% and occurs near 444 K and 9.7 mol·dm<sup>-3</sup>. The data of Tufeu *et al*.<sup>110</sup> show 2–3.5% deviations along the 500 K isotherm, but give systematic deviations to 6.2% along the critical isochore between 307 and 500 K. The low density isothermal data of Fleeter et al.<sup>54</sup> seem to indicate a smaller slope than those of Roder, as indicated above, and have systematic deviations to nearly 3%. For the single isotherm from Gilmore and Comings<sup>108</sup>, a deviation of about 9% occurs at the highest pressure studied, 304 MPa. The lower pressure isotherm of Leng and Comings<sup>109</sup> agrees with our correlation to within 3.2%.

Note added in final revision: We have examined, but not illustrated, the deviations between our correlation and very recent thermal conductivity data published by Millat et al.<sup>111</sup> and the critical region data from Mostert et al.<sup>112,113</sup> The AAD-% for the 64 data from Ref. 111 is 2.6% with deviations up to 4% at the highest temperatures and lowest densities. The AAD-% for the 109 data in the critical region reported in Ref. 113 is 3.8%. The maximum deviation is 18% for the measurement closest to the critical point; deviations are comparable to those for the correlation developed in Ref. 113. The 9 preliminary measurements tabulated in Ref. 112 have an AAD-% of 5.1% from our correlation and have a maximum deviation of 14% for the point at 305.455 K, closest to the critical temperature. These new data agree with our correlations within combined uncertainties.

We have estimated the uncertainty associated with our correlation for the thermal conductivity of ethane based on these comparisons and on consideration of the experimental technique and error estimates of the researchers. In the gas phase, the large uncertainty in the dilute gas

332

contribution certainly dominates at temperatures below 200 K: the uncertainty in the total thermal conductivity is 10% for the gas at these low temperatures. For temperatures from 200 K to  $T_c$ , the uncertainty in the gas phase thermal conductivity is about 3%. For the compressed liquid, we estimate uncertainties of 2% to the upper pressure limit of 70 MPa. The asymptotically critical region extending from 305 to 306 K with densities between 6 and 7.5 mol·dm<sup>-3</sup> can still give large errors; our value of the thermal conductivity at the critical point is about 1.5  $kW \cdot m^{-1} \cdot K^{-1}$ , not infinity. In the more general critical region bounded by 305 and 307 K from 5 to 8.5 mol·dm<sup>-3</sup>, we estimate the uncertainty as 10%. At supercritical temperatures, the uncertainty is 4% for densities less than 2 mol·dm<sup>-3</sup> and 2% for higher densities. For temperatures between 307 and 360 K, and densities between 5 and 8.5  $mol dm^{-3}$ , the critical enhancement contribution remains substantial, and an uncertainty of 5% is estimated.

#### 5. Conclusions

The correlations we have presented represent improved empirical algebraic representations of the thermodynamic surface and transport properties over a broad range of the fluid states of ethane. The abundance of data for this important and relatively simple molecule has enabled us to determine these correlating equations and to establish limits on their accuracy. Unfortunately, there continues to be some disagreement among experimental measurements, to the extent that data are too often inconsistent when we consider the error bounds reported by the experimenters. We have used careful, conservative judgment when deciding on the relative weights of inconsistent data, and we think that our error estimates of Sec. 4 reflect the actual uncertainties of the correlations. In certain instances, a sparsity of data or a problem with the correlating function itself causes an increase in the uncertainty of predictions based on the correlation. These problems have been discussed in Sec. 4.

Additional measurements on fluid ethane, especially in some of the problem regions and for some of the specific properties mentioned above, will further refine future correlations. For instance, additional PVT data in the highly compressed liquid, derivative and transport data in the critical region, low temperature gas-phase viscosity measurements, and also measurements near and on both liquid and vapor phase boundaries would be extremely useful. For greatest utility in developing correlations, such measurements should adhere strictly to appropriate guidelines concerning the acquisition of primary data. Samples should be well characterized and very pure; absolute determination of all experimental quantities, traceable to relevant standards, is preferred; and the tabulation of original, unsmoothed data, with all experimental correction factors fully discussed or referenced is imperative. Theoretical advances, including improved treatment of the critical region, calculation and incorporation of precise intermolecular potential functions, and a rigorous theory of energy transfer problems for the

thermal conductivity and initial density dependences for both transport properties, will also assist in the development of future correlations.

Within the specified ranges and tolerances, these correlations will allow the user to calculate the important thermodynamic and transport properties of ethane. They should also be useful for additional studies, such as for generalized corresponding states models and mixture calculations, especially in conjunction with our previous study of methane.

## 6. Acknowledgments

We gratefully acknowledge support from the Standard Reference Data Program of the National Institute of Standards and Technology. We thank R.D. McCarty and R.T. Jacobsen for helpful discussions and their careful review of parts of this manuscript. We thank G.C. Straty for discussions concerning some of the experimental data and D.E. Diller for providing some data prior to their publication. We also thank B.S. Coultrip for assistance with data compilation. The anonymous reviewers selected by the Journal have provided numerous suggestions which have greatly improved this paper.

### 7. References

- <sup>1</sup>D. G. Friend, J. F. Ely, H. Ingham, J. Phys. Chem. Ref. Data 18, 583 (1989).
- <sup>2</sup>D. G. Friend, J. F. Ely, H. Ingham, "Tables of the Thermophysical Properties of Methane," National Institute of Standards and Technology, U. S., Technical Note 1325 (1989).
- <sup>3</sup>D. G. Friend, J. F. Ely, H. Ingham, "Tables of the Thermophysical Properties of Ethane," National Institute of Standards and Technology, U. S., Technical Note, 1346, to be published.
- <sup>3a</sup>D. G. Friend, J. F. Ely, H. Ingham, "Tables of Experimental Data Used for the Correlation of the Thermophysical Properties of Ethane," National Institute of Standards and Technology, U. S., Internal Report 3953, to be published.
- <sup>4</sup>R. D. Goodwin, H. M. Roder, and G. C. Straty, National Bureau of Standards, U. S., Technical Note 684 (1976).
- <sup>5</sup>V. V. Sychev, A. A. Vasserman, A. D. Kozlov, V. A. Zagoruchenko, G. A. Spiridonov, and V. A. Tsymarny, *Thermodynamic Properties of Ethane* (Hemisphere Publishing, Washington, 1987), T. B. Selover Jr., English Language Edition Ed.
- <sup>6</sup>R. Schmidt and W. Wagner, Fluid Phase Equil. 19, 175 (1985).
- <sup>6a</sup>U. Setzmann and W. Wagner, Int. J. Thermophys. 10, 1103 (1989).
- <sup>7</sup>J. O. Hirschfelder, C. F. Curtiss, and R. B. Bird, *Molecular Theory of Cases and Liquids* (Wiley, New York, 1967).
- <sup>8</sup>M. Klein, H. J. M. Hanley, F. J. Smith, and P. Holland, Natl. Bur. Stand. (U. S. ), Monogr. 47 (1974).
- <sup>9</sup>G. A. Olchowy and J. V. Sengers, Phys. Rev. Lett. **61**, 15 (1988), Int. J. Thermophys. **10**, 417 (1989) and personal communication, Univ. Maryland, Inst. Phys. Sci. and Tech., College Park, Md, 1988.
- <sup>10</sup>B. A. Younglove and J. F. Ely, J. Phys. Chem. Ref. Data 16, 577 (1987).
- <sup>11</sup>J. Chao, R. C. Wilhoit, and B. J. Zwolinski, J. Phys. Chem. Ref. Data 2, 427 (1973).
- <sup>12</sup>R. D. Goodwin, J. Research Nat. Bur. Stand. 75A, 15 (1971).
- <sup>13</sup>H. J. M. Hanley and M. Klein, National Bureau of Standards, U. S., Technical Note 628 (1972).
- <sup>14</sup>R. C. Reid, J. M. Prausnitz, and B. E. Poling, *The Properties of Gases and Liquids* (McGraw-Hill, New York, 4th Ed., 1987).
- <sup>14a</sup>J. C. Rainwater and D. G. Friend, Phys. Rev. A 36, 4062 (1987).

- <sup>15</sup>J. V. Sengers, R. S. Basu, and J. M. H. Levelt Sengers, National Aeronautics and Space Administration, U. S., NASA Contractor Report 3424 (1981).
- <sup>16</sup>E. R. Cohen and B. N. Taylor, J. Research Nat. Bur. Stand. **92**, 85 (1987).
- <sup>17</sup>IUPAC Commission on Atomic Weights and Isotopic Abundances, Pure Appl. Chem. **60**, 842 (1988).
- <sup>18</sup>IUPAC Commission on Atomic Weights and Isotopic Abundances, Pure Appl. Chem. **52**, 2349 (1980).
- <sup>19</sup>F. Pavese, J. Chem. Thermodynamics 10, 369 (1978).
- <sup>20</sup>R. E. Bedford, G. Bonnier, H. Maas, and F. Pavese, Metrologia 20, 145 (1984).
- <sup>21</sup>G. C. Straty and R. Tsumura, J. Chem. Phys. 64, 859 (1976).
- <sup>22</sup>D. R. Douslin and R. H. Harrison, J. Chem. Thermodynamics 5,491 (1973).
- <sup>23</sup>J. V. Sengers, personal communication, Univ. Maryland, Inst. Phys. Sci. and Tech., College Park, Md, 1988.
- <sup>24</sup>H. J. Strumpf, A. F. Collings, and C. J. Pings, J. Chem. Phys. **60**, 3109 (1974).
- <sup>24a</sup>M. Burton and D. Balzarini, Canad. J. Phys. 52, 2011 (1974).
- <sup>24b</sup>F. J. Wegner, Phys. Rev. B 5, 4529 (1972).
- <sup>24c</sup>M. W. Pestak, R. E. Goldstein, M. H. W. Chan, J. R. de Bruyn, D. Balzarini, and N. W. Ashcroft, Phys. Rev. B 36, 599 (1987).
- <sup>25</sup>L. J. Van Poolen, R. T Jacobsen, and M. Jahangiri, Int. J. Thermophys. 7, 513 (1986).
- <sup>26</sup>H. J. M. Hanley, K. E. Gubbins, and S. Murad, J. Phys. Chem. Ref. Data, 6, 1167 (1977).
- <sup>27</sup>A. Boushehri, J. Bzowski, J. Kestin, and E. Mason, J. Phys. Chem. Ref. Data 16, 445 (1987).
- <sup>28</sup>F. M. Mourits and F. H. Rummens, Canad. J. Chem. 55, 3007 (1977).
- <sup>29</sup>D. D. Wagman, W. H. Evans, V. B. Parker, R. H. Schumm, I. Halow, S. M. Bailey, K. L. Churney, and R. L. Nuttall, J. Phys. Chem. Ref. Data 11, Sup. 2 (1982).
- <sup>30</sup>K. M Pamidimukkala, D. Rogers, and G. B. Skinner, J. Phys. Chem. Ref. Data 11, 83 (1982).
- <sup>30a</sup>TRC Thermodynamic Tables-Hydrocarbons, Tables 23–3–1. 200 r-x, pp. 1350 r-x, Thermodynamics Research Center, Texas A&M University, College Station, TX (Oct. 31, 1985).
- <sup>31</sup>A. K. Pal, G. A. Pope, Y. Arai, N. F. Carnahan, and R. Kobayashi, J. Chem. Eng. Data 21, 394 (1976); unsmoothed data of Pal reported in Ref. [32] were preferentially used; additional very small corrections based on personal communication (ca. 1976) were also applied. The complete set of data used is reported in Ref. [3].
- <sup>32</sup>G. A. Pope, Ph. D. Thesis (Dept. Chem. Eng. ), Rice University (1971).
- <sup>33</sup>W. T. Ziegler, B. S. Kirk, J. C. Mullins, and A. R. Berquist, Tech. Report No. 2, Proj. A-764, Eng. Expt. Sta., Georgia Inst. Tech., Atlanta, GA (Dec. 1964); the data were used as adjusted in Ref. [4].
- <sup>34</sup>G. C. Straty and R. Tsumura, J. Research Nat. Bur. Stand. 80A, 35 (1976).
- <sup>35</sup>P. Sliwinski, Z. Phys. Chem. Neue Folge 63, 263 (1969).
- <sup>36</sup>C. H. Chui and F. B. Canfield, Trans. Faraday Soc. 67, 2933 (1971); M. Y. Shana'a and F. B. Canfield, Trans. Faraday Soc. 64, 2281 (1968).
- <sup>37</sup>J. B. Rodosevich and R. C. Miller, A. I. Ch. E. J. 19, 729 (1973).
- <sup>38</sup>W. M. Haynes and M. J. Hiza, J. Chem. Thermodynamics 9, 179 (1977).
   <sup>39</sup>C. R. McClune, Cryogenics 16, 289 (1976).
- <sup>40</sup>J. E. Orrit and J. M. Laupretre, Adv. Cryo. Eng. 23, 573 (1978); J. E. Orrit and J. F. Olives, "*Density of liquefied natural gas and its components*," distributed at Fourth International Conference on Liquefied Natural Gas, Algeria (1974).
- <sup>41</sup>S. L. S. Jacoby, J. S. Kowalilc, and J. T. Pizzo, *Iterative Methods for Nonlinear Optimization* (Prentice-Hall, Englewood Cliffs, N. J. 1972).
- <sup>42</sup>A. Michels, W. Van Straaten, and J. Dawson, Physica 20, 17 (1954).
  <sup>43</sup>G. A. Pope, P. S. Chappelear, and R. Kobayashi, J. Chem. Phys. 59, 423 (1973).
- <sup>44</sup>H. Mansoorian, K. R. Hall, and P. T. Eubank, Proc. 7th Symp. on Thermophysical Properties, A. Cezairliyan, ed., Amer. Soc. Mech. Engin., New York (1977) p. 456.
- <sup>45</sup>M. L. McGlashan and D. J. Potter, Proc. Roy. Soc. (London) A267, 478 (1962).

