Determining noble gas partitioning within a CO$_2$–H$_2$O system at elevated temperatures and pressures

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Abstract

Quantifying the distribution of noble gases between phases is essential for using these inert trace gases to track the processes controlling multi-phase subsurface systems. Here we present experimental data that defines noble gas partitioning for two phase CO$_2$–water systems. These are at the pressure and temperature range relevant for engineered systems used for anthropogenic carbon capture and geological storage (CCS) technologies, and CO$_2$-rich natural gas reservoirs (CO$_2$ density range 169–656 kg/m$^3$ at 323–377 K and 89–134 bar). The new partitioning data are compared to predictions of noble gas partitioning determined in low-pressure, pure noble gas–water systems for all noble gases except neon and radon. At low CO$_2$ density there was no difference between measured noble gas partitioning and that predicted in pure noble gas–water systems. At high CO$_2$ density, however, partition coefficients express significant deviation from pure noble gas–water systems. At 656 kg/m$^3$, these deviations are $-35\%$, $74\%$, $113\%$ and $319\%$ for helium, argon, krypton and xenon, respectively. A second order polynomial fit to the data for each noble gas describes the deviation from the pure noble gas–water system as a function of CO$_2$ density. We argue that the difference between pure noble gas–water systems and the high density CO$_2$–water system is due to an enhanced degree of molecular interactions occurring within the dense CO$_2$ phase due to the combined effect of inductive and dispersive forces acting on the noble gases. As the magnitude of these forces are related to the size and polarisability of each noble gas, xenon followed by krypton and argon become significantly more soluble within dense CO$_2$. In the case of helium repulsive forces dominate and so it becomes less soluble as a function of CO$_2$ density.

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1. INTRODUCTION

Noble gases are used as geochemical tracers to investigate a wide range of key geological systems (Porcelli et al., 2002; Burnard, 2013). Their inert natures prevent these noble gas tracers from chemically interacting with their surroundings. Thus their distribution within a geological system is a result of physical interaction between solids and different fluid phases. Consequently they can be used to interpret the physical processes which are often poorly constrained by other, more reactive, tracers. For example noble gases have proven their potential in topics as diverse as understanding the source, migration and sinks of fluids throughout the crust (e.g., Ballentine et al., 2002;
Sherwood-Lollar and Ballentine, 2009; Brennwald et al., 2013; Pinti et al., 2013) through to aiding our understanding of the origin and evolution of our planet (e.g., Holland et al., 2009; Marty, 2012; Halliday, 2013).

Noble gas utility is intrinsically linked to an understanding of their physical chemistry within the geological systems of interest. For systems where noble gas partitioning is undefined, estimations are typically made by approximating to experimentally determined low pressure systems which tend towards ideal gas behaviour. Although this approach provides a viable short term resolution, this creates uncertainties which restrict the usefulness of quantitative interpretations. The supercritical CO$_2$–water binary phase system in geological systems is a prime example as this can occur naturally (e.g., Ballentine et al., 2001; Ballentine and Holland, 2008) or anthropogenically for Enhanced Oil Recovery (EOR) (e.g., Blunt et al., 1993; Ravagnani et al., 2009). Additionally storage of anthropogenic CO$_2$ in geological structures as part of national CCS (Carbon Capture and Storage) strategies has further stimulated much recent interest in supercritical CO$_2$–water systems (Kharkaka et al., 2006; Bickle et al., 2007; Cole et al., 2010). With this focus noble gases have been used extensively to understand the processes controlling CO$_2$ behaviour in the subsurface (Gilfillan et al., 2008, 2009; Dubaçq et al., 2012; Zhou et al., 2012; Kampman et al., 2013) and they offer the potential to trace in-reservoir processes and subsurface leakage from engineered systems (Nimz and Hudson, 2005; Mackintosh and Ballentine, 2012).

Partitioning of noble gas $i$ within a CO$_2$–water system is described using Henry’s law:

$$\Phi_i p_i = \gamma_i K_i x_i$$

where $p_i$ is the partial gas pressure of the noble gas, $\Phi_i$ is the fugacity, $K_i$ is Henry’s constant at a given temperature ($T$), $x_i$ is the molar fraction of $i$ within the liquid phase and $\gamma_i$ is the activity coefficient. By combining system-specific fugacities and activity coefficients with the Henry’s constant, Henry’s coefficients ($k_i$) for noble gases at trace levels can be determined for the gas–liquid systems of interest:

$$k_i = \frac{\gamma_i K_i}{\Phi_i}$$

To date, water–CO$_2$ noble gas partitioning calculations within natural and engineered geological systems have been based on Henry’s coefficients determined from noble gas–water experiments at low pressures (Crovetto et al., 1982; Smith and Kennedy, 1983; Smith, 1985) with the implicit assumption that a dense carbon dioxide phase will have no effect on noble gas fugacity. Additionally both the effect of aqueous CO$_2$ and pressure acting on the water phase are also assumed to have a negligible effect on the activity coefficient (Ballentine et al., 2002).

We assess here the effect of a dense CO$_2$ phase on Henry’s coefficients by experimentally determining noble gas partitioning within high-pressure high-temperature supercritical carbon dioxide–water systems. These are compared with Henry’s coefficients derived from the low pressure data sets (Crovetto et al., 1982; Smith, 1985). We discuss the physical causes for observed differences and we also provide a correlation that allows the calculation of Henry’s coefficients for noble gases in the presence of carbon dioxide.

### 2. EXPERIMENTAL METHODS

#### 2.1. Rationale

At optimal conditions for CCS storage, CO$_2$ densities typically range between 600 and 800 kg/m$^3$ (Bachu, 2003; IPCC, 2005; Friedmann, 2007). Where natural CO$_2$ systems are involved, assuming hydrostatic pressure, CO$_2$ densities range between 200 and 850 kg/m$^3$ (Bachu, 2003; Gilfillan et al., 2008). In this work a CO$_2$ density range between 169 and 656 kg/m$^3$ was selected to represent a wide variety of natural and anthropogenic crustal CO$_2$–water systems. This density range was achieved using a temperature and pressure grid which ranged between 323–377 K and 89–134 bar, respectively. The CO$_2$ density was thermodynamically derived via the NIST website (NIST, 2014). These are presented graphically on a CO$_2$ phase diagram (Fig. 1) and numerically (Table 1).

Binary phase CO$_2$–water systems, enriched with a noble gas spike, were created at the British Geological Survey (BGS) in Keyworth. Aliquots of each phase were subsequently sampled, sealed in pressurised containers, and transported to the School of Earth, Atmospheric and Environmental Sciences (SEAES), University of Manchester, where the respective noble gas components were isolated and their concentrations measured (Warr, 2013).

