Below we first show our detailed responses to the comments of the reviewers, followed by a revised manuscript version showing the changes marked in blue.

In addition to the changes discussed in detail in our response to the reviewers, we also added a co-author, Carl Percival, who was unfortunately not listed in our initial submission.

We thank reviewer 1 for the positive feedback and her/his suggestions. Below, we answer the specific comments one by one.

1) The treatment of elevated RH in EDB is not very clear. The authors presented the relationship between water uptake by some PEGs and water activity (Fig. 1), and mentioned this in line 14-16, page 10. But in the description of the EDB setups, there is no information about whether possible water evaporation was taken into account for the saturation vapor pressure measurements or not. Does the error associated with RH in EDBs include this effect or after water evaporation is taken into account?

Both reviewers are missing a detailed description on how we treat the data taken at elevated RHs in the EDB setups. We presented this in detail in previous publications, e.g. (???). We agree with the reviewers that the manuscript becomes more accessible when this analysis is presented again. Water evaporation is taken into account as the composition of the binary particle is constant with fixed relative humidity and temperature. Hence with each PEG evaporating the corresponding number of water molecules is evaporating from the particles as well to maintain constant composition. The error caused by the uncertainty of RH within the EDB is small compared to the error introduced by the uncertainty of the gas phase diffusivity of PEG and is neglected in our analysis.

We will add the following Appendix to the manuscript:

0.1 Appendix: Treatment of EDB data taken at elevated relative humidities

Evaporation rates in the EDB setups were measured not only under dry conditions but some also at elevated relative humidities. After the particle was equilibrated with respect to temperature and RH, any change in size, $\frac{dr^2}{dt}$, was attributed to evaporative loss at a constant composition, and the partial pressure of the PEG, $p_{PEG}$, under those conditions was calculated using Eq. (1) of ?:

$$p_{PEG} = \frac{-1}{2} \frac{dr^2}{dt} \frac{x\rho RT}{(xM_{PEG} + (1-x)M_{H_2O})D},$$  \hspace{1cm} (1)$$

where $r$ is the particle radius, $x$ is the mole fraction of PEG, $\rho$ is the density of the condensed phase, $R$ is the gas constant, $M$ is molar mass, and $D$ is the diffusivity of the PEG vapor in the gas phase. The mole fraction of PEG in an aqueous particle was taken from the RH in the EDB (see Fig. 1 of the manuscript) by assuming that the particle water activity is equivalent to measured RH, i.e., that the particle is homogeneous and in thermodynamic equilibrium (which is a good approximation as long as evaporation is sufficiently slow). Density was estimated by assuming ideal mixing. As an example we show data obtained by the EDB setup of Union College at 286.5 K for PEG4 in Fig. ??.

Extrapolation of the data to a PEG activity equal one yields the saturation pressure of PEG at this temperature.
Figure 1. Partial pressures of PEG4 versus PEG4 activity deduced from evaporation rates of single, levitated, aqueous PEG4 particles in the EDB setup of Union College at a temperature of 286.5 K. Black symbols: experiment data, red line and shaded area: linear fit to these data with the 95% confidence interval. Extrapolation to $a_{\text{PEG}}$ equal one yields the saturation vapor pressure at this temperature, i.e. $p^0 = (5.69 \pm 0.55) \cdot 10^{-3}$ Pa at this temperature.

2) Page 3, Line 12 – 16. It is totally justifiable that the authors used PEGs for the purpose of this study. However, apart from the advantages presented here, I suggest the authors also mention a few caveats of using PEGs: atmospheric occurrence would not be very likely, but what about structural similarity with commonly found components in organic aerosols?

We agree with the reviewer that the occurrence of PEGs in the atmosphere is unlikely. However, polyols in general have been observed frequently in the atmosphere as well as polyethers. Clearly, the hydroxyl group of the PEGs is a common functional group in organic aerosol compounds. However, we feel there is no need in the present context for direct atmospheric relevance of the compounds. More important for the selection of the PEGs are the three arguments listed at the end of the introduction section. We will add a fourth point to the revised manuscript:

PEGs contain ether and alcohol functional groups which are both abundant in the organic fraction of atmospheric aerosols. Although we do not expect the presence of larger PEGs in the atmosphere there functional groups renders them nevertheless relevant for the atmosphere.
3) Page 15, line 5: the use of accommodation coefficient is not clearly stated. Is it unity in all cases needed? In line 32 on page 7, it was only stated that “for all measurements the accommodation coefficient is assumed to be identical between samples.” Please specify.

We thank the reviewer for pointing out that our description is not quite precise here. For the KEMS setup it is important that the accommodation (or more precise the evaporation) coefficient is identical between reference samples and measured compounds, but its value may differ from unity. It effectively cancels out when relating the measured to the reference samples. For the FT-TDMA setup we assume it to be unity. The close agreement between the FT-TDMA data with those of the other techniques let us conclude that the accommodation coefficient is close to one.

We will change line 32 on page 7 to: For all measurements the accommodation coefficient is assumed to be identical between reference samples and the PEGs.

4) If the authors suggest that this dataset can be used to validate vapor pressure measurements for other techniques by other researchers, a note on data accessibility would be helpful: included in supporting information? Contact authors?

We agree. We made the data available already with the AMTD submission. It is available under the Assets button on the AMT web-page. The direct link is https://doi.org/10.3929/ethz-b-000168628.

Technical corrections.

We will change the manuscript accordingly.
References


We thank reviewer 2 for the positive feedback and her/his thoughtful suggestions. Below, we answer the specific comments one by one, but start with question #2 as the answer to it provides the framework for answering question #1.

2) **Change in evaporation enthalpy with temperature** One of the issues (related to the previous comments) that the authors avoid discussing in any detail is the change in evaporation enthalpy with temperature. This change depends on the difference of the heat capacities of the liquid and vapor and of course the temperature change. Reasonable estimates of this effect can be obtained. How important are these effects for the 260-550 K temperature range and the compounds discussed here? Could these effects explain the tendency of the linear regressions used here to exceed the measured values in a number of cases?

We thank the reviewer for raising this question. The enthalpy of evaporation is a function of temperature; the first question being how much this influences the values provided in Table 2 of the manuscript. Let us analyze the data of PEG3 following e.g. Chickos and Acree Jr. (2003). If the heat capacities of the liquid and gas phase are known, \( C_{pl} \) and \( C_{pg} \), respectively, then the vaporization enthalpy, \( \Delta H_{\text{vap}}(T^\circ) \), at \( T^\circ = 298.15 \text{ K} \) can be related to the experimental measurements by using Eq. 1, generally referred to as Kirchhoff’s equation.

\[
\Delta H_{\text{vap}}(T^\circ) = \Delta H_{\text{vap}}(T) + \int_{T^\circ}^{T} (C_{pl} - C_{pg}) dT,
\]

where \( T^\circ \) refers to the mean temperature of all measurements, which is equal to 357 K for our PEG3 data when calculating the mean of the inverse temperature range covered by the data. If we estimate \( (C_{pl} - C_{pg}) \) by using the method suggested by Chickos and Acree Jr. (2003) (which should be accurate to about \( \pm 30 \text{ J} \cdot \text{mol}^{-1} \text{K}^{-1} \)):

\[
(C_{pl} - C_{pg}) = (10.56 + 0.26C_{pl}),
\]

it yields \( 101\pm30 \text{ J} \cdot \text{mol}^{-1} \text{K}^{-1} \) for the difference between heat capacity of the liquid and heat capacity of the gas phase, when we take the value for the heat capacities of the liquid phase, \( C_{pl} \), from Stephens and Tamplin (1979). Using Eq. 1 with this average value for \( (C_{pl} - C_{pg}) \) yields an increase to the uncorrected enthalpy of vaporization of \( 6.0\pm1.8 \text{ kJ} \cdot \text{mol}^{-1} \) when using the temperature difference between mean temperature, \( T^\circ \), and 298.15 K. Therefore, the enthalpy at the mean temperature of the measurements obtained by the linear regression, which is \( \Delta H_{\text{vap}}(\text{exp}, T^\circ) = 78.3\pm0.7 \text{ kJ} \cdot \text{mol}^{-1} \), should be corrected to \( \Delta H_{\text{vap}} = 84.3\pm1.9 \text{ kJ} \cdot \text{mol}^{-1} \) at 298.15 K. This corresponds to a shift of 7.5 %, which is clearly non-negligible. The magnitude is somewhat smaller for the higher mass PEGs. In the revised manuscript we calculate the enthalpy for 298 K for all PEGs and will provide the additional data in an updated Table 2 as given below and correct the values plotted in Fig. 11 of the manuscript as shown in Fig. 1 and add a few sentences explaining how we calculated the correction.

