Techno-Economic Aspects of Noble Gases as Monitoring Tracers

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Abstract: A comprehensive monitoring program is an integral part of the safe operation of geological CO2 storage projects. Noble gases can be used as geochemical tracers to detect a CO2 anomaly and identify its origin, since they display unique signatures in the injected CO2 and naturally occurring geological fluids and gases of the storage site complex. In this study, we assess and demonstrate the suitability of noble gases in source identification of CO2 anomalies even when natural variability and analytical uncertainties are considered. Explicitly, injected CO2 becomes distinguishable from shallow fluids (e.g., subsea gas seeps) due to its inheritance of the radiogenic signature (e.g., high He) of deep crustal fluids by equilibration with the formation water. This equilibration also results in the CO2 inheriting a distinct Xe concentration and Xe/noble gas elemental ratios, which enable the CO2 to be differentiated from deep crustal hydrocarbon gases that may be in the vicinity of a storage reservoir. However, the derivation has uncertainties that may make the latter distinction less reliable. These uncertainties would be best and most economically addressed by coinjection of Xe with a distinct isotope ratio into the CO2 stream. However, such a tracer addition would add significant cost to monitoring programs of currently operating storage projects by up to 70% (i.e., from 1 $US/t to 1.7 $US/t).

Keywords: carbon capture and storage; monitoring and verification; noble gases; leakage detection; inherent distinctiveness; tracer addition; cost calculation

1. Introduction

Measurement, monitoring, and verification (MMV) programs are an essential part of the safe operation of geological CO2 storage projects. Noble gases are one chemical tracer group that has received attention due to their key properties of being inert and having a large number of isotopes that can be measured with high precision [5,6]. This makes them safe to utilize in industrial processes and their use does not have additional negative environmental consequences. The concentrations of He, Ne, Ar, Kr, Xe, and their isotopic ratios constitute the noble gas signature of a fluid. This practically labels most environmental reservoirs with an identifiable atmospheric [7], crustal (radiogenic) [8], or mantle [9] signature [10]. Further, injected CO2 from various capture processes (anthropogenic) has an inherent noble gas signature, which depends on the CO2 source and capture process [11–13].

The conservative behavior of noble gases and the knowledge of the baseline noble gas signatures of the storage system allows end-member mixing calculations to be performed, such as those used in a study that refused an alleged leak from a CO2 storage site [14].
Further, in CO₂ injection for enhanced oil recovery (EOR), changes in the noble gas signature of the produced fluids quantified the contribution of the injected CO₂ [15–17]. It is also possible to label the injected CO₂ with deliberately added noble gases to distinguish the signature of the injected CO₂ further [15,18]. However, this has only been conducted in small-scale injection experiments, but allowed for identification of the CO₂ gas plume distribution and of mixing and degassing processes, e.g., [19].

In the study by Roberts et al. [18], the addition of noble gas tracers is discussed from a practicality and cost perspective for large-scale storage. Tracer addition was analyzed to differentiate the noble gas signature of the injected, and eventually, leaking CO₂ from the background signatures of seawater and atmosphere. They conclude that He and ¹²⁴,¹²⁹Xe isotopes are most suitable as added tracers with costs in the order of 100–100,000 £/MtCO₂, similar to earlier results by Nimz et al. [15].

In [12,13], we outlined that the injected CO₂ is likely to inherit a radiogenic noble gas signature from the storage formation waters through equilibration or mixing with native formation gases. The calculation based on phase partitioning data for a high-pressure CO₂-H₂O-system [20] led to the hypothesis that injected CO₂ becomes differentiable from atmospheric signatures without the need of adding tracers.

The altered, radiogenic signature of the injected CO₂ is similar to natural hydrocarbons [21]. Hydrocarbon reservoirs are often found in the vicinity of a number of active or suggested CO₂ storage sites, e.g., [22–24]. A leak into such hydrocarbon reservoirs may not have the environmental impact of a leak to the seabed, but the contamination of these resources can have significant economic consequences. Further, if the CO₂ was to be fed back into the atmosphere, subsequent to gas or oil production, the climate mitigation value of a CCS project would be reduced. Therefore, there is the need to expand the evaluation of the distinctiveness of noble gas signatures and tracer addition to include the discrimination of CO₂ and hydrocarbon gases.

There are several possible scenarios of how leakage could occur to the atmosphere or seabed or into hydrocarbon reservoirs [25]. An overview of some possible pathways that may result in varying exchange of the injected CO₂ with the storage site fluids and mixture with different background reservoirs is shown in Figure 1. An injection well leak, case 1 in Figure 1, could leave the CO₂ with its initial composition; meanwhile, cases 2, 3, 4, and 5 would likely lead to significant interaction of the CO₂ with crustal fluids. Therefore, including the phase partitioning between the injected CO₂ and formation water data derived in [20] is an essential addition in the assessment of noble gas tracers, both from a technical and economic perspective.

In this work, we aim to validate the hypothesis of the inherent differentiability of the injected CO₂ from the atmospheric signatures quantitatively drawn in [12]. We will conduct calculations based on current analytical capabilities for noble gases and the natural variability of noble gas signatures. In the same way, we evaluate distinctiveness to other environmental signatures, i.e., natural gas and gas hydrates, that have to be considered in the leakage scenarios 2–5 in Figure 1. Then, we derive the necessary amounts and the associated cost for tracer addition to increase distinctiveness between the injected CO₂ and hydrocarbons, i.e., natural gas. Finally, we discuss the practical and economic roles of noble gases in a monitoring scheme and the implications for the viability of a storage project.
Figure 1. Overview sketch of a CO₂ storage site with areas typically attributed to a type of noble gas (NG) signature [10]. CO₂ will have a different NG signature for different plant types, such as postcombustion, natural gas processing, or oxyfuel [11,12]. Possible leakage pathways are (1) injection well leakage, (2) migration into and contamination of hydrocarbon reservoirs that may subsequently be produced, (3) fault leakage, (4) diffuse leakage, and (5) leakage through an abandoned well. Leaking CO₂ may enter the atmosphere or dissolve in seawater.

