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Phase Behavior of System Methane + Hydrogen Sulfide
Stefano Langè\textsuperscript{a}, Marco Campestrini\textsuperscript{b}, Paolo Stringari\textsuperscript{b,*}

\textsuperscript{a}Politecnico di Milano– Dipartimento di Chimica, Materiali e Ingegneria Chimica"Giulio Natta", Piazza Leonardo da Vinci 32, I-20133, Milan, Italy

\textsuperscript{b}MINES ParisTech, PSL Research University, CTP – Centre of Thermodynamics of Processes, 35 rue St Honoré 77300 Fontainebleau, France

* Corresponding author. Tel.: +33 (0)1 64 69 4857. E-mail address: paolo.stringari@mines-paristech.fr
Abstract

An accurate description of the phase behavior of the CH$_4$+H$_2$S system is given for temperatures from 70 K to the critical temperature of H$_2$S and pressures up to 250 MPa. The study includes the solid phases of CH$_4$ and H$_2$S. A global pressure-temperature diagram is presented. The types of temperature-composition and pressure-composition phase diagrams that can be encountered in the studied temperature and pressure ranges have been described. The temperature and pressure ranges where the phase behavior of the system changes have been identified and a representative phase diagram is presented for each range. Phase diagrams have been obtained through the solid-liquid-vapor equation of state proposed by Yokozeki. The parameters of the equation of state have been regressed on all the available phase equilibrium data for the considered system.

Keywords: Natural gas purification, Methane-hydrogen sulfide mixture, Equation of state, Phase diagram, Solid-liquid-vapor equilibrium
Introduction

The global energy demand is expected to increase rapidly over the next twenty years. The rise of the primary energy demand is predicted to be 41% between 2012 and 2035. Among fossil fuels consumptions, the natural gas ones are projected to have the most rapid growth trend.

Recent studies have shown that 40% of the remaining natural gas reserves are sour and/or with high CO\textsubscript{2} content (15%-80%). In these low quality gas fields, the hydrogen sulfide content in certain gas reservoirs could be high, even up to about 15%.

In this scenario, a detailed description of the phase behavior of systems containing methane and hydrogen sulfide is necessary to perform the correct process design of new gas purification technologies that are required to allow the profitable production of commercial-grade gas from low quality natural gas reserves.

In the range of temperature and pressure investigated in this work (T from 70 K to the critical temperature of H\textsubscript{2}S and P up to 250 MPa), the system CH\textsubscript{4} (1) + H\textsubscript{2}S (2) can present two liquid phases (L\textsubscript{1}, rich in CH\textsubscript{4}, and L\textsubscript{2}, rich in H\textsubscript{2}S) and two solid phases (S\textsubscript{1}, rich in CH\textsubscript{4}, and S\textsubscript{2}, rich in H\textsubscript{2}S) in addition to the vapor phase (V).

Several authors have studied the methane - hydrogen sulfide system from the experimental point of view. The first literature work presenting experimental values for the methane - hydrogen sulfide mixture was published by Reamer et al. in 1951. The authors investigated the critical locus and the vapor-liquid equilibrium of the mixture at 277.59 K, 310.93 K and 344.26 K.

A total of 6 experimental values of VL\textsubscript{2}E were measured by Robinson and coauthors: the correspondent temperatures are 310 K, Robinson and Bailey, and 277 K, Robinson et al.

A more complete experimental investigation of the phase behavior of the system down to 170 K was carried out by Kohn and Kurata in 1958. The different mixture compositions under study leaded the authors inferring the presence of two different solid-liquid-vapor equilibrium loci and a solid-liquid-liquid-vapor Quadruple Point (QP). The first S\textsubscript{2}L\textsubscript{VE} locus has a methane rich liquid phase (S\textsubscript{2}L\textsubscript{1}VE), the second a hydrogen sulfide rich liquid phase (S\textsubscript{2}L\textsubscript{2}VE). From the QP, a
L₂L₁VE and a S₂L₂L₁E curves originate. The L₂L₁VE curve ends to a first Upper Critical EndPoint (UCEP₁), L₂(L₁=V). Kohn and Kurata treated the solid phase S₂ as pure H₂S. ²⁵

Furthermore, Kohn and Kurata argued the presence of a L₂=V critical locus, which is continuous with a L₁=L₂ critical locus at high pressure. This latter ends to a second Upper Critical EndPoint at high pressure (UCEP₂), S₂(L₁=L₂), where the critical curve meets the S₂L₁L₂E curve. In addition to these points, same authors claimed the presence of a second quadruple point at a temperature close to the triple point temperature of methane, where two solid phases (S₁ and S₂), a CH₄-rich liquid phase, and a vapor phase coexist in equilibrium.²⁵

The system described by Kohn and Kurata is then of Type - III according to the classification of phase diagrams of Van Konynenburg and Scott,²⁶ or 1⁰PAl₅Q according to the one by Bolz et al.²⁷ Furthermore, the mixture presents two totally immiscible solid phases (S₁ and S₂) in the low temperature region.

Cheung and Zander²⁸ reported few experimental points for the S₂L₁E region, while in 1991 supplementary VL₂E values were obtained by Yarim-Agaev et al.²⁹ for temperatures ranging from 222 K up to 273 K. More recently, Coquelet et al.³⁰ have experimentally investigated the VL₂E behavior together with the immiscibility gap in the liquid phase at 186.25 K and 203.4 K, thus proposing the first literature values of L₁L₂E.

From the modeling point of view, different approaches have been used in the literature to represent the phase behavior of the mixture of interest in this work. Equations of state have been largely used to this scope. Examples are the works by Huron et al.,³¹ Adachi et al.,³² Mohsen-Nia et al.,³³ and Sakoda and Uematsu.³⁴-³⁵

Huron et al.³¹ used the Soave-Redlich-Kwong Equation of State (EoS); Adachi et al.³² adopted a cubic EoS with four parameters; Mohsen-Nia et al.³³ made use of a two constant cubic EoS; an EoS in the Helmholtz free energy form was used by Sakoda and Uematsu³⁴-³⁵ for representing the phase behavior and thermal properties of the system. Same authors investigated the Pressure-Temperature
(PT) diagram including only equilibria between fluid phases (namely VL₂E, L₂L₁VE and the critical locus of the system), without any extension to phase equilibria involving solid phases.\textsuperscript{34-35} Privat et al.\textsuperscript{36} used a temperature dependent $k_{ij}$ parameter within the group contribution model PPR78 and compared the calculated values with literature data of equilibria involving fluid phases.

Tsivintzelis et al.\textsuperscript{37} studied the methane-hydrogen sulfide system with the CPA EoS considering four different association schemes for H₂S. As in the works by Sakoda and Uematsu,\textsuperscript{34-35} and Privat et al.,\textsuperscript{36} Tsivintzelis et al.\textsuperscript{37} represented only fluid phases.

In 2010, Tang and Gross\textsuperscript{28} have published the first modeling work involving also the apparition of the solid phase at low temperature. The authors used the PCP-SAFT EoS and the PC-SAFT EoS for the fluid phases and a fugacity equation for the solid phase considered as pure H₂S. Nevertheless, the presented study is limited to temperatures higher than the temperature of the $S_2L_2L_1V$ quadruple point.\textsuperscript{38}

In this work the detailed phase behavior of the CH₄ + H₂S system has been investigated using the analytical equation of state for Solid, Liquid and Vapor phases, SLV EoS.\textsuperscript{39} The phase diagram has been studied for temperatures from 70 K up to the critical temperature of H₂S and for pressures up to 250 MPa. The CH₄ + H₂S phase behavior has been investigated for the first time in a so wide range of temperature and pressure. With respect to the cited literature works on the phase behavior of the CH₄ + H₂S system, this work adds new information about the phase diagram, especially for the low temperature region and the high pressure region. Isothermal P-x diagrams and isobaric T-x diagrams are presented for all the P-T regions in which the system changes qualitatively the phase behavior. For a chosen temperature and pressure in the investigated P-T range, the reader can find a P-x or T-x diagram representative of the behavior of the system. In practice, this work provides a map of the CH₄ + H₂S phase behavior that can be used for the process design, crystallization risk evaluation, or other purposes that need a reliable representation of the phase behavior of the system.
Solid Transitions in Pure CH\textsubscript{4} and H\textsubscript{2}S

From the authors’ knowledge, the work by Kohn and Kurata\textsuperscript{25} is the first dealing with the solid phases of the methane-hydrogen sulfide system. As stated in the previous section, these authors described the phase diagram of the mixture taking into account two solid phases made of pure CH\textsubscript{4} (S\textsubscript{1}) and pure H\textsubscript{2}S (S\textsubscript{2}), respectively.

Nevertheless, these components present different phase transitions in the solid phase. The orders and the types of the transitions that occur in the molecular crystals of methane and hydrogen sulfide have been widely discussed in the literature\textsuperscript{40-43}, and agreements have been reached about the solid-solid transition temperatures and the related thermodynamic properties.