- <sup>46</sup>W. R. Parrish, Fluid Phase Equil. 18, 279 (1984).
- <sup>47</sup>H. M. Roder, J. Research Nat. Bur. Stand. 80A, 739 (1976).
- <sup>48</sup>A. Furtado, Ph. D. Thesis (Dept. Chem. Eng.), University of Michigan, Ann Arbor, MI (1973).
- <sup>49</sup>K. Bier, J. Kunze, and G. Maurer, J. Chem. Thermodynamics 8, 857 (1976).
- <sup>49a</sup>K. Bier, J. Kunze, G. Maurer, and H. Sand, J. Chem. Engin. Data 21, 5 (1976).
- <sup>50</sup>R. Tsumura and G. C. Straty, Cryogenics 17, 195 (1977).
- <sup>51</sup>V. E. Terres, W. Jahn, and H. Reissmann, Brennstoff-Chemie 38, 129 (1957).
- <sup>52</sup>J. Kestin, H. E. Khalifa, and W. A. Wakeham, J. Chem. Phys. 66, 1132 (1977).
- <sup>53</sup>J. Kestin, S. T. Ro, and W. A. Wakeham, Trans. Far. Soc. 67, 2308 (1971).
- <sup>54</sup>R. Fleeter, J. Kestin, and W. A. Wakeham, Physica 103A, 521 (1980).
- <sup>55</sup>D. E. Diller and J. M. Saber, Physica 108A, 143 (1981); and D. E. Diller, Proc. 8th Symp. on Thermophysical Properties, J. V. Sengers, ed., Amer. Soc. Mech. Engin., New York (1982) p. 219.
- <sup>56</sup>D. E. Diller, High Temp.-High Pressures 21, 613 (1989).
- <sup>57</sup>L. T. Carmichael and B. H. Sage, J. Chem. Engin. Data 8, 94 (1963).
- <sup>58</sup>H. M. Roder and C. A. Nieto de Castro, High Temp.-High Pressures 17, 453 (1985).
- <sup>59</sup>R. C. Prasad and J. E. Venart, Int. J. Thermophys. 5, 367 (1984).
- <sup>60</sup>H. M. Roder, National Bureau of Standards, U. S., Interagency Report, NBSIR 84–3006, (1984).
- <sup>61</sup>P. Desmarest and R. Tufeu, Int. J. Thermophys. 8, 293 (1987).
- <sup>62</sup>R. D. Goodwin, National Bureau of Standards, U. S., Interagency Report, NBSIR 74–398, (1974).
- <sup>63</sup>G. F. Carruth and R. Kobayashi, J. Chem. Engin. Data 18, 115 (1973).
   <sup>64</sup>J. Regnier, J. Chimie Physique 69, 942 (1972).
- <sup>65</sup>L. Djordjevich and R. A. Budenholzer, J. Chem. Engin. Data 15, 10 (1970).
- <sup>66</sup>V. M. Miniovich and G. A. Sorina, Russ. J. Phys. Chem. 45, 306 (1971).
- <sup>67</sup>P. T. Eubank, Adv. Cryo. Engin. 17, 270 (1971).
- <sup>68</sup>J. Klosek and C. McKinley, *Proc. 1st Inter. Conf. on LNG*, Paper 22, Chicago, (1968).
- <sup>69</sup>L. C. Kahre, J. Chem. Engin. Data 18, 267 (1973).
- <sup>70</sup>J. R. Tomlinson, Natural Gas Processors Association Tech. Publ. TP-1, Tulsa, OK (1971).
- <sup>71</sup>R. A. Perkins, H. M. Roder, D. G. Friend, and C. A. Nieto de Castro, "The Thermal Conductivity and Heat Capacity of Fluid Nitrogen," Physica, to be published.
- <sup>72</sup>F. Porter, J. Am. Chem. Soc. 48, 2055 (1926).
- <sup>73</sup>K. Buhner, G. Maurer, and E. Bender, Cryogenics 21 157 (1981).
- <sup>74</sup>A. Eucken and A. Parts, Z. Phys. Chem. **B20**, 184 (1933).
- <sup>75</sup>J. H. Dymond and E. B. Smith, *The Virial Coefficients of Pure Gases* and *Mixtures*, Clarendon Press, Oxford (1980).
- <sup>76</sup>A. E. Hoover, I. Nagata, T. W. Leland, and R. Kobayashi, J. Chem. Phys. **48**, 2633 (1968).
- <sup>77</sup>R. D. Gunn, M. S. Thesis, Univ. Calif. (Berkeley) 1958.
- <sup>78</sup>J. A. Huff and T. M. Reed III, J. Chem. Engin. Data, 8, 306 (1963).
   <sup>79</sup>V. K. Strein, R. N. Lichtenthaler, B. Schramm, and Kl. Schäfer, Ber. Bunsenges. Phys. Chem. 75, 1308 (1971).
- <sup>80</sup>R. L. Powell, W. J. Hall, C. H. Hyink, L. L. Sparks, G. W. Burns, M. G. Scroger, and H. H. Plumb, *Thermocouple Reference Tables Based on the IPTS-68*, National Bureau of Standards, (U. S. ), Monograph 125 (1974).
- <sup>81</sup>J. V. Sengers and J. M. H. Levelt Sengers, Ann. Ref. Phys. Chem. 37, 189 (1986).
- <sup>82</sup>J. A. Beattie, C. Hadlock, N. Poffenberger, J. Chem. Phys. 3, 93 (1935).
- <sup>82a</sup>G. J. Besserer and D. B. Robinson, J. Chem. Engin. Data 18, 137 (1973).
- <sup>83</sup>Ye. A. Golovskiy, E. P. Mitsevich, V. A. Tsymarnyy, "Measurement of the density of ethane at 90.24-270. 21 K and at pressures to 604. 09 bar," VNIIEGazprom Dep. No. 39M USSR (1978); data have been tabulated in Ref. [5].

- <sup>84</sup>W. W. R. Law, "A continuously weighed pycnometer providing densities for CO<sub>2</sub> + Ethane mixtures between 240 and 350 K at pressures to 35 MPa," Ph. D. Thesis, Texas A&M University, (1986).
- <sup>85</sup>N. E. Khazanova and E. E. Sominskaya, Russ. J. Phys. Chem. 45, 87 (1971).
- <sup>86</sup>V. M. Miniovich and G. A. Sorina, Teplofiz. Svoistva Veshchestv Mater., 6, 134 (1973).
- <sup>86</sup><sup>a</sup>H. H. Reamer, R. H. Olds, B. H. Sage, and W. N. Lacey, Ind. Engin. Chem. 36, 957 (1944)
- <sup>87</sup>J. R. Tomlinson, "Liquid Densities of Ethane, Propane, and Ethane-Propane Mixtures," Tech. Pub. TP-1, Natural Gas Processors Ass., Tulsa, OK (1971).
- <sup>88</sup>C. B. Wallace Jr., I. H. Silberberg, J. J. McKetta, Petrol. Refiner, Hydrocarbon Processing 43 No. 10, 177 (1964).
- <sup>89</sup>T. Miyazaki, A. V. Hejmaki, and J. E. Powers, J. Chem. Thermodynamics 12, 105 (1980).
- <sup>50</sup>R. Wiebe, K. H. Hubbard, and M. J. Brevoort, J. Am. Chem. Soc. **52**, 611 (1930).
- <sup>91</sup>R. K. Witt and J. D. Kemp, J. Am. Chem. Soc. 59, 273 (1937).
- <sup>92</sup>E. Vangeel, Katholieke Universiteit, Leuven, Belgium, private communication to D. Diller, NIST., (1974); data were tabulated in Table 17 of Ref. [4].
- <sup>93</sup>G. R. Poole and R. A. Aziz, Can. J. Phys. 50, 721 (1972).
- <sup>94</sup>V. M. Trautz and K. G. Sorg, Ann. Phys. 10, 81 (1931).
- <sup>95</sup>Y. Abe, J. Kestin, and H. E. Khalifa and W. A. Wakeham, Physica 93A, 155 (1978).
- <sup>96</sup>H. Iwasaki and M. Takahashi, J. Chem. Phys. 74, 1930 (1981).
- <sup>97</sup>A. G. De Rocco and J. O. Halford, J. Chem. Phys. 28, 1152 (1958).
- 98P. M. Craven and J. D. Lambert, Proc. Roy. Soc. 205A, 439 (1951).
- <sup>99</sup>J. D. Lambert, K. J. Cotton, M. W. Pailthorpe, A. M. Robinson, J. Scrivins, W. R. F. Vale, and R. M. Young, Proc. Roy. Soc. 231A, 280 (1955).

- <sup>100</sup>E. Eakin, K. E. Starling, J. P. Dolan, and R. T. Ellington, J. Chem. Engin. Data 7, 33 (1962).
- <sup>101</sup>Y. S. Touloukian, S. C. Saxena, and P. Hestermans, *Viscosity*, Thermo-physical Properties of Matter, Vol. 11, Plenum, New York (1975).
- <sup>102</sup>J. D. Baron, J. G. Roof, and F. W. Wells, J. Chem. Engin. Data 4, 283 (1959).
- <sup>103</sup>G. W. Swift, J. Lohrenz, and F. Kurata, A. I. Ch. E. J. 6, 415 (1960).
- <sup>104</sup>K. Stephan and K. Lucas, *Viscosity of Dense Fluids*, (Plenum, N. Y., 1979).
- <sup>104a</sup>A. A. Clifford, E. Dickinson, and P. Gray, J. Chem. Soc. (London), Faraday Trans. **172**, 1997 (1976).
- <sup>105</sup>L. V. Yakush, N. A. Vanicheva, and L. S. Zaitseva, Inz.-Fiz. Zhurnal, **37**, 1071 (1979); tr. in J. Eng. Phys. **37**, 1071 (1979).
- <sup>106</sup>B. Le Neindre, R. Tufeu, P. Bury, P. Jahannin, and B. Vodar, in *Proc.* 8th Int. Conf. Therm. Conduct., C. Y. Ho and R. E. Taylor, eds. (Plenum, N. Y., 1969), p. 229.
- <sup>107</sup>L. T. Carmichael, V. Berry, and B. H. Sage, J. Chem. Engin. Data 8, 281 (1963).
- <sup>108</sup>T. F. Gilmore and E. W. Comings, A. I. Ch. E. J. 12, 1172 (1966).
- <sup>109</sup>D. E. Leng and E. W. Comings, Ind. Engin. Chem. 49, 2042 (1957).
   <sup>110</sup>R. Tufeu, Y. Garrabos, and B. Le Neindre, in *Proc. 16th Conf. Therm. Conduct.*, D. C. Larsen, ed. (Plenum, N. Y., 1983), p. 605.
- <sup>111</sup>J. Millat, M. Ross, W. A. Wakeham, and M. Zalaf, Int. J. Thermophys. 9, 481 (1988).
- <sup>112</sup>R. Mostert, H. R. van den Berg, and P. S. van der Gulik, Int. J. Thermophys. 10, 409 (1989).
- <sup>113</sup>R. Mostert, H. R. van den Berg, P. S. van der Gulik, and J. V. Sengers, J. Chem. Phys. **92**, 5454 (1990).

## FRIEND, INGHAM, AND ELY

# 8. Appendix

 T K	A <sup>id</sup> kI-mol <sup>-1</sup>	H <sup>id</sup> kJ·mol <sup>-1</sup>	S <sup>id</sup> J·mol <sup>-1</sup> ·K <sup>-1</sup>	$C_p^{id}$ J·mol <sup>-1</sup> ·K <sup>-1</sup>	η₀ μPa·s	λ <sub>0</sub> mW·m <sup>-1</sup> ·K <sup>-1</sup>	
100.	-15.846	3.384	183.99	35.698	3.32	3.46	
110.	-17.786	3.744	187.41	36.249	3.60	4.04	
120.	-19.759	4.109	190.59	36.817	3.90	4.65	
130.	-21.763	4.480	193.56	37.401	4.19	5.28	
140.	-23.796	4.857	196.36	38.003	4.50	5.94	
150.	-25.856	5.241	199.00	38.628	4.80	6.62	
160.	-27.942	5.630	201.51	39.279	5.11	7.33	
170.	-30.053	6.026	203.91	39.961	5.42	8.08	
180.	-32.186	6.429	206.22	40.680	5.73	8.85	
190.	-34.343	6.840	208.44	41.439	6.04	9.65	
200.	-36.521	7.258	210.58	42.243	6.35	10.49	
210.	-38.721	7.685	212.66	43.092	6.66	11.36	
220.	-40.941	8.120	214.69	43.989	6.97	12.28	
230.	-43.181	8.565	216.67	44.934	7.28	13.23	
240.	-45.440	9.019	218.60	45.924	7.59	14.23	
250.	-47.719	9.484	220.49	46.959	7.89	15.26	
260.	-50.016	9.959	222.36	48.036	8.20	16.35	
270.	-52.332	10.444	224.19	49.151	8.50	17.47	
280.	-54.666	10.942	226.00	50.302	8.80	18.65	
290.	-57.018	11.451	227.79	51.484	9.09	19.86	
300.	-59.388	11.971	229.55	52.693	9.39	21.13	
310.	-61.776	12.505	231.30	53.926	9.68	22.43	
320.	-64.180	13.050	233.03	55.178	9.97	23.78	
330.	-66.602	13.608	234.75	56.446	10.25	25.17	
340.	-69.042	14.179	236.45	57.727	10.54	26.60	
350.	-71.498	14.763	238.14	59.017	10.82	28.07	
360.	-73.971	15.359	239.82	60.313	11.10	29.58	
370.	-76.460	15.969	241.49	61.612	11.38	31.12	
380.	-78.967	16.592	243.15	62.913	11.65	32.70	
390.	-81.490	17.227	244.81	64.212	11.92	34.31	
400.	-84.029	17.876	246.45	65.507	12.19	35.95	
410.	-86.585	18.537	248.08	66.798	12.46	37.63	
420.	-89.157	19.212	249.71	68.082	12.72	39.33	
430.	-91.745	19.899	251.32	69.357	12.99	41.05	
440.	-94.350	20.599	252.93	70.624	13.25	42.81	
450.	-96.970	21.311	254.53	71.880	13.50	44.58	
460.	-99.607	22.036	256.13	73.126	13.76	46.38	
470.	-102.259	22.774	257.71	74.360	14.01	48.20	
480.	-104.927	23.524	259.29	75.582	14.26	50.04	
490.	-107.611	24.285	260.86	76.791	14.51	51.90	
500.	-110.311	25.059	262.43	77.987	14.76	53.78	

TABLE A1. Properties of ideal gas at 0.1 MPa and dilute gas transport properties

The ideal gas values of the molar Helmholtz energy, enthalpy, entropy, and isobaric heat capacity are evaluated from Eq. (3). The conversion from atmospheric pressure to 0.1 MPa affects the values of  $A^{1d}$  and  $S^{1d}$ . The dilute gas viscosity is from Eq. (10a) and the dilute gas thermal conductivity is from Eq. (13a).

