



Cite this: DOI: 10.1039/d5lc00208g

## A review of microfluidic approaches for carbon capture and storage research

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The reliance on fossil fuels for the increasing global energy demand necessitates the advancement of carbon capture and storage technologies to mitigate anthropogenic CO<sub>2</sub> emissions. However, conventional experimental platforms have reached their practical limitations, stagnating further advancements in optimizing carbon capture and storage (CCS) processes. Microfluidic technologies have emerged as promising tools for investigating and optimizing CCS processes at the microscale, offering precise control over gas–liquid interactions, reaction kinetics, and multiphase flow dynamics, which would be very challenging with conventional macroscale platforms. This review uniquely consolidates carbon capture and storage advancements, providing a structured approach that starts from fundamental mechanisms and systematically progresses to advanced microfluidic-assisted CCS strategies. Unlike prior reports, this review demonstrates how microfluidics complements conventional macroscale approaches and outperforms in certain aspects for studying CO<sub>2</sub> capture and storage through specific experimental examples. The carbon capture section explores microfluidic approaches for physical absorption, chemical absorption, and adsorption, highlighting their advantages over conventional methods. The discussion extends to key mass transfer models, microchannel geometries, and flow regimes that dictate CO<sub>2</sub> dissolution rates and interfacial transport phenomena. The carbon storage section examines microfluidic investigations into carbon mineralization and geological sequestration, providing insights into pore-scale trapping mechanisms, mineralization kinetics, and enhanced storage efficiency. The integration of real-time analytical techniques has further facilitated high-resolution quantification of CO<sub>2</sub> transport and reaction dynamics. Despite significant progress, challenges remain in scaling microfluidic findings to field applications, accurately mimicking reservoir conditions, and developing reactive microfluidic platforms for *in situ* mineralization studies. This review aims to show the pivotal role of microfluidics in accelerating CCS innovation.

Received 28th February 2025,  
Accepted 22nd April 2025

DOI: 10.1039/d5lc00208g

rsc.li/loc

## 1. Introduction

Addressing global warming is among the most critical challenges faced by the scientific community today. With technological advancements driving a steady increase in global energy demand, projected to rise by 28% by 2040, fossil fuels are expected to remain a primary energy source for the foreseeable future.<sup>1</sup> This reliance on fossil fuels results in the emission of substantial greenhouse gases, particularly carbon dioxide (CO<sub>2</sub>), which intensifies climate change by contributing to severe environmental consequences such as rising sea levels and increasingly extreme weather events. According to the 2022 International Energy Agency (IEA) report, global anthropogenic CO<sub>2</sub> emissions reached an

alarming 36.8 gigatons (Gt), highlighting the urgent need for emission mitigation and carbon removal solutions.<sup>2</sup>

In recent decades, researchers have increasingly focused on preventing further CO<sub>2</sub> emissions to the atmosphere. Carbon capture and storage (CCS) technologies have been developed to capture emissions from industries such as coal-based thermal power plants, cement production plants, and steel manufacturing plants, storing the captured CO<sub>2</sub> in stable geological formations to prevent atmospheric release. CCS has made significant strides since the 1970s, with early efforts rooted in amine scrubbing – a chemical absorption process that remains foundational in industrial CO<sub>2</sub> capture due to its high capture efficiency and reliability.<sup>3</sup> Another landmark project, the Sleipner Project in Norway, initiated in 1996, demonstrated the feasibility of large-scale CO<sub>2</sub> storage by successfully storing over 1 million tons of CO<sub>2</sub> annually in an underground sandstone formation, setting a precedent for modern geological storage initiatives.<sup>4</sup> Building on these foundational technologies, the field of CCS has been expanded to somewhat contemporary methods such as oxy-

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fuel combustion, membrane separation, metal–organic frameworks (MOFs), and the use of ionic liquids and nanofluids.<sup>5–9</sup> Additionally, storage technologies of captured carbon, such as CO<sub>2</sub>-based enhanced oil recovery (EOR) and carbon storage in saline aquifers, have parallelly been developed.<sup>10</sup> EOR is a dual-purpose approach, where injected CO<sub>2</sub> boosts oil production while facilitating long-term underground CO<sub>2</sub> sequestration. Each technique offers unique advantages and challenges, but all share a common goal of reducing atmospheric CO<sub>2</sub> levels. These methods have been comprehensively documented and are being improved constantly. Most researchers in these fields have raised concerns about some drawbacks despite their advancements.

In the field of carbon capture, most of the solvents or chemicals are corrosive, non-selective, and require large amounts of energy to regenerate the captured carbon.<sup>11</sup> Conventional macroscale experimental methods in CCS research would be impractical in resolving these issues as they would require comprehensive parametric studies for characterization using an extensive number of resources and labor work. Additionally, the conventional macroscale experimental methods face challenging analytical issues like uncertainties in the characterization of the CO<sub>2</sub> capture process due to the indistinct gas–liquid interface, inconvenient or impossible measurements of physical dissolution, and chemical reactions of CO<sub>2</sub> in solvent due to complex kinetic and thermodynamic properties. Similarly, in the carbon storage field, the existing methods face the challenges of slow reaction kinetics in carbon mineralization, the uncertainty of long-term safe CO<sub>2</sub> storage stability due to the risk of leakage, and huge capital and monitoring costs.<sup>12–14</sup> Conventional research methods like core tests, field pilot studies, computation studies, and batch reactor experiments are not fully qualified to work on these challenges because these methods would require extensive parametric studies, substantial resources, and intensive labor for characterization. Additionally, they would face analytical difficulties in accurately monitoring the CO<sub>2</sub> trapping mechanism due to complex multiphase flow dynamics, indistinct fluid–rock interactions, and the inability to capture real-time pore-scale dissolution and mineralization processes in geological formations. These limitations highlight the urgent need to adopt advanced technological research methods like microfluidics to overcome existing challenges and move towards a more sustainable and effective solution for carbon capture and storage.

Adopting microfluidic technology improves research efficiency in both carbon capture and storage. For carbon capture, microfluidics enables a well-defined gas–liquid interface at the microscale, allowing precise analysis of CO<sub>2</sub> absorption and desorption.<sup>15</sup> It facilitates rapid solvent screening and optimization with minimal material usage, significantly reducing time and resource requirements compared to conventional methods. For carbon storage, microfluidic platforms replicate a wide range of complex geological structures, enabling the study of key mechanisms

such as mineralization, capillary trapping, and dissolution under controlled conditions. These systems provide real-time monitoring of multiphase flow and reactive transport, offering insights that are difficult to achieve with traditional experimental methods. Overall, microfluidics accelerates CCS research by enhancing control, significantly reducing time and cost, and improving experimental observation techniques. Acknowledging these advantages over conventional methods, researchers are increasingly shifting toward microfluidic platforms for CCS-based research.

This increasing interest in microfluidic-based CCS research is reflected in the growing number of publications over the past decade. According to our trend search from Scopus, the number of research articles on microfluidic approaches for CCS has steadily increased from 2017 to 2024, as shown in Fig. 1. This trend highlights the growing recognition of microfluidics as a valuable tool for advancing carbon capture and storage technologies. Given the growing amount of literature in this field, there is a need to consolidate this work and provide a comprehensive understanding of microfluidic contributions to CCS research. Several recent review articles have addressed various aspects of CCS technology-based microfluidic research. For example, Abolhasani *et al.* explored early applications of segmented flow microfluidics in CO<sub>2</sub> dissolution studies, especially focused on chemical absorption processes, CO<sub>2</sub> utilization in materials science, and supercritical CO<sub>2</sub> as a solvent.<sup>16</sup> Cheng *et al.* provided an in-depth review of microfluidic-based chemical absorption technology for CO<sub>2</sub> capture, discussing mass transfer dynamics, gas–liquid flow patterns, and the optimization of absorber design for improved performance.<sup>17</sup> Liu *et al.* highlighted the application of high-throughput droplet microfluidics in producing monodisperse microcapsules for carbon capture, focusing on fabrication techniques and the scalability of microfluidic approaches for large-scale CO<sub>2</sub> mitigation.<sup>18</sup> Pan *et al.* reviewed the

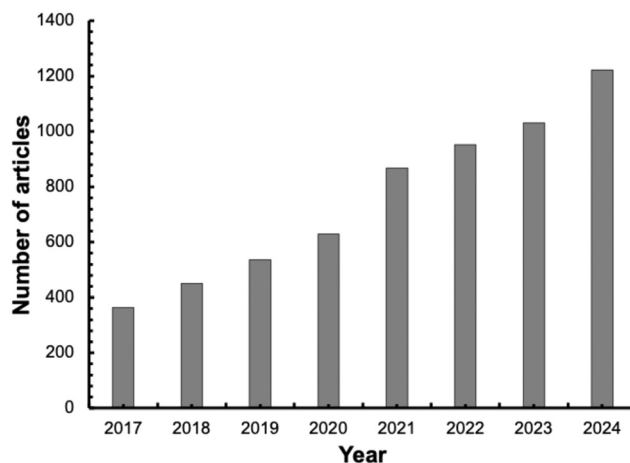


Fig. 1 A number of literature reports produced on microfluidic-based carbon capture and storage research (Scopus analytics results with the keywords “CCS”, “microfluidic”, “carbon capture”, and “carbon storage”).



advancement in micro and nano scale studies for CCS applications, summarizing the potential of *in situ* visualization and understanding of gas–liquid interactions in confined domains using microfluidics, which is crucial for optimizing CO<sub>2</sub> capture and storage processes.<sup>19</sup> While these reviews provide valuable insights into specific aspects of microfluidic technology, a well-balanced study consolidating the primary advancements is needed to present a comprehensive perspective on how microfluidic technology outperforms conventional methods across all key aspects of CCS research. This review aims to consolidate the overall microfluidic research in both fields of carbon capture and carbon storage research. Unlike prior studies that solely focused on solvent selection, gas–liquid interaction, microfluidic fabrication, flow dynamics, or comprehensive collection of experimental research on microfluidics, our review integrates these aspects with experimental examples, directly comparing microfluidic approaches to conventional macroscale platforms. By systematically evaluating the advantages of microfluidics, we provide a clear framework for its benefits over traditional methods. Additionally, we incorporate qualitative analyses of key studies demonstrating how advancements in solvent optimization, surface wettability manipulation, and real-time reaction monitoring have driven significant breakthroughs in CCS research. This work not only contextualizes existing research but also identifies emerging challenges for future innovations.

In this review, we have presented a structured approach, starting from the fundamental mechanism of CCS and progressing systematically to the advanced methodologies of microfluidic platform applications, which ensures an accessible resource to most of the researchers in these fields. We divide this review into two parts: microfluidics-assisted carbon capture and microfluidics-assisted carbon storage research. Section 2 discusses the mechanisms of carbon capture elucidated by microfluidic approaches. After introducing specific examples and strategies for capturing CO<sub>2</sub>, such as physical absorption, chemical absorption, and adsorption, the review transitions into detailed discussions of mass transport theories, channel designs, and measurement techniques that influence gas–liquid interaction efficiency. These subsections provide practical guidance for selecting appropriate theoretical models (*e.g.*, Fickian diffusion, two-film theory) and interpreting experimental observations based on microchannel geometry and flow behavior. These insights not only clarify how microfluidics can be optimized for carbon capture but also serve as a bridge to section 3, where the same tools and principles are adapted to study CO<sub>2</sub> behavior in porous media and geological environments. Section 3 focuses on carbon storage research using microfluidic platforms, covering different CO<sub>2</sub> storage approaches like carbon mineralization, storage in saline aquifers, and enhanced oil recovery. In this section, we demonstrate how the core ideas developed in section 2, such as interfacial transport analysis, flow visualization techniques, and channel design, are

extended to evaluate mineralization kinetics, pore-scale trapping, and flow behavior in realistic subsurface analogs. Furthermore, we review various microfluidic models used in these studies, along with their flow dynamics and measurement techniques. Lastly, we provide a comprehensive summary, highlighting challenges and future directions for microfluidic applications in CCS research.

## 2. Role of microfluidics in carbon capture

### 2.1. Carbon capture mechanisms

Many carbon separation methods include chemical absorption, physical absorption, adsorption, membrane separation, chemical-looping combustion, and biological and cryogenics.<sup>20</sup> However, carbon capture mechanisms can be primarily categorized into three types: physical absorption, chemical absorption, and adsorption.<sup>21</sup> Each operates on distinct principles to retain CO<sub>2</sub> molecules in a capturing medium and has advantages and limitations. Advancing research in these mechanisms to address their limitations using macroscale platforms presents major challenges, such as high solvent consumption, inefficient control over mass transfer dynamics, and difficulty in precisely monitoring reaction kinetics. These constraints hinder the development of optimized or novel capture materials and techniques. Given these limitations, microfluidic technology has been increasingly adopted to advance carbon capture research. It offers several advantages, including precise control over fluid dynamics, high surface-area-to-volume ratios, well-defined gas–liquid interfaces for effective interaction studies, efficient material usage, and improved accessibility for analyzing reaction kinetics. To provide a strong foundation for understanding the role of microfluidics in carbon capture, it is essential to first discuss the primary CO<sub>2</sub> capture mechanisms and their associated research limitations, as well as how microfluidic technology offers solutions to these challenges. This section presents a detailed explanation of these mechanisms and examines how conventional macroscale research is adapted to microfluidic-based approaches to overcome existing constraints.

**2.1.1. Physical absorption.** Physical absorption is the process of dissolving CO<sub>2</sub> in a liquid solvent through physical contact without altering the chemical composition of the CO<sub>2</sub> molecule, which is mainly driven by pressure and temperature.<sup>22</sup> This process occurs naturally in open bodies of water, where CO<sub>2</sub> dissolves at the surface and establishes concentration equilibrium with the environment.<sup>23</sup> The dissolved CO<sub>2</sub> can easily return to the atmosphere when exposed to heat (*e.g.*, sunlight), lower pressure, or reduced atmospheric CO<sub>2</sub> concentration, and the absorption rate is constrained by the solvent's surface area.<sup>24</sup> While physical absorption is straightforward and cost-effective, it is only efficient in industrial settings at high pressures, significantly increasing operational costs. However, one of the main advantages of physical absorption in the industrial CO<sub>2</sub> capture



process is its reversibility.<sup>22</sup> Since the CO<sub>2</sub> remains chemically unchanged, it can be removed from the solvent relatively easily, allowing the solvent to be reused. This process is energy-efficient compared to chemical absorption methods like amine scrubbing, which requires an average of four times more energy for solvent regeneration.<sup>25</sup> Established physical absorption methods, such as the Selexol, Rectisol, and Purisol processes, use various solvents and operate under different temperature and pressure conditions, but all share the limitation of high-pressure requirements for effective CO<sub>2</sub> capture.<sup>26</sup> To overcome this major limitation, the need to develop new physical solvents that can operate under lower pressure with improved or at-par absorption capacity has been raised. Recent research shows the potential of several innovative solvents, including ionic liquids, deep eutectic solvents (DESs) derived from choline chloride and urea, nanofluid-based solvents, where nanoparticles like silicon dioxide (SiO<sub>2</sub>) and aluminum oxide (Al<sub>2</sub>O<sub>3</sub>) are dispersed in conventional solvents, and hybrid solvents, combining physical and chemical absorption properties, such as dimethyl ether of polyethylene glycol (DEPG) mixed with amines.<sup>27,28</sup> Conducting rigorous studies on these new solvents using traditional macroscale platforms would have been challenging as it would require large solvent volumes, reduced precision and control during the mass transfer and reaction kinetics studies, and long real-time monitoring would lead to poor reproducibility. In contrast, microfluidic platforms have proven more practical for studying these solvents.

For example, Lefortier *et al.* conducted a study to screen the CO<sub>2</sub> solubility and diffusivity in various ionic solvents such as poly(ethylene glycol)dimethyl ether (Depeg), 1-octanol, 1-methyl-2-pyrrolidinone (NMP), propylene carbonate (PC) and 1-methyl-3-butylimidazolium bis(trifluoromethylsulfonyl)imide.<sup>29</sup> As shown in Fig. 2, they used a segmented flow-based microchannel with a T-junction to generate monodisperse CO<sub>2</sub> bubbles in proposed solvents; they achieved rapid determination of CO<sub>2</sub> solubility and diffusivity in each solvent. By adopting a microfluidic platform, they reduced the experimental time to 5 minutes per sample, which typically takes hours per solvent batch with macroscale platforms. Moreover, the microfluidic

experimental measurements were found to be accurate compared with the literature values for pure solvents. This research demonstrates the superiority of microfluidic techniques with respect to time and volume requirements to conduct such studies.

With a similar microfluidic experimental technique, Voicu *et al.* explored a novel approach to CO<sub>2</sub> capture using frustrated Lewis pairs (FLPs) within a microfluidic system.<sup>30</sup> FLP theory involves a Lewis acid and a Lewis base that are too bulky to bond, so they stay “frustrated” and reactive. This reactivity allows them to capture and activate small molecules like CO<sub>2</sub>. In this microfluidic study, Voicu *et al.* demonstrated an efficient, low-energy CO<sub>2</sub> capture by enhancing control over various reaction conditions. FLPs provide a promising route for CO<sub>2</sub> capture from direct emissions and for direct air capture (DAC), where the CO<sub>2</sub> concentrations are significantly low. To have an efficient FLP system for different acids and bases, the conventional methods would take days, along with bulk solvents, to produce a similar outcome.

Similarly, Zhao *et al.* conducted a rigorous experimental study on DES solvents with the enzyme carbonic anhydrase to increase the reaction rate (Fig. 3).<sup>31</sup> They developed a microfluidic encapsulation platform to produce microcapsules of DES solvents with precisely controlled amounts of carbonic anhydrase. They found that the homogenous distribution of enzymes is key to gaining higher mass transfer efficacy with the DES solvent. Previously, similar attempts were made to use enzymes like carbonic anhydrase in DES using a macroscale. However, due to the heterogeneous distortion of enzymes in DES solvent in a macroscale setup, poor reaction rates for carbon capture were yielded. The microcapsules showed a 253-fold increase in CO<sub>2</sub> absorption rate and a 74-fold increase in absorption capacity compared to neat DES.

**2.1.2. Chemical absorption.** Chemical absorption captures CO<sub>2</sub> by forming chemical bonds with a solvent, typically an alkaline solution, which results in stable compounds such as bicarbonates.<sup>32</sup> This method is highly effective in separating CO<sub>2</sub> even at low pressure, making it a popular choice for post-combustion capture applications.<sup>33</sup> However, the chemical absorption process requires specific conditions,

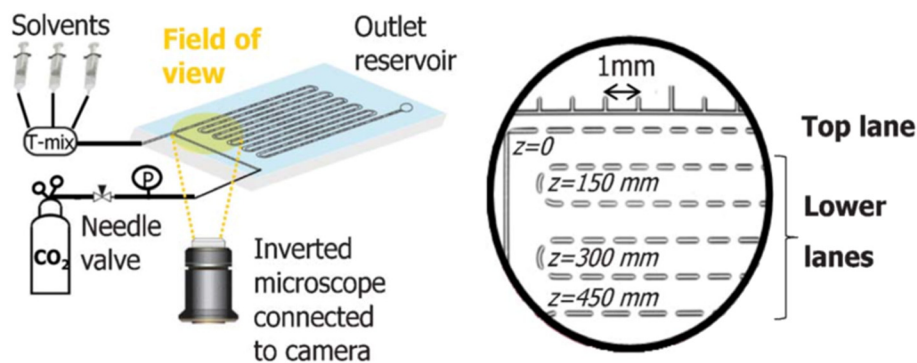
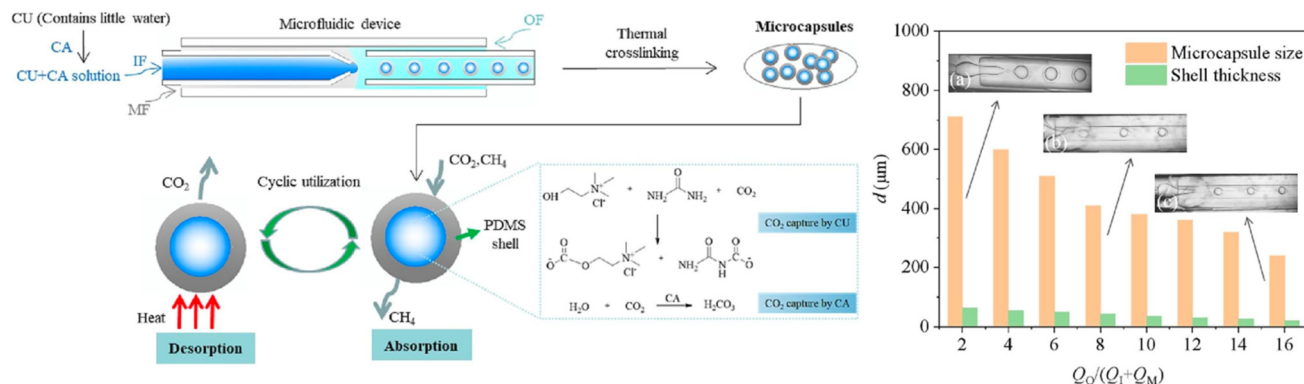


Fig. 2 Schematic representation of the experimental setup with the field of view. In the field of view, the top lane shows bubble shrinkage due to absorption; the lower lanes show slow expansion due to pressure drop.<sup>29</sup>







**Fig. 3** [Left] Schematic diagram of the flow-focusing microchannel, encapsulating DES and CA. IF – inner fluid; MF – middle fluid; OF – outer fluid. [Right] Effect of the flow rates on the microcapsule size and shell thickness in a 3D glass-capillary microfluidic device (in the graph, “d” as a function of  $Q_O/(Q_I + Q_M)$  – flow rate of inner fluid;  $Q_M$  – flow rate of middle fluid;  $Q_O$  – flow rate of outer fluid: figures (a)–(c) show actual experimental images of microcapsules with varying sizes and shell thicknesses).<sup>31</sup>

including controlled temperature and concentration, and is highly energy-intensive due to the need for solvent regeneration.<sup>34</sup> Traditional chemical absorption methods, like amine scrubbing with monoethanolamine (MEA), consume significant energy and materials, limiting their sustainability at industrial scales.<sup>35</sup> To overcome these limitations, new chemical absorption-based solvents have been developed with lower volatility and environmental impact, such as sterically hindered amines, functionalized ionic liquids, and polymeric amine-based solvents.<sup>36</sup> The thorough study of these newly developed solvents can be challenging with a macroscale platform, as it will require more time and solvent volumes. Additionally, chemical absorption-based solvents pose more challenges on the macroscale for several reasons. First, the chemical absorption process goes through a series of chemical reactions, often with multi-step kinetics, such as the formation of carbamates, carbonates, or bicarbonates. In macroscale reactors, tracking reaction kinetics becomes more challenging due to the heterogeneous mixing and variable gas-liquid contact. On the other hand, using a microfluidic approach, uniform mixing conditions and accurate monitoring of reaction kinetics can be achieved.<sup>37</sup> Second, chemical absorption reactions are exothermic, which leads to localized temperature increases, which can cause poor reproducibility and unreliable measurements. Third, due to bulk volume and long-time scale requirements of macroscale reactors, side reactions and unwanted byproducts can be generated during the  $\text{CO}_2$  dissolution study, leading to inconsistent results. Using microfluidic platforms with a shorter time and smaller volume requirement, the side reactions and unwanted byproduct generations can be reduced or avoided.<sup>38</sup> Because of these benefits, chemical absorption-based  $\text{CO}_2$  capture research has adopted microfluidic platforms.