According to Colwell et al.\textsuperscript{40} methane has two possible crystal lattices (α and β) resulting in a solid\textsubscript{α}-solid\textsubscript{β} transition and a related solid\textsubscript{α}-solid\textsubscript{β}-vapor triple point located at 20.5 K and vapor pressure (P\texttextless;0). Crystal lattices α and β have been stated to have a face-centered cubic (fcc) structure, and the solid transition involves uniquely changes in the degree of orientational order of the fcc structure.\textsuperscript{40} In 1957, Stevenson\textsuperscript{41} used a strong cylindrical container for measuring pressure-volume isotherms and found two additional solid phases (γ and δ). As a consequence, the pressure-temperature diagram of pure methane presents other three solid-solid transition boundaries (β-γ, α-γ, and γ-δ) and a solid\textsubscript{α}-solid\textsubscript{β}-solid\textsubscript{γ} triple point located at about 32 K and 260 MPa. In the low temperature region, the β-γ and γ-δ phase equilibrium curves do not match any other phase transition curve, and they go toward absolute zero at about 90 MPa and 210 MPa, respectively. Furthermore, the γ-δ boundary extends in the high pressure region remaining always located at temperatures lower than 35 K.\textsuperscript{41}

In 1936, Giauque and Blue\textsuperscript{42} carried out experimental measurements for pure H\textsubscript{2}S by means of a calorimetric apparatus and observed two solid-solid transitions in addition to the freezing transition. Authors observed a region of considerable energy absorption at 126 K and a sharp transition at 103.5 K. The former solid-solid transition was explained as an orientational ordering with respect to the rotational vibrations of the hydrogen atoms, whereas a drop in the dielectric constant was
associated to the latter. Giauque and Blue\textsuperscript{42} underlined then the presence of three fcc structures ($\alpha$, $\beta$, and $\gamma$) for solid H\textsubscript{2}S, and suggested three triple points (solid$_\alpha$-liquid-vapor, solid$_\alpha$-solid$_\beta$-vapor, and solid$_\beta$-solid$_\gamma$-vapor) at vapor pressure (P=0). Successively, Stevenson\textsuperscript{43} showed the presence of other two solid phases ($\delta$ and $\kappa$) of H\textsubscript{2}S in the high pressure region by a volume discontinuity method. The slopes of the related solid-solid boundaries are such as to originate four solid-solid-solid triple points between 110 K and 160 K, and 400 MPa and 900 MPa.\textsuperscript{43}

All the solid-solid-solid triple points of pure methane and hydrogen sulfide occur at pressures higher than 250 MPa,\textsuperscript{40-43} then this limit has been considered for the application of the thermodynamic model used in this work. With respect to the low pressure region and the solid-solid-vapor triple points, only pure H\textsubscript{2}S presents more than one solid phase in the range of temperature chosen in this work (70 – 373K). Nevertheless, these solid-solid transitions do not involve a meaningful change in the solid-fluid equilibria of the mixture seeing that the solid phase S\textsubscript{2} is always made of fcc crystals of pure H\textsubscript{2}S. This aspect has been pointed out in the last section of this work, where the presence of two additional solid phases for pure H\textsubscript{2}S have been qualitatively taken into account in a temperature-composition diagram and compared with the case of considering a single solid phase.
**Thermodynamic Model**

The pressure-explicit equation of state proposed by Yokozeki in 2003 is presented in Eq. 1.  

\[
P(T, v) = \frac{RT}{v-b} v - c - \frac{a}{v^2 + qbv + rb^2} 
\]

In Eq. 1, \(P\) is the pressure, \(R\) is the gas constant, \(T\) is the temperature, \(v\) is the molar volume, \(a\) keeps into account the attractive forces among molecules, \(c\) is the liquid covolume, \(b\) is the solid covolume, \(d\) is molar volume for which the repulsive term in Eq. 1 is null, \(q\) and \(r\) are parameters of the attractive term. The van der Waals attractive term \((q, r = 0)\) has been considered in this work.

According to Yokozeki, parameters \(a\) and \(b\) in Eq. 1 are temperature dependent functions whose functional forms are given in Eqs. 2 and 3:

\[
a(T) = \left(\frac{RT_c}{P_c}\right)^2 \left[a_0 + a_1 T_r \exp\left(-a_2 T_r^n\right)\right] 
\]

\[
b(T) = v_c \left[b_0 + b_1 \exp\left(-b_2 T_r^m\right)\right] 
\]

where \(T_r = T/T_c\) is the reduced temperature, \(a_0, a_1, a_2, n, b_0, b_1, b_2, m\) and \(m\) are parameters, \(R\) is the gas constant, and \(T_c, P_c\), and \(v_c\) are temperature, pressure, and volume at the critical point.

The application of Eqs. 1-3 for the representation of the phase equilibrium behavior of a pure component requires the previous evaluation of 11 parameters (the critical volume \(v_c\) or equivalently the critical compressibility coefficient \(Z_c\), \(a_0, a_1, a_2, n, b_0, b_1, b_2, m, c, \) and \(d\)). The procedure deeply illustrated by Stringari et al. has been used to this scope.

This procedure is based on the analytical representation of the critical and triple points of a pure component, and on the minimization of an objective function based on the iso-fugacity conditions along the saturation, sublimation, and melting curves. The fugacities of the coexisting phases are evaluated at the experimental conditions (temperature and pressure) of equilibrium.

For CH\(_4\), pseudo-experimental temperature-pressure couples for vapor-liquid, solid-vapor, and solid-liquid equilibria have been generated from accurate auxiliary equations considering these values as true equilibrium values. Therefore, it has been considered that comparing the model with
the values obtained from the auxiliary equations (referred to as auxiliary values) is equivalent to
compare the model to experimental data, seeing that these auxiliary equations are precise
correlations of the available experimental data.

For H$_2$S, only the auxiliary equation for the vapor-liquid equilibrium is available\textsuperscript{45}. Experimental
values of sublimation have been proposed by Stevenson,\textsuperscript{43} and Clark et al.,\textsuperscript{46} while temperature-
pressure couples of solid-liquid equilibrium have been generated by means of Eq. 4.

\[
P^{SLE} = P_t + \frac{\Delta H^{SLE}}{\Delta v^{SLE}} \left( 1 + \frac{T_s}{T} \right)
\]  

Eq. 4 has been obtained from the expression relating the solid and the liquid fugacities of a pure component, also known as “classic approach”.\textsuperscript{47} In Eq. 4, \(P_t\) is the triple point pressure, \(T_t\) is the
triple point temperature, \(\Delta H^{SLE}=H_L-H_S\) and \(\Delta v^{SLE}=v_L-v_S\) are latent heat of fusion and the volume change upon melting at the triple point, whose values are 2.377 kJ/mol,\textsuperscript{48} and 3.892 cm$^3$/mol,\textsuperscript{49} respectively.

The references of VLE, SLE, and SVE data needed for the regression of parameters of the pure components are summarized in Table 1.

<table>
<thead>
<tr>
<th>Substance</th>
<th>VLE</th>
<th>SLE</th>
<th>SVE</th>
</tr>
</thead>
<tbody>
<tr>
<td>CH$_4$</td>
<td>auxiliary values\textsuperscript{45}</td>
<td>auxiliary values\textsuperscript{45}</td>
<td>auxiliary values\textsuperscript{45}</td>
</tr>
<tr>
<td>H$_2$S</td>
<td>auxiliary values\textsuperscript{45}</td>
<td>generated values, Eq. 4</td>
<td>experimental values\textsuperscript{43,46}</td>
</tr>
</tbody>
</table>

For sake of simplicity, the term data has been henceforth used to group all the kind of values: auxiliary values obtained from auxiliary equations,\textsuperscript{45} experimental values of SVE for H$_2$S,\textsuperscript{43,46} and SLE values calculated by Eq. 4 for H$_2$S,\textsuperscript{47}.

Values of pressure and temperature at critical and triple points for methane and hydrogen sulfide are presented in Table 2.\textsuperscript{45}
Table 2. Temperature and pressure of triple and critical points of methane and hydrogen sulfide.\(^{45}\)

<table>
<thead>
<tr>
<th>Substance</th>
<th>(P_t/ \text{kPa})</th>
<th>(T_t/ \text{K})</th>
<th>(P_c/ \text{MPa})</th>
<th>(T_c/ \text{K})</th>
</tr>
</thead>
<tbody>
<tr>
<td>CH(_4)</td>
<td>11.697</td>
<td>90.694</td>
<td>4.5992</td>
<td>190.56</td>
</tr>
<tr>
<td>H(_2)S</td>
<td>22.3</td>
<td>187.7</td>
<td>9</td>
<td>373.1</td>
</tr>
</tbody>
</table>

For each substance, the P-T range used for the regression of the parameters within the SLV EoS extends from a minimum on the sublimation branch to a maximum on the melting branch. These values are presented in Table 3.

Table 3. Temperature and pressure ranges for the regression of the parameters for methane and hydrogen sulfide.

<table>
<thead>
<tr>
<th>Substance</th>
<th>SVE</th>
<th>SLE</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(T_{\text{min}}/ \text{K})</td>
<td>(P_{\text{min}}/ \text{kPa})</td>
</tr>
<tr>
<td>CH(_4)</td>
<td>60</td>
<td>0.0164</td>
</tr>
<tr>
<td>H(_2)S</td>
<td>130</td>
<td>0.0417</td>
</tr>
</tbody>
</table>

Regressed parameters are \(Z_c\), \(a_0\), \(a_1\), \(a_2\), \(n\), \(b_0\), \(b_1\), \(b_2\), \(m\), \(c\), and \(d\) for CH\(_4\) and H\(_2\)S are reported in Table 4. Values for CH\(_4\) are the same presented by Stringari et al.,\(^{44}\) and here reported for the reader convenience.

Table 4. Parameters of Eqs. 1-3 for methane and hydrogen sulfide.

<table>
<thead>
<tr>
<th>Substance</th>
<th>(Z_c/ (x10^3))</th>
<th>(a_0/ (x10^6))</th>
<th>(a_1)</th>
<th>(a_2)</th>
<th>(n)</th>
<th>(b_0)</th>
<th>(b_1)</th>
<th>(b_2)</th>
<th>(m/ (x10^3))</th>
<th>(c/ (x10^3))</th>
<th>(d/ (x10^3))</th>
</tr>
</thead>
<tbody>
<tr>
<td>CH(_4)</td>
<td>37.50400</td>
<td>3.903</td>
<td>21.983</td>
<td>4.051</td>
<td>0.381</td>
<td>0.335</td>
<td>−0.334</td>
<td>4.201</td>
<td>0.682</td>
<td>4.468</td>
<td>4.422</td>
</tr>
<tr>
<td>H(_2)S</td>
<td>37.50023</td>
<td>0.998</td>
<td>11.043</td>
<td>3.289</td>
<td>0.503</td>
<td>0.334</td>
<td>−0.236</td>
<td>5.160</td>
<td>1.370</td>
<td>4.430</td>
<td>4.426</td>
</tr>
</tbody>
</table>

The comparison of the phase equilibrium values calculated with Eq. 1 and data is presented in Table 5 in terms of Absolute Average Deviation, Bias, and Maximum Absolute Deviation. Table 5 represents the deviations obtained calculating the equilibrium pressure at fixed temperature; for each kind of equilibrium, N is the number of data used for the comparison. Values for CH\(_4\) are the same presented by Stringari et al.\(^{44}\)
Table 5. Summary of the statistical errors in calculating equilibrium pressures at fixed temperatures for CH\(_4\) and H\(_2\)S.