To answer the question whether the temperature dependence of the enthalpy of evaporation can explain the deviations seen in the linear regression, we integrate the Clausius-Clapeyron equation, eq. (3) in a more general form, by including Kirchhoff’s equation, eq. (1), in the same manner as above, namely assuming a constant, temperature independent, heat capacity difference
Figure 1. Enthalpies for all PEGs, open circles this study, open squares Chickos and Acree Jr. (2003). Filled circles: computational predictions from this study. Dashed line is a fit through the data of PEG4 to PEG8.

Table 1. Saturation vapor pressures at 298.15 K and enthalpies of vaporization of the polyethylene glycols measured in this study as well as estimated enthalpies of vaporization at 298.15 K, together with the results of the quantum chemistry calculations. In addition we provide experimental data from the literature for ethylene glycol and diethylene glycol.

<table>
<thead>
<tr>
<th>n</th>
<th>p(exp) [Pa]</th>
<th>ΔH_vap(exp, T) [kJ · mol⁻¹]</th>
<th>T [K]</th>
<th>ΔH_vap(exp, 298 K) [kJ · mol⁻¹]</th>
<th>p(calc) [Pa]</th>
<th>ΔH_vap(calc, 298 K) [kJ · mol⁻¹]</th>
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</thead>
<tbody>
<tr>
<td>triethylene glycol</td>
<td>3</td>
<td>(6.68 ± 1.10) × 10⁻²</td>
<td>78.3 ± 0.7</td>
<td>349</td>
<td>84.3 ± 1.9</td>
<td>2.65 × 10⁰</td>
</tr>
<tr>
<td>tetraethylene glycol</td>
<td>4</td>
<td>(1.69 ± 0.11) × 10⁻²</td>
<td>77.1 ± 0.4</td>
<td>345</td>
<td>82.1 ± 1.6</td>
<td>1.82 × 10⁻¹</td>
</tr>
<tr>
<td>pentaethylene glycol</td>
<td>5</td>
<td>(5.38 ± 0.71) × 10⁻¹</td>
<td>90.6 ± 1.1</td>
<td>339</td>
<td>94.6 ± 1.6</td>
<td>2.10 × 10⁻²</td>
</tr>
<tr>
<td>hexaethylene glycol</td>
<td>6</td>
<td>(3.15 ± 0.58) × 10⁻⁵</td>
<td>102.1 ± 1.5</td>
<td>351</td>
<td>107.2 ± 2.1</td>
<td>–</td>
</tr>
<tr>
<td>heptaethylene glycol</td>
<td>7</td>
<td>(1.38 ± 0.47) × 10⁻⁶</td>
<td>113.7 ± 2.7</td>
<td>357</td>
<td>119.2 ± 3.2</td>
<td>–</td>
</tr>
<tr>
<td>octaethylene glycol</td>
<td>8</td>
<td>(9.2 ± 20.4) × 10⁻⁸</td>
<td>124.2 ± 9.7</td>
<td>363</td>
<td>130.2 ± 9.9</td>
<td>–</td>
</tr>
<tr>
<td>ethylene glycol</td>
<td>1</td>
<td>11.7 ± 0.6⁷</td>
<td>–</td>
<td>–</td>
<td>65.6 ± 0.3⁶</td>
<td>3.98 × 10¹</td>
</tr>
<tr>
<td>diethylene glycol</td>
<td>2</td>
<td>0.6 ± 0.03⁶</td>
<td>66.9 ± 0.3⁶</td>
<td>420</td>
<td>80.2 ± 4.0⁶</td>
<td>3.80 × 10⁰</td>
</tr>
</tbody>
</table>

Figure 2. $\Psi$ versus inverse temperature difference, $(1/T - 1/T^\circ)$, filled circles: all data but Wise et al. (1950), red line: linear regression between liquid and gas phase.

$$\frac{dp}{p} = \frac{\Delta H_{\text{vap}}(T)}{R} \frac{dT}{T^2}, \text{ with } \Delta H_{\text{vap}}(T) = \Delta H_{\text{vap}}(T^\circ) - (C_{p_l} - C_{p_g})(T^\circ - T)$$

Integration of eq. (3) leads to:

$$\ln \left( \frac{p}{p^\circ} \right) = \frac{\Delta H_{\text{vap}}(T^\circ) - T^\circ (C_{p_l} - C_{p_g})}{R} \left( \frac{1}{T} - \frac{1}{T^\circ} \right) + \frac{(C_{p_l} - C_{p_g})}{R} \ln \left( \frac{T}{T^\circ} \right).$$

If we now plot $\Psi := \left\{ R \ln \left( \frac{p}{p^\circ} \right) - (C_{p_l} - C_{p_g}) \ln \left( \frac{T}{T^\circ} \right) \right\}$ versus $1/T - 1/T^\circ$, we expect a straight line with $\Psi(0) = 0$. In Fig. 2 we show such a plot for the PEG3 data.

Clearly, considering the temperature dependence of the enthalpy of evaporation helps to reconcile the high temperature data with those taken by us at roughly room temperatures. However, its effect close to 298 K is minimal by default, see eq. 4. In particular, it does not significantly reduce the differences between our individual measurements. Hence, we prefer to not include this detailed discussion in the main part of the revised manuscript, but we will put it into an appendix.

However, we will use this approach to estimate the vapor pressures at 298.15 K from the literature data at high temperatures without taking our new data into account. This will answer question #1 of the reviewer:

1) **Extrapolation from higher temperatures.** The first conclusion for the paper is that at least for the compounds studied
Figure 3. Saturation vapor pressure at 298.15 K plotted for all PEGs. Open black circles with error bars: results of this study. Filled orange circles: extrapolation of high temperature literature data to 298.15 K using an estimation of the temperature dependence of the enthalpy of evaporation as detailed in appendix B. Open orange circles: extrapolation of high temperature literature data to 298.15 K assuming a constant enthalpy of evaporation. Filled black circles: computational predictions from this study. Open squares with error bars: Ambrose and Hall (1981). Dashed gray line is a linear regression to all the experimental data, dashed red lines are separate linear regressions for n=1 to n=3 and n=4 to n=8.

Extrapolation from higher temperatures can be used instead of the challenging measurements at the lower temperatures. However, the authors miss the opportunity of providing some additional guidance given the availability of data from other studies. Could one obtain accurate estimates for PEG3 extrapolating from the 400-550 K range to room temperature? How about PEG4? Some additional analysis without using the new data would be helpful.

We have two options for this analysis now: first, assuming a temperature independent enthalpy of vaporization (shown as open orange circles in the revised figure of all vapor pressures, i.e. Fig. 3). Second, taking the difference in heat capacity between the two phases temperature independent, i.e. using eq. (4) (shown as filled orange circles in Fig. 3). Extrapolation using the high temperature data to room temperature taking into account the temperature dependence of the enthalpy of evaporation yields very consistent data with the new measurements: both agree within uncertainty estimates for PEG3, PEG7 and PEG8 and agree better than a factor of two for PEG4, PEG5 and PEG6. Taking a constant enthalpy of vaporization yields very good agreement for PEG4, PEG5 and PEG6, but substantial deviation - up to a factor of 10 - for PEG3, PEG7 and PEG8. We conclude that extrapolation to lower temperatures of high temperature data should take Kirchhoff’s law into account.
We will add extrapolated values to Fig. 10 of the manuscript as shown in Fig. 3 here. Also, we will add the following paragraph to page 15 line 10 of the manuscript: Very satisfyingly, extrapolation of the high temperature literature data to 298.15 K incorporating the temperature dependence of the enthalpy of vaporization as detailed in Appendix B (orange filled circles in Fig. 3) yields values agreeing well within error with the newly derived saturation vapor pressures for PEG3, PEG7 and PEG8 of this study and agree better than within a factor of two for PEG4, PEG5 and PEG6. In contrast, using a constant enthalpy of vaporization for the extrapolation yields very good agreement for PEG4, PEG5 and PEG6, but substantial deviation - up to a factor of 10 - for PEG3, PEG7 and PEG8. We conclude that extrapolation to lower temperatures of high temperature data should take the temperature dependence of the enthalpy of vaporization (Kirchhoff’s law) into account.