#### 2.2. Sample generation

Samples were generated using a stainless steel pressurised batch reactor (Fig. 2). The reactor itself is initially

![Fig. 1. Figure plotting the pressure and temperature of the experimental charges in this study and displayed in the context of the CO$_2$ phase diagram. All experimental data points were collected within the supercritical regime and are given in Table 1. Low pressure verification data point within gaseous CO$_2$ phase (Experiment 0) has also been included for reference.]
charged with 150 cm$^3$ of deionised water and a magnetic stirrer bead was added and the system placed within an oven.

Three 1/8” Swagelok tube connections allow CO$_2$ injection and sampling, and the remaining two to act as inlet/outlet ports for water phase cycling and sampling. The oven was heated to the experiment temperature and laboratory grade CO$_2$ used to displace the air within the batch reactor. The sampler valve was closed and the headspace pressurised to the experiment pressure using an ISCO 260D syringe pump. A 1.5 cm$^3$ noble gas spike at ~0.6 bar containing 32% helium, 32% neon, 17% argon, 11% krypton, and 8% xenon was added to the headspace. After initial pressurisation the system was isolated from the CO$_2$ pump and the magnetic stirrer was turned on. An external Jasco PU-2086 HPLC pump circulated the water phase through the water samplers at 1 cm$^3$/min. During this time, the pressure of the system was logged to ensure the system had no significant loss of CO$_2$. This was used as an indication that a closed system had been attained. Samples of the CO$_2$ and water phases were collected in triplicate.

Prior to sampling, experimental temperatures and pressures were recorded from a calibrated platinum-resistance thermometer connected to an Isotech calibration unit thermometer and an ESI Technology transducer that had been checked against a second, externally-calibrated pressure transducer. For the water and CO$_2$ 3 cm$^3$ aliquots were sampled respectively using 1/8” stainless steel Swagelok needle valves connected in series. For simplicity only one sampler for each phase is depicted here.

To sample the water phase the magnetic stirrer and HPLC pump were stopped and the valves on the water samplers were closed sequentially in order of proximity to the pump, starting with the closest. For the CO$_2$ phase the samplers, initially filled with air at atmospheric pressure, were connected in series with all valves closed to the batch reactor at the CO$_2$ inlet/outlet valve (Fig. 2). To displace the air in the samplers ~120 cm$^3$ of CO$_2$ at atmospheric pressure and laboratory temperature (19°C) (i.e., ~20× the total sampler and buffer volumes) was vented through the final sampler. All valves were then closed in order of furthest to nearest from the oven and the samplers removed. Due to the pressures involved and sampling at room temperature the CO$_2$ was sampled as a liquid phase. As noble gases exist in uniform trace amounts within the CO$_2$ phase the ratio of noble gas to CO$_2$ is independent to the phase of collection within a closed system.

### 2.3. Sample analysis

CO$_2$ and water phase samples were each prepared and analysed in an all metal UHV system using a Hiden Analytical 1–200 AMU (HAL200) quadrupole mass spectrometer (Fig. 3). Through a process of expansions, dilutions and chemical reactions the noble gas component was isolated prior to measurement (Warr, 2013).

All samples were connected to the analytical line using 1/8” stainless steel Swagelok fittings (Fig. 3). After standard operating pressures were reached in the quadrupole volume (6 × 10$^{-11}$ bar) the CO$_2$ sample buffer volume was evacuated using the roughing pump. The buffer valve was closed and the system was again pumped down to reach operating pressure. To check for leakage on mounting each sample the pressure was monitored by opening to the closed volume containing the 0–13.33 bar Baratron$^\text{®}$ after reopening the buffer volume. Negligible deflection on the Baratron$^\text{®}$ for all samples indicated a contribution of <0.03% to all measured noble gases due to leakage from the sample connection, with full procedural blanks confirming negligible noble gas contributions from sources other than the sample aliquots.
The CO₂ sample was expanded to the cleanup section and left to equilibrate for 2 min. Aliquot A1 (0.81 cm³) was used to reduce the volume of CO₂ and the remainder of the vacuum line opened to the pump until operating pressures were once again reached. The CO₂ aliquot was exposed to a hot SAES NP10 getter for 10 min to remove all active gases via chemical reaction before being expanded to the quadrupole analyser for noble gas determination. The noble gas isotopes measured are listed in Table 2.

The noble gas content of each sample was measured for 10 min after which the sample was pumped away. In addition, a baseline reading for each isotope was collected for 10 min before and after sample analysis.

Water sample preparation involved a two stage cryogenic approach to remove the water phase. The water sample was released into the underlying stainless steel sample receptacle where it was degassed using an ultrasonic bath to ensure that the helium and neon occupied the headspace. The receptacle containing the water was then cooled with liquid nitrogen leaving helium and neon and >5% of argon in the headspace. The headspace was expanded into the clean-up section and the Ampule aliquot (Fig. 3) was used to reduce the volume by a factor of 0.014. The clean-up section was then pumped to standard operating pressures after which the Ampule aliquot was reopened to the cleanup section and exposed to the hot SAES NP10 getter prior to analysis.

Argon, krypton and xenon were released into the headspace by warming the sample receptacle to room temperature followed by ultrasonic degassing. The sample receptacle was then immersed in an acetone slush (178 K) to re-freeze the water. The heavy noble gases, now in the headspace, followed an identical procedure as described for helium and neon. Post-degassing redissolution resulted in a maximum, negligible (<0.2%) reduction in headspace values.

2.4. Data analysis

System stability and linearity are provided in Appendix 2. Initial raw signal intensities for each isotope were determined for the CO₂ and water phases ($C_{i(raw)}$ and $W_{i(raw)}$, respectively). A baseline measurement ($C_{i(back)}$ and $W_{i(back)}$) was also measured which was then subtracted from these raw signal intensities. Procedural blanks indicated no discernible background above baseline values and therefore the baseline correction alone was applied. Next, a series of corrections were applied to negate measured differences in both sample and analytical aliquot sizes, as well as increases to sample density due to sampling both experimental phases at room temperature. Consequently signals were standardised for each noble gas isotope ($i$) for the CO₂ and water phases (at pressures ($P$) and temperatures ($T$) ($C_{i(P,T)}$ and $W_{i(P,T)}$, respectively). These corrected, dimensionless, noble gas peak intensities of each phase were then used to generate a phase ratio for each noble gas isotope:

$$D_{i(P,T)} = \frac{C_{i(P,T)}}{W_{i(P,T)}}$$ (3)

where $D_{i(P,T)}$ is the dimensionless partitioning ratio of noble gas isotope $i$ between each phase at a given pressure ($P$) and temperature ($T$).
The partitioning ratios were determined three times in each experiment. An arithmetic mean of these three data points was used to generate partitioning ratios for each isotope. Through comparison with predicted partitioning based on pure noble gas–water Henry’s coefficients (Crovetto et al., 1982; Smith, 1985) Henry’s coefficients were obtained (see Table 4). The method for deriving the uncertainty associated with each data point is provided in Appendix 1.