2. Materials and Methods
2.1. Background Concentrations

Noble gas tracers have widespread application in deciphering physical and chemical processes in geosciences [10] since they are only influenced by physical processes. There are three main groups of noble gas signatures in the environment, which are of atmospheric [7], crustal [8], or mantle [9] character and allow researchers to attribute the source of a fluid. In practice, these categories in themselves can have wide isotopic concentration ranges or be mixtures derived during the history of origin of a fluid. Characterizing the background fluids of a storage site is key for the applicability of noble gases as tracers.

Here, we collect and discuss the known signatures from samples relevant to a possible storage site in the North Sea (Table 1). This is due to the topicality of the Norwegian large-scale storage project ‘Longship’, which is currently under development [26]. Additionally, in this project, a contamination of the stratigraphically overlying Troll gas and oil field could be considered a risk [27]. Analogue studies may be used and discussed where samples are not available. These signatures may have to be adapted for other sites but reflect typical continental shelf settings.

For natural gas, sample analyses for the North Sea are currently available for the Sleipner Vest and the Magnus fields [28,29]. The results show a typical crustal, radiogenic character, derived from the accumulation of ⁴He, ²¹Ne, and ⁴⁰Ar through radioactive decay of the elements K, U, and Th in the rocks [8]. The radioactive accumulation is a function of time and the content of radioactive elements in the rock. The radiogenic signature acquires an atmospheric gas component from the in situ formation water that was in atmospheric equilibrium during sedimentation, or meteoric recharge. The additional amounts of stripped gas then depend on the gas/water volume ratio, \( V_g/V_w \), and if the hydrocarbon system is closed or open, allowing for significant gas volumes to migrate through a reservoir [28].
Table 1. Observed noble gas concentrations for different environmental fluids. Natural gas for He, Ne, and Ar compiled from the North Sea Sleipner and Magnus Fields in [28,29], and Kr and Xe from the Sleipner Field [28]. Natural, geologic CO$_2$ compiled from [30,31]. Sediment values from the South Pacific Ocean [32] and the Mediterranean Ridge for He [33]. Gas hydrates from [34]. Seawater concentrations after [35] (assuming a temperature of 7.5 °C and a salinity of 34.2 g/L). Atmospheric values from [7]. Captured CO$_2$ from several sites [11–13]. In cm$^3$/cm$^3$ STP, if not otherwise stated. Rounded to nearest tenth asides atmosphere.

<table>
<thead>
<tr>
<th></th>
<th>He</th>
<th>Ne</th>
<th>Ar</th>
<th>Kr</th>
<th>Xe</th>
</tr>
</thead>
<tbody>
<tr>
<td>Natural Gas/Oil</td>
<td>1.2E-05–1.3E-04</td>
<td>7.0E-07–2.7E-08</td>
<td>1.1E-05–4.1E-05</td>
<td>4.1E-09–5.2E-09</td>
<td>1.3E-09–1.6E-09</td>
</tr>
<tr>
<td>Natural, geologic CO$_2$</td>
<td>1.0E-04–1.0E-02</td>
<td>7.0E-08–1.8E-06</td>
<td>1.5E-05–2.7E-05</td>
<td>1.0E-10–1.0E-08</td>
<td>5.0E-12</td>
</tr>
<tr>
<td>Gas Hydrates</td>
<td>7.0E-10–9.0E-10</td>
<td>2.5E-09–6.2E-08</td>
<td>3.5E-05–5.3E-04</td>
<td>1.8E-08–3.0E-07</td>
<td>2.7E-09–9.4E-08</td>
</tr>
<tr>
<td>Seafloor Sediments (cm$^3$/g)</td>
<td>1.0E-04–1.0E-02</td>
<td>5.0E-09–5.0E-07</td>
<td>1.0E-05–1.0E-08</td>
<td>5.0E-08–1.5E-09</td>
<td>8.0E-09–1.7E-08</td>
</tr>
<tr>
<td>Seawater (cm$^3$/g)</td>
<td>4.0E-08</td>
<td>5.7E-09</td>
<td>1.1E-06</td>
<td>8.5E-08</td>
<td>1.1E-08</td>
</tr>
<tr>
<td>Atmosphere</td>
<td>5.24E-06</td>
<td>1.82E-05</td>
<td>9.34E-03</td>
<td>1.13E-06</td>
<td>8.7E-08</td>
</tr>
<tr>
<td>Captured CO$_2$</td>
<td>2.3E-09–2.9E-06</td>
<td>5.9E-11–4.3E-07</td>
<td>3.7E-10–1.4E-04</td>
<td>3.6E-12–6.3E-09</td>
<td>1.1E-13–1.9E-09</td>
</tr>
</tbody>
</table>

Natural, geologic CO$_2$ shows dominantly magmatic, and hence, mantle signatures with subsequent equilibration with radiogenic formation water. The values in Table 1 are from large natural CO$_2$ systems in the US [30,31]. Natural CO$_2$ in the North Sea context is typically a minor constituent of natural gas accumulations, e.g., [22], and would therefore have very similar noble gas signatures as the associated hydrocarbons.

The values for both natural gas and natural, geologic CO$_2$ depict that the noble gas concentration within what is considered a radiogenic signature can vary by orders of magnitude (see Table 1). For the ‘Longship’ project, the most relevant radiogenic signature to compare to with respect to traceability of CO$_2$ is storage reservoir formation water and gas and oil in the Troll Field stratigraphically overlying the storage formation. Even though the Troll gas and oil reservoirs are shallower than in the Magnus and Sleipner fields, the age and the proposed origin of the Troll hydrocarbons in the Oseberg area [36] would suggest that noble gas signatures in the Troll field are similar to those in Table 1.

Close to the seabed, gas hydrates may occur. Only total gas concentration, not isotopic composition, have been analyzed for these [34]. Those samples were collected outside the coast of Oregon, US, and showed a quite specific fractionation pattern with preferential incorporation of heavier noble gases and suppression of the lighter ones. This fractionation process should be independent of the location, making the concentrations applicable for the North Sea if gas hydrates were to be found.

Analysis of the dissolved noble gas distribution in the pore-water of shallow, unconsolidated sediments are relatively rare [32,33,37] and, for the North Sea, there is no dataset available. Sediment pore-water concentrations are dependent on the temperature during sedimentation and if there are leakage sites, e.g., shallow biogenic gas or deep fluid release, such as black smokers, nearby. The data from [32,33,37] in Table 1 has to be seen in the context of plate boundaries and is therefore not representative for the continental crust of the North Sea. However, there have been methane leakages above the Troll field in the past [38], which could have left a minor radiogenic component in the pore-water signature.