TABLE A2. Properties along saturation boundary

Т к	P <sub>o</sub> MPa	$\rho_{\sigma L}$	ρ <sub>σ</sub> ν mol·dm <sup>-3</sup>	$C_{\sigma L}$ J·mol <sup>-1</sup> ·K <sup>-1</sup>	$W_{\sigma L}$ m·s <sup>-1</sup>	Պσ∟ µPa∙s	$\lambda_{\sigma L}$ mW·m <sup>-1</sup> ·K <sup>-1</sup>
92.	0.17E-5	21.61	0.227E-5	67.74	1987.2	1193.00	254.4
94.	0.28E-5	21.54	0.364E-5	68.59	1975.7	1099.01	252.8
96.	0.46E-5	21.47	0.573E-5	69.24	1963.5	1016.05	251.3
98.	0.72E - 5	21.39	0.883E-5	69.73	1950.8	942.48	249.7
100.	0.11E-4	21.32	0.133E-4	70.09	1937.6	876.96	248.1
102.	0.17E-4	21.25	0.198E-4	70.34	1924.2	818.40	246.4
104	0.25E - 4	21.18	0.289E - 4	70.51	1910.4	765.84	244.7
106	0.37E - 4	21.11	0.415E - 4	70.61	1896.5	718.52	243.0
108	0.57E - 4	21.11	0.586E - 4	70.66	1887.4	675 76	241 3
110	0.75E 4	21.05	0.90012 + 0.817E - 4	70.68	1868 7	637.00	239.5
110.	0.75E-4	20.90	0.0172 - 4	70.00	1000.2	057.00	207.0
112.	0.10E - 3	20.89	0.112E-3	70.66	1853.9	601.75	237.7
114.	0.14E - 3	20.82	0.152E - 3	/0.63	1839.5	569.61	235.9
116.	0.20E - 3	20.74	0.204E – 3	70.58	1825.1	540.22	234.1
118.	0.27E - 3	20.67	0.271E - 3	70.52	1810.6	513.27	232.2
120.	0.35E-3	20.60	0.356E - 3	70.46	1796.0	488.50	230.4
122.	0.47E-3	20.52	0.462E-3	70.39	1781.3	465.67	228.5
124.	0.61E-3	20.45	0.594E-3	70.33	1766.7	444.57	226.6
126.	0.79E - 3	20.38	0.756E-3	70.27	1752.0	425.04	224.6
128.	0.10E - 2	20.30	0.955E-3	70.22	1737.2	406.92	222.7
130.	0.13E - 2	20.23	0.120E - 2	70.18	1722.4	390.06	220.8
132	0.16E - 2	20.15	0.149 E - 2	70 14	1707 6	374 36	218.8
132.	0.10E = 2	20.15	0.183E - 2	70.12	1692.8	359.69	216.9
134.	0.20L = 2	20.00	0.225 - 2	70.12	1677.0	345.08	210.9
130.	0.23E - 2	20.00	0.223E - 2	70.10	1677.9	242.20	214.5
156.	0.31E - 2	19.95	0.273E - 2	70.09	1649.1	333.12	212.9
140.	0.38E - 2	19.85	0.330E-2	/0.10	1048.1	321.05	210.9
142.	0.47E-2	19.78	0.397E-2	70.11	1633.2	309.70	209.0
144.	0.56E - 2	19.70	0.474E – 2	70.14	1618.2	299.00	207.0
146.	0.68E - 2	19.62	0.562E - 2	70.17	1603.2	288.91	205.0
148.	0.81E - 2	19.55	0.664 E - 2	70.22	1588.2	279.38	203.0
150.	0.97E – 2	19.47	0.780E - 2	70.27	1573.2	270.35	201.0
152.	0.011	19.39	0.912E – 2	70.34	1558.2	261.80	199.0
154.	0.013	19.32	0.011	70.42	1543.2	253.68	197.0
156	0.016	19.74	0.012	70.51	1528.2	245.96	195.0
158	0.010	19.24	0.012	70.60	1513.1	238.62	193.0
160.	0.021	19.08	0.016	70.71	1498.1	231.63	191.0
160	0.025	10.00	0.010	70.92	1492.0	224.05	190.0
102.	0.025	19.00	0.019	70.65	1465.0	224.93	189.0
164.	0.029	18.92	0.021	70.95	1467.9	218.58	187.1
166.	0.033	18.84	0.024	71.09	1452.8	212.48	185.1
168.	0.038	18.76	0.027	71.24	1437.7	206.65	183.1
170.	0.043	18.68	0.031	71.39	1422.6	201.06	181.2
172.	0.049	18.60	0.035	71.56	1407.5	195.69	179.2
174.	0.055	18.52	0.039	71.73	1392.4	190.55	177.3
176.	0.062	18.44	0.044	71.91	1377.2	185.60	175.3
178.	0.070	18.36	0.049	72.11	1362.1	180.84	173.4
180.	0.079	18.28	0.054	72.31	1346.9	176.25	171.4
182	0.088	18.19	0.060	72.52	1331 7	171 83	169 5
184	0.098	18 11	0.066	72.74	1316.6	167 57	167.6
194	0.000	18.02	0.000	72.07	1301 4	167.57	165 7
199.	0.109	17.03	0.075	72.71	1796 1	103.40	162.9
100.	0.122	17.94	0.001	13.21	1200.1	159.48	103.8
190.	0.135	17.80	0.089	/3.40	12/0.9	155.04	101.9
192.	0.149	17.77	0.098	73.72	1255.6	151.92	160.0
194.	0.164	17.69	0.107	73.99	1240.4	148.32	158.1
196.	0.181	17.60	0.117	74.27	1225.1	144.83	156.2
198.	0.198	17.51	0.127	74.56	1209.7	141.45	154.4
200.	0.217	17.42	0.139	74.86	1194.4	138.17	152.5

TABLE A2. Properties along saturation boundary - Continued

				_			
Т	$P_{\sigma}$	ρ <sub>σ</sub> ι	ρσν	CoL	WoL	η <sub>σL</sub>	λσL
K	MPa	mol·dm <sup>-3</sup>	mol·dm <sup>-3</sup>	J-mol <sup>-1</sup> ·K <sup>-1</sup>	m·s <sup>-1</sup>	μPa-s	mW·m <sup>−1</sup> ·K <sup>−1</sup>
202.	0.238	17.33	0.151	75.17	1179.0	134.98	150.7
204	0.260	17.24	0 164	75 50	1163.6	131.89	148.8
206	0.200	17.15	0.177	75.93	1149.0	100.00	147.0
200.	0.205	17.15	0.177	75.65	1140.2	120.00	147.0
208.	0.308	17.06	0.192	76.18	1132.7	125.95	145.2
210.	0.334	16.97	0.208	76.53	1117.2	123.10	143.4
212.	0.362	16.88	0.224	76.90	1101.7	120.32	141.6
214.	0.392	16.78	0.241	77 29	1086 1	117.62	139.8
216	0.423	16 69	0.260	77.69	1070.4	114 98	138.0
210.	0.457	16.50	0.270	79.10	1054.9	117.70	136.0
210.	0.407	10.39	0.279	78.10	1034.0	112.41	150.2
220.	0.492	10.50	0.300	/8.52	1039.0	109.90	134.5
222	0.530	16.40	0.322	78.96	1023.3	107.45	132.7
224.	0.569	16.30	0.345	79.42	1007.4	105.06	131.0
226.	0.611	16.20	0.369	79.90	991.5	102.72	129.3
228	0.654	16 10	0 305	80 30	075.6	100.43	127.5
220.	0.004	16.00	0.333	80.09	975.0	100.45	127.5
250.	0.700	10.00	0.422	80.90	939.0	98.19	125.8
232.	0.749	15.90	0.451	81 43	943 5	96.00	124 1
234	0.800	15 70	0.491	01.45	077.3	02.00	100 4
254.	0.000	15.79	0.401	01.90	921.5	95.65	122.4
230.	0.853	15.68	0.512	82.55	911.0	91.74	120.7
238.	0.909	15.58	0.546	83.14	894.7	89.68	119.0
240.	0.967	15.47	0.581	83.76	878.3	87.65	117.4
242.	1.028	15.36	0.618	84 41	861.8	85.67	115.7
244.	1.092	15 24	0.657	85.08	845 1	83 72	114.1
246	1 150	15 13	0.608	85.70	878 4	81 80	117.1
240.	1.139	15.15	0.030	05.19	020.4	70.01	112.4
240.	1.229	15.01	0.741	80.52	811.0	79.91	110.8
250.	1.301	14.89	0.787	87.29	794.6	78.06	109.1
252	1 377	14 77	0.835	88 11	777 5	76 23	107.5
254	1 456	14.65	0.000	88.06	760.3	70.25	107.5
254.	1.430	14.50	0.000	80.70	700.5	74.43	103.9
250.	1.538	14.52	0.938	89.86	742.9	/2.66	104.3
258.	1.623	14.39	0.994	90.81	725.4	70.91	102.7
260.	1.712	14.26	1.053	91.82	707.7	69.19	101.1
262.	1.804	14.13	1,116	92.89	689.8	67.49	99.5
264	1 900	13.99	1 182	94.03	671 7	65.80	07.0
264.	1,000	12.95	1.102	05.05	652 4	64.14	06.2
200.	1.999	13.03	1.232	95.25	033.4	04.14	90.3
208.	2.103	13.70	1.325	96.55	634.9	62.48	94.8
270.	2.210	13.55	1.404	97.96	616.2	60.85	93.2
272.	2.321	13.40	1.487	99.49	597.2	59.22	91.6
274.	2,436	13.24	1.575	101.15	577.9	57.61	90.0
276	2 555	13.07	1.670	102.07	558.2	56.00	80.0
270.	2.333	12.07	1.0/0	102.97	520.2	50.00	00.J
270.	2.070	12.70	1.//1	104.70	330.3	34.39	00.9
280.	2.806	12.72	1.879	107.22	517.9	52.79	85.4
282.	2.938	12.54	1.995	109.73	497.1	51.18	83.8
284.	3,075	12.34	2,121	112.58	475.8	49.56	82.2
286	3 216	12 13	2 257	115 87	454.0	17.02	80 7
200.	2 262	11.02	2.231	110.07	A21 4	71.73	70.1
200.	3.303	11.94	2.400	119.72	431.0	40.28	/9.1
290.	3.314	11.08	2.570	124.32	408.5	44.60	77.6

TABLE A2.	Properties	along	saturation	boundary	-	Continued
-----------	------------	-------	------------	----------	---	-----------

T K	Р <sub>о</sub> MPa	ρ <sub>σL</sub> mol·dm <sup>-3</sup>	ρ <sub>σ</sub> ν mol·dm <sup>-3</sup>	C <sub>σL</sub> J•mol <sup>−1</sup> •K <sup>−1</sup>	<i>W</i> <sub>σL</sub> m·s <sup>−1</sup>	η₀⊥ μ₽a•s	$\lambda_{\sigma L}$ mW·m <sup>-1</sup> ·K <sup>-1</sup>
292.	3.671	11.43	2.753	129.97	384.7	42.87	76.1
294.	3.834	11.16	2.959	137.15	360.1	41.09	74.6
296.	4.002	10.85	3.195	146.73	334.4	39.22	73.2
298.	4.176	10.51	3.472	160.40	307.4	37.22	72.0
300.	4.356	10.10	3.813	182.06	278.4	35.01	71.3
302.	4.543	9.59	4.262	223.66	246.4	32.44	72.0
304.	4.738	8.82	4.969	354.78	209.4	28.97	79.0

Values of the pressure, density of the saturated liquid, density of the saturated vapor, heat capacity, sound speed, viscosity, and thermal conductivity along the two phase liquid-vapor coexistence curve. The quantities  $P_{\sigma}$ ,  $\rho_{\sigma\nu}$ , and  $\rho_{\sigma L}$  are from the ancillary equations, Eqs.(4-6). The heat capacity along the saturated boundary is from the equation in Table 7; the sound speed is also taken from Table 7 but the density argument is for the saturated liquid and is taken from column 3 of this table. The viscosity and thermal conductivity at saturation are from Eqs. (8) and (9) [with the terms evaluated from Eqs. (10-20)]; again the density input is from column 3 of this table.

TABLE A.3. Properties of ethane in the single-pl	hase regio	on
--	------------	----

T	P	0	 H	S	 Сv	C <sub>P</sub>	W	m	λ
K	MPa	mol·dm <sup>-3</sup>	kJ/mol	J/(mol·K)	J/(mol·K)	J/(mol·K)	m·s <sup>-1</sup>	µPa·s	mW/(m·K)
100.	0.1	21.33	- 14.221	83.60	48.15	70.11	1938.7	878.68	248.2
100.	0.5	21.33	- 14.206	83.57	48.15	70.09	1940.5	881.47	248.4
100.	1.0	21.34	- 14.186	83.53	48.16	70.07	1942.6	884.97	248.6
100.	2.0	21.35	- 14.147	83.45	48.17	70.02	1947.0	892.03	249.1
100.	5.0	21.39	- 14.030	83.22	48.22	69.89	1959.8	913.68	250.5
100.	10.0	21.45	13.835	82.84	48.28	69.67	1980.8	951.47	252.7
100.	20.0	21.57	- 13.444	82.10	48.42	69.27	2021.6	1034.29	257.2
100.	30.0	21.68	-13.052	81.39	48.56	68.92	2060.7	1128.62	261.6
100.	40.0	21.79	- 12.660	80.71	48.70	68.61	2098.4	1237.38	265.8
100.	50.0	21.90	- 12.268	80.05	48.84	68.32	2134.9	1364.46	270.0
100.	60.0	22.00	- 11.876	79.42	48.97	68.06	2170.3	1515.27	274.1
110.	0.1	20.96	- 13.515	90.33	47.39	70.82	1868.7	637.50	239.6
110.	0.5	20.97	- 13.500	90.30	47.41	70.80	1870.5	639.34	239.8
110.	1.0	20.98	- 13.480	90.26	47.42	70.78	1872.7	641.66	240.0
110.	2.0	20.99	- 13.442	90.17	47.45	70.74	1877.2	646.31	240.5
110.	5.0	21.03	- 13.326	89.93	47.55	70.62	1890.3	660.54	242.1
110.	10.0	21.10	- 13.133	89.53	47.71	70.44	1911.8	685.13	244.6
110.	20.0	21.23	- 12.745	88.76	48.01	70.11	1953.4	737.99	249.5
110.	30.0	21.36	- 12.356	88.02	48.30	69.84	1993.3	796.59	254.2
110.	40.0	21.47	- 11.967	87.31	48.58	69.60	2031.7	862.12	258.9
110.	50.0	21.59	- 11.577	86.64	48.86	69.40	2068.8	936.09	263.5
110.	60.0	21.69	- 11.187	85.98	49.12	69.23	2104.8	1020.48	267.9
120.	0.1	20.60	- 12.808	96.48	46.15	70.52	1796.0	488.49	230.4
120.	0.5	20.60	- 12.793	96.45	46.16	70.50	1797.8	489.83	230.6
120.	1.0	20.61	- 12.774	96.40	46.18	70.48	1800.2	491.52	230.9
120.	2.0	20.63	- 12.736	96.32	46.22	70.43	1804.8	494.90	231.4
120.	5.0	20.67	- 12.621	96.06	46.34	70.30	1818.7	505.21	233.1
120.	10.0	20.75	- 12.430	95.65	46.53	70.10	1841.3	522.90	235.8
120.	20.0	20.89	- 12.045	94.85	46.91	69.75	1884.8	560.39	241.1
120.	30.0	21.03	- 11.659	94.09	47.26	69.47	1926.3	601.13	246.2
120.	40.0	21.16	- 11.272	93.36	47.60	69.24	1966.2	645.73	251.3
120.	50.0	21.28	- 10.885	92.66	47.92	69.04	2004.7	694.92	256.2
120.	60.0	21.39	- 10.496	92.00	48.24	68.88	2041.7	749.59	261.0
130.	0.1	20.23	- 12.105	102.11	44.98	70.15	1722.3	390.02	220.8
130.	0.5	20.23	-12.090	102.07	44.99	70.12	1724.4	391.08	221.0