3) Measurements at higher relative humidity It is not quite clear which measurements were performed at higher RH and how they were analyzed. Figure 1 should be the key here, but it is too early in the manuscript and is not well connected to the corresponding results.

As reviewer 1 raised the same question we refer to our answer to this question of reviewer 1.

4) Mass accommodation coefficient How close to unity can the mass accommodation coefficients be based on this data set? More than 0.9, 0.5, 0.1, something else? This is a nice result of the study and a little extra analysis could be helpful.

We thank reviewer 2 for the suggestion. We reanalyzed all FT-TDMA data by fixing the surface tension to the values provided by Gallaugher (1932), taking the vapor pressures of Table 1 as reference pressure and varying the accommodation coefficient, $\alpha$, until the calculated vapor pressure of an FT-TDMA experiment including its error exceeds the reference pressure confidence interval at the temperature of the FT-TDMA experiment. This yields a lower limit for $\alpha$ for each FT-TDMA experiment, with a mean value for $\alpha$ of all experiments equal to 0.5 and no discernible difference between the different PEGs. We will add the following sentences at line 5 page 15 in the revised manuscript:

To retrieve a lower limit for the accommodation coefficient, we analyzed the FT-TDMA experiments varying the accommodation coefficient until the calculated vapor pressure including its error exceeds the reference pressure confidence interval given in Table 1 at the temperature of each FT-DMA experiment. The mean value of these accommodation coefficients for all experiments is equal to 0.5 and there is no discernible difference between the experiments for different PEGs. We conclude that the mass accommodation coefficient of the PEGs based on our study is greater than 0.5.

5) PEG8 melting point The PEG8 melting point is quite close to some of the measurements in Fig. 9. Could this be one more factor contributing to the variability of these data points.

When taking the data at temperatures close to the melting point with the EDB setup there was no indication for any crystallization happening. The light scattering data in the EDB setup are very sensitive to phase transitions, see e.g. (Braun and
Krieger, 2001). In addition, crystallization should yield lower apparent vapor pressures, but the temperature data with the KEMS setup show a reverse temperature trend.

6) **Surface free energies** The authors measured the surface free energies in some of the systems but these measurements are not included in Table 2. Were these estimates compared to literature values for these compounds?

For the analysis of the FT-TDMA data, surface tensions are needed to take into account the Kelvin effect. The data shown in the manuscript used surface tensions derived from the FT-TDMA experiments. However, as precise surface tension data are available (Gallaugher and Hibbert, 1937; Gallaugher, 1932), now we use those for the analysis of the FT-TDMA experiments in the revised version and include them in the revised version of Table 2. (Since those are approximately - within 5‰- the same for PEG5 to PEG7, we used the one for PEG7 in our calculations.) The figures and tables are revised accordingly (Table 1 and Fig. 3 show already the revised data.) Switching to the surface tensions reported in literature changes the values of the estimated vapor pressures by a very small amount.

7) **Presentation of results** I do like the idea of having cut-outs of the figures but I was confused by the fact that these were different figures in different pages. It is probably better to combine Figures 2 and 3 into a two-part Figure 2, Figures 4 and 5 into a two-part Figure 3. This will also allow the addition of cut-outs of the rest of the figures and help the discussion of the results. It took me some effort to see the difference of a factor of 2 mentioned in the text. It would also be helpful if the authors used filled color symbols to separate the measurements performed in the present work with those existing in the literature. The use of large symbols together with the small uncertainties in Figure 11 result in confusion. It appears that there are different symbols in the first 4 cases.

We will follow the advice and put the cut-outs into two-part figures. We tried filled symbols, however, then it became even more difficult to clearly relate the symbols to a specific data set as they tended to cover each other. We will increase the symbol edge thickness to make the distinction between our data and literature data more clear. We re-size the symbols for Fig. 11.
References


A reference data set for validating vapor pressure measurement techniques: Homologous series of polyethylene glycols

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Abstract. To predict atmospheric partitioning of organic compounds between gas and aerosol particle phase based on explicit models for gas phase chemistry, saturation vapor pressures of the compounds need to be estimated. Estimation methods based on functional group contributions require training sets of compounds with well established saturation vapor pressures. However, vapor pressures of semi- and low volatile organic molecules at atmospheric temperatures reported in the literature often differ by several orders of magnitude between measurement techniques. These discrepancies exceed the stated uncertainty of each technique which is generally reported to be smaller than a factor of two. At present, there is no general reference technique for measuring saturation vapor pressures of atmospherically relevant compounds with low vapor pressures at atmospheric temperatures. To address this problem we measured vapor pressures with different techniques over a wide temperature range for intercomparison and to establish a reliable training set. We determined saturation vapor pressures for the homologous series of polyethylene glycols (H−(O−CH₂−CH₂)n−OH) for n=3 to n=8 ranging in vapor pressure at 298 K from 10⁻⁷ Pa to 5·10⁻² Pa and compare them with quantum chemistry calculations. Such a homologous series provides a reference set that covers several orders of magnitude in saturation vapor pressure, allowing a critical assessment of the lower limits of detection of vapor pressures for the different techniques as well as permitting the identification of potential sources of systematic error. Also, internal consistency within the series allows to reject outlying data more easily. Most of the measured vapor pressures agreed within the stated uncertainty range. Deviations mostly occurred for vapor pressures values approaching the lower detection limit of a technique. The good agreement between the measurement techniques (some of which are sensitive to the mass accommodation coefficient and some not) suggest that the mass accommodation coefficients of the studied compounds are close to unity. The quantum chemistry calculations were about one order of magnitude higher than the measurements. We find that extrapolation of vapor pressures from elevated to atmospheric temperatures is permissible over a range of about 100 K for
these compounds, suggesting that measurements should be performed best at temperatures yielding the highest accuracy data allowing subsequent extrapolation to atmospheric temperatures.

1 Introduction

Atmospheric oxidation of organic vapors can lead to low- and semivolatile organic compounds (LVOCs and SVOCs) which are multifunctional in nature with molar masses typically between 150 g · mol⁻¹ and 300 g · mol⁻¹ and saturation vapor pressures between 0.1 Pa and 10⁻⁷ Pa (Jimenez et al., 2009; O’Meara et al., 2014). In this range of vapor pressures individual compounds partition dynamically between the gas and particle phases depending on total aerosol loading and temperature. Recently, highly oxidized organic compounds with even lower saturation vapor pressures (ELVOCs), which are essentially non-volatile, have also been detected in both laboratory studies and in the atmosphere (Ehn et al., 2012). These compounds are also produced primarily in the gas phase and their saturation vapor pressures are needed to constrain their flux onto the atmospheric particulate phase. However, not all highly oxidized multifunctional compounds (HOMs) show extremely low saturation vapor pressures and hence HOMs may still partition between gas and particle phase (Kurtén et al., 2016).

If we aim to predict atmospheric partitioning using a bottom-up approach using explicit or near explicit models for gas phase oxidation of volatile organic compounds (VOCs), reliable model estimations of saturation vapor pressure for individual compounds are needed. Most estimation methods are constrained by databases heavily biased toward monofunctional compounds with saturation vapor pressures more in the range of ~ 10³ - 10⁵ Pa (Bilde et al., 2015). Experimental data of multifunctional compounds are required to improve estimation methods for atmospheric applications. However, even for the straight-chain dicarboxylic acids the experimental saturation vapor pressures reported in the literature deviate by up to four orders of magnitude between different measurement techniques (Bilde et al., 2015) and the difference can become as large as six orders of magnitude when adding additional functional groups to the straight-chain dicarboxylic acids (Huisman et al., 2013). These differences are strikingly larger than the error estimates for the individual techniques, which are at most stated as a factor of two. A very interesting observation of the Bilde et al. (2015) study when comparing different data sets was: “...that the temperature dependence of the saturation vapor pressure, i.e., the slopes of the individual data sets (the enthalpies of sublimation and vaporization), agree almost always better with each other than the reported saturation vapor pressures themselves.” Obviously, there are systematic biases of the different techniques, which are neither fully understood nor characterized.

Another important aspect of the differences observed between different datasets was identified to be caused by the physical state of the compound studied (Soonsin et al., 2010; Bilde et al., 2015). As multicomponent aerosol particles are expected often to be liquid or amorphous (glassy) under atmospheric conditions (Marcolli et al., 2004), the reference state for atmospheric applications is in general the subcooled liquid. If the vapor pressure is measured in the solid crystalline state, it needs to be converted to that of the subcooled liquid by taking the enthalpy of fusion and the change in molar heat capacity upon the crystalline solid to liquid transition into account (e.g. Bilde et al. (2015)), introducing corresponding uncertainties. In addition, it is not always certain that a solid crystalline compound is purely crystalline without any amorphous content. Any amorphous material present can enhance the vapor pressure significantly (Soonsin et al., 2010), as the saturation vapor pressure over an
amorphous solid resembles more that of the subcooled liquid than that of the crystalline solid. Therefore, a reference material for instrument testing should be preferable in liquid state at the temperatures at which the measurements are performed.