Seawater, if not in the vicinity of specific fluid releases, is typically air equilibrated water at a given temperature and salinity [35]. This means atmospheric gas, including noble gases, is dissolved based on the solubility of the respective gas species at these conditions in water. In case there are seepages, ocean currents may redistribute them quickly, so anomalies can be local. The atmosphere is typically well mixed and has a specific noble gas signature [7].

The noble gas signature of captured CO$_2$ has been analyzed for several capture plants in Norway [12,13] and other countries [11]. Noble gas concentrations are typically low and the isotopic signatures are either airlike for plants with combustion prior to capture (e.g., waste incineration) or natural-gas-like (e.g., natural gas processing). For captured CO$_2$,
the observed range spans several orders of magnitude. Captured CO\(_2\) from oxyfuel plants, for example, are significantly enriched in Kr and Xe [6].

With regards to isotopic ratios, there are the three main signatures and fluids that represent mixtures of these to various degrees. Typical values for the mainly affected isotopes are shown in Table 2.

Noble gas ratios associated with hydrocarbons are, as previously discussed for concentrations, dependent on the geological setting of a reservoir and the provenance of the hydrocarbons. In Table 2, isotopic ratios are shown for the North Sea [28]. However, the \(\frac{\text{He}}{\text{He}}\) and \(\frac{\text{Ar}}{\text{Ar}}\) ratio especially can vary significantly, e.g., natural gas from Snøhvit has a He-ratio an order of magnitude lower [12].

Table 2. Isotopic ratios for the main signatures: atmospheric; crustal, hence radiogenic; and mantle. Atmospheric values from [7]. Natural gas from the Sleipner field [28]. Mantle data from [9].

<table>
<thead>
<tr>
<th>Isotopic Species</th>
<th>Atmospheric Signature</th>
<th>Crustal Signature</th>
<th>Mantle signature (MORB)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\frac{\text{He}}{\text{He}}) (R/R(_A))</td>
<td>1</td>
<td>(\sim 0.1)</td>
<td>8</td>
</tr>
<tr>
<td>(\frac{\text{Ne}}{\text{Ne}})</td>
<td>0.029</td>
<td>(\sim 0.033)</td>
<td>0.06</td>
</tr>
<tr>
<td>(\frac{\text{Ar}}{\text{Ar}})</td>
<td>296</td>
<td>(\sim 455)</td>
<td>(\sim 30,000)</td>
</tr>
<tr>
<td>(\frac{\text{Kr}}{\text{Kr}})</td>
<td>0.303</td>
<td>(\sim 0.303)</td>
<td>-</td>
</tr>
<tr>
<td>(\frac{\text{Xe}}{\text{Xe}})</td>
<td>6.61</td>
<td>(\sim 6.61)</td>
<td>-</td>
</tr>
</tbody>
</table>

For captured CO\(_2\), the ratios are dependent on the source of the CO\(_2\) and if combustion has taken place prior to capture, e.g., at heat and power plants. Combustion introduces noble gases from the air shifting ratios towards atmospheric values. At natural gas processing sites, the radiogenic ratios are maintained [12]. However, one has to consider that the CO\(_2\) can undergo interaction with, for example, in situ formation water or gases, subsequently altering the noble gas fingerprint, as we discuss next.

### 2.2. Phase Partitioning

Following injection, the equilibration of the injected CO\(_2\) with the formation water of the storage reservoir will significantly modify its composition, especially given that captured and injected CO\(_2\) will be significantly depleted in noble gases [12]. We calculate this phase partitioning by applying Henry’s law following the derivation for the thermodynamic conditions governing in the crust in [39]. The gas concentration \(c_i\) of a gas \(i\) in the gas phase calculates to

\[
c_{ig} = c_{iT} \left( \frac{22400T \rho_{H_2O} V_{H_2O}}{18 \cdot 273 \cdot K_i,\text{CO}_2\_\text{H}_2\text{O} \cdot V_{\text{CO}_2} + 1} \right)^{-1},
\]

with the total concentration \(c_T\) of gas \(i\); water density \(\rho_{H_2O}\) at system pressure; temperature \(T\); activity \(\gamma_i\); and fugacity \(\phi_i\) of the gas at system pressure, temperature, and volume of water \(\text{CO}_2\), \(V_{\text{H}_2\text{O}}\), and \(V_{\text{CO}_2}\), respectively.

Values of \(K_i,\text{H}_2\text{O}\), \(\gamma_i\), and \(\phi_i\) are collected in [39]. However, Henry’s constants \(K_i\) of noble gases were found to deviate significantly for a high-density CO\(_2\)-H\(_2\)O system [20]. Warr et al. [20] experimentally derived deviation factors \(\kappa_i\) for all noble gases besides Ne and Rn. This correction allows calculation of the deviation of Henry’s constant with gas-specific constants in the dependency of CO\(_2\) density:

\[
K_{i,\text{CO}_2\_\text{H}_2\text{O}} = \kappa_i K_{i,\text{H}_2\text{O}} = \left( a_i \rho_{\text{CO}_2}^2 + b_i \rho_{\text{CO}_2} + c_i \right) K_{i,\text{H}_2\text{O}}.
\]

\(K_{i,\text{H}_2\text{O}}\) is Henry’s constant at a given temperature in units of pressure for a low-pressure air–water system. The upper limit for density in the experiments of [20] was 656 kg/m\(^3\). Though explicitly advised against, we also apply this deviation for densities up to 700 kg/m\(^3\) due to the lack of data.
The densities in Equations (1) and (2) are also functions of temperature and pressure, which can be expressed as gradients of depth. Geothermal and pressure gradients vary in different geological settings. Here, we use 30 K/km and 10.18 MPa/km. We derive the densities for CO\(_2\) from [40] and for H\(_2\)O from [41].

Since Equation (1) is a function of the volume ratio \(V_{H_2O}/V_{CO_2}\), phase partitioning can be calculated independent of actual storage site volume. In this work, we derive phase-partitioned values at the ratio of maximum saturation \(S_{max}\) of CO\(_2\) for residual trapping in a reservoir rock containing CO\(_2\) and water. The CO\(_2\) saturation was found experimentally to be between 13 and 92\% for several reservoir rock samples, with a mean of 61\% [42], which is subsequently used.