For instance, Hallenbeck *et al.* used a glass microfluidic reactor to compare the  $\text{CO}_2$  absorption capacity and absorption rate of several amino acid salt-based solvents like lysinate (LYS), glycinate (GLY), taurine (TAU), and

proline (PRO). Using a tailored  $\text{CO}_2$  microbubble flow in solvent (Fig. 2), each solution was tested and compared with MEA for its  $\text{CO}_2$  absorption capacity. They found that potassium lysinate (LYS) exhibited a 50% higher  $\text{CO}_2$  absorption capacity than MEA, with a maximum  $\text{CO}_2$  loading of 1.32 mol  $\text{CO}_2$  per mol lysine *versus* 0.68 mol for MEA. Additionally, the time to reach 90% of maximum  $\text{CO}_2$  loading was the shortest for MEA (0.30 seconds), followed closely by GLY and PRO (0.34 seconds), while LYS took longer (0.47 seconds) but showed superior long-term absorption stability. With a simple microfluidic platform, this time-dependent study can be done, which otherwise would require a very complex macroscale setup to analyze such a minute time-dependent analysis.

**2.1.3. Adsorption.** Adsorption captures  $\text{CO}_2$  by allowing it to adhere to the surface of solid adsorbents, such as zeolites, MOFs, and activated carbon.<sup>39</sup> Adsorption is valued for its reversibility, as it allows  $\text{CO}_2$  to be released and the adsorbent to be reused with minimal energy input. Conventional research on adsorption-based carbon capture has been predominantly conducted on macroscale platforms like batch reactors, packed beds, and pressure swing adsorption (PSA) systems.<sup>40,41</sup> In these systems, the pore structure is made from basic functional groups or Lewis basic sites, prone to adsorb  $\text{CO}_2$  gas molecules by weak intermolecular bonds. Adsorption-based carbon capture research poses a few challenges, like poor reproducibility due to non-uniform adsorption of  $\text{CO}_2$  gas. Microfluidic devices have been adopted to ensure more homogeneous active sites in synthesizing adsorbent materials for  $\text{CO}_2$  adsorption. We want to clarify that microfluidics is primarily utilized for the synthesis of adsorbent materials (such as N-doped carbon spheres and porous metal oxides) rather than for the fabrication of large-scale carbon capture structures onto which these materials are mounted or coated, such as packed beds or monolithic substrates. While microfluidic platforms enable precise control over particle morphology, porosity, and surface chemistry during material synthesis, the integration of these materials onto larger surfaces for practical deployment still relies on conventional manufacturing



techniques.<sup>42,43</sup> This distinction is often overlooked in the literature, leading to misconceptions about the role of microfluidics in adsorption-based carbon capture research.

Several studies have demonstrated the substantial benefits of microfluidic platforms for enhancing CO<sub>2</sub> capture on adsorption-based mechanisms. Jin *et al.* conducted a study in which a microfluidic channel was used to synthesize N-doped microporous carbon spheres for CO<sub>2</sub> capture.<sup>44</sup> N-doping (nitrogen doping) introduces nitrogen atoms into the carbon framework, enhancing the material's CO<sub>2</sub> capture capacity by creating basic sites that interact strongly with acidic CO<sub>2</sub> molecules. This modification not only improves CO<sub>2</sub> selectivity but also enhances the thermal stability of the adsorbent. To obtain it, they used nanodroplet templating, where tiny oil droplets acted as templates for creating a network of uniform micropores (Fig. 4). After synthesis and carbonization, the oil droplets were removed, leaving behind highly porous structures that increased the surface area and active sites for CO<sub>2</sub> adsorption. In conventional macroscale systems, such as batch reactors, synthesizing porous carbon typically results in particles with inconsistent sizes and pore structures. This inconsistency reduces adsorption efficiency and leads to non-uniform gas–solid interactions. Furthermore, macroscale processes often require chemical activation and additional post-synthesis steps, which are time-consuming and environmentally harmful. In contrast, Jin *et al.* produced highly uniform carbon spheres with a Brunauer–Emmett–Teller (BET) surface area of 576 m<sup>2</sup> g<sup>−1</sup> and a microporous volume of 0.22 cm<sup>3</sup> g<sup>−1</sup>, significantly improving the adsorption capacity from 0.42 mmol g<sup>−1</sup> (untreated carbon spheres) to 1.20 mmol g<sup>−1</sup>, a nearly 200% improvement (Fig. 4).<sup>44</sup> Additionally, the microfluidic process enabled continuous production with real-time control over the synthesis, making it more scalable and sustainable.

Similarly, Xie *et al.* advanced the concept by incorporating a self-templated synthesis of N-doped hierarchical porous carbon spheres (in a self-templated synthesis, the precursor

decomposes or transforms into carbon, creating multiple types of pores).<sup>45</sup> This process used ZIF-8 (a type of metal–organic framework) as a sacrificial template, enabling *in situ* nitrogen doping during pyrolysis (Fig. 5). The resulting carbon spheres exhibited a hierarchical pore structure with a specific surface area (SBET) of 886.9 m<sup>2</sup> g<sup>−1</sup> and superior CO<sub>2</sub> adsorption capacity of 1.29 mmol g<sup>−1</sup> at 30 °C and 1 bar.<sup>45</sup> Compared to macroscale synthesis, which often yields particles with a wide size distribution and high diffusion resistance, the microfluidic system ensured monodisperse particles with optimized mesoporous structures (Fig. 5), enhancing selective CO<sub>2</sub> adsorption in mixed gas streams.

Another innovative example is the work published by Kurlov *et al.*, who developed CaO-based CO<sub>2</sub> sorbents with a hierarchical porous structure using microfluidic droplet templating.<sup>46</sup> In calcium looping processes commonly used for industrial CO<sub>2</sub> capture, the sintering and attrition of CaO particles reduce their long-term stability and adsorption capacity. By employing microfluidic emulsification to create monodisperse oil droplets as templates, they synthesized a sorbent with a stable hierarchical structure, which enhanced mass transfer and improved cyclic stability. The CO<sub>2</sub> uptake capacity of the microfluidic-fabricated sorbent was 140% higher than that of conventional limestone-based sorbents, making it highly suitable for large-scale applications.<sup>46</sup> Similarly, Yu *et al.* presented a novel in-capsule synthesis of MOFs for CO<sub>2</sub> capture using a double-capillary microfluidic assembly (Fig. 6).<sup>47</sup> In conventional systems, deploying MOFs as loose particles leads to operational challenges, such as high-pressure drops and particle aggregation. The microfluidic approach allowed MOFs to be synthesized directly inside gas-permeable microcapsules, preserving their high surface area while providing protection from moisture and preventing particle agglomeration. This configuration significantly improved CO<sub>2</sub> adsorption efficiency and operational stability, addressing the limitations of traditional MOF-based systems.<sup>47</sup>

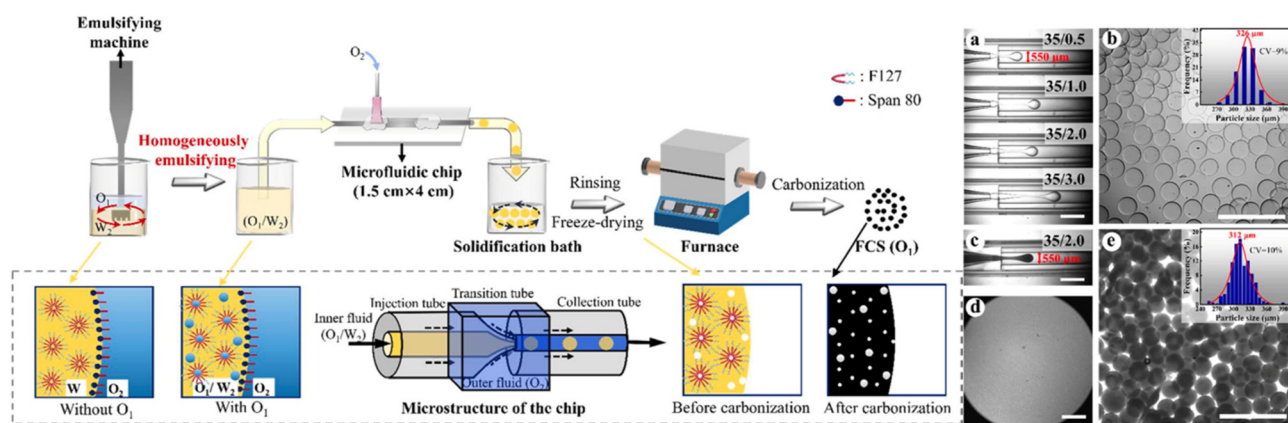


Fig. 4 [Left] Illustrative image of the fabrication of microporous carbon spheres. [Right] (a) Experimental images showing the controllable formation of emulsions at varying flow rates (values in each photo indicate the outer-to-inner phase ratio), (b) optical micrographs of solidified 2% FS with an inset displaying the particle size distribution, (c) formation of double emulsions, (d) optical micrographs of nanodroplets dispersed in water, and (e) optical micrographs of solidified 2% FS with 10% oil phase, with insets showing the corresponding particle size distributions. Scale bars: 1000 μm.<sup>44</sup>



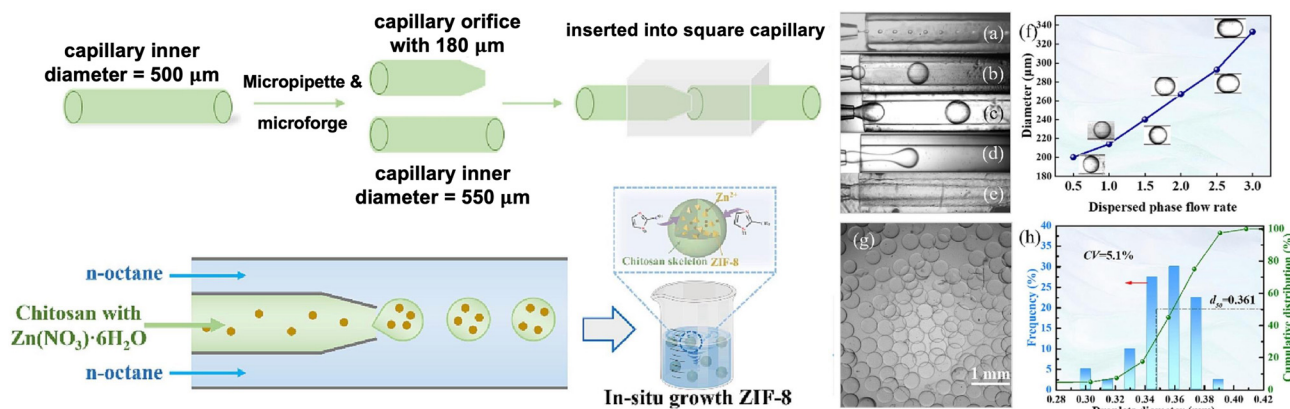


Fig. 5 [Left] Fabrication of chitosan-based microparticles using a capillary microfluidic device and the formation of chitosan/ZIF-8 composites. [Right] The influence of different dispersed phase flow rates on droplet formation, with optical images shown for (a) 0.2  $\text{mL h}^{-1}$ , (b) 1  $\text{mL h}^{-1}$ , (c) 2  $\text{mL h}^{-1}$ , (d) 4  $\text{mL h}^{-1}$ , and (e) 6  $\text{mL h}^{-1}$ . (f) The effect of the continuous and dispersed phase flow rate ratio on droplet diameter (continuous phase flow rate = 10  $\text{mL h}^{-1}$ ). (g) Optical micrograph and (h) particle size distribution of chitosan microdroplets.<sup>45</sup>

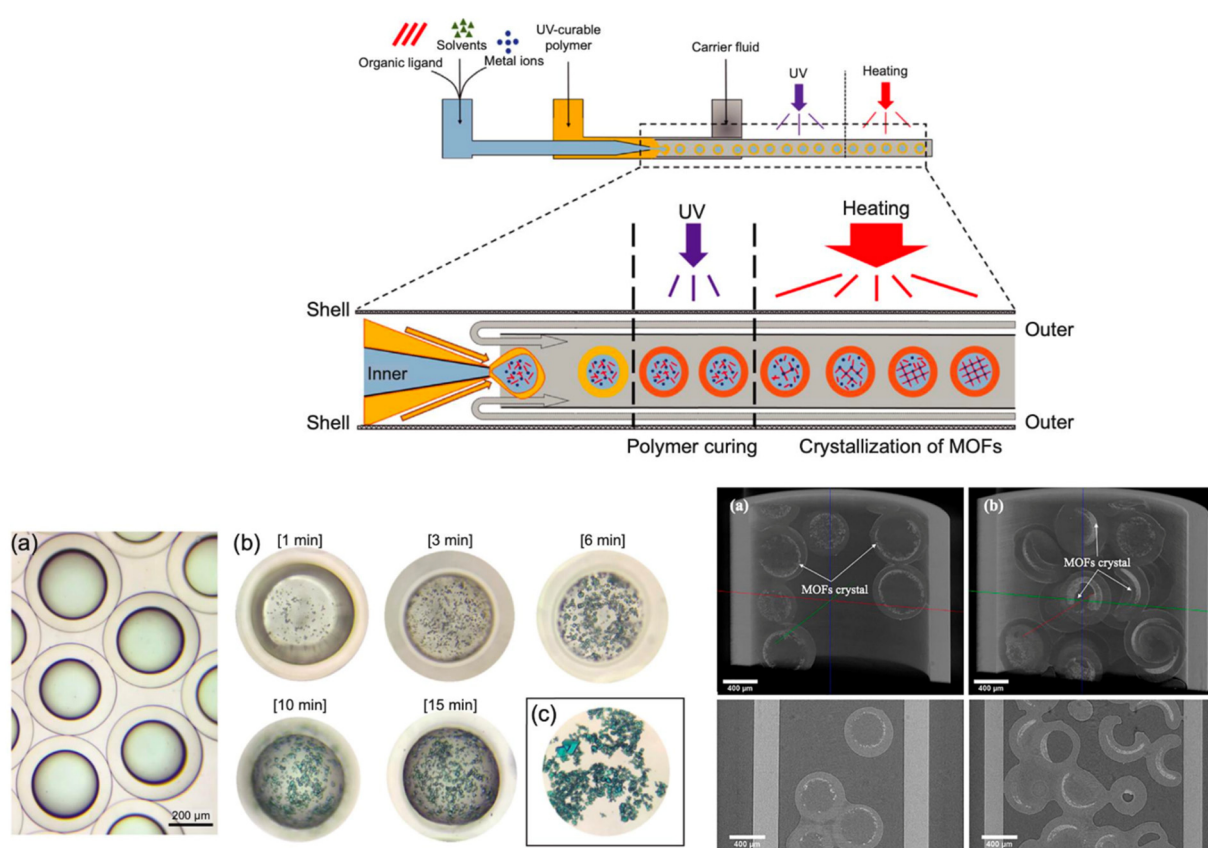


Fig. 6 [Top] Schematic representation of the microfluidic system used for the *in situ* encapsulated synthesis of MOFs, where the shell material TEGO Rad 2650 is structurally similar to TEGO Rad 2100 and TEGO Rad 2500. [Bottom left] Microscopy images of encapsulated HKUST-1 crystals: (a) microcapsules containing precursor solutions of copper(II) nitrate trihydrate ( $\text{Cu}(\text{NO}_3)_2 \cdot 3\text{H}_2\text{O}$ ) and 1,3,5-benzenetricarboxylic acid ( $\text{H}_3\text{BTC}$ ) dissolved in *N,N*-dimethylformamide (DMF) before heating, (b) sequential stages of HKUST-1 crystal formation during the *in situ* thermal reaction within the microcapsules over a reaction time of 1–15 minutes, and (c) HKUST-1 crystals synthesized *via* the conventional solvothermal method in bulk solution for comparison. [Bottom right] three-dimensional X-ray microtomography of encapsulated HKUST-1 within a glass tube: (a) microcapsules containing HKUST-1 and residual DMF solvent, and (b) dried microcapsules after solvent removal.<sup>47</sup>

**2.1.4. Distinct microfluidic roles across  $\text{CO}_2$  capture mechanisms.** While previous subsections have outlined microfluidic strategies for physical absorption, chemical

absorption, and adsorption, it is equally important to highlight how these approaches are tailored to address uniquely different core challenges in each mechanism. In



physical absorption, microfluidics is primarily used to improve gas–liquid mass transfer and rapidly screen solvents through controlled interfacial dynamics. In chemical absorption, where reaction kinetics and solvent–CO<sub>2</sub> interactions are central, microfluidics acts as a miniaturized reactor that enables precise flow control and real-time monitoring. For adsorption, the focus shifts to material

design, and microfluidic platforms serve as synthesis tools for tuning particle morphology and surface area to enhance CO<sub>2</sub> uptake. Despite sharing the same underlying platform, the role of microfluidics diverges significantly across this CO<sub>2</sub> capture mechanism, serving as a measurement tool in physical absorption, a reaction environment in chemical absorption, and a fabrication system in adsorption. This

**Table 1** Comparison of different microfluidic studies based on their carbon capture mechanism

Mechanism	Microfluidic strategy	Microfluidic purpose and outcomes	Functional category	
Physical absorption	Segmented flow microchannel (T-junction) with CO <sub>2</sub> bubbles in ILs and PC	CO <sub>2</sub> solubility and diffusivity determined in less than 5 min per sample. The measurements were validated against literature values. CO <sub>2</sub> loading findings were from 0.8–1.3 mmol g <sup>-1</sup> in different solutions	Mass transfer enhancement and solvent selection	29
	Bubble dissolution analysis in oil-filled microchannels	Dissolution patterns were used to derive interfacial mass transfer rates. The modeled interfacial transfer rate was approximately 10 <sup>-4</sup> m s <sup>-1</sup>		48
	Reservoir-specific diffusivity studies using microfluidic chips	CO <sub>2</sub> diffusivity was quantified in brine and oil systems using <i>in situ</i> visualization. Diffusivity values were found to be 1.5 × 10 <sup>-9</sup> m <sup>2</sup> s <sup>-1</sup> in brine and 3.2 × 10 <sup>-10</sup> m <sup>2</sup> s <sup>-1</sup> in oil		49
	Microbubble visualization in DES and PEGDME solvents	K <sub>La</sub> values were compared across five solvents. Deep eutectic solvents (DES) showed a 1.5× improvement in transfer rate. The K <sub>La</sub> of the DES system was 0.0031 s <sup>-1</sup> , with visualization completed in 90 seconds		24
	Model-based IL flow simulation in pressure-tunable microchannels	A trade-off between CO <sub>2</sub> solubility and energy efficiency was analyzed under pre-combustion conditions. The estimated energy cost for absorption using pressurized ionic liquids was 32 MJ kg <sup>-1</sup> of CO <sub>2</sub>		25
Chemical absorption	Microencapsulation of carbonic anhydrase in DES <i>via</i> glass microfluidics	Microfluidic encapsulation of carbonic anhydrase in DES led to a 253-fold increase in CO <sub>2</sub> absorption rate. The enzyme was uniformly distributed, and the CO <sub>2</sub> loading reached 2.8 mmol g <sup>-1</sup>	Reaction kinetics of solvent and CO <sub>2</sub>	31
	Frustrated Lewis pairs tested under segmented microdroplet flow	Low-energy CO <sub>2</sub> activation was demonstrated at approximately 0.1 bar using frustrated Lewis pairs under segmented droplet flow in a microfluidic setup		30
	SpectIR-fluidic reactor for FTIR absorption tracking	Real-time monitoring of CO <sub>2</sub> bond conversion kinetics was performed using FTIR within a microfluidic reactor. Bond changes were observed within 5 seconds, and the reaction yield was tracked spectroscopically		38
	Functionalized ILs in microchannels with CO <sub>2</sub> gas flow	Thermodynamic performance of functionalized ionic liquids was evaluated in microchannels. The reaction enthalpy was -35 kJ mol <sup>-1</sup> , and the viscosity was measured at 79 cP at 25 °C		35
	SH-amine testing under microfluidic flow	Sterically hindered amines achieved faster CO <sub>2</sub> uptake than MEA, completing absorption within 3 seconds. The solvents also showed reduced degradation and reached a CO <sub>2</sub> capture of 1.5 mol mol <sup>-1</sup>		36
Adsorption	Real-time spectroscopy in glass-PDMS reactor	Absorption curves for test solvents were plotted in under 10 seconds using real-time spectroscopy. The feedback loop resolved dynamic changes with an accuracy of ±3%		37
	Nanodroplet-templated synthesis of N-doped microporous carbon	N-doped carbon produced <i>via</i> droplet templating achieved a BET surface area of 576 m <sup>2</sup> g <sup>-1</sup> . CO <sub>2</sub> uptake increased threefold compared to unmodified carbon, reaching 1.2 mmol g <sup>-1</sup>	Material synthesis with pore/morphology control	44
	Self-templated porous carbon spheres in flow-focusing chip	Self-templated porous carbon spheres generated in a flow-focusing device achieved a surface area of 886.9 m <sup>2</sup> g <sup>-1</sup> . CO <sub>2</sub> adsorption was measured at 1.29 mmol g <sup>-1</sup> at 30 °C		45
	Hierarchical CaO sorbents <i>via</i> monodisperse emulsion templates	Hierarchical CaO sorbents created using monodisperse emulsion templates showed a 140% increase in CO <sub>2</sub> capacity compared to conventional CaO and remained stable for 25 operational cycles		46
	In-capsule MOF synthesis <i>via</i> a capillary double emulsion device	MOFs synthesized within capillary double-emulsion droplets retained 95% of their surface area and showed a twofold improvement in humidity resistance compared to bulk materials		47
	MOF-coated fibers synthesized in microreactors	MOF-coated fibers fabricated in microreactors achieved a surface area of 1745 m <sup>2</sup> g <sup>-1</sup> and demonstrated CO <sub>2</sub> uptake of 2.1 mmol g <sup>-1</sup> at 298 K		42
	Double-emulsion capsule with internal MOF synthesis	Microcapsules with internally synthesized MOFs retained 95% porosity after five humid cycles and exhibited CO <sub>2</sub> uptake up to 1.9 mmol g <sup>-1</sup> under ambient conditions		47





targeted functional use can be seen in the experimental studies discussed in the previous subsections. To further illustrate this distinction, we have compiled representative microfluidic studies in Table 1, each demonstrating how the strategy employed aligns with the specific objective of the corresponding CO<sub>2</sub> capture mechanism.