At present, there is no reference instrument nor technique which could serve as a “gold standard” for measuring the saturation vapor pressures of atmospheric multifunctional compounds. Without such a standard the lower limit of detection of a particular technique is difficult to access with reference materials available only with saturation vapor pressures above the targeted pressure range. Reference data of a homologous series could help to assess measurement techniques and instruments, and allow to establish detection limits. Internal consistency within a series increases its trustworthiness for establishing training sets needed for developing estimation methods.

Here, we use five different experimental setups based on three different techniques as well as high temperature data reported in the literature to build such a reference data set. We determine saturation vapor pressures and enthalpies of vaporization for the homologous series of polyethylene glycols (H−(O−CH2−CH2)n−OH) for n=3 to n=8 ranging in vapor pressure at 298 K from $10^{-7}$ Pa to $5 \cdot 10^{-2}$ Pa. Polyethylene glycols are chosen for four reasons: first, they are liquids at room temperature, so one of the potential sources of disagreement between different data sets identified by Bilde et al. (2015) is removed. Second, high temperature saturation vapor pressures have been reported for some of the compounds in the literature and can be combined with our measurements performed at room temperature. Third, they span over a large range of saturation vapor pressures relevant for atmospheric applications. Fourth, PEGs contain ether and alcohol functional groups which are both abundant in the organic fraction of atmospheric aerosols. Although we do not expect the presence of larger PEGs in the atmosphere there functional groups renders them nevertheless relevant for the atmosphere.

2 Materials

Polyethylene glycols with an oligomer purity of $> 98\%$ (penta- to octaethylene glycol from Polypure AS, Oslo, Norway) and an oligomer purity of $> 98.5\%$ (tri- and tetraethylene glycol, Sigma-Aldrich), were used as received without any additional purification. Clearly, oligomer purity poses a potential problem for evaporation measurements: impurities of more volatile oligomers will lead to faster evaporation, while oligomers with lower volatility will bias toward lower evaporation. Samples of the same batch were shared for measurements with all five experimental systems. Melting points measured by differential scanning calorimetry and refractive indices measured with an Abbe-type refractometer are given in Table 1 as well as measured (Crespo et al., 2017) and estimated (Poling et al., 2001) densities. Also given are estimations for the gas phase diffusivity in air. Gas phase diffusivities of the organics are needed to calculate vapor pressures from measured evaporation rates by the electrodynamic balance (EDB) setups as well as those of the laminar flow tube - tandem differential mobility analyzer (FT-TDMA) set-up. Since data are only available for triethylene glycol, diffusivities were estimated following Bird et al. (2007) and using $T_c$ from UManSysProp (Topping et al., 2016). In addition, water activity measurements of a PEG400 mixture with mean molecular weight of 400 g·mol$^{-1}$ have been performed with a water activity meter (AquaLab, Model 3B, Decagon Devices, USA) to estimate PEG activities for aqueous PEG-mixtures, see Figure 1.
Table 1. Physical properties of the polyethylene glycols (H−(O−CH₂−CH₂)n−OH) used in this study: n, molar mass M, melting temperature Tₘ, gas phase diffusivity in air, Dₚ, estimated as described in Bird et al. (2007) using Tc from UManSysProp (Topping et al., 2016), refractive index nD, surface tension σ. If not indicated otherwise, the data are from this study.

<table>
<thead>
<tr>
<th></th>
<th>n</th>
<th>M [g · mol⁻¹]</th>
<th>Tₘ [K]</th>
<th>Dₚ [m² · s⁻¹]</th>
<th>nD</th>
<th>ρ [g · cm⁻³]</th>
<th>σ [mN · m⁻¹]</th>
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<td></td>
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<td></td>
<td>at 25°C, 1013 hPa</td>
<td></td>
<td>at 25°C</td>
<td>at 25°C</td>
<td></td>
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<tr>
<td>triethylene glycol</td>
<td>3</td>
<td>150.2ᵃ</td>
<td>266.0ᵇ</td>
<td>5.95 · 10⁻⁶</td>
<td>1.4557ᶜ</td>
<td>1.108</td>
<td>45.15ʰ</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>(5.90 ± 0.059) · 10⁻⁶ᵈ</td>
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<tr>
<td>tetraethylene glycol</td>
<td>4</td>
<td>194.2ᵃ</td>
<td>267.6ᶠ</td>
<td>5.20 · 10⁻⁶</td>
<td>1.4593ᶜ</td>
<td>1.132</td>
<td>45.08ʰ</td>
</tr>
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<tr>
<td>pentaethylene glycol</td>
<td>5</td>
<td>238.4ᵍ</td>
<td>270.3± 1.0</td>
<td>4.66 · 10⁻⁶</td>
<td>1.4592</td>
<td>1.155</td>
<td>44.95ʰ</td>
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<tr>
<td>hexaethylene glycol</td>
<td>6</td>
<td>282.3ᵍ</td>
<td>281.2± 1.0</td>
<td>4.26 · 10⁻⁶</td>
<td>1.4623</td>
<td>1.180</td>
<td>44.87ʰ</td>
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<tr>
<td>heptaethylene glycol</td>
<td>7</td>
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<td>289.2± 1.0</td>
<td>3.94 · 10⁻⁶</td>
<td>1.4636</td>
<td>1.206</td>
<td>44.73ʰ</td>
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<td></td>
</tr>
<tr>
<td>octaethylene glycol</td>
<td>8</td>
<td>370.4ᵍ</td>
<td>297.4± 1.0</td>
<td>3.68 · 10⁻⁶</td>
<td>1.4643</td>
<td>1.234</td>
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</tr>
</tbody>
</table>

ᵃ Sigma-Aldrich,ᵇ Curme and Johnston (1952),ᶜ Gallaugher and Hibbert (1936) at 20 °C,ᵈ Lugg (1968),ᵉ Crespo et al. (2017),ᶠ Oakwood Chemical,ᵍ Polypure AS, Norway,ʰ Gallaugher (1932).

3 Methods

We used five experimental systems based on three different techniques to determine saturation vapor pressures. Detailed descriptions of the instruments have been published previously, but we provide brief descriptions here focusing on the uncertainties of vapor pressures derived in the following sections. Three systems use single, micrometer-size particles levitated in an electrodynamic balance (EDB) to measure diffusion-controlled evaporation rates, namely the EDB setup at ETH Zurich (see Sect. 3.1), the EDB setup at Union College (see Sect. 3.2) and the EDB setup at the University of Bristol (see Sect. 3.3). Besides slightly different electrode configurations, the main difference of the EDB setups is the injection procedure, the accessible temperature range and how the particles are sized during evaporation. For deducing vapor pressures from evaporation rates, all make use of Maxwell’s quasi-stationary approximation (Maxwell, 1877) for evaporation of a motionless spherical particle relative to a uniform gaseous medium, with the particle’s radius being large compared to the mean free path of the evaporating species, referred to as the continuum regime. The fourth setup, the Aarhus FT-TDMA instrument (see Sect. 3.4) uses a differential mobility analyzer (DMA) to select monodisperse particles in the accumulation mode size range and uses a scanning mobility particle sizer (SMPS) to measure the size distribution of the particles before and after partial evaporation in a laminar flow tube. A key parameter needed for quantitative analysis for the EDB experiments as well as the FT-TDMA setup is the diffusivity of the evaporating species in the gaseous medium. In the FT-TDMA setup the particles radii are of similar magnitude as the mean free path of the evaporating molecules (referred to as the transition regime). In this case, obtaining
Figure 1. Water and PEG activities versus mole fraction of PEG for PEGs with three different mean molar masses: PEG200 (black), PEG400 (red) and PEG600 (purple). Filled data points are from this study, open symbols from Ninni et al. (1999). The dashed lines black lines illustrate ideal behavior, the short dashed lines are van Laar fits to the water activity data, the solid lines are the PEG activities obtained from the van Laar fits using the Gibbs-Duhem relation.

vapor pressures from measured evaporation rates requires knowledge of the mass accommodation coefficient (e.g. Bilde et al. (2015)). The fifth instrument, the University of Manchester KEMS, measures the gas phase concentration of the vapor effusing from a macroscopic sample in a Knudsen cell using mass spectrometry, see Sect. 3.5. Here, it is assumed that the sample establishes an equilibrium within the cell.

