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Shales at all scales: Exploring coupled processes in mudrocks

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Shales at all scales: Exploring coupled processes in mudrocks

Abstract

Fine-grained sedimentary rocks – namely mudrocks, including their laminated fissile variety — shales — make up about two thirds of all sedimentary rocks in the Earth's crust and a quarter of the continental land mass. Organic-rich shales and mudstones are the source rocks and reservoirs for conventional and unconventional hydrocarbon resources. Mudrocks are relied upon as natural barriers for geological carbon storage and nuclear waste disposal. Consideration of mudrock multi-scale physics and multi-scale spatial and temporal behavior is vital to address emergent phenomena in shale formations perturbed by engineering activities. Unique physical characteristics of shales arise as a result of their layered and highly heterogeneous and anisotropic nature, low permeability fabric, compositional complexity, and nano-scale confined chemical environments. Barriers of lexicon among geoscientists and engineers impede the development and use of conceptual models for the coupled thermal-hydraulic-mechanical-chemical-biological (THMCB) processes in mudrock formations. This manuscript reviews the THMCB process couplings, resulting emergent behavior, and key modeling approaches. We identify future research priorities, in particular fundamental knowledge gaps in understanding the phase behavior under nano-scale confinement, coupled chemo-mechanical effects on fractures, the interplay between physical and chemical processes and their rates, and issues of non-linearity and heterogeneity. We develop recommendations for future research and integrating multi-disciplinary conceptual models for the coupled multi-scale multi-physics behavior of mudrocks. Consistent conceptual models across disciplines are essential for predicting emergent processes in the subsurface, such as self-focusing of flow, time-dependent deformation (creep), fracture network development, and wellbore stability.

Keywords
Mudrock
Shale
Coupled processes
Hydraulic fracturing
1. Introduction: shale architecture, heterogeneity, and coupled processes

1.1. Significance of mudrocks, and laminated mudrocks — shales

Sedimentary rock containing more than 50% (by weight or volume) of particles less than 62.5 μm in size is known variously as shale, siltstone, claystone, mudstone, and is cumulatively referred to as mudrocks (Folk, 1980, Milliken, 2014, Tucker, 2009). Some workers apply “shale” narrowly to refer to the visibly laminated, fissile variety of this sedimentary rock, but in this paper we apply this term as the overall name for the broad class of fine-grained layered sedimentary rocks, and, where appropriate, use it interchangeably with the term “mudrock” (Boggs, 2006). Shale constitutes around two-thirds of the sedimentary record of planet Earth (Garrels and Mackenzie, 1969, Blatt, 1982), and a quarter of the continental land mass (Jin et al., 2014). In some portions of sedimentary basins, distant from the principal axes of sediment transport, the abundance of mudrocks may approach 90% of the local sediment volume (Galloway et al., 1982). Shales are volumetrically dominant in both marine and terrigenous successions, and host significant portions of the fluid-rock interactions controlling fate and transport of elements in the upper crust (Milliken, 2004), and atmosphere – e.g., chemical weathering of shale has been shown to serve as a long-term global sink for carbon dioxide (CO₂) (Jin et al., 2014). Thus, shale properties are key controls on interactions of the atmosphere, hydrosphere, and sedimentary lithosphere in many contexts, and are crucial reservoirs for unconventional oil and gas production, top seals for conventional hydrocarbon traps and geological CO₂ storage (GCS), and isolation of nuclear and other high-level wastes.

In each case, shales are of interest both as barriers to fluid flow (seals) and as rock units that support flow of their contained fluids (reservoirs). Accelerated growth of information on Earth’s most abundant sedimentary rock highlights key gaps in our understanding of this rock type. The development of conceptual models for the coupled thermal-hydraulic-mechanical-chemical-biological (THMCB) processes in shale formations presents a major scientific challenge. In this paper we assess outstanding and fundamental issues in shale science that present obstacles to practical management of shales as seals, reservoirs, and source rocks. We develop
recommendations for future research and integrating multi-disciplinary data for models appropriate for multi-scale, multi-physics coupled processes in shale.