The total concentration of a gas in the system is derived from the sum of the product of concentration in the single phases and their volumes, hence,

\[
c_{i,T} = c_{i,CO_2,\text{inj}} \cdot V_{CO_2} + c_{i,H_2O} \cdot V_{H_2O},
\]

where \(c_{i,CO_2,\text{inj}}\) are the inherent concentrations in captured CO\(_2\) (see Table 1). There are currently no formation water analyses on noble gases for the North Sea. Therefore, we derive the formation water concentrations, \(c_{i,H_2O}\), assuming natural gas from the Sleipner field in [28] had been in equilibrium with formation water. For the calculation of tracer addition, the concentrations in \(c_{i,CO_2,\text{inj}}\) can be altered such that Equation (1) results in the desired \(c_{i,g}\).

### 2.3. Distinctiveness

The distinctiveness of the noble gas signature is constrained by analytical capabilities and natural variability. These have to be considered when analyzing the reliability of mixing calculations. In [18], it is assumed that “10 times the detectable perturbation above background levels are required for reliable tracer detection in the atmosphere or seawater”, hence, a leak would then be identified. As discussed in Section 2.1, the signatures in the atmosphere and seawater that were analyzed in [18] can be reasonably assumed to be fixed.

Tables 1 and 2 show that this is not the case for hydrocarbons and gas hydrates, such that the natural variability has to be considered to further constrain the distinctiveness. We use the values in [28] to derive the natural variability that can be expected for a gas field. The resulting ranges are smaller than observed in the Magnus Field [29]; however, it appears that analytical capabilities have increased significantly, such that the samples from [28] are likely to reflect state-of-the-art precision and accuracy.

Our criterion for being distinct is that the resulting concentration in a mixture of 90\% background fluid and 10\% equilibrated CO\(_2\) has to be larger than the natural variability plus the 3\(\sigma\)-interval of the typical measurement error in [28].

### 3. Results

#### 3.1. Equilibration with Formation Water

The phase partitioning of injected CO\(_2\) with the formation water leads to a significant change of the initially depleted signature of the captured CO\(_2\). The results of the calculation at 2000 m depth following Section 2.2 for the respective isotopes and ratios are shown in Table 3. Ne is not further considered since there are no correction terms for a CO\(_2\)-H\(_2\)O-system available.

Injected CO\(_2\) acquires a specific signature. With the high He (\(~3.90E-05 \text{ cm}^3\text{STP} / \text{cm}^3\text{STP}\)), it may be considered a radiogenic signature; however, the correction term for the solubilities for a CO\(_2\)-H\(_2\)O-system (Equation (2)) affects each gas species individually. For example, the phase partitioning leads to a signature that is \(~500\%\) different compared to natural gas in Xe. It is also very different compared to atmosphere, \(~750\%\) in He, and compared to gas hydrates \(~5E+07\%\) in He Table 3.

Results are calculated both for the highest and lowest values in captured CO\(_2\) (see Table 1). The concentrations in the injected CO\(_2\) are most influential for Ar results with a resulting \(~50\%\) difference for \(^{36}\text{Ar}\).
These values can be compared to the analytical uncertainty of an element's measurement and the natural variability in natural gas for the results of the Sleipner field [28], which can be as high as 45%, as stated in Table 3.

| Noble gas concentration in cm$^3$ STP/cm$^3$ STP and isotopic ratios after equilibration with formation water for low and high concentrations in captured CO$_2$, as in Table 1. Analytical uncertainty and natural gas variability from the Sleipner field in [28]. Concentrations and ratios are relative to the respective background fluids; the stated ranges are for $c_{g,low,S_{max}}$ and $c_{g,high,S_{max}}$, respectively. |
|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
|                  | $c_{g,low,S_{max}}$ | $c_{g,high,S_{max}}$ | Analytical Variability | Variability in Natural Gas | $c_{g,S_{max}}$ rel. to Average Natural Gas | $c_{g,S_{max}}$ rel. to Atmosphere | $c_{g,S_{max}}$ rel. to Hydrates |
| **He**           | 3.90E-05         | 3.92E-05         | 2%              | 8%              | 62–63%            | 744–749%          | 4,900,000%       |
| $^4$He           | 3.90E-05         | 3.92E-05         | 2%              | 8%              | 62–63%            | 744–749%          | -                |
| $^3$He           | 5.86E-12         | 6.49E-12         | 3%              | 35%             | 62–68%            | 80–88%            | -                |
| $^3$He/$^4$He    | 1.50E-07         | 1.65E-07         | 3%              | 13%             | 104–115%          | 11–12%            | -                |
| **Ar**           | 1.13E-07         | 1.75E-07         | 1%              | 45%             | 166–258%          | 1%               | 52–71%           |
| $^{36}$Ar        | 5.21E-05         | 7.07E-05         | 2%              | 30%             | 166–225%          | 1%               | -                |
| $^{40}$Ar        | 4.63E+02         | 4.05E+02         | 1%              | 5%              | 99–87%            | 156–137%          | -                |
| **Kr**           | 1.06E-07         | 1.16E-07         | 2%              | 18%             | 239–262%          | 1%               | 13–14%           |
| $^{84}$Kr        | 2.46E+02         | 2.69E+02         | 1%              | 17%             | 239–262%          | 1%               | -                |
| $^{86}$Kr/$^{84}$Kr | 3.03E-01         | 3.03E-01         | 1%              | 1%              | 100%              | 99%              | -                |
| **Xe**           | 6.36E-09         | 6.67E-09         | 5%              | 10%             | 458–480%          | 7–8%             | 13%              |
| $^{124}$Xe       | 6.37E-12         | 7.46E-12         | 3%              | 8%              | 458–536%          | 8–9%             | -                |
| $^{126}$Xe       | 6.67E-12         | 6.67E-12         | 3%              | 11%             | 458–540%          | 7–9%             | -                |
| $^{128}$Xe       | 1.16E-10         | 1.38E-10         | 3%              | 17%             | 458–544%          | 7–8%             | -                |
| $^{130}$Xe       | 1.68E-09         | 1.98E-09         | 3%              | 11%             | 458–540%          | 7–9%             | -                |
| $^{131}$Xe       | 2.56E-10         | 3.03E-10         | 3%              | 12%             | 458–541%          | 7–9%             | -                |
| $^{132}$Xe       | 1.36E-09         | 1.60E-09         | 3%              | 10%             | 458–539%          | 7–9%             | -                |
| $^{133}$Xe       | 2.17E-09         | 2.02E-09         | 3%              | 9%              | 458–540%          | 7–9%             | -                |
| $^{134}$Xe       | 6.36E-10         | 7.82E-10         | 3%              | 11%             | 458–540%          | 7–9%             | -                |
| $^{136}$Xe       | 5.64E-10         | 6.65E-10         | 3%              | 11%             | 458–540%          | 7–9%             | -                |
| $^{124}$Xe/$^{130}$Xe | 2.49E-02         | 2.46E-02         | 5%              | 9%              | 101–100%          | 106–105%         | -                |
| $^{126}$Xe/$^{130}$Xe | 2.21E-02         | 2.20E-02         | 5%              | 3%              | 101%              | 101–101%         | -                |
| $^{128}$Xe/$^{130}$Xe | 4.52E-01         | 4.55E-01         | 5%              | 5%              | 101–102%          | 96–96%           | -                |
| $^{130}$Xe/$^{130}$Xe | 6.55E+00         | 6.54E+00         | 5%              | 1%              | 101%              | 101%             | -                |
| $^{131}$Xe/$^{130}$Xe | 5.31E+00         | 5.30E+00         | 5%              | 2%              | 101%              | 102%             | -                |
| $^{132}$Xe/$^{130}$Xe | 6.68E+00         | 6.67E+00         | 5%              | 1%              | 101%              | 101%             | -                |
| $^{134}$Xe/$^{130}$Xe | 2.59E+00         | 2.58E+00         | 5%              | 1%              | 101%              | 101%             | -                |
| $^{136}$Xe/$^{130}$Xe | 2.20E+00         | 2.20E+00         | 5%              | 1%              | 101%              | 101%             | -                |