## FRIEND, INGHAM, AND ELY

## TABLE A3. Properties of ethane in the single-phase region - Continued

T	Р	ρ	H	S	Cv	$C_{p}$	W	η	λ
К	MPa	mol·dm <sup>-3</sup>	kJ/mol	J/(mol·K)	J/(mol·K)	J/(mol·K)	m•s <sup>−1</sup>	µPa∙s	mW/(m·ŀ
130.	1.0	20.24	- 12.071	102.03	45.01	70.10	1726.9	392.40	221.3
130.	2.0	20.26	- 12.033	101.94	45.06	70.04	1731.8	395.05	221.9
130	5.0	20.31	- 11 920	101 67	45 19	69.88	1746.6	403 10	223.7
120	10.0	20.31	_ 11 721	101.07	45.10	60.64	1770.6	405.10	225.1
120.	20.0	20.55	- 11.751	101.24	45.40	(0.22	1016.6	410.05	220.0
130.	20.0	20.55	- 11.550	100.41	45.80	09.23	1810.0	445.01	232.3
130.	30.0	20.70	- 10.968	99.62	46.18	68.90	1860.4	476.41	237.8
130.	40.0	20.84	- 10.583	98.88	46.54	68.63	1902.2	509.58	243.2
130.	50.0	20.97	- 10.197	98.17	46.88	68.40	1942.3	545.55	248.4
130.	60.0	21.10	- 9.810	97.48	47.21	68.22	1980.9	584.80	253.5
140.	0.1	19.85	- 11.404	107.30	44.05	70.01	1648.3	321.12	211.0
140	0.5	19.86	- 11 390	107 26	44 07	60 08	1650.4	322.00	211.2
140.	10	19.87	- 11 371	107.20	44.09	69.94	1653 1	323 10	211.2
140	2.0	10.80	11.371	107.22	44.02	60.99	1655.1	225.10	211.5
140.	2.0	17.07	- 11.554	107.12	44.13	09.00	1058.5	323.29	212.2
140.	5.0	19.95	- 11.223	106.84	44.27	69.68	16/4.4	331.95	214.1
140.	10.0	20.04	- 11.036	106.39	44.49	69.38	1700.1	343.24	217.2
140.	20.0	20.21	- 10.660	105.52	44.90	68.89	1749.1	366.68	223.3
140.	30.0	20.37	- 10.281	104.71	45.29	68.48	1795.4	391.44	229.1
140.	40.0	20.52	- 9.899	103.94	45.66	68.16	1839.4	417.79	234.8
140.	50.0	20.67	-9.516	103.21	46.01	67.89	1881.4	445.97	240.3
140.	60.0	20.80	-9.131	102.52	46.34	67.67	1921.7	476.29	245.7
150	0.1	10 47	- 10 704	112 14	12 28	70.19	1572.9	270 54	201.1
150.	0.1	19.47	- 10.704	112.14	43.38	70.16	1575.8	270.34	201.1
150.	0.5	19.46	- 10.089	112.10	43.40	70.14	1576.2	271.30	201.3
150.	1.0	19.49	- 10.671	112.05	43.42	70.10	1579.1	272.25	201.7
150.	2.0	19.51	- 10.635	111.95	43.47	70.01	1584.9	274.15	202.4
150.	5.0	19.58	- 10.526	111.65	43.61	69.77	1602.1	279.89	204.4
150.	10.0	19.68	- 10.342	111.17	43.83	69.40	1629.8	289.59	207.7
150.	20.0	19.87	- 9.972	110.27	44.25	68.80	1682.2	309.55	214.2
150.	30.0	20.04	- 9.597	109.43	44.65	68.31	1731.4	330.41	220.4
150	40.0	20.21	-9219	108 64	45.02	67.92	1777.8	352 36	226.3
150.	50.0	20.36	- 8.839	107.89	45.36	67.60	1822.0	375.59	232.1
150	60.0	20.51	9 156	107 17	45 60	67 31	1964 1	400.32	727 8
1.0	00.0	20.31	- 8.430	107.17	43.09	07.34	1004.1	400.52	237.0
100.	0.1	19.08	- 10.000	116.68	42.95	70.65	1499.0	231.87	191.1
160.	0.5	19.09	- 9.986	116.64	42.97	70.60	1501.6	232.55	191.4
160.	1.0	19.11	- 9.968	116.58	42.99	70.55	1504.8	233.40	191.8
160.	2.0	19.13	- 9.933	116.48	43.04	70.44	1511.1	235.10	192.5
160.	5.0	19.20	-9.826	116.16	43.18	70.14	1529.8	240.22	194.7
160.	10.0	19.31	- 9.647	115.66	43.41	69.69	1559.7	248.84	198.2
160.	20.0	19.52	-9.283	114 72	43.83	68.95	1616.0	266.42	205.1
160	30.0	19 72	- 8 914	113.84	44 23	68 38	1668 3	284 62	211 6
160.	40.0	19.89	- 8.540	113.02	44.60	67.91	1717.3	303.59	217.8
160	50.0	20.04	0 162	112.25	44.04	67 51	1762 0	272 49	222.0
160	20.0	20.00	- 0.105	114.40	44.34	07.34	1907.0	343.40 744 45	223.9
100.	0.00	20.22	- 1.184	111.52	45.27	0/.23	1007.9	344.43	229.8
170.	0.1	18.09	- 9.290	120.98	42.12	/1.40	1423.7	201.30	181.3
1/0.	0.5	18.70	- 9.276	120.94	42.74	71.34	1426.5	201.92	181.6
170.	1.0	18.71	- 9.259	120.88	42.76	71.27	1430.0	202.70	182.0
170.	2.0	18.74	- 9.225	120.77	42.81	71.14	1437.0	204.27	182.8
170.	5.0	18.82	-9.122	120.43	42.95	70.77	1457.4	208.96	185.1
170.	10.0	18.94	- 8.948	119.90	43.18	70.22	1489.9	216.81	188.8
170.	20.0	19.17	- 8 592	118 91	43 61	69 33	1550.4	232.71	196.0
170.	30.0	19.39	- 8.229	117.99	44.01	68.65	1606.1	249.01	202.8
170	40.0	10 59	7 020	117 14	11 27	60 11	1650 0	265 06	200.4
170.	40.0	19.30	~ 7.000	117.14	44.3/	00.11	1036.0	203.80	209.4
170.	50.0	19.70	- 7.487	116.34	44.72	0/.07	1/06.8	283.38	215.7
1/0.	60.0	19.92	- 7.111	115.59	45.04	67.32	1753.1	301.71	221.8
180.	0.1	18.28	- 8.571	125.09	42.65	72.41	1347.8	176.42	171.5
180.	0.5	18.29	- 8.558	125.04	42.67	72.34	1351.0	177.01	171.9

TABLE A3. Properties of ethane in the single-phase region - Continued

T K	P MPa	ρ mol·dm <sup>-3</sup>	<i>H</i> kJ/mol	S J/(mol·K)	C <sub>V</sub> J∕(mol·K)	C <sub>p</sub> J/(mol·K)	W m·s <sup>-1</sup>	η μPa·s	λ mW/(m·K)
190	1.0	18 31	-8 542	124 98	42 70	72.26	1354.8	177.75	172.3
100.	2.0	18 34	- 8 509	124.96	42.76	72.09	1362.5	179.21	173.1
100.	5.0	18 43	8 410	124.50	42.89	71.63	1384.9	183.61	175.6
180.	10.0	18 57	- 8 242	123.93	43.12	70.96	1420.3	190.92	179.5
180.	20.0	18.82	- 7.896	122.89	43.55	69.89	1485.4	205.60	187.1
180	30.0	19.05	- 7 540	121 93	43 95	69.09	1544.8	220.50	194.2
180.	40.0	19.05	-7178	121.04	44.32	68.47	1599.7	235.79	201.0
180.	50.0	19.46	- 6 809	120.22	44.66	67.97	1651.1	251.56	207.6
180.	60.0	19.63	-6.437	119.45	44.97	67.56	1699.5	267.95	213.9
190.	0.1	0.07	6.704	207.97	32.55	41.80	251.8	6.06	9.9
100	0.5	17 97	- 7 820	128.00	42 75	73 61	1274 7	156.26	162.3
190.	0.5	17.07	- 7.813	128.97	42.78	73.50	1279.0	156.97	162.7
190.	2.0	17.02	-7783	128.72	42.70	73.29	1287.5	158 37	163.6
190.	2.0	18.03	-7.689	128.75	42.02	72 71	1312.2	162.56	166.2
190.	10.0	18 18	7 528	120.40	43.20	71.89	1350.9	169.48	170.4
190.	10.0	10.10	1.520	127.73	45.20	/1.0/	1550.5	105.10	170.1
190.	20.0	18.47	- 7.194	126.68	43.64	70.62	1421.2	183.26	178.3
190.	30.0	18.72	- 6.847	125.68	44.03	69.68	1484.6	197.12	185.8
190.	40.0	18.95	- 6.490	124.76	44.40	68.96	1542.7	211.21	192.9
190.	50.0	19.15	- 6.128	123.91	44.74	68.39	1596.6	225.66	199.6
190.	60.0	19.35	- 5.759	123.11	45.05	67.94	1647.3	240.56	206.1
200.	0.1	0.06	7.130	210.15	34.06	43.27	258.1	6.37	10.7
200.	0.5	17.44	- 7.085	132.80	42.96	75.15	1197.6	138.62	152.8
200.	1.0	17.46	- 7.071	132.73	42.98	75.01	1202.4	139.31	153.3
200.	2.0	17.50	- 7.043	132.58	43.03	74.75	1211.9	140.67	154.3
200.	5.0	17.61	- 6.955	132.16	43.17	74.02	1239.2	144.72	157.0
200.	10.0	17.79	- 6.804	131.51	43.41	73.00	1281.7	151.38	161.5
200.	20.0	18.10	- 6.483	130.33	43.84	71.48	1357.8	164.49	169.8
200.	30.0	18.38	- 6.146	129.27	44.24	70.38	1425.3	177.54	177.6
200.	40.0	18.63	- 5.798	128.31	44.60	69.57	1486.8	190.71	184.9
200.	50.0	18.85	- 5.441	127.43	44.94	68.93	1543.4	204.12	191.9
200.	60.0	19.06	- 5.078	126.61	45.25	68.42	1596.4	217.87	198.6
210.	0.1	0.06	7.567	212.29	35.08	44.18	264.3	6.68	11.6
210.	0.5	16.98	-6.325	136.51	43.28	77.02	1119.2	123.35	143.6
210.	1.0	17.00	- 6.312	136.43	43.30	76.84	1124.6	124.03	144.1
210.	2.0	17.05	- 6.287	136.27	43.35	76.49	1135.3	125.37	145.1
210.	5.0	17.18	- 6.208	135.81	43.49	75.56	1165.8	129.36	148.1
210.	10.0	17.38	- 6.067	135.10	43.72	74.29	1212.7	135.85	152.8
210.	20.0	17.74	- 5.764	133.84	44.16	72.46	1295.1	148.47	161.6
210.	30.0	18.04	- 5.439	132.72	44.55	71.19	1367.2	160.89	169.6
210.	40.0	18.31	- 5.099	131.72	44.92	70.26	1432.1	173.32	177.2
210.	50.0	18.55	- 4,749	130.80	45.25	69.55	1491.5	185.90	184.4
210.	60.0	18.77	- 4.391	129.96	45.55	69.00	1546.8	198.72	191.2
220.	0.1	0.06	8.013	214.36	35.99	44.99	270.4	6.99	12.5
220.	0.5	16.50	5.543	140.14	43.71	79.31	1039.1	109.91	134.5
220.	1.0	16.53	- 5.533	140.05	43.73	79.07	1045.3	110.59	135.0
220	20	16 58	- 5 512	139.88	43 78	78.60	1057.4	111 95	136 1
220	5.0	16.73	- 5.443	139.37	43.91	77.38	1091.8	115.92	139.4
220	10.0	16.97	-5.317	138.59	44.13	75.78	1143.7	122.32	144.4
220	20.0	17.36	- 5.034	137.23	44.57	73.57	1233.3	134.60	153.6
220.	30.0	17.70	- 4.722	136.06	44.96	72.10	1310.2	146.54	162.0
220	40.0	17 00	- 1 202	125 01	15 22	71.04	1378 7	158 27	160 9
220.	40.0 50.0	12.77	- 4.592	133.01	45.55 A5 66	71.04	1441 0	170.37	107.0
220.	50.0	18.49	- 3 602	132.10	45.00	60.64	1498 5	187 33	184 7
220.	00.0 A 1	0.70	9.070 8 467	216 28	36 01	45 87	276.2	7 30	13.4
230.	0.1	0.05	8 031	201.68	38 51	50.89	262.3	740	14.2
<i></i>	0.0	0.4.7	0.001			2 3.07			- ··-

TABLE A3. Properties of ethane in the single-phase region - Continued

T K	P MPa	ρ mol·dm <sup>-3</sup>	H kJ/mol	S J/(mol·K)	C <sub>V</sub> J/(mol∙K)	Cp J/(mol·K)	W m·s <sup>−1</sup>	ຖ µPa∙s	λ mW/(m·K)
230	10	16.02	-4 729	143.63	44.27	81.87	963 7	98 59	126.2
230.	2.0	16.08	-4.713	143.42	44.31	81.18	977.7	99,98	127.4
230.	5.0	16.26	- 4.659	142.85	44.43	79.54	1017.0	104.01	130.8
230.	10.0	16.53	- 4.551	142.00	44.64	77.48	1074.9	110.41	136.2
230.	20.0	16.98	- 4.292	140.53	45.07	74.79	1172.4	122.47	145.9
230.	30.0	17.35	- 3.996	139.28	45.46	73.08	1254.5	134.02	154.6
230.	40.0	17.66	- 3.678	138.18	45.82	71.90	1326.6	145.37	162.7
230.	50.0	17.94	- 3.343	137.20	46.15	71.02	1391.8	156.70	170.2
230.	60.0	18.19	- 2.998	136.29	46.45	70.35	1451.7	168.12	177.4
240.	0.1	0.05	8.930	218.35	37.87	46.70	281.8	7.61	14.4
240.	0.5	0.27	8.539	203.84	39.15	50.78	269.5	7.71	15.0
240.	1.0	15.47	- 3.894	147.18	44.93	85.36	878.8	87.70	117.4
240.	2.0	15.55	- 3.886	146.94	44.95	84.42	895.4	89.15	118.8
240.	5.0	15.77	-3.851	146.29	45.05	82.13	940.8	93.32	122.6
240.	10.0	16.08	- 3.767	145.33	45.24	79.43	1006.0	99.79	128.3
240.	20.0	16.59	- 3.537	143.74	45.65	76.13	1112.4	111.75	138.5
240.	30.0	16.99	- 3.260	142.41	46.04	74.14	1200.0	123.01	147.6
240.	40.0	17.34	-2.954	141.26	46.40	72.81	1276.0	133.96	155.8
240.	50.0	17.64	- 2.629	140.24	46.72	71.84	1344.1	144.81	163.6
240.	60.0	17.91	-2.290	139.30	47.02	71.12	1406.2	155.70	170.9
250.	0.1	0.05	9.401	220.27	38.87	47.64	287.3	7.91	15.4
250.	0.5	0.26	9.048	205.92	39.92	51.02	276.3	8.01	16.0
250.	1.0	0.56	8.525	198.66	42.07	58.31	260.3	8.19	17.0
250.	2.0	14.96	- 3.021	150.47	45.72	88.72	808.9	79.18	110.2
250.	5.0	15.24	- 3.014	149.71	45.77	85.33	862.7	83.59	114.5
250.	10.0	15.61	- 2.961	148.62	45.93	81.68	936.9	90.25	120.7
250.	20.0	16.19	-2.769	146.88	46.31	77.59	1053.5	102.20	131.5
250.	30.0	16.64	-2.513	145.46	46.69	75.28	1146.9	113.24	140.9
250.	40.0	17.01	-2.221	144.25	47.05	73.77	1226.9	123.86	149.4
250.	50.0	17.34	- 1.906	143.19	47.37	72.71	1297.8	134.31	157.3
250.	60.0	17.62	- 1.575	142.22	47.66	71.93	1362.3	144.73	164.7
260.	0.1	0.05	9.883	222.16	39.92	48.63	292.5	8.22	16.5
260.	0.5	0.25	9.560	207.93	40.79	51.49	282.7	8.32	17.0
260.	1.0	0.53	9.100	200.91	42.31	56.89	268.8	8.49	17.9
260.	2.0	14.31	- 2.105	154.07	46.64	94.93	716.0	69.78	101.7
260.	5.0	14.66	-2.141	153.13	46.60	89.46	781.7	74.61	106.5
260.	10.0	15.11	-2.132	151.87	46.69	84.31	867.4	81.59	113.4
260.	20.0	15.78	- 1.985	149.95	47.04	79.17	995.7	93.65	124.8
260.	30.0	16.28	- 1.755	148.44	47.41	76.48	1095.3	104.53	134.5
260.	40.0	16.68	- 1.479	147.17	47.76	74.79	1179.3	114.88	143.2
260.	50.0	17.03	- 1.175	146.06	48.08	73.63	1253.1	124.97	151.3
260.	60.0	17.34	-0.852	145.06	48.37	72.78	1319.8	134.99	158.8
270.	0.1	0.05	10.374	224.02	41.00	49.68	297.7	8.52	17.6
270.	0.5	0.24	10.078	209.88	41.74	52.13	288.8	8.62	18.1
270.	1.0	0.50	9.666	203.05	42.90	56.38	276.6	8.78	18.9
270.	2.0	1.20	8.589	194.29	47.40	76.57	245.2	9.33	21.6
270.	5.0	14.01	- 1.220	156.61	47.57	95.16	696.3	66.16	98.7
270.	10.0	14.58	- 1.273	155.11	47.55	87.46	797.4	73.66	106.3
270.	20.0	15.35	- 1.185	152.97	47.83	80.88	939.0	85.94	118.4
270.	30.0	15.91	- 0.983	151.35	48.19	77.74	1045.2	96.72	128.5
270.	40.0	16.35	- 0.725	150.01	48.54	75.85	1133.3	106.83	137.4
270.	50.0	16.73	-0.434	148.85	48.85	74.58	1210.1	116.63	145.6
270.	60.0	17.05	-0.119	147.82	49.14	73.67	1278.9	126.29	153.3
280.	0.1	0.04	10.876	225.84	42.13	50.77	302.6	8.82	18.8
280.	0.5	0.23	10.603	211.79	42.76	52.89	294.6	8.92	19.2