1.2. Key physical and chemical characteristics of shales

The physical and chemical properties of shales are controlled by their depositional environment, post-depositional diagenetic history, and engineering activities. In general terms, shales are commonly distinguished by their layered low-permeability fabric and composed of fine-grained minerals. One of the primary components of source rock shale is organic matter (e.g., kerogen). Porosity and pore sizes of shale are variable; however, nano-pores comprise a large fraction of the total porosity (Ross and Bustin, 2009, Loucks et al., 2009, Chalmers et al., 2012, Nelson, 2009).

Historically, shales have been strongly identified by their clay mineral content because clay minerals are a component that controls the physical behavior of shale at scales ranging from field and laboratory to the nano-scale (Potter et al., 2005). Although most shales contain at least a few volume percent of clay minerals, petrographic inspection by high-resolution methods shows that non-phylosilicate minerals such as quartz, feldspar, and calcite dominate in many shale lithologies and that organic components contribute additional complexity (Cook and Sherwood, 1991, Dean et al., 1985, Aplin and Macquaker, 2011). Compositional classification of shales has not, to date, benefited from a level of community consensus that is analogous to the widely applied classifications of sandstones and limestones (Milliken, 2014, Lazar et al., 2015, Bourg, 2015). It is clear however, that shales display a range of mineralogical compositions that encompass that of sandstones and limestones in addition to actual clay-mineral-rich rocks (Milliken, 2014).

Recent advances in understanding shale heterogeneity at many scales have been founded, in part, on a heightened appreciation of the nature of fundamental components (grains, cements, and grain replacements) as revealed by high resolution electron microbeam imaging (Wawak et al., 2013). Both observational (Lazar et al., 2015) and experimental (Schieber et al., 2007) approaches demonstrate that the physical sedimentology of fine-grained sediments entails a complex set of advective transport mechanisms as well as gravity settling, leading to distinctive textural and fabric characteristics at the bed scale that can be used to infer depositional conditions (Lazar et al., 2015). In this paper we use the term “texture” (rock microstructure) to refer to the combination of properties, including maturity, pore characteristics, fragment shape, roughness, composition, sorting, and diagenetic features — matrix and cement (Vernon, 2004).
Pores in shale manifest at a variety of sizes within kerogen (organic material), the inorganic mineral matrix, and fractures formed during natural processes or engineering activities (Akkutlu and Fathi, 2012, Saraji and Piri, 2015). The structures of pores reflect depositional processes and also the chemical and mechanical diagenetic processes associated with burial. In organic-rich shales, thermal maturation results in kerogen cracking and production of liquid hydrocarbons and gas, which also modifies the nature of porosity within organic components associated with these rocks. Pore waters comprise approximately 20% by volume of most sedimentary basins (Mondol et al., 2007, Kharaka and Hanor, 2014). The salinity of pore waters in petroleum reservoir rocks, including values reported for shale and tight reservoirs with in situ temperatures of ~ 20–150 °C and fluid pressures of ~ 100–1000 bar, varies widely from approximately 1000 mg L⁻¹ to over 400,000 mg L⁻¹ total dissolved solids (TDS) (Kharaka and Hanor, 2014). Detailed inorganic and organic chemical analyses, together with measurements of stable and radioactive water and solute isotopes have shown that the formation waters in sedimentary basins are dominantly of local meteoric or marine connate origin. However, bittern — residual evaporated seawater, geologically old meteoric water, and especially waters of mixed origin are important components in most sedimentary basins (Kharaka and Hanor, 2014). During diagenesis, the original waters of deposition evolve to Na–Cl, Na–Cl–CH₃COO⁻, or Na–Ca–Cl-type waters by a combination of several processes including (1) dissolution of evaporites, especially halite; (2) diffusion and advection, especially in and near salt domes; (3) reflux and incorporation of bitterns; (4) dissolution, precipitation and transformation of minerals other than evaporites; (5) interaction with clay minerals, principally mudrocks (and shales) behave as geologic membranes and have high ion exchange capacities; (6) interactions with organics, including petroleum and solid organic matter, as well as bacteria that can survive in sedimentary basins at temperatures of up to ~ 80 °C; and (7) mixing of different waters. The important processes responsible for the chemical evolution of water in each basin can be identified using chemical markers and isotopic tracers (Kharaka and Hanor, 2014, Hitchon et al., 1971, Hanor, 1994, Kharaka et al., 1987, Capo et al., 2014).