### 3.2. Tracer Addition

Needed concentrations in the injection stream to achieve additional distinction range from 1E-11 cm$^3$ STP/cm$^3$ STP for some Xe isotopes to 1E-03 cm$^3$ STP/cm$^3$ STP for elemental He. The results for the amounts of a specific gas isotope needed for deliberate tracer addition are shown in Table 4. Calculations are conducted in order to increase distinctiveness of the injected, equilibrating CO$_2$ from natural gas, following the description in Section 2.2. The results fulfill the condition set in Section 2.3. Then, the necessary concentrations...
that the injected CO$_2$ needs to be labeled with to achieve the given concentrations after equilibration are given.

The necessary concentrations in the injected CO$_2$ are scaled up to volumes of the respective tracer gas needed in a 1MtCO$_2$/a storage project, depicting the large range of concentrations needed for the different gases. These volumes range from 1E+01 to 1E+08 l/a (Table 4). From the amounts needed, the cost of an isotope addition is derived to a wide range of 0.7 to 37,000 $US/tCO$_2$ for the different gases. For reference, the price calculation is performed to fulfill the method used in [18] (see Section 2.3). Here, tracer addition would be up to 3 orders of magnitude cheaper.

Table 4. Necessary noble gas concentrations in cm$^3$/cm$^3$ STP and isotopic ratios to fulfill the detection condition described in Section 2.3 both in the equilibrated and the injected CO$_2$. For isotopic ratios, calculations are conducted for the isotope in the numerator of the ratio. As shown in Section 3.1, the distinctiveness condition for Xe and its isotopes is inherently met in our calculations. For Xe ratios, a concentration of 1.50E-11 $^{130}$Xe inherent in the injected CO$_2$ is assumed. The volume is calculated for 1MtCO$_2$/a storage project, and the associated cost for a tracer addition is derived. For comparability, the cost is additionally derived with the method in [18] and gas prices as in [18], where applicable. Exchange rate used for prices in [18]: £ = 1.3 $US.

<table>
<thead>
<tr>
<th>Gas</th>
<th>Nec. Conc./Ratio in Equilibrated CO$_2$</th>
<th>Nec. Conc. in Injected CO$_2$</th>
<th>Nec. Volume in a 1MtCO$_2$/a Project (l/a)</th>
<th>Gas Price ($US/l)</th>
<th>Cost ($US/tCO$_2$)</th>
<th>Cost; Method as in [18] ($US/tCO$_2$)</th>
<th>Gas Price Source</th>
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<tr>
<td>He</td>
<td>1.53E-04</td>
<td>1.26E-03</td>
<td>6.43E+08</td>
<td>57.2</td>
<td>37,000</td>
<td>18,000</td>
<td>[18]</td>
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<td>$^3$He</td>
<td>3.02E-11</td>
<td>2.72E-10</td>
<td>1.39E+02</td>
<td>3800</td>
<td>0.69</td>
<td>750</td>
<td>[18]</td>
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<tr>
<td>$^3$He/$^4$He</td>
<td>7.72E-07</td>
<td>2.74E-10</td>
<td>1.40E+02</td>
<td>3800</td>
<td>0.69</td>
<td>0.07</td>
<td>[18]</td>
</tr>
<tr>
<td>Ar</td>
<td>1.43E-04</td>
<td>7.60E-04</td>
<td>3.88E+08</td>
<td>64.38</td>
<td>32,000</td>
<td>13,400</td>
<td>[18]</td>
</tr>
<tr>
<td>$^{36}$Ar</td>
<td>3.91E-07</td>
<td>2.09E-06</td>
<td>1.07E+06</td>
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<td>1066</td>
<td>450</td>
<td>Shell</td>
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<tr>
<td>$^{36}$Ar/$^{40}$Ar</td>
<td>3.79E-03</td>
<td>1.17E-06</td>
<td>5.97E+05</td>
<td>1000</td>
<td>597</td>
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<tr>
<td>Kr</td>
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<td>1.84E+04</td>
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<td>3.16</td>
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<td>$^{86}$Kr</td>
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<td>5.55E-09</td>
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<td>10.16</td>
<td>[18]</td>
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<td>$^{86}$Kr/$^{84}$Kr</td>
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<td>1.08E-08</td>
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<td>1490</td>
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<tr>
<td>Xe</td>
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<td>Condition met inherently</td>
<td>340</td>
<td>-</td>
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<tr>
<td>$^{124}$Xe/$^{130}$Xe</td>
<td>8.48E-02</td>
<td>9.70E-11</td>
<td>4.95E+01</td>
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<td>1.84</td>
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<td>6.48E-11</td>
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<td>1.45E-09</td>
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<td>-</td>
<td>-</td>
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<td>Shell</td>
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<td>1.58E-08</td>
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<td>1.25</td>
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<td>Shell</td>
</tr>
<tr>
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<td>1.60E-08</td>
<td>8.16E+03</td>
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<td>17.96</td>
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<td>Shell</td>
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<tr>
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<td>6.33E-09</td>
<td>3.23E+03</td>
<td>4585</td>
<td>19.25</td>
<td>0.0044</td>
<td>[18]</td>
</tr>
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<td>2200</td>
<td>6.09</td>
<td>0.0014</td>
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</tbody>
</table>