## 2.2. Mass transfer models

With an understanding of various carbon capture methods and the adoption of microfluidics for research advancement, we now discuss specific aspects of the microfluidic approach, such as mass transfer models to analyze CO<sub>2</sub> dissolution, flow patterns, microfluidic channel design, and measurement techniques for CO<sub>2</sub> dissolution. Selecting the appropriate transport model is critical for accurately quantifying dissolution rates, as different models account for varying transport mechanisms, including diffusion, convection, and interfacial dynamics. Improper model selection can lead to significant deviations in mass transfer predictions, affecting the optimization of microfluidic devices for CO<sub>2</sub> capture. Various models have been developed to describe CO<sub>2</sub> transport in microfluidic systems, each addressing different aspects of mass transfer varying from a basic diffusion process to complex interactions at the gas–liquid interface. This section presents key models used in CO<sub>2</sub> microfluidic capture, along with experimental research examples demonstrating their applications. This section aims to help researchers determine the most suitable model for their specific system parameters and experimental conditions in the microfluidic approach.

**2.2.1. Fick's laws of diffusion.** Fick's laws describe the process of molecular diffusion, which is fundamental in understanding mass transfer in diffusion-dominated environments such as microchannels with low or stagnant flow. Here, eqn (1) shows steady-state diffusion, whereas eqn

(2) shows transient diffusion. Here,  $J$ : diffusion flux (mol m<sup>-2</sup> s<sup>-1</sup>),  $D$ : diffusion coefficient (m<sup>2</sup> s<sup>-1</sup>),  $C$ : concentration of the diffusing species (mol m<sup>-3</sup>).<sup>50</sup>

$$J = -D \frac{\partial C}{\partial x} \quad (1)$$

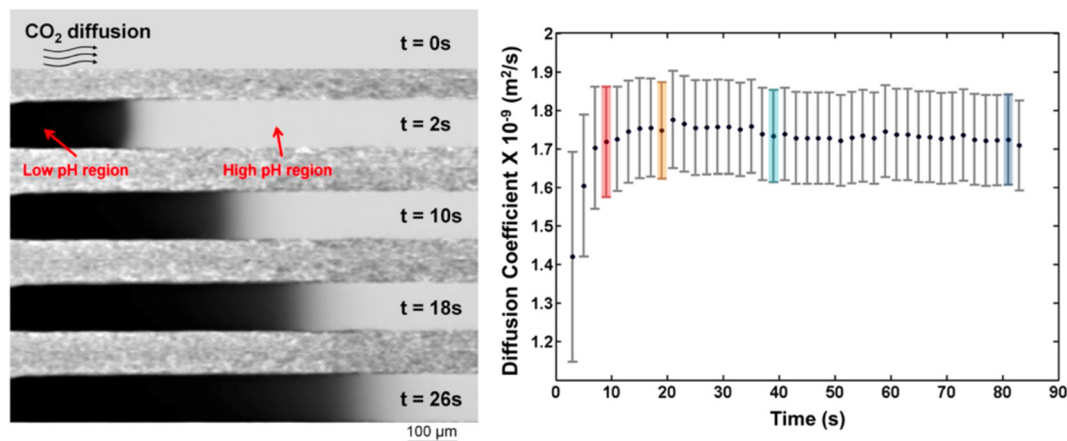
$$\frac{\partial C}{\partial t} = D \frac{\partial^2 C}{\partial x^2} \quad (2)$$

Several studies have applied Fick's laws of diffusion to quantify mass transfer in microfluidic systems for CO<sub>2</sub> capture and dissolution. Sell *et al.* used a microfluidic approach to measure the diffusion coefficient of CO<sub>2</sub> in water and brine for reservoir-specific carbon sequestration (Fig. 7).<sup>49</sup> They found that the diffusion coefficient in water was  $1.86 \times 10^{-9}$  m<sup>2</sup> s<sup>-1</sup> at 26 °C and constant across pressures from 5 to 50 bar, while in brine (0–5 M NaCl), it varied up to threefold, highlighting the effect of salinity. Sauzade and Cubaud modeled the dissolution dynamics of CO<sub>2</sub> bubbles in viscous silicone oils using Fick's first law.<sup>48</sup> At low capillary numbers ( $Ca \approx 10^{-2}$ ), bubbles dissolved steadily, while at high capillary numbers ( $Ca \approx 1.1$ ), dissolution became uneven due to bubble shape variations, offering insights for CO<sub>2</sub> injection in viscous environments (Fig. 8).<sup>48</sup>

**2.2.2. Convective–diffusive transport model.** The convective–diffusion model accounts for both diffusion and convective flow, which is particularly relevant in microchannels where laminar flow conditions are controlled to influence CO<sub>2</sub> transport.<sup>51</sup>

$$\frac{\partial C}{\partial t} + u \frac{\partial C}{\partial x} = D \frac{\partial^2 C}{\partial x^2} \quad (3)$$

Frost *et al.* chose the convective diffusion model to analyze a bilayer microfluidic device with a porous membrane



**Fig. 7** [Left] Visualization and analysis of CO<sub>2</sub> diffusion into water. Time-lapse images showing fluorescence quenching as CO<sub>2</sub> diffuses into water, leading to solution acidification. Diffusion coefficients were obtained at a pressure of 5 bar. A diffusion coefficient is determined for each frame captured during the experiment, with each data point spaced 2 s apart. The error bars account for uncertainties in measurements (e.g., initial pH, temperature) and their effect on the calculated diffusion coefficient ( $D$ ). [Right] The overall diffusion coefficient at this pressure is determined as  $D = 1.74 \times 10^{-9}$  m<sup>2</sup> s<sup>-1</sup>, obtained by averaging individual  $D$  values.<sup>49</sup>



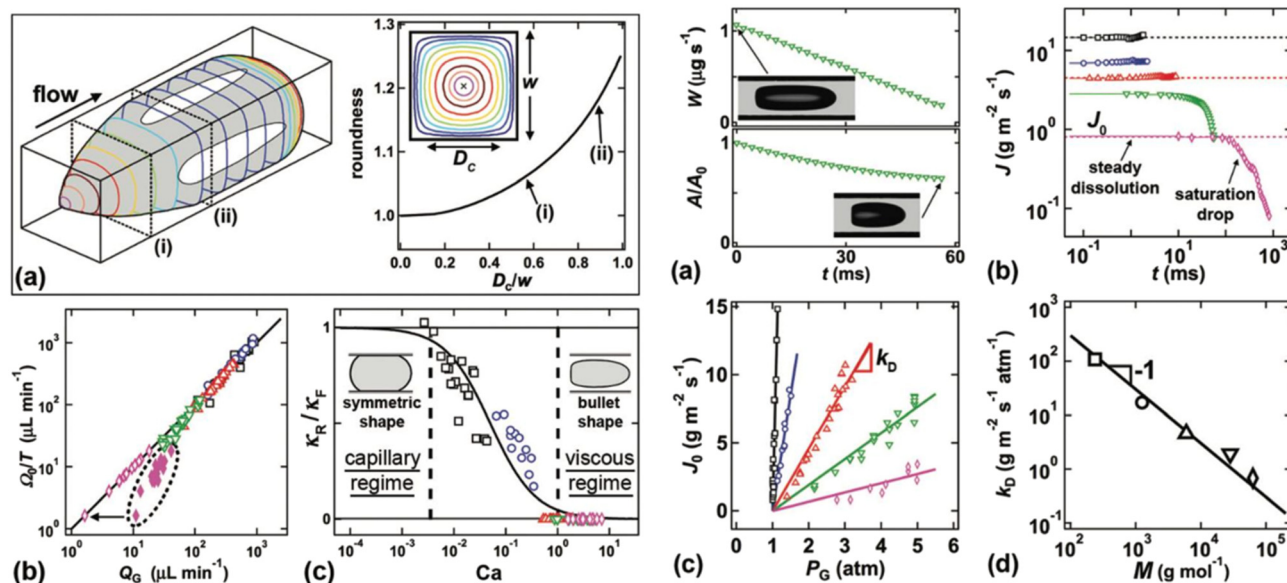


Fig. 8 [Left] (a) Schematic representation and analysis of CO<sub>2</sub> bubble dynamics in microfluidic systems with volume reconstruction where (i) and (ii) represent two random sections illustrating the roundness of the reconstructed CO<sub>2</sub> bubble, (b) comparison between the local gas flow rate, determined from *in situ* measurements, and the inlet gas flow rate, showing discrepancies at high viscosities where small flow rates are calibrated using the local measurement. (c) The ratio of rear cap curvature to front cap curvature as a function of the capillary number, highlighting the transition from capillary-dominated to viscous-dominated regimes, is modeled by  $K_R/K_F = (1 + Ca/Ca_c)^{-1}$  with  $Ca_c = 5 \times 10^{-2}$ . [Right] Representation of the diffusive mass flux of CO<sub>2</sub> bubbles in silicone oils under varying viscosity and pressure conditions. (a) Temporal evolution of the diffusive mass flow rate and interfacial area for a viscosity of 10<sup>3</sup> cS, showing an initial linear phase followed by a drop as saturation occurs. (b) Mass diffusion flux at different viscosities, demonstrating that higher viscosity fluids exhibit a more significant decline in flux due to increased diffusion resistance. (c) Early mass flux  $J_0$  as a function of gas pressure, following the relation  $J_0 = k_D(P_G - P_{atm})$ , indicating that higher pressures enhance CO<sub>2</sub> transfer rates. (d) Dissolution coefficient  $k_D$  as a function of oil molecular weight, showing an inverse relationship modeled by  $k_D = 3 \times 10^4 M^{-1}$ , where higher molecular weight oils impose greater resistance to CO<sub>2</sub> diffusion.<sup>48</sup>

because of its ability to capture molecular transport influenced by simultaneous flow (Fig. 9).<sup>52</sup> This model allowed for a detailed analysis of how the flow rate and membrane porosity affected concentration distributions. The study found that at higher flow rates, convection dominated, minimizing diffusion effects, while at lower flow rates, diffusion significantly influenced transport. This balance was critical for optimizing CO<sub>2</sub> permeation efficiency in diffusion-convective environments.<sup>52</sup>

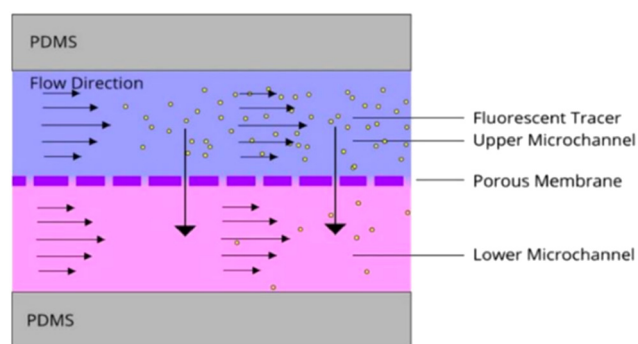


Fig. 9 Schematic representation of convection-diffusion mass transport in a microfluidic device. Molecules from the high-concentration stream (purple) diffuse into the low-concentration stream (pink) while convecting downstream, illustrating the simultaneous influence of advection and diffusion on mass transfer within the microchannel.<sup>52</sup>

**2.2.3. Two film theory.** Two-film theory proposes that mass transfer resistance occurs within two stagnant layers on either side of the gas-liquid interface, making it highly applicable to segmented or Taylor flow systems in microchannels (Fig. 10(a) and eqn (4)). Here,  $N_A$ : molar flux of component A (mol m<sup>-2</sup> s<sup>-1</sup>),  $k$ : mass transfer coefficient.<sup>53</sup>

$$N_A = k_G(C_{A,G} - C_{A,i}) = k_L(C_{A,i} - C_{A,L}) \quad (4)$$

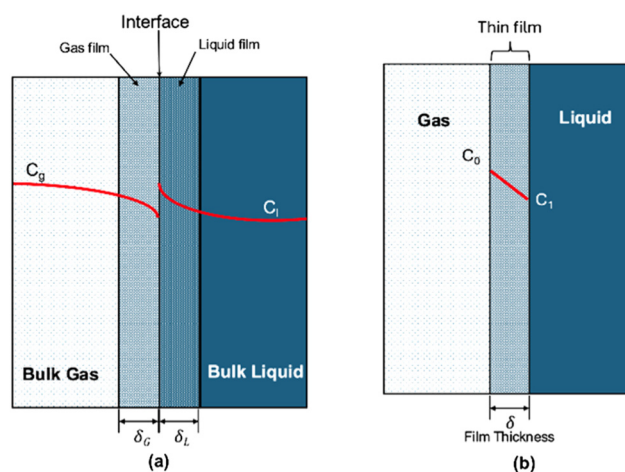


Fig. 10 Illustrative images of mass transfer models. (a) Two film mass transfer; (b) thin film mass transfer.



Pang *et al.* applied the two-film theory to quantify CO<sub>2</sub> absorption under Taylor flow in serpentine mini channels because it accurately estimated mass transfer coefficients ( $k_{La}$ ) in systems with alternating gas–liquid segments. Secondary flow patterns generated in the curved sections of the channels enhanced mass transfer, leading to volumetric mass transfer coefficients ranging from 0.1 to 0.4 s<sup>-1</sup>, significantly higher than those observed in straight channels. This theory was chosen to capture the complex dynamics of segmented flow and bubble-induced mixing.<sup>54</sup>

**2.2.4. Thin film theory.** Thin film theory is similar to two-film theory but assumes a single stagnant film near the gas–liquid interface in the liquid phase (Fig. 10(b)). It is particularly useful in confined microchannels where the diffusion layer thickness significantly impacts mass transfer. Eqn (5) is used to find the molar flux of the stagnate film. Here,  $\delta$  = thickness of the film (m).<sup>55</sup>

$$N_A = \frac{D}{\delta} (C_{A,i} - C_{A,L}) \quad (5)$$

Ibrahim *et al.* implemented thin film theory to study mass transfer in electrochemical energy conversion systems. The model helped precisely control mass transport phenomena in microchannels, improving efficiency in CO<sub>2</sub> reduction reactions by minimizing diffusion layer thickness. This theory was chosen because of the highly confined geometry of the channels, where diffusion-limited processes governed overall performance.<sup>56</sup>

**2.2.5. Surface renewal theory.** Surface renewal theory assumes that fresh liquid continuously replaces the interface, enhancing mass transfer. This model is particularly relevant in channels that generate vortices, such as serpentine or spiral microchannels. Eqn (6) shows the derived equation used for surface renewal theory, where  $t$  is characteristic renewal time (s).<sup>57</sup>

$$k_L = \sqrt{\frac{D}{\pi t}} \quad (6)$$

Li *et al.* applied surface renewal theory to study CO<sub>2</sub> absorption in rectangular microchannel reactors (Fig. 11).<sup>58</sup> The model was chosen due to its ability to account for the rapid replacement of fluid elements at the interface, significantly enhancing mass transfer rates. The experiments showed that  $k_{La}$  values reached up to 21 s<sup>-1</sup> at optimal gas flow rates, far exceeding the performance of conventional reactors. This high mass transfer efficiency made surface renewal theory the ideal model for quantifying CO<sub>2</sub> absorption in dynamic microfluidic systems.<sup>58</sup>

**2.2.6. Penetration theory.** Penetration theory describes mass transfer where liquid elements at the interface interact briefly with the CO<sub>2</sub> gas phase before fresh liquid replaces them, making it particularly relevant for Taylor flow systems with alternating gas and liquid segments. Fig. 12 explains the penetration theory, while eqn (7) is derived from Fick's second law with the assumption of an infinite medium with a time-dependent boundary condition.<sup>59,60</sup>

$$k_L = \frac{2D}{\pi t} \quad (7)$$

The study by Ho *et al.* used penetration theory to analyze CO<sub>2</sub> mass transfer in a microfluidic system under high-pressure conditions relevant to carbon sequestration in deep saline aquifers.<sup>61</sup> A microfluidic T-junction was employed to generate CO<sub>2</sub> microbubbles, and their dissolution dynamics were tracked to determine the liquid-side mass transfer coefficient. The characteristic exposure time of the liquid elements was estimated from the pressure of CO<sub>2</sub> and the flow rate. In their experiment, characteristic time was calculated to be approximately 30 milliseconds at a pressure of 5 MPa and a flow rate of 0.2 mL min<sup>-1</sup>. The study demonstrated that penetration theory is particularly useful for short-lived mass transfer events, making it ideal for systems where rapid gas–

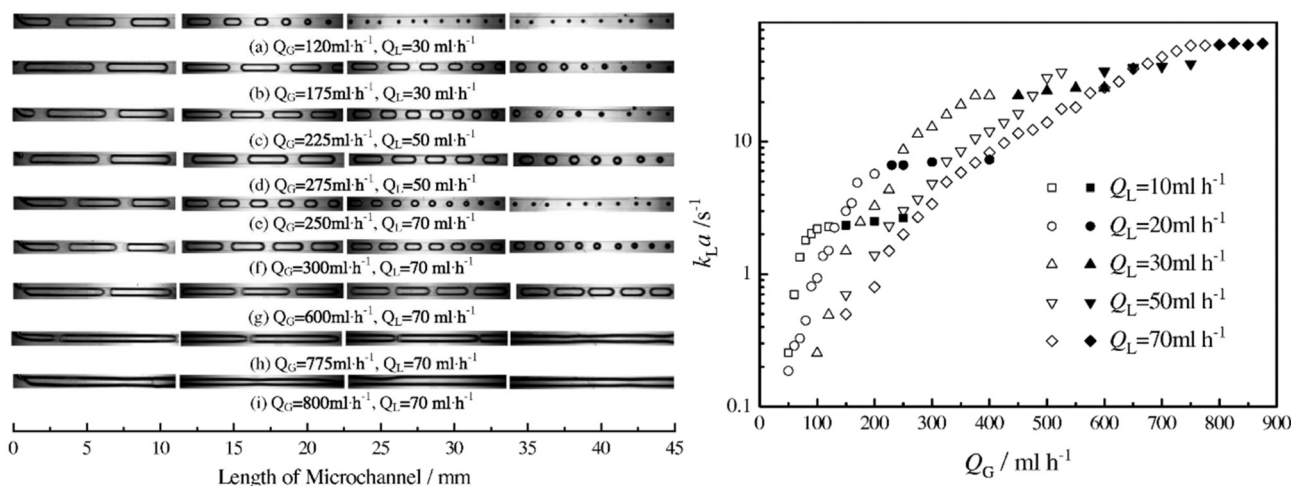


Fig. 11 [Left] Experimental image of gas–liquid flow at different flow rates (from a to i). [Right] The calculated volumetric mass transfer coefficient ( $k_{La}$ ) for different flow rates.<sup>58</sup>





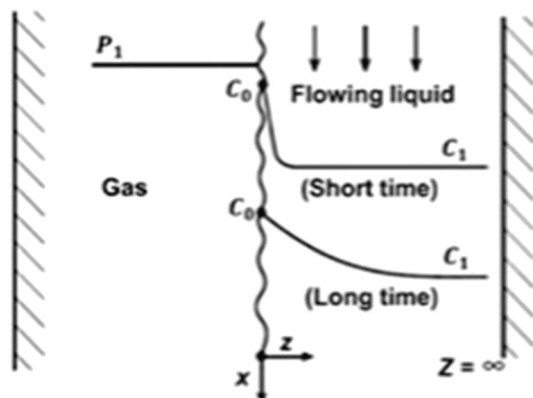


Fig. 12 Illustrative image showing gas-liquid interaction representing penetration mass transfer theory.<sup>62</sup>

liquid interactions occur, such as in enhanced oil recovery and carbon capture applications. This method provides a reliable framework for optimizing operating conditions in high-pressure microfluidic environments.<sup>61</sup>

### 2.3. Microchannel geometries and flow patterns

Similar to the transport model, selecting the appropriate microchannel geometry and flow pattern is essential for ensuring flow stability and reproducibility in microfluidic CO<sub>2</sub> capture studies. The microchannel geometry directly impacts the mass transfer efficiency, gas-liquid interfacial area, and overall experimental precision. Improper design selection can lead to significant errors and poor reproducibility. This subsection aims to provide the reviewers with overall knowledge of different geometries and flow

patterns, highlighting their impact on experimental outcomes and helping researchers select optimal designs for improved performance.

Common cross-sectional shapes of microchannels include rectangular, circular, square, triangular, trapezoidal, and elliptical (Fig. 13). Each geometry offers distinct advantages, influencing the liquid film thickness, bubble formation, and overall gas-liquid interaction. Among these, rectangular channels are the most widely used due to their high surface-to-volume ratio and the ability to generate thin, stable liquid films that improve mass transfer. Al-Rawashdeh *et al.* showed that rectangular channels produced a spatially uniform liquid film caused by the corner capillary effect, which stabilized the flow pattern and enhanced Taylor bubble formation, ensuring uniform CO<sub>2</sub> absorption.<sup>63</sup> Quantitative studies further highlight the role of microchannel dimensions in optimizing performance. Reducing the hydraulic diameter ( $d_h$ ) from 762  $\mu\text{m}$  to 254  $\mu\text{m}$  resulted in a 2.6-times increase in the volumetric mass transfer coefficient ( $k_{La}$ ), confirming that smaller channels significantly boost mass transfer efficiency.<sup>64</sup>

In contrast, circular channels are preferred for Taylor flow systems, offering symmetrical flow profiles and consistent bubble generation. Their design reduces pressure drops and ensures steady flow, making them ideal for gas-liquid contact applications. However, circular channels often require specialized molds or capillary assembly for fabrication, which can be more challenging than rectangular channels. Square cross-sections balance rectangular and circular geometries, ensuring moderate mass transfer efficiency and flow stability.<sup>65</sup> Triangular and trapezoidal channels are less common but serve specific purposes where higher velocity gradients and enhanced mixing are required, particularly in low Reynolds number flows.<sup>66</sup> Trapezoidal

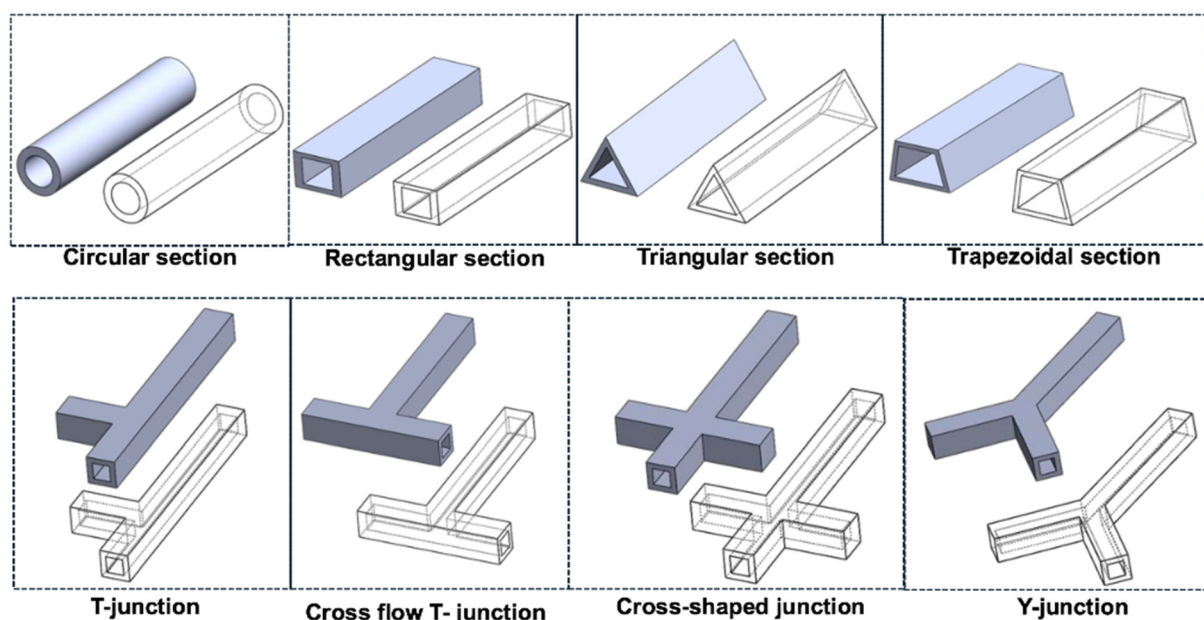


Fig. 13 Illustration of different cross sections of microchannels, different junctions in microchannel geometries, and their wireframe views.