2.5. Experimental verification

In order to verify the experimental procedure, experiments were conducted at ambient temperature and pressure (Experiment 0, Table 1). Under these conditions Henry’s coefficients in a CO2–water system are expected to be indistinguishable from pure noble gas–water values.

Ambient pressure and temperature results are presented in Table 3. For both argon and krypton the pure noble gas–water values were experimentally reproduced within 1σ uncertainty. Although xenon partitioning was slightly over-predicted the pure noble gas–water values are within 2σ (95% confidence) of the experimental values. Uncertainty in the helium determination was significantly higher due to the low solubility of helium in water resulting in a yet smaller signal size. Helium partitioning values are nevertheless within 2σ of predicted values. Low signal size was a concern and these experiments resulted in an increase in the amount of noble gas being added into the system during initial pressurisation of the system (see Section 2.2) to ensure helium was present in greater abundances resulting in lower, acceptable, uncertainties (see Table 6).

3. RESULTS

The experimentally measured Henry’s coefficients for each noble gas are given in Table 4. All values are presented here as deviation from pure noble gas water Henry’s coefficients as well as absolute values. The pure noble gas–water values are reproduced in Table 5 and experimental deviation from predicted values are given in Table 6.

3.1. Quantifying density-deviation trends

Physical properties of CO2 such as density, enthalpy and viscosity do not deviate from their current trends when migrating across the supercritical–subcritical “boundary” (Span and Wagner, 1996; NIST, 2014). It is reasonable to assume noble gas trends are similarly unaffected across the supercritical and subcritical CO2 transition. As such, any deviations in noble gas partitioning behaviour from low pressure behaviour are reasonably controlled by increased molecular interactions and related to density. Indeed, for all noble gases, deviations from pure noble gas–water systems as a function of CO2 density are observed. This deviation trend is negative for helium and positive for argon, krypton and xenon (Table 6, Fig. 4).

### Table 3
Comparing experimental Henry’s coefficients with their pure noble gas–water counterparts under ambient conditions. All Henry’s coefficients are given in GPa to 2 decimal places. Ratios were derived from 4He, 40Ar, 82Kr, 84Kr, 132Xe & 136Xe. For reasons outlined in Section 3.5 Henry’s coefficients for neon could not be determined. Noble gas–water values are taken from Crovetto et al. (1982) and Smith (1985). Uncertainties for noble gas–water values are calculated using values in Table 7 with Eq. (4). All uncertainty is given as 1σ confidence.

| Noble gas | Noble gas–water Henry’s coefficient (GPa) | Experimental Henry’s coefficient (GPa) |
|-----------|------------------------------------------|---------------------------------------|
| Helium    | 14.59 ± 1.38                             | 8.28 ± 4.81                           |
| Argon     | 4.02 ± 0.49                              | 3.77 ± 0.59                           |
| Krypton   | 2.22 ± 0.21                              | 2.14 ± 0.22                           |
| Xenon     | 1.31 ± 0.21                              | 1.89 ± 0.23                           |

### Table 4
Experimentally derived Henry’s coefficients for all noble gases. Values are given in GPa and are quoted to 2 decimal places. Uncertainty is given as 1σ confidence. *Indicates no viable partition coefficient was generated.

| Experiment | Helium (GPa) | Neon (GPa) | Argon (GPa) | Krypton (GPa) | Xenon (GPa) |
|------------|--------------|------------|-------------|---------------|-------------|
| 0          | 8.28 ± 4.81  | *          | 3.77 ± 0.59 | 2.14 ± 0.22   | 1.89 ± 0.23 |
| 1          | 13.2 ± 2.35  | 18.82 ± 6.91 | 6.02 ± 0.87  | 4.16 ± 0.38   | 3.69 ± 0.23 |
| 2          | 11.39 ± 2.01 | 23.48 ± 28.5 | 8.07 ± 1.18  | 7.23 ± 0.68   | 8.44 ± 0.64 |
| 3          | 13.55 ± 2.38 | 19.49 ± 17.66 | 4.58 ± 0.72  | 6.06 ± 0.6    | 7.90 ± 1.04 |
| 4          | 6.73 ± 1.19  | 16.42 ± 20.98 | 9.81 ± 1.43  | 6.93 ± 0.65   | 8.64 ± 0.68 |
| 5          | 13.76 ± 2.44 | 22.77 ± 8.13 | 8.46 ± 1.26  | 5.17 ± 0.47   | 4.19 ± 0.26 |
| 6          | 10.19 ± 1.81 | 19.47 ± 22.69 | 6.46 ± 0.96  | 5.01 ± 0.46   | 5.42 ± 0.4  |
| 7          | 13.05 ± 2.29 | 22.06 ± 10.25 | 6.94 ± 1.08  | 5.86 ± 0.53   | 6.76 ± 0.43 |
| 8          | 9.64 ± 1.71  | 8.28 ± 5.69  | 7.96 ± 1.27  | 5.27 ± 0.48   | 4.92 ± 0.35 |
| 9          | 12.08 ± 2.13 | 22.02 ± 17.07 | 5.50 ± 0.83  | 5.27 ± 0.49   | 4.92 ± 0.32 |
| 10         | 12.72 ± 2.24 | 22.61 ± 13.86 | 5.98 ± 0.93  | 5.47 ± 0.51   | 5.76 ± 0.38 |
Table 5
Pure noble gas–water Henry’s coefficients for all noble gases. Helium values taken from Smith (1985) and neon, argon, krypton and xenon values taken from Crovetto et al. (1982). Values are given in GPa and are quoted to 2 decimal places.