In a Knudsen cell the mean free path is so large that mass transport can be described using kinetic gas theory, again with mass accommodation being a key parameter.

3.1 EDB setup ETH Zurich

The electrodynamic balance setup at ETH Zurich uses a double ring configuration (Davis et al., 1990) to levitate a charged particle in an environmental cell with a gas flow free of the evaporating species under investigation, but allowing precise conditioning of temperature and relative humidity (Zardini et al., 2006; Soonsin et al., 2010; Huisman et al., 2013). Evaporation rates are measured at fixed temperature and relative humidity using optical resonance spectroscopy in backscattering geometry with a broadband LED source and applying Mie theory for the analysis (Zardini et al., 2006). The accuracy of the rate measurement is estimated to be $\pm 1 \cdot 10^{-6} \, \mu m^2 s^{-1}$. Huisman et al. (2013) did a detailed error analysis for the setup and evaluation scheme for
determining vapor pressures from the measured evaporation rates and concluded that the largest systematic uncertainty arises from estimating the gas phase diffusivity (±20% uncertainty in vapor pressure), while possible drifts in relative humidity could lead to some error (2%) when measuring at higher humidity. Obviously, the uncertainty associated with a measurement depends also on the total time span observing the evaporation. Previously, we estimated the lower limit of quantification for our setup to be about $4 \cdot 10^{-7}$ Pa for a 24 h measurement period. Conservatively, we estimated the uncertainty for each individual measurement to be ±35% plus the lower limit of quantification based on the total time spent to measure the rate. To avoid being biased by impurities of higher volatility, measurements were started typically 6 hours after injecting the particle in the EDB.

### 3.2 EDB setup Union College Schenectady

The electrodynamic balance at Union College uses a double ring configuration similar to that of the ETH Zurich unit. Temperature and relative humidity are monitored via a precision PT100 temperature sensor (Omega PRTF-10-3-100-1/4-3/4-E) integrated into the chamber and a chilled-mirror dewpoint hygrometer (General Eastern Optisonde-1-1-5-2-2-1-A-0) immediately after the chamber. All gas transfer lines are heated to at least 33°C but no more than 45°C to prevent condensation of water vapor. Evaporation rates are measured at fixed temperature and relative humidity by optical resonance spectroscopy using a broadband LED ($\lambda = 850\pm 20$ nm) at a 90 degree scattering angle, applying Mie theory for the analysis following Zardini et al. (2006). Propagation of error was used to estimate uncertainty in the calculated vapor pressures; estimated uncertainty in temperature is set as 0.25 K; estimated uncertainty in humidity is conservatively set at 2%; uncertainty in density and diffusivity are set at 20%. The estimated uncertainty that results is driven almost entirely by the uncertainty in RH, except at low RH (e.g., <25%) when all other terms contribute approximately equally.

### 3.3 EDB setup University of Bristol

The EDB instrument has been described extensively in previous publications (Davies et al., 2012, 2013; Rovelli et al., 2016). EDB measurements at the University of Bristol were performed with a charged droplet confined between two cylindrical electrodes, in a temperature (248 to 330 K) and relative humidity (RH, 0% to >90%) controlled chamber. In these measurements, the organic droplet was always trapped in a dry flow of nitrogen gas (assuming an RH equal to 0%). The organic evaporating droplet was illuminated with a 532 nm laser and the evaporation profile determined from the detected phase function using the geometric optics approximation ($\pm 100$ nm) (Glatschnig and Chen, 1981). Several evaporating droplets were collected to ensure measurement reproducibility, with a minimum of five droplets collected for each PEG compound at each temperature. The derived vapor pressure is an a average of all measurements taken, which have very good reproducibility. The error associated with this measurement therefore corresponds to the error estimated for the diffusion constant used in the determination of vapor pressure (20%). At the temperatures studied here, the uncertainty in temperature is smaller than the size of the points shown, increasing from 1°C at 15°C to 2°C at 40°C.
3.4 FT-TDMA setup University of Aarhus

ARAGORN (Aarhus Aerosol Gas evapORatioN flow tube) is a Tandem Differential Mobility Analyzer (TDMA) set-up including a laminar Flow Tube (Bilde et al., 2003; Emanuelsson et al., 2016) allowing for studies under dry (e.g. (Bilde et al., 2003; Frosch et al., 2010)) or humid (e.g. (Riipinen et al., 2006; Zardini et al., 2010)) conditions. In this work dried aerosol particles from nebulized aqueous solution were size selected to monodisperse size distributions (69 to 285 nm) using a differential mobility analyzer (DMA), diluted with dry clean air, and allowed to evaporate in a temperature controlled (282–322 K) laminar flow tube at ambient pressure. The peaks remained monodisperse during evaporation and the initial and final peak sizes (typically after 40 s of evaporation) were measured using a SMPS consisting of a DMA (Hauke Vienna short type, negative power supply FUG HCE) connected to a condensation particle counter (CPC TSI 3776). Saturation vapor pressures for PEG5, PEG6, and PEG7 were derived from experimental data as described in Bilde et al. (2003), using a mass accommodation coefficient equal to one, gas phase diffusivities reported in Table 1, and subcooled liquid densities at the corresponding temperature using Poling et al. (2001). For the analysis of the FT-TDMA data, surface tensions are needed to take into account the Kelvin effect. As precise surface tension data are available (Gallaugher and Hibbert, 1937; Gallaugher, 1932) (see Table 1), we use those for the analysis of the FT-TDMA experiments. (Since they are approximately the same for PEG5 to PEG7 - within 5‰-, we used the one for PEG7 in our calculations.) The relative uncertainty on the individual measured saturation vapor pressures is estimated to be around 50% (Bilde et al., 2003).

3.5 KEMS setup University of Manchester

The Knudsen Effusion Mass Spectrometer (KEMS) has been previously described in multiple publications (e.g.(Booth et al., 2009)). Errors for solid-state measurements with the KEMS have previously been determined by calculating the standard deviation of measured species using different reference samples. For example Booth et al. (2009) used 3 different reference compounds to calculate the error for oxalic acid. Based on the numerous repeat runs presented in Booth et al. (2010) over a range of 20 K, a maximum error of ±40% was determined in the solid state. Deriving the subcooled liquid pressure from solid state vapor pressure measurements increased the error to a maximum of ±75% and is representative of all previously published measurements on solid samples. The maximum uncertainty associated with the PEG series is estimated to be ±40%, as the sample is measured in the liquid state where no subcooled liquid correction is required. Additional uncertainties are expected to increase this general error as we approach the lower limit of the instrument. In these regions, the mass spectrometry technique will result in decreased signal to noise ratios, introducing a higher error in the measurements. It is also possible that impurities in the sample and the provenance of sample preparation and storage would have an increasing impact on error as the vapor pressure decreases. Random errors are possible as a result of the variation in sensitivity of the mass spectrometer (ionizer and/or detector). This factor is limited by ensuring that the instrumental background signal is consistent between both the reference samples and the PEGs. We therefore feel that this random effect will cause very little error in final results. Systematic errors can be introduced by the choice of reference compounds as measurements, in all cases, are relative to this choice and from calculation of the ionization cross section. For all measurements the accommodation coefficient is assumed to be identical.
between reference samples and the PEGs. Such an assumption may introduce unquantifiable errors, but it is expected that they are minimized by the appropriate choice of similar reference and sample compounds (e.g (Booth et al., 2012)). These sources of uncertainty require further elucidation with an extended set of compounds. For this study, based on the standard deviation of 4 runs of PEG7 an associated error of ±29% is initially calculated, yet we prescribe a value of ±40% based on the aforementioned issues.

