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Towards improved monitoring of offshore carbon storage: A real-world field experiment detecting a controlled sub-seafloor CO₂ release

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ABSTRACT

Carbon capture and storage (CCS) is a key technology to reduce carbon dioxide (CO₂) emissions from industrial processes in a feasible, substantial, and timely manner. For geological CO₂ storage to be safe, reliable, and accepted by society, robust strategies for CO₂ leakage detection, quantification and management are crucial. The STEMM-CCS (Strategies for Environmental Monitoring of Marine Carbon Capture and Storage) project aimed to provide techniques and understanding to enable and inform cost-effective monitoring of CCS sites in the marine environment. A controlled CO₂ release experiment was carried out in the central North Sea, designed to mimic an unintended emission of CO₂ from a subsurface CO₂ storage site to the seafloor. A total of 675 kg of CO₂ were released into the shallow sediments (~3 m below seafloor), at flow rates between 6 and 143 kg/d. A combination of novel techniques, adapted versions of existing techniques, and well-proven standard techniques were used to...
detected, characterised and quantified gaseous and dissolved CO₂ in the sediments and the overlying seawater. This paper provides an overview of this ambitious field experiment. We describe the preparatory work prior to the release experiment, the experimental layout and procedures, the methods tested, and summarise the main results and the lessons learnt.

1. Introduction

Human activities, including fossil fuel burning, land-use changes, and cement manufacture, have caused the atmospheric carbon dioxide (CO₂) concentration to rise from a pre-industrial level of 277 parts per million (ppm) to a current level of ~412 ppm in 2020 (e.g., Friedlingstein et al., 2019; Drügken and Tans, 2020). This atmospheric accumulation of anthropogenic CO₂ has been linked to the rise of the global mean temperature, presently approximately 1.0 °C above pre-industrial levels (IPCC, 2018). The United Nations Framework Convention on Climate Change (UNFCCC) agreed to take action and to keep global warming below 1.5–2 °C above pre-industrial levels (UNFCCC, 2015; IPCC, 2014, 2018). CO₂ capture and storage (CCS) is a key technology in many of the recent mitigation scenarios that would meet this goal (IPCC, 2005, 2018). CCS involves the capture of CO₂ – principally from large point sources such as industrial power plants – and its injection into geological storage formations such as deep saline aquifers or depleted oil and gas reservoirs for permanent storage (IPCC, 2005). Compared to other CO₂ mitigation strategies, such as improving energy efficiency and use of renewable energy, the crucial benefit of CCS lies in its potential to significantly (at giga-tonne scale) and rapidly reduce CO₂ emissions while making use of infrastructure that already exists for oil and gas production (IPCC, 2005).

It is estimated that the majority of Western Europe’s potential CO₂ storage capacity is located offshore (IEAGHG, 2008; Vangkilde-Pederson, 2009). There are a small number of active (Sleipner, North Sea, Norway; Snøhvit, Barents Sea, Norway) and completed (K12-B, North Sea, Netherlands) offshore CO₂ injection projects in Europe that provide confidence in the performance of offshore gas injection and storage (Vandeweijer et al., 2011; Hansen et al., 2015; Van der Meer, 2013; Furre et al., 2017; Ringrose and Meckel, 2019). These and previous CO₂ storage demonstration projects have improved our understanding of the strengths and development needs of various monitoring approaches (IEAGHG, 2006, 2012, 2015; Dixon and Romanak, 2015; Jenkins et al., 2015).

Leakage of injected CO₂ from well-selected, designed and managed geological storage sites back into the atmosphere is generally considered unlikely (IPCC, 2005). However, robust strategies for leak detection and management of offshore CCS projects are a regulatory requirement to comply with international marine legislation (e.g., EU CCS Directive (EU, 2009), London Protocol (IMO, 2006) and OSPAR Convention (OSPAR, 2007)), and are essential for public acceptance of CCS as a safe and reliable technology for the long-term mitigation of elevated anthropogenic CO₂ emissions (e.g., Mabon et al., 2014, 2015, 2017).

Previous research projects, including ECO2 (Sub-seabed CO₂ Storage: Impact on Marine Ecosystems), QICS (Quantifying and Monitoring Potential Ecosystem of Geological Carbon Storage) and ETI MMV (Energy Technologies Institute Measurement, Monitoring and Verification of CO₂ Storage), have advanced technologies for detection of leakage at the seafloor and have improved knowledge of the local impacts that CO₂ leakage may have on marine ecosystems (Blackford et al., 2014, 2015; Jones et al., 2015; Taylor et al., 2015). Building on this, an overall aim of the Strategies for Environmental Monitoring of Marine Carbon Capture and Storage (STEMM-CCS) project was to deliver new approaches, methodologies and tools for cost-effective environmental monitoring and leakage quantification at offshore CO₂ storage sites.

The specific aims of the STEMM-CCS project were to:

- develop new tools and techniques for the monitoring, quantification, and assessment of a potential CO₂ leakage from an offshore CO₂ storage site;
- demonstrate the suitability of these tools and techniques in a representative field experiment mimicking a CO₂ leakage from a subsea CO₂ storage site;
- demonstrate a CCS-specific implementation of an ecological baseline measurement, incorporating physical, chemical and biological variability;
- understand the role of natural geological and artificial features as potential pathways for subsurface CO₂ migration;
- develop and evaluate modelling tools to inform monitoring strategies and provide a framework to interpret the experimental results, and translate them to other settings.

To achieve these aims, the STEMM-CCS project conducted a unique CO₂ release experiment designed to mimic a CO₂ leakage from an offshore CO₂ storage site and to demonstrate and evaluate existing and new approaches for characterising and quantifying this release. To enable this, a controlled mixture of CO₂ and tracer gases was released into the sediments at 3 m below the seafloor. A range of novel and standard methods were applied to detect and characterise the release of CO₂ (Fig. 1). The field experiment involved two research vessels and was carried out in the British sector of the central North Sea during 27 April - 27 May 2019.

This paper provides a detailed overview of this field experiment and an introduction to the STEMM-CCS special issue by summarising (i) the preparatory and baseline work that was conducted prior to the release experiment, (ii) the experimental set up and the methods evaluated, and (iii) the main results and the lessons learnt. For detailed results, the reader is referred to the specific papers published as part of this special issue.

1.1. Experiment location and setup

1.1.1. Experiment location

The STEMM-CCS experimental CO₂ release was conducted within the surface sediments overlying the proposed Goldeneye CO₂ storage reservoir, a depleted gas condensate field situated offshore from Scotland in the Outer Moray Firth, in the British sector of the central North Sea (56°–60° N) (Dean and Tucker, 2017) (Fig. 2). Located in a sandstone formation of the Lower Cretaceous (Albian-Aptian) period, the reservoir is 10–15 km wide and 60–90 m thick (e.g., Law et al., 2000).

The Goldeneye reservoir was discovered in 1996 and produced 16 × 10⁹ m³ of gas and 23 million barrels of condensate after production by Shell commenced in 2004 (Wilson et al., 2005; Shell, 2015). After gas production ended in 2011, the depleted reservoir was identified as a potential CO₂ storage site for the Peterhead CCS demonstration project. This project was envisaged to capture CO₂ from the Peterhead power plant with subsequent injection of about 15 million tonnes of CO₂ into the depleted Goldeneye reservoir (Shell, 2015; Cotton et al., 2017; Dean and Tucker, 2017). The Peterhead CCS demonstration project was cancelled after the withdrawal of funding by the UK government in 2015 (Dean and Tucker, 2017). The associated scientific research projects STEMM-CCS and CHIMNEY went ahead nonetheless, to underpin the scientific and technological developments necessary for other or future subsea CO₂ storage work. The CHIMNEY project (Characterisation of Major Overburden Leakage Pathways above Sub-sea floor CO₂ Storage Reservoirs in the North Sea; https://www.southampton.ac.uk/oes/
eath/projects/chimney.page) developed new techniques to predict natural and anthropogenically-induced permeability of the reservoir overburden, including intensive studies of naturally occurring “seismic chimney” structures. While this paper focusses on the in-situ release experiment, the paper by Robinson et al. (2020) examines the work done on the experiment, the paper by Robinson et al. (2020) examines the work done on the experiment.

1.1.2. Permissions and legal considerations for the CO₂ release experiment

Any work in the Scottish sector of the North Sea requires permission from Marine Scotland, the statutory body of the Scottish Government responsible for the protection of Scotland’s coastal waters and sea. Acquiring the required authorisation for the STEMM-CCS experiment was complex because injecting CO₂ into the seabed could be viewed as ocean dumping. Following consultation with OSPAR representatives, the experiment was designated as placement of tracers (CO₂ was considered a tracer in this context) below the seabed. Following the submission of the application for the controlled CO₂ release experiment, there was a three-month public consultation period after which the licence for the work was published. In addition, Crown Estate Scotland conducted a spatial conflict check, because a commercial company had been granted a license for future storage of CO₂ in the Goldeneye reservoir. Crown Estate identified no conflicts. Consequently, since there was no storage of CO₂, no permanent presence, and no conflict with other users on the seabed, permission was granted. The permit allowed the limited release of up to 3 tonnes of CO₂ with associated tracers; we released just over 20 % of this. In addition, a model was developed and executed that showed the limited spatial and temporal impact the release would have on the environment, given the low quantity of gas released and the impact of dilution from tidal flushing. Prior to, and periodically during the release experiment, a trained marine mammal observer performed regular surveys from the ship. This was required due to concerns around the use of acoustic sensor-based systems.

1.1.3. Baseline

To differentiate with confidence between CO₂ leakage and any natural variability in the local environment, monitoring needs to be supported by a robust baseline (OSPAR, 2007; EU, 2009; IEAGHG, 2012, 2015), which provides a longer-term picture of the regional spatio-temporal variability. This is in contrast to “background measurements”, which refers here to samples taken at non-affected sites near the experimental release site during the experiment. STEMM-CCS therefore conducted baseline surveys to characterise the spatio-temporal variability of all relevant processes in the region and to provide a basis against which anomalies could be identified. Guided by knowledge gained from the precursor project QICS (Blackford et al., 2014), STEMM-CCS aimed to provide best practice optimisation for creating ecological and environmental baselines for offshore CO₂ storage sites. To this end, two baseline cruises to the Goldeneye site were coordinated by GEOMAR Helmholtz Centre for Ocean Research Kiel: cruise number POS518 in October 2017 and cruise number POS527 in August 2018, both aboard RV Poseidon (Table 1). The main goals were to deploy baseline landers for continuous physical and biogeochemical measurements prior to the release experiment and to take samples for further characterisation of the water column, the sediments and the benthic communities (Section 2). Additional background work was carried out prior to and throughout the CO₂ injection during the JC180 and POS534 cruises (Section 2).