The significant compositional and textural heterogeneity that arises from the integrated effects of depositional, biologic, and diagenetic processes in shales is observed on the scale of nanometers (Macquaker et al., 2010, Schieber, 2004) to meters (Lazar et al., 2015) to kilometers. Methods to honor this multi-scale heterogeneity in models that predict the response of shales to natural and induced physical and chemical changes is one of the key challenges of shale science.
1.3. Why is THM CB process coupling important for shale?

The compositional and textural complexity of shale (Fig. 1) is manifested in non-linear progression of physical and chemical processes, and hard-to-predict response to natural and anthropogenic perturbations. The time-dependence and the interplay of chemical, mechanical, and transport processes have been directly observed, in particular in engineered systems. These coupled processes develop in highly nonlinear fashion and can range across length scales from nanometers to kilometers (Fig. 1), and across time scales from geological time scale to nano-seconds. The two-way process coupling in shale is presented in Table 1. There are numerous unknowns about how THCMB processes in shale are coupled and at what spatial and temporal scales.
Fig. 1. Structures, processes, and the relevant length scales for **shale** formations.

Table 1. Two-way coupling of thermal, hydraulic, mechanical, **chemical**, and **biological processes** relevant to natural and engineered **shale**.

<table>
<thead>
<tr>
<th>Process</th>
<th>Thermal</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Hydraulic</strong></td>
<td>Changes in fluid buoyancy and viscosity, change in fluid phase – evaporation and condensation. Thermal diffusion and fluid flow. Heat convection by moving fluid.</td>
</tr>
<tr>
<td><strong>Mechanical</strong></td>
<td>Conversion of mechanical energy into heat; thermal stress and thermal expansion, damage and deformation.</td>
</tr>
<tr>
<td><strong>Chemical</strong></td>
<td>Temperature control on chemical reaction rates and mineral stability fields; heat release from exothermal chemical reactions, involving inorganic and organic components.</td>
</tr>
<tr>
<td><strong>Biological</strong></td>
<td>Thermal effects on metabolism of microorganisms.</td>
</tr>
</tbody>
</table>

Diagenesis is one of the classic examples of process coupling in shale. A volumetrically important diagenetic reaction is the transformation of **mixed layer** clays (MLC),
containing smectite layers, to MLCs containing increasing proportions of illite (Hower and Mowatt, 1900). This diagenetic reaction is often accompanied by a volume change and is the source of significant dissolved silica that results in quartz cementation of proximal reservoir sands (Towe, 1962, Lynch et al., 1997), and possibly, the shales themselves (Thyberg et al., 2010). This reaction also produces significant volumes of water that can cause the development of geologic overpressures (Osborne and Swarbrick, 1997, Rask et al., 1997). The absolute value of volume change during the transformation of smectite to illite is not a constant, and depends on the chemical pathway. Osborne and Swarbrick (1999) calculated the volume change associated with 10 possible smectite to illite reaction pathways (Osborne and Swarbrick, 1999). The volume change ranges from an increase of 4.1% to a decrease of 8.4% depending on reactants and products (see also Swarbrick et al., 2001). A positive volume change and developed overpressure may cause a decrease of horizontal stress and induce fracturing in cases where the fluid pressures exceed the local minimum principal stress. Shale fracturing takes place anisotropically, resulting in irreversible changes in shale fabric (Gale et al., 2014), and increases, also anisotropically, the permeability by orders of magnitude. Shale architecture shaped by diagenesis controls the preferential flow across scales.