3.6 Quantum chemistry calculations

Saturation vapor pressures were calculated for the the PEG series with \( n=1–5 \) using multiple low energy conformers. All conformers were first obtained with MMFF force-field and B3LYP density functional (Becke, 1993a, b) using the Spartan ’14 program (Spartan, 2014). For PEGs with \( n=1–3 \) we performed systematic conformer sampling using the MMFF force-field in order to produce a representative set of unique conformers. We selected all conformers of PEG1 and PEG2 and the conformers within 5 kcal/mol of the lowest MMFF energy conformer of PEG3 and optimized these using the B3LYP/6-31+G(d) level of theory (Hehre et al., 1972). For PEG4 we performed full Monte Carlo sampling using the MMFF force-field finding over 50 000 conformers. We selected the conformers within 5 kcal/mol of the lowest MMFF energy conformer and computed the B3LYP/6-31+G(d) single-point energies of these structures. Next, we optimized the 100 lowest B3LYP/6-31+G(d) energy conformations at the same level of theory. Due to the large number of different conformers for PEG5, we set up the maximum number of conformers to 100 000 and performed Monte Carlo sampling using the MMFF force-field. Then we selected the 100 lowest MMFF energy conformers and optimized the structures at the B3LYP/6-31+G(d) level of theory. The lowest energy conformers were used in gas-phase and COSMO calculations with BP_TZVP_C30_1701 (BP/TZVP) (Schäfer et al., 1994) parametrization using TmoleX version 4.2 (Steffen et al., 2010) and TURBOMOLE version 7.1 (TURBOMOLE, 2016). The BP/TZVP refers to a B88-VWN-P86 functional with a resolution-of-the-identity (RI) approximation together the TZVP basis set and corresponding auxiliary basis set for the RI computation. Calculations yielded the cosmo and energy files which were used as an input for COSMOtherm (Cosmotherm, 2017; Eckert et al., 2002) saturation vapor pressure calculations.

4 Results and discussion

Figures 2 through 7 show the pure compound saturation vapor pressures as a function of inverse temperature for each polyethylene glycol measured with the experimental systems described above together with available literature data. The vapor pressures measured with our setups cover a range from 1 Pa to \( 10^{-8} \) Pa, with literature data extending this range to pressures up to \( 10^{4} \) Pa. Only pentaethylene glycol could be measured with all three techniques, some instruments not being able to measure the glycols with high saturation vapor pressures, because of fast evaporation; other instruments reached their lower limit of detection for glycols with larger molecular weight and low saturation vapor pressure.

Triethylene glycol was measured with the three EDB setups, as shown in Fig. 2(a) and Fig. 2(b) for a cut-out to show our data in more detail. The setup at the University of Bristol allows measurements at elevated temperatures, which agree very well with extrapolated high temperature literature data. While the Union College EDB show slightly larger saturation pressures the
Figure 2. Saturation vapor pressures versus temperature of triethylene glycol (PEG3). Bold open symbols: this work, open symbols: data reported in the literature. Red line: linear regression to all data below 500 K, red shaded area: 95% confidence band of the regression. (a) overview of all data, (b) cut-out showing the pressure range of the experimental data of this study.
agreement with the Bristol EDB data is within error for almost all data points. The setup at ETH Zurich requires a time span of about 30 minutes to establish constant conditions of temperature and relative humidity after injecting a particle, which limits its ability to measure fast evaporation rates. Hence, the measurements for PEG3 were performed at temperatures below room temperature and at relative humidities ranging from almost 0% at the lowest temperatures to 94% relative humidity at 288 K.

As discussed in 3.1 measurements at elevated relative humidity increases the uncertainty in differentiating between evaporation and shrinkage due to drifts in humidity, which has been taken into account by a factor of 2 increase in estimated uncertainty. However, while the data taken at 288 K agrees with the other two EDB data, the lower temperature data are clearly below those of the other two EDBs. The bias could arise from higher mass oligomeric impurities in the sample, since the measurements at lower temperatures required a considerably longer time to reach equilibrium with respect to temperature and humidity during which a large volume of the particle already evaporated with corresponding potential enrichment of such impurities.

Jakubczyk et al. (2010) used an EDB setup as well and their saturation vapor pressures measured at 298 K compares very favorably with the EDB data of the University of Bristol measurement at this temperature. There is one more data set at lower temperatures available (Wise et al., 1950) obtained by observing the formation of condensates of the vapor on a polished mirror. These data seem to significantly overestimate the vapor pressure and was discarded for the following evaluations. In addition, Grenier et al. (1981) have measured gas saturation in an inert carrier gas, and Steele et al. (2002) using an ebulliometric technique have measured the saturation vapor pressure at much higher temperatures from about 400 K up to 500 K.

The data at the highest and lowest temperatures seem to deviate from a Clausius-Clayperon temperature dependence, assuming the enthalpy of evaporation to be independent of temperature. Nevertheless, we included a part of this data when performing a linear regression using the data between 260 K and 500 K yielding a saturation vapor pressure, $p^0$, of $(6.68^{+1.10}_{-0.95}) \cdot 10^{-2}$ Pa at 298.15 K and an enthalpy of evaporation, $\Delta H_{vap}$, of $84.3\pm1.9$ kJ·mol$^{-1}$ at this temperature, see Table 2. Here the error marks the 95% confidence interval of the regression.

Tetraethylene glycol was measured with the three EDB setups and the KEMS, as shown in Fig. 3. These measurements cover a temperature range from 263 K to 318 K, together with data available in the literature at higher temperatures, the complete data set covers a range from 263 K to 482 K. All data fit well to a temperature independent Clausius-Clayperon relationship yielding a saturation vapor pressure, $p^0$, of $(1.69^{+0.11}_{-0.10}) \cdot 10^{-2}$ Pa at 298.15 K and an enthalpy of evaporation, $\Delta H_{vap}(298K)$, of $82.1\pm1.6$ kJ·mol$^{-1}$. Fig. 3(b) shows a cut-out of Fig. 3(a) to allow closer inspection of the data measured with our setups.

Pentaethylene glycol could be measured with all three techniques used in this study, as shown in Fig. 4. Again, our data cover a considerable temperature range of over 50 K: from 266 K to 318 K. The vapor pressure measured over these temperatures range increases from $10^{-5}$ Pa to $10^{-2}$ Pa. A number of measurements performed with the EDB setup of ETH Zurich at elevated relative humidities ranging from dry (less than 3%) to 50% RH at a temperature of 291 K show that the scatter in the deduced pure component vapor pressure increases when measuring at elevated humidities. The data differ by up to a factor of two, which is beyond the estimated error of $\pm35\%$. Of course, when measuring a binary mixture of PEG and water, the activity of the PEG at the relative humidity of the measurement is needed to evaluate the vapor pressure from evaporation rates (Krieger et al., 2012). Since the water activities for polymeric mixtures with mean molecular weights of 200 g·mol$^{-1}$ (PEG200), 400 g·mol$^{-1}$ (PEG400), and 600 g·mol$^{-1}$ (PEG600) have been extensively measured, see Fig. 1, we are able to constrain the
Figure 3. Saturation vapor pressures versus temperature of tetraethylene glycol (PEG4). Bold open symbols: this work, open symbols: data reported in the literature. Red line: linear regression of all data below 495 K. (a) overview of all data, (b) cut-out showing the pressure range of the experimental data of this study.
activity of the PEG in this mass range to better than 20%. In particular at relative humidities lower than 30% the binary system is close to ideal behavior which further reduces any potential error in the activity estimate. Also, there is no systematic trend in vapor pressure with water content of the particle. Hence it is unlikely that the scatter is due to the uncertainty in activity. Rather, there exist other systematic errors when measuring at elevated humidities which are not yet identified.

A linear regression for all pentaethylene data including the high temperature data of Grenier et al. (1981) yield a saturation vapor pressure, $p^0$, of $(5.38^{+0.71}_{-0.63}) \times 10^{-4}$ Pa at 298.15 K and an enthalpy of evaporation, $\Delta H_{\text{vap}}(298\text{K})$, of $94.6 \pm 1.6$ kJ · mol$^{-1}$.

Hexaethylene glycol was measured with the EDB Zurich setup, the Manchester KEMS and the FT-TDMA setup of the University of Aarhus, as shown in Fig. 5. Vapor pressures were measured over a temperature range of 45 K covering a range in pressures from $10^{-6}$ Pa to $10^{-3}$ Pa. All data agree within error, but there appears to be a slightly different temperature trend in the KEMS data and the EDB data. The linear regression for all data including the high temperature data of Grenier et al. (1981) yield a saturation vapor pressure, $p^0$, of $(3.15^{+0.58}_{-0.49}) \times 10^{-5}$ Pa at 298.15 K and an enthalpy of evaporation, $\Delta H_{\text{vap}}(298\text{K})$, of $107.2 \pm 2.1$ kJ · mol$^{-1}$.