Errors are evaluated with respect to the auxiliary values of VLE, SLE, SVE of CH\(_4\) and of VLE of H\(_2\)S, to the experimental values of SVE of H\(_2\)S, and to the values generated from Eq. 4 for SLE of H\(_2\)S.

<table>
<thead>
<tr>
<th>Substance</th>
<th>VLE</th>
<th>SLE</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>AAD</td>
<td>Bias</td>
</tr>
<tr>
<td>CH(_4)</td>
<td>0.11</td>
<td>-0.07</td>
</tr>
<tr>
<td>H(_2)S</td>
<td>0.19</td>
<td>0.85</td>
</tr>
</tbody>
</table>

According to Table 5, the SLV EoS agrees well with the auxiliary values\(^{45}\) of melting, saturation, and sublimation obtained for CH\(_4\). Considering all the kind of equilibrium, the AAD is always lower than 0.3%, the Bias is close to zero, and the highest MAD of about 1.4% is obtained for the SLE.

Similar results are related to the comparison between the SLV EoS and the auxiliary values\(^{45}\) of saturation of H\(_2\)S; the AAD is 0.19%, the Bias lower than 1%, the MAD is about 4%.

Higher deviations are obtained for the quantitative comparison between calculated and experimental pressures of SVE and SLE of H\(_2\)S. With respect to the SVE, the highest deviation (12.38%) occurs at 130K where the experimental pressure is lower than 5\times10^{-5} MPa. For the SLE, the AAD, Bias and MAD are about 8%, 7%, and 13%, respectively. It should be remembered that the pressure of melting have been calculated by means of Eq. 4, which is a simple approximation of the solid-liquid equilibrium.

Pressure-temperature equilibrium behaviors obtained with the SLV EoS are represented in Figures 1-2 for CH\(_4\) and H\(_2\)S, respectively; empty symbols are selected data of VLE, SLE, and VLE.
Figure 1. Pressure-temperature phase diagram of CH₄.
—: SLV EoS; □: auxiliary values for SVE, SLE, and VLE.

Figure 2. Pressure-temperature phase diagram of H₂S.
—: SLV EoS; □: VLE auxiliary values; ○: SLE generated values, Eq. 4; Δ: SVE experimental values.
Extension of Eq. 1 to the binary mixture has been considered using the mixing rules proposed by Yokozeki\textsuperscript{39}, Eqs. 5-8. The consistency test for the adopted mixing rules has been presented in Ref. 44.

\[ a = \sum_{i,j=1}^{NC} \sqrt{a_i a_j} (1 - k_{ij}) x_i x_j \]  
\[ b = \sum_{i=1}^{NC} b_i x_i \]  
\[ c = \sum_{i=1}^{NC} c_i x_i \]  
\[ d = \sum_{i=1}^{NC} d_i x_i \]

The binary interaction parameter \( k_{ij} \) has been regressed by minimization of an objective function defined as the sum of the absolute difference between the calculated and experimental methane compositions, averaged with respect to the total number \( N \) of VL\(_2\)E, L\(_2\)L\(_1\)E and S\(_2\)L\(_1\)E data, Eq. 9.

\[ f_{ob} = \left[ \frac{1}{N} \sum_{i=1}^{N} |x_{CH_4}^{calc} - x_{CH_4}^{exp}| \right] \]  

In Eq. 9, each calculated composition of methane refers to the equilibrium composition at VL\(_2\)E or L\(_2\)L\(_1\)E obtained by the model by means of a PT flash calculation at imposed experimental temperature and pressure.

In spite of the complex phase equilibrium behavior, a not temperature dependent value for \( k_{ij} \) (\( k_{ij} = 0.058 \)) has been found to allow the complete description of the methane-hydrogen sulfide system from 70 K up to the critical temperature of H\(_2\)S.

Section 4 presents the comparisons between experimental values of equilibrium and the SLV EoS, while Section 5 illustrates the complete phase diagram of the binary mixture in the temperature and pressure ranges of this work.

As it is possible to observe in the figures presented in the following sections, the solid phases S\(_2\) and S\(_1\) calculated by the model are not always pure phases. This discrepancy with what is expected
according to the literature depends heavily on the functional form of Eqs. 5-8. Because the mixing rules used for the volumetric parameter \( b, c, \) and \( d \) are of the type developed for the fluid phase, it follows that the solid phase is treated as a liquid-like phase, which entails the solubility in the solid phase. The fact that the solid phases are not constituted by a pure component does not affect the quality of the representation of the temperature-pressure-composition data of the fluid phases in equilibrium with the solid.
Model Validation

This section has the scope to assess the accuracy and reliability of the developed SLV EoS in representing the existing phase equilibrium data. Once the SLV EoS validated, it will be used in Section 5 for predicting the phase equilibrium behavior of the CH$_4$ + H$_2$S system in the considered range of temperature and pressure.

The quantitative comparison between the values calculated by means of Eq. 1, with the mixing rules of Eqs. 5-8, and the experimental values proposed in the literature is illustrated in Table 6. The first part of Table 6 presents the comparison between calculated and experimental values of VL$_2$E, L$_1$L$_2$E, and S$_2$L$_1$E; errors are in terms of the compositions of methane in the liquid and the vapor phase, obtained by means of a PT flash calculation at imposed experimental temperature and pressure.

The second part of Table 6 portrays the comparison with respect to the experimental values of temperature and pressure related to the three-phase equilibria S$_2$L$_1$VE, S$_2$L$_2$VE, L$_2$L$_1$VE, and S$_2$L$_2$L$_1$E. Errors in terms of temperature have been obtained comparing the experimental value with the temperature calculated imposing the experimental pressure of three-phase equilibrium. Errors in terms of pressure have been obtained comparing the experimental value with the pressure calculated imposing the experimental temperature of three-phase equilibrium.

Finally, the last two rows present the comparison along the Critical Curve (CC) of the mixture; errors have been evaluated comparing experimental and calculated values of temperature and pressure at fixed composition of methane.
Table 6. Quantitative comparison of equilibrium compositions, temperatures and pressures of three-phase equilibria, and critical temperatures and pressures for the system xCH₄+(1-x)H₂S.

<table>
<thead>
<tr>
<th>Reference</th>
<th>N</th>
<th>Kind of data</th>
<th>T</th>
<th>P</th>
<th>x</th>
<th>y</th>
<th>PT FLASH</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>K</td>
<td>MPa</td>
<td></td>
<td></td>
<td>x</td>
</tr>
<tr>
<td>Reamer et al. 1951</td>
<td>59</td>
<td>VL₂E TPxy</td>
<td>277-344</td>
<td>1.4-13.4</td>
<td>0.0057-0.55</td>
<td>0.059-0.73</td>
<td>N calc</td>
</tr>
<tr>
<td>Robinson and Bailey 1957</td>
<td>3</td>
<td>VL₂E TPxy</td>
<td>310</td>
<td>4.1-12.4</td>
<td>0.033-0.26</td>
<td>0.29-0.51</td>
<td>N calc</td>
</tr>
<tr>
<td>Kohn and Kurata 1958</td>
<td>54</td>
<td>VL₂E TPxy</td>
<td>188-344</td>
<td>1.4-11.0</td>
<td>0.005-0.24</td>
<td>0.10-0.97</td>
<td>N calc</td>
</tr>
<tr>
<td>Robinson et al. 1959</td>
<td>3</td>
<td>VL₂E TPxy</td>
<td>277</td>
<td>2.8-11.0</td>
<td>0.023-0.26</td>
<td>0.51-0.72</td>
<td>N calc</td>
</tr>
<tr>
<td>Cheung and Zander 1968</td>
<td>6</td>
<td>S₂L₁E Tx</td>
<td>119-162</td>
<td></td>
<td>0.98-0.99</td>
<td></td>
<td>N calc</td>
</tr>
<tr>
<td>Yarim-Agaev et al. 1991</td>
<td>42</td>
<td>VL₂E TPxy</td>
<td>222-273</td>
<td>0.7-11.8</td>
<td>0.001-0.36</td>
<td>0.14-0.90</td>
<td>N calc</td>
</tr>
<tr>
<td>Coquelet et al. 2014</td>
<td>14</td>
<td>VL₂E TPxy</td>
<td>186-203</td>
<td>0.5-5.8</td>
<td>0.0091-0.13</td>
<td>0.86-0.98</td>
<td>N calc</td>
</tr>
<tr>
<td></td>
<td>10</td>
<td>L₁L₂E TPxx</td>
<td>186-203</td>
<td>6.7-10.8</td>
<td>0.090-0.14</td>
<td>0.85-0.98</td>
<td>N calc</td>
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<tr>
<td>Reference</td>
<td></td>
<td></td>
<td>T</td>
<td>P</td>
<td></td>
<td></td>
<td>THREE-PHASE EQ.</td>
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<td></td>
<td></td>
<td></td>
<td>K</td>
<td>MPa</td>
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<td></td>
<td>MAD%</td>
</tr>
<tr>
<td>Kohn and Kurata 1958</td>
<td>3</td>
<td>S₂L₁VE¹</td>
<td>167-168</td>
<td>2.05-2.2</td>
<td></td>
<td></td>
<td>N calc</td>
</tr>
<tr>
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<td></td>
<td></td>
<td></td>
<td>MAD%</td>
</tr>
</tbody>
</table>
According to Table 6, the critical values of Reamer et al. and Kohn and Kurata are well represented by the model; the maximum MAD considering the two sources of experimental values is about 6%.

The model agrees well also with the values of S$_2$L$_1$E of Cheung and Zander, and the L$_2$L$_1$E values of Coquelet et al. In this last case, a MAD of about 17% is found at 186 K and 3.7 MPa for a composition of methane in the liquid phase lower than 0.1.

Considering all the VL$_2$E experimental values, the methane composition in the vapor phase is represented within an overall AAD of about 5%, the Bias% is between about -3% and 4%, while the highest values of MAD of about 30% are found with respect to the data of Reamer et al. and Kohn and Kurata for two compositions lower than 0.1. Taking into account the deviations in Table 6 related to the composition of the liquid phase mole fraction, it is possible to state that the AAD is always lower than about 21%, and that the MAD is higher than in the vapor phase. Higher percentage deviations are encountered in the liquid phase because of the low CH$_4$ mole fraction. For instance, the MAD of about 34% (Reamer et al.), 87% (Kohn and Kurata), and 260% (Yarim-Agaev) are related to experimental liquid compositions of methane of 0.0057, 0.0125, and 0.001, respectively.