1.2. Behaviour of CO₂ in the marine environment

When CO₂ is injected into marine sediments, part of it remains in the gas phase, where it either accumulates in gas pockets within the sediments or percolates through the sediments to the seabed where it emerges as bubbles that rapidly dissolve in the overlying seawater (e.g., McGinnis et al., 2011; Gros et al., 2019; Vielstädte et al., 2019). When CO₂ dissolves in seawater, it forms carbonic acid (H₂CO₃) that subsequently dissociates to form bicarbonate (HCO₃⁻) and carbonate (CO₃²⁻) ions. The total concentration of dissolved CO₂ in seawater is therefore given by the sum of the concentrations of H₂CO₃, HCO₃⁻, and CO₃²⁻, which is referred to as dissolved inorganic carbon (DIC) (Zeebe and Wolf-Gladrow, 2001). Some part of the CO₂ gas also dissolves and potentially reacts with the sediment pore waters (i.e., the water surrounding the solid sediment particles), increasing the DIC content of the pore waters. During the in-situ release experiment, a wide range of

Fig. 1. Schematic overview of the CO₂ release experiment and a selection of deployed detection methods.
methods was used to target these different phases, i.e., the gaseous and dissolved CO₂ in both the sediment pore water and the overlying seawater. These methods were used to detect, characterise, and quantify the released CO₂. Each method performed one or more tasks: detection of the CO₂ release; attribution of the release (distinguishing between CO₂ leaking from a reservoir and naturally emitted CO₂); or quantification of the release (estimating a flux or a mass flow rate) (Table 2).

1.3. Experimental design and overall implementation

The CO₂ release experiment involved two research vessels: the UK royal research ship RRS James Cook (cruise number JC180; Connelly, 2019; Table 1) and the German research vessel RV Poseidon (cruise number POSS34; Schmidt, 2019). It was carried out near the Goldeneye platform, at 120 m water depth, in the central North Sea (Fig. 2) between 27 April and 27 May 2019. Two underwater vehicles were also used: the NOC work-class ROV (Remotely Operated Vehicle) Isis (German et al., 2003; Fig. 3a) was used to carry equipment and maneuver it on the seafloor and an AUV (Autonomous Underwater Vehicle) Gavia (Teledyne Gavia, Iceland; German et al., 2003; Fig. 3b) was used for surveying. The CO₂ release equipment and these vehicles were deployed from the larger vessel RRS James Cook.

The release experiment was designed to enable all of the detection techniques (Table 2) to be tested and their sensitivity to be quantified over a range of gas release flow rates. This was achieved by deploying as much of the seafloor-based equipment as possible around the centre of the experiment site (defined as the point on the seafloor above the end of the inserted gas injection pipe). An overview of the experiment setup is shown in Fig. 1; the relative locations of all equipment and timing of deployments are shown in Fig. 4. Most of the equipment was deployed by the ROV Isis.

For safe operation of the ROV and maneuverability at the experimental site, equipment was only deployed outside the ROV landing zone (Fig. 4). Equipment with limited battery life was recovered and redeployed by the ROV every few days over the course of the experiment, with like-for-like replacements minimising gaps in the dataset. Equipment that had to be sequentially moved in a transect across the release site (e.g., the sediment microprofiler and ROV chemical mapping surveys) was deployed on alternate nights in order to maximise the operational ROV dive time for each survey. Whenever possible an AUV survey was conducted over the experiment site before increasing the rate of CO₂ release, in order to characterise any build-up or changes of gaseous CO₂ within the sediment. For comparison, initial AUV surveys had been conducted over the site before deploying any equipment and following the insertion of the gas release pipe. An AUV survey was also carried out after the experiment, and after all equipment was removed from the site.

The release of CO₂ was started at 15:19 (UTC time format used throughout the manuscript) on 11 May 2019, thus 00:00 on 11 May 2019 is considered the start of day 0 of the experiment. The gas flow rate was initially set to 6 kg/d (equivalent to the lowest achievable flow rate of 2 normal L/min; “normal” conditions are defined here as 0 °C and 1.013 bar). The CO₂ flow rate was sequentially increased over the duration of the experiment to a maximum of 143 kg/d (equivalent to a flow rate of 50 normal L/min) (Fig. 4), adding up to a total release of 675 kg of CO₂ over the course of the experiment. In most instances each flow rate was maintained for a minimum of 2 days to enable all different types of equipment to be tested under those conditions. The only exception to this was the flow rate of 14 kg/d, which was only maintained for 15 h to check that the sediments and operational setup were able to accommodate changes in CO₂ flow rate (i.e., that there was no abrupt release of gas from within the sediment).

When the CO₂ was first turned on, bubbles were observed emerging from the sediments as soon as the ROV had travelled from the CO₂ rig to the release site (within 30 min). At the lowest flow rate, three separate bubble streams were observed, and further bubble streams formed at higher flow rates. The dissolution of bubbles within the overlying water column was observed using the ROV. No bubbles were visible with the ROV camera at >8 m above the seabed.

The CO₂ injection was turned off at 11:17 on 22 May 2019, with all infrastructure recovered from the seafloor onto the RRS James Cook over the following 3 days. Sediment cores were taken by the RV Poseidon from the release site during this interval, before final ROV and AUV surveys were carried out.

2. Baseline site characterisation

To characterise the natural variability at the study site, the chemical,
Table 1
Research cruises for baseline and main experiments of the STEMM-CCS project.

<table>
<thead>
<tr>
<th>Vessel</th>
<th>Cruise</th>
<th>Date</th>
<th>Purpose</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>RV Maria S. Merian</td>
<td>MSM63</td>
<td>May 2017</td>
<td>Baseline: bathymetry and subsurface imaging</td>
<td>Berndt et al. (2017)</td>
</tr>
<tr>
<td>RV Poseidon</td>
<td>POS518</td>
<td>October 2017</td>
<td>Baseline: water chemistry, chemical and physical properties of the sediments, benthic chamber incubations (in-situ)</td>
<td>Linke and Haeckel (2018)</td>
</tr>
<tr>
<td>RV Poseidon</td>
<td>POS527</td>
<td>August 2018</td>
<td>Baseline: water chemistry, chemical and physical properties of the sediments, benthic biology, microbial activity</td>
<td>Achterberg and Esposito (2018)</td>
</tr>
<tr>
<td>RV Maria S. Merian</td>
<td>MSM78</td>
<td>October 2018</td>
<td>Baseline: bathymetry and subsurface imaging</td>
<td>Karstens et al. (2019)</td>
</tr>
</tbody>
</table>

biological, and physical properties of the area were determined prior to the CO2 release experiment (Section 1.1.3, Table 1). These baseline data were used to differentiate between the effect of the released CO2 and the variability caused by natural phenomena or unrelated human activity. Customised sediment and hydrodynamic-chemical models of the Goldeneau site were developed to understand how the experimental CO2 release would manifest in shallow sediments and water column.

2.1. Water column characteristics and chemistry

The natural dynamics of the physical and biogeochemical parameters of the water column were examined by discrete water column sampling (Section 2.1.1) and by baseline landers for continuous physical and biogeochemical measurements (Section 2.2.1).

2.1.1. Discrete water sampling

Water column sampling for the characterisation of physical background conditions was conducted during POS518 (October 2017) and POS527 (August 2018) (Table 1; Achterberg and Esposito, 2018; Linke and Haeckel, 2018). Environmental baseline sampling was undertaken in an approximately 10 km by 20 km area over the Goldeneu complex with the Goldeneu platform (58° 0’ 10.8” N, 0° 22’ 48” W) as centre point. Water was collected using Niskin bottles located on a rosette frame equipped with a conductivity, temperature, depth (CTD) probe. Concentrations of dissolved oxygen (O2), dissolved organic and inorganic carbon (respectively, DOC and DIC), the stable carbon and nitrogen isotope composition of DIC (δ13C and δ15N) and the oxygen isotopic composition of seawater (δ18O) were established after collection and analysis by standard procedures (e.g., Murphy and Riley, 1962; Clayton and Byrne, 1993; Grasshoff et al., 1999; Dickson et al., 2007; Hansen and Koroleff, 2007; Aßmann et al., 2011). These data were compared to water column chemistry data available on the GLODAP and Cefas databases (latitude-longitude box of 56° N, 2° W to 59° N, 2° E) (GLODAP Reference Group, 2020; Cefas Data Hub, 2020) to broaden the duration and seasonal coverage of the baseline data.

To study baseline O2, nutrient and CO2 fluxes across the sediment-water interface from in-situ incubations, two short-term deployments of the Biogeochemical Observatory (BIGO, Sommer et al., 2016) were conducted during cruise POS518. These incubation chamber landers are similar to the benthic chambers used during the CO2 release experiment (Section 4.3.2).

2.1.2. Landers for autonomous collection of in-situ data

To obtain baseline data over a longer time period and with higher temporal resolution, baseline landers equipped with autonomous in-situ sensors were deployed to gather continuous physical and biogeochemical baseline data. A long-term seafloor baseline lander was deployed prior to the main CO2 release experiment (Develogic GmbH Subsea Systems, Hamburg, Germany). It combined commercially available sensors and newly developed technology for autonomous gathering of in-situ biogeochemical and hydrodynamic baseline data. The lander was deployed during expedition POS518 on 16 October 2017 approximately 100 m north of the actual injection site (Linke and Haeckel, 2018) at 120 m water depth. All data were logged centrally on the lander and mirrored to a number of expendable pop-up data pods. The lander was programmed to release a pod to the surface every 3 months to relay collected data via Iridium satellite. The pop-up system did not work well but the lander was recovered during expedition JC180 in May 2019 and several months of data were found to have been recorded for the majority of the sensors.

As it was not possible to recover the Develogic long-term lander before the release experiment, additional landers for experimental background measurements were prepared by NOC and GEOMAR Helmholtz Centre for Ocean Research Kiel and were deployed for the duration of the release experiment. The landers were positioned approximately 350 m southeast and northeast of the experiment site respectively (see Fig. 4), sufficiently far to provide unperturbed background data. These landers provided measurements of physical (temperature, salinity and water currents) and biogeochemical (O2, carbonate chemistry and nutrients) parameters of the bottom part of the water column, with some parameters measured at high resolution (up to 0.1 Hz). With automated data collection, the three observational systems were able to provide a sound characterisation of the marine environment at the experiment site.

2.2. Baseline environmental study

The environmental baseline was assessed to establish conditions prior to the experimental work, to distinguish environmental impacts as a result of CO2 storage from those produced by other factors. To this end, the sediment and pore water characteristics were determined (Section 2.2.1), the benthic community was characterised (Section 2.2.2), and the area was mapped (Sections 2.2.3 and 2.2.4) prior to the controlled CO2 release experiment.