TABLE A	A3.	Propertie	es of	ethane	in 1	the sir	igle-ph	ase re	egion		Continued
---------	-----	-----------	-------	--------	------	---------	---------	--------	-------	--	-----------

T K	<i>P</i> MPa	ρ mol∙dm <sup>-3</sup>	H kJ/mol	S J/(mol·K)	C <sub>V</sub> J/(mol⋅K)	$C_{\rho}$ J/(mol·K)	W m·s <sup>−1</sup>	η μPa·s	λ mW/(m·K)
280	1.0	0.48	10.230	205 10	13 68	56 37	282.7	0.07	10.0
280	2.0	1 10	0.250	106.03	46 57	69.55	263.7	9.07	19.9
280	5.0	13.26	-0.228	160.21	48.37	103.93	603 7	57.96	00 7
280	10.0	14.01	-0.380	158 36	48.50	91 30	726.4	66 31	99.5
280.	20.0	14.92	- 0.367	155.94	48.69	82.72	883.7	78.96	112.4
280.	30.0	15.54	-0.200	154.20	49.03	79.05	996.8	89.68	122.8
280.	40.0	16.02	0.038	152.79	49.36	76.94	1089.0	99.60	132.0
280.	50.0	16.42	0.317	151.58	49.68	75.56	1168.6	109.13	140.3
280.	60.0	16.77	0.622	150.52	49.96	74.59	1239.6	118.48	148.0
290.	0.1	0.04	11.390	227.64	43.29	51.90	307.5	9.11	20.0
290.	0.5	0.22	11.136	213.66	43.83	53.76	300.2	9.21	20.4
290.	1.0	0.45	10.795	207.08	44.59	56.68	290.5	9.36	21.0
290.	2.0	1.02	9.992	199.30	46.67	66.30	268.1	9.81	22.8
290.	5.0	12.33	0.884	164.11	50.26	120.55	497.6	49.56	82.4
290.	10.0	13.38	0.556	161.65	49.55	96.18	654.2	59.42	92.9
290.	20.0	14.47	0.470	158.88	49.60	84.71	829.9	72.61	106.8
290.	30.0	15.16	0.598	157.00	49.91	80.42	950.2	83.32	117.5
290.	40.0	15.69	0.814	155.51	50.24	78.07	1046.5	93.08	126.9
290.	50.0	16.12	1.078	154.25	50.55	76.57	1128.8	102.37	135.3
290.	60.0	16.49	1.372	153.15	50.83	75.53	1201.9	111.44	143.1
300.	0.1	0.04	11.914	229.42	44.49	53.06	312.2	9.41	21.2
300.	0.5	0.21	11.678	215.50	44.95	54.70	305.6	9.50	21.6
300.	1.0	0.43	11.364	209.01	45.59	57.20	296.8	9.65	22.2
300.	2.0	0.96	10.646	201.52	47.20	64.69	277.3	10.07	23.8
300.	5.0	10.91	2.283	168.85	53.19	172.16	360.6	39.64	73.1
300.	10.0	12.68	1.548	165.01	50.73	102.60	580.6	52.88	86.4
300.	20.0	14.00	1.327	161.79	50.57	86.84	777.6	66.82	101.4
300.	30.0	14.78	1.409	159.75	50.85	81.83	905.4	77.56	112.6
300.	40.0	15.36	1.600	158.17	51.17	79.22	1005.7	87.18	122.1
300.	50.0	15.82	1.849	156.86	51.47	77.60	1090.8	96.26	130.7
300.	60.0	16.21	2.133	155.73	51.75	76.50	1165.7	105.07	138.6
310.	0.1	0.04	12.451	231.18	45.71	54.26	316.9	9.70	22.5
310.	0.5	0.20	12.231	217.31	46.10	55.71	310.8	9.80	22.9
310.	1.0	0.42	11.939	210.89	46.65	57.87	302.9	9.94	23.4
310.	2.0	0.91	11.289	203.63	47.94	63.94	285.6	10.33	24.8
310.	5.0	4.13	7.602	186.23	59.31	263.20	210.7	14.98	45.4
310.	10.0	11.88	2.616	168.51	52.07	111.37	506.3	46.56	80.0
310.	20.0	13.52	2.207	164.67	51.59	89.10	727.2	61.52	96.5
310.	30.0	14.39	2.234	162.45	51.82	83.27	862.5	72.33	108.0
310.	40.0	15.02	2.398	160.79	52.13	80.39	966.8	81.83	117.7
310.	50.0	15.51	2.630	159.43	52.42	78.65	1054.4	90.71	126.4
310.	60.0	15.93	2.902	158.25	52.70	77.48	1131.3	99.29	134.3
320.	0.1	0.04	13.000	232.92	46.95	55.48	321.4	9.99	23.9
320.	0.5	0.19	12.793	219.10	47.29	56.78	315.9	10.08	24.2
320.	1.0	0.40	12.522	212.74	47.76	58.67	308.7	10.22	24.7
320.	2.0	0.87	11.927	205.65	48.83	63.72	293.3	10.59	25.9
320.	5.0	3.18	9.258	191.50	54.72	123.27	236.9	13.55	36.4
320.	10.0	10.94	3.787	172.23	53.59	123.47	434.1	40.37	73.8
320.	20.0	13.02	3.110	167.54	52.64	91.49	679.1	56.68	91.9
320.	30.0	14.00	3.074	165.12	52.83	84.73	821.7	67.57	103.8
320.	40.0	14.68	3.208	163.36	53.12	81.57	929.9	76.97	113.6
320.	50.0	15.21	3.422	161.94	53.41	79.71	1019.9	85.67	122.4
320.	60.0	15.65	3.682	160.73	53.68	78.48	1098.4	94.03	130.3
330.	0.1	0.04	13.561	234.65	48.20	56.72	325.9	10.28	25.2
330.	0.5	0.19	13.366	220.86	48.51	57.89	320.8	10.37	25.6

## FRIEND, INGHAM, AND ELY

				······					
T K	Р MPa	ρ mol·dm <sup>-3</sup>	<i>H</i> kJ/mol	S J/(mol·K)	C <sub>V</sub> J/(mol⋅K)	C <sub>p</sub> J/(mol·K)	W m·s <sup>-1</sup>	η μPa·s	λ mW/(m·K)
220	1.0	0.20	10 110	014.50	40.01	50.56	214.0	10.51	26.0
330.	1.0	0.39	13.113	214.56	48.91	59.50	314.2	10.51	26.0
330.	2.0	0.83	12.304	207.02	49.01	05.65	300.3	10.00	27.1
330.	5.0	2.77	10.347	194.85	53.90	98.73	253.9	13.22	34.7
330.	10.0	9.81	5.098	176.26	55.21	139.15	369.9	34.34	67.8
330.	20.0	12.51	4.037	170.39	53.74	93.95	633.6	52.24	87.7
330.	30.0	13.61	3.929	167.75	53.87	86.20	783.1	63.25	99.9
330.	40.0	14.35	4.030	165.89	54.14	82.76	894.8	72.54	109.9
330.	50.0	14.91	4.224	164.41	54.43	80.77	987.0	81.09	118.7
330.	60.0	15.38	4,472	163.16	54.70	79.48	1067.1	89.24	126.7
340.	0.1	0.04	14.134	236.36	49.48	57.97	330.3	10.56	26.7
240	0.5	0.19	12.051	222 (1	40.74	50.02	225 (	10.65	27.0
340.	1.0	0.18	13.951	222.01	49.74	59.03	325.0	10.05	27.0
340.	1.0	0.37	13.713	210.55	50.09	00.52	319.0	10.79	27.4
540.	2.0	0.79	13.205	209.53	50.86	64.22	307.2	11.12	28.4
340.	5.0	2.51	11.277	197.63	53.99	88.51	267.5	13.14	34.5
340.	10.0	8.52	6.562	180.63	56.59	151.90	321.0	28.79	62.2
340.	20.0	11.97	4.989	173.23	54.86	96.40	591.4	48.20	83.8
340.	30.0	13.21	4,798	170.35	54.93	87.65	746.9	59.31	96.4
340.	40.0	14.01	4.863	168.38	55.19	83.94	861.7	68.52	106.5
340.	50.0	14.61	5.037	166.84	55.47	81.84	955.9	76.90	115.3
340.	60.0	15.10	5.272	165.55	55.73	80.49	1037.5	84.87	123.4
250	0.1	0.02	14 500	000.07	50 54	50.04	224.6	10.04	00.1
350.	0.1	0.03	14.720	238.06	50.76	59.24	334.6	10.84	28.1
350.	0.5	0.18	14.547	224.33	50.99	60.20	330.2	10.93	28.4
350.	1.0	0.36	14.323	218.12	51.30	61.54	324.8	11.06	28.8
350.	2.0	0.76	13.849	211.40	51.97	64.77	313.5	11.39	29.7
350.	5.0	2.32	12.133	200.11	54.49	83.15	279.1	13.17	34.8
350.	10.0	7.25	8.080	185.03	57.58	148.87	294.3	24.46	57.3
350.	20.0	11.43	5.965	176.06	56.00	98.73	553.0	44.52	80.3
350.	30.0	12.81	5.682	172.91	56.02	89.07	713.1	55.73	93.2
350.	40.0	13.67	5,708	170.83	56.26	85.10	830.6	64.85	103.4
350.	50.0	14.32	5.861	169.22	56.53	82.90	926.6	73.08	112.3
350	60.0	14 83	6 082	167.00	56 70	<b>81 5</b> 0	1000 4	80.86	120.3
360	01	0.03	15 210	220.75	52.05	60.57	228.8	11 17	20.5
360.	0.1	0.03	15.319	239.73	52.05	61.40	330.0	11.12	29.0
300.	0.5	0.17	15.155	226.05	52.20	01.40	334.0	11.21	29.9
300. 260	1.0	0.55	14.944	219.87	52.55	62.60	329.8	11.34	30.3 21.1
500.	2.0	0.75	14.500	215.25	55.11	03.40	519.0	11.05	51.1
360.	5.0	2.17	12.948	202.40	55.20	80.07	289.4	13.26	35.5
360.	10.0	6.23	9.499	189.03	58.24	134.09	286.6	21.65	53.4
360.	20.0	10.87	6.962	178.87	57.15	100.80	519.1	41.21	77.2
360.	30.0	12.41	6.580	175.44	57.12	90.44	681.8	52.48	90.4
360.	40.0	13.34	6.565	173.24	57.34	86.25	801.5	61.50	100.6
360.	50.0	14.02	6.695	171.57	57.61	83.96	899.0	69.59	109.5
360.	60.0	14.56	6,902	170.21	57.86	82.51	982.9	77.20	117.5
370.	0.1	0.03	15 931	241 42	53 34	61.80	343.0	11 40	31.2
370	0.5	0.17	15 775	227 75	53 53	62 61	339 3	11 49	31.4
370.	1.0	0.34	15.575	221.60	53.77	63.70	334.7	11.61	31.8
270	2.0	0.71	15 150	015.00	54.00	(( ))	205.4	44.04	22 (
370.	2.0	0./1	13.139	215.03	54.28	00.24	343.4	11.91	32.0
370.	5.0	2.04	15.759	204.57	50.05	/8.20	298.0	13.39	30.4
370.	10.0	5.48	10.763	192.49	58.80	119.38	288.9	19.97	50.9
370.	20.0	10.31	7.979	181.66	58.29	102.40	490.0	38.27	74.5
370.	30.0	12.01	7.490	177.93	58.23	91.72	653.2	49.53	87.8
370.	40.0	13.01	7.433	175.62	58.44	87.36	774.5	58.45	98.1
370.	50.0	13.73	7.540	173.89	58.70	85.00	873.1	66.40	107.0
370.	60.0	14.30	7.732	172.48	58.95	83.52	957.9	73.84	115.0
380.	0.1	0.03	16.555	243.09	54.64	63.09	347.1	11.67	32.8

TABLE A3. Properties of ethane in the single-phase region - Continued

0.16

16.407

229.43

63.83

54.80

343.7

11.76

33.0

0.5

380.