With respect to phase equilibria involving three phases, a good agreement is achieved along the S$_2$L$_1$E and L$_2$L$_1$E boundaries. The extent to which the pressures of S$_2$L$_2$L$_1$E and S$_2$L$_2$E calculated by the SLV EoS deviate from the experimental pressures is a direct consequence of the
difference between the calculated and the experimental temperature at the quadruple point, as discussed later in this section.

Table 7 presents the quantitative comparison between the SLV EoS and the experimental values (composition of methane, temperature, and pressure) proposed by Kohn and Kurata\textsuperscript{25} concerning the Quadruple Point (S\textsubscript{2}L\textsubscript{2}L\textsubscript{1}VE) and the Upper Critical EndPoint L\textsubscript{2}(L\textsubscript{1}=V).

With respect to the QP, the relative percentage errors between calculated and literature values are about -0.6\%, -32\%, and 2\% for the composition of CH\textsubscript{4} in the liquid L\textsubscript{1}, liquid L\textsubscript{2}, and vapor phase, respectively; the temperature and pressure of the QP are reproduced by the model within 1\% of deviation. It should be noted that in Ref. 25 the solid phase S\textsubscript{2} is treated as pure H\textsubscript{2}S, whereas the calculated mole fraction of methane in the S\textsubscript{2} phase is 0.046.

With respect to the UCEP\textsubscript{1}, the deviations related to the experimental temperature and pressure are about 1.3\% and 2.6\%, respectively. For the composition of methane, the errors are -1.28\% for the L\textsubscript{1}=V phase and 14\% for the L\textsubscript{2} phase.

Improved representations could have been obtained using more complex mixing rules involving binary interaction parameters instead of Eqs. 6-8 and/or introducing temperature dependent binary interaction parameters, but authors decided to use a single binary interaction parameter, $k_{ij}$, constant with temperature, to increase the prediction capability of the model. The model is applied in the next section for representing the whole and complex phase diagram of the mixture in a wide range of temperature and pressure even for regions where data are scarce or not at all available. The chosen parameterization of the SLV EoS is then a compromise between the accuracy in representing the existing experimental values and the robustness of the model with respect to extrapolation for predicting the phase diagrams at temperatures and pressures for which experimental values are not available.
Table 7. Comparison between calculated and the experimental\textsuperscript{25} singular points for the methane-hydrogen sulfide system.

<table>
<thead>
<tr>
<th>QP (S\textsubscript{2}L\textsubscript{1}L\textsubscript{2}VE)</th>
<th>xCH\textsubscript{4} in S\textsubscript{2}</th>
<th>xCH\textsubscript{4} in L\textsubscript{1}</th>
<th>xCH\textsubscript{4} in L\textsubscript{2}</th>
<th>xCH\textsubscript{4} in V</th>
<th>T / K</th>
<th>P / MPa</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ref. 25</td>
<td>0</td>
<td>0.935</td>
<td>0.104</td>
<td>0.967</td>
<td>182.21</td>
<td>3.38</td>
</tr>
<tr>
<td>SLV EoS</td>
<td>0.046</td>
<td>0.929</td>
<td>0.071</td>
<td>0.988</td>
<td>183.92</td>
<td>3.40</td>
</tr>
<tr>
<td>err%</td>
<td>/</td>
<td>-0.64</td>
<td>-31.73</td>
<td>2.17</td>
<td>0.94</td>
<td>0.59</td>
</tr>
<tr>
<td>UCEP\textsubscript{1} L\textsubscript{2}(L\textsubscript{1}=V)</td>
<td>Ref. 25</td>
<td>/</td>
<td>0.935</td>
<td>0.1</td>
<td>0.935</td>
<td>199.76</td>
</tr>
<tr>
<td>SLV EoS</td>
<td>/</td>
<td>0.923</td>
<td>0.114</td>
<td>0.923</td>
<td>202.25</td>
<td>5.44</td>
</tr>
<tr>
<td>err%</td>
<td>/</td>
<td>-1.28%</td>
<td>14.00</td>
<td>-1.28%</td>
<td>1.25</td>
<td>2.64</td>
</tr>
</tbody>
</table>

Figures 3-7 portray some qualitative comparisons between selected experimental values and the model.

**Figure 3. Pressure-composition diagram at 252 K, 277 K, and 311 K.**
Data: ◊: Reamer et al.\textsuperscript{22}; ●: Kohn and Kurata\textsuperscript{25}; ■: Robinson et al.\textsuperscript{24}, Robinson and Bailey\textsuperscript{23}; +: Yarim-Agaev\textsuperscript{29}. SLV EoS: —: VL\textsubscript{2}E, --: L\textsubscript{1}L\textsubscript{2}E.

The pressure-composition cross sections in Figure 3 show the agreement of the SLV EoS with VL\textsubscript{2}E data in the range 252 – 311 K. Data in Figure 3 are rather consistent among them, and the
model slightly deviates from the experimental composition of methane in the L₂ phase when the temperature increases approaching the critical temperature of pure H₂S. In Figure 3, the isotherms at 277 K and 311 K end at a V=L₂ critical point. To the contrary, at 252 K the VL₂E becomes a L₁L₂E at about 13.8 MPa and the isotherm ends at a L₁=L₂ critical point, thus explaining the odd shape of the isotherm at this temperature with respect to the behaviors at the higher temperatures shown in Figure 3. The dashed line in Figure 3 represents the L₁L₂E at 252 K.

Figure 4 presents the qualitative comparison at 186 and 203 K; data are from Coquelet et al. At 203 K the system presents immiscibility between two liquid phases (L₁ and L₂), and the one rich in methane (L₁) becomes a vapor phase in the low pressure region. Being this temperature higher than the calculated temperature of the UCEP₁ (202.25 K from Table 7), the transition from L₁ to the vapor does not involve neither a VL₁E nor a critical point L₁=V.
A VL$_2$E occurs at any temperature lower than the critical temperature of methane (190.56 K) and higher than the calculated temperature of the quadruple point QP$_1$ (183.92 K from Table 7), as for 186 K in Figure 4. Furthermore, a L$_2$L$_1$VE appears because of the immiscibility gap in the liquid phase. The isotherm at 186K has not been presented in Figure 4 in the low pressure region seeing that a solid phase and related solid-fluid equilibria are involved there. Nevertheless, next section presents the complete phase equilibrium behavior.

![Figure 5. Temperature-composition diagram at 1.4, 4.1, 5.5, and 11 MPa.](image)

The temperature-composition cross sections of Figure 5 show the phase equilibrium behavior of the CH$_4$+H$_2$S system between 1.4 MPa and 11 MPa. Only the VL$_2$E has been presented in Figure 5, whereas the low temperature equilibria have been added and discussed in the next section. According to Figure 5, the SLV EoS is in a quite good agreement with data, except for the deviation with respect to the composition of methane in the L$_2$ phase at 11 MPa. As a consequence, the
calculated $L_2=V$ critical point appears to have a higher methane content than what can be inferred from the experimental values.

Figure 6. Vapor-liquid and liquid-liquid critical loci for the CH$_4$+H$_2$S system.

Data: $V=L_1$ and $V=L_2$: ■: Kohn and Kurata$^{25}$; △: Reamer et al.$^{22}$; $L_2(L_1=V)$: ◊: Kohn and Kurata$^{25}$.

SLV EoS: ———: VLE of CH$_4$ and H$_2$S; ——: $V=L_1$, $V=L_2$ and $L_1=L_2$; ×: $L_2(L_1=V)$.

Figure 6 illustrates the comparison between calculated and experimental values of the critical points of the mixture. Considering the left-branch of the critical curve (mixtures at higher methane content), deviations concerning the calculated (cross) and experimental (empty square) UCEP$_1$ have been already presented in Table 7. The calculated $L_1=V$ critical curve exits the critical point of methane (see Table 2) and rises up to the UCEP$_1$.

The calculated $L_2=V$ critical curve originates from the critical point of H$_2$S (see Table 2) and reaches firstly a local maximum and then a local minimum in pressure while extending toward the low temperature region. These values (respectively 13.9 MPa and 13.87 MPa) are located at temperatures higher than about 266 K, temperature where, in the critical curve, the vapor phase
becomes a liquid phase rich in methane (phase L₁). As a consequence, the L₂=L₁ critical curve replaces the L₂=V one for temperature lower than 266 K, and it develops in the high pressure region for decreasing temperatures. This L₂=L₁ critical curve ends in the singular point S₂(L₂=L₁), UCEP₂, as discussed in the next section.

With respect to the data of Reamer et al.,²² the calculated critical curve develops at pressures slightly higher than the experimental values before reaching the inflection point and the L₂=L₁ – L₂=V transition.

![Figure 7. Pressure-temperature phase diagram of the CH₄+H₂S system in proximity of the QP₁.](image)

Figure 7 groups all the three-phase equilibria of the binary mixture occurring at temperatures higher than 160 K and pressures up to 16 MPa. It should be remarked that being the data in Figure 7 not available numerically, they have been obtained from graphs²⁵ by means of a specific software.
Filled square and circle represent the experimental temperature-pressure couples for the quadruple point and the UCEP\textsubscript{1}, respectively; correspondent calculated values are the filled triangle and the empty triangle. The dashed curves in Figure 7 are the saturation curves of CH\textsubscript{4} and H\textsubscript{2}S, and the melting curve of H\textsubscript{2}S as calculated by the SLV EoS.

Four three-phase boundaries originate from the calculated QP; the S\textsubscript{2}L\textsubscript{1}VE curve which extends down to 160 K remaining close to the saturation curve of methane; the L\textsubscript{2}L\textsubscript{1}VE curve ending at the calculated UCEP\textsubscript{1}; the S\textsubscript{2}L\textsubscript{2}L\textsubscript{1}E curve which rapidly extends in the high pressure region; the S\textsubscript{2}L\textsubscript{2}VE curve which joins the triple point of H\textsubscript{2}S (▼). For sake of completeness, the V=L\textsubscript{1} critical curve has been added in Figure 7; it exits the critical point of CH\textsubscript{4} (+) and joins the UCEP\textsubscript{1} (Δ).