2.2.1. Sediment coring

To understand the effects of CO2 injection on sediment geochemistry, baseline values for relevant parameters, as well as their spatio-temporal variability, were determined. Baseline data were collected during the baseline cruises POS518 and POS527 to the study site (Linke and Haeckel, 2018; Achterberg and Esposito, 2018) (Table 1).

Sediment cores were obtained using push, gravity and multi corers and sub-sampled for biogeochemical analyses of the pore waters (nutrients, cations, anions, TA, DIC, δ13C and δ18O) and the solid phase (porosity, particle size, TOC, TON, δ13C of organic and inorganic carbon, chemical composition).

2.2.2. Benthic biology

Benthic organisms living on or in sediments can affect the sediment structure and biogeochemical processes locally as well as over larger
Table 2 (continued)

<table>
<thead>
<tr>
<th>Measurement target</th>
<th>Technique / method</th>
<th>Objective (section in paper)</th>
<th>Detection (D), Attribution (A), Quantification (Q)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Benthic chambers</td>
<td></td>
<td></td>
<td>DA</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Benthic fluxes of DIC, total oxygen uptake, nutrients, and other parameters (Section 4.3.1)</td>
</tr>
<tr>
<td>Eddy covariance</td>
<td></td>
<td></td>
<td>DA</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>Quantify vertical flux and total DIC content seawater in the vicinity of bubble streams with point measurements of current and pH (Section 4.3.3)</td>
</tr>
<tr>
<td>Chemical gradient measurements</td>
<td></td>
<td></td>
<td>DA</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Quantify vertical flux and total DIC content seawater in the vicinity of bubble streams with lab-on-chip sensors (Section 4.3.4)</td>
</tr>
<tr>
<td>Chemical mapping with ROV</td>
<td></td>
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<td>DA</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Mapping spatial extent and DIC content seawater in the vicinity of bubble streams with real-time data (Section 4.3.5)</td>
</tr>
<tr>
<td>pH sensing on an AUV</td>
<td></td>
<td></td>
<td>D</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Mapping spatial extent and DIC content seawater in the vicinity of bubble streams (Section 4.3.6)</td>
</tr>
</tbody>
</table>

Note: Table 2 provides a summary of targeted phases, techniques, and objectives of the release experiment for the detection and/or attribution and/or quantification of the released CO₂.

**Table 2**

Summary of targeted phases, techniques and objectives of the release experiment for the detection and/or attribution and/or quantification of the released CO₂.

<table>
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<tr>
<th>Measurement target</th>
<th>Technique / method</th>
<th>Objective (section in paper)</th>
<th>Detection (D), Attribution (A), Quantification (Q)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Automated image collection and annotation for baseline and identification of CO₂-related changes in seafloor biota (Section 2.2.2)</td>
<td>D</td>
<td></td>
<td>DA</td>
</tr>
<tr>
<td>Collection and chemical analysis of sediments and pore water around bubble streams (Section 4.1.1)</td>
<td>DA</td>
<td></td>
<td>DA</td>
</tr>
<tr>
<td>In-situ profiles of temperature, pH, oxygen, and other parameters in the sediments (Section 4.1.2)</td>
<td>D</td>
<td></td>
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</tr>
<tr>
<td>pH and temperature measurement within the sediment pore water (Section 4.1.3)</td>
<td>D</td>
<td></td>
<td>DA</td>
</tr>
<tr>
<td>Measuring physical structure of seafloor and imaging of potential gas pockets (Section 4.1.4)</td>
<td>DQ</td>
<td></td>
<td>DA</td>
</tr>
<tr>
<td>Direct optical measurement of bubble size and gas flow rate of a bubble stream (Section 4.2.1)</td>
<td>DQ</td>
<td></td>
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</tr>
<tr>
<td>Direct measurement of gas flow rate of a bubble stream (Section 4.2.2)</td>
<td>DQ</td>
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<tr>
<td>Collection of gas bubbles from a bubble stream for tracer analysis (Section 4.2.2)</td>
<td>DQ</td>
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<tr>
<td>Direct acoustic measurements of bubble size and gas flow rate (Section 4.2.3)</td>
<td>DQ</td>
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<tr>
<td>Location of bubble stream sites from the ship and AUV (Section 4.2.4)</td>
<td>DQ</td>
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<tr>
<td>Optical observation of bubble streams and measurement of dissolved CO₂ and related seawater parameters by using a towed and video-guided CTD probe (Section 4.3.1)</td>
<td>D</td>
<td></td>
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<tr>
<td>Chemical analysis of sampled water (Section 4.3.1)</td>
<td>D</td>
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<tr>
<td>Pumping of seawater in the vicinity of bubble streams directly to the ship for</td>
<td>D</td>
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</tbody>
</table>

**Note:** The categorisation (D, A, Q) is specific to this study and does not preclude use of the method in other ways. Shaded cells represent methods/analyses performed in-situ, and unshaded cells represent methods that relied on sample collection and subsequent analysis on board the ship or onshore.

Additional areas (Van Hoey et al., 2008; Montserrat et al., 2009). Additionally, the release of CO₂ may affect the benthic fauna near or at a release site. Sampling for benthic macrofauna (body size 0.5–10 mm) of the Goldeneye area was carried out in August 2018 during the RV Poseidon cruise POS527 using a winch operated box-corer (50 cm width × 50 cm depth) (Achterberg and Esposito, 2018). A total of 76 samples were collected from 75 sites selected according to sediment type, fishing pressure and pockmark location. All fauna were sorted and identified to the lowest possible taxonomic level under a stereomicroscope, with individuals and species abundances per sample recorded. Identification and quantification of interactive effects of sediment type, fishing pressure and distance from pockmark on community abundance and diversity measures were determined.

Benthic megafauna (>10 mm; sensu Grassle et al., 1975) at the experimental site were assessed using seabed photography before the experiment started. Photographic surveys were conducted with the Gavia AUV during the JC180 cruise; one survey was conducted at the experimental site and another at a site distant from it. Seabed photographs were captured using a GRAS-145M-C camera with a Tamron × 50 cm lens mounted to the AUV (Connelly, 2019). The camera captured photographs at a temporal frequency of 1.875 frames per second, a resolution of 1280 × 960 pixels, from a target altitude of 2 m above the seafloor. The total seafloor area represented, and the extent of overlap between photographs, were estimated using the camera specifications, location and altitude of the camera, and any overlapping photographs were removed from the analysis. The photographic
datasets used for analysis were comprised of 446 images representing 1053 m² of seabed from the experimental area, and 981 images representing 2366 m² seabed from the distant site. Megafaunal specimens were enumerated and classified to morphotypes using the BIIGLE 2.0 annotation software (Langenkämper et al., 2017) through a combination of manual annotation and implementation of the MAIA unsupervised machine learning algorithm (Zurowietz et al., 2018). Megafaunal density, diversity and community composition were calculated for the test site, and compared with those from the distant site.

### 2.2.3. Site mapping

The baseline mapping work for the Goldeneye site comprised three related phases, covering different extents and levels of detail as proposed by Widdicombe et al. (2018). Phase 1 collated existing industry and public data to establish a broad overview of the study area and to identify data gaps requiring survey effort. This analysis pointed to sediment type as the main habitat driver for the area; however, no detailed, full-coverage maps of this parameter existed for the Goldeneye site. Phase 2 combined the existing data with spatial statistical modelling techniques (Random Forest, Generalised Additive Models) to predict the distribution of sediment type and the density of a key crustacean benthic species, *Nephrops norvegicus*, across the experiment site. In addition, shipboard multibeam bathymetry data were collected during three expeditions at sea, mapping the seafloor to a 10 × 10 m or 5 × 5 m resolution (Berndt, 2017; Achterberg and Esposto, 2018; Connelly, 2019). Phase 3 involved dedicated data collection at even finer resolution (as described as “Tier 4” in Widdicombe et al., 2018) to characterise fine-scale bathymetric variability at the Goldeneye site, and to monitor the experimental site before, during and after the controlled release experiment (during cruise JC180). These activities included the acquisition of high-resolution (0.25 × 0.25 m) bathymetry and sidescan sonar, photographic, and environmental data with the Gavia AUV, and photogrammetry data collected by the ROV Isis (Connelly, 2019).

### 2.2.4. Anthropogenic activities

The two most significant impacts of anthropogenic activities visible at the seafloor around the Goldeneye site are from ceased oil and gas (O&G) extraction (and its associated structures) and commercial fishing activity. All anthropogenic structures, such as the oil and gas infrastructure, were mapped within the project GIS, based on data from O&G infrastructure databases (UKOilandGasData, 2007; http://www.ukoilandgasdata.com/). The feature locations were buffered according to the physical dimensions of the structures (e.g., pipe diameters and typical well dimensions) and by typical near-field modifications (e.g., scour pits, etc.) using standard buffer values to visualise actual footprints. The Goldeneye platform was also included and buffered to account for the potential for historical contamination according to literature values for persistent contaminants. Field studies in the North Sea have demonstrated that the effects of dumping of cuttings (physical smothering, organic enrichment and chemical contamination) are persistent and principally confined to benthic communities within a 1–2 km radius of platform sites (see Breuer et al., 2004 and references therein; Gates and Jones, 2012; Jones et al., 2012). Vessel Monitoring System layers for 2007–2015 (vessels over 15 m), sourced from the UK Marine Management Organisation (data obtained from the MMO via: http://data.gov.uk/; accessed January 2019), were used to represent the intensity of fishing effort across the site. Trawl scar densities were estimated from the Gavia AUV Geoswath surveys and used as a proxy for the cumulative pressures in specific areas.

### 2.3. Modelling to inform experimental design and predict impact

In order to understand, a priori, how the experimental CO₂ release would manifest in shallow sediments and the water column, customised sediment and hydrodynamic-chemical models of the Goldeneye site were developed. These models had high spatial and temporal resolution to resolve chemical changes at the centimeter to meter scale (in sediments) and meter to kilometer scale (in water column). Models of sediment chemistry were used to estimate the change in solute concentrations, including DIC and O₂, due to redox processes in the top 30 cm of the sediments (Dale et al., 2015). Models of two-phase geophysical flow were used to estimate the degree of CO₂ retention, flow pathways in the sediments, and the mode of exchange between sediments and the water column using modified Navier-Stokes-Darcy equations. These estimates can predict the footprint of CO₂ release at the seabed, the gas breakthrough time, but most vitally, the required injection rate for sustained gas breakthrough to the water column.