TABLE A3. Properties of ethane in the single-phase region - Continued

$ \begin{array}{c c c c c c c c c c c c c c c c c c c $										
K         MPa         moldm <sup>-73</sup> k/mol $J/(mol+K)$ $J/(mol+K)$ $ms^{-1}$ $\mu$ Pars         m W           380.         1.0         0.33         16.218         223.32         55.02         64.82         339.5         11.88         3           380.         5.0         1.94         14.515         206.64         56.99         77.23         307.1         13.553         3           380.         10.0         4.94         11.901         195.33         59.34         92.289         627.3         46.687         8           380.         30.0         11.61         6.414         80.39         59.54         82.49         627.3         46.687         8           380.         50.0         13.44         8.352         170.6         59.55         88.44         70.4         15           380.         60.0         14.03         8.572         174.72         60.05         84.51         934.4         12.03         3           390.         0.1         0.32         11.6872         223.111         56.68         66.60         348.1         12.03         3           390.         5.0         1.587         12.281         23.66	Т	Р	ρ	Н	S	$C_V$	$C_p$	W	η	λ
	K	MPa	mol·dm <sup>-3</sup>	kJ/mol	J/(mol·K)	J/(mol·K)	J/(mol·K)	m·s <sup>-1</sup>	µPa∙s	mW/(m·K)
	380	10	0.33	16 218	223.32	55.02	64.82	339.5	11.88	33.3
	380	2.0	0.68	15.826	216.81	55.47	67.11	331.0	12.17	34.1
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	380	5.0	1 94	14 515	206.64	56.99	77 23	307.1	13 55	37.5
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	380	10.0	4 94	11 901	195.53	59.42	108.70	295.0	18.96	49.4
300.         300.         11.61         8.414         180.39         59.34         92.89         627.3         46.87         8           300.         40.0         12.68         8.312         177.96         59.55         88.44         749.4         55.68         90           300.         50.0         13.44         8.352         174.17         59.00         84.51         93.44         70.76         11           390.         0.5         0.16         17.051         22.11.1         56.07         65.06         44.81         12.15         3           390.         2.0         0.66         16.501         218.57         56.66         66.03         33.64         12.43         3           390.         2.0         0.66         16.501         20.86.4         58.00         7.72         315.0         13.72         3           390.         2.00         9.23         10.045         187.09         60.42         103.75         447.0         33.45         7           390.         2.00         9.23         10.245         187.09         60.47         39.34         60.42         44.47         8           390.         2.00         13.13         9.	380.	20.0	9.76	9.008	184.40	59.41	103.39	466.0	35.68	72.2
380.         300.         11.61         8.414         180.39         59.34         92.89         627.3         46.87         8           380.         40.0         12.68         8.312         177.66         59.55         88.44         707.67         11           380.         60.0         14.03         8.572         174.72         60.05         84.51         934.4         707.67         11           390.         0.1         0.03         17.192         244.74         55.93         64.38         351.2         11.94         3           390.         0.5         0.16         17.051         221.11         56.08         65.06         348.1         12.03         3           390.         2.0         0.66         16.501         218.57         56.68         68.03         336.4         12.43         3           390.         10.0         4.52         12.849         198.25         60.13         101.44         302.5         18.33         4           390.         10.0         4.52         12.849         180.27         60.66         89.47         72.64         60.80         100           390.         0.0         13.15         9.260										
380.         400.         12.68         8.12         177.96         59.55         88.44         749.4         55.68         9           380.         50.0         11.403         8.572         174.72         60.05         84.51         93.44         10.43         31.2         11.94         33           390.         0.1         0.03         17.192         244.74         55.93         64.38         351.2         11.94         33           390.         0.5         0.16         17.051         225.01         56.67         65.97         344.2         12.43         33           390.         2.0         0.66         16.501         218.57         56.68         68.03         336.4         12.43         3           390.         1.0.0         4.52         12.949         198.25         60.13         101.44         302.5         18.33         4           390.         2.0.0         9.23         10.045         187.09         60.52         193.75         447.0         33.45         7           390.         40.0         12.35         9.20         193.42         60.91         87.02         82.64         60.89         100         10.99         10.99	380.	30.0	11.61	8.414	180.39	59.34	92.89	627.3	46.87	85.6
380.         50.0         13.44         8.395         176.17         59.80         86.02         84.81         90.43.4         70.76         11           390.         0.1         0.03         17.192         244.74         55.93         64.38         351.2         11.94         3           390.         0.5         0.16         17.051         231.11         56.08         65.06         348.1         12.03         3           390.         1.0         0.32         16.672         25.01         56.27         65.97         344.2         12.15         3           390.         2.0         0.66         16.501         218.57         56.66.8         68.03         336.4         12.43         3           390.         2.0.0         9.23         10.045         187.09         60.52         103.75         447.0         33.45         7           390.         40.0         12.35         9.202         18.02         16.031         87.02         82.64         60.801         13.15         9           390.         60.0         13.15         9.224         176.93         111         40.726         83.47         12.42         33         33.47         12.42	380.	40.0	12.68	8.312	177.96	59.55	88.44	749.4	55.68	95.9
	380.	50.0	13.44	8.395	176.17	59.80	86.02	848.9	63.48	104.8
390.         0.1         0.03         17.192         244.74         55.93         64.38         351.2         11.94         3           390.         0.5         0.16         17.051         221.11         56.08         65.06         348.1         12.03         3           390.         2.0         0.66         16.501         218.57         56.68         68.03         336.4         12.15         3           390.         5.0         1.851         55.285         208.44         58.00         76.72         315.0         13.72         3           390.         10.0         4.52         12.949         198.25         60.13         101.44         302.5         18.33         4           390.         40.0         11.21         9.48         182.82         60.47         73.64         60.42         44.47         8           390.         60.0         13.75         9.221         176.93         61.15         85.50         912.4         67.93         11           400.         0.1         0.03         17.843         246.39         57.23         65.66         355.2         12.21         93           400.         0.0         1.0         0.31 <td>380.</td> <td>60.0</td> <td>14.03</td> <td>8.572</td> <td>174.72</td> <td>60.05</td> <td>84.51</td> <td>934.4</td> <td>70.76</td> <td>112.7</td>	380.	60.0	14.03	8.572	174.72	60.05	84.51	934.4	70.76	112.7
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	390.	0.1	0.03	17.192	244.74	55.93	64.38	351.2	11.94	34.4
390.         0.5         0.16         17.051         231.11         36.08         65.06         344.2         12.15         3           390.         2.0         0.66         16.501         218.57         56.68         66.03         336.4         12.43         3           390.         5.0         1.85         15.285         208.64         58.00         76.72         315.0         13.72         3           390.         20.0         9.23         10.045         187.09         60.52         103.75         447.0         33.45         78           390.         30.0         11.21         9.348         182.82         60.47         93.94         604.2         44.47         8           390.         40.0         12.35         9.202         178.42         60.91         87.02         82.4         60.80         10           390.         60.0         13.17         9.422         176.93         61.15         85.50         912.4         67.93         11           40.         0.1         0.03         17.843         246.39         57.23         65.66         355.2         12.21         33           400.         0.0         0.10         0.31 </td <td></td>										
390.         1.0         0.32         16.872         223.01         56.27         56.397         344.2         12.15         3           390.         5.0         1.85         15.285         208.64         58.00         76.72         315.0         13.72         3           390.         10.0         4.52         12.949         198.25         60.13         101.44         302.5         18.33         4           390.         20.0         9.23         10.045         187.09         60.52         103.75         447.0         33.45         7           390.         40.0         11.21         9.348         182.82         60.47         93.94         604.2         44.47         8           390.         60.0         13.15         9.260         178.42         60.91         87.02         826.4         60.80         10           390.         60.0         13.78         9.422         176.93         61.15         85.50         912.4         67.93         11           400.         0.1         0.31         17.783         226.70         57.33         67.13         348.7         12.42         33           400.         1.0         4.31         17	390.	0.5	0.16	17.051	231.11	56.08	65.06	348.1	12.03	34.6
390.         2.0         0.06         16.301         218.37         36.08         68.03         35.64         12.43         3           390.         10.0         4.52         12.949         198.25         60.13         101.44         302.5         18.33         4           390.         20.0         9.23         10.045         187.09         60.52         103.75         447.0         33.45         7           390.         30.0         11.21         9.348         182.82         60.047         93.34         60.42         44.47         8           390.         50.0         13.15         9.200         176.42         60.91         87.02         826.4         60.80         10           390.         60.0         13.78         9.422         176.93         61.15         85.50         912.4         67.93         11           400.         0.1         0.031         17.843         246.39         57.23         65.66         355.2         12.21         3           400.         2.0         0.44         17.186         220.70         57.53         67.13         348.7         12.42         33           400.         1.0         0.31         1	390.	1.0	0.32	16.872	225.01	56.27	65.97	344.2	12.15	34.9
390.         3.0         1.3.2         1.3.29         1.3.29         1.3.2         3.4.3           390.         10.0         4.52         12.949         198.25         60.13         101.44         302.5         18.33         4           390.         20.0         9.23         10.045         187.09         60.52         103.75         447.0         33.45         7           390.         30.0         11.21         9.348         182.82         60.47         93.94         604.2         44.47         8           390.         50.0         13.15         9.202         180.27         60.66         89.47         726.4         63.51         17.31         11           400.         0.1         0.03         17.843         246.39         57.23         65.56         355.2         12.21         3           400.         1.0         0.31         17.708         232.77         57.36         67.13         348.7         12.42         3           400.         2.0         0.64         17.186         220.30         57.73         322.4         13.91         4           400.         2.0         8.73         11.062         189.72         61.61 <td< td=""><td>390.</td><td>2.0</td><td>0.66</td><td>16.501</td><td>218.57</td><td>50.08</td><td>68.03</td><td>330.4</td><td>12.43</td><td>35.6</td></td<>	390.	2.0	0.66	16.501	218.57	50.08	68.03	330.4	12.43	35.6
390.         10.0         4.5.2         12.494         198.25         60.13         101.44         302.5         18.3.3         4           390.         20.0         9.23         10.045         187.09         60.52         103.75         447.0         33.45         7           390.         30.0         11.21         9.348         182.82         60.047         93.94         60.42         44.47         8           390.         40.0         12.35         9.202         176.93         61.15         85.50         912.4         67.93         11           400.         0.1         0.03         17.843         246.39         57.23         65.66         355.2         12.21         3           400.         1.0         0.31         17.737         22.670         57.73         66.29         352.3         12.30         3           400.         2.0         0.44         17.186         220.30         57.89         69.00         341.6         12.69         3           400.         10.0         4.20         13.937         200.75         60.92         96.53         310.4         17.94         4           400.         10.0         8.33         1	390.	5.0	1.85	15.285	208.64	58.00	/6./2	315.0	13.72	38.7
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	390.	10.0	4.52	12.949	198.25	60.13	101.44	302.5	18.33	48.8
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	300	20.0	0.23	10.045	187 09	60.52	103 75	447.0	33 45	70.4
350.         360.         11.21         3.2.47         102.32         60.72         77.26.4         53.17         93.07           390.         50.0         13.15         9.202         180.27         60.66         89.47         726.4         53.15         9.203           400.         0.1         0.03         17.843         246.39         57.23         65.66         355.2         12.21         3           400.         0.1         0.31         17.373         226.70         57.35         66.29         352.3         12.30         3           400.         1.0         0.31         17.373         226.70         57.53         67.13         348.7         12.42         3           400.         2.0         0.64         17.178         220.30         57.89         69.00         341.6         12.69         3           400.         10.0         4.20         13.937         200.75         60.92         96.53         310.4         17.94         44           400.         30.0         10.83         10.852.1         61.61         1003.57         352.5         58.36         100           400.         60.0         13.52         10.282         199.11	300.	20.0	11 21	0 348	187.87	60.32	03.04	604.2	55.45 AA A7	83.7
300.         500         12.35         32.60         100.21         60.91         87.02         826.4         60.80         10           390.         60.0         13.78         9.422         176.93         61.15         85.50         912.4         67.93         11           400.         0.1         0.03         17.843         246.39         57.23         65.66         355.2         12.20         3           400.         1.0         0.31         17.537         226.70         57.53         67.13         348.7         12.42         3           400.         2.0         0.64         17.186         200.30         57.89         69.00         341.6         12.69         3           400.         1.0.0         4.20         13.937         200.75         60.92         96.53         310.4         17.94         4           400.         20.0         8.73         11.082         189.72         61.61         103.57         432.5         31.55         6           400.         30.0         10.83         10.292         185.21         61.59         94.85         583.8         42.31         8           400.         60.0         13.52         10	200	30.0 40.0	12.21	9.540	180.27	60.66	80 47	776 4	53.15	04.0
300.         60.0         13.78 $2.423$ 176.92         60.15         85.55         912.4         67.93         11           400.         0.1         0.03         17.843         246.39         57.23         65.66         355.2         12.21         3           400.         0.5         0.15         17.708         232.77         57.36         66.29         352.3         12.20         3           400.         2.0         0.64         17.186         232.77         57.36         67.13         348.7         12.42         3           400.         2.0         0.64         17.186         220.30         57.89         69.00         341.6         12.69         3           400.         1.0         4.20         13.937         200.75         60.92         96.53         310.4         17.94         4           400.         30.0         10.83         10.292         185.21         61.59         94.85         583.8         42.31         8           400.         40.0         12.03         10.101         182.55         61.78         90.45         58.36         10           400.         60.0         13.52         10.282         1	390.	40.0 50.0	13.15	9.202	178 42	60.00	87.02	826.4	60.80	102.8
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	390.	60.0	13.78	9.422	176.93	61.15	85.50	912.4	67.93	110.7
$\begin{array}{cccccccccccccccccccccccccccccccccccc$										