The deviation between the calculated S\textsubscript{2}L\textsubscript{2}L\textsubscript{1}E and S\textsubscript{2}L\textsubscript{2}VE boundaries and the experimental trends which can be inferred from the correspondent available data (respectively empty squares and diamonds) can be related to the deviation between the calculated and experimental temperature of the QP (see Table 7). Furthermore, the S\textsubscript{2}L\textsubscript{2}VE data\textsuperscript{25} seem not to extend in the low pressure region towards the triple point temperature of pure H\textsubscript{2}S (187.7 K, 22.3 kPa). In fact, the experimental points tend to a temperature of about 185 K in the zero-pressure limit, temperature that is about 3 degrees lower than the triple point temperature of H\textsubscript{2}S proposed in Ref. 45. This explains the remarkable deviations for the S\textsubscript{2}L\textsubscript{2}L\textsubscript{1}E and S\textsubscript{2}L\textsubscript{2}VE presented in Table 7.

As previously stated, next section aims at presenting an overview of the phase equilibrium behavior of the CH\textsubscript{4}+H\textsubscript{2}S system by means of several isotherms and isobars. With respect to this section, solid phases and correspondent equilibria have been added in cross sections and P-T diagrams in order to present the whole phase equilibrium behavior.
Description of the Phase Diagram of the Methane-Hydrogen Sulfide System

This section has the scope of describing the pressure-temperature phase diagram and all the possible types of pressure-composition and temperature-composition phase diagrams that can be encountered in the investigated range of temperature and pressure. This analysis aims to be helpful to the process engineer who needs to understand how the CH₄+H₂S system behaves at a certain temperature and pressure.

Figure 8 presents the pressure-temperature phase diagram of the CH₄+H₂S mixture for temperatures from 70 K up to the critical temperature of H₂S and pressures up to 250 MPa.

According to the available experimental values and the results of the SLV EoS, in these ranges the mixture presents two Quadruple Points and two Upper Critical EndPoints. The QPs are related to the S₂S₁L₁VE (QP₂) and the S₂L₂L₁VE (QP₁); the UCEPs are related to the singular points L₂(L₁=V), UCEP₁, and the S₂(L₂=L₁), UCEP₂.

The phase equilibrium behavior involves then 7 three-phase equilibrium boundaries (S₂S₁VE, S₂S₁L₁E, S₁L₁VE, S₂L₁VE, S₂L₂VE, S₂L₂L₁E, and L₂L₁VE) and 5 phases (S₂, S₁, L₂, L₁, and V).

Taking into account the huge range of pressure considered in this work, the y-axis in Figure 8 has been cut twice (at 0.5 MPa and 19 MPa) for easing the comprehension of the equilibria occurring in proximity of the QP₂, of the points QP₁-UCEP₁, and the high-pressure equilibria located near the UCEP₂. Furthermore, a zoom of Figure 8 has been presented in Figure 9 in order to show the phase equilibrium behavior in the low-pressure region, namely in proximity of the QP₂ and the triple point of CH₄.

In the low pressure region of Figure 8, 0-0.5 MPa, the QP₂ almost overlaps the triple point of pure methane, than the saturation, melting, and sublimation curves of pure CH₄ are not clearly visible being overlapped by the S₂S₁VE, S₂S₁L₁E, and S₂L₁VE curves. The S₂L₂VE curve originates at the triple point of H₂S, and extends at higher pressures with a negative slope.

The more complex part of the phase equilibrium behavior concentrates between 0.5 MPa and 19 MPa. The S₂S₁L₁E remains close to the SLE of CH₄, while the S₂L₁VE boundary distances itself
from the saturation curve of CH₄ while approaching the filled triangle, which represents the QP₁. The S₂L₂VE, curve originating at the triple point of H₂S, also ends at the QP₁. The other three-phase equilibria joining the QP₁ are the L₂L₁VE and the S₂L₂L₁E. The former reaches the UCEP₁, namely the empty triangle in Figure 8, the latter extends rapidly at high pressures with a positive slope. In addition to that, two critical boundaries occur: the V=L₁ loci leaving the critical point of CH₄ to end in the UCEP₁, and the V=L₂ loci originating from the critical point of H₂S and becoming the L₂=L₁ loci at low temperatures.

In the high pressure region of Figure 8, the S₂S₁L₁E starts to deviate from the SLE of CH₄, and the L₂=L₁ loci meets the S₂L₂L₁E curve at the empty square, namely the UCEP₂ (208.98 K and 224.09 MPa). As a consequence, the L₂ and L₁ phases become a unique liquid phase for any pressure higher than the pressure at the UCEP₂, S₂(L₂=L₁).
Figure 8. Calculated pressure-temperature phase diagram of the CH$_4$+H$_2$S system.

---: three-phase equilibrium boundaries; ···: critical curves; ---: SVE, VLE and SLE of CH$_4$ and H$_2$S;

■: QP$_2$; ▲: QP$_1$; △: UCEP$_1$; □: UCEP$_2$. 
The three-phase equilibrium boundaries related to the QP$_2$, calculated at 90.75 K and 11.77 kPa, can be appreciated in Figure 9. From the QP$_2$, the S$_2$S$_1$VE curve extends in the low temperature region, the S$_2$S$_1$L$_1$E extends in the high pressure region, the S$_2$L$_1$VE curve develops in such a way to end in the QP$_1$, the S$_1$L$_1$VE joins the triple point of CH$_4$. Dashed curves represent the sublimation, saturation, and melting curves of pure CH$_4$.

It should be stated that any experimental evidence has been provided in the literature concerning the precise position of the QP$_2$, thus the calculated QP$_2$ is a simple result of the SLV EoS with a binary interaction parameter regressed with respect to the available data at high temperatures. The calculated QP$_2$ is at a temperature slightly higher than the triple point temperature of methane, and this feature results in a solid-fluid equilibrium behavior of the peritectic type, as it has been discussed in the following.
Figures 10-15 present a series of cross sections at constant temperature and pressure that have been selected in the PT phase diagram (Figure 8) in order to describe all the phase equilibrium behaviors of the mixture in the wide ranges of temperature and pressure considered in this work.

As it has already been stated, the objective is to provide the reader with a mapping of the phase diagram and the possibility of rapidly obtaining an overview of the phase equilibrium behavior according to the system temperature and pressure.

A set of temperature and pressure couples has been selected in order to follow the qualitative changes of the phase behavior, as summarized in Tables 8 and 9.

Table 8 presents all the main regions of temperature (listed from 1T to 13T) where the correspondent pressure-composition cross section does not change from a qualitative point of view. A lower and an upper temperature limit have been fixed for each region of temperature, and a representative temperature ($T_{ref}$) has been chosen for each interval in order to describe its qualitative phase diagram. The last column of Table 8 indicates the figure presenting the cross section correspondent to the chosen representative temperature. In some cases, either the lower or the upper limit or both temperature limits are represented by the temperature where two phase boundaries meet or cross each other. For instance for the region 4T, the lower limit is the temperature at the quadruple point $Q_{P1}$, whereas the upper is the temperature where the saturation curve of CH$_4$ (VLE CH$_4$) crosses the $S_2L_2L_1E$ curve.
Table 8. Temperature ranges presenting different qualitative pressure-composition phase equilibrium behaviors for the methane-hydrogen sulfide system, and the selected representative temperature ($T_{ref}$).

<table>
<thead>
<tr>
<th>Region</th>
<th>Lower limit (T / K)</th>
<th>Upper limit (T / K)</th>
<th>$T_{ref}$ / K</th>
<th>Figure</th>
</tr>
</thead>
<tbody>
<tr>
<td>1T</td>
<td>Low-temperature limit (70)</td>
<td>T, CH$_4$ (90.694)</td>
<td>85</td>
<td>10A</td>
</tr>
<tr>
<td>2T</td>
<td>T, CH$_4$ (90.694)</td>
<td>T, QP$_2$ (90.75)</td>
<td>90.7</td>
<td>10B</td>
</tr>
<tr>
<td>3T</td>
<td>T, QP$_2$ (90.75)</td>
<td>T, QP$_1$ (183.92)</td>
<td>140</td>
<td>10C</td>
</tr>
<tr>
<td>4T</td>
<td>T, QP$_1$ (183.92)</td>
<td>T, cross VLE CH$_4$/S$_2$L$_2$L$_1$E (184.13)</td>
<td>183.94</td>
<td>10D</td>
</tr>
<tr>
<td>5T</td>
<td>T, cross VLE CH$_4$/S$_2$L$_2$L$_1$E (184.13)</td>
<td>T, H$_2$S (187.7)</td>
<td>186.25</td>
<td>10E</td>
</tr>
<tr>
<td>6T</td>
<td>T, H$_2$S (187.7)</td>
<td>T, cross VLE H$_2$S/L$_2$L$_1$VE (188.47)</td>
<td>188</td>
<td>10F</td>
</tr>
<tr>
<td>7T</td>
<td>T, cross VLE H$_2$S/L$_2$L$_1$VE (188.47)</td>
<td>T, cross SLE H$_2$S/L$_2$L$_1$VE (188.56)</td>
<td>188.48</td>
<td>11A</td>
</tr>
<tr>
<td>8T</td>
<td>T, cross VLE CH$_4$/SLE H$_2$S (188.56)</td>
<td>T, CH$_4$ (190.56)</td>
<td>189.5</td>
<td>11B</td>
</tr>
<tr>
<td>9T</td>
<td>T, CH$_4$ (190.56)</td>
<td>T, UCEP$_1$ (202.25)</td>
<td>195</td>
<td>11C</td>
</tr>
<tr>
<td>10T</td>
<td>T, UCEP$_1$ (202.25)</td>
<td>T, UCEP$_2$ (208.98)</td>
<td>203.4</td>
<td>11D</td>
</tr>
<tr>
<td>11T</td>
<td>T, UCEP$_2$ (208.98)</td>
<td>T, cross SLE H$_2$S/L$_2$L$_1$E (210.71)</td>
<td>210</td>
<td>11E</td>
</tr>
<tr>
<td>12T</td>
<td>T, cross SLE H$_2$S/L$_2$L$_1$E (210.71)</td>
<td>T, H$_2$S (373.1)</td>
<td>230</td>
<td>11F</td>
</tr>
<tr>
<td>13T</td>
<td>T, cross SLE H$_2$S/L$_2$L$_1$E (210.71)</td>
<td>T, H$_2$S (373.1)</td>
<td>300</td>
<td>12</td>
</tr>
</tbody>
</table>

A S$_2$VE, a S$_1$VE, and a S$_2$S$_1$E occur at any temperature higher than 70 K and lower than the triple point temperature of methane, as shown in the Pressure-composition (P$_x$) cross section at 85 K, Figure 10A. At this temperature, the binary mixture presents a region of S$_2$VE for all pressures lower than 4.911 kPa down to the sublimation pressure of pure H$_2$S. An equilibrium between the phases S$_2$, S$_1$, and V occurs at 4.911 kPa. Then, the S$_2$S$_1$E extends up to 250 MPa, while the S$_1$VE ends at the sublimation pressure of pure CH$_4$ (4.913 kPa).