This abundance of data was then exploited in models of multi-scale hydrodynamic transport in the water column, including multi-phase bubble stream dynamics (Andeland and Drange, 2001; Dewar et al., 2015; Gros et al., 2019) linked to local and regional General Circulation Models coupled to a model of carbonate chemistry (Artioli et al., 2012). These models simulated the transport and dispersion of CO₂ in the water column and the resulting chemical changes detectable by sensors (Cazenave et al., 2018). These estimates of the physio-chemical footprints of the released CO₂ enabled informed planning of the observational layout and the choice of sensors and parameters measured from fixed and moving platforms (Hvidfeldt et al., 2015; Greenwood et al., 2015; Hvidevold et al., 2016; Alendal, 2017; Oleynik et al., 2020). These predictions also estimated the footprint of increased acidification (Blackford et al., 2020), assisting in environmental impact assessments.

Medium resolution models (Blackford et al., 2017) were used to spatially and temporally extrapolate our knowledge of natural variability as a key question was: can we distinguish potentially small signals of a CO₂ release from the complex natural variability in the marine
system? The baseline and release experiment data were used to evaluate the performance of the various models, to assess their utility as useful tools for studies of impact and monitoring strategy, and to assess other methods for analysing data streams to detect leakage, such as the stoichiometric $C_{\text{deep}}$ method (Botnen et al., 2015).

3. CO$_2$ release equipment

A fundamental objective of the field experiment was to release gas from a point within the sediments approximately 3 m below the seafloor surface. To do this, the gas needed to be released from tanks positioned directly on the seabed, with no connection to the surface, to allow safe ROV operations and to increase ship mobility. A central design consideration was the ability to change the injection flow rates during the field experiment to simulate different leakage rate scenarios. Consequently, it was necessary to insert a gas release pipe on site (Section 3.1) and to connect it to a gas storage and delivery system (Section 3.2) that was custom-built for the STEMM-CCS field experiment.

### 3.1. Pipe insertion setup

The injection of gas into shallow sediments in the North Sea at 100 km offshore from Scotland presented a major engineering challenge. Directional drilling from land, as done in the QICS project (Taylor et al., 2015), was not an option. The solution was to draw on well-known cone penetrometer techniques, whereby an instrumented rod is pushed vertically downwards into the seabed using a subsea drive unit. This technique was adapted to push a pre-curved steel pipe downwards into the seabed such that the end of the pipe was located 3 m below the seabed with an upward attitude (Fig. 5a). The upward attitude of the outlet was necessary to prevent the gas from tracking back along the outside of the pipe rather than finding a natural pathway up through the

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**Fig. 4.** STEMM-CCS experimental geometry and timeline. (Top) Schematic overview of representative positions of all deployed equipment around the experimental site. All observed bubble streams are indicated on the map although not all were active simultaneously. Equipment sizes are approximately to scale. (Bottom) Timing of the ships, vehicles, deployed equipment, sampling, and CO$_2$ injection flow rates after the initial setup phase of the experiment.
sediment. The design and manufacture of the pipe insertion rig was contracted to Cellula Robotics (Cellula Robotics Ltd., Burnaby, Canada), following an open tender exercise.

The pipe insertion rig (Fig. 5b) was a 2.3 m cubic steel frame. It housed a hydraulic power pack to drive a set of clamp rollers that firmly held the pipe and slowly rotated to drive the pipe along its own axis into the sediment. Control and electrical power to the hydraulic unit was provided via an umbilical cable from the ship. The umbilical also carried live video back to the ship (i) to verify the rig had landed on a suitable site, (ii) to confirm the orientation of the rig (and hence the direction that the pipe would be inserted), and (iii) to monitor progress of the pipe as it was pushed into the sediment. The rig had sufficient push force to jack up its 6 t mass should the pipe encounter an impenetrable object. An inclinometer was included to provide early warning of any change of frame angle that would result from such an occurrence.

The carbon steel pipe (9 m length, 38.1 mm outer diameter, 12.7 mm inner diameter) was pre-curved to a radius of 6.9 m. A retractable ‘goose-neck’ was included to support the pipe during deployment. The outlet end of the pipe was closed with a pointed tip to aid penetration through the sediment. Just behind the tip were a number of 12.7 mm diameter gas exit holes drilled through the pipe wall at a 45° backward slant to prevent them becoming clogged with sediment. Inside the drilled portion of the pipe was a 460 mm long sintered stainless-steel diffuser with a pore size of 9 μm to ensure the gas flow was distributed evenly across the outlet holes. The pipe inlet, which remained above the seabed, had a quick-connect fitting for connection to the gas supply via a flexible hose.

3.2. Gas release system

It was originally intended that the gases would be supplied in standard industrial cylinders and delivered to the seabed on pallets. However, commercially available cylinders are not designed for a subsea environment and establishing leak-proof connections between the pallets would have been problematic. It was therefore necessary to design and build a custom gas storage system.

A pair of custom-made bulk CO$_2$ storage tanks, connected to act as one single storage volume of $5.6 \text{ m}^3$, was procured from City Gas (City Gas EOOD, Stara Zagora, Bulgaria). This storage volume was sufficient to accommodate 3 t of liquid CO$_2$ with a 1.7 m$^3$ vapour headspace at 20 °C. It is usual for bulk CO$_2$ to be stored cryogenically to reduce storage pressure to around 20 bar. However, this requires insulated tanks and a cooling plant to maintain the temperature at around −20 °C, which was not viable in a marine setting. The tanks were therefore uninsulated and designed for a maximum working pressure of 80 bar. This was sufficient to allow for an ambient temperature in excess of 30 °C, which was unlikely to be encountered during springtime in the UK. At the seabed, the ambient temperature would be around 8 °C, resulting in a storage pressure of 42 bar(a) (absolute pressure). The bulk storage tanks, along with the other equipment described below, were mounted in a steel deployment frame (5.5 m length, 2.55 m width, 2 m height) (Fig. 6) that had a gross weight of 13 t (including a total of 3.3 t of liquid CO$_2$).

The gas rig also housed 200 L of a tracer gas mixture (BOC, UK) comprised of molar fractions of 58.98 % krypton (Kr), 1.77 % sulfur hexafluoride (SF$_6$) and 0.11 % octafluoropropane (C$_3$F$_8$), with the balance consisting of gaseous CO$_2$ (Section 4.2.2). The tracer gas mixture was decanted into four manifolded bladder accumulators (QHP, England) for deployment. The accumulators were kept charged to a constant pressure of 30 bar(a) via a regulated gas feed from the bulk CO$_2$ tanks. This was necessary to aid stability of flow and to ensure that nearly all of the mixture could be extracted when submerged at 120 m water depth with an external pressure of approximately 12 bar. The tracer gas mixture was fed into a custom-designed control unit (Fig. 7) where the flow was regulated through a mass flow controller (MFC) (Bronkhorst, UK) and then mixed into the main CO$_2$ line. This was necessary to aid stability of flow and to ensure that nearly all of the mixture could be extracted when submerged at 120 m water depth with an external pressure of approximately 12 bar. The tracer gas mixture was fed into a custom-designed control unit (Fig. 7) where the flow was regulated through a mass flow controller (MFC) (Bronkhorst, UK) and then mixed into the main CO$_2$ line. This was necessary to aid stability of flow and to ensure that nearly all of the mixture could be extracted when submerged at 120 m water depth with an external pressure of approximately 12 bar. The tracer gas mixture was fed into a custom-designed control unit (Fig. 7) where the flow was regulated through a mass flow controller (MFC) (Bronkhorst, UK) and then mixed into the main CO$_2$ line. This was necessary to aid stability of flow and to ensure that nearly all of the mixture could be extracted when submerged at 120 m water depth with an external pressure of approximately 12 bar. The tracer gas mixture was fed into a custom-designed control unit (Fig. 7) where the flow was regulated through a mass flow controller (MFC) (Bronkhorst, UK) and then mixed into the main CO$_2$ line. This was necessary to aid stability of flow and to ensure that nearly all of the mixture could be extracted when submerged at 120 m water depth with an external pressure of approximately 12 bar. The tracer gas mixture was fed into a custom-designed control unit (Fig. 7) where the flow was regulated through a mass flow controller (MFC) (Bronkhorst, UK) and then mixed into the main CO$_2$ line. This was necessary to aid stability of flow and to ensure that nearly all of the mixture could be extracted when submerged at 120 m water depth with an external pressure of approximately 12 bar. The tracer gas mixture was fed into a custom-designed control unit (Fig. 7) where the flow was regulated through a mass flow controller (MFC) (Bronkhorst, UK) and then mixed into the main CO$_2$ line. This was necessary to aid stability of flow and to ensure that nearly all of the mixture could be extracted when submerged at 120 m water depth with an external pressure of approximately 12 bar.
10,000:1 to yield molar fractions of 58.98 ppm Kr, 1.77 ppm SF$_6$ and 0.11 ppm C$_2$F$_6$ in the final injection gas. This target ratio was kept constant throughout the release experiment.

The overall flow rate was adjustable from 0 to 100 normal litres per minute which is equivalent to a mass flow range of 0–285 kg/d. Remote adjustment of the MFCs and feedback of engineering data for flow rate, pressure and temperature were achieved using an optical modem that enabled communication to the research ship via the ROV’s umbilical cable. The control unit was powered by a pair of 24 V lead-acid batteries housed in stainless steel cylinders. From the control unit the mixed gas flowed to a valve panel where an outlet could be selected. These outlets housed in stainless steel cylinders. From the control unit the mixed gas flowed to a valve panel where an outlet could be selected. These outlets included a sample port from which ‘raw’ samples could be collected to verify the composition of the gas mixture, a 100 m long flexible hose that could be laid on the seabed to release gas directly into the water column should the buried pipe be blocked and not allow gas to flow.

After the successful insertion of the curved pipe and deployment of the CO$_2$ storage rig, the ROV carried the 100 m long flexible hose from the CO$_2$ rig to the pipe inlet. The ROV’s manipulator arm was used to connect the hose to the pipe inlet. The gas was turned on using an optical modem connection between the ROV and the CO$_2$ source and for quantifying the rate of reactions that occur in the sediments. Gaseous CO$_2$ migrating through sediments will dissolve in the pore water and will react with the sediment’s mineral phases such as carbonates and silicates. Thus, sediment cores were examined for parameters indicative of CO$_2$ dissolution, mineral dissolution, and changes in the physical properties and composition of the sediment.