| Experiment | Helium (GPa) | Neon (GPa) | Argon (GPa) | Krypton (GPa) | Xenon (GPa) |
|------------|--------------|------------|-------------|---------------|-------------|
| 1          | 14.50        | 13.49      | 5.57        | 3.36          | 2.21        |
| 2          | 14.50        | 13.49      | 5.56        | 3.35          | 2.21        |
| 3          | 14.49        | 13.50      | 5.58        | 3.37          | 2.21        |
| 4          | 14.50        | 13.49      | 5.56        | 3.35          | 2.21        |
| 5          | 13.92        | 13.43      | 6.09        | 3.80          | 2.59        |
| 6          | 13.19        | 13.10      | 6.39        | 4.11          | 2.86        |
| 7          | 13.16        | 13.09      | 6.39        | 4.12          | 2.87        |
| 8          | 11.18        | 11.79      | 6.49        | 4.43          | 3.18        |
| 9          | 10.93        | 11.61      | 6.46        | 4.44          | 3.19        |
| 10         | 11.26        | 11.85      | 6.49        | 4.43          | 3.17        |

Table 6
Experimental Henry’s coefficients displayed as percentage deviations from values calculated for a pure noble gas water system (Table 5). Negative values indicate lower Henry’s coefficients than predicted i.e., a greater solubility within the water phase (Eq. 1) and vice versa. All values are given to 2 decimal places. *Indicates no viable partition coefficient was generated.

| Exp. | Helium (%) | Neon (%) | Argon (%) | Krypton (%) | Xenon (%) |
|------|------------|----------|-----------|-------------|-----------|
| 0    | –43.24 ± 58.11 | *        | –6.24 ± 15.77 | –3.97 ± 10.15 | 44.48 ± 11.97 |
| 1    | –8.98 ± 17.79  | 39.46 ± 36.74 | 8.15 ± 14.52  | 23.93 ± 9.06  | 66.83 ± 6.20  |
| 2    | –21.48 ± 17.68 | 74.04 ± 121.37 | 45.14 ± 14.66 | 115.87 ± 9.33 | 282.36 ± 7.59 |
| 3    | –6.48 ± 17.55  | 44.44 ± 90.60  | –17.77 ± 15.71 | 80.19 ± 9.81  | 255.91 ± 13.20 |
| 4    | –53.63 ± 17.65 | 21.71 ± 127.72 | 76.51 ± 14.56 | 106.84 ± 9.41 | 291.60 ± 7.83 |
| 5    | –1.18 ± 17.76  | 69.57 ± 35.71  | 39.00 ± 14.83  | 36.01 ± 9.12  | 61.65 ± 6.29  |
| 6    | –22.76 ± 17.74 | 48.59 ± 116.53 | 1.24 ± 14.88   | 21.87 ± 9.24   | 89.39 ± 7.42  |
| 7    | –0.83 ± 17.57  | 68.59 ± 46.46  | 8.53 ± 15.61   | 42.40 ± 9.12   | 135.56 ± 6.40 |
| 8    | –13.72 ± 17.70 | –29.76 ± 68.70 | 22.76 ± 15.91  | 18.83 ± 9.17   | 54.79 ± 7.15  |
| 9    | 10.52 ± 17.61  | 89.73 ± 77.54  | –14.87 ± 15.06 | 18.71 ± 9.21   | 53.91 ± 6.44  |
| 10   | 12.96 ± 17.61  | 90.85 ± 61.31  | –7.87 ± 15.55  | 23.52 ± 9.37   | 81.35 ± 6.55  |

Without a detailed mechanistic understanding of the role of CO₂ density in determining the noble gas data any fit of the data to a function cannot be used to extrapolate the data beyond the physical conditions of these experiments. Nevertheless, such a fit provides a pragmatic tool for correcting noble gas partition data over the range of densities covered by this work.

At very low densities ideal gas behaviour can reasonably be assumed, and is demonstrated during our experimental validation. Consequently a full reconciliation is expected with pure noble gas–water systems where as density → 0; deviation from pure noble gas–water values → 0. For each of the noble gases an error weighted second-order polynomial fit line which passes through the origin is has been determined for the form:

\[ y = ax^2 + bx + c \]  

where \( x \) is the density, \( a, b \) and \( c \) are coefficients (with \( c \) fixed at 0) and \( y \) is percentage deviation. We also determined the ±1σ confidence envelope for each of the polynomial fits with associated \( a, b \) and \( c \) coefficients. Accordingly a single equation is provided which characterises the deviation in Henry’s coefficient from a low pressure water–gas system for each noble gas for the 0–656 kg/m³ CO₂ density range (Table 7).

The polynomial fit can be very simply used to directly derive the deviation coefficient from a low pressure water–gas system by applying the equation:

\[ \kappa_i = \frac{(a_i x^2 + b_i x + c_i)}{100} + 1 \]  

where \( a, b \) and \( c \) are the fitted coefficients for noble gas \( i \) (Table 7), \( x \) is the CO₂ density in kg/m³ and \( a_i x^2 + b_i x + c_i \) gives the percentage deviation from pure noble gas–water values. Using this deviation coefficient, Henry’s coefficients for a pure noble gas–water system \( (k_i^{(pub)}) \), taken from Crovetto et al. (1982) and Smith (1985), can be recalculated for application to dense CO₂-rich systems \( (k_i^{(CO2)}) \):

\[ k_i^{(CO2)} = k_i^{(pub)} \]  

3.2. Xenon

Experimental Henry’s coefficients were all significantly larger than predicted with values increased by between 53.9% and 291.6%. From Fig. 4A, a clear, positive CO₂ density-deviation trend from pure noble gas–water partitioning was observed. A high \( R^2 \) (0.98) indicates that the polynomial can be used with reasonable confidence to predict the CO₂ density-deviation relationship within the
experimental density range. Using this fit, Henry’s coefficients reconcile with those for a pure noble gas–water system (≤5% deviation) at CO₂ densities lower than 28 kg/m³. This polynomial is considered to suitably define deviation from a pure noble gas–water system as a function of CO₂ density for the 0–656 kg/m³ range.

3.3. Krypton

All experimentally-defined Henry’s coefficients exceed those predicted for pure noble gas–water systems, with deviations ranging from 18.7% to 115.9% at the highest density. By plotting deviation as a function of density a positive CO₂ density-deviation trend is again observed (Fig. 4B). When extrapolated to lower densities the polynomial predicts progressive reconciliation with pure noble gas–water values, with deviations less than 5% (i.e., within typical experimental uncertainty) at densities below 69 kg/m³. Given the good fit to the experimental data (R² = 0.94) and the reconciliation with partitioning for noble gas–water systems at low densities, the polynomial fit provides a viable method of defining deviation from pure noble gas–water behaviour as a function of CO₂ density between 0 and 656 kg/m³. R² values are given to indicate confidence of polynomial fit to experimental values. Coefficients are also given for calculating 1σ uncertainty via Eq. (4). All values are given to three significant figures.