As indicated by Dusseault, 2004, shale is the only “common” rock type, where all four (Darcian, Fickian, Fourier and Ohmic) diffusion processes can co-exist as processes of first-order importance (Dusseault, 2004). Presence of the nano-channel and nano-pore structures in shale, as well as mixed wettability (e.g. water- and oil-wetting) networks control transport behavior (Javadpour, 2009), while reactive transport in turn affects the pore structures (Milliken and Curtis, 2016, Milliken and Day-Stirrat, 2013). Additionally, the combined effects of high salt content and large proportion of water bound at interfaces (water films) relative to free water, result in the overall low chemical potential (limited activity) of water, and chemical behavior differing from the bulk-phase behavior. In coupled processes, observed in shale formations, one of the key variables is volume change (Osborne and Swarbrick, 1999, Swarbrick et al., 2001). Microscopic processes of swelling and shrinking of clay minerals results in macroscopically observed expansion and contraction of shale beds. Volume changes control the state of stress, which leads to yield (shearing and fracturing) and changes in geometry of pore and fracture networks, and resulting changes in permeability and diffusivity (Milliken and Curtis, 2016). Understanding the THCMB process coupling and feedbacks necessitates quantifying the volume changes, governing processes and their rates and incorporating them into numerical models.
Chemical controls on fractures are common for various rock types, including shale. Most commonly, chemically-induced fracturing in shale is due to (1) cation-exchange reactions involving swelling clay minerals, which may result in positive or negative volume change (Dusseault, 2004), (2) mineral phase transformations, also resulting in volume change, and (3) chemical weakening of the silica- or metal-oxygen bonds due to the chemical attack (by water, proton or hydroxyl) on the accessible substrates – mineral grain surfaces and cements – contributing to fracture initiation and growth (Dove, 1995).

In some cases, biologic interactions in fine-grained systems can be profound (Macquaker et al., 2010, Schieber, 2004, Petsch et al., 2005). They are, however, limited by the “tightness” of the rocks: it has been shown that in shale with pore throat sizes < 0.2 μm, biological activity is limited (Fredrickson et al., 1997). However, in perturbed shale systems – for example, in gas wells in the Barnett shale – significant biological activity, manifested as biogenic sulfide production and microbi ally-induced corrosion, have been documented (Struchtemeyer et al., 2011). These introduced microorganisms in oil and gas fields can cause reservoir plugging, decline in the resource quality, and corrosion of metal-containing equipment (Struchtemeyer et al., 2011).

Coupled THCMB processes in engineered, or perturbed, systems are of particular interest in various applications ranging from wellbore stability, to performance of geological CO$_2$ storage reservoirs, and repositories for used nuclear fuel (UNF). In these applications, removal (oil and gas production, construction of wellbores or underground repository shafts) or emplacement (CO$_2$, nuclear waste packages) of materials interfere with the original either true or pseudo-steady-state, or, local equilibrium conditions of the system. For example, when CO$_2$ is injected into a geologic formation, some of the CO$_2$ dissolves in the pore brine and forms carbonic acid, lowering the brine pH, initiating geochemical re-equilibration through mineral dissolution and re-precipitation (Ilgen and Cygan, 2016, Kharaka et al., 2006). Other examples of process coupling during engineering activities are: (1) when a stainless steel UNF waste package is placed in a repository, gas generation from the canister corrosion can cause fracturing of the porous media, contributing to the positive feedback loop in the coupled fracture-transport (Olivella and Alonso, 2004); and (2) borehole instability, which has been observed for both smectite-rich shales with porosities in excess of 10%, as well as in low-porosity (less than 10%), highly fractured, quartz/illite-rich shale (Dusseault, 2004). These types of shale are common and are encountered in most deep drilling operations. The borehole instability is largely attributed to process coupling – when volume is added or
withdrawn as occurs with swelling clay minerals in smectite-rich shale, the state of stress around the borehole is modified, creating the potential for further yield, channeling and dilation (Choi et al., 2004). The dynamic volumetric dilation is manifested all the way to the processes occurring at the pore throats (Choi et al., 2004). Quantitative predictive models are limited and still under development due to the complexity of coupled processes and their manifestations over the large range of length and temporal scales. The contemporary challenge is to understand process couplings and emergent phenomena such as preferential flow path development, time-dependent deformation, fracture network development, and wellbore stability across spatial and temporal scales in natural as well as in perturbed systems. We suggest that the complexity of the THCMB process coupling in shale requires a systematic approach involving a broad range of disciplines. This systematic approach could create