Three different types of sediment cores were collected during different phases of the CO$_2$ release experiment: push cores, gravity cores, and multicores. Push cores (up to 30 cm length) were retrieved prior, during, and after the CO$_2$ release experiment using the ROV manipulator arm. The advantage of this core sampling strategy is that the core locations can be targeted on a cm scale, e.g., close to bubble streams or pockmarks. Up to six push cores were taken on selected dives at each stage of the CO$_2$ release experiment (see indicative locations on Fig. 4 and photo in Fig. 8a). The sandy nature of the sediment meant that the holes caused by the coring were rapidly infilled to only leave a shallow indent on the seafloor surface that quickly disappeared. After recovery the push cores were processed on board the ship (RRS James Cook) in a chamber filled with nitrogen gas to minimise contact with oxygen within a controlled temperature lab set to the seabed temperature (~8 °C). Gravity cores were also taken from the RV Poseidon to investigate deeper layers of the sediments (up to 4.5 m below seabed) as well as multicores (up to 60 cm below the seabed) taken at similar locations, which enabled sampling of the sediment-water interface. The gravity and multicores were obtained after the CO$_2$ injection had stopped and all the equipment had been recovered (Fig. 4) to ensure the integrity of the sediments was maintained during CO$_2$ release. The gravity cores and multicores were processed under ambient conditions on board the RV Poseidon. All cores were sub-sampled for biogeochemical analyses of the pore waters (nutrients, cations, anions, TA, DIC, δ$^{13}$C$_{DIC}$ and δ$^{18}$O$_{H2O}$) and the sediments (geochemistry, mineralogy, total carbon and nitrogen content, total organic carbon, δ$^{13}$C of organic carbon, particle size and porosity) (Connelly, 2019; Schmidt, 2019).

### 4.1.2. Sediment microprofiler

In-situ sampling and analysis of pore water has significant advantages over core sampling and ex-situ pore water extraction because concentrations of many chemical species may change during sampling or cannot be sampled accurately as they have steep vertical gradients near the sediment-water interface (Revsbech and Jørgensen, 1986; de Beer, 2000; Kühl and Revsbech, 2001; Gieske and de Beer, 2004). In-situ analyses of sediment pore waters can be achieved using micro-sensors: miniaturised electrochemical or optical sensors. The sediment microprofiler was equipped with several microsensors (Fig. 8b and c) that measure the concentrations of different chemical...

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**Fig. 7. Schematic of the gas control unit.**
species using electrochemical techniques. As the tip diameter is <50 μm, the microsensors impose minimal disturbance to the sediments and provide high resolution vertical profiles of O₂, pH, hydrogen sulphide (H₂S), redox state, and temperature within the sediment. The microprofiler can be deployed for up to 72 h at 2 °C. The sensors were calibrated on the ship, before and after each deployment.

The microprofiler was deployed along transects from outside the bubble stream area towards the experiment epicentre. It was positioned on the seafloor by the ROV (Fig. 8b), and the profiling was switched on by activating a magnetic switch with the ROV manipulator arm. The device was programmed to make measurements at 250 μm intervals downwards through the sediments. Measurements began a few cm above the seabed and extended to 12 cm depth into the sediment. Each profile took approximately 50 min to record. The O₂ profiles were used to calculate vertical O₂ fluxes using Fick’s law of diffusion (Jørgensen and Revsbech, 1985).

4.1.3. Sediment optodes

Four stand-alone optical sensors (optodes) continuously monitored the pH in the sediment pore waters at 20 cm depth. They were placed at distances of 1, 1.4, 4 and 7 m away from the bubble streams and recorded pH prior to, during and after the CO₂ release (Fig. 8a).

The optodes utilised the pH-dependent fluorescence of an indicator dye immobilised into a proton-permeable polymeric matrix (hydrogel). An additional material with pH-independent fluorescent properties provided a reference measurement (Klimant et al., 2001). Trials of an earlier version of the system (Staudinger et al., 2019) showed that the response of the optodes was very slow at Goldeneye, so a new optode system that utilised polymeric microparticles was deployed and this substantially improved the optode response time. The sediment optodes used commercially available opto-electronics from PyroScience (FireStingO2), a custom-built logger unit (Max Planck Institute for Marine Microbiology, Bremen, Germany) and rechargeable batteries, all placed in a titanium housing. An optical fibre guided the light between the logger and the pH-sensing element. The distal end of the fibre housed the sensing material and was enclosed in a stainless-steel sleeve for rigidity in the sediment. The temperature was recorded by a PT100 sensor. The temperature was needed to calculate the pH accurately and incidentally demonstrated that reactions between the sediments and CO₂ generated heat.

For the deployment, the sediment optode loggers were mounted on a perforated metal sheet with an ROV handle attached on top and the pH and temperature probes extending 20 cm below the sheet. Prior to the deployment the pH sensors were calibrated and set to measure every 5 s for 1 min, then to sleep for 15 min. The four sediment optode loggers were deployed by the ROV four days before the CO₂ release was initiated. They were left in place throughout the experiment and recovered shortly after the CO₂ release stopped (Fig. 4).

4.1.4. AUV-mounted chirp sub-bottom profiling

An AUV-mounted chirp sub-bottom profiler was used to image the subsurface in and around the release site. This sonar-based system is sensitive to density and velocity changes in the sediments, which permits imaging of sediment stratification and to identify and characterise gas pockets in the seafloor. Data were collected in a dense grid centred above the CO₂ release point with a line spacing of between 2 and 5 m from elevations of 7.5 and 2 m above the seabed. The chirp sonar used a digitally produced 14–21 kHz acoustic transmission and matched filter processing to achieve both high resolution and good penetration of soft sediments up to 10 m below the seabed. Surveys were carried out before the gas release began, during the release at rates of 6, 29, and 143 kg/d, and after the release was stopped to observe the migration of CO₂ within the sediments and any lasting effects it may have had.

4.2. Detection of gaseous CO₂ in the water column

A range of custom developed equipment was deployed to detect, characterise, and quantify the CO₂ gas bubbles in the water column. Physical bubble characteristics were measured with optical and acoustic methods (Sections 4.2.1, 4.2.3 and 4.2.4). Gas chemistry was analysed from collected gas samples and physical measurements of flow rates were made using funnel-based methods (Sections 4.2.1 and 4.2.2).

4.2.1. Gas bubble imaging (Optical Lander)

An Optical Lander was custom-built to obtain optical and physical measurements of the gas flow rate into the water column. The design for the optical measurements incorporated two underwater cameras (Sony FDR-X3000 Action Cam) in custom housings opposite an illuminated scale board. The frame was placed directly over an individual seep by the ROV, such that the bubble stream passed directly between the cameras and the screen. The visible scale on the screen enabled the gas flow to be quantitatively estimated by using the camera footage to measure the size of each bubble passing in a given time period and then estimating the total volume of gas observed. The frame was deployed successfully for one measurement period, collecting 15 min of useable footage at 90 frames per second before being recovered. An alternative mechanical measurement of the flow rate of an individual seep was accomplished using an “inverted rain gauge”, which was placed over the centre of the Optical Lander, roughly 0.5 m above the seabed (Fig. 9b). This gauge collected gas from the underlying seep until a known volume of gas was collected, at which point it tilted upwards emptying the gas and resetting itself. By observing how often the gauge tilted, a measurement of gas flow rate could be made.

4.2.2. Gas bubble sampling

In monitoring for potential leakage from an offshore CO₂ storage site, it is important to be able to demonstrate that an observed anomaly
originates from a leakage rather than a natural source (Dixon and Romanak, 2015). One way to do this is to label the CO2 in the storage reservoir with tracers. These are chemical constituents that are either inherent to the injected CO2, arising from natural processes within the reservoir, or have been purposely added to the injected CO2. In the STEMM-CCS release experiment inherent tracers (δ13C,C02, δ18O,C02) and added tracers (Kr, SF6) were tested (Section 3.2). The added tracers had a significantly lower solubility in seawater than CO2, so the change in gas composition could be used to estimate the fraction of CO2 that dissolved in the sediment pore waters and in the water column at different injection flow rates.

Gas was sampled using custom-built gas bubble samplers (GBS) (Corsyde, Germany) that were operated by the ROV manipulator arms. The GBS consisted of an inverted transparent funnel (0.7 l internal volume), inlet valve, stainless steel sample cylinder (0.5 l internal volume) and outlet valve (Fig. 9a). Markings on the funnel helped to identify the volume of gas collected over a given time period, providing an approximate flow rate from the bubble stream. Gas was usually collected once a day from (i) the gas rig sample point, (ii) ~10–15 cm above seabed bubble streams, and (iii) occasionally from ~0.9–2.7 m above seabed. The rig gas samples were used to identify any temporal variability of the CO2 tracer ratio in the injection gas so that changes in the gas composition of the bubble stream samples could be accurately computed. Gas samples from ~0.9–2.7 m above the seabed were collected to quantify the rate of CO2 dissolution in the water column. In addition to the added tracer gases, methane (CH4), which was present in trace quantities in the released CO2 gas, provided an independent tracer unaffected by potential variability of the tracer mixture injection system. An onboard flow-through Fourier-Transmission Infra-Red (FTIR) analyser (atmosFIR, Protea Ltd., UK) was used to measure CO2, CH4, SF6 and CF6 concentrations in the gas samples. Sub-samples for determination of the isotopic signatures (δ13C,C02, δ18O,C02) and Kr concentration were taken for analyses back onshore.

4.2.3. Passive acoustics (hydrophone walls)

Measurements of the sounds of bubbles as they emerge from the seabed into the water column allows determination of the gas flux and the sizes of bubbles (Leighton and White, 2012).

In previous underwater bubble acoustic experiments, one hydrophone was used to take passive acoustic time series data, and the temporal data were used to quantify the gas flux (Berges et al., 2015). In the STEMM-CCS experiment, this technique was extended by using an array of hydrophones. This allowed the location of the bubble streams to be identified. It also increased the range of the technique by enhancing the signal-to-noise ratio because a single-hydrophone-based passive acoustic technique is susceptible to underwater background noise, e.g., ship noise and sea surface noise (Leighton and White, 2012; Li et al., 2019). During the CO2 release experiment the application of a hydrophone array and beamforming technique for potential CCS leakage monitoring was tested. The hydrophone array comprised five hydrophones fixed at different spatial positions on a frame with a height of 1.10 m and a width of 1.36 m (Fig. 9c). The five hydrophones were linked to an acoustic recorder (RS-ORCA), which was used to archive the sound of bubbles emerging from the seabed. The passive system was programmed to make measurements at predetermined time intervals of 5 min on and 5 min off during the gas release experiment. These developments made it possible to improve the signal-to-noise ratio for flow rate quantification and single bubble detection, to localise the gas seep site, and to quantify the bubble size and gas flow rate for each seep.