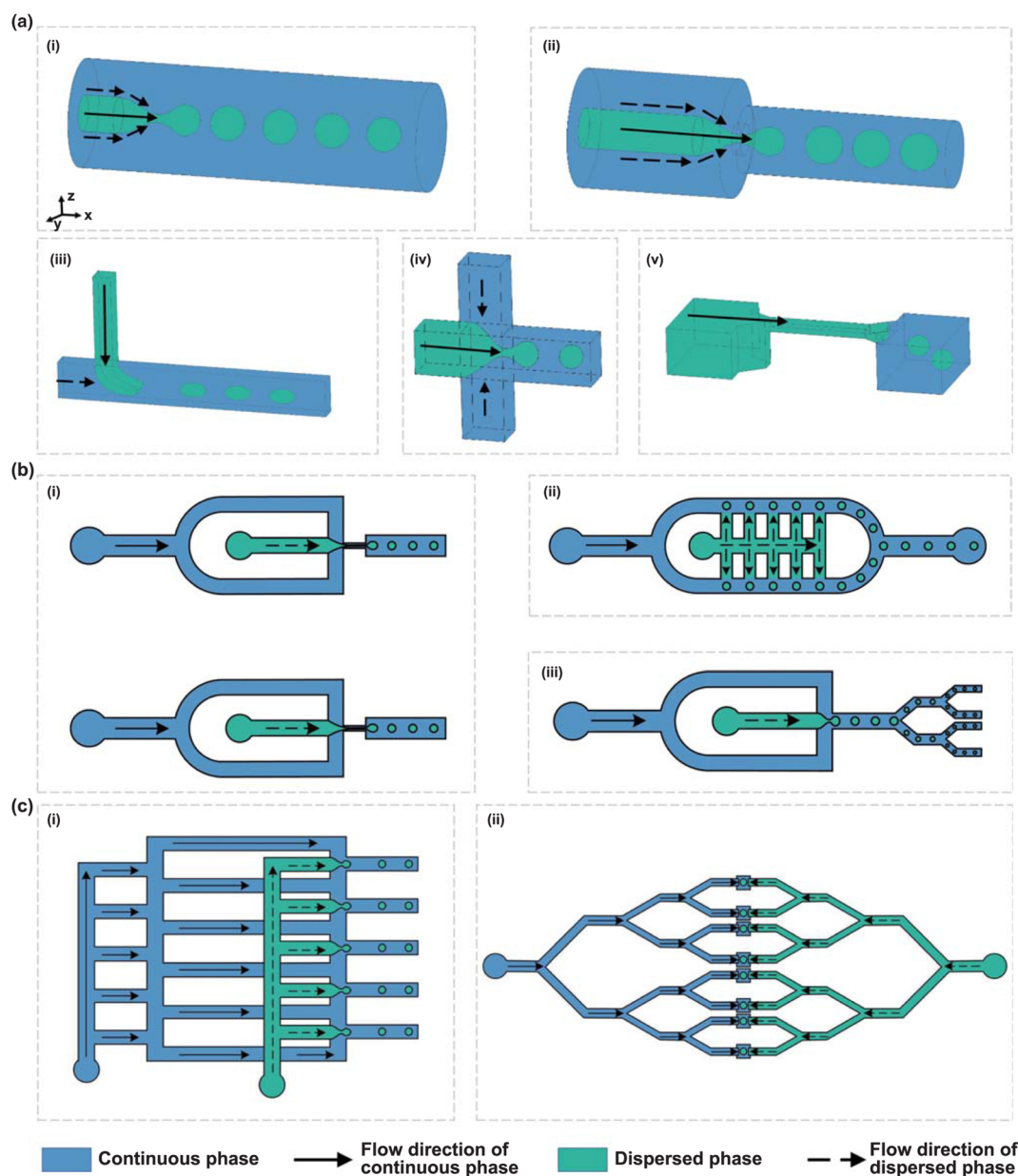




cross-sections provide smoother flow transitions, reducing flow disturbances and improving selectivity in chemical absorption processes. Meanwhile, elliptical channels offer reduced shear stress and smooth flow profiles, making them suitable for gentle flow applications, such as enzymatic CO<sub>2</sub> conversion or biological reactions. The hydraulic diameter ( $d_h$ ) of microchannels used for CO<sub>2</sub> absorption generally ranges between 200  $\mu\text{m}$  and 1000  $\mu\text{m}$ , while the channel length ( $L_c$ ) in laboratory experiments is usually limited to 400 mm for easier visualization and flow control.<sup>17</sup> Channel size

directly affects energy dissipation and mass transfer, with smaller channels promoting slug flow patterns in broader gas velocity ranges and significantly enhancing mass transfer performance.<sup>67–69</sup>

In addition to the channel cross-section, the junction types and their configurations significantly affect gas–liquid interaction and bubble formation (Fig. 13). Common microfluidic CO<sub>2</sub> capture system junctions include T-junctions, Y-junctions, and cross-junctions. Among these, T-junctions are the most frequently used due to their



**Fig. 14** Schematic illustration of microchannel characteristics. (a) Flow behaviors in different droplet generators: (i) co-flow, (ii) flow-focusing, (iii) T-junction, (iv) cross-junction, and (v) step emulsification, demonstrating various methods for generating emulsions in microfluidic systems. (b) The scale-up strategies for microfluidic processes: (i) parallel numbering-up of both dispersed and continuous phase nozzles, (ii) parallel numbering-up of dispersed phase nozzles only, and (iii) splitting of emulsion droplets to achieve higher throughput. (c) The fluid distribution network in two common layouts: (i) ladder network and (ii) tree network, illustrating different methods for ensuring uniform flow distribution across multiple microchannels.<sup>18</sup>



reliable blocking-squeezing mechanism, which generates stable Taylor bubbles at low liquid flow rates.<sup>70,71</sup> This mechanism ensures consistent bubble formation and improved mass transfer performance. Tan *et al.* demonstrated that T-junctions achieve a superior mass transfer rate due to comparatively high shear forces. It was found that the slug formation stage was responsible for 30–40% of the total mass transfer.<sup>72</sup> Y-junctions are preferred for their smoother bubble generation, particularly at higher flow rates, while cross-junctions facilitate intensive mixing between gas and liquid phases but are associated with higher pressure drops.<sup>73,74</sup>

Flow patterns in microchannels significantly affect the mass transfer dynamics and efficiency of CO<sub>2</sub> absorption. These patterns depend on the gas and liquid flow rates, channel geometry, and physical properties of the fluids (Fig. 14). The most common flow patterns observed in gas-liquid systems are bubble flow, slug (Taylor) flow, annular flow, stratified flow, and churn flow, each with distinct characteristics. In bubble flow, small, dispersed gas bubbles are randomly distributed within a continuous liquid phase.<sup>75</sup> Typically, the bubble shape is spherical in bubble flow, whereas slug flow, also known as Taylor flow, consists of elongated bubbles (plugs or slugs) with a comparable diameter to the channel or tube in the liquid phase.<sup>76</sup> This flow type is the most desirable for CO<sub>2</sub> absorption experiments due to its high mass transfer efficiency. While bubble flow has a greater surface area-to-volume ratio, which may intuitively suggest a higher mass transfer, the stagnate layer surrounded by the bubble only limits the diffusion mass transfer, whereas, in Taylor flow, the thin liquid layer between gas and surface walls provides faster diffusion rate by preventing any stagnate layer buildup. Studies have shown that slug flow provides up to 50% higher mass transfer rates than bubble flow, particularly in rectangular channels where uniform bubble size and stability are maintained.<sup>77,78</sup> Bubble flow is observed at low gas flow rates and is characterized by dispersed bubbles in the continuous liquid phase. It is commonly used in systems that require gentle flow conditions to prevent shear-sensitive reactions. Annular flow occurs at high gas flow rates, where the gas forms a core surrounded by a thin liquid film along the channel walls. This pattern offers good gas-liquid contact but comes with increased pressure drop and reduced stability. Stratified flow, where gas and liquid form separate layers, is more common in horizontal large-diameter channels than in microchannels.<sup>79</sup> Churn flow represents an unstable transition between slug and annular flow, with highly irregular gas-liquid interactions, making it less suitable for controlled CO<sub>2</sub> capture processes. The choice of flow patterns directly impacts the volumetric mass transfer coefficient. These findings demonstrate the importance of optimizing channel size and flow pattern to maximize CO<sub>2</sub> absorption efficiency. Additionally, spiral and serpentine channels have been known to intensify CO<sub>2</sub> absorption by promoting radial mixing and extending the gas-liquid interface.<sup>80,81</sup> These

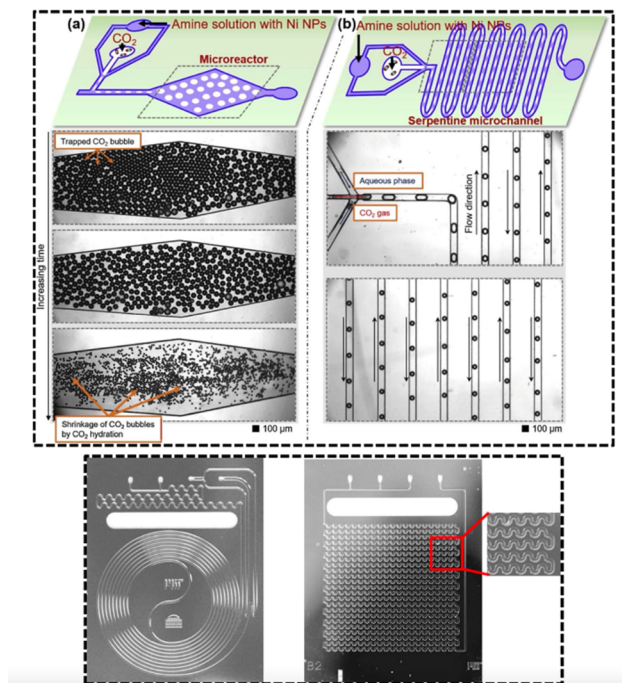


Fig. 15 [Top] Schematic of the microfluidic approaches for evaluating CO<sub>2</sub> absorption. The top panel illustrates two designs. (a) Microreactor operating under limited mixing conditions, along with sequential images showing changes in CO<sub>2</sub> microbubble size. (b) A long serpentine microchannel under high mixing conditions, with images of CO<sub>2</sub> microbubbles at two different locations. The scale bar represents 100 μm.<sup>81</sup> [Bottom] Actual photograph of the spiral channel and meandering channel geometries.<sup>80</sup>

designs reduce residence time while maintaining high mass transfer coefficients, offering continuous and efficient CO<sub>2</sub> absorption in high-throughput systems (Fig. 15).

For large-scale CO<sub>2</sub> capture, multi-channel integration and high-throughput systems are essential to achieve scalable performance. Conventional single-channel systems have limitations in throughput, but recent advancements in parallel microchannel reactors and tree-like manifolds have enabled high-throughput absorption while maintaining uniform flow.<sup>82</sup> Li *et al.* developed a 16-parallel-channel reactor, which achieved volumetric mass transfer coefficients ( $k_L$ ) ranging from 1.59 to 90 s<sup>-1</sup>, ensuring stable slug flow across all channels (Fig. 16).<sup>83</sup> Tree-like structures and fractal designs are commonly employed to reduce flow resistance and ensure uniform distribution across multiple channels, further enhancing performance in multi-channel systems.

#### 2.4. Measurement techniques of CO<sub>2</sub> capture in microfluidics

Along with the mass transfer models, microfluidic channel geometry and flow pattern selection play a crucial role in microfluidic CO<sub>2</sub> capture studies, and the ability to accurately measure CO<sub>2</sub> dissolution and transport dynamics is equally essential. Accurate and reliable measurement of CO<sub>2</sub> capture efficiency in microfluidic systems is critical for optimizing system performance, validating mass transfer models, and



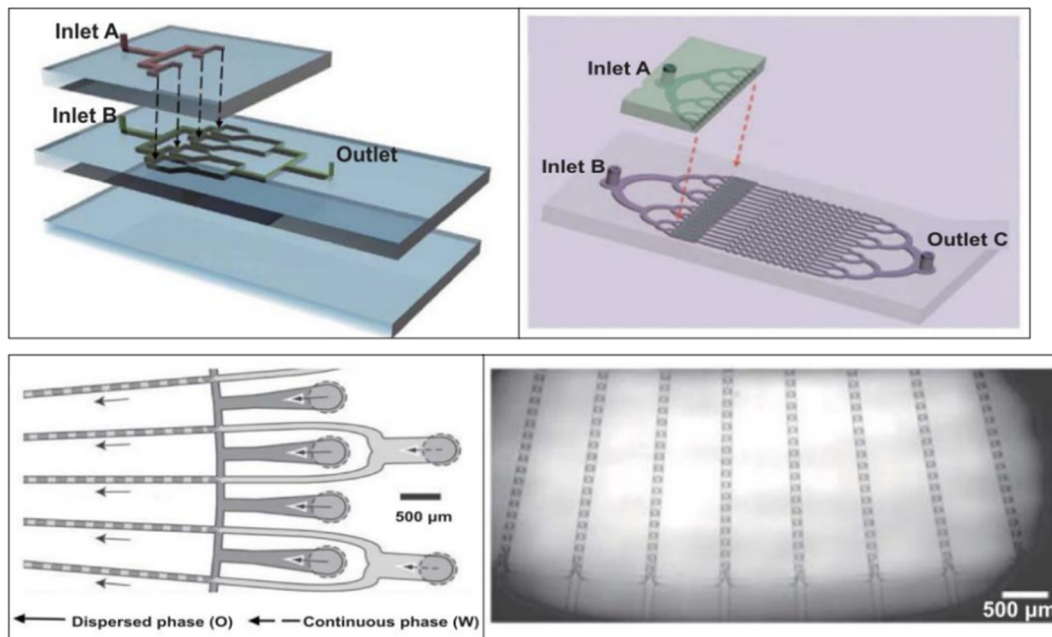


Fig. 16 [Top left] Schematic of parallelization of a flow-focusing device. [Top right] Droplets formed in four integrated flow-focusing devices, demonstrating the simultaneous generation of emulsions in multiple channels.<sup>82</sup> A channel consists of 16 flow-focusing junctions.<sup>83</sup> [Bottom] A schematic of a microfluidic geometry where 128 cross junctions are integrated together and its magnified view.<sup>84</sup>

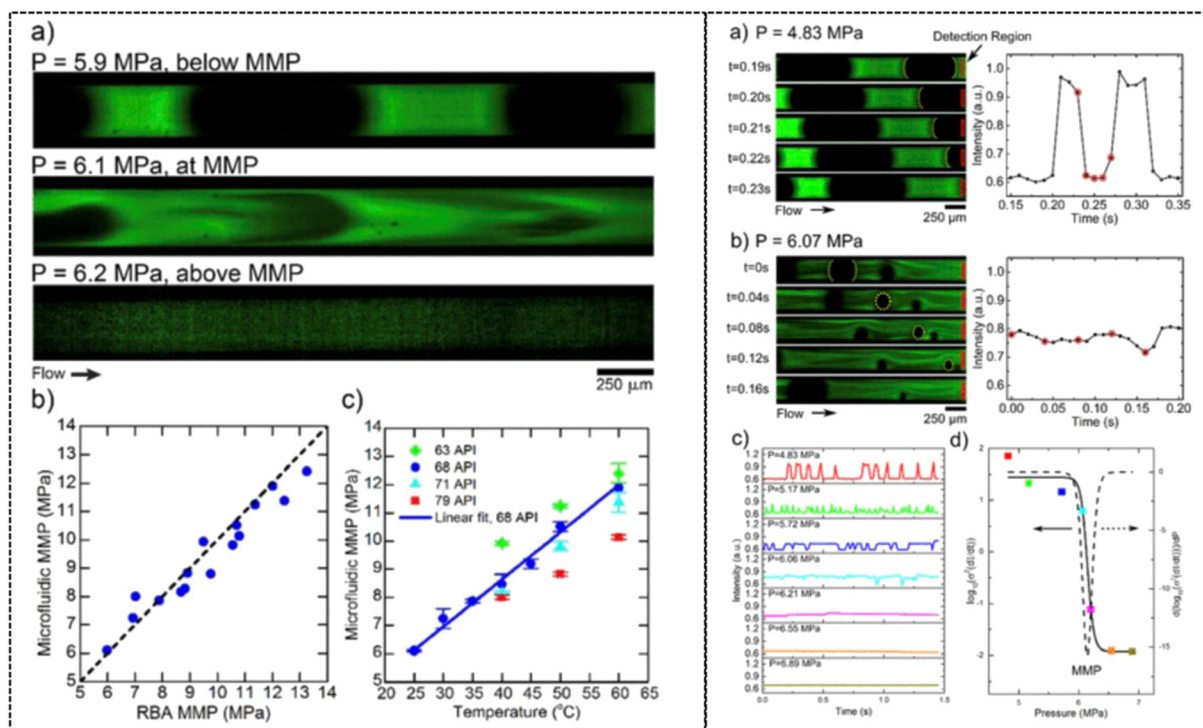


Fig. 17 Fluorescence-based microfluidic measurement of MMP for CO<sub>2</sub> in crude oils. [Left] (a) Fluorescence images of CO<sub>2</sub> bubbles at different pressures: below MMP (5.9 MPa), at MMP (6.1 MPa), and above MMP (6.2 MPa), showing the transition from distinct CO<sub>2</sub> bubbles to complete mixing. (b) Correlation between microfluidic MMP measurements and rising bubble apparatus (RBA) MMP values showing strong agreement. (c) Temperature dependence of MMP for synthetic oil mixtures of varying API gravities, demonstrating a linear increase in MMP with temperature. [Right] Operator-independent fluorescence measurements showing intensity fluctuations at (a) 4.83 MPa and (b) 6.07 MPa. Red-circled data points correspond to selected images. (c) Average intensity within the detection region over time at various pressures. (d) Logarithmic variance of the intensity derivative reveals the steepest change.<sup>85</sup>





scaling for real-world applications. Advanced measurement techniques combined with microfluidic platforms enable real-time monitoring of key parameters such as localized gas concentration, pH profiles across the interface, and ionic composition, which are often impractical or highly challenging in conventional systems. This section outlines key methods for quantifying CO<sub>2</sub> capture in microfluidics, including optical methods, electrochemical sensors, gas chromatography, and Raman spectroscopy, along with details on accuracy, time scales, and applications.

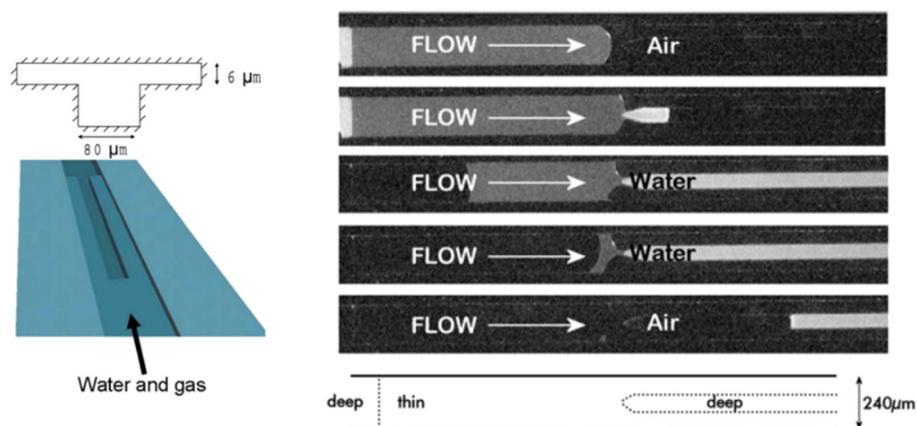
Optical methods are widely used in microfluidic systems due to their non-invasive nature and ability to provide real-time measurements of CO<sub>2</sub> concentration and dissolution behavior. Among the optical techniques, brightfield microscopy and fluorescence microscopy are the most used for visualizing gas-liquid interfaces and tracking CO<sub>2</sub> dissolution in microchannels. Brightfield microscopy offers a simple and direct imaging approach, ideal for observing bubble dynamics and flow patterns. In contrast, fluorescence microscopy is used for more quantitative analysis, such as changes in pH, temperature, and phase concentration. A recent study by Mohaddes *et al.* developed a fluorescence-based microfluidic method for measuring the minimum miscibility pressure (MMP) of CO<sub>2</sub> in crude oil (Fig. 17).<sup>85</sup> This method leverages the inherent fluorescence of crude oils and achieves precise, operator-independent measurements. Compared to conventional techniques, such as the rising bubble apparatus, which can take days, this microfluidic method provides quantitative MMP values in under 30 minutes with an accuracy difference of less than 0.5 MPa. Such advancements highlight the potential of fluorescence-based approaches for rapid, real-time CO<sub>2</sub> measurement in microfluidic CO<sub>2</sub> capture systems, offering a significant improvement in both speed and accuracy over traditional methods.<sup>85</sup>

Infrared absorption spectroscopy is a popular approach for measuring CO<sub>2</sub> concentration by detecting the absorption of infrared light at specific wavelengths. For instance, Tavernier *et al.* developed an inline microfluidic system for measuring

CO<sub>2</sub> concentration using infrared absorption at 4.24  $\mu\text{m}$  (Fig. 18).<sup>86</sup> The system operated under pressures ranging from 0 to 70 bars, offering precise real-time measurements, even in challenging environments such as oil fields.<sup>86</sup>

Gas chromatography (GC) has been utilized in analyzing the composition of CO<sub>2</sub> and its reaction by-products in microfluidic systems. While less suitable for real-time monitoring, it provides high-resolution and quantitative analysis of gas-phase products. Rudyk *et al.* used GC to characterize CO<sub>2</sub> reduction products during EOR experiments, demonstrating its importance in identifying and quantifying multiple gas species.<sup>87</sup>

Raman spectroscopy is increasingly used for real-time monitoring of CO<sub>2</sub> absorption and isotopic analysis. One significant application of Raman spectroscopy is in measuring  $\delta^{13}\text{C}$  values—a ratio representing the relative abundance of the heavier isotope <sup>13</sup>C compared to <sup>12</sup>C in CO<sub>2</sub>. This value is commonly used in geochemical studies to trace the source of CO<sub>2</sub> or analyze carbon cycling processes. Li *et al.* used micro-laser Raman spectroscopy to quantitatively determine  $\delta^{13}\text{C}$  values in natural CO<sub>2</sub> gas samples. The method showed relative errors ranging from 0.076% to 1.154%, with results that matched those from Isotope Ratio Mass Spectrometry (IRMS).<sup>88</sup> Unlike IRMS, however, Raman spectroscopy offered faster results within minutes and required less sample preparation. In another study, Wells *et al.* used Raman spectroscopy to monitor CO<sub>2</sub> hydrate formation in high-pressure microfluidic devices. Their findings revealed that hydrate conversion reached 47% within 1 hour, compared to days or weeks required for similar results in bulk systems.<sup>89</sup> These analytical methods collectively contribute to advancing microfluidic CO<sub>2</sub> capture research. Optical techniques provide rapid and non-invasive measurement capabilities; electrochemical sensors enable direct chemical monitoring, and gas chromatography is a reliable validation tool. Combining these approaches ensures a comprehensive evaluation of CO<sub>2</sub> absorption processes, guiding the development of efficient carbon capture technologies.