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	400.	0.1	0.03	17.843	246.39	57.23	65.66	355.2	12.21	36.0
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	400.	0.5	0.15	17.708	232.77	57.36	66.29	352.3	12.30	36.2
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	400.	1.0	0.31	17.537	226.70	57.53	67.13	348.7	12.42	36.5
400. $5.0$ $1.77$ $16.051$ $210.58$ $59.05$ $76.57$ $322.4$ $13.91$ $44$ $400.$ $10.0$ $4.20$ $13.937$ $200.75$ $60.92$ $96.53$ $310.4$ $17.94$ $44$ $400.$ $20.0$ $8.73$ $11.082$ $189.72$ $61.61$ $103.57$ $432.5$ $31.55$ $66$ $400.$ $30.0$ $10.83$ $10.292$ $185.21$ $61.59$ $94.85$ $583.8$ $42.31$ $88$ $400.$ $40.0$ $12.03$ $10.101$ $182.55$ $61.78$ $90.45$ $705.3$ $50.85$ $99$ $400.$ $50.0$ $12.87$ $10.135$ $180.63$ $62.02$ $88.00$ $805.5$ $58.36$ $10$ $400.$ $60.0$ $13.52$ $10.282$ $179.11$ $62.26$ $86.47$ $891.8$ $65.33$ $100$ $410.$ $0.1$ $0.03$ $18.506$ $248.03$ $58.51$ $66.94$ $359.2$ $12.48$ $37$ $410.$ $1.0$ $0.30$ $18.215$ $228.37$ $58.79$ $68.30$ $353.2$ $12.68$ $33$ $410.$ $2.0$ $0.62$ $17.881$ $222.02$ $59.11$ $70.01$ $346.7$ $12.95$ $33$ $410.$ $50$ $1.69$ $16.817$ $212.47$ $60.14$ $76.68$ $239.4$ $14.10$ $44$ $410.$ $10.0$ $3.93$ $14.885$ $20.30.9$ $61.79$ $93.18$ $318.2$ $17.70$ $44$ $410.$ $50.0$ $12.59$ $1$	400.	2.0	0.64	17.186	220.30	57.89	69.00	341.6	12.69	37.2
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	400.	5.0	1.77	16.051	210.58	59.05	76.57	322.4	13.91	40.0
400.10.04.2013.937200.75 $60.92$ $96.53$ $310.4$ $17.94$ 4400.20.0 $8.73$ 11.082189.72 $61.61$ $103.57$ $432.5$ $31.55$ $66$ 400.30.010.8310.292185.21 $61.59$ $94.85$ $583.8$ $42.31$ $88$ 400.40.012.0310.101182.55 $61.78$ $90.45$ $705.3$ $50.85$ $99$ 400.60.013.5210.282179.11 $62.26$ $86.47$ $891.8$ $65.33$ $100$ 410.0.10.0318.506 $248.03$ $58.51$ $66.94$ $359.2$ $12.48$ $35$ 410.0.50.1518.377 $234.42$ $58.64$ $67.53$ $356.5$ $12.56$ $33$ 410.1.00.3018.215 $228.37$ $58.79$ $68.30$ $353.2$ $12.68$ $33$ 410.2.00.6217.881 $222.02$ $59.11$ $70.01$ $346.7$ $12.95$ $33$ 410.5.01.6916.817 $212.47$ $60.14$ $76.68$ $329.4$ $14.10$ $44$ 410.20.0 $8.25$ 12.115 $192.27$ $62.68$ $103.00$ $41.9$ $29.95$ $66$ 410.30.010.4511.244 $187.56$ $62.71$ $95.60$ $566.0$ $40.39$ $88$ 410.50.012.5911.020 $182.82$ $63.13$ $88.96$ $786.1$ $48.76$ $99$ <tr< td=""><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td></tr<>										
400.       20.0       8.73       11.082       189.72       61.61       103.57       432.5       31.55       66         400.       30.0       10.83       10.292       185.21       61.59       94.85       583.8       42.31       8         400.       40.0       12.03       10.101       182.55       61.78       90.45       705.3       50.85       9         400.       60.0       13.52       10.282       179.11       62.26       86.47       891.8       65.33       10         400.       60.0       13.52       10.282       179.11       62.26       86.47       891.8       65.33       10         410.       0.1       0.03       18.516       248.03       58.51       66.94       359.2       12.48       33         410.       1.0       0.30       18.215       22.837       58.79       68.30       353.2       12.68       33         410.       2.0       0.62       17.881       222.02       59.11       70.01       346.7       12.95       33         410.       20.0       8.25       12.15       192.77       62.68       103.00       421.9       29.95       66	400.	10.0	4.20	13.937	200.75	60.92	96.53	310.4	17.94	48.7
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	400.	20.0	8.73	11.082	189.72	61.61	103.57	432.5	31.55	68.9
400.       40.0       12.03       10.101       182.55       61.78       90.45       705.3       50.85       9         400.       50.0       12.87       10.135       180.63       62.02       88.00       805.5       58.36       10         400.       60.0       13.52       10.282       179.11       62.26       86.47       891.8       65.33       10         410.       0.1       0.03       18.506       248.03       58.51       66.94       359.2       12.48       3         410.       1.0       0.30       18.215       228.37       58.79       68.30       353.2       12.68       33         410.       2.0       0.62       17.881       222.02       59.11       70.01       346.7       12.95       33         410.       10.0       3.93       14.885       203.09       61.79       93.18       318.2       17.70       44         410.       20.0       8.25       12.115       192.27       62.68       103.00       421.9       29.95       67         410.       30.0       10.45       11.244       187.56       62.71       95.60       566.1       48.76       96	400.	30.0	10.83	10.292	185.21	61.59	94.85	583.8	42.31	82.1
400. $50.0$ $12.87$ $10.135$ $180.63$ $62.02$ $88.00$ $805.5$ $58.36$ $10$ $400.$ $60.0$ $13.52$ $10.282$ $179.11$ $62.26$ $86.47$ $891.8$ $65.33$ $100.11$ $410.$ $0.1$ $0.03$ $18.506$ $248.03$ $58.51$ $66.94$ $359.2$ $12.48$ $33.100.11$ $410.$ $0.5$ $0.15$ $18.377$ $234.42$ $58.64$ $67.53$ $356.5$ $12.56$ $33.110.10.10.10.10.10.10.10.10.10.10.10.10$	400.	40.0	12.03	10.101	182.55	61.78	90.45	705.3	50.85	92.3
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	400.	50.0	12.87	10.135	180.63	62.02	88.00	805.5	58.36	101.1
400.00.015.3210.20217.1102.2006.7707.1300.3316410.0.10.0318.506248.0358.5166.94359.212.4833410.1.00.3018.215228.3758.7968.30353.212.6833410.2.00.6217.881222.0259.1170.01346.712.9533410.5.01.6916.817212.4760.1476.68329.414.1044410.10.03.9314.885203.0961.7993.18318.217.7044410.20.08.2512.115192.2762.68103.00421.929.956410.30.010.4511.244187.5662.7195.60566.040.3986410.40.011.7211.010184.8062.9091.37686.148.769410.60.013.2711.152181.2563.3787.43872.562.93107420.0.10.0319.181249.6559.7968.21360.612.8339420.0.50.1419.059236.0659.9168.76360.612.8339420.1.03.7115.804205.3162.7290.90325.917.5644420.2.07.8213.41194.7463.74102.18414.528.6267420.	400	60.0	13 52	10 282	170 11	67.76	86 17	801.8	65 22	109.0
410.0.10.0316.500246.0355.100.54359.212.485410.0.50.1518.377234.4258.6467.53356.512.563'410.1.00.3018.215228.3758.7968.30353.212.683'410.2.00.6217.881222.0259.1170.01346.712.953'410.5.01.6916.817212.4760.1476.68329.414.104'410.10.03.9314.885203.0961.7993.18318.217.704'410.20.08.2512.115192.2762.68103.00421.929.956'410.30.010.4511.244187.5662.7195.60566.040.398'410.40.011.7211.010184.8062.9091.37686.148.769'410.60.013.2711.152181.2563.3787.43872.562.9310'420.0.10.0319.181249.6559.7968.21363.112.743'420.0.50.1419.059236.0659.9168.76360.612.833'420.1.00.2918.904230.0360.0569.48357.612.943'420.2.00.6018.587223.7260.3471.05351.613.204'420.<	400.	01	13.52	10.202	248.03	58 51	66.04	250.2	03.33	108.9
410. $0.3$ $0.157$ $10.77$ $274.72$ $50.67$ $61.35$ $10.55$ $12.56$ $33$ $410.$ $2.0$ $0.62$ $17.881$ $222.02$ $59.11$ $70.01$ $346.7$ $12.95$ $33$ $410.$ $5.0$ $1.69$ $16.817$ $212.47$ $60.14$ $76.68$ $329.4$ $14.10$ $44$ $410.$ $10.0$ $3.93$ $14.885$ $203.09$ $61.79$ $93.18$ $318.2$ $17.70$ $44$ $410.$ $20.0$ $8.25$ $12.115$ $192.27$ $62.68$ $103.00$ $421.9$ $29.95$ $66$ $410.$ $30.0$ $10.45$ $11.244$ $187.56$ $62.71$ $95.60$ $566.0$ $40.39$ $86$ $410.$ $40.0$ $11.72$ $11.010$ $184.80$ $62.90$ $91.37$ $686.1$ $48.76$ $96$ $410.$ $60.0$ $13.27$ $11.152$ $181.25$ $63.37$ $87.43$ $872.5$ $62.93$ $100$ $420.$ $0.1$ $0.03$ $19.181$ $249.65$ $59.79$ $68.21$ $363.1$ $12.74$ $39$ $420.$ $0.5$ $0.14$ $19.059$ $236.06$ $59.91$ $68.76$ $360.6$ $12.83$ $39$ $420.$ $2.0$ $0.60$ $18.587$ $223.72$ $60.34$ $71.05$ $351.6$ $13.20$ $44$ $420.$ $2.0$ $0.60$ $18.587$ $223.72$ $60.34$ $71.05$ $351.6$ $13.20$ $44$ $420.$ $2.0$ $0.60$ $18.587$	410.	0.1	0.05	18 377	234 42	58.64	67.53	356.5	12.40	37.7
10. $1.0$ $1.0$ $1.0$ $1.01$ $1.203$ $1.203$ $50.77$ $50.75$ $50.75$ $50.75$ $12.05$ $31.2$ $410.$ $2.0$ $0.62$ $17.881$ $222.02$ $59.11$ $70.01$ $346.7$ $12.95$ $31.2$ $410.$ $5.0$ $1.69$ $16.817$ $212.47$ $60.14$ $76.68$ $329.4$ $14.10$ $41.10$ $410.$ $10.0$ $3.93$ $14.885$ $203.09$ $61.79$ $93.18$ $318.2$ $17.70$ $44.10$ $410.$ $20.0$ $8.25$ $12.115$ $192.27$ $62.68$ $103.00$ $421.9$ $29.95$ $67.10$ $410.$ $30.0$ $10.45$ $11.244$ $187.56$ $62.71$ $95.60$ $566.0$ $40.39$ $86.1$ $410.$ $40.0$ $11.72$ $11.010$ $184.80$ $62.90$ $91.37$ $686.1$ $48.76$ $96.10$ $410.$ $60.0$ $13.27$ $11.152$ $181.25$ $63.37$ $87.43$ $872.5$ $62.93$ $107.100$ $420.$ $0.1$ $0.03$ $19.181$ $249.65$ $59.79$ $68.21$ $363.1$ $12.74$ $39.420$ $420.$ $1.0$ $0.29$ $18.904$ $230.03$ $60.05$ $69.48$ $357.6$ $12.94$ $35.16$ $420.$ $2.0$ $0.60$ $18.587$ $223.72$ $60.34$ $71.05$ $351.6$ $13.20$ $44.20$ $420.$ $2.0$ $7.82$ $13.41$ $194.74$ $63.74$ $102.18$ $414.5$ $28.62$	410.	1.0	0.15	18 215	224.42	58 79	68 30	353.7	12.50	37.9
110. $10.0$ $10.01$ $11.01$ $121.02$ $0.011$ $10.01$ $11.01$ $11.01$ $11.01$ $410.$ $10.0$ $3.93$ $14.885$ $203.09$ $61.79$ $93.18$ $318.2$ $17.70$ $44$ $410.$ $20.0$ $8.25$ $12.115$ $192.27$ $62.68$ $103.00$ $421.9$ $29.95$ $66$ $410.$ $30.0$ $10.45$ $11.244$ $187.56$ $62.71$ $95.60$ $566.0$ $40.39$ $88$ $410.$ $40.0$ $11.72$ $11.010$ $184.80$ $62.90$ $91.37$ $686.1$ $48.76$ $96$ $410.$ $50.0$ $12.59$ $11.020$ $182.82$ $63.13$ $88.96$ $786.1$ $56.12$ $99$ $410.$ $60.0$ $13.27$ $11.152$ $181.25$ $63.37$ $87.43$ $872.5$ $62.93$ $100$ $420.$ $0.1$ $0.03$ $19.181$ $249.65$ $59.79$ $68.21$ $363.1$ $12.74$ $39$ $420.$ $0.5$ $0.14$ $19.059$ $236.06$ $59.91$ $68.76$ $360.6$ $12.83$ $39$ $420.$ $1.0$ $0.29$ $18.904$ $230.03$ $60.05$ $69.48$ $357.6$ $12.94$ $35$ $420.$ $2.0$ $0.60$ $18.587$ $223.72$ $60.34$ $71.05$ $351.6$ $13.20$ $44$ $420.$ $5.0$ $1.63$ $17.585$ $214.32$ $61.26$ $76.98$ $336.0$ $14.30$ $42$ $420.$ $20.0$ $7.82$	410.	2.0	0.50	17 881	220.57	59 11	70.01	346 7	12.00	38.8
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	110.	2.0	0.02	17.001	222.02	57.11	70.01	540.7	12.75	50.0
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	410.	5.0	1.69	16.817	212.47	60.14	76.68	329.4	14.10	41.4
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	410.	10.0	3.93	14.885	203.09	61.79	93.18	318.2	17.70	49.0
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	410.	20.0	8.25	12.115	192.27	62.68	103.00	421.9	29.95	67.7
410.40.0 $11.72$ $11.010$ $184.80$ $62.90$ $91.37$ $686.1$ $48.76$ $94.96$ 410.50.0 $12.59$ $11.020$ $182.82$ $63.13$ $88.96$ $786.1$ $56.12$ $94.96$ 410. $60.0$ $13.27$ $11.152$ $181.25$ $63.37$ $87.43$ $872.5$ $62.93$ $10'$ 420. $0.1$ $0.03$ $19.181$ $249.65$ $59.79$ $68.21$ $363.1$ $12.74$ $39.96$ 420. $0.5$ $0.14$ $19.059$ $236.06$ $59.91$ $68.76$ $360.6$ $12.83$ $39.94$ 420. $1.0$ $0.29$ $18.904$ $230.03$ $60.05$ $69.48$ $357.6$ $12.94$ $39.94$ 420. $2.0$ $0.60$ $18.587$ $223.72$ $60.34$ $71.05$ $351.6$ $13.20$ $40.4420$ 420. $5.0$ $1.63$ $17.585$ $214.32$ $61.26$ $76.98$ $336.0$ $14.30$ $42.9420$ 420. $20.0$ $7.82$ $13.141$ $194.74$ $63.74$ $102.18$ $414.5$ $28.62$ $67.9420$ 420. $20.0$ $7.82$ $13.141$ $194.74$ $63.74$ $102.18$ $414.5$ $28.62$ $67.9420$ 420. $40.0$ $11.41$ $11.928$ $187.01$ $64.01$ $92.24$ $668.8$ $46.86$ $89.946$ 420. $50.0$ $12.32$ $11.914$ $184.97$ $64.25$ $89.88$ $768.3$ $54.06$ $99.96$	410.	30.0	10.45	11.244	187.56	62.71	95.60	566.0	40.39	80.8
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	410.	40.0	11.72	11.010	184.80	62.90	91.37	686.1	48.76	90.9
$410.$ $50.0$ $12.59$ $11.020$ $182.82$ $63.13$ $88.96$ $786.1$ $56.12$ $99.4$ $410.$ $60.0$ $13.27$ $11.152$ $181.25$ $63.37$ $87.43$ $872.5$ $62.93$ $10^{\circ}$ $420.$ $0.1$ $0.03$ $19.181$ $249.65$ $59.79$ $68.21$ $363.1$ $12.74$ $39.420.5$ $420.$ $0.5$ $0.14$ $19.059$ $236.06$ $59.91$ $68.76$ $360.6$ $12.83$ $39.420.5$ $420.$ $1.0$ $0.29$ $18.904$ $230.03$ $60.05$ $69.48$ $357.6$ $12.94$ $39.420.5$ $420.$ $2.0$ $0.60$ $18.587$ $223.72$ $60.34$ $71.05$ $351.6$ $13.20$ $40.420.5$ $420.$ $5.0$ $1.63$ $17.585$ $214.32$ $61.26$ $76.98$ $336.0$ $14.30$ $42.420.5.0$ $10.0$ $3.71$ $15.804$ $205.31$ $62.72$ $90.90$ $325.9$ $17.56$ $49.420.5.20.5$ $420.5.31$ $62.72$ $90.90$ $325.9$ $17.56$ $49.420.5.20.5$ $420.5.31.6$ $12.204$ $189.87$ $63.82$ $96.22$ $550.8$ $38.67$ $77.56$ $420.$ $40.0$ $11.41$ $11.928$ $187.01$ $64.01$ $92.24$ $668.8$ $46.86$ $89.420.5$ $50.0$ $12.32$ $11.914$ $184.97$ $64.25$ $89.88$ $768.3$ $54.06$ $99.69.5$										
410. $60.0$ $13.27$ $11.152$ $181.25$ $63.37$ $87.43$ $872.5$ $62.93$ $10'$ 420.0.10.03 $19.181$ $249.65$ $59.79$ $68.21$ $363.1$ $12.74$ $39$ 420.0.50.14 $19.059$ $236.06$ $59.91$ $68.76$ $360.6$ $12.83$ $39$ 420.1.00.29 $18.904$ $230.03$ $60.05$ $69.48$ $357.6$ $12.94$ $39$ 420.2.00.60 $18.587$ $223.72$ $60.34$ $71.05$ $351.6$ $13.20$ $44$ 420.5.01.63 $17.585$ $214.32$ $61.26$ $76.98$ $336.0$ $14.30$ $42$ 420.5.01.63 $17.585$ $214.32$ $61.26$ $76.98$ $336.0$ $14.30$ $42$ 420.20.0 $7.82$ $13.141$ $194.74$ $63.74$ $102.18$ $414.5$ $28.62$ $67$ 420.30.010.08 $12.204$ $189.87$ $63.82$ $96.22$ $550.8$ $38.67$ $79$ 420.40.0 $11.41$ $11.928$ $187.01$ $64.01$ $92.24$ $668.8$ $46.86$ $89$ 420. $50.0$ $12.32$ $11.914$ $184.97$ $64.25$ $89.88$ $768.3$ $54.06$	410.	50.0	12.59	11.020	182.82	63.13	88.96	786.1	56.12	99.6
420. $0.1$ $0.03$ $19.181$ $249.65$ $59.79$ $68.21$ $363.1$ $12.74$ $34.74$ $420.$ $0.5$ $0.14$ $19.059$ $236.06$ $59.91$ $68.76$ $360.6$ $12.83$ $35.76$ $420.$ $1.0$ $0.29$ $18.904$ $230.03$ $60.05$ $69.48$ $357.6$ $12.94$ $35.76$ $420.$ $2.0$ $0.60$ $18.587$ $223.72$ $60.34$ $71.05$ $351.6$ $13.20$ $44.76$ $420.$ $5.0$ $1.63$ $17.585$ $214.32$ $61.26$ $76.98$ $336.0$ $14.30$ $47.76$ $420.$ $5.0$ $1.63$ $17.585$ $214.32$ $61.26$ $76.98$ $336.0$ $14.30$ $47.76$ $420.$ $10.0$ $3.71$ $15.804$ $205.31$ $62.72$ $90.90$ $325.9$ $17.56$ $49.76$ $420.$ $20.0$ $7.82$ $13.141$ $194.74$ $63.74$ $102.18$ $414.5$ $28.62$ $67.79$ $420.$ $30.0$ $10.08$ $12.204$ $189.87$ $63.82$ $96.22$ $550.8$ $38.67$ $79.79$ $420.$ $40.0$ $11.41$ $11.928$ $187.01$ $64.01$ $92.24$ $668.8$ $46.86$ $89.768.3$ $54.06$ $99.79.768$ $420.$ $50.0$ $12.32$ $11.914$ $184.97$ $64.25$ $89.88$ $768.3$ $54.06$ $99.88.768.3$	410.	60.0	13.27	11.152	181.25	63.37	87.43	872.5	62.93	107.3
420. $0.5$ $0.14$ $19.059$ $236.06$ $59.91$ $68.76$ $360.6$ $12.83$ $39.94$ $420.$ $1.0$ $0.29$ $18.904$ $230.03$ $60.05$ $69.48$ $357.6$ $12.94$ $39.94$ $420.$ $2.0$ $0.60$ $18.587$ $223.72$ $60.34$ $71.05$ $351.6$ $13.20$ $40.4$ $420.$ $5.0$ $1.63$ $17.585$ $214.32$ $61.26$ $76.98$ $336.0$ $14.30$ $42.0$ $420.$ $10.0$ $3.71$ $15.804$ $205.31$ $62.72$ $90.90$ $325.9$ $17.56$ $49.4$ $420.$ $20.0$ $7.82$ $13.141$ $194.74$ $63.74$ $102.18$ $414.5$ $28.62$ $67.420$ $420.$ $30.0$ $10.08$ $12.204$ $189.87$ $63.82$ $96.22$ $550.8$ $38.67$ $79.420$ $420.$ $40.0$ $11.41$ $11.928$ $187.01$ $64.01$ $92.24$ $668.8$ $46.86$ $89.420$ $420.$ $50.0$ $12.32$ $11.914$ $184.97$ $64.25$ $89.88$ $768.3$ $54.06$ $99.90$	420.	0.1	0.03	19.181	249.65	59.79	68.21	363.1	12.74	39.4
420. $1.0$ $0.29$ $18.904$ $230.03$ $60.05$ $69.48$ $357.6$ $12.94$ $39.66$ $420.$ $2.0$ $0.60$ $18.587$ $223.72$ $60.34$ $71.05$ $351.6$ $13.20$ $44.66$ $420.$ $5.0$ $1.63$ $17.585$ $214.32$ $61.26$ $76.98$ $336.0$ $14.30$ $42.66$ $420.$ $10.0$ $3.71$ $15.804$ $205.31$ $62.72$ $90.90$ $325.9$ $17.56$ $49.662$ $420.$ $20.0$ $7.82$ $13.141$ $194.74$ $63.74$ $102.18$ $414.5$ $28.62$ $67.662$ $420.$ $30.0$ $10.08$ $12.204$ $189.87$ $63.82$ $96.22$ $550.8$ $38.67$ $79.662$ $420.$ $40.0$ $11.41$ $11.928$ $187.01$ $64.01$ $92.24$ $668.8$ $46.86$ $89.67$ $420.$ $50.0$ $12.32$ $11.914$ $184.97$ $64.25$ $89.88$ $768.3$ $54.06$ $90.662$	420.	0.5	0.14	19.059	236.06	59.91	68.76	360.6	12.83	39.6
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	420.	1.0	0.29	18.904	230.03	60.05	69.48	357.6	12.94	39.8
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	420	20	0.60	19 597	222 <b>72</b>	60.24	71.05	251 6	12 20	40.4
420. $5.0$ $1.03$ $17.363$ $214.32$ $61.20$ $76.98$ $536.0$ $14.30$ $4.4.90$ $420.$ $10.0$ $3.71$ $15.804$ $205.31$ $62.72$ $90.90$ $325.9$ $17.56$ $49.90$ $420.$ $20.0$ $7.82$ $13.141$ $194.74$ $63.74$ $102.18$ $414.5$ $28.62$ $67.96$ $420.$ $30.0$ $10.08$ $12.204$ $189.87$ $63.82$ $96.22$ $550.8$ $38.67$ $79.96$ $420.$ $40.0$ $11.41$ $11.928$ $187.01$ $64.01$ $92.24$ $668.8$ $46.86$ $89.96$ $420.$ $50.0$ $12.32$ $11.914$ $184.97$ $64.25$ $89.88$ $768.3$ $54.06$ $90.90$	420.	2.0	0.00	10.30/	223.72	61 26	76.09	331.0 226.0	13.20	40.4
120. $10.0$ $3.71$ $13.004$ $200.31$ $02.72$ $90.90$ $325.9$ $17.56$ $44$ $420.$ $20.0$ $7.82$ $13.141$ $194.74$ $63.74$ $102.18$ $414.5$ $28.62$ $67$ $420.$ $30.0$ $10.08$ $12.204$ $189.87$ $63.82$ $96.22$ $550.8$ $38.67$ $79$ $420.$ $40.0$ $11.41$ $11.928$ $187.01$ $64.01$ $92.24$ $668.8$ $46.86$ $89$ $420.$ $50.0$ $12.32$ $11.914$ $184.97$ $64.25$ $89.88$ $768.3$ $54.06$ $90$	420.	5.U 10.0	1.03	11.303	214.32	62 72	70.98	225.0	14.30	42.8
120. $20.0$ $1.02$ $10.141$ $127.74$ $03.74$ $102.16$ $414.3$ $28.02$ $0$ $420.$ $30.0$ $10.08$ $12.204$ $189.87$ $63.82$ $96.22$ $550.8$ $38.67$ $79$ $420.$ $40.0$ $11.41$ $11.928$ $187.01$ $64.01$ $92.24$ $668.8$ $46.86$ $89$ $420.$ $50.0$ $12.32$ $11.914$ $184.97$ $64.25$ $89.88$ $768.3$ $54.06$ $90$	420.	20.0	3./1 7.87	13.004	104.74	63 74	50.50 102 18	525.9 414 5	1/.30	49./ 67.0
420.         40.0         11.41         11.928         187.01         64.01         92.24         668.8         46.86         89           420.         50.0         12.32         11.914         184.97         64.25         89.88         768.3         54.06         69	420.	20.0	10.02	12 204	124./4	62.87	102.10	414.J 550 Q	20.02	07.0
420.         40.0         11.41         11.928         187.01         64.01         92.24         668.8         46.86         89           420.         50.0         12.32         11.914         184.97         64.25         89.88         768.3         54.06         69	720.	50.0	10.00	12.204	107.07	05.02	30.22	220.0	30.07	19.1
420. 50.0 12.32 11.914 184.97 64.25 89.88 768.3 54.06 05	420.	40.0	11.41	11.928	187.01	64.01	92.24	668.8	46.86	89 7
	420.	50.0	12.32	11.914	184.97	64.25	89.88	768.3	54.06	98.3
420, 60.0 13.02 12.031 183.37 64.48 88.38 854.6 60.73 100	420.	60.0	13.02	12.031	183.37	64.48	88.38	854.6	60.73	106.0
430. 0.1 0.03 19.870 251.27 61.07 69.48 367.0 13.01 44	430.	0.1	0.03	19.870	251.27	61.07	69.48	367.0	13.01	41.1
430. 0.5 0.14 19.752 237.69 61.17 69.99 364.7 13.09 4	430.	0.5	0.14	19.752	237.69	61.17	69.99	364.7	13.09	41.3
## FRIEND, INGHAM, AND ELY