4.2.4. Active acoustics: Ship-based sonar and AUV-based sidescan sonar

Figure 9. Selection of methods employed to detect and/or characterise gaseous CO2 in the water column. (a) Gas bubble sampling, with the ROV arm holding gas bubble sampler connected to an inverted funnel over a bubble stream. (b) Self-illuminated optical lander with bubble imaging and “rain gauge” bubble collector closed (left) and open (right) with gas escaping. (c) Hydrophone wall housing the passive acoustic setup. Given the strong impedance contrast between water and gas, hydro-acoustics are very sensitive to the presence of gas bubbles in the water column. Hydro-acoustic sonar systems can be both ship- and AUV-based, depending on the required physical scale and resolution.

Hydro-acoustic data were collected during the JC180 cruise using the ship-mounted multi-beam Kongsberg EM710 and the single-beam Simrad EK60. The EM710 transmits a high frequency pulse in a fan shape beneath the vessel, surveying a large portion of the seafloor perpendicular to the ship’s track. The EK60 echo sounder transmits a single beam of 5 different monochromatic frequencies surveying a small portion of the seafloor and the water column directly beneath the ship. The multi-beam echosounder was used to survey a wide area around the experimental site to ensure all bubble streams were identified. The calibrated single-beam system was used to determine bubble stream properties, i.e., bubble size and mass flow rate, via modelling. Similar ship-based active acoustic measurements were made by RV Poseidon, which was a Simrad EK80 and EC150-3C transducer were mounted through the ship’s moon pool for echosound and current profile measurements. This system was used to observe bubble streams as well as to identify the location of the towed Video-CTD system.

To examine the surface features of the seafloor in greater detail, a Geoswath bathymetric sidescan sonar was mounted on the AUV. Surveys using the AUV were carried out prior, during and after the CO2 release (Fig. 4). Similar to the multibeam system described above, an even wider, fan-shaped, outgoing sound pulse is transmitted by the sidescan sonar, and the backscatter returns are recorded as continuous signals on both sides of the instrument, in two sets of slightly offset receivers, enabling the calculation of water depth and seafloor reflectivity across the swath. The bubble streams could also be identified in the ‘water column’ section of the sidescan sonar records. The system operated at higher frequency than the ship systems, providing a finer resolution and more detail close to the seabed though with smaller footprint.
4.3. Detection of dissolved CO\(_2\) in the water column

Dissolved CO\(_2\) in the water column was measured in collected samples, in seawater pumped onto the ship (Section 4.3.1) and with in-situ instruments in the water column on mobile and fixed equipment. The flux of dissolved CO\(_2\) from the seafloor was measured using incubation chambers deployed on the seafloor (Section 4.3.2). Dissolved CO\(_2\) concentration and flux across the sediment-water interface in the vicinity of bubble streams was quantified with eddy covariance and the lab-on-chip gradient method (Sections 4.3.3 and 4.3.4). Measurements were conducted with in-situ sensors mounted onto a towed frame (Section 4.3.1) and mounted directly onto the ROV and AUV (Sections 4.3.5 and 4.3.6).

4.3.1. Discrete and continuous water sampling

Discrete water column samples were collected from both ships to detect the influence of the injected CO\(_2\) gas on carbonate system parameters and/or on the dissolved tracer concentrations.

On board the RRS James Cook, the water column was sampled using Niskin bottles (6 × 1.7 L) mounted at the back of the ROV (Fig. 10a). Once a day, up to four Niskin bottles were fired between 1.5 and 2.5 m above seabed bubble streams and a further two bottles were fired close to the gas rig to establish a background value far from the bubble streams. These samples were used to determine concentrations of dissolved inorganic nutrients, TA, DIC, carbon and oxygen isotopes (\(\delta^{13}\text{C}_{\text{DIC}}, \delta^{18}\text{O}_{\text{H}_2\text{O}}\), SF\(_6\), C\(_3\)F\(_8\) and Kr.

On board the RV Poseidon, a towed multipurpose Video-CTD water sampler rosette was used to sample the water column. This equipment was developed for detecting and monitoring gas-rich fluid seepage from the seafloor (McGinnis et al., 2013; Schmidt et al., 2015; Sommer et al., 2015) and for leakage rate estimation (Gros et al., 2019). The Video-CTD was equipped with Niskin bottles (10 × 10 L), a HD video camera, and additional sensors for pH (SBE27, Sea-Bird Electronics, Inc.) and dissolved CO\(_2\) (HydroC-CO\(_2\), Kongsberg Maritime Contros). The towed Video-CTD water sampler rosette was used for (i) discrete water sampling using the Niskin bottles, (ii) continuous water sampling by pumped supply, and (iii) in-situ measurements of the partial pressure of CO\(_2\) (pCO\(_2\)). The Video-CTD was towed in bottom view mode at ~1–5 m above seafloor) while the ship traveled at low speed (0.2–1 knots) following a random track pattern for ~10 h. The system was deployed during CO\(_2\) release rates of 0, 6, 29, and 143 kg/day (Fig. 4). The natural background pCO\(_2\) during deployments was determined by sampling in areas 700–2700 m from the release site. All data sets from in-situ sensor measurements were recorded online and related to the ship’s navigation data and weather data. Live HD video-streams and images were recorded for subsequent benthic characterisation. Discrete water samples were taken from the Niskin bottles for analyses of DIC, TA, \(\delta^{13}\text{C}_{\text{DIC}}\), DOC, DON (dissolved organic nitrogen) and inorganic nutrients. In addition, seawater was continuously pumped from specific depths into an onboard Membrane Inlet Mass Spectrometer (MIMS) for real-time analysis of dissolved gases. Discrete samples for DIC, TA, \(\delta^{13}\text{C}_{\text{DIC}}\) and inorganic nutrient analyses were collected hourly from the pumped water supply and when the Video-CTD was above active bubble streams (confirmed by video image) to help calibrate the in-situ pH and pCO\(_2\) sensor measurements.

4.3.2. Benthic chambers

Benthic chambers are in-situ incubation chambers, which were used to quantify the flux of DIC across the sediment-water interface near the CO\(_2\) bubble streams before and during the release experiment and to monitor the potential effect of the CO\(_2\) release on the fluxes of other solutes. The chambers enclose and gently mix a known volume of benthic water and automatically collect water samples into glass syringes throughout the deployment for later analyses of a wide variety of solutes.

During the field experiment, two benthic chambers were deployed (Fig. 10b; McGinnis et al., 2014) for a total of 5 deployments (Fig. 4). The chambers were deployed by the ROV and inserted into the top few centimeters of the sediments to seal a fixed ~6 L volume of water above the sediment. By monitoring the evolution of solute concentrations within this incubated volume over 27–38 h, their fluxes across the sediment-water interface were quantified. The benthic chambers included an O\(_2\) optode (Aanderaa, Xylem, USA) and temperature sensors for continuous collection of in-situ data. During each deployment, eight water samples of ~46 ml were automatically collected into glass syringes by the chamber system. These samples were later analysed in the laboratory for dissolved gases, nutrients, stable isotopes (\(\delta^{13}\text{C}_{\text{DIC}}, \delta^{18}\text{O}_{\text{H}_2\text{O}}\)), and TA.

4.3.3. Eddy covariance

The rapid dissolution of CO\(_2\) bubbles as they emerge from the seabed generates a strong vertical gradient of hydrogen ions (which cause low pH) and DIC (Zeebe and Wolf-Gladrow, 2001). The interaction of currents with this source induces turbulent mixing. The vertical component of this mixing drives a net upwards flux of hydrogen ions and DIC from the seafloor. The eddy covariance technique quantifies the upward flux by simultaneously measuring the pH and 3D velocity of a small volume of water at high frequency (e.g., 5 Hz). The implementation for the STEMM-CCS project followed published work which used this technique for benthic biological O\(_2\) (Berg et al., 2003) and CO\(_2\) (Long et al., 2015) production. Because pH eddy covariance is sensitive enough to quantify the naturally-occurring benthic biotic CO\(_2\) flux, it was expected to be exceedingly sensitive to a CO\(_2\) bubble stream. During the release experiment, both O\(_2\) and pH eddy covariance fluxes were determined. Turbulent fluctuations in velocity were measured with an acoustic Doppler velocimeter (Nortek, Norway), pH with a fast-response pH ion sensitive field effect transistor (ISFET, Microsens Switzerland), and O\(_2\) with an optode minisensor (PyroScience, Germany). The instruments were mounted onto the fiberglass EC/gradens lander frame (Fig. 10d). The technical development for the application of eddy covariance to the quantification of the CO\(_2\) release included a flow-through housing with an integrated, minimally stirring-sensitive reference developed for the

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Fig. 10. Selection of methods employed for measuring dissolved CO\(_2\) in the water column. (a) Niskin bottles for water sampling from the ROV. (b) Benthic chamber for measuring benthic fluxes. (c) pH optode sensors held in the bubble stream by the ROV’s manipulator arm. (d) EC/gradens lander for eddy covariance and lab-on-chip chemical gradient measurements.
ISFET sensor. Flow was driven by an electronically-controlled gear pump that periodically reversed to flush debris that would otherwise clog the flow. Measurements were made 16 cm above the seafloor, 2.6 m south of the centre of the release site (Fig. 4). Complementary carbonate system parameters were obtained from lab-on-chip sensors on the same instrument frame (Section 4.3.4). The payload limitation of the ROV limited the maximum battery capacity of the landers, so two identical instrument frames were developed and swapped every 48 h before the batteries ran down. This allowed us to acquire a near-continuous time-series of observations during the CO₂ release.

The eddy covariance technique measured vertical flux at a single point located downstream of the source. Therefore, the measured flux represents only a small portion of the total release. To estimate the total release of DIC, the measured velocity, flow direction, and pH were combined with a modelled distribution of CO₂ dissolution above the seafloor.

4.3.4. In-situ chemical gradient measurements

As CO₂ bubbles dissolve in the water column, they cause a strong drop in pH and an increase in DIC near the seafloor. Miniaturised spectrophotometric sensors measuring pH (Rérolle et al., 2013) and TA, and a conductometric sensor for measuring DIC directly, monitored these carbonate parameters. Spectrophotometric sensors for measuring nitrate (Beaton et al., 2012) and phosphate (Clinton-Bailey et al., 2017) were included to characterise any background changes in DIC due to biological activity.

These sensors were all “lab-on-chip” (LOC) devices: autonomous miniaturised instruments which perform chemical assays in-situ with low power and reagent consumption. The sensors draw in seawater, mix it with chemical reagents, and measure the reaction products either optically or electrochemically. The TA and DIC sensors were custom-developed for this CO₂ release experiment.

One LOC sensor for each of the above parameters was mounted on each EC/gradients lander (Fig. 10d). Each sensor sampled from two inlets: a lower inlet located 18 cm above the seafloor and a higher inlet located 87 cm above the seafloor. Each sensor continuously alternated between the two sample inlets to characterise the vertical concentration gradient in each parameter at sample intervals of 5–15 min depending on the sensor. The EC/gradients landers were deployed throughout the release experiment, yielding a nearly-continuous dataset from before the gas was turned on until the day it was turned off (Fig. 4). The data from these LOC sensors, along with measurements of the current, can be used to quantify the dissolved fraction of the CO₂ release by estimating the total excess DIC content of the water as it passes over the lander.