### Table 7

| Noble Gas | Coefficient a | Coefficient b | Coefficient c | R²   |
|-----------|---------------|---------------|---------------|------|
| Helium    | −1.213E−04    | 2.564E−02     | 0             | 0.48 |
|           | +1σ           | −8.195E−05    | 1.430E−03     | 9.467E+00 |
|           | −1σ           | −1.607E−04    | 4.985E−02     | −9.467E+00 |
| Argon     | 2.218E−04     | −3.297E−02    | 0             | 0.73 |
|           | +1σ           | 2.841E−04     | −6.945E−02    | 1.226E+01 |
|           | −1σ           | 1.595E−04     | 3.510E−03     | −1.226E+01 |
| Krypton   | 1.685E−04     | 6.142E−02     | 0             | 0.94 |
|           | +1σ           | 2.099E−04     | 3.617E−02     | 9.680E+00 |
|           | −1σ           | 1.271E−04     | 8.667E−02     | −9.680E+00 |
| Xenon     | 4.873E−04     | 1.671E−01     | 0             | 0.98 |
|           | +1σ           | 5.771E−04     | 1.146E−01     | 1.626E+01 |
|           | −1σ           | 3.976E−04     | 2.195E−01     | −1.626E+01 |

Fig. 4. Deviations of experimental Henry’s coefficients for xenon (A), krypton (B), argon (C) and helium (D) from pure noble gas–water behaviour (Crovetto et al., 1982; Smith, 1985) as a function of CO₂ density. CO₂ density was calculated via the NIST website (NIST, 2014). Coefficients for each polynomial and their 1σ confidence envelopes are given in Table 7. Low pressure samples (Experiment 0) are provided for reference and discussed fully in Section 2.5.
noble gas–water systems as a function of CO₂ density for the 0–656 kg/m³ range.

3.4. Argon

Most argon Henry’s coefficients were close to or within pure noble gas–water values. However, at sufficiently high CO₂ densities, a general positive deviation emerges (Fig. 4C). At low density no overall deviation from the pure noble gas–water phase partition coefficients was observed. Experiments 5 and 8 (169 & 252 kg/m³) showed high relative CO₂ phase concentrations. Argon is susceptible to air contamination and it is possible that some of the variance at both low and high CO₂ density may be due to trapped air in the sampling process. However, as they broadly supported the trends seen for argon and the heavier noble gases (i.e., positive deviation) Experiments 5 and 8 were included in the polynomial fit. In contrast, Experiment 3 (602 kg/m³) did not fit the overall positive trend. Given that this otherwise positive density-deviation trend is strongly reinforced by both closest sister elements, krypton and xenon, the validity of this data point was doubtful and has been omitted from the polynomial fit. This resultant fit reached reasonable agreement with the data ($R^2 = 0.73$) and expressed deviations greater than 5% of pure noble gas–water values at 242 kg/m³. This polynomial can therefore be used to predict deviation from pure noble gas–water systems as a function of CO₂ density for the 0–656 kg/m³ range for argon.

3.5. Neon

$^{20}$Ne, $^{21}$Ne, and $^{22}$Ne were measured in both phases. However the quadrupole is unable to resolve contributions from doubly charged interference ions $^{40}$Ar$^{++}$, and CO₂$^+$ contributing to masses 20 and 22, respectively. Although the relationship between singly and doubly charged ions for the Hiden Quadrupole Mass Spectrometer is constrained, these interferences are substantial and, the propagated error when correcting for doubly charged ions results in unresolvable Henry’s coefficients for $^{20}$Ne and $^{22}$Ne. The remaining isotope, $^{21}$Ne, is only present in minor quantities and the signal size from the water phase resulted in large uncertainties. While we report the Ne data for completeness (Tables 4–6), no conclusions are reached for neon behaviour within dense CO₂–water systems in this work and future experiments will need either a larger relative Ne spike or to adopt analytical instruments with the ability to resolve or accurately correct for the interference peaks. This issue is specific to neon and therefore all other noble gas Henry’s coefficients remained unaffected.

3.6. Helium

At low density most experimental values are within uncertainty indistinguishable from pure noble gas/water values. Overall, a progressive negative deviation trend was observed as a function of increasing density, characterised by the second order polynomial with deviations greater than 5% of pure noble gas–water systems (i.e., exceeding experimental uncertainty) at CO₂ densities of 335 kg/m³ (Fig. 4D). The polynomial fit allows correction for the CO₂ density-derived deviation from pure noble gas–water systems to be interpolated for a CO₂ density range of 0–656 kg/m³.

4. DISCUSSION

From Eq. (2), Henry’s coefficients for noble gases in both CO₂–water and noble gas water systems are the net effect of non-ideality in both phases acting on the Henry’s constant. Differences between the two Henry’s coefficients therefore are due to changes in these non-ideal terms which require evaluation.

4.1. Non-ideality in the water phase

The temperature range presented here fell within the original pure noble gas–water range (Crovetto et al., 1982; Smith, 1985). Therefore thermal controls on solubility within the water phase were already accounted for. At pressures relevant to the experiment designed here, pressure can have an effect on Henry’s coefficients (Hou et al., 2013) and can be quantified using the Poynting correction (Enns et al., 1965; Prini and Crovetto, 1989).

$$\Phi_P = \gamma K_0^* \frac{\tau_{\text{m}}(P_e - P)}{P}$$

where $\tau_{\text{m}}$ is the partial molar volume of $i$ at infinite dilution at reference pressure $P^*_0$, $P$ is the experimental pressure, $K_0^*$ is Henry’s constant at the reference pressure, $R$ is the ideal gas constant and $T$ is the temperature. Partial molar volume is considered pressure independent (Enns et al., 1965; Gerth, 1983; Prini and Crovetto, 1989; Poling et al., 2001). Values of $\tau_{\text{m}}$, for each noble gas are given in Table 8.

The Poynting correction was applied to the experimental conditions which yielded the highest density (Experiment 4, 656 kg/m³) to determine the maximum effect pressure-derived non-ideality in the water phase has on Henry’s coefficients (Table 9).