**Fig. 18** [Left] Microfluidic gas-liquid phase separation for CO<sub>2</sub> measurement. Schematic of the microchannel design, featuring deep and thin regions for controlled phase separation. [Right] Experimental images showing sequential air and water flow, demonstrating effective gas isolation for inline CO<sub>2</sub> analysis with infrared absorption spectroscopy.<sup>86</sup>





### 3. Role of microfluidics in carbon storage

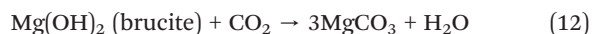
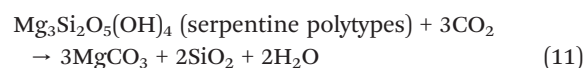
#### 3.1. CO<sub>2</sub> storage approaches

Successful carbon capture must be followed and combined with suitable carbon storage and/or sequestration technologies to prevent the captured CO<sub>2</sub> from returning to the atmosphere. According to the definition by Scott *et al.*, carbon sequestration generally refers to the permanent locking of carbon with a lifetime of greater than 100 000 years, whereas carbon storage often refers to the temporary containment of CO<sub>2</sub> with a lifetime of 1000 years or less, with the later maintaining the possibility of re-accessing the stored CO<sub>2</sub> for future utilization purposes.<sup>90,91</sup> Nevertheless, the two terms, storage and sequestration, are frequently used interchangeably in the literature, and no distinction will be made between them in this review, as both timescales are hardly meaningful in lab-scale experiments using microfluidics.

The National Academies of Sciences, Engineering, and Medicine, in their report published in 2019, laid out six major technical approaches to CO<sub>2</sub> removal and storage, including (i) coastal blue carbon, (ii) terrestrial carbon removal and sequestration, (iii) bioenergy with carbon capture and sequestration (BECCS), (iv) DAC, (v) carbon mineralization, and (vi) geologic CO<sub>2</sub> sequestration.<sup>92</sup> Coastal blue carbon aims to take advantage of improved land use and management practices that increase the carbon stored in living plants or sediments in mangroves, tidal marshlands, seagrass beds, and other tidal or salt-water wetlands, whereas terrestrial carbon removal and sequestration aims to enhance soil carbon storage through afforestation/reforestation, changes in forest management, or changes in agricultural practices. BECCS typically refers to the integration of trees and crops that extract CO<sub>2</sub> from the atmosphere as they grow, the use of this biomass in power plants, and the application of carbon capture and sequestration *via* CO<sub>2</sub> injection into geological formations. Similarly, DAC aims to capture CO<sub>2</sub> from ambient air and concentrate it so that it can be injected into a storage reservoir. Finally, carbon mineralization aims to take the CO<sub>2</sub> captured through BECCS or DAC and bond it with reactive minerals such as peridotite and basaltic lava for permanent sequestration, whereas geological sequestration looks to inject captured CO<sub>2</sub> into geological formations such as saline aquifers. In addition to these six pathways, CO<sub>2</sub> conversion technologies, such as electrochemical, photocatalytic, thermocatalytic, and biological processes, also contribute to carbon sequestration.<sup>93–95</sup> These methods serve as post-capture pathways that transform CO<sub>2</sub> into valuable products like fuels, chemicals, and solid materials. Although these pathways have been identified for carbon storage, most of them are merely combinations of carbon storage with various carbon capture technologies. For “pure” carbon storage, there are essentially only two approaches, namely carbon mineralization and geological sequestration, which, along with their overlap with microfluidics, will be our

particular focus in this review.<sup>91</sup> The readers are referred to the work by Kazemifar and the National Academies of Science, Engineering, and Medicine, and the review articles on CO<sub>2</sub> conversion for more generic reviews of technologies for carbon sequestration, utilization, and their technological readiness.<sup>91,92,96,97</sup>

**3.1.1. Carbon mineralization.** Carbon mineralization was proposed as a CO<sub>2</sub> reduction strategy in the 1990s.<sup>98,99</sup> It aims to bring captured CO<sub>2</sub> of different forms (gas, liquid, water-bearing, or supercritical) to react with mine tailings as well as surface and subsurface rocks that are rich in calcium (Ca) and magnesium (Mg) to form solid carbonate minerals. Another source of Mg and Ca are industrial byproducts (cement kiln dust, steel slag, and fly ash).<sup>100</sup> CO<sub>2</sub> reacts with minerals rich in Ca and Mg to form carbonates, such as calcite (CaCO<sub>3</sub>), magnesite (MgCO<sub>3</sub>), dolomite (CaMg(CO<sub>3</sub>)<sub>2</sub>), and often quartz (SiO<sub>2</sub>), all of which are minimally soluble in aqueous solution. Some idealized reactions are as follows:



All these reactions are spontaneous and exothermic as carbonate minerals are the “ground state” for near-surface rock systems like Mg–Ca–C–O–H and Mg–Ca–Si–C–O–H. Since the final reaction products are in stable solid forms, they are considered one of the safest and most secure storage mechanisms for minimizing leakage.<sup>101</sup> In fact, carbon mineralization occurs naturally during the weathering of silicate materials such as olivine and peridotite, which compose Earth's upper mantle and basaltic lava formed by partially melting the upper mantle. In that sense, this approach as a CO<sub>2</sub> storage technology essentially represents nothing more than an accelerated “weathering” process, where concentrated CO<sub>2</sub> is allowed to react with ultramafic and/or basaltic rocks at a rate that is fast enough to contribute to climate change mitigation, adding another layer of safety.<sup>102</sup> Additionally, these natural and spontaneous processes make use of the abundant chemical potential energy that is vastly available at and near the earth's surface, offering the potential to accommodate a large amount of CO<sub>2</sub> at competitively low cost.

Depending on the location where carbon mineralization is performed, two approaches have been proposed and exercised: *ex situ* carbon mineralization and *in situ* carbon mineralization. In the former, CO<sub>2</sub>-bearing air and surface waters are brought to react with crushed and/or ground mine tailings, alkaline industrial wastes, or sedimentary



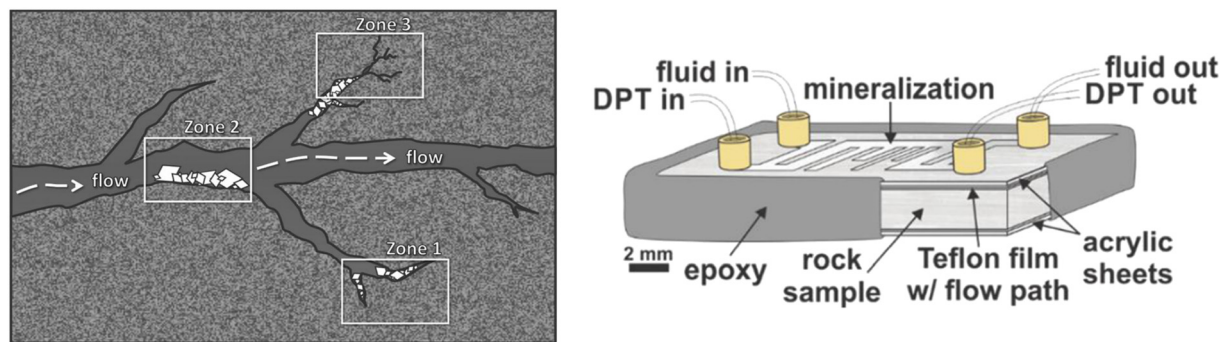
formations above the earth's surface, whereas in the latter, CO<sub>2</sub>-bearing fluids are injected and circulated through subsurface porosity in geological formations, and reactions occur *in situ* in the subsurface environment. While both methods potentially offer giant storage capacities at similar costs, *in situ* carbon mineralization has attracted more scientific interest, as it involves uncertain feedback between permeability, reactive surface area, and reaction rate, providing a fascinating topic for fundamental research.<sup>103</sup>

So far, most experiments investigating carbon mineralization processes are based on core samples.<sup>104–110</sup> Microfluidic studies of mineralization are challenging because, in carbon mineralization, CO<sub>2</sub> directly reacts with solids, meaning that the microfluidic devices would have to be fabricated or functionalized with reactive minerals, which causes significant challenges in fabrication and pore-scale quantification. Nevertheless, one recent study by Neil *et al.* attempted to identify key processes for carbon mineralization in fractured mafic and ultramafic rocks using microfluidic experiments. They employed a reactive microfluidic system to observe coupled reactions and flow in a simple fracture network containing both dead ends and a primary flow pathway for natural rock samples (Fig. 19).<sup>111</sup> The microfluidic device allowed for the direct observation of transport and chemistry for realistic mineralization systems, comparison of reaction *vs.* transport, and measurement of how reaction affects flow and can potentially lead to clogging. As shown in Fig. 19, rectangular chips of crystalline gypsum were cut, and a comb-like flow path was defined by a laser cut. A Teflon sheet is sandwiched between acrylic sheets and held together by epoxy (Loctite metal/concrete). Their results depict a visible difference in mineralization within the main channel at different flow rates, with calcites more dominant at higher flow rates. Channel clogging was not observed, presumably because the reaction of gypsum to calcite and vaterite is a volume-reduction reaction. A noticeable difference was observed in the amount and type of precipitation found in the dead-end channels of the flow path, where the faster flow rate resulted in more mineralization within the dead ends and a greater abundance of calcite. The experiments, however, cannot reveal whether

the calcite forms as a result of vaterite transformation or direct precipitation, calling for further studies in that regard.

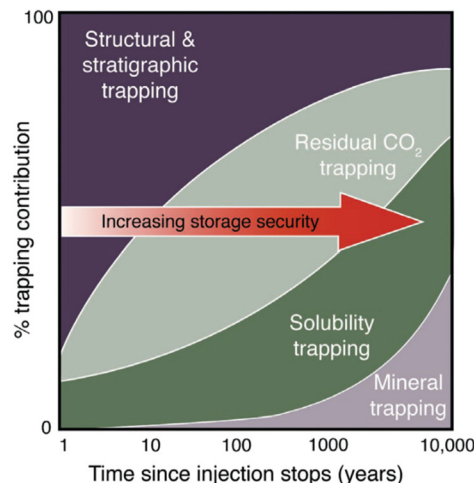
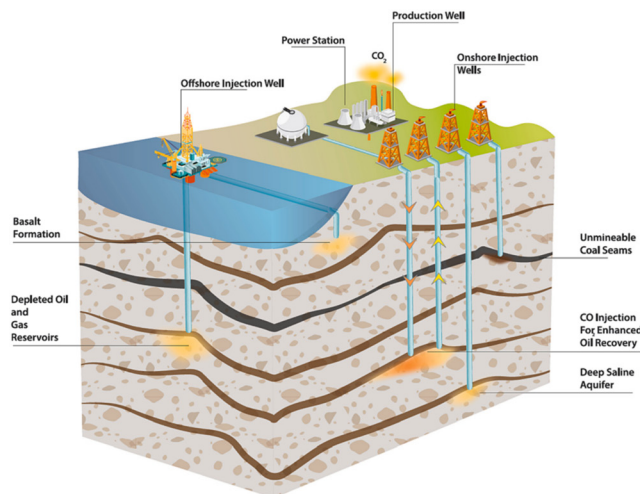
**3.1.2. Geological CO<sub>2</sub> sequestration.** Geological CO<sub>2</sub> sequestration in deep sedimentary formations typically involves the capture of CO<sub>2</sub> from large point sources (*e.g.*, thermal power plants), compression into a liquid or supercritical form, transportation to storage sites, and injection into a closed geological formation at depth (*i.e.*, a saline aquifer, a depleted oil or gas field, or an inaccessible coal bed) as illustrated in Fig. 20 [left].<sup>112</sup> As the National Academies of Sciences, Engineering, and Medicine pointed out, geological CO<sub>2</sub> sequestration benefits from nearly a half-century of experience with CO<sub>2</sub> injection for EOR and nearly two decades of commercial experience with saline aquifer sequestration.<sup>113,114</sup> Moreover, various aspects of CO<sub>2</sub> sequestration, including methods for monitoring the fate and transport of CO<sub>2</sub> on the surface and leakage, have been tested through many pilot-scale experiments in various formations.<sup>115–121</sup> For a general knowledge of geological CO<sub>2</sub> sequestration, the readers are referred to those aforementioned reviews. It is worth noting that microfluidics has played a crucial role in advancing our understanding of the foundational physics and trapping mechanisms, as discussed in section 3.3.

The safety and security of geological CO<sub>2</sub> sequestration are enabled by effective trapping mechanisms, including structural trapping, residual trapping, solubility trapping, and mineral trapping. Under typical reservoir conditions, compressed and supercritical CO<sub>2</sub> is less dense than the resident fluids and thus tends to rise through the formations driven by buoyancy forces. Therefore, as part of the requirement for site selection, a low-permeability rock, typically called a reservoir seal or a caprock, is needed to structurally contain the rising CO<sub>2</sub>. This mechanism is referred to as structural trapping, which plays a crucial role in the initial stage of CO<sub>2</sub> injection, as shown in Fig. 20.<sup>122</sup> However, structuring trapping doesn't eliminate the risk of CO<sub>2</sub> leakage, as any pre-existing faults or cracks in the caprock can cause catastrophic leakage, rendering all the efforts in vain. Therefore, secondary trapping mechanisms such as residual trapping, solubility trapping, and mineral trapping have been explored. Residual trapping, also known



**Fig. 19** [Left] The schematic of carbon mineralization occurring in different zones of a fractured mafic/ultramafic rock. [Right] The schematic showing a microfluidic model system used to investigate carbon mineralization in a fracture.<sup>111</sup>





**Fig. 20** [Left] Options for storing CO<sub>2</sub> in deep underground geological formations.<sup>112</sup> [Right] A general representation of the evolution of trapping mechanisms over time.<sup>114</sup>

as capillary trapping, relies on capillary pressure and surface tension between the injected CO<sub>2</sub> and the resident brine to immobilize the CO<sub>2</sub>.<sup>123</sup> Solubility trapping is due to the dissolution of CO<sub>2</sub> into the pore brine, which results in a denser CO<sub>2</sub>-bearing brine fluid that tends to sink rather than rise. Mineral trapping occurs when dissolved CO<sub>2</sub> reacts with minerals in the geologic formation, promoting precipitation of carbonate minerals.<sup>124</sup> Fig. 20 shows the contribution of each trapping mechanism depending on the formation characteristics and injection stage.

Given the long-term implications of CO<sub>2</sub> injections, developing a holistic understanding and predictive models capable of providing accurate and reliable predictions of large-scale CO<sub>2</sub> migration through the reservoir is considered a prerequisite for site selections and post-injection operations. One big challenge in this regard is the extensive range of spatial (from km to sub-micrometer) and temporal scales (from milliseconds to years) involved in these geophysical and geochemical processes.<sup>122,125</sup> Evidence shows that the nature of the large-scale mobility of the fluids (*i.e.*, flow displacement patterns) is linked to pore-scale physics and interfacial instabilities. It is also increasingly accepted that developing subgrid-scale models capable of accurately representing pore-scale processes is critical for improving the accuracy of reservoir-scale simulations.<sup>126–128</sup> In this regard, extensive microfluidic studies have been conducted to probe, observe, quantify, and understand various pore-scale mechanisms at the microscopic scale. These studies have been pivotal to advancing our understanding of multiphase flow in porous media, miscible and immiscible displacement processes, flow regimes, interfacial instabilities, Haines jumps, capillary pressure and hysteresis, and wettability effects.<sup>97,122–160</sup> As mentioned previously, one significant advantage, among others, of microfluidics is the valuable optical access it offers, which allows for direct observation and quantification of pore-scale mechanisms enabled by advanced optical flow diagnostics

such as epi-fluorescent microscopy, multi-color microscopy, high-speed imaging, and particle imaging velocimetry (PIV).<sup>129–133,136,138</sup> Microfluidic experiments combined with those imaging techniques have characterized microscopic processes with great temporal and spatial resolutions, which would not have been possible in core sample experiments and field observations.<sup>161</sup> These microfluidic studies and findings are reviewed in detail in section 3.3.

### 3.2. Microfluidic models for CO<sub>2</sub> storage studies

A microfluidic porous model system, so-called micromodel, is typically defined as an artificial representation of a porous medium made of a transparent material. Such a device must: (i) bear a flow network, (ii) contain features of micrometers to sub-millimeters in size, and (iii) offer a certain level of optical access. The flow network consists of connected throats and pores through which fluids of one or more phases and/or components flow. Since the advent of microfluidic devices, arguably by Chatenever *et al.* and later by Lenormand *et al.*, numerous micromodels have been developed and constructed to study the physical, chemical, and biological processes during carbon storage in porous media.<sup>162,163</sup> This section provides a brief description of the different types of microfluidic devices used in carbon storage studies, with a focus on a few unconventional devices that have been reported just recently. For a comprehensive knowledge of micromodels and the relevant fabrication and imaging techniques, the readers are referred to several reviews by Karadimitriou *et al.*, Gerami *et al.*, and Jahanbakhsh *et al.*<sup>132,133,164</sup> This review is to update and complement them from a microfluidics perspective.

**3.2.1. Conventional micromodels.** Common materials used to produce micromodels include glass and silicon, which are inorganic, and polymers, such as polydimethyl siloxane (PDMS) and polymethyl methacrylate (PMMA), which are organic.<sup>131,133</sup> Glass has been widely used to fabricate



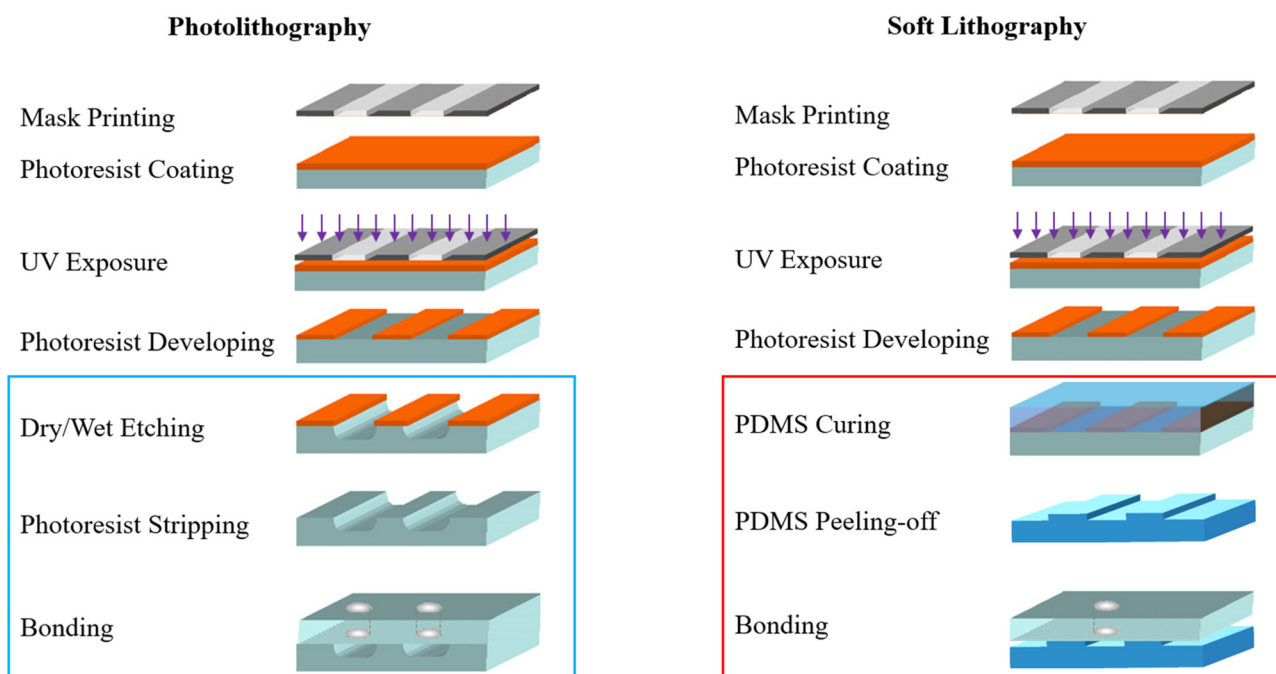
micromodels for its excellent optical transparency, chemical compatibility, and good thermal and mechanical properties.<sup>129,163</sup> A pre-designed porous pattern is etched into a glass slide, which can then be bonded thermally or adhesively to a second glass slide to form a closed porous flow network. Despite the complexity of the fabrication of glass-based micromodels, a couple of recent studies have creatively taken advantage of the isotropic etching behavior of glass to create so-called 2.5D micromodels, whose depth was purposely designed to vary, allowing for the study of pore-scale mechanisms that were not achievable with regular 2D micromodels.<sup>156,165</sup>

Silicon is another common material used to fabricate micromodels. As silicon is not transparent in the visible spectrum, an etched silicon wafer that carries the flow network is usually anodically bonded with a glass wafer, through which the internal flow can be observed optically.<sup>143,146,166,167</sup> The etching of silicon using DRIE is highly directional, enabling close to 90° side walls in etched microchannels, which is crucial to producing 2D micromodels. Additionally, PDMS and PMMA have been extensively used to fabricate 2D micromodels for their ease of fabrication and cost-effectiveness, with a possible fabrication of 3D micromodels.<sup>168</sup> However, PDMS and PMMA are mechanically weak compared to silicon and glass and incompatible with many chemicals and common organic solvents, rendering them unsuitable for high-pressure and high-temperature experiments.<sup>169,170</sup> Other materials that have been successfully used to fabricate micromodels include UV-sensitive polymers and

geomaterials such as calcite and rock slices.<sup>171–173</sup> Several studies reported the fabrication and use of high-pressure micromodels, which allowed direct measurement of multiphase flow at reservoir-relevant pressures.<sup>174–176</sup> The readers are referred to the review by Jahanbakhsh *et al.* (2020) for common materials as well as their advantages and disadvantages for micromodel fabrication.<sup>133</sup>

The fabrication method of a micromodel depends on the material selected. The two most common techniques include photolithography, which is used to process glass and silicon, and soft lithography, which is used to process PDMS and many other polymers. These methods are further complemented by different techniques, including micromachining, direct laser writing, mineral coating and deposition, and, more recently, 3D printing, as summarized in Gerami *et al.* and Jahanbakhsh *et al.*<sup>132,133</sup> As illustrated in Fig. 21, a standard photolithography process involves mask making, photoresist spin coating, UV exposing, and developing, which is followed by etching, cleaning, and bonding. On the other hand, soft lithography, which is also initiated with mask-making and photoresist processing, finishes with PDMS molding.