4. Discussion

4.1. Inherent Distinctiveness

The phase-partitioning calculations show that inherited noble gas signatures of injected CO$_2$ are significantly distinct compared with all background signatures, given the presented analytical uncertainty and database (Table 3). This is further demonstrated when mixing He, Ar, Kr, and Xe concentrations in the CO$_2$ with the respective gas concentrations in natural gas, atmosphere, and gas hydrates (yellow lines in Figure 2).
Figure 2. Mixing of the equilibrated CO$_2$ with background concentrations in natural gas, atmosphere, and gas hydrates in (cm$^3$/cm$^3$). For seawater, gas-specific solubilities have to be applied. The range of equilibrated CO$_2$ is the orange line, with the values as stated in Table 3. Mixing is calculated for the lower of these values. Background concentrations are shown with error bars (blue)—being dominated by the natural variability for natural gas and gas hydrates, and the analytical uncertainty for the atmosphere.

In the case of equilibrated CO$_2$ mixing with natural gas, case 2 in Figure 1, Xe is most distinct and would allow identification of injected CO$_2$ in gas mixtures at percentages of CO$_2$ as low as 3% (Figure 2). For He, Ar and Kr distinctiveness is worse. This is mainly caused by the natural variability in the natural gas (blue error bars in Figure 2). Further,
for Ar and Kr, the range of the concentrations in the equilibrated CO$_2$ is relatively large, resulting from the concentration range in the captured CO$_2$ (orange line in Figure 2).

Even though different to the natural gas, the equilibrated signature of injected CO$_2$ exhibits some characteristics of a radiogenic signature. Primarily, high He makes the signature very distinct compared to the atmosphere and gas hydrates (Figure 2). For mixing of injected and equilibrated CO$_2$ with the atmosphere, all gases are significantly different (note the logarithmic scale for Ar and Kr in the second row of Figure 2). He will be most reliable since He concentration is the only gas that is higher in the equilibrated CO$_2$ than in the atmosphere.

For seawater, the assumption can be made that the distinctiveness will be the same as for an atmospheric signature, considering that the same Henry’s constants would be applied to the dissolution of noble gases for both air and leaking CO$_2$.

The very distinct depletion of He in gas hydrates allows unambiguous identification of an injected and equilibrated CO$_2$ component in a mixture of these two components (note the logarithmic scale for He in the third row of Figure 2). The other gases exhibit large natural variability and therefore do not add distinctiveness.

4.2. Limitations of the Method

4.2.1. Calculation Method

In the calculations of phase partitioning, we calculated the specific solubility in a CO$_2$-H$_2$O-system using the correction term of [20] (Equation (2)). For deriving the formation water concentrations of noble gases in the storage reservoir—currently not directly measured—we assumed equilibrium with the analyzed natural gas. Here, we used solubilities for a high-pressure H$_2$O-system, i.e., not applying $\kappa$ in Equation (2). Such specific correction terms for the noble gas solubility in a CH$_4$-CO$_2$-system have yet to be derived. The CO$_2$-solubility correction, i.e., the preferential partitioning into the CO$_2$ phase, is exactly what makes Xe values in particular differ significantly (∼500% in Table 3). Deriving correction terms for a high-pressure CH$_4$-CO$_2$-system may alter the concentrations and lead to the signatures that the injected CO$_2$ adopts to be less or more distinct. Further, impurities in the CO$_2$ injection stream could impact the applicability of solubilities for a pure CO$_2$-H$_2$O-system; however, such deviations are anticipated to be minor given the relatively pure nature of injected CO$_2$. For example, the injected CO$_2$ at the Sleipner storage site has a CH$_4$ constituent of only 1–2% [22].

We further assumed that phase partitioning is the only process influencing the signature before mixing. Mixing with native formation gases, such as minor CH$_4$ accumulations, could alter the mixtures significantly, i.e., including a signature of natural gas. Thereby, the injected CO$_2$ would be more similar to natural gas, reducing distinctiveness towards it. The inclusion of the radiogenic signature from native formation gases would, however, further increase distinctiveness towards shallow signatures.

4.2.2. Leakage Scenarios

Reaching initial equilibrium with the formation water may depend on the type of leakage. The later a leak occurs after injection, the more likely equilibrium is achieved. For “catastrophic” leakages—e.g., at the injection well—the CO$_2$ could stay in its depleted state (case (1) in Figure 1). Such a leak can be detected based on a suit of other geophysical and geochemical tools that are installed along the well [1]. The depleted noble gas concentrations could, however, be used for leakage attribution. For other advective leakage types—e.g., cases 2, 3, 4, and 5 in Figure 1—the CO$_2$ will likely have equilibrated during prior migration in the storage reservoir making our calculations applicable. This could also depend on leakage rates. Those have been constrained for natural CO$_2$ analogues [43], but it is unclear how that translates to other settings. Leakage for such fields showed that, for example, the elevated He concentrations are maintained up until the surface [44].

Our calculations assume full equilibration of the injected CO$_2$ with formation water. In [13], we argued that equilibrium is likely to be reached, considering that at the front
of a moving CO₂-plume, the concentration gradients between the two phases, formation water, and CO₂ are high. For a dedicated storage case, such a behavior was, for example, observed at Aquistore, where the injected CO₂ had adopted a radiogenic signature one year after injection [11]. In contrast, the formation water of a system that is continuously flushed may become depleted in the noble gases over time, leading to domination of the depleted, original signature of the captured CO₂. Such a behavior was observed for the specific case of the Cranfield EOR field [16]. Although, in that case, the presence of a hydrocarbon phase meant low residual water saturation and a limited volume of water in contact with the CO₂ phase, leading to more rapid noble gas stripping than would be observed for CO₂ storage in saline aquifers.