From Table 9 although pressure has a minor effect on solubility the majority of the observed deviations remain unaccounted for. Additionally for helium, contrary to the observed trend, the Poynting correction indicates a lower solubility as a function of increasing pressure. Both observations are indicative that pressure acting on the

| Noble gas | Partial molar volume (cm³/mol) |
|-----------|--------------------------------|
| Helium    | 29.7                           |
| Argon     | 32.2                           |
| Krypton   | 32.8                           |
| Xenon     | 42.7                           |

Table 8
Partial molar volumes of each noble gas at infinite dilution. Helium and argon values taken from Enns et al. (1965), krypton value taken from Moore et al. (1982) and xenon value taken from Biggerstaff and Wood (1988).
Table 9
Assessing the effects of the Poynting correction on Henry’s coefficients for the highest experimental density (656 kg/m³) where deviations were at their greatest. Partial molar volumes for each noble gas are taken from Table 8. The outstanding deviations are the deviations derived using Eq. (4) minus those which can be attributed to pressure effects.

| Noble gas | Change to partitioning due to pressure (%) | Experimental deviation (%) | Outstanding deviation (%) |
|-----------|-------------------------------------------|---------------------------|--------------------------|
| Helium    | 16                                        | –35                       | –51                      |
| Argon     | 17                                        | 74                        | 57                       |
| Krypton   | 18                                        | 113                       | 95                       |
| Xenon     | 24                                        | 319                       | 295                      |

water phase has a relatively minor effect. Consequently this is discounted as the driving force affecting partitioning.

Currently the effect of CO₂(aq) on noble gas solubility remains unquantified (Ballentine et al., 2002). However, it is reasonable to assume that CO₂(aq) will reduce noble gas solubility given the finite capacity of water for non-polar solute particles at fixed P,T conditions. Experimental CO₂(aq) concentrations range between 0.73 and 1.13 mol/l (Spycher et al., 2003). Under these concentrations the CO₂(aq) activity coefficient remains at unity (Diamond and Akinfeiev, 2003) therefore no significant change in overall behaviour or internal structure of the water phase can be assumed. Accordingly, a negligible effect on the chemical potential of a dissolved inert gas is expected.

Neither pressure nor CO₂(aq) adequately account for the observed deviations from pure noble gas–water Henry’s coefficients. With no additional factors attributable to water phase non-ideality, non-ideality in the water phase can be ruled out as the dominant force driving the observed deviations in Henry’s coefficients.

4.2. Non-ideality in the CO₂ phase

At sufficiently high densities a departure from the ideal gas law due to enhanced molecular interactions within the CO₂ phase is expected. CO₂–noble gas interactions are similarly expected to become significant at higher densities. With all water phase factors discounted it is these enhanced interactions that are expected to be the source of deviation. For argon, krypton and xenon this non-ideality is expressed as a greater affinity for the CO₂ phase (Φ_i < 1), while with helium the opposite is observed (Φ_i > 1).

Presently the effect of water within the CO₂ phase on Henry’s coefficients is undocumented. However, under the conditions where deviations are at their greatest (Experiment 4) the molar fraction of water within the CO₂ phase is extremely low (0.006, Briones et al., 1987). Additionally, for any given pressure, the molar fraction of water increases with temperature (Spycher et al., 2003); this corresponds with decreasing CO₂ density and therefore does not fit with the observed trends. Consequently it is reasonable to assume water within this phase has a negligible effect on partitioning.

Dense CO₂ acts as a fluid solvent (e.g., Kamihira et al., 1987; Black, 1996; Morgenstern et al., 1996). This allows application within a wide range of industrial processes such as decaffeination (e.g., Mohamed et al., 2002; Kim et al., 2008), Enhanced Oil Recovery (EOR) (e.g., Blunt et al., 1993; Ravagnani et al., 2009) or in pharmaceutical processes (e.g., Ferrieri, 2003; Reverchon et al., 2009). As the solvation power of a fluid increases as a function of density (Giddings et al., 1968; Yonker et al., 1986) a link between CO₂ density (i.e., non-ideality) and solvation power can be assumed (Baiker, 1999). In addition there will be dispersion effects, related to atomic size, which also increase as a function of density. Coupled with this, dense CO₂ phases possess polar properties (e.g., Reynolds et al., 1996; Kaufman, 2001; Raveendran et al., 2005). Therefore, noble gas solubility within the CO₂ phase is expected to be enhanced in order of both molecular size and polarisability (i.e., xenon → helium). Similar induced polarisation solubility trends are observed within the water phase (Ballentine et al., 2002; Kipfer et al., 2002). Helium solubility however is reduced as a function of CO₂ density. This is considered to be due to the low polarisability of helium resulting in the net repulsive forces in the CO₂ phase at higher density outweighing the enhanced attractive forces. The positive fugacity is already observed in the pure noble gas phase at low temperature high pressure (Ballentine et al., 2002, Table 1). Thus helium behaviour is also satisfactorily described via this mechanism.

4.3. Extrapolating observed deviation trends?

Overall, deviations increased as a function of density for all noble gases and this trend in the data is well-defined using a second order polynomial. To extrapolate these polynomials beyond the range presented here would presume that the forces driving deviation continue to respond accordingly to changes in CO₂ density. Additionally the assumption that no other factor will significantly affect noble gas partitioning would have to be maintained, regardless of changes to the thermodynamic conditions outlined in this work. At present no data supports either assumption. Consequently the experimentally defined trend lines cannot reasonably be extrapolated beyond the range presented.

4.4. Impacts of study on geological processes

Noble gas phase partitioning within dense CO₂ environments deviates strongly from low-density predictions. Consequently, where low pressure partitioning has been assumed, a moderate degree of recalculation will be required to maintain mass-balance as in the case study example. For example processes which increase the noble gas content of a CO₂ phase (e.g., degassing of groundwater) will require scaling down. Additionally key isotopic ratios such as ^4He/^36Ar, ^84Kr/^36Ar will also be affected. The CO₂ density must therefore be considered where such ratios are used to quantify geological processes. Similarly, due to this change in relative Henry’s coefficients, a re-evaluation
of all models involving open Rayleigh degassing in CO$_2$ environments is required.

The Henry’s coefficients presented here can be used to re-interpret previous, related studies. As an example we consider the paper of Gilfillan et al. (2008) which uses noble gases to consider the processes controlling CO$_2$ emplacement and removal in naturally occurring CO$_2$ in the Sheep Mountain and Bravo Dome gas fields (Colorado and New Mexico, USA). Here we re-evaluate the models employed in order to take into account the effect of dense CO$_2$. For a full geological history and analytical methodology the reader is directed to the original text (Gilfillan et al., 2008). As a summary, Air-Saturated Water (ASW) noble gases are initially considered to be completely stripped out of the water phase due to sustained injection of magmatic CO$_2$. This results in a gas field that is dominated by CO$_2$ containing noble gases in ASW proportions. Following reservoir filling, a period of redissolution occurs. This redissolution was modelled as open system noble gas loss from the CO$_2$ phase. The fractionation of noble gases remaining within the CO$_2$ phase was modelled for both fields as Raleigh loss using Henry’s coefficients. Graphs showing original and re-evaluated fractionation via redissolution curves for Bravo Dome and Sheep Mountain are presented in Fig. 5A and B.