**3.2.2. Unconventional micromodels.** In contrast to a few recent reviews, herein, we highlight a few innovative micromodels that have enabled new measurements but are not sufficiently covered and discussed in other reviews.<sup>131–133,164</sup> One trend of micromodel-based CO<sub>2</sub> storage research is to create “non-reactive” artificial micromodels with more complex and realistic porous geometries to better represent the real geology of the storage



**Fig. 21** [Left] Major fabrication steps carried out for photolithography and [Right] soft lithography. While both methods start with mask making and photoresist processing, photolithography typically finishes with etching and bonding of silicon and/or glass, but soft lithography finishes with PDMS molding.





formations. The 2.5D micromodels and dual-permeability micromodels are good examples in this regard. Zhang *et al.* developed a dual-permeability pore network micromodel, which contains two distinct permeability zones, each occupying one-half of the micromodel width, to better emulate permeability contrasts that exist in multilayer geological formations under consideration for carbon sequestration.<sup>177</sup> Their results showed that heterogeneous permeability at the microscopic pore scale may directly influence aquifer storage capacity. In the dual-permeability pore network, a permeability contrast factor of approximately two resulted in the preferential displacement of water by liquid CO<sub>2</sub> in the high-permeability zone. Buchgraber *et al.* tested an etched-silicon micromodel with dual porosity, mimicking those found in certain carbonate reservoir rocks.<sup>178</sup> This dual porosity has reportedly enabled the observations of pore-level mechanisms of multiphase flow and the interpretation of petrophysical properties. Later, Yun *et al.* developed an optimized sequential photolithography protocol to etch micropores less than the depth of wider macropores to improve the structural realism of a single-depth micromodel with a carbonate-derived pore structure, which paves the way for the systematic investigation of the effect of various dimensions on multiphase flow in porous media (Fig. 22).<sup>165,179</sup>

While these non-reactive micromodels have been serving as excellent platforms to investigate physical processes, they are not suitable for real carbon storage studies where erosive reactions are expected. For such cases, it is critical to fabricate micromodels using chemically reactive materials that are geologically representative of the subsurface environment. Several studies have explored “bottom-up” or “top-down” approaches to creating micromodels with reactive minerals such as calcite.<sup>171,172,180,181</sup> Bottom-up approaches have grown reactive minerals (*e.g.*, CaCO<sub>3</sub>) *in situ* within a traditional non-reactive micromodel (Fig. 23). Lee *et al.* developed a technique to selectively grow CaCO<sub>3</sub> in a glass

microchannel to form a porous section.<sup>182</sup> The method is enabled by a UV-curable precursor solution, which creates preferential sites for CaCO<sub>3</sub> growth. Upon delicate controls of the rinsing and precipitating flows, CaCO<sub>3</sub> posts were successfully grown with pre-defined geometry, as shown in Fig. 23. This approach, while novel and precise, requires sophisticated control of various parameters and has a low throughput. Wang *et al.*, Song *et al.*, and Alzahid *et al.* took a similar approach by fabricating a standard micromodel in glass, silicon, and PDMS, respectively, and grew a thin layer or grains of CaCO<sub>3</sub> crystals within.<sup>180,181,183</sup> The approach is relatively straightforward to implement, but the major drawback is that the original surfaces (*i.e.*, glass, silicon, or PDMS) of the micromodels are either not fully covered with CaCO<sub>3</sub> or only by a thin layer of CaCO<sub>3</sub> of 0 (1 μm) thick, making them unsuitable for studies of the dissolution rate or prolonged fluid–mineral interactions.

Top-down strategies often employ traditional micro-milling and etching combined with laser cutting and patterning for better precision.<sup>172,184</sup> The work by Song *et al.* is among the first studies to successfully fabricate a calcite-based micromodel using laser cutting and wet etching (Fig. 23).<sup>172</sup> The process started with thin sectioning a large block of natural calcite crystal, following which a porous pattern was generated with the aid of a layer of beeswax and a laser cutter. The micromodel was completed by wet etching in hydrochloric acid, drilling, and bonding to another piece of glass. While this approach is innovative, its precision is relatively limited, with a minimum feature size of approximately 140 μm. Soulaire *et al.* and Rembert *et al.* adopted a similar idea of using a thin calcite slice that is pre-machined, and the thin calcite of a pre-defined shape was then directly embedded in a straight PDMS microchannel.<sup>185,186</sup> However, the entire micromodel consisted of only one calcite post, hardly justified as a simulation of geologic media. To take one step further, Singh *et al.* embedded a real rock slice 500 μm thick into a PDMS

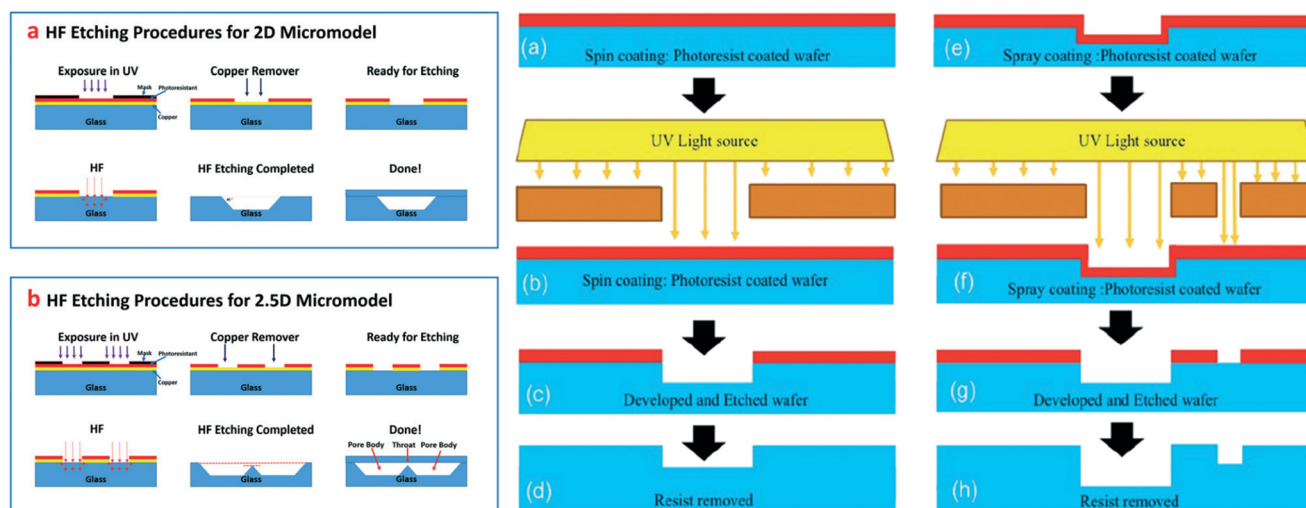
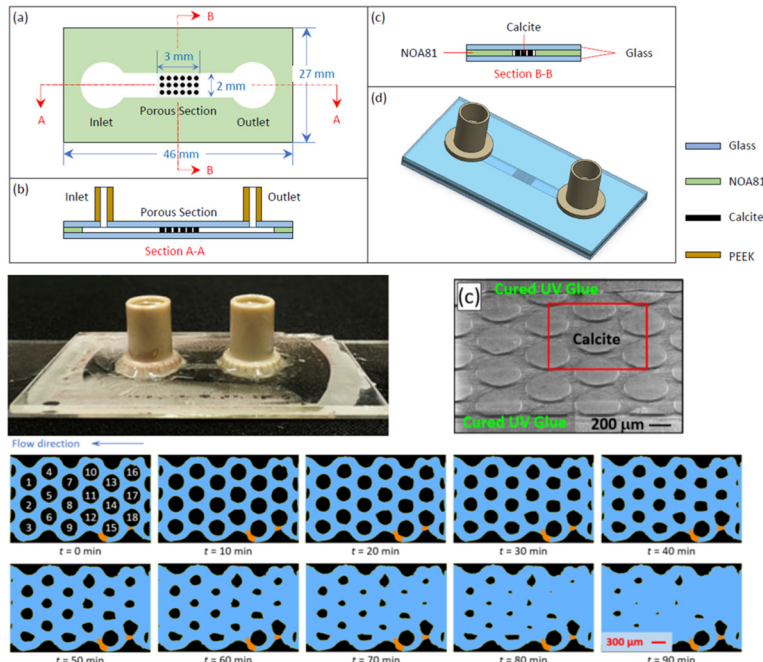
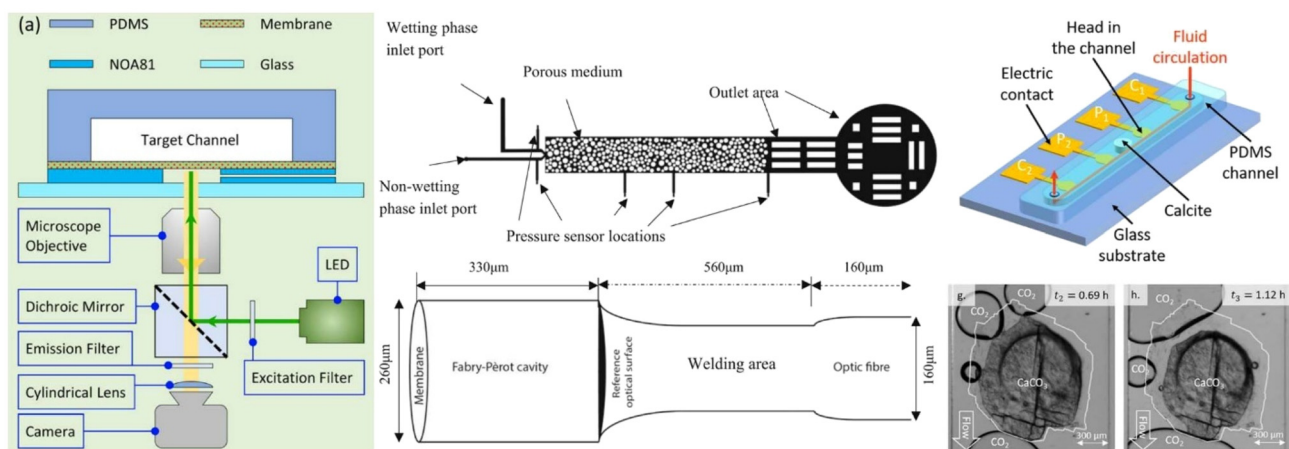


Fig. 22 [Left] Fabrication steps of a 2.5D micromodel.<sup>165</sup> [Right] Fabrication steps of a dual-porosity dual-depth micromodel.<sup>179</sup>



channel.<sup>171</sup> Thanks to the use of real rock slices, this micro model is faithfully representative of real geometry and geochemistry, thus being ideal when quantifying global quantities such as reaction rates and pressure drops. However, the use of such natural material makes each micromodel different and unique in terms of physical, geological, and chemical properties, hindering the necessary repeatability test at the pore scale as well as the possibility of parametric studies of pore flow using control variables. Moreover, the opaqueness of the rock sample renders the optical-based measurement challenging and sub-optimal.

Most recently, Rahman *et al.* reported an innovative microfluidic approach that includes novel 2D reactive porous media and advanced pore flow diagnostics for studying pore-scale dissolution in porous media with unprecedented details (Fig. 23).<sup>184</sup> The 2D micromodels were fabricated in calcite by combining photolithography and wet etching directly on the calcite sample, which facilitates unobstructed optical access to the pore flow, improving over previous methods. Additionally, by molding the microchannel with UV glue, the authors could eliminate unwanted reactions outside the porous section, thus achieving more controlled flow and



**Fig. 24** [Left] A schematic diagram illustrating the on-chip pressure sensor enabled by astigmatic particle tracking.<sup>187</sup> [Middle] Micromodel with integrated fiber-optical pressure sensors.<sup>188</sup> [Right] 3D sketch of the microfluidic chip with integrated SIP sensors and sample results of calcite dissolution.<sup>186</sup>

reactions. Their results revealed the crucial roles of reactive transport and local concentration gradients in mineral dissolution in porous media and called for reconsidering many assumptions (Fig. 24).

In addition, several recent developments attempted to integrate on-chip sensors into the porous section to enable *in situ* measurement of pore-scale quantities, such as pore pressure. Zarikos *et al.* reported the manufacturing procedure of a micromodel with integrated fiber optic pressure sensors with a circular measurement window of 260  $\mu\text{m}$  in diameter, which measures pressure at the pore scale.<sup>188</sup> The integrated micromodel was tested for both single- and two-phase flows, which led to valuable insights into the link between fast pressure changes and pore-scale events. More recently, Raventhiran *et al.* reported the design and fabrication of an on-chip sensor that quantifies pressure in microfluidic devices based on a novel technique called astigmatic particle tracking.<sup>187</sup> With this technique, thin membranes that sense pressure variations in the fluid flow can be characterized conveniently by imaging the shapes of the particles embedded in the membranes. This innovative design only relies on the reflected light from the back of the microchannel, rendering the sensor separate and noninvasive to the flow of interest. This sensor was then applied to characterize the pressure drop in single-phase flows with an accuracy of  $\sim 70$  Pa. Rembert *et al.* developed a complex electrical conductivity acquisition using the spectral-induced polarization (SIP) method on a microfluidic chip equipped with electrodes. Enabled by high-speed microscopy, the micromodel provides direct observation and monitoring of microscopic reactive transport processes in porous media, highlighting the strong correlation between SIP response and dissolution.<sup>186</sup>

### 3.3. Flow dynamics in CO<sub>2</sub> storage within microfluidics

Micromodels are well-suited to identify, observe, and characterize pore-scale processes when combined with advanced optical diagnostic techniques such as epifluorescence microscopy and micro-PIV.<sup>137,140,147,163,166,189–192</sup> In fact, the application of micromodels and microfluidics has directly contributed to the discovery of numerous pore-scale mechanisms in carbon storage, which is the primary focus of this section.

**3.3.1. Flow regime and displacement efficiency.** Many microfluidic studies in carbon storage are directly related to the immiscible displacement processes of multiple fluids in porous media, as flow displacement patterns and regimes directly determine the saturation of the injected CO<sub>2</sub>, further defining the storage capacity. In particular, the drainage process (as opposed to imbibition) has arguably attracted more attention, where a wetting resident fluid (the resident brine in saline aquifers, for example) is displaced by a non-wetting invading phase (CO<sub>2</sub>). Based on our current understanding, at short-time scales, where chemical processes such as dissolution and precipitation are not

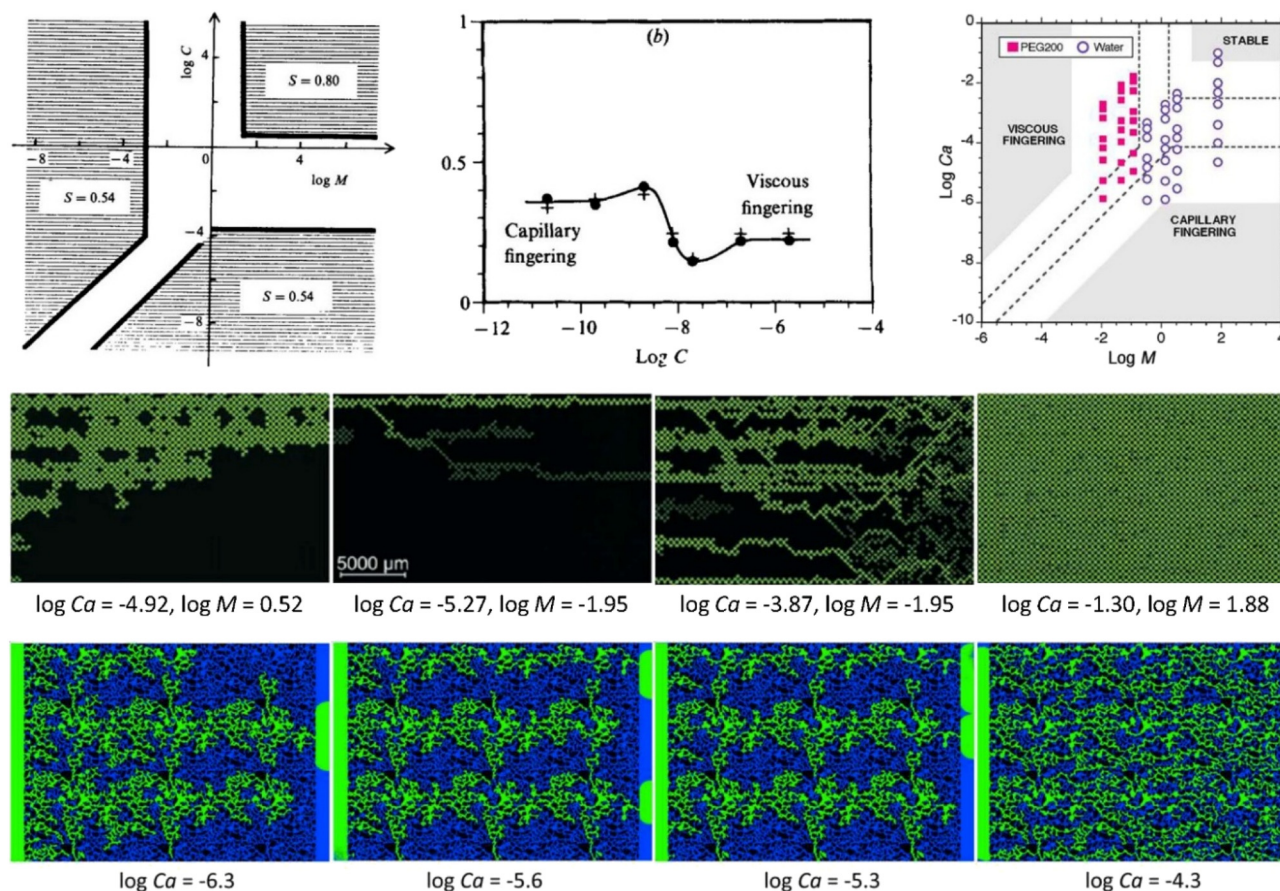
significant and in the absence of gravitational forces, the pore-scale displacement of a wetting phase by a non-wetting phase in porous media is inherently governed by two forces: capillary forces and viscous forces.<sup>137,163</sup> The relative importance of these two forces is, in turn, characterized by two dimensionless numbers: the viscosity ratio  $M = \frac{\mu_n}{\mu_w}$ , where  $\mu_n$  and  $\mu_w$  are the dynamic viscosities of the nonwetting and wetting phases, respectively, and the capillary number  $\text{Ca} = \frac{\mu_n U}{\sigma \cos \theta}$ , where  $\sigma$  is the surface tension between the two fluids,  $\theta$  is the contact angle between the fluids and the solid surfaces, and  $U$  is the bulk flow velocity. Here,  $U = Q/(A\phi)$ , where  $Q$  is the volumetric flow rate,  $A$  is the cross-sectional area of the flow domain, and  $\phi$  is the porosity of the porous media. Depending on the balance of viscous and capillary forces, three distinct flow regimes exist: (1) stable displacement, (2) viscous fingering, and (3) capillary fingering. For the specific case of CO<sub>2</sub> injection into deep saline aquifers, the flow falls into either the viscous fingering or capillary fingering regime, depending on the injection rate, the distance from the injection well, and the stage of the injection process (Fig. 25).

Lenormand *et al.* are among the first ones to investigate the flow regimes or so-called “phase diagram” of multiphase flow in porous media using a series of experiments in a homogeneous micromodel (*i.e.*, featuring a regularly arranged porous structure) for a large parameter space, which essentially laid the foundation of the current understanding of pore-scale multiphase flow in porous media.<sup>137</sup> Their experiments clearly showed the existence of the three basic domains (*i.e.*, capillary fingering, viscous fingering, and stable displacement), as shown in Fig. 25, within which the patterns remain unchanged.<sup>137</sup> Cottin *et al.* illustrated the role of the viscous forces in the invasion process and their competition with the capillary force heterogeneities.<sup>141</sup> Later, Zhang *et al.* conducted displacement experiments in a water-wet pore network micromodel with microfabricated pore structures.<sup>189</sup> Using seven wetting–non-wetting fluid pairs, they explored viscosity ratios ( $\log M = -1.95$  to  $1.88$ ) and capillary numbers ( $\log \text{Ca} = -5.88$  to  $-1.02$ ) across four orders of magnitude. In the experiments with PEG200 as the resident phase ( $M < 1$ ), unstable displacement was observed by viscous fingering over all capillary numbers. For the experiments with water as the resident phase, unstable displacement occurred by capillary fingering at low capillary number, whereas crossover into stable displacement was observed for the fluid pairs with  $M > 1$  when the viscous forces were increased by increasing the injection rate.<sup>137</sup>

Wang *et al.* continued the work by Zhang *et al.* with two sets of experiments: discontinuous-rate injection, where the micromodel was saturated with water before each injection rate was imposed, and continuous-rate injection, where the rate was increased after quasi-steady conditions were reached.<sup>189,193</sup> For the discontinuous-rate experiments, capillary fingering and viscous fingering are the dominant mechanisms for low ( $\log \text{Ca} \leq -6.61$ ) and high injection rates







**Fig. 25** [Top left] The original phase diagram achieved by Lenormand *et al.*, demonstrating the existence of capillary fingering, viscous fingering, stable displacement regimes, and the crossover between them.<sup>137</sup> [Top middle] Plot of the invading fluid at breakthrough, showing a decrease of the nonwetting phase saturation at the crossover between capillary and viscous fingering.<sup>137</sup> [Top right] An updated phase diagram obtained by Zhang *et al.*<sup>189</sup> [Middle row] Representative displacement patterns for capillary fingering, crossover, viscous fingering, and stable displacement, respectively, captured in homogeneous micromodels.<sup>189</sup> [Bottom row] Representative displacement patterns going from capillary fingering to crossover, and finally to viscous fingering as the capillary number increases in 2D heterogeneous micromodels.<sup>166</sup>

( $\log Ca \geq -5.21$ ), respectively. However, crossover from capillary to viscous fingering was observed for  $\log Ca$  ranging from  $-5.91$  to  $-5.21$ , resulting in a large decrease in  $\text{CO}_2$  saturation, in agreement with the numerical predictions by Lenormand *et al.*<sup>137</sup> Recently, Li *et al.* studied the pore-scale flow of  $\text{CO}_2$  and water in 2D heterogeneous porous micromodels over a  $Ca$  range of nearly three orders of magnitude.<sup>143</sup> It was reported that under conditions relevant to  $\text{CO}_2$  sequestration, final  $\text{CO}_2$  saturation first decreases and then increases logarithmically with  $Ca$  within the capillary and viscous-fingering regimes, respectively, with a minimum occurring during the crossover regime. A recent study by Rabbani *et al.* demonstrated with microfluidic experiments that gradual and monotonic variation of pore sizes along the front path suppresses viscous fingering during immiscible displacement, which not only provides insights into ways for suppressing unwanted interfacial instabilities in porous media but also opens the door to a new way to understand flow instability in porous media.<sup>194</sup>

While a tentative consensus has been achieved regarding the flow regimes during drainage, conflicting findings have

been reported in the literature regarding the relationship between the final saturation of the invading phase and  $Ca$  at different  $M$ . Some recent studies reported a monotonically increasing relation, whereas others indicated either a decreasing saturation or nonmonotonic behavior.<sup>135,141,189</sup> For a fluid pair with  $\log M = -4.7$ , Lenormand *et al.* found that the saturation decreases during the crossover, with two high-saturation plateaus for low and high  $Ca$  corresponding to capillary and viscous fingering, respectively.<sup>137</sup> Wang *et al.* noted a substantial decrease in  $\text{CO}_2$  saturation between  $\log Ca = -5.91$  and  $-5.21$ , which is consistent with the findings of Lenormand *et al.*<sup>137,193</sup> Chen *et al.* investigated the crossover during water displacing oil ( $\log M = -3, -2.7, -2, -1.7$ ) in a hydrophobic rough fracture for  $-7.07 \leq \log Ca \leq -3.07$  and observed that the saturation of the invading fluid first decreases and then increases with increasing  $Ca$ , with the minimum value occurring at  $\log Ca = -4.07$  or  $\log Ca = -5.07$ , depending on  $M$ .<sup>139</sup> This agrees with the study in a 2D heterogeneous micromodel by Li *et al.*, where again a minimum  $\text{CO}_2$  saturation was observed during the crossover from capillary fingering to viscous



fingering.<sup>143</sup> The existence of a minimum value was attributed to the fact that both fingering propagation toward the outlet and void filling in the transverse/backward directions were suppressed during the crossover. The inconsistent findings certainly call for additional endeavors for more insights and deterministic measurements.