In the phase-partitioning and subsequent mixing calculations, the dissolution of CO₂ into the brine over time is not taken into account. This effect would tend to increase the residual noble gas concentrations and reduce the gas water volume ratio, $V_g/V_{H_2O}$. Our calculations do not evaluate the migration of a dissolved CO₂ phase. It is known (from natural analogues) that ex-solution of dissolved CO₂ tends to lead to quantitative partitioning of the dissolved noble gas load into the CO₂ phase and, thus, the reservoir signature is likely to be inherited [44]. However, the complexities of multistage dissolution-ex-solution have not been assessed in this study, and further investigation of the preservation of the noble gas signature in such systems is merited.

4.2.3. Mixing Calculations

Mixing calculations of end-members will not allow the complete exclusion of a minor contribution of one end-member to a mixture, due to analytical uncertainty typically being an unsystematic error in the sample measurement [45]. Further, noble gases are found in every natural system and exhibit significant natural concentration variability, thereby introducing uncertainty in end-member calculations. The use of element or isotope ratios helps to reduce this uncertainty in mixing calculations because of the covariance of element or isotope concentrations, which removes dilution or concentration effects that otherwise create variability in absolute element concentrations. Further, the calculations become more constrained the more accurately the end-member compositions are known from baseline characterization of the storage system or via measurements of the captured and injected CO₂. For example, above hydrocarbon systems, methane releases may imprint fluids in seafloor sediments with a radiogenic signature, reducing the distinctiveness of leaking CO₂ into the surrounding environment (Figure 1). Further, the gas hydrates composition is only known for the element and not the isotopic ratios, resulting in an incomplete description of one possible end-member.

Despite these uncertainties, noble gas signatures will deliver strong evidence when analyzing the fluids of a potential leak, as our calculations have shown. In Figure 2, only one element was used. By combining the analysis of a number of isotopes or ratios, the vector of the mixing line representing possible solutions to the mixing calculation increases; thereby, measurement errors become less influential. This is exemplified for a mixing with natural gas, where using both Kr and Xe increases differentiability (Figure 3a). Meanwhile, for atmosphere, the inclusion of the $^{3}$He/$^{4}$He ratio has the same effect. Besides, noble gases such as this could include additional geochemical constraints, such as the stable isotopic composition of the CO₂. For example, injected CO₂ with an atmospheric $^{13}$C isotope signature provides an additional distinct parameter with which to discriminate the CO₂ from natural gas [6].
A further approach to reduce uncertainty introduced by natural variability is the addition of artificial tracers, e.g., SF₆ or PFCs. Since these are not present in the natural environment, uncertainties are reduced to the analytical uncertainty alone. Partly, however, these tracers have other disadvantages, such as toxicity or functionality as a greenhouse gas [18]. For details and examples, the reader may be referred to [15,18], where the addition of artificial trace gases is also analyzed from a cost perspective.

4.3. Tracer Addition

Tracer addition could help to manage the limitations of the calculation, as described in the previous Section 4.2, i.e., increase the altering of the injected CO₂ signature of a mixture against the natural variability and the analytical uncertainty. The additional distinction through the addition is depicted for Xe and $^{126}$Xe/$^{130}$Xe in Figure 4.

The necessary concentrations to be added into the injection stream are lowest for $^3$He and $^{124,126}$Xe isotopes with concentrations of $\sim$1E-10 cm$^3_{\text{STP}}$/cm$^3_{\text{STP}}$, or of 50–100 l/MtCO₂. These He and Xe isotopes are among the most expensive in terms of cost per liter, at 1000–10,000 $\text{US}/l$ (Table 4). Despite this, due to the small amounts required, these would be the cheapest tracers to add with a cost of $\sim$0.7 $\text{US}/t$CO₂ (Table 4).

Xe tracer addition appears to be the most suitable noble gas tracer in CO₂ storage, as both lab and field experiments suggest that Xe could even have an early warning function, in contrast to He [46,47]. This is due to the molecule size, i.e., allowing He to enter smaller, dead-end pores and thereby getting retarded. Adding $^3$He would also shift the $^3$He/$^4$He ratio towards atmospheric values, thereby interfering with the distinction to atmospheric signatures. Further, $^3$He has seen a price increase and supply shortage in the last decades [48] that may make its application unfeasible within a few decades. In contrast, availability of Xe isotopes is estimated to be sufficient [15].
Figure 4. The impact of tracer addition: The signatures of natural gas, equilibrated CO$_2$, and atmosphere in comparison to the equilibrated CO$_2$ with tracer addition. The mean concentration is surrounded by the respective error ellipses.

The cheapest tracer addition would cost \( \sim 0.7 \) $US/tCO$_2$, given our assumptions. In comparison to the cost of noble gas tracer addition derived with the method in [18], i.e., “10 times the detectable perturbation above background levels”, the cost estimates derived in our study are typically higher (Table 4). Even though the same isotopes are assigned the lowest cost in the derivation of [18], the cost presented here is several orders of magnitude more expensive, e.g., for $^{126}$Xe 0.00014 $US/tCO$_2$ after the method in [18] vs. 0.76 $US/tCO$_2$ for the derivation presented here. Our results are also higher, but more in line with calculations for detection in soil gas made in [15], which resulted in a cost of \( \sim 0.1 \) $US/tCO$_2$ for both $^3$He and $^{136}$Xe.

Our criterion for distinctiveness of CO$_2$ in leakage detection schemes and considering the phase partitioning estimates leads to higher volumes of tracers needed and increased costs compared to previous studies [15,18]. However, defining when a signature is sufficiently distinct is, to a certain degree, a subjective choice. It appears that this will have to be defined by decision makers, i.e., how soon and how certain one wants to know a leakage/contamination in a trade-off between cost and risk.

In comparison to total operational costs of a CCS project, \( \sim 0.7 \) $US/tCO$_2$ is a low cost, for example, compared to a CO$_2$ capture cost of 15–120 $US/tCO$_2$ [49]. However, considering that the current cost of the whole monitoring scheme of running sites is in the order of 1–4 $US/a [1], the added cost would be a significant contribution to the total monitoring cost, especially considering the generally low leakage likelihood [50]. Further, on the one hand the estimated cost does not yet include sampling, analysis, and evaluation costs, on the other hand there may be a price reduction through bulk orders of the respective tracer gas.