For the Sheep Mountain gas field an additional initial krypton component 400% in excess of ASW values was proposed, in order to be consistent with the physical model. In the absence of Henry’s coefficients for CO$_2$–water systems the effect of CO$_2$ was estimated using virial coefficients to calculate pressure-derived fugacities (Ballentine et al., 2002). For this case study the Henry’s coefficients, calculated for reservoir temperatures and salinities, are recalculated to incorporate the CO$_2$-derived density deviations presented in this work. For neon values the virial-derived Gilfillan et al., values are used as these remain the best estimation of neon non-ideal behaviour (Fig. 5).

In the case of Bravo Dome, although a slightly shallower fractionation gradient is observed, the curve still reaches good agreement with the measured data (Fig. 5A). This indicates that after including the effect of dense CO$_2$ (273 kg/m$^3$) the straightforward redissolution model remains viable for Bravo Dome. However, in the case of Sheep Mountain, where CO$_2$ density was significantly higher (572 kg/m$^3$), a greater effect is observed (Fig. 5B). Where originally an excess of 400% initial krypton was required to reconcile the dissolution model with the data, the new, significantly steeper, fractionation gradient requires an initial excess of ~700%. This increased excess therefore requires a greater external input of krypton e.g. excess air or organic sediments.

4.5. Related systems

Re-evaluation is required of closely related binary systems. As the CO$_2$ density is responsible for the observed deviations, similar deviations should also be expected wherever a dense CO$_2$ phase is encountered. The CO$_2$–magma system is a prime example. During the ascent of magma, depressurisation results in the formation of separate phases (i.e., vesicles) within the melt (e.g., Paonita and Martelli, 2006, 2007). The vesicles consist of almost pure CO$_2$ (Carroll and Draper, 1994) which at those temperatures and pressures exists as a supercritical fluid (Sparks, 1978; NIST, 2014). The noble gases partition between the CO$_2$ and melt with a strong preference for the CO$_2$ phase. This partitioning is used to constrain variables such as degassing rates and magmatic origins. However, the partitioning coefficients used for this were derived from low pressure experiments and assume only melt composition affects noble gas partitioning (Jambon et al., 1986; Carroll and Draper, 1994; Hilton and Porcelli, 2003), and the effect of dense CO$_2$ is therefore considered negligible. Our study indicates the weakness in this assumption. Indeed recent simulation-based studies of CO$_2$-magma systems suggest that CO$_2$ density is an important model component (Sarda and Guillot, 2005; Guillot and Sarda, 2006; Aubry et al., 2013).

Our work indicates CO$_2$ density is liable to affect partitioning within all CO$_2$-based systems. These potentially include CO$_2$-hydrocarbon, CO$_2$-mantle and CO$_2$-hydrothermal fluid environments. It is therefore advised that wherever a dense CO$_2$ phase is present, noble
gas partitioning is liable to be poorly predicted by low pressure analogues.

Additionally, as differences are due to changes in the non-ideal terms derived for low densities, an argument can be made that noble gas behaviour within all dense binary phase geological systems, not just those involving CO₂, may require a degree of re-examination. A prime example is the methane–water system for which noble gases provide industrial and academic insights (e.g., Zhou et al., 2005; Hunt et al., 2012). However, noble gas behaviour within this system remains experimentally uncharacterised. Traditionally pure noble gas–water partitioning would be considered comparable for this system, but our work highlights the issue in making such an assumption. Accordingly, independent experimental investigation of noble gas behaviour is required.

5. SUMMARY

Noble gas Henry’s coefficients for supercritical CO₂–water systems do not replicate pure noble gas–water system values. This is due to density-derived non-ideality within the CO₂ phase. As density increases so too do deviations from predicted partition coefficients for all noble gases. These deviations are positive for argon, krypton and xenon. Helium possesses a negative deviation as a function of density and consequently decreases as a function of density. Deviations are most likely caused by dispersion effects related to atomic size coupled with CO₂ acting as a polar solvent with the solvation power increasing as a function of density. Polarisation is induced in each noble gas in line with their respective polarisabilities. Xenon affinity for the dense CO₂ phase is increased the most followed by krypton and then argon. This therefore results in positive deviations to the pure noble gas–water partition coefficients. Helium being far less polarisable and having the smallest atomic size has a reduced solubility compared with a near-ideal gas, thereby producing negative deviations from predicted partition coefficients. All of the deviation trends are suitably described using a second order polynomial fit for CO₂ densities up to 656 kg/m³. This fit provides a means to correct the existing pure noble gas–water partition coefficients derived from pure noble gas/water systems for use within high density CO₂ systems.

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APPENDIX 1. CALCULATING MEASUREMENT UNCERTAINTIES

Uncertainty was assessed using a two-tiered approach. Firstly all relative measurement uncertainties for each isotopic Henry’s constant were combined via addition in quadrature. All measurement uncertainties are described in Table A1 and the detailed methodology for calculating this uncertainty is given in this appendix.

However not all sources of uncertainty (e.g., the accuracy of NIST-derived densities) were satisfactorily included within this measurement uncertainty. All remaining sources of uncertainty would therefore affect reproducibility of each paired sample. Consequently, by determining the reproducibility uncertainty (ΔD(Proc(P,T))), all additional uncertainties could be reasonably incorporated. The process of deriving the reproducibility uncertainty for each isotope and combining it with the measurement uncertainty is outlined in this appendix.

To generate overall uncertainty for noble gases where multiple isotopes were analysed, arithmetic means were taken of each overall uncertainty and were divided by √n where n is the number of isotopes being combined.