**3.3.2. Inertia effect.** During capillary-dominated fingering, the fluid–fluid interfaces are susceptible to strong instabilities. The interface dynamics are crucial to a deeper understanding of fluid migration in porous media. Theories, such as the percolation theory, are typically based on the assumption that the invasion of pores is solely controlled by the local capillarity (and thus the porous media geometry), and the dynamics are often not considered. However, from a pore-scale perspective, the macroscopic movement of the drainage process is essentially an ensemble of high-speed interfacial burst events occurring within individual pores, termed Haines jumps.<sup>195</sup> Typical Haines jumps occur at milliseconds and can appreciably affect a length scale of up to 30 pore diameters.<sup>166,196</sup> Several studies have shown that Haines jumps play a key role in drainage processes by effectively creating nonlocal velocities of large magnitudes, enhancing inertial effects, affecting displacement regimes, and eventually shaping macroscopic fluid morphology.<sup>143,145,146,148,166,196,197</sup> Microfluidic experiments have greatly contributed to the revelation and quantification of inertial effects.

Moebius and Or studied the rapid interfacial dynamics during drainage, showing interfacial velocities exceeding 50

times the mean front velocity.<sup>148</sup> Armstrong and Berg imaged pore drainage events in a glass micromodel system and analyzed the dependency of interfacial velocity on the bulk flow rate and spatial fluid configurations, as shown in Fig. 26.<sup>196</sup> Their results indicated that pore drainage events are cooperative, revealing that capillary pressure differences over multiple pores affect fluid topology and menisci dynamics, which highlight the inertial effect. Additionally, it was discovered that the velocity of a pore drainage is controlled by its intrinsic dynamics and is, therefore, independent of the bulk flow rate. Kazemifar *et al.* quantified the flow dynamics associated with water displacement by CO<sub>2</sub> in a 2D homogeneous porous micromodel.<sup>146</sup> Employing dual-color fluorescent microscopy and micro-PIV techniques, they were able to capture the pore-scale burst events (Haines jumps) with sufficient time resolution, which showed that velocity jumps 20–25 times larger in magnitude than the bulk velocity and that these bursts of water flow occurred both in-line with and against the bulk flow direction. With the velocity data captured with micro-PIV, they calculated that the maximum local Reynolds number *Re* for CO<sub>2</sub> reaches as high as 25, which is well beyond the limit of *Re* = 10 for the validity of Darcy's law, highlighting the effects of inertia. Li *et al.* extended the study to a 2D heterogeneous micromodel whose porous structures were inspired by sandstone CO<sub>2</sub> storage reservoirs. They measured a local *Re* of 21 in the CO<sub>2</sub> phase.<sup>166</sup> Furthermore, these drainage events were observed to be cooperative, extending beyond tens of pores, confirming that Haines jumps are non-local phenomena.<sup>167</sup> Some

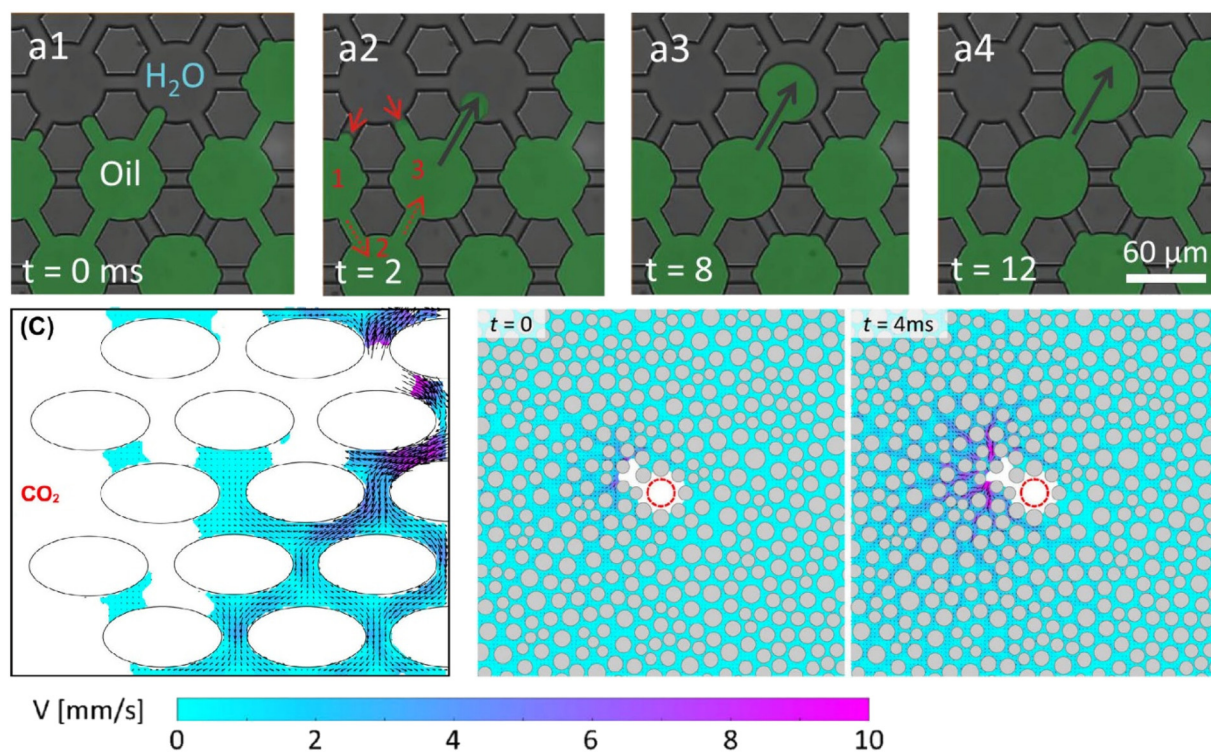


Fig. 26 [Top] High-speed images of a Haines jump event at  $Ca = 3.4 \times 10^{-6}$ .<sup>196</sup> [Bottom] Velocity vector fields during a Haines jump event in linear drainage and radial drainage.<sup>146,167</sup>





representative velocity fields captured during the occurrences of Haines jumps are shown in Fig. 26. Informed by these microfluidic studies, several numerical studies attempted to incorporate an inertial effect in the model and noted that the capillary number is not sufficient to characterize the two-phase flow, as suggested by Lenormand *et al.*<sup>137,140,198–200</sup> Instead, the Ohnesorge number, which gives the relative importance of viscous forces over inertial and capillary forces, is required to fully describe the fluid flow, along with the viscosity ratio.<sup>137</sup>

### 3.3.3. Capillary pressure, hysteresis, and interfacial area.

Capillary pressure ( $P_c$ ) is the pressure difference across the interface between two immiscible fluids arising from the surface tension or interfacial tension. In conventional models of two-phase flow in porous media, a  $P_c$ – $S_w$  relation is necessary to enable the closure of the equation system that is derived based on conservation laws.<sup>201–203</sup> Here, saturation ( $S_w$ ) denotes the amount of one fluid phase present in the porous section at an instance relative to the other phase, and  $P_c$ – $S_w$  relations are often developed empirically.  $P_c$ – $S_w$  relations are long known to be hysteretic, *i.e.*, a single saturation could correspond to multiple capillary pressures and, hence, multiple states of a system, significantly limiting the accuracy of available predictive tools. Several recent theoretical studies have shown that hysteresis behavior can potentially be eliminated by accounting for the pore-scale topological features of the flow, such as interfacial area, interfacial curvature, and Euler characteristics.<sup>204–206</sup> Additionally, characterization of the fluid–fluid interfacial area is critical for understanding many heat, momentum, and mass transfer processes, including shear-induced flow, dissolution, exsolution, and subsequent chemical reactions, all occurring at the fluid–fluid interface.<sup>143,146,152,166,189,191</sup> Several microfluidic studies have focused on characterizing the behavior of fluid–fluid interfaces and pore-scale capillary pressure in 2D porous micromodels.<sup>143,150–154,189</sup>

Micromodels are well suited for characterizing interfacial areas thanks to their excellent optical access and the availability of various optical flow diagnostic tools. In 2D micromodels, the interfacial area reduces to interfacial length due to the 2D nature of the flow, assuming the depth of the micromodel is constant. Tsakiroglou *et al.* performed unsteady displacement experiments in a glass-etched pore network to examine the impact of the capillary number on relative permeability and capillary pressure.<sup>154</sup> They found that these properties are sensitive to the transient displacement pattern, which changes from the invasion percolation to frontal drive as the capillary number increases, highlighting the influence of the fluid configuration and interfacial area on capillary pressure. Cheng *et al.* studied interfacial areas in the  $P_c$ – $S_w$  relation during quasi-static drainage and found that the specific interfacial area lifts the hysteresis in the capillary pressure and saturation relationship in porous media.<sup>207</sup> Their work provided the first experimental validation of theoretical predictions, showing that the capillary-dominated subset plays a role like

a state variable and that the interfacial area is essential for describing multiphase flow properties, as well as capillary pressure and fluid saturation.<sup>201</sup> Karadimitriou *et al.* studied interface movement under transient conditions using a 2D PDMS micromodel.<sup>151</sup> While their results produced a unique  $P_c$ – $S_w$ – $a_{wn}$  surface for both drainage and imbibition, different relationships were obtained under steady state and transient conditions. This suggests that one interfacial area surface cannot sufficiently describe two-phase flow under transient and quasi-static conditions. This mismatch was attributed to the disconnection of the non-wetting phase, and the interfacial areas between the solid and the two fluid phases were not included in the capillarity theory, which warrants further research.

Zhang *et al.* calculated the total interfacial length in 2D micromodels at different flow stages and found a linear relationship between specific interfacial length and non-wetting fluid saturation.<sup>177</sup> In a dual-permeability micromodel, Zhang *et al.* reported that while the linearity between interfacial length and nonwetting fluid saturation is preserved for both low and high flow rates, the slope for high flow rates was much larger than that for low flow rates. This behavior was attributed to a geometrical effect: the smaller pores, which are invaded only at high flow rates, are associated with higher interfacial length.<sup>177</sup> Liu *et al.* confirmed this linear relationship for both homogeneous and heterogeneous pore networks using LBM simulations and observed a higher slope for higher flow rates.<sup>152</sup> It was conjectured that the higher slope results from viscous fingering in which some fingers are stretched very thinly and even broken into smaller ganglia, resulting in a higher interfacial length than in capillary fingering and stable displacement. To that end, Li *et al.* performed high-speed measurements in 2D heterogeneous micromodels and confirmed that the total specific interfacial length scales linearly with CO<sub>2</sub> saturation, which is in agreement with what was previously reported in the literature.<sup>143</sup> Higher slopes were noted for high Ca cases, likely because CO<sub>2</sub> fingers stretch or even break into ganglia in the viscous-fingering regime. They also noted the critical role of inertia in creating significant numbers of isolated CO<sub>2</sub> ganglia *via* snap-off events at high Ca, which in turn increases total capillary-associated interfaces and reduces film-associated interfaces. The reduction and potential elimination of capillary hysteresis through developing new functional relationships that include a more complete set of variables is highly desirable. Future research is needed to enable more accurate determination of the interfacial area and other functional variables (*e.g.*, interfacial curvature and Euler characteristics) and direct measurement of pore-scale pressure using on-chip microscale pressure sensors.<sup>187,188</sup>

**3.3.4. Effect of wettability.** Wettability refers to the “affinity” of a solid surface for one fluid in the presence of one or more other immiscible fluids, and it is often characterized by the contact angle,  $\theta$ . Wettability is difficult to control experimentally and model numerically and poses a

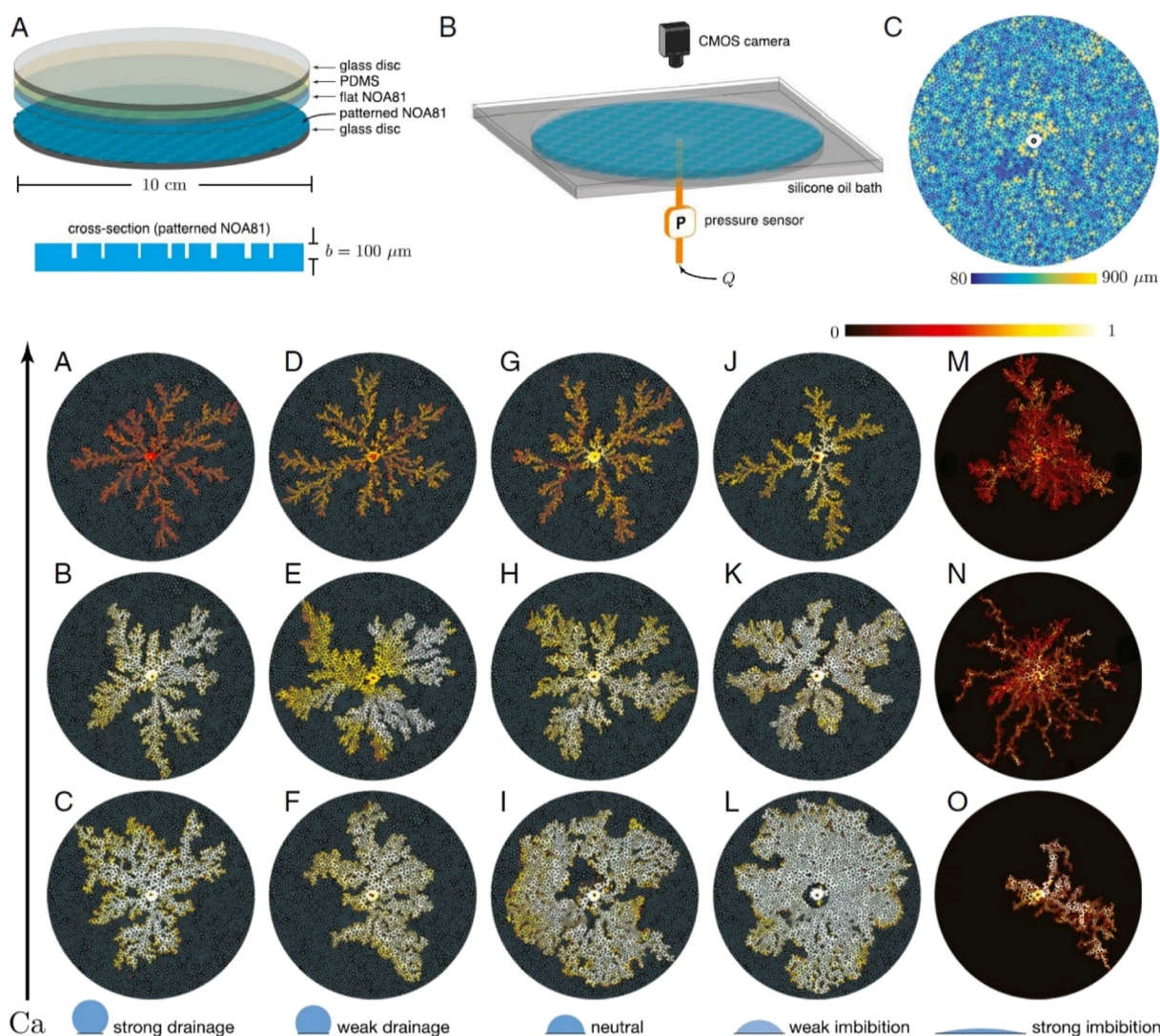




further layer of complexity in the study of multiphase flow in porous media.<sup>157,167,173,204,208,209</sup> Based on wettability, a fluid–fluid displacement process in porous media can be classified into two types: drainage and imbibition. Although a wide range of wettability conditions is possible, a practical CO<sub>2</sub> storage application typically encounters a drainage process.<sup>156</sup> For instance, while deep saline aquifers are typically water-wet to CO<sub>2</sub> injections, depleted hydrocarbon reservoirs can be intermediate-wet or mixed-wet.<sup>210,211</sup> These conditions may change over time due to rock reactions modifying the surface properties of the porous matrix.<sup>205,207</sup> Wettability is important as it is well known to dramatically impact the efficiency of the displacement process and, therefore, affect the efficiency and capacity of CO<sub>2</sub> storage.<sup>212</sup>

From a pore scale perspective, wettability governs the capillary force and directly impacts the interface stability and

displacement efficiency. Most previous studies on the wettability effects on flow behavior in porous media reported that increasing the contact angle of the resident wetting phase (*i.e.*, making the wetting phase less wetting) stabilizes the displacement interface under various flow conditions, leading to more compact displacement patterns and thus increasing the displacement efficiency.<sup>153,209</sup> Herein, we provide an overview of several representative microfluidic studies. Trojer *et al.* experimentally studied the wettability impacts on fluid–fluid-displacement patterns in granular media using a thin bed of glass beads.<sup>209</sup> They varied the glass surface contact angle from 5° (drainage) to 120° (imbibition) through a combination of cleaning and salinization procedures. Holtzman conducted micromodel experiments in a plastic micromodel to understand the role of pore-scale heterogeneity and wettability in fluid



**Fig. 27** [Top left] Schematic showing the micromodel used for the study of wettability effects by Zhao *et al.* (2016). [Top right] The micromodel was made of a photocurable polymer (NOA81) patterned with circular posts. [Bottom] displacement patterns for various wettability conditions (left to right:  $\theta = 150^\circ$ ,  $120^\circ$ ,  $90^\circ$ ,  $60^\circ$ ,  $7^\circ$ ) and capillary numbers (bottom to top:  $Ca = 2.9 \times 10^{-3}$ ,  $2.9 \times 10^{-2}$ ,  $2.9 \times 10^{-1}$ ); note the contact angles were defined with reference to the invading fluid ( $\theta > 90^\circ$  and  $\theta < 90^\circ$  correspond to drainage and imbibition, respectively).<sup>173</sup>



displacement.<sup>157</sup> Their results show that increasing the contact angle suppresses both trapping and fingering, hence reducing the sensitivity of the displacement to the underlying geometry disorder. Jung *et al.* studied the wettability effects in microfluidic Hele-Shaw cells where the contact angle was varied from 0° to 134°. <sup>213</sup> The results indicated a consistent crossover between stable interfacial displacement at contact angles greater than 100° and capillary fingering at low contact angles less than 60°. Hu *et al.* used a high-pressure micromodel-microscopy system to study supercritical CO<sub>2</sub> invasion into brine-saturated water-wet and intermediate-wet micromodels and observed a smaller number of fingers with larger finger width under intermediate-wet conditions.<sup>208,214</sup> Zhao *et al.* performed a more comprehensive investigation of wettability effects by systematically varying the wettability of the porous flow cell over a wide range of contact angles.<sup>173</sup> They found that increasing the solid matrix affinity to the invading fluid results in a more efficient displacement of the resident fluid up to a critical wetting transition, beyond which the trend is reversed (Fig. 27). This behavior was attributed to two pore-scale mechanisms: cooperative pore filling (increasing displacement efficiency) and corner flow (decreasing displacement efficiency). This work suggests that wettability is a necessary dimension, in addition to the capillary number,  $Ca$ , and viscosity ratio,  $M$ , to fully describe two-phase flow in porous media.

Li *et al.* investigated the pore-scale flow dynamics of liquid CO<sub>2</sub> and water in 2D micromodels with different wetting properties using high-speed micro-PIV under reservoir-relevant conditions for drainage and imbibition scenarios.<sup>167</sup> They noted that when CO<sub>2</sub> displaces water in a hydrophilic micromodel (*i.e.*, drainage), unstable capillary

fingering occurs, and the pore flow is dominated by successive pore-scale burst events (*i.e.*, Haines jumps). However, when the same experiment was repeated in a nearly neutral wetting micromodel (*i.e.*, weak imbibition where capillary forces are minimized), flow instability and fluctuations were virtually eliminated, leading to a more compact displacement pattern. Enabled by high-speed microscopy, their measurements further revealed a significant alteration of the pore-filling mechanisms during drainage and imbibition, as shown in Fig. 28. While the former typically featured burst events, which often occur only at one of the several throats connecting a pore, the latter is typically dominated by a cooperative filling mechanism involving simultaneous invasion of a pore from multiple throats. This cooperative filling mechanism leads to the merging of two interfaces and releases surface energy, causing instantaneous high-speed events that are similar yet fundamentally different from burst events.