TABLE A3. Properties of ethane in the single-phase region $-$ C	Continued
---	-----------

T K	P MPa	ρ mol∙dm <sup>−3</sup>	H kJ/mol	S J/(mol·K)	C <sub>V</sub> J∕(mol•K)	C <sub>p</sub> J/(mol·K)	₩ m·s <sup>-1</sup>	η μPa·s	λ mW/(m·K)
430	1.0	0.29	10 604	231 68	61 30	70.66	361.0	13 20	
430.	2.0	0.59	19.302	225.40	61.56	72.11	356.5	13.46	41.5
430.	5.0	1.57	18.357	216.14	62.39	77.43	342.3	14.50	44 3
430.	10.0	3.52	16.705	207.43	63.70	89.37	333 3	17.49	50.5
430.	20.0	7.42	14.158	197.14	64.79	101.22	409.8	27.52	66.5
430.	30.0	9.73	13,168	192.14	64 93	96 71	537 9	37 15	78.9
430.	40.0	11.10	12.855	189.19	65.13	93.05	653 3	45.14	88.8
430.	50.0	12.05	12.818	187.10	65.36	90.78	751.9	52.19	97.2
430.	60.0	12.78	12.919	185.46	65.59	89.30	838.0	58.70	104.8
440.	0.1	0.03	20.571	252.89	62.33	70.74	370.8	13.27	42.8
440	0.5	0.14	20 459	730 37	67 43	71 22	368 7	13 35	43.0
440.	1.0	0.28	20.317	233.32	62.55	71.84	366 1	13.35	43.0
440.	2.0	0.57	20.029	227.07	62.79	73.18	361.2	13.71	43.8
440.	5.0	1 52	19 134	217.93	63 53	78.00	348.4	14 71	45.0
440.	10.0	3.35	17.594	209.47	64.71	88.37	340.4	17.48	51.5
440	20.0	7.06	15 165	100.45	(5.97	100.04	407.1	26.62	
440.	20.0	/.00	13.103	199.45	05.83	100.24	407.1	20.62	66.3
44U. 440	<i>3</i> 0.0	9.39 10.91	14.157	194.37	00.04	97.09	521.2	33.81	78.4
440. 440	40.0	10.81	13.720	191.34	00.24	93.80	037.J	43.38	88.0
440.	60.0	12.54	13.817	189.19	66.70	90.21	822.5	56.84	103.9
450	0.1	0.02	<b>01 005</b>	<b>05</b> 4 40	(a 50			10.50	
450.	0.1	0.03	21.285	254.49	63.59	71.99	374.6	13.52	44.6
450.	0.5	0.13	21.1//	240.93	63.67	72.44	372.7	13.60	44.8
450.	1.0	0.27	21.041	234.95	63.78	73.02	370.3	13.71	45.0
450.	2.0	0.56	20.766	228.73	64.00	74.26	365.7	13.96	45.5
450.	5.0	1.47	19.917	219.69	64.68	78.65	354.2	14.92	47.5
450.	10.0	3.21	18.474	211.45	65.75	87.77	347.2	17.50	52.6
450.	20.0	6.73	16.163	201.69	66.86	99.31	406.2	25.88	66.3
450.	30.0	9.07	15.110	196.56	67.13	97.40	518.5	34.62	78.0
450.	40.0	10.52	14.731	193.45	67.35	94.51	627.3	42.17	87.5
450.	50.0	11.54	14.651	191.26	67.58	92.48	723.4	48.89	95.7
450.	60.0	12.31	14.723	189.56	67.81	91.10	808.3	55.12	103.1
460.	0.1	0.03	22.011	256.09	64.83	73.23	378.4	13.78	46.4
460.	0.5	0.13	21.907	242.54	64.91	73.65	376.6	13.86	46.6
460.	1.0	0.27	21.777	236.56	65.01	74.20	374.4	13.96	46.8
460.	2.0	0.54	21.514	230.37	65.22	75.35	370.2	14.20	47.3
460.	5.0	1.43	20.707	221.42	65.83	<b>79.37</b>	359.8	15.14	49.1
460.	10.0	3.08	19.350	213.38	66.80	87.47	353.8	17.55	53.8
460.	20.0	6.43	17.152	203.87	67.90	98.50	406.5	25.27	66.5
460.	30.0	8.76	16.085	198.70	68.22	97.64	511.4	33.58	77.8
400.	40.0	10.25	12.0/2	195.54	08.45	95.17	010.0	40.90	87.1
460.	50.0	11.29	15.579	193.30	68.68	93.29	711.0	47.46	95.2
460.	60.0	12.08	15.639	191.57	68.91	91.97	795.1	53.53	102.4
470.	0.1	0.03	22.749	257.67	66.06	74.46	382.1	14.03	48.2
470.	0.5	0.13	22.650	244.13	66.14	74.85	380.5	14.11	48.4
470.	1.0	0.26	22.525	238.17	66.23	75.37	378.5	14.22	48.6
470.	2.0	0.53	22.273	232.00	66.42	76.45	374.6	14.45	49.0
470.	5.0	1.39	21.505	223.14	66.99	80.14	365.3	15.35	50.8
470.	10.0	2.97	20.224	215.26	67.87	87.40	360.2	17.63	55.2
470.	20.0	6.16	18.133	205.98	68.94	97.82	407.9	24.78	67.0
470.	30.0	8.47	17.063	200.80	69.31	97.86	505.8	32.67	77.8
470.	40.0	9.98	16.634	197.59	69.55	95.79	607.3	39.75	86.9
470.	50.0	11.04	16.516	195.32	69.78	94.07	699.9	46.14	94.8
470.	60.0	11.85	16.563	193.56	70.00	92.82	783.0	52.07	102.0
480.	0.1	0.03	23,500	259.25	67.28	75.67	385.8	14.28	50.1
480	0.5	0.13	23 404	245 72	67.25	76.05	204.2	14.76	50.2

## THERMOPHYSICAL PROPERTIES OF ETHANE

T K	Р MPa	ρ mol·dm <sup>-3</sup>	H kJ/mol	S J/(mol·K)	C <sub>V</sub> J/(mol⋅K)	$\frac{C_p}{J/(\text{mol}\cdot K)}$	W m·s <sup>-1</sup>	η μPa·s	λ mW/(m·K)
480.	1.0	0.25	23.284	239.77	67.44	76.53	382.4	14.46	50.4
480.	2.0	0.52	23.043	233.63	67.62	77.54	379.0	14.69	50.9
480.	5.0	1.35	22.310	224.83	68.14	80.95	370.6	15.56	52.5
480.	10.0	2.86	21.099	217.10	68.95	87.51	366.4	17.72	56.6
480.	20.0	5.92	19.109	208.03	69.98	97.30	409.9	24.38	67.6
480.	30.0	8.20	18.042	202.86	70.38	98.05	501.5	31.87	78.0
480.	40.0	9.72	17.595	199.61	70.63	96.37	599.2	38.72	86.9
480.	50.0	10.81	17.461	197.31	70.87	94.83	689.9	44.94	94.6
480.	60.0	11.63	17.495	195.52	71.09	93.66	771.9	50.73	101.7
<b>490.</b>	0.1	0.02	24.263	260.83	68.49	76.88	389.4	14.53	51.9
490.	0.5	0.12	24.171	247.30	68.56	77.23	388.1	14.61	52.1
490.	1.0	0.25	24.056	241.36	68.64	77.69	386.4	14.71	52.3
490.	2.0	0.50	23.824	235.24	68.80	78.64	383.2	14.93	52.7
490.	5.0	1.31	23.124	226.51	69.28	81.80	375.7	15.78	54.3
490.	10.0	2.77	21.975	218.90	70.03	87.76	372.3	17.83	58.0
490.	20.0	5.69	20.080	210.03	71.02	96.92	412.6	24.06	68.3
490.	30.0	7.94	19.024	204.89	71.45	98.23	498.4	31.17	78.4
490.	40.0	9.47	18.561	201.61	71.71	96.94	592.4	37.78	87.0
490	50.0	10.58	18.413	199.27	71.95	95.56	680.9	43.84	94.6
490.	60.0	11.42	18.436	197.46	72.17	94.48	761.8	49.49	101.5
500.	0.1	0.02	25.037	262.39	69.69	78.07	393.1	14.78	53.8
500.	0.5	0.12	24.949	248.87	69.75	78.41	391.8	14.85	54.0
500.	1.0	0.24	24.838	242.94	69.82	78.84	390.3	14.95	54.1
500.	2.0	0.49	24.616	236.84	69.98	79.73	387.4	15.17	54.5
500.	5.0	1.28	23.946	228.17	70.43	82.67	380.7	15.99	56.0
500.	10.0	2.68	22.854	220.68	71.12	88.12	378.1	17.95	59.6
500.	20.0	5.49	21.048	211.99	72.07	96.68	415.7	23.80	69.2
500.	30.0	7.69	20.007	206.87	72.51	98.43	496.2	30.56	78.8
500.	40.0	9.23	19.533	203.57	72.79	97.48	586.5	36.95	87.2
500.	50.0	10.35	19.372	201.21	73.02	96.28	672.9	42.83	94.6
500.	60.0	11.21	19.385	199.38	73.24	95.28	752.5	48.34	101.4

TABLE A3. Properties of ethane in the single-phase region - Continued

Values of the density, enthalpy, entropy, isochoric and isobaric heat capacities, speed of sound, viscosity, and thermal conductivity in the singlephase region of the ethane fluid. The independent variables were chosen to be temperature and pressure. The density was evaluated by inverting the pressure equation in Table 7; the quantities  $H, S, C_{\nu}, C_{p}$ , and W were then evaluated directly from the expressions in Table 7. The viscosity and thermal conductivity are from Eqs. (8) and (9) [with the terms evaluated from Eqs. (10-20)]; the density input is from column 3 of this table.