4.3.5. Chemical mapping with the ROV

The ROV was used as a vehicle for deployment and positioning of equipment at the seafloor as well as a survey instrument. To determine the spatial extent of the plume, several overnight dives were used to perform three-dimensional chemical surveys of the experiment site with LOC sensors and optodes. (“Plume” here refers to water from the vicinity of the bubble streams that is characterised by elevated DIC concentration and reduced pH as a result of the CO₂ gas dissipating in the seawater.) While ROVs cannot be used as a long-term monitoring approach without a ship present, these measurements enabled the creation of maps of the plume which could be used to validate models of the release.

A suite of five spectrophotometric LOC sensors were used to measure carbonate system parameters and nutrients (see Section 4.3.4). These were mounted to a removable rack on the back of the ROV, visible in the bottom-left corner of the vehicle in Fig. 3a. A pump at the front of the ROV supplied seawater to the LOC sensors for analysis. A live data stream was delivered via the ROV tether to provide real-time results from the LOC sensors to the ship allowing the operators to adjust survey plans in real-time during the surveys if required.

The pH optodes were similar to those utilised for sediment pore water analysis (Section 4.1.3). Unlike the sedimt optodes, however, these pH optodes utilised sensor spots on a screw-on cap adaptor attached to the logger in a titanium pressure housing (63 mm diameter, 270 mm length) designed by PyroScience GmbH (Germany). The optodes were held by the ROV’s robotic arm during the surveys (Fig. 10c). Oxygen optodes were also deployed in the same set up for reference purposes.

Chemical surveys were performed with LOC sensors and pH optodes operating simultaneously. Longitudinal (north-south) surveys were performed with the ROV moving from directly above the bubbles to 10 m downstream at altitudes of 1.5 and 3.5 m. A lateral (east-west) survey was performed at 6 m downstream from the bubbles and 1.2 m altitude to map the distribution of the plume perpendicular to the current. The LOC sensors also gathered data on all ROV dives while the ROV was completing other tasks in and around the experimental site.

4.3.6. pH sensors on AUV

To collect spatial data on the pH distribution over the larger experimental region a high temporal resolution pH sensor (Deep SeaFET™, Sea-Bird Scientific, USA) was mounted on to the AUV (Teledyne, USA) (Fig. 3b). This pH sensor uses an ion sensitive field effect transistor (ISFET) to measure pH changes. This sensor was selected because of its high sample frequency of 1 Hz and its depth rating of 2000 m. The sensor was mounted externally on the Gavia AUV using a custom-made bracket, fabricated from syntactic foam, designed to minimise drag.

The SeaFET recorded data on seven dives over the course of the experiment using a variety of dive track patterns. To avoid any interactions with the CO₂ release equipment at the seafloor, the AUV had to remain at a relatively high altitude (4–7.5 m) when it was close to the CO₂ release site.

5. Outcomes and discussion

This section summarises the general outcomes of the release experiment activity and the lessons learnt. An overview of the position and timing of the ships, vehicles, deployed equipment, and CO₂ injection flow rates throughout the experiment is given in Fig. 4. For detailed results, the reader is referred to the specialised papers, which can largely be found in this special issue.

5.1. Baseline measurements and site characteristics

In environmental monitoring a robust understanding of the systems’ natural variability is essential to differentiate between a potential leak signal and an anomaly caused by natural phenomena or unrelated human activity.

Results from the environmental baseline studies showed that the experimental site, and wider Goldeneye area, was located on a relatively flat seabed with a gentle slope, dipping east-north-east towards a broad depression termed the Witch Ground Basin (Strong et al., 2020). The near-surface sediments at Goldeneye had a sand content of ~70 %, decreasing to ~30 % by 120 cm depth below seabed, and a porosity between 0.4 and 0.6 (Dale et al., 2020). The organic carbon content was low (0.6 %). Benthic respiration rates at Goldeneye were typical for coastal margins (~6 mmol/m²/d of O₂). Stoichiometric relationships between the fluxes of sulfate, ammonium and alkalinity in the pore waters have the potential to serve as diagnostic indicators for CO₂ leakage at Goldeneye (Dale et al., 2020).

The benthic fauna studies revealed that the experimental site was a fairly typical habitat for the continental shelf surrounding the British Isles (Strong et al., 2020). The benthic macrofauna study identified 10, 207 individuals from 264 taxa in 13 phyla in 76 box cores. Based on abundance, 70 % of individuals were annelids, 11 % molluscs, 8 % arthropods, 7.5 % echinoderms and 2.3 % nemerteans. The most significant factor influencing benthic community structure was sediment type, in particular whether the substratum was muddy sand or sandy mud.
Lesser impacts on the benthic macrofauna from trawling and the presence of pockmarks were also evident. The baseline imaging survey revealed that heterogeneity in the megafaunal community at the Goldeneye area was low. A total of 24 morphotypes were observed at these sites (Hosking et al., 2020; Schmidt et al., 2020; STEMM-CCS D2.7, 2020). The crustacean, *Neprops norvegicus*, the subject of some mapping work (see Section 2.2.3), was one of the most common megafaunal organisms observed. The diversity and composition of the megafaunal community was not significantly different between the experimental site and the background site distant from it. These data suggest that a robust baseline assessment design should consider spatial variability for different organism size classes (e.g., macrofauna and megafauna).

Hydrographical data gathered before and during the release experiment by the NOC baseline lander (Fig. 13a) show that the experimental region is dominated by tides (tidal range 1.7 m) and by associated currents with a very narrow tidal ellipse with the major axis aligned in the north-south direction. The maximum current ranged from approximately 10–25 cm/s at 16 cm and 15–30 cm/s at 120 cm above the seafloor (Fig. 13b). The currents remained within 30 degrees of north to south 80% of the time, allowing for occasional quasi-steady-state conditions for measurements. Consequently, landers and stationary instruments were located on the seabed upstream and downstream of bubble streams for half of their deployment time. The instruments thus effectively measured “background” data during the upstream period, and CO2 release during the downstream period. The varying direction of current meant that physically stationary landers could examine the spatial aspects of the plume from a temporal signal, as the currents caused the plume of dissolved CO2 to sweep across the instruments. This is evident in the chemical time series data from the experiment, described in other papers within this special issue (Koopmans et al., 2020; Schaap et al., 2020). This underscores the importance of knowledge of the local hydrography when determining optimal lander positioning for environmental monitoring.

Water column carbonate system baseline data acquired during three expeditions along with historical data from the Goldeneye area show that near the seafloor the carbonate system varies as a function of seasonality, tidal cycles, and weather events such as storms. On the seasonal scale, drops in pH and build-up of pCO2 in the near-bottom waters are dominated by cumulative remineralisation of sinking organic matter over the course of the year while the breakup of water column stratification reduced bottom water pCO2 and increased pH. The annual range in pH was ~0.15 units and the annual range in pCO2 was about 160 μatm. High-resolution pH and pCO2 measurements obtained from the seabed landers suggest that tides drive short-term pH and pCO2 variability in bottom waters of, respectively, ±0.008 and ±1.5 μatm. Storm events led to abrupt and pronounced changes in carbonate chemistry parameters of up to 0.02 pH units and 30 μatm in pCO2 over a relatively short time (6 h) interval. Crucially, the results suggest that the natural short-term variations in pH are smaller than the proposed theoretical ΔpH criteria to effectively detect anomalies (Blackford et al., 2017). The application of multilinear regression analysis and co-variance relationships between water column parameters allowed the establishment of robust pCO2:O2 seasonal thresholds capable of assisting with the identification of anomalies that would be indicative of non-natural sources of CO2 (Esposito et al., 2020). The comparison of STEMM-CCS data with historical data from the Goldeneye area show that the carbonate system baseline in seawater has shifted due to invasion of increasing levels of anthropogenic atmospheric carbon dioxide. This would also need to be taken into account with, for example, stoichiometric analysis (Esposito et al., 2020; Martínez-Cabanas et al., 2020; Omar et al., 2020).

### 5.2. Outcomes of release experiment

The location of the experimental site was guided by the baseline site mapping of the oil and gas infrastructure in the Goldeneye area such as wellheads and pipelines (Strong et al., 2020, Fig. 2). Ultimately, the CO2 container and the experimental site were located approximately 900 m south of the Goldeneye platform, i.e., beyond the range of influence of oil and gas infrastructure.

The properties of the sediment were conducive for inserting the pipe and releasing the gas. The pipe insertion occurred without technical complications, taking only 24 min, and the insertion rig was back on deck within ~1 h. After maintenance of the gas release system to address gas release performance issues, the CO2 rig was successfully re-deployed, the gas supply was tested, and the flexible hose was connected to the pipe inlet within a day and without major issues.

The CO2 was turned on at 15:19 on 11 May 2019 and the flow rate was set to 6 kg/d. Bubbles were observed emerging from the sediments within the 30 min it took the ROV to arrive at the release site. The gas injection flow rate was turned up incrementally over the course of the experiment to 14, 29, 86, and finally 143 kg/d (Fig. 11). The CO2 release system was built to allow maximum injection flow rates of 286 kg/d (100 normal L/min) but following issues with the performance of the gas regulator it was decided to keep the maximum injection flow rate at 143 kg/d. The technical challenge of injecting very low amounts of tracer gas into the CO2 gas flow meant that the CO2-tracer ratio fluctuated significantly at low injection rates, but stabilised at higher injection rates (>29 kg/d).

Three separate streams of bubbles were visible at the lowest flow rate. Over the course of the experiment several further bubble streams were identified as the gas flow was increased, with some bubble streams occurring intermittently (see examples of bubble streams visible in Fig. 12a). All of the observed bubble streams occurred within a 4 m radius of the expected position of the pipe outlet, though most were clustered ~2 m to the south of this point (Fig. 4) (Strong et al., 2020). Dissolution of the bubbles in the water column was observed with the video footage from the ROV; no bubbles were visible to the camera at heights of >8 m above the seabed.

Most sediment cores were collected away from the pre-existing points of gas release and coring did not appear to modify the flux of CO2 gas across the seabed. Some cores were collected from a point where there was already gas contained within the sediments (i.e., immediately below or next to pre-existing bubble streams). In these cases, the collection of the cores was associated with a slight change in where the CO2 was emitted from the seafloor but no significant change in flow rate.