4.4. Technical Aspects of Tracer Addition

The addition of a noble gas tracer into a CO$_2$ injection stream represents an impurity, which potentially changes the thermodynamic properties of the CO$_2$. Even though indi-
vidual for each CO$_2$-tracer mixture, depending on the type and amount of an impurity, typically, the critical pressure is increased [51]. This implies changes of the transport properties and could affect transportation and injection setup, and is thus likely to add costs. Impurities also tend to decrease the viscosity of the fluid mixture [51], reducing migration properties of the CO$_2$ in the reservoir. Therefore, it seems desirable to have the lowest concentrations of the added tracer as possible. For the trace isotopes we derive as most economical, i.e., elemental He and Ar being excluded, the added impurity is very low (below 1E-06 vol%) compared to those typically found after the CO$_2$ capture process, such as N$_2$, O$_2$, H$_2$S, and H$_2$O which can collectively make up 1–5 vol%) [52].

On another technical note, achieving continuous tracer concentrations in a gas stream as low as 1E-11–1E-10 cm$^3$STP/cm$^3$STP is not trivial. Nimz and Hudson [15] outline how a side-track with flow regulation and an additional compressor could be used. It is also suggested that the low volumes of the gas necessary, e.g., tenths of liter per year, could better be applied through prior mixture with CO$_2$ as the carrier gas, i.e., stepwise dilution [15]. It might also be considered to have pulsed injection patterns and assume effective distribution after injection.

4.5. Noble Gases in Monitoring Schemes

In case of an observed or suspected CO$_2$ anomaly or leakage, noble gases are very likely to give an indication of the CO$_2$ source due to the distinct signatures in the environment. Therefore, noble gases’ main application will—also offshore—lie in leakage attribution as it has in the past, e.g., [14].

In contrast, high-frequency monitoring of a storage site for leakage detection, does seem less attractive given that noble gas analysis for high sensitivity, precision, and accuracy—i.e., full isotopic analysis including the least abundant isotopes—currently has to be conducted using a lab-based multicollector mass spectrometers with a cost of up to several thousand $US per sample. Further, collecting and transporting a sample to the lab does introduce a significant time lag between sampling and interpreting results.

However, in recent years, in situ, continuous, and remotely controllable noble gas analysers have been developed [53,54]. Currently these techniques cannot deliver full isotopic resolution, typically covering the most abundant elements of a noble gas, e.g., $^4$He, but not $^3$He. Further, the limit of quantification is usually in the ppm range. Still, with on-site measurements, not only leakage attribution but also leakage detection may become possible. Especially, the elevated concentrations of He (see Table 3) could be targeted with a mobile mass spectrometer when sampling offshore seabed fluid releases, as demonstrated in the vicinity of black smokers [53]. Both water and gas analysis could be conducted on board a ship to scan if elevated He concentrations indicate leaking CO$_2$ from a deep CO$_2$ source with a radiogenic signature. Simultaneous measurement of O$_2$, N$_2$, CH$_4$, and CO$_2$, which the instrument is capable of, add valuable information.

In contrast to the previous case, the application of these technologies for the monitoring of the production stream at an operating gas field is limited. Considering our results, e.g., for the mixture of CO$_2$ and natural gas, where Xe is the most distinctive gas, the portable analysis systems do currently not deliver sufficient sensitivity for detection of Xe concentrations of 1E-09 cm$^3$STP/cm$^3$STP, even with tracer addition (see Table 3). Further advances of these technologies may allow for wider applicability and for continuous, remote leakage detection.

5. Conclusions

The partitioning of the formation water noble gas signature into injected CO$_2$ is destined to allow the differentiation from natural geological fluids or gases due to the specific fractionation through the phase partitioning in a CO$_2$-H$_2$O-system. Meanwhile, there are uncertainties attached to the distinction towards natural gas, i.e., unknown solubilities in a CH$_4$-H$_2$O-system, the difference between the adopted radiogenic signature and that of shallow signatures is very pronounced.
The presented approach also allows us to evaluate the active addition of a noble gas tracer to the injection stream with the goal of maintaining a significantly different noble gas signature after the equilibration. Xe isotopes are the most relevant choice for noble gas tracer addition, since they have the lowest concentrations in the background (see Table 1) and do not impact the main features of the radiogenic signature, in contrast to He. Such an addition will increase identifiability but adds significant cost to a monitoring scheme of a storage project.

Further, baseline characterization is fundamental, and even though we use relevant analogues in this study, each storage site and CO₂ capture plant has to be expected to be different. For the inherent recognition through Xe, specific fractionation patterns in captured CO₂, such as elevated Kr and Xe at oxyfuel plants, could significantly increase detectability. Further, formation water concentrations are only indirectly derived.

Considering that CCS is currently only feasible with government subsidies, adding cost to CCS projects endangers its large-scale application [3]. Therefore, implementation of a noble gas tracer addition and other MMV methods will, on the one hand, depend on the legal framework regarding long-term liability of safe containment, and on the other hand, on widespread and meaningful implementation of carbon pricing options, such as carbon taxes or a carbon trading system. Further, (i) with the involved techniques maturing over the past and coming decades, uncertainties attached to such advanced green investments should reduce and thereby improve the green finance framework [55]; (ii) and increased MMV expenditures could increase the environmental reputation of the involved entities [55].

The planned, continuous MMV program in the Norwegian storage project ‘Longship’ currently does not include noble gases [27]. We believe that noble gas sampling should be considered during routine environmental surveys and should definitely be the content of ‘triggered’ environmental monitoring surveys. The content of such a triggered environmental monitoring survey has yet to be defined.

This work is another step towards upscaling and realizing the utilization of noble gas tracers, quantitatively and qualitatively describing how noble gases can be applied, which behavior is to be expected, and how they may be useful in deploying CCS. Implementation in case-specific leakage models, including secondary processes such as dissolution and mineralization of CO₂ combined with advances in defining solubilities of noble gases in the relevant systems and further developments in field-applicable measurement devices, will constrain our findings further and allow for wider application of noble gases in monitoring schemes for CO₂ storage sites, enhancing detectability and source attribution.


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