Chang *et al.* performed supercritical CO<sub>2</sub> injection into heterogeneous micromodels while considering three different wetting conditions: uniformly water-wet, uniformly intermediate-wet, and mostly water-wet with intermediate-wet patches.<sup>156</sup> High storage efficiency and wide interconnections of CO<sub>2</sub> flow paths were observed in reservoirs containing more and uniformly distributed intermediate-wet and water-wet patches, whereas hindered storage efficiency and channelized CO<sub>2</sub> flow paths were observed in reservoirs containing heterogeneously distributed intermediate-wet patches. Most recently, Irannezhad *et al.* studied the displacement of silicone oil by water in a mostly oil-wet porous medium patterned with discrete water-wet clusters that have precisely controlled

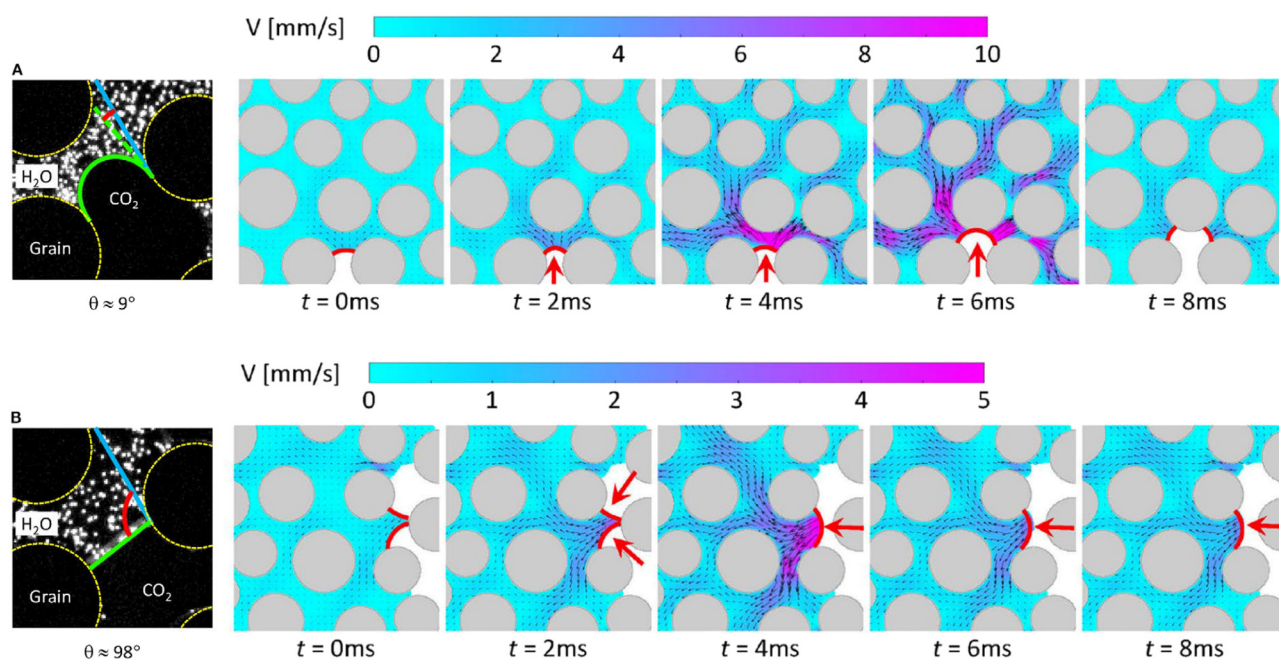


Fig. 28 [Top] Evolution of velocity fields during a Haines jump event in drainage. [Bottom] Evolution of velocity fields during a cooperative filling event in weak imbibition.<sup>167</sup>



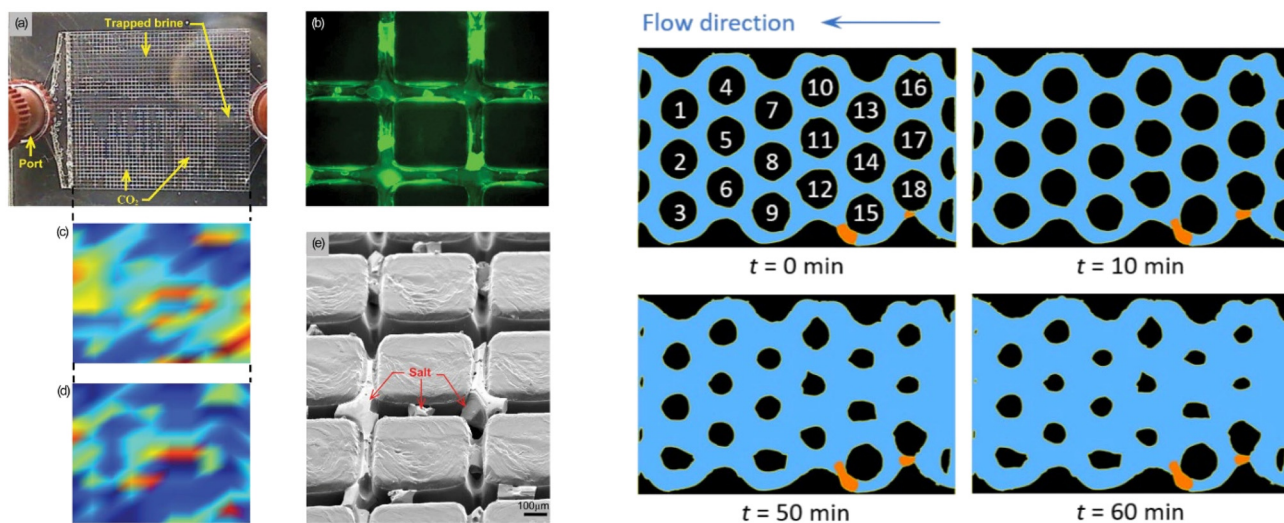


wettability.<sup>215</sup> It was observed that the macroscopic displacement pattern varies dramatically depending on the details of wettability alteration. The invading water preferentially fills strongly water-wet clusters but encircles weakly water-wet clusters instead, resulting in significant trapping of the defending oil. The experimental observation was explained with pore-scale simulations, which revealed that the fluid–fluid interfaces at mixed-wet pores resemble an S-shaped saddle with mean curvatures close to zero. The studies reviewed above have demonstrated the complex nature of wettability and its effects on multiphase flow in porous media. While numerical simulation still faces challenges in accurately modeling wettability effects, microfluidic experiments are expected to continue to play a key role in the identification of new physics as well as the development and validation of new models.<sup>216</sup>

**3.3.5. Mineral precipitation and dissolution in microfluidics.** Mineral precipitation and dissolution are both important processes in the geological storage of CO<sub>2</sub> during all stages of the operation, including CO<sub>2</sub> injection, migration, trapping, and mineralization. On the one hand, precipitation is expected to lock CO<sub>2</sub> in a solid form, significantly alleviating concerns about the safety and security of CO<sub>2</sub> storage. On the other hand, precipitation of solids within the porous matrix tends to clog pore space, blocks CO<sub>2</sub> flow pathways, and hinders CO<sub>2</sub> injectivity and storage capacity. Similarly, while dissolution increases porous matrix connectivity and potentially enhances storage capacity, it simultaneously comprises reservoir rocks, potentially creating leakage pathways that threaten the safety and security of CO<sub>2</sub>. As noted by a few studies, mineral precipitation and dissolution often co-exist in the same system.<sup>217,218</sup> In the literature, the saturation index (SI) is widely used to define whether precipitation or dissolution is preferred. At SI = 0, the system is in equilibrium, indicating that precipitation is dynamically equal to dissolution. Mineral precipitation and dissolution

dominate at SI > 0 and SI < 0, respectively.<sup>217,219</sup> It is crucial to understand the fundamental process, pore-scale mechanisms, and the underlying physics associated with mineral precipitation and dissolution in porous media. Both mineral precipitation and dissolution in porous media can be due to physical (*e.g.*, drying of pore brine and dissolving of minerals by injected water), chemical (*e.g.*, the reaction between native minerals and injected CO<sub>2</sub>), and biological (*e.g.*, microbial-induced calcium carbonate precipitation or MICP) processes.<sup>192,219–221</sup> Each topic represents a vast area of study, which is virtually impossible for this review to cover comprehensively. Instead, a few representative microfluidics studies are discussed below.

Salt precipitation due to drying of brine is a physical process where the injected CO<sub>2</sub>, which is often low in the water component, extracts water molecules from the resident brine, causing the dissolved solutes to saturate, crystallize, separate from the brine, and eventually deposit in pore spaces. Kim *et al.* conducted a pore-scale experiment on salt precipitation using micromodels (Fig. 29).<sup>169</sup> Their results indicate that the porosity decreases by ~20% due to salt precipitation and that the salt precipitation front moves at a constant velocity, which is 2% of the superficial CO<sub>2</sub> flow speed. Two different types of salt formation were observed, with large bulk crystals on the same order of the pore size (20–50 μm) forming during the early stage of the drying process and polycrystalline aggregated structures ranging over broad length scales forming in late stages of the drying process and aggregating from the CO<sub>2</sub>–brine interface. Rufai & Crawshaw studied the evaporation of brine in 2.5D silicon micromodels and noted that the evaporation rate of brine is strongly impacted by salt precipitation, which was attributed to salt deposition at the matrix surface progressively impeding hydraulic connectivity to the evaporating surface.<sup>222</sup> Nooraiepour *et al.* investigated the precipitation of salt crystals in a microfluidic vessel under various reservoir-



**Fig. 29** [Left] Experiments of salt precipitation in the regular grid-like microfluidic network in PMMA.<sup>169</sup> [Right] Evolution of the calcite grains over time, subject to dissolution by HCl. The dissolution rate is shown to be strongly dependent on the pore flow and the presence of gas bubbles.<sup>184</sup>





relevant pressure and temperature conditions.<sup>223</sup> They showed that the magnitude, distribution, and precipitation patterns of salt accumulations are influenced by the phase of the injected CO<sub>2</sub> (*i.e.*, gaseous, liquid, or supercritical), presumably due to the differences in CO<sub>2</sub> density and the solubility of water in CO<sub>2</sub> for different CO<sub>2</sub> phases. Injection of gaseous CO<sub>2</sub> resulted in higher salt precipitation than liquid and supercritical CO<sub>2</sub>, and micrometer-sized halite crystals are thought to have the potential to partially or entirely clog fracture apertures. Ho & Tsai conducted experiments using a microfluidic device and noted the two types of crystal patterns, bulk crystal and polycrystalline aggregate, which are in agreement with the results by Kim *et al.*<sup>169,224</sup> He *et al.* studied the morphology and distribution of salt precipitation in porous structures and the effects of wettability and CO<sub>2</sub> flow rate using silica micromodels.<sup>225,226</sup> It was found that for hydrophilic and neutral porous surfaces, *ex situ* precipitation occurs, which completely blocks the throats and pore bodies and results in a significant reduction in permeability. However, for a hydrophobic porous surface, *in situ* precipitation occurs and occupies a much smaller pore volume, which only causes a mild reduction in permeability compared to the hydrophilic case at the same injection rate, highlighting the important role played by surface wettability in salt precipitation.

Mineral precipitation in porous media can be more complicated, as it further involves complex mixing and reactive transport behaviors, in addition to the challenges observed in drying. The relative importance of reaction, diffusive transfer, and advective transport is often characterized by two dimensionless parameters, the Péclet number (Pe) and the Damköhler number (Da), defined as  $Pe = Vl/D$  and  $Da_I = kl/V$ , respectively.<sup>227</sup> Here,  $V$  is the fluid velocity,  $l$  is the characteristic length scale (*e.g.*, pore diameter),  $D$  is the diffusion coefficient, and  $k$  is the reaction rate constant. Physically, Pe defines the ratio of advective to diffusive transport rates, and  $Da_I$  defines the ratio of the overall chemical reaction rate to the advective mass transport rate. When, however, the transport is dominated by diffusion, the second Damköhler number,  $Da_{II}$ , which compares the chemical reaction rate to the diffusive mass transfer rate, is defined as  $Da_{II} = Da_I \cdot Pe = kl^2/D$ .<sup>227</sup> The definitions of Pe and Da provide a convenient basis for characterizing reactive transport and predicting local reaction potentials in porous media.

Several studies investigated reactive transport and mineral precipitation processes in 2D micromodels. For example, Zhang *et al.* studied mineral precipitation in a 2D micromodel to observe how a porous structure affects the mixing processes between CaCl<sub>2</sub> and Na<sub>2</sub>CO<sub>3</sub>.<sup>217</sup> They considered four different saturation states and revealed that in certain conditions, pore spaces were occluded by CaCO<sub>3</sub> precipitates along the transverse mixing zone, thus substantially modifying pore connectivity and mixing patterns. Fanizza *et al.* evaluated the abiotic precipitation of uranium (U(VI)) in a microfluidic pore network and suggested

that uranium precipitation can block pores, alter fluid flow paths, and potentially limit mixing, thus slowing further precipitation.<sup>228</sup> Willingham *et al.* investigated the effects of porous media structure on mixing-controlled reactions using micromodel experiments.<sup>229</sup> Their results indicated that grain orientation significantly affects mixing and the extent of reaction at the pore scale and that the interfacial contact area between reactive species plumes is a controlling factor for mixing and the extent of chemical reaction.

Several other studies conducted very similar micromodel experiments focusing on various effects, such as the combined effects of evaporation and reaction, and the effects of Mg<sup>2+</sup> concentration on the morphology and polymorphs of CaCO<sub>3</sub> and calcite nucleation and growth.<sup>230</sup> For more information on the geochemical reactions in carbon sequestration, the readers are referred to several previous reviews.<sup>231–234</sup> More recently, several studies have been conducted specifically to support the development of pore-scale modeling.<sup>219,234–237</sup> Their experiments, where a barium chloride solution was injected into PDMS micromodels, leading to the dissolution of celestine and growth of barite, marked 4 stages of the process: the induction period, the crystal growth, the clogging, which prevented the mixing of the reactant solutions, and finally the dissolution of celestine crystals. Additionally, they showed that the evolution of the porosity–diffusivity relationship in response to precipitation reactions displays a behavior deviating from Archie's law. Although applying an extended power law improves the description of the evolving porosity–diffusivity, the model still neglects post-clogging features, calling for further studies. It is also worth noting that many microfluidic studies have been devoted to the understanding of microbially induced calcite precipitation (MICP) in porous media, which unfortunately cannot be covered in this review due to space constraints. The readers are referred to the following references for the recent advances in MICP research.<sup>192,221,238–240</sup>

Several other studies focused on dissolution to understand the pore-scale mechanisms that lead to pore structure changes due to dissolution.<sup>241</sup> In this scenario, it is often required that the micromodels be fabricated out of reactive/dissolvable materials, such as calcite, which is reactive with acids, and gypsum, which slowly dissolves in water.<sup>220,242,243</sup> Song *et al.* and Soulaire *et al.* studied the dissolution of calcite reservoirs by weak acids using calcite-based micromodels.<sup>181,244</sup> Their results illustrate highly flow-directed dissolution and associated positive feedback wherein acid preferentially invades high conductivity flow paths, resulting in higher dissolution rates, called “wormholing”. Song *et al.* later reported a new microscale mechanism that dictates the overall behavior of the reactive dissolution, where the reaction product, CO<sub>2</sub>, due to carbonate rock dissolution, forms a separate, protective phase that engulfs the carbonate rock grain and reduces further dissolution.<sup>181</sup> Agrawal *et al.* conducted experiments by flowing an acidic solution through a microscopic capillary



channel in a calcite crystal at two different flow rates.<sup>245</sup> Two key stages in pore shape evolution were observed: a transient phase and a quasi-steady-state phase. Their results also showed that the common assumption of constant pore shape in pore network modeling (*i.e.*, ignoring dissolution) may lead to an overestimation of pore conductance by up to 80%. Jiménez-Martínez *et al.* used a combination of pore-scale experiments and numerical simulation to study the effects of single and multiphase flow on dissolution and precipitation in a geomaterial-based micromodel.<sup>218</sup> They found that the presence of CO<sub>2</sub> bubbles significantly changes the flow dynamics and the resulting reaction patterns from a single-phase system, spatially homogenizing the rock dissolution. Additionally, bubbles redirect oversaturated fluid into low-velocity regions, thereby enhancing carbonate precipitation therein. Xu *et al.* studied reaction-infiltration instability, specifically the dissolution finger growth in radial geometries, using gypsum-based microfluidic devices.<sup>243</sup> In their experiments, pure water was injected to dissolve circular gypsum samples, which showed a few features consistent with theoretical and numerical predictions on the finger growth dynamics, such as screening and selection between the fingers. More recently, Rahman *et al.* conducted dissolution experiments using a 2D micromodel fabricated out of a pure calcite crystal and noted that the dissolution rate strongly depends on the local pore flow and the presence of gas bubbles that are either initially trapped or generated *in situ* (Fig. 29).<sup>184</sup> Combined, mineral dissolution in porous media is allured by the coupled process of reaction, solute transport, and pore flow, which is further complicated by instability and multiphase flow. Although microfluidics has enabled the discovery and characterization of numerous pore-scale mechanisms, direct modeling and predicting mineral dissolution in porous media is still challenging, even in single-phase flow. Additionally, mineral dissolution coupled with multiphase flow remains minimally explored, providing vast research opportunities for microfluidics to contribute.<sup>244</sup>

#### 4. Challenges and future directions in microfluidic-based carbon capture and storage

Even with significant advancements in microfluidic research for carbon capture and storage (CCS), there is still much to be done to bridge the gap between laboratory-scale innovations and real-world deployment. While microfluidic platforms have demonstrated excellent application benefits over conventional macroscale methods, several challenges remain that must be addressed to fully leverage their potential. Therefore, in this section, we have discussed key obstacles in microfluidic CCS research, which presents the scope of future research.

- Microfluidic platforms offer highly controlled environments and precise analysis of gas-liquid interactions

and reaction kinetics. However, their small scale presents a significant challenge for handling large CO<sub>2</sub> outputs in industrial applications. Some studies suggest that multi-channel parallelization and hybrid modeling, which integrates microfluidic insights with large-scale CCS simulations, could enable microfluidics to contribute to large-scale carbon capture. However, concerns remain regarding high-pressure drops and the practicality of large-scale fabrication, which could limit their feasibility. Addressing these challenges presents a promising research direction for advancing microfluidic-based CCS technologies.

- A literature review reveals that most microfluidic studies operate at ambient pressure and temperature, with limited variations to simulate real-world conditions. However, key CCS processes often occur under extreme conditions. For instance, CO<sub>2</sub> capture *via* chemical absorption, such as amine scrubbing, typically operates at around 70 °C, while CO<sub>2</sub> sequestration in deep geological formations takes place under high-pressure ( $\geq 100$  bar) and high-temperature ( $\geq 50$  °C) environments. This disparity between experimental and real-world conditions limits the direct applicability of microfluidic findings to large-scale implementation. Recent advancements in high-pressure, high-temperature microfluidic systems, including developing pressure-resistant materials and thermally stable microchannels, offer promising solutions. Implementing these innovations in CCS research could enable more accurate assessments of CO<sub>2</sub> dissolution, trapping mechanisms, and mineralization reactions, ensuring that microfluidic insights are more relevant for industrial applications.

- This review presents studies that have incorporated reservoir-mimicking surfaces in microfluidic CO<sub>2</sub> storage experiments. However, most research still relies on PDMS or glass-based microfluidic platforms, limiting the understanding of reactive transport behavior, long-term mineralization, and multiphase interactions such as capillary trapping. To address these gaps, more rigorous studies using reservoir-relevant materials are needed. Additionally, advancements in microfabrication techniques, such as 3D printing and laser lithography, could enable more accurate replication of fracture networks and permeability variations observed in natural reservoirs.

- Furthermore, our review highlights a significant gap in understanding mineral precipitation and its effects on pore clogging and flow diversion during CO<sub>2</sub> sequestration. While carbon capture mechanisms are well studied, the long-term stability of CO<sub>2</sub> storage remains poorly characterized. More comprehensive research incorporating reactive substrates and *in situ* imaging techniques could provide valuable insights into these processes, improving predictions of CO<sub>2</sub> retention and migration in geological formations.

- Most microfluidic devices used for CCS bear a quasi-2D nature, *i.e.*, the depth is usually much smaller than the other two dimensions, causing deviations from realistic 3D porous media. It is still a topic of scientific debate regarding how much of the knowledge that has been learned from those 2D



systems can be directly applicable to realistic 3D systems.<sup>242</sup> Therefore, more research is needed to ensure a faithful translation of the understanding obtained in 2D micromodels to realistic 3D processes.

- The unique advantages of microfluidic platforms discussed throughout this review, such as precise control of flow, tunable environmental conditions, high-resolution imaging, and material adaptability, can also be extended to a wide range of CO<sub>2</sub> utilization strategies that were not covered in this review. In particular, while CO<sub>2</sub> conversion was not a primary focus here, processes such as electrochemical reduction, photocatalytic transformation, and biological fixation stand to benefit significantly from microfluidic integration, enabling improved control over reaction conditions, deeper mechanistic understanding, and more efficient catalyst screening for compact and scalable CO<sub>2</sub> utilization systems.

- Finally, proper upscaling from microscales to large scales has remained a big challenge.<sup>243–247</sup> As described in section 2.3, recent progress in multi-channel and high-throughput microfluidic designs is a few examples toward large-scale implementation. However, there is a lack of extensive work in this field, and future work should aim to bridge this gap. Additionally, new approaches are needed to ensure the proper incorporation of pore-scale physics in large-scale modeling and predictions by developing novel constitutive relations.

## 5. Conclusion

Microfluidic technologies have emerged as a revolutionary tool for advancing CCS research, offering precise control over gas–liquid interactions, real-time *in situ* monitoring, and enhanced analytical capabilities while significantly reducing resource requirements. This review provides a comprehensive understanding of how microfluidic platforms have redefined experimental approaches in CCS research, replacing traditional macroscale setups and driving innovation in both carbon capture and storage. The first section focuses on microfluidic-based carbon capture, beginning with fundamental CO<sub>2</sub> capture mechanisms and detailing how microfluidic advancements address current challenges in solvent-based absorption, adsorption, and gas–liquid reaction kinetics. Additionally, mass transport theories are explored in relation to different microfluidic models, providing insights into their applicability across various capture processes supported by relevant experimental studies. The influence of flow patterns and microchannel geometries on CO<sub>2</sub> dissolution, reaction kinetics, and mass transport efficiency is examined, along with advanced measurement techniques such as micro-PIV, fluorescence imaging, and spectroscopy, which enhance gas–liquid interaction analysis. The second section, dedicated to microfluidic-based carbon storage research, explores recent advancements in various storage methods, particularly mineralization and geological sequestration. It covers pore-scale trapping mechanisms, mineralization kinetics, and permeability evolution,

highlighting the role of microfluidics in understanding multiphase flow dynamics and reactive transport. However, fabricating microfluidic devices for carbon storage presents greater challenges than carbon capture micromodels, as it requires precise surface modifications to mimic the complex heterogeneity of geological formations. This review addresses conventional and emerging fabrication techniques, discussing their applicability and limitations with contextual examples. Lastly, we have provided challenges like replicating geological conditions, scaling issues with microfluidics to handle large amounts of carbon, and a limited pressure and temperature range compared to micromodels. We also discussed some future scopes with possible solutions for these challenges.

## Data availability

The data analyzed in this review were obtained from publicly available sources, including PubMed, Web of Science, and various journal databases. All relevant data can be accessed through the provided citations within the review. Where data was not readily available, attempts were made to contact the original researchers for further information.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

AR and MK acknowledge the National Science Foundation for support of this work under award number 2207642. YL acknowledges the partial support by the NSF Career Award under award number 2144802 with Dr. Fangyu Cao as the program officer. Acknowledgement is also made by YL to the donors of the American Chemical Society Petroleum Research Fund for partial support of this research under Doctoral New Investigator Grant 62687-DNI9.

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