The main findings of each of the different techniques for CO2 detection, attribution, and quantification are as follows:

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**Fig. 11.** Gas injection during the release experiment. Injected gas mass flow rates (black solid line, left y-axis) and cumulative gas mass (grey solid line, right y-axis) injected over time (day of experiment (top x-axis) and respective date (bottom x-axis)).
Sediment geochemistry: In the sediments close to the bubble streams, the impact of injected CO$_2$ was detectable based on changes in the pore water chemistry, mainly associated with carbonate dissolution (Lichtschlag et al., 2020a).

Sediment microprofiler: The microprofiles in the bubble streams were highly variable, with porewater pH values as low as 5. An increase in temperature could be attributed to exergonic reactions between CO$_2$ and sediment pore waters and CO$_2$-sediment weathering (reactions with calcite and silicate). The pore water chemistry and temperature effects were only detectable within a few cm from the bubble streams (de Beer et al., 2021).

Sediment optodes: The sediment optodes showed a clear increase in temperature and a decrease in pH at a distance of 1 m and 1.4 m from the leakage. At a distance of 4 m and 7 m the values stayed constant. The deviation in temperature was easier to detect because the background values did not drift.

Gas bubble imaging: Cameras on the seabed lander identified bubbles from a single seep with ~80% of the bubbles ranging in size from 0.2 to 0.5 cm in radius, with most around 0.3 cm. The optically derived bubble size distribution was successfully used to quantify the amount of CO$_2$ that remained in the sediment due to CO$_2$ dissolution in pore waters. The total leakage rates were successfully determined using gas bubble samplers (Flohr et al., 2020).

Passive acoustics: The frequency of the bubbles recorded by passive acoustic methods allowed the determination of bubble radii, with a range of 0.15 to 0.30 cm, consistent with gas bubble imaging. Inversion of the hydrophone data facilitated an estimate of the total gas flux into the water column (Li et al., 2019, 2020).

Active acoustics: The chirp sub-bottom profiler on the Gavia AUV resolved the evolution of gas migration pathways throughout the experiment via enhanced reflectors, shadowing, variations in attenuation and root mean square amplitude and unit thickness. Particularly noteworthy was the imaging of the development of a gas pocket as injection rate increased, and its later decline, with the development of open fluid flow conduits (Roche et al., 2020).

Water sampling: Analyses of carbonate chemistry parameters throughout the water column detected drops in pH values and sharp increases in pCO$_2$ within 8 m of the bubble streams at low tide. The influence of the injected CO$_2$ was confirmed by anomalies in the stoichiometric ratio of O$_2$ to pCO$_2$ relative to baseline values (Esposito et al., 2020; Martínez-Cabanas et al., 2020).

Video-CTD: Video imaging and pCO$_2$ sensor data acquired with the Video-CTD detected visible bubble streams and peaks in sensor signal resulting from the experimentally released CO$_2$. Deviations from the time-varying baseline were related to the injected CO$_2$ flow...
rate, and combined with modelling, which could potentially be used to quantify leakage rate (Gros et al., 2020).

- Benthic chambers: Benthic chambers deployed 0.5–1.0 m away from bubble streams did not detect any flux of experimentally-derived CO₂ in the form of excess DIC across the sediment-water interface, within the limit of detection of the analytical methods. This is tentatively interpreted as a sign that during the early period of CO₂ release, most of the CO₂ escaping into the water column was in gas form (Gros et al., 2020).

- Eddy covariance: The eddy covariance system detected a clear signal of hydrogen ion production due to CO₂ gas dissolution at the lowest CO₂ injection rate and throughout the experiment. It also quantified naturally-occurring biotic CO₂ production in surrounding sediments. Therefore, pH eddy covariance is a highly sensitive technique for the detection of a seafloor source of CO₂ (Koopmans et al., 2020).

- In-situ chemical gradient measurements: This technique showed a strong drop in pH whenever the current direction caused the CO₂-enriched water from the plume to contact the sensors. When coupled with simple models, this technique could quantify the emission of CO₂ into the water column at all but the lowest release rate (Schamp et al., 2020).

- Chemical mapping with the ROV: Each of the three types of pH sensors mounted on the ROV detected the plume. Plume mapping confirmed the highly-localised nature of the plume and the impact of currents on the shape and orientation of the plume. This highlights the importance of considering currents when designing a survey path for vehicles searching for plumes (Monk et al., 2020).

- pH sensors on the AUV: The AUV collected pH data over a much wider spatial area than the ROV but had to maintain an altitude of >4 m above the seabed to avoid the gas rig infrastructure. The pH sensors were not able to detect the plume at this height. This result is consistent with models, which predicted the plume would not be detectable at distances of >2 m above the seabed. However, when the SeafET pH sensor was removed from the AUV and mounted on the ROV it detected the plume, demonstrating that the method should work in an environment where the AUV can maintain a lower altitude.

In summary, during the experiment, the released CO₂ was detectable in its gaseous and dissolved form in both the sediments and the water column. In the sediments close to the bubble streams, the impact of injected CO₂ was detectable based on changes in the pore water chemistry, as a temperature increase in the pore water and in gas form from chrip measurements. Gas bubbles in the water column were detectable optically and acoustically, and CO₂ that dissolved in the water column created a distinctive signal that was detectable chemically by in-situ and lab-based methods.

5.3. Relevance of experimental outcomes for environmental monitoring of offshore subseafloor CO₂ storage

While no acceptable leakage rates have been legislated yet for offshore CO₂ storage, a range of acceptable leakage rates could be estimated using the rate of 0.01 % reservoir loss per year proposed in the literature (e.g., Hepple and Benson, 2005). If applied to the proposed injection rate (~1 Mt/yr) and duration (20 years) at Goldeneye (Dean and Tucker, 2017), this would yield a range of acceptable leakage rates of 274 kg/d after the first year of injection and 5480 kg/d after 20 years of injection when full storage capacity is reached. During the STEMM-CCS release experiment CO₂ gas was injected at flow rates from 6 to 143 kg/d. Around 50 % of the injected CO₂ escaped across the seabed into the water column, which was well below these estimated acceptable leakage rates. The injected CO₂ was detected in all its forms, i.e., gaseous and dissolved in the sediments, and gaseous and dissolved in the water column. As CO₂ leakage is unlikely to be continuous over the whole reservoir area but is rather expected to be preferentially transported through small focused fractures and faults or through poorly-sealed, abandoned wells (IPCC, 2005), both the flow rates and the type (point-release) of our simulated leakage are relevant for real-world scenarios.

A major factor in the design of the experiment was the use of a range of approaches, which mitigated risk and offered a balance between the quantity and quality of the gathered data. The combination of mobile and fixed-position methods provided both high-accuracy data and spatial coverage. Measuring the CO₂ within the sub-seafloor and in its gaseous and dissolved form in the water column allowed quantification of the CO₂ fraction remaining in the sediment versus the CO₂ fraction that escaped into the water column. This knowledge is important for designing future monitoring programmes as well as for informing and improving models.

The radically different approaches, relying on complementary signals from optical, acoustic, and chemical changes, meant that biases or limitations in any one technique could be identified and accounted for. Crucially, in a real-world scenario, the use of complementary approaches will help to maximise the chances of detecting false positives. The combination of novel techniques, adapted versions of existing techniques, and well-proven standard techniques allowed for high-frequency and high-spatial coverage data to be cross-checked and quality-controlled and also made the release experiment a suitable demonstration opportunity for new technologies developed specifically for STEMM-CCS. The use of in-situ technology meant informed decisions could be made in a responsive manner. We anticipate that any future environmental monitoring programme may wish to consider a similarly balanced set of approaches. However, the monitoring technologies developed in STEMM-CCS are just one element (the environmental) of a comprehensive risk-based Measurement, Monitoring and Verification (MMV) programme an operator has to develop, which includes the subsurface and wells (Dean and Tucker, 2017). Consequently, any deployment of the STEMM-CCS technologies will also depend largely on the risk features identified in the subsurface. Many techniques utilised in this experiment are complex and not all the technology is, as yet, commercially available or familiar to the wider research community. To support its dissemination, an online tool for supporting decisions about monitoring techniques has been prepared by the project consortium (Lichtschlag et al., 2020b) and is available online at www.stemm-ccs.eu/monitoring-tool. This tool indicates the level of operational expertise and cost of individual techniques, as well as their spatial and temporal resolution and overall utility.

The plume modelling conducted prior to the field experiment indicated that any CO₂ that escaped across the seabed and into the overlying water column would be dispersed and diluted very rapidly by natural mixing driven by turbulence and tidal motion. As a result, it was expected that the resulting chemical changes would be below the threshold which would present any environmental risk (Blackford et al., 2020). Whilst this result facilitated the consenting process for the experiment, it also highlighted that the deployment of sensors had to be spatially meticulous in order to maximise the utility of the experimental outcomes (see photo of seafloor equipment layout, Fig. 12b).

Given the likely small spatial extent of a plume and its rapid dilution away from its seabed source, future monitoring strategies will need to be supported by model simulations, e.g., coupled hydrodynamic-biogeochemical models (e.g., Blackford et al., 2017; Lessin et al., 2016, 2018; Vielstätte et al., 2019) which can provide optimal deployment strategies for sensors and efficient vehicle trajectories for detection, location, and quantification of leakages (Alendal et al., 2017; Alendal, 2017; Hvidvold et al., 2015; Gundersen et al., 2018; Oleynik et al., 2018).

6. Conclusions

The STEMM-CCS project completed a large-scale field experiment in the central North Sea designed to simulate and detect an unintended
emission of CO₂ from a CO₂ storage site. A unique experimental design enabled the release of CO₂ into the shallow sediments and a range of novel and standard methods were applied to detect and quantify the release of CO₂. The aim of this paper was to synthesise the work done prior to and during the CO₂ release experiment. Overall, the experiment was a major success. The CO₂ release system was able to deliver flow rates over more than an order of magnitude (6–143 kg/d) to test relevant leakage rate scenarios. The outcomes of the experiment demonstrate that a sub-seafloor release of CO₂, mimicking a leakage from an offshore CO₂ storage site, can be reliably achieved and can serve as a test bed for trialing new technologies and techniques. It further demonstrates that such a release can be detected, attributed, and quantified using multiple different approaches.

Author contributions

Conceptualisation and management of overall project was done by EA, GA, CBe, JBl, JBu, BC, AD, DdB, MDea, MHa, MHe, VH, RJ, TL, AL, PL, SL, JM, MM, HH, CS, KS, MS, SS, PW, SW, and DC.


AF and AS co-led the writing of the paper; contributions to writing and revising were also provided by EA, GA, JBl, CBg, SB, JBu, BC, AD, DdB, MDeW, JD, ME, MG, RH, MHe, VH, RJ, DK, EK, TL, JL, AL, SM, SP, CP, BR, KS, JS, PW, and DC.

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Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

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