Again, heptaethylene glycol could be measured with the EDB Zurich setup, the Manchester KEMS and the FT-TDMA setup of the University of Aarhus, as shown in Fig. 6. The deviation in temperature trends between the three different measurement setups becomes more apparent at the lower vapor pressures of this compound at about room temperature. Nevertheless, all data seem to be consistent with the 95% confidence interval of a linear regression to all data plus the high temperature data of Grenier et al. (1981). The regression yielded a saturation vapor pressure, $p^0$, of $(1.38^{+0.47}_{-0.35}) \times 10^{-6}$ Pa at 298.15 K and an enthalpy
**Figure 5.** Saturation vapor pressures versus temperature of hexaethylene glycol (PEG6). Symbols as in Fig. 2.

**Figure 6.** Saturation vapor pressures versus temperature of heptaethylene glycol (PEG7). Symbols as in Fig. 2.
of evaporation, $\Delta H_{\text{vap}}(298K)$, of $119.2\pm2.7$ kJ·mol$^{-1}$. Note, that a correct temperature trend is crucial for extrapolations to lower temperatures when measuring above room temperature for atmospheric applications. Clearly, relying on one of our datasets alone would yield unrealistic values for the enthalpy of evaporation.

The highest molecular weight compound in the homologous series of PEGs for which a saturation vapor pressure at about ambient temperatures could be measured with our techniques was octaethylene glycol. Both the KEMS instrument as well as the EDB Zurich instrument were used to determine vapor pressures with temperatures ranging from 296 K to 313 K. The vapor pressure at these temperatures range from less than $10^{-7}$ Pa to about $10^{-6}$ Pa. The EDB data scatter over one order of magnitude and are about one order of magnitude smaller than those measured using the KEMS instrument. While the KEMS data do not show scatter, they show almost no temperature dependence, which may indicate that the lower limit of detection is reached at these vapor pressures. Again we did a linear regression including the high temperature data of Grenier et al. (1981) resulting in a saturation vapor pressure, $p^0$, of $(9.2^{+20.4}_{-6.4})\cdot10^{-8}$ Pa at 298.15 K and an enthalpy of evaporation, $\Delta H_{\text{vap}}(298K)$, of $130.2\pm9.9$ kJ·mol$^{-1}$.

A summary of the derived saturation vapor pressures and enthalpies of vaporization are given in Table 2.

Looking at all data it is evident that the high temperature literature data agree quite well with our data at about room temperature. Hence, extrapolation of experiments performed at elevated temperatures toward atmospheric temperatures is possible with high accuracy as long as the temperature trend is measured accurately enough. More quantitatively, extrapolation of the high temperature literature data to 298.15 K incorporating the temperature dependence of the enthalpy of vaporization as
Table 2. Saturation vapor pressures at 298.15 K and enthalpies of vaporization of the polyethylene glycols measured in this study as well as estimated enthalpies of vaporization at 298.15 K, together with the results of the quantum chemistry calculations. In addition we provide experimental data from the literature for ethylene glycol and diethylene glycol.

<table>
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<th>n</th>
<th>p(exp)</th>
<th>$\Delta H_{\text{vap}}$(exp, $T$)</th>
<th>$T$</th>
<th>$\Delta H_{\text{vap}}$(exp, 298 K)</th>
<th>p(calc)</th>
<th>$\Delta H_{\text{vap}}$(calc, 298 K)</th>
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<td></td>
<td>[Pa]</td>
<td>[kJ · mol$^{-1}$]</td>
<td></td>
<td>[kJ · mol$^{-1}$]</td>
<td>[Pa]</td>
<td>[kJ · mol$^{-1}$]</td>
</tr>
<tr>
<td>triethylene glycol</td>
<td>3 (6.68$^{+1.10}_{-0.95}$) · $10^{-2}$</td>
<td>78.3±0.7</td>
<td>349</td>
<td>84.3±1.9</td>
<td>2.65·10$^{0}$</td>
<td>68.81</td>
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<tr>
<td>tetraethylene glycol</td>
<td>4 (1.69$^{+0.11}_{-0.10}$) · $10^{-2}$</td>
<td>77.1±0.4</td>
<td>345</td>
<td>82.1±1.6</td>
<td>1.82·10$^{-1}$</td>
<td>72.78</td>
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<tr>
<td>pentaethylene glycol</td>
<td>5 (5.38$^{+0.71}_{-0.63}$) · $10^{-4}$</td>
<td>90.6±1.1</td>
<td>339</td>
<td>94.6±1.6</td>
<td>2.10·10$^{-2}$</td>
<td>81.60</td>
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<tr>
<td>hexaethylene glycol</td>
<td>6 (3.15$^{+0.58}_{-0.49}$) · $10^{-5}$</td>
<td>102.1±1.5</td>
<td>351</td>
<td>107.2±2.1</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>heptaethylene glycol</td>
<td>7 (1.38$^{+0.47}_{-0.35}$) · $10^{-6}$</td>
<td>113.7±2.7</td>
<td>357</td>
<td>119.2±3.2</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>octaethylene glycol</td>
<td>8 (9.2$^{+0.35}_{-0.49}$) · $10^{-8}$</td>
<td>124.2±9.7</td>
<td>363</td>
<td>130.2±9.9</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>ethylene glycol</td>
<td>1 11.7±0.6$^a$</td>
<td>–</td>
<td>–</td>
<td>65.6±0.3$^b$</td>
<td>3.98·10$^1$</td>
<td>60.75</td>
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<td>diethylene glycol</td>
<td>2 0.6±0.03$^a$</td>
<td>66.9±0.3$^b$</td>
<td>420</td>
<td>80.2±4.0$^b$</td>
<td>3.80·10$^0$</td>
<td>67.41</td>
</tr>
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</table>


detailed in Appendix B (orange filled circles in Fig. 8) yields values agreeing well within error with the newly derived saturation vapor pressures for PEG3, PEG7 and PEG8 of this study and agree better than within a factor of two for PEG4, PEG5 and PEG6. In contrast, using a constant enthalpy of vaporization for the extrapolation yields very good agreement for PEG4, PEG5 and PEG6, but substantial deviation - up to a factor of 10 - for PEG3, PEG7 and PEG8. We conclude that extrapolation to lower temperatures of high temperature data should take the temperature dependence of the enthalpy of vaporization (Kirchhoff’s law) into account. Also, it helps tremendously when data cover a large temperature range. It is obvious that each technique used in our study has an optimal pressure range for obtaining high quality data. The optimal pressure range should be utilized in the future by rather performing measurements at elevated temperatures with extrapolation to atmospheric temperatures and avoid pressure ranges where the accuracy of the derived saturation vapor pressures is limited.

In addition, the good agreement between the FT-TDMA data and all other techniques points towards the accommodation coefficient being close to one as this was assumed for the data analysis. To retrieve a lower limit for the accommodation coefficient, we analyzed the FT-TDMA experiments varying the accommodation coefficient until the calculated vapor pressure including its error exceeds the reference pressure confidence interval given in Table 2 at the temperature of each FT-DMA experiment. The mean value of these accommodation coefficients for all experiments is equal to 0.5 and there is no discernible difference between the experiments for different PEGs. We conclude that the mass accommodation coefficient of the PEGs based on our study is greater than 0.5.

In Fig. 8 we plot the saturation vapor pressures on a logarithmic scale at 298.15 K versus the number of PEG units. The saturation pressures show an exponential dependence on the number of PEG units with a reduction in pressure by a factor of 13.8 per PEG unit added, as indicated by the linear regression (dashed gray line) over the complete range of PEGs in the figure. Closer inspection reveals that there is a change in slope between PEG3 and PEG4. In particular the tetraethylene glycol...
Figure 8. Saturation vapor pressure at 298.15 K plotted for all PEGs. Open black circles with error bars: results of this study. Filled orange circles: extrapolation of high temperature literature data to 298.15 K using an estimation of the temperature dependence of the enthalpy of evaporation as detailed in appendix B. Open orange circles: extrapolation of high temperature literature data to 298.15 K assuming a constant enthalpy of evaporation. Filled black circles: computational predictions from this study. Open squares with error bars: Ambrose and Hall (1981). Dashed gray line is a linear regression to all the experimental data, dashed red lines are separate linear regressions for n=1 to n=3 and n=4 to n=8.

deviates from the linear relationship by a factor of three to higher pressures, out of the 95% confidence interval uncertainty of the derived vapor pressures. As the measurements of this compound is particularly well constrained by numerous data not only from this study, see Fig. 3, we conclude that this deviation is real. Since there is no fundamental reason for the homologous series to follow a strict exponential behavior, it is of interest whether computational prediction methods are able to reproduce this behavior. In Fig. 8 we also plot the results of the COSMOtherm calculations given in Table 2. Clearly, these calculations overpredict the vapor pressure by a factor of 3–40. Interestingly, they show a similar deviation from the linear trend as the experimental data, but here the change in slope occurs at one PEG-unit lower than what is observed in the experimental data.