Calculating measurement uncertainties

Experimental and sampling densities for the density correction were calculated using the respective pressures and temperatures via the NIST website (NIST, 2014). Therefore the corresponding uncertainty was considered a function of each of these values. Thus these were added

| Uncertainty factor | Symbol | Calculated uncertainty | Units |
|--------------------|--------|------------------------|-------|
| Noble gas isotope i in CO₂ phase | ΔC_{(P,T)}^i | 1σ of average peak intensity | bar |
| Noble gas isotope i in H₂O phase | ΔW_{(P,T)}^i | 1σ of average peak intensity | bar |
| Volume of CO₂ sampler | ΔV | 1 S.E. of mean volume | cm³ |
| Volume of H₂O sampler | ΔW_A | 1 S.E. of mean volume | cm³ |
| Aliquot correction factor | ΔF | 0.1275 (1 S.E. of mean) | N/A |
| Density correction for CO₂ phase | Δρ_C | Measurement uncertainty in pressure and temperature | kg/m³ |
| Density correction for H₂O phase | Δρ_W | Measurement uncertainty in pressure and temperature | kg/m³ |
| Uncertainty in pressure | ΔP | 1 | bar |
| Uncertainty in temperature | ΔT | 0.2 | °C |
in quadrature to calculate the relative uncertainty from the density correction:

\[
\frac{\Delta P}{\rho_d} = \sqrt{\left(\frac{\Delta T_{\text{Samp}}}{T_{\text{Samp}}} \right)^2 + \left(\frac{\Delta P_{\text{Samp}}}{P_{\text{Samp}}} \right)^2 + \left(\frac{\Delta T_{\text{Exp}}}{T_{\text{Exp}}} \right)^2 + \left(\frac{\Delta P_{\text{Exp}}}{P_{\text{Exp}}} \right)^2}
\]

(A1)

where \(\rho_d\) is the density correction factor for phase \(A\), \(T_{\text{Samp}}\) and \(P_{\text{Samp}}\) are sampling temperatures and pressures, \(T_{\text{Exp}}\) and \(P_{\text{Exp}}\) are experimental temperatures and pressures and \(\Delta\) denotes the corresponding uncertainty.

To calculate the net relative uncertainty for each partition coefficient calculated from a paired set of samples these relative uncertainties were added in quadrature:

\[
\frac{\Delta D_{\text{Exp}(P,T)}}{D_{\text{Exp}(P,T)}} = \sqrt{\left(\frac{\Delta C_i(P,T)}{C_i(P,T)} \right)^2 + \left(\frac{\Delta W_i(P,T)}{W_i(P,T)} \right)^2 + \left(\frac{\Delta C_f}{C_f} \right)^2 + \left(\frac{\Delta W_f}{W_f} \right)^2 + \left(\frac{\Delta F_i}{F_i} \right)^2 + \left(\frac{\Delta \rho_C}{\rho_C} \right)^2 + \left(\frac{\Delta \rho_w}{\rho_w} \right)^2}
\]

(A2)

By multiplying this relative uncertainty by the partition coefficient the absolute uncertainty could be determined. Thus uncertainty for each partition coefficient was derived for a CO\(_2\)–water paired set of samples.

To combine the paired samples for each noble gas at a given density each relative uncertainty was added in quadrature. These were then divided by the number of paired samples used for creating this average \((n_{P,T})\). This was then square-rooted to give the average relative uncertainty for a given density:

\[
\frac{\Delta D_{\text{Proc}}}{D_{\text{Proc}}} = \sqrt{\frac{\sum n_{P,T} \left(\frac{\left|\Delta D_{\text{Exp}(P,T)} - D_{\text{Exp}(P,T)}\right|}{D_{\text{Exp}(P,T)}}\right)^2}{n_{\text{tot}}}}
\]

(A3)

Assessing the reproducibility uncertainty

This was calculated using the average deviation of partition coefficients from the mean values. For this, each partition coefficient for a given isotope \(i\) \((D_{i(P,T)})\) was subtracted from the mean \((\bar{D}_{i(P,T)})\) to give the absolute deviation. By dividing each deviation by the mean value a relative deviation of each individual data point from the mean was calculated. These relative uncertainties for every partition coefficient were then all added in quadrature and divided by the total number of samples \((n_{\text{tot}})\). Finally this was square-rooted to generate the average the relative procedural reproducibility uncertainty for each isotope:

\[
\frac{\Delta D_{\text{Proc}}}{D_{\text{Proc}}} = \sqrt{\frac{\sum n_{P,T} \left(\frac{\left|\Delta D_{\text{Exp}(P,T)} - D_{\text{Exp}(P,T)}\right|}{D_{\text{Exp}(P,T)}}\right)^2}{n_{\text{tot}}}}
\]

(A4)

APPENDIX 2. SYSTEM STABILITY AND LINEARITY

System stability

System stability was evaluated by measuring the \(^{40}\text{Ar}\) content of six independent 0.2264 cm\(^3\) air calibrations over the course of a day. The mean and standard error was compared with 4 identical measurements collected 6 months previously. The mean and standard error are shown in Table A2 and demonstrate sufficient system stability for quantitative analysis.

| Table A2 | Assessment of system stability using air calibrations. The average \(^{40}\text{Ar}\) content of six single 0.2264 cm\(^3\) air calibrations taken over a day are compared with four single 0.2264 cm\(^3\) calibrations analysed over the course of a day 6 months prior. Percentage change is the increase in average values over this period. Uncertainty is quoted as one standard error. |
|-----------------|-----------------|-----------------|-----------------|
| 6 \(^{40}\text{Ar}\) air calibrations (bar) | 4 \(^{40}\text{Ar}\) air calibrations 6 months prior (bar) | Percentage change (%) | Combined \(^{40}\text{Ar}\) values (bar) |
| 9.37 (±0.09) \times 10^{-11} | 9.58 (±0.10) \times 10^{-11} | 2.23 (±1.45) | 9.45 (±0.09) \times 10^{-11} |

| Table A3 | Assessment of signal intensity calibration. The average signal response at mass 40 \(^{40}\text{Ar}\) of six single 0.2264 cm\(^3\) air standards compared to the average \(^{40}\text{Ar}\) content of three 4 \times 0.2264 cm\(^3\) air standards. Deviation from linearity was calculated as the percentage by which measured four-shot values differed from the expected values. Uncertainty is quoted as one standard error. |
|-----------------|-----------------|-----------------|-----------------|
| \(^{40}\text{Ar}\) in 1 \times air standard (bar) | Expected \(^{40}\text{Ar}\) in 4 \times air standard (bar) | Measured \(^{40}\text{Ar}\) in 4 \times air standard (bar) | Deviation from linearity (%) |
| 6.67 (±0.10) \times 10^{-11} | 2.67 (±0.04) \times 10^{-10} | 2.73 (±0.16) \times 10^{-10} | 2.2 (±6.3) |
System linearity

At higher pressures, ionisation and signal response in quadrupole mass spectrometers can become non-linear. This was assessed for the system using 0.2264 cm$^3$ aliquots of air standards, corrected for temperature, pressure, altitude and humidity. Single aliquots were analysed for four noble gases, diluted and expanded into known volumes (Ampule and aliquot A1) on the analytical line (Fig. 3). The relative signal intensities were compared for all noble gases and all were found to produce a linear response across the experimental range comparable to argon which was also measured as a control.

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