With respect to the range 1T, any P$_x$ cross section in the range 2T presents one additional pressure of three-phase equilibrium. This is related to the S$_1$L$_1$VE.

Figure 10B shows the phase equilibrium behavior at 90.7 K: the S$_2$S$_1$V$_1$E and S$_1$L$_1$VE are at 11.694 kPa and 11.705 kPa, respectively. The S$_2$VE extends again from the sublimation pressure of H$_2$S (1.3×10$^{-6}$ kPa) up to the pressure of S$_2$S$_1$VE, and the S$_2$S$_1$E extends up to 250 MPa and beyond. In this case, the S$_1$VE ends at the S$_1$L$_1$VE rather than at the sublimation of CH$_4$; the V$_1$E and the S$_1$L$_1$E end respectively at the saturation (11.708 kPa) and melting (40.13 kPa) pressures of pure CH$_4$.

Any P$_x$ cross section in the range 3T is qualitatively represented in Figure 10C, which has been calculated at 140 K. The S$_2$VE originating at the sublimation pressure of H$_2$S ends now at the pressure of S$_2$L$_1$VE (0.635 MPa): from this pressure, the V$_1$E joins the saturation pressure of CH$_4$. 

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(0.647 MPa), while the $S_2L_1E$ extends in the high pressure region up to the $S_2S_1L_1E$ (198 MPa). From this three-phase equilibrium, the $S_1L_1E$ ends at the melting pressure of CH$_4$ (230 MPa), the $S_2S_1E$ extends in the high pressure region.

When the system temperature exceeds 183.92 K, namely the temperature of QP$_1$, a liquid phase L$_2$ appears in order to provide the $S_2L_2VE$, the $L_2L_1VE$, the $S_2L_2L_1E$, and the related two-phase equilibria. The $P_x$ cross section at 183.94 K in Figure 10D presents a $S_2VE$ ending at the pressure of $S_2L_2VE$ (3.374 MPa). The system is at $S_2L_2E$ between this pressure and the pressure of $S_2L_2L_1E$ (3.645 MPa) only for compositions of CH$_4$ between about 0.04 and 0.07. To the contrary, the VL$_2E$ and L$_2L_1E$ occur for higher contents of CH$_4$, and the transition between these equilibria involves the L$_2L_1VE$ (3.402 MPa). The $S_2L_1E$ originates from the $S_2L_2L_1E$ and extends up to 250 MPa and beyond, while a VL$_1E$ runs from the pressure of L$_2L_1VE$ and the saturation pressure of CH$_4$ (3.768 MPa).

In the region 4T the saturation pressure of CH$_4$ is higher than the pressure of $S_2L_2L_1E$, while the contrary occurs in the region 5T, where the phase L$_2$ develops and reaches higher pressures thus making the $S_2L_2L_1E$ occurring at higher pressures. For instance at 186.25 K (Figure 10E), the pressure of $S_2L_2L_1E$ is 23.449 MPa, higher than the saturation pressure of CH$_4$ (4.045 MPa).

In general, all the solid-liquid-vapor equilibria occur once the temperature is higher than the triple point temperature of CH$_4$; the $S_2L_2E$ develops between the pressures of $S_2L_2VE$ and $S_2L_2L_1E$ in regions 4T and 5T, whereas it starts at the melting pressure of H$_2$S in all the PT cross sections from the region 6T to higher temperatures. For instance at 188 K, Figure 10F, the $S_2L_2E$ exits the melting pressure of H$_2$S (1.632 MPa) and ends at the $S_2L_2L_1E$ (37.439 MPa); furthermore, the VL$_2E$ extends from the L$_2L_1VE$ (3.808 MPa) down to the saturation pressure of H$_2$S (0.0228 MPa).

The underlying difference between the phase equilibrium behaviors in the regions 6T-8T is the reciprocal positions of the melting pressure of H$_2$S, the pressures of L$_2L_1VE$, and the saturation pressure of CH$_4$. 

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In the region 6T, the melting pressure of H$_2$S is lower than the pressure of L$_2$L$_1$VE and the saturation pressure of CH$_4$. In the region 7T, the melting pressure of H$_2$S exceeds the pressure of L$_2$L$_1$VE remaining lower than the saturation pressure of CH$_4$. For instance, in the Px cross section at 188.48 K, Figure 11A, these values are 4.222 MPa, 3.859 MPa, and 4.326 MPa, respectively.
Figure 10. Calculated pressure-composition (Px) cross sections for the regions 1T-6T.
The melting pressure of H\(_2\)S exceeds also the saturation pressure of pure CH\(_4\) in the region 8T; for instance, in the Px cross section at 189.5 K (Figure 11B) these values are 9.786 MPa (SLE H\(_2\)S), 4.458 MPa (VLE CH\(_4\)), and 4.326 MPa (L\(_2\)L\(_1\)VE), respectively.

The upper temperature related to the region 8T is the critical temperature of CH\(_4\). As a consequence, the VL\(_1\)E ends at the saturation pressure of pure CH\(_4\) in all the Px cross sections in the range of temperatures corresponding to the regions from 2T to 8T. The VL\(_1\)E ends instead at a critical point L\(_1\)=V in region 9T, as it can be appreciated in Figure 11C which shows the Px cross section at 195 K. At this temperature, the calculated L\(_2\)L\(_1\)VE and the S\(_2\)L\(_2\)L\(_1\)E are at 4.57 MPa and 94.718 MPa, respectively.

According to the PT diagram of Figure 9, the L\(_2\)L\(_1\)VE boundary ends at the singular point UCEP\(_1\). Consequently, the transition between the V and L\(_1\) phases becomes continuous for temperatures higher than 202.25 K. For instance, the Px cross section at 203.4 K portrayed in Figure 11D illustrates the absence of the VL\(_1\)E and of the critical point L\(_1\)=V. The difference between the VL\(_2\)E and the L\(_2\)L\(_1\)E is the consequence of a change in density of the phase at equilibrium with the phase L\(_2\). Furthermore, the S\(_2\)L\(_2\)L\(_1\)E occurs at 169.717 MPa.

No critical point exists for the mixture in the region 10T; to the contrary, the L\(_2\)=L\(_1\)/V=L\(_2\) critical curve is crossed at each temperature higher than the temperature at the UCEP\(_2\) (208.98 K).

At 210 K, Figure 11E, the VL\(_2\)E originates at the saturation pressure of H\(_2\)S, and the L\(_2\)L\(_1\)E ends at a critical point L\(_2\)=L\(_1\) (176.4 MPa). The S\(_2\)L\(_2\)E exits the melting pressure of H\(_2\)S (139.74 MPa) and extends in the high pressure region. It should be noted that the melting pressure of H\(_2\)S is lower than the pressure of the critical point L\(_2\)=L\(_1\) at 210 K, and so for all the temperatures in the region 11T.

In the region 12T the critical point of the system is placed at a pressure lower than the melting pressure of pure H\(_2\)S; for instance at 230 K, Figure 11F, the L\(_2\)L\(_1\)E ends at about 23 MPa, which represents a critical point L\(_2\)=L\(_1\); the solid-liquid equilibria are located at pressures higher 250 MPa.
Finally, the extent of the \( L_2L_1E \) reduces for temperatures approaching the critical temperature of \( \text{H}_2\text{S} \), and the critical point \( L_2=L_1 \) becomes a critical point \( L_2=V \), as shown in Figure 12 by the Px cross section at 300 K.
Figure 11. Calculated pressure-composition (P-x) cross sections for the regions 7T-12T.
Figure 12. Calculated pressure-composition (Px) cross section at 300 K, i.e. the reference temperature for the region 13T.
A similar study has been carried out with respect to the qualitative changes of the temperature-composition cross sections with pressure. Table 9 summarizes the main pressure regions listed from 1P to 13P. Lower and upper pressure limits and a reference pressure have been fixed for each region of pressure, whereas the last column of Table 9 indicates the number of the figure presenting the correspondent cross section.

In some case, either the lower or the upper limit or both pressure limits are represented by the pressure where two phase boundaries cut cross each other. For instance, the lower limit for the region 5P is the pressure at the quadruple point \( QP_1 \), whereas its upper limit is the pressure where the melting curve of \( H_2S \) (SLE \( H_2S \)) crosses the \( L_2L_1VE \) curve.

Table 9. Pressure ranges presenting different qualitative temperature-composition phase equilibrium behaviors for the methane-hydrogen sulfide system, and the selected representative pressure (\( Pref \)).

<table>
<thead>
<tr>
<th>Region</th>
<th>Lower limit (( P / MPa ))</th>
<th>Upper limit (( P / MPa ))</th>
<th>( Pref / MPa )</th>
<th>Figure</th>
</tr>
</thead>
<tbody>
<tr>
<td>1P</td>
<td>Low-pressure limit (0)</td>
<td>( P_t ) ( CH_4 ) (0.011697)</td>
<td>0.009</td>
<td>13A</td>
</tr>
<tr>
<td>2P</td>
<td>( P_t ) ( CH_4 ) (0.011697)</td>
<td>( P_t ) ( QP_1 ) (0.01177)</td>
<td>0.0117335</td>
<td>13B</td>
</tr>
<tr>
<td>3P</td>
<td>( P_t ) ( QP_1 ) (0.01177)</td>
<td>( P_t ) ( H_2S ) (0.0223)</td>
<td>0.02</td>
<td>13C</td>
</tr>
<tr>
<td>4P</td>
<td>( P_t ) ( H_2S ) (0.0223)</td>
<td>( P_t ) ( QP_1 ) (3.4)</td>
<td>1.37</td>
<td>13D</td>
</tr>
<tr>
<td>5P</td>
<td>( P_t ) ( QP_1 ) (3.4)</td>
<td>( P_t ) ( QP_1 ) (3.4)</td>
<td>1.37</td>
<td>13D</td>
</tr>
<tr>
<td>6P</td>
<td>( P_t ) ( QP_1 ) (3.4)</td>
<td>( P_t ) ( QP_1 ) (3.4)</td>
<td>1.37</td>
<td>13D</td>
</tr>
<tr>
<td>7P</td>
<td>( P_t ) ( QP_1 ) (3.4)</td>
<td>( P_t ) ( QP_1 ) (3.4)</td>
<td>1.37</td>
<td>13D</td>
</tr>
<tr>
<td>8P</td>
<td>( P_t ) ( QP_1 ) (3.4)</td>
<td>( P_t ) ( QP_1 ) (3.4)</td>
<td>1.37</td>
<td>13D</td>
</tr>
<tr>
<td>9P</td>
<td>( P_t ) ( QP_1 ) (3.4)</td>
<td>( P_t ) ( QP_1 ) (3.4)</td>
<td>1.37</td>
<td>13D</td>
</tr>
<tr>
<td>10P</td>
<td>( P_t ) ( QP_1 ) (3.4)</td>
<td>( P_t ) ( QP_1 ) (3.4)</td>
<td>1.37</td>
<td>13D</td>
</tr>
<tr>
<td>11P</td>
<td>( P_t ) ( QP_1 ) (3.4)</td>
<td>( P_t ) ( QP_1 ) (3.4)</td>
<td>1.37</td>
<td>13D</td>
</tr>
<tr>
<td>12P</td>
<td>( P_t ) ( QP_1 ) (3.4)</td>
<td>( P_t ) ( QP_1 ) (3.4)</td>
<td>1.37</td>
<td>13D</td>
</tr>
<tr>
<td>13P</td>
<td>( P_t ) ( QP_1 ) (3.4)</td>
<td>( P_t ) ( QP_1 ) (3.4)</td>
<td>1.37</td>
<td>13D</td>
</tr>
</tbody>
</table>

At 9 kPa (Figure 13A) the system \( CH_4+H_2S \) presents a \( S_2S_1VE \) at 88.906 K; a \( S_2S_1E \) extends down to 70 K, whereas the \( S_2VE \) and the \( S_1VE \) ends at the sublimation temperature of pure \( H_2S \) (175.957 K) and \( CH_4 \) (88.902 K), respectively. The Temperature-composition (\( Tx \)) cross section of Figure 13A is representative of the region 1P, whose upper pressure limit is the triple point of \( CH_4 \).

At any pressure in the region 2P, namely between the triple point pressure of \( CH_4 \) (11.697 kPa) up to the pressure of the \( QP_2 \) (11.77 kPa), the \( S_1VE \) exists between a \( S_1L_1VE \) and a \( S_2S_1VE \). For instance at 11.7335 kPa, Figure 13B, the \( S_1VE \) is confined in the high-methane content region.
between 90.72 K (temperature of S₁L₁VE) and 90.722 K (temperature of S₂S₁VE). From the S₁L₁VE, a S₁L₁E and a VL₁E originate and join the melting temperature (90.695 K) and the saturation temperature (90.718 K) of pure CH₄, respectively. The S₂VE originating at the S₂S₁VE always ends at the sublimation temperature of H₂S (179.22 K in the case of Figure 13B).

A S₂S₁L₁E and a S₂L₁VE appear when the system pressure exceeds the pressure at QP₂ remaining lower than the triple point pressure of H₂S, as shown in the Tx cross section at 0.02 MPa portrayed in Figure 13C. Figure 13C is representative of the phase equilibrium behavior in the region 3P, and the temperatures of S₂S₁L₁E and a S₂L₁VE at 0.02 MPa are 90.748 K and 95.14 K, respectively.

The S₁VE does not exist for pressures higher than the pressure of the QP₂, and for increasing pressures the S₂S₁L₁E remains located almost at the same temperature while the S₂L₁VE moves to higher values. Furthermore, a S₂L₁E appears for the first time in the region 3P and it will never disappear in the Tx cross sections at higher pressures. With respect to the behavior in the region 2P (Figure 13B) where the S₁L₁E and a VL₁E both join the temperature of S₁L₁VE, in the region 3P the S₁L₁E and a VL₁E reach two different three-phase equilibria. According to Figure 13C, the S₁L₁E ranges from the melting temperature of CH₄ (90.694 K) and the temperature of S₂S₁L₁E, while the VL₁E extends from the saturation temperature of CH₄ (95.136 K) and the temperature of S₂L₁VE.

A third three-phase equilibrium, the S₂L₂VE, is encountered for pressures in the region 4P, namely from the triple point pressure of H₂S (22.3 kPa) up to the pressure of QP₁ (3.4 MPa). This additional equilibrium and its related two-phase equilibria (S₂L₂E and VL₂E) are the consequence of the appearance of the liquid phase L₂ in the Tx cross section, as shown at 1.37 MPa in Figure 13D. Thus, in the region 4P, the phase equilibrium behavior changes qualitatively in the high temperature region. With respect to Figure 13C, in Figure 13D the S₂VE reaches the S₂L₂VE (157.132 K), from which the S₂L₂E and VL₂E continue up to the melting temperature (186.112 K) and saturation temperature (282.753 K) of H₂S, respectively.
The $S_2L_1$VE and $S_2L_2$VE cease to occur when the system pressure becomes higher than the pressure of $QP_1$, while the $S_2S_1L_1E$ persists at low temperatures. To the contrary, two different three-phase equilibria turn out, at least for pressures lower than the pressure of $UCEP_1$ (5.44 MPa).

For instance at 3.75 MPa, whose Tx cross section is presented in Figure 13E, the system is characterized by a $S_2S_1L_1E$ (91.778 K), a $S_2L_2L_1E$ (183.95 K), and a $L_2L_1$VE (187.428 K), and related two-phase equilibria. Consequently, it can be stated that the immiscibility between the $L_1$ and $L_2$ phases occurs only for pressures higher than the pressure of $QP_1$, according obviously to the value of the system temperature. The Tx cross sections in Figures 13F, 14A, and 14B are qualitatively of the same kind of Figure 13E. The difference among the correspondent phase equilibrium behaviors (respectively at 3.75 MPa, 3.825 MPa, 4.14 MPa, and 4.5 MPa) is the position of the saturation and/or melting temperatures of the pure components.

At 3.75 MPa, Figure 13E, the melting temperature of $H_2S$ (188.393 K) is higher than the temperature of $L_2L_1$VE (187.428 K), which in turns is higher than the temperatures of $S_2L_2L_1E$ (183.95 K) and saturation of $CH_4$ (183.787 K).
Figure 13. Calculated temperature-composition (T-x) cross sections for the regions 1P-6P.
At 3.825 MPa, Figure 13F, the melting temperature of H$_2$S (188.407 K) is still higher than the temperature of L$_2$L$_1$VE (188.156 K), and the saturation temperature of CH$_4$ (184.425 K) is in this case lower than the temperature of L$_2$L$_1$VE but higher than the temperature of S$_2$L$_2$L$_1$E (183.958 K).

At 4.14 MPa, Figure 14A, the temperature of L$_2$L$_1$VE (191.09 K) exceeds the melting temperature of H$_2$S (188.464 K); the saturation temperature of CH$_4$ (186.99 K) and the temperature of S$_2$L$_2$L$_1$E (183.98 K) are again lower than the melting temperature of H$_2$S.

At 4.5 MPa, Figure 14B, the temperatures of L$_2$L$_1$VE (194.379 K) and saturation of CH$_4$ (189.815 K) are both higher than the temperatures of melting of H$_2$S (188.531 K) and of S$_2$L$_2$L$_1$E (184.022 K).

The region 9P involves the appearance of the critical point L$_1$=V. The Tx cross section at 5 MPa in Figure 14C is representative of the phase equilibrium behavior in this region of pressure. In this case, the VL$_1$E joining the L$_2$L$_1$VE at 198.672 K does not end at the saturation temperature of CH$_4$ seeing that the system pressure is now higher than the critical point pressure of same component (4.5992 MPa). Therefore the VL$_1$E ends at a critical point L$_1$=V at about 197.093 K.

It should be noted that in Figures 13D-13F and 14A-14C the S$_1$L$_1$E is confined in the region of high-methane content, and that the temperature of S$_2$L$_2$L$_1$E changes from 91.125 K at 1.37 MPa (Figure 13D) to 92.12 K at 5 MPa (Figure 14C).

In the Tx cross section, a L$_1$=V critical point occurs at each pressure between the critical point pressure of CH$_4$ and the pressure of UCEP$_1$ (5.44 MPa). It means that for higher pressures, as in the region 10P represented by the diagram at 7.5 MPa in Figure 14D, the transition between the L$_1$ and the vapor is continuous, thus the L$_2$L$_1$VE ceases to exist.

Nevertheless, a critical point exists when the pressure is higher than the critical pressure of H$_2$S (9 MPa) and lower than the pressure of UCEP$_2$ (224.09 MPa). This critical point represents either a L$_2$=V or a L$_2$=L$_1$ equality according to pressure. For instance, at 15 MPa, Figure 14E, the VL$_2$E
ends at a critical point \( L_2=V \) (363.3 K), while at 200 MPa, Figure 14F, the VL\(_2\)E has disappeared and the L\(_2\)L\(_1\)E ends directly at a critical point \( L_2=L_1 \) (209.53 K).

In Figures 14C-14F, the solid-liquid equilibria \( S_2L_2E \) and \( S_1L_1E \) join the \( S_2L_2L_1E \) and the \( S_2S_1L_1E \), respectively. The \( S_2L_2E \) ends at the melting temperature of \( H_2S \), the \( S_1L_1E \) ends at the melting temperature of \( CH_4 \).

Because the immiscibility in the liquid phase ceases at pressures higher than the pressure of UCEP\(_2\) (224.09 MPa), it follows that the \( S_2L_2L_1E \) does not persist in the region 13P, as it happens at 235 MPa, Figure 15. This feature leads to have a homogeneous liquid phase and a \( S_2LE \) extending in the low temperature region down to the \( S_2S_1L_1E \).
Figure 14. Calculated temperature-composition (Tx) cross sections for the regions 7P-12P.
Figure 15. Calculated temperature-composition (Tx) cross sections at 235 MPa, i.e. the reference pressure for the region 13P.

Miscibility of the Solid Phases and Allotropic Behavior of H$_2$S

The first comment concerning the Px and Tx cross sections calculated by the SLV EoS concerns the purity of the solid phases S$_1$ and S$_2$. As stated in Section 3, using the mixing rules developed for fluid phases presented in Eqs. 5-8 results in obtaining not pure solid phases, as it can be observed for instance in Figure 15. The Tx cross section at 235 MPa presents a solid phase S$_2$ rich in H$_2$S with a maximum mole fraction of methane of 0.18, and a solid phase S$_1$ rich in CH$_4$ where the mole fraction of hydrogen sulfide reaches a maximum of 0.13. As already stated, any evidence has been
provided yet in the literature concerning the purity of the solid phases in the methane + hydrogen sulfide system, and the results here presented resulted from the mere application of the cited mixing rules in the SLV EoS.

Moreover, as previously introduced in this section, also the position of the $S_2 S_1 L_1$ VE ($QP_2$) has not been determined yet. The kind of the solid-liquid equilibrium of the system of interest strictly depends on the exact position of the quadruple point, $QP_2$. The binary mixture of $CH_4$ and $H_2S$ presents immiscibility in the solid phase, but this feature could result in a solid-liquid equilibrium either of the eutectic type or the peritectic type. The discerning aspect is comparison between the mole fraction for instance of $CH_4$ in the liquid phase $L_1 (x_{L_1})$ and those in the solid phases $S_1$ and $S_2 (x_{S_1}$ and $x_{S_2})$: $x_{L_1}$ is lower than $x_{S_1}$ but higher than $x_{S_2}$ in the eutectic behavior, it is higher than both $x_{S_1}$ and $x_{S_2}$ in the peritectic behavior.

According to the results presented in Figures 10C, 14C-14F, and 15, it clearly appears that the mole fraction of $CH_4$ in the liquid phase $L_1$ is higher than the mole fractions in the solid phases $S_1$ and $S_2$. This is a direct consequence of the position of the singular point $QP_2$ in the PT diagram. No binary interaction parameter has been regressed in order to match an eutectic behavior seeing that there is no literature evidence of this feature for the $CH_4+H_2S$ system.

The last analysis concerning the results of the SLV EoS is related to the three different crystal structures of $H_2S$ that should be considered for obtaining the real behavior of the system. As introduced in Section 2, $H_2S$ presents a triple point of solid$_\gamma$-solid$_\beta$-vapor equilibrium at 103.5 K and triple point of solid$_\beta$-solid$_\alpha$-vapor equilibrium at 126 K in addition to the common triple point of solid$_\alpha$-liquid-vapor at 187.7 K.\textsuperscript{42} The triple points at low temperature are stated to occur at the zero-pressure limit, while the solid-liquid-vapor equilibrium is at 22.3 kPa.\textsuperscript{45}

Being the SLV EoS in Eq. 1 a fourth-degree polynomial in the variable volume, a single root can be associated to the solid volume, being the other roots related to the fluid phases. In dealing with mixtures, this solid volume correspond to a solid phase rich in either $H_2S$ or $CH_4$, which explains
the possibility of representing the phases $S_2$ and $S_1$. To the contrary, the possibility of representing several solid phases for each pure component is outside the range of applicability of this EoS.

Nevertheless, the way the pure $H_2S$ solid phases $\gamma$ and $\beta$ affect the phase equilibrium behavior of the mixture can be added relatively easily in both $T_x$ and $P_x$ cross sections. For instance, Figures 16A and 16B present the $T_x$ cross sections at 4.14 MPa in the range 91.8 K – 196 K considering a single phase for the solid $H_2S$, $S_2$, and two additional solid-solid transitions occurring in pure $H_2S$ at 103.5 K and 126 K, respectively.

One can notice at once that the phase $S_\alpha$ in Figure 16B takes the role of the phase $S_2$ in Figure 16A. In addition to that, the $S_2S_1L_1E$ is replaced by the $S_\gamma S_1L_1E$ at about 91.9 K, while two new three-phase equilibria appear. A $S_\gamma S_\beta L_1E$ originating at 103.5 K ends at the temperature of the former, the $S_\gamma S_\beta L_1E$ at about 92 K, and a $S_\beta S_\alpha E$ originating at 126 K ends at the temperature of the latter, namely the $S_\beta S_\alpha L_1E$ placed at about 108 K. The temperatures of these SSL$_1E$ have been randomly fixed and supposed lower than the correspondent solid-solid transitions of pure $H_2S$.

As it is possible to observe in Figure 16, the inclusion of different crystal structures of $H_2S$ does not modify the solid-fluid equilibria, although the different solid phases imply the presence of supplementary solid-solid and solid-solid-fluid equilibria.

Figure 16. Comparison between the $T_x$ cross sections at 4.14 MPa without (A) and with (B) the solid$_\alpha$-solid$_\beta$ and solid$_\beta$-solid$_\gamma$ transitions of $H_2S$. 

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Conclusions

The increasing global energy demand and the availability of highly sour natural gas reserves establish the need of a detailed description of the phase behavior of systems containing methane and hydrogen sulfide in order to properly tune thermodynamic models and, most of all, to perform the correct process design of new gas purification technologies (particularly the low temperature ones) that are required to allow the commercialization of such a kind of low quality natural gas reserves.

Different authors studied the phase behavior of the system methane - hydrogen sulfide considering only fluid phases, whereas few works considered the presence of a pure solid phase of $\text{H}_2\text{S}$. The experimental works proposed in the last 60 years investigated the phase behavior of the system in a large proportion of the Pressure-Temperature (PT) diagram down to about 180 K. In particular, the work by Kohn and Kurata presented qualitative representations of the PT diagram and highlighted the complexity of the phase equilibrium behavior.

Nevertheless, no work has been proposed to reproduce the PT diagram of the mixture down to the triple point of methane by means of a unified thermodynamic model for both fluid and solid phases. In addition to that, literature data never exceed 14 MPa.

In this work, the complete PT diagram of the methane - hydrogen sulfide system has been investigated down 70 K and for pressures up to 250 MPa thanks to a model based on the analytic Solid-Liquid-Vapor Equation of State (SLV EoS) proposed in 2003 by Yokozeki. The parameters within the model have been tuned and validated against the available experimental data.

The calculated PT diagram and isobaric/isothermal cross sections show the complexity of the phase equilibrium behavior; immiscibility in both the liquid and solid phases characterizes the system and results in the presence of two quadruple points and two upper critical endpoints.

In addition to that, this work aims at proposing a map of the phase equilibrium behavior thus providing a rapid access to the evaluation of the change of the equilibria as function of pressure and temperature. 13 regions of both pressure and temperature have been found to resume all the possible
schemes of the equilibrium behavior that can be encountered in dealing with the process simulation and/or plant design.

One of the strength of the model here applied is the possibility of representing the rather complex phase equilibrium behavior of the system by means of a single binary interaction parameter. This point better highlights also the simplicity of the approach, enforcing the validity and the effective industrial interest of a simple thermodynamic tool, reliable in a wide region of possible operating conditions.

According to model results for the phase behavior of the system CH$_4$ – H$_2$S, it is possible to identify areas where further experimental investigations are needed. A region of interest extends from temperatures below the UCEP1 and pressures up to 15 MPa. These experimental studies will be useful to better investigate the L$_2$VE, L$_1$VE, L$_1$L$_2$E, S$_2$L$_2$E, S$_2$L$_1$E and S$_2$VE regions where complete datasets (T, P and fluid phase compositions) are missing and phase behavior is complex. Particularly, it can be of interest to determine experimentally the existence of the narrow L$_2$ area present at temperatures between the QP$_1$ and the triple point of pure H$_2$S. TP$_{x_2x_1y}$ data along the L$_1$L$_2$VE locus, TP$_{xy}$ data along the S$_2$L$_2$VE and S$_2$L$_1$VE loci and TP$_{x_2x_1}$ data along the S$_2$L$_2$L$_1$E locus are also of paramount importance. These experimental data will provide useful information also for the validation and proper design of low-temperature process technologies for the purification of highly acid and/or sour natural gases and biogas.

**Nomenclature**

*List of symbols*

- $a$  Equation of state parameter
- $a_0$  Parameter in Eq. 2
- $a_1$  Parameter in Eq. 2
- $a_2$  Parameter in Eq. 2
- $b$  Solid covolume
$b_0$ Parameter in Eq. 3

$b_1$ Parameter in Eq. 3

$b_2$ Parameter in Eq. 3

c Liquid covolume

d Equation of state parameter

$k$ Binary interaction parameter

$N$ Number of points (experimental or auxiliary)

$NC$ Number of components in the mixture

$P$ Pressure

$R$ Gas constant

$T$ Temperature

$v$ Molar volume

$x$ Mole fraction

$Z$ Compressibility factor

*Subscript*

c Critical point property

$i$ Relative to the substance $i$

$j$ Relative to the substance $j$

$ij$ Relative to the interaction between substance $i$ and the substance $j$

$r$ Reduced property

$t$ Triple point property

$1$ Relative to methane

$2$ Relative to hydrogen sulfide

*Superscript*

calc Calculated value

exp Experimental value
\[ m \text{ Parameter in Eq. 3} \]
\[ n \text{ Parameter in Eq. 2} \]

**Acronyms**

AAD Average Absolute Deviation  
EoS Equation of State  
L Liquid phase  
LLE Liquid - Liquid Equilibrium  
LLVE Liquid - Liquid - Vapor Equilibrium  
MAD Maximum Absolute Deviation  
QP Quadruple Point  
S Solid phase  
SLE Solid - Liquid Equilibrium  
SLL Solid - Liquid - Liquid Equilibrium  
SLVE Solid - Liquid - Vapor Equilibrium  
SSE Solid - Solid Equilibrium  
SSLE Solid - Solid - Liquid Equilibrium  
SSVE Solid - Solid - Vapor Equilibrium  
SVE Solid - Vapor Equilibrium  
V Vapor phase  
VLE Vapor - Liquid Equilibrium  
UCEP Upper Critical EndPoint

**Literature Cited**