Separate linear regressions for ethylene glycol to PEG3 and for PEG4 to PEG8 yield regression lines within the 95% confidence interval of the individual data for each PEG (red dashed lines in Fig. 8). Hence, we predict a saturation vapor pressure for nonaethylene glycol with some confidence as \((3.7^{\pm 4.1 \times 10^{-9}})\) Pa at 298.15 K. All our instruments were not able to measure a vapor pressure that low.
In the plot of the enthalpies at 298.15 K for the homologous series in Fig. 9 there is a general trend for an increase in enthalpy of vaporization with number of PEG units, but a strict linear dependence is observed only from the PEG4 to higher PEGs, with an average 12.3 kJ·mol\(^{-1}\) increase per PEG unit. The lower molecular weight PEGs show a more irregular behavior, while the COSMOherm calculations reveal again a change in slope between diethylene glycol and PEG4 while the increase for ethylene glycol to diethylene glycol and from PEG4 to PEG5 is similar. Again, this may be an indication that there is a transition occurring in the homologous series at triethylene glycol. Note that the transition does not need to be related to changes in the condensed phase but might also be related to the gas phase, e.g. internal hydrogen bonds that stabilize larger oligomers.

5 Conclusions

A reference data set for validating vapor pressure measurement techniques is provided, spanning a range in pressure at room temperature from \(5 \times 10^{-2}\) Pa to \(10^{-7}\) Pa based on experimental data and extending to \(10^{-9}\) Pa with high confidence based on the observed trend in the homologous series of polyethylene glycols. Our data reveal clearly that (at least for the compounds studied) extrapolations to lower temperatures are possible as long as the temperature dependence is correctly measured. Therefore, it seems to be more appropriate to perform measurements at (slightly) elevated temperatures and pressures with high accuracy and then extrapolate to lower temperatures compared to measuring at lower temperature with low accuracy. Each
of our experimental setups has a pressure range in which high quality data can be obtained. Future measurements should be
done preferably in the appropriate pressure range. We suggest to use our dataset of the homologous series of polyethylene
glycols to determine the lower detection limit of saturation vapor pressures for each experimental setup, to use it for instru-
ment calibration, for estimating systematic errors in experimental setups and for comparison with vapor pressure estimation
methods.

Appendix A: Data treatment of EDB data taken at elevated relative humidities

Evaporation rates in the EDB setups were measured not only under dry conditions but some also at elevated relative humidities.
After the particle was equilibrated with respect to temperature and RH, any change in size, $\frac{dr^2}{dt}$, was attributed to evaporative
loss at a constant composition, and the partial pressure of the PEG, $p_{\text{PEG}}$, under those conditions was calculated using Eq. (1)
of Soonsin et al. (2010):

$$p_{\text{PEG}} = -\frac{1}{2} \frac{dr^2}{dt} \frac{x \rho R T}{(x M_{\text{PEG}} + (1-x) M_{\text{H}_2\text{O}}) D}, \quad (A1)$$

where $r$ is the particle radius, $x$ is the mole fraction of PEG, $\rho$ is the density of the condensed phase, $R$ is the gas constant, $M$
is molar mass, and $D$ is the diffusivity of the PEG vapor in the buffer atmosphere. The mole fraction of PEG in an aqueous particle
was taken from the RH in the EDB (see Fig. 1 of the manuscript) by assuming that the particle water activity is equivalent to
measured RH, i.e., that the particle is homogeneous and in thermodynamic equilibrium (which is a good approximation as long
as evaporation is sufficiently slow). Density was estimated by ideal mixing. As an example we show data obtained by the EDB
setup of Union College at 286.5 K for PEG4 in Fig. A1.

Extrapolation of the data to a PEG activity equal one yields the saturation pressure of PEG at this temperature.

Appendix B: Change in evaporation enthalpy with temperature

Generally, the enthalpy of evaporation is a function of temperature. Let us analyze the data of PEG3 as an example following
Chickos and Acree Jr. (2003). If the heat capacities of the liquid and gas phase are known, $C_{p_l}$ and $C_{p_g}$, respectively, then
the vaporization enthalpy, $\Delta H_{\text{vap}}(T^\circ)$, at $T^\circ = 298.15$ K can be related to the experimental measurements by using Eq. B1,
generally referred to as Kirchhoff’s equation.

$$\Delta H_{\text{vap}}(T^\circ) = \Delta H_{\text{vap}}(\bar{T}) + \int_{T^\circ}^{\bar{T}} (C_{p_l} - C_{p_g}) dT, \quad (B1)$$

where $\bar{T}$ refers to the mean temperature of all measurements, which is equal to 357 K for our PEG3 data when calculating
the mean of the inverse temperature range covered by the data. If we estimate $(C_{p_l} - C_{p_g})$ by using the method suggested by
Chickos and Acree Jr. (2003) (which should be accurate to about $\pm 30$ J mol$^{-1}$K$^{-1}$):

$$(C_{p_l} - C_{p_g}) = (10.56 + 0.26C_{p_l}), \quad (B2)$$
Figure A1. Partial pressures of PEG4 versus PEG4 activity deduced from evaporation rates of single, levitated, aqueous PEG4 particles in the EDB setup of Union College at a temperature of 286.5 K. Black symbols: experiment data, red line and shaded area: linear fit to these data with the 95% confidence interval. Extrapolation to \( a_{\text{PEG}} \) equal one yields the saturation vapor pressure at this temperature, i.e. \( p^0 = (5.69 \pm 0.55) \cdot 10^{-3} \text{Pa} \) at this temperature.

it yields 101±30 J · mol\(^{-1}\)K\(^{-1}\) for the difference between heat capacity of the liquid and heat capacity of the gas phase, when we take the value for the heat capacities of the liquid phase, \( C_{p_l} \), from Stephens and Tamplin (1979). Using Eq. B1 with this average value for \( (C_{p_l} - C_{p_g}) \) yields an increase to the uncorrected enthalpy of vaporization of 6.0±1.8 kJ · mol\(^{-1}\) when using the temperature difference between mean temperature, \( \overline{T} \), and 298.15 K. Therefore, the enthalpy at the mean temperature of the measurements obtained by the linear regression, which is \( \Delta H_{\text{vap}}(\text{exp, } T) = 78.3\pm0.7 \text{kJ · mol}^{-1} \), should be corrected to \( \Delta H_{\text{vap}} = 84.3\pm1.9 \text{kJ · mol}^{-1} \) at 298.15 K. This corresponds to a shift of 7.5 %, which is clearly non-negligible. The magnitude is somewhat smaller for the higher mass PEGs. In Table 2 we provide the enthalpies at both temperature, \( T \) and \( T^\circ \).

Now we may integrate the Clausius-Clapeyron equation, eq. (B3) in a more general form, by including Kirchhoff’s equation, eq. (B1), in the same manner as above, namely assuming a constant, temperature independent, heat capacity difference between liquid and gas phase.

\[
\frac{dp}{p} = \frac{\Delta H_{\text{vap}}(T)}{R} \frac{dT}{T^2}, \text{ with } \Delta H_{\text{vap}}(T) = \Delta H_{\text{vap}}(T^\circ) - (C_{p_l} - C_{p_g})(T^\circ - T) \tag{B3}
\]

Integration of eq. (B3) leads to:

\[
\ln \frac{p}{p^\circ} = -\frac{\Delta H_{\text{vap}}(T^\circ) - T^\circ(C_{p_l} - C_{p_g})}{R} \left( \frac{1}{T} - \frac{1}{T^\circ} \right) + \frac{(C_{p_l} - C_{p_g})}{R} \ln \left( \frac{T}{T^\circ} \right). \tag{B4}
\]
If we now plot $\Psi := \left\{ R \ln \left( \frac{p}{p_\infty} \right) - (C_{p_l} - C_{p_g}) \ln \frac{T}{T_\infty} \right\}$ versus $1/T - 1/T^\circ$, we expect a straight line with $\Psi(0) = 0$. In Fig. A2 we show such a plot for the PEG3 data.

Clearly, considering the temperature dependence of the enthalpy of evaporation helps to reconcile the high temperature data with those taken by us at roughly room temperatures. However, its effect close to 298 K is minimal by default if data at room temperature are available, see eq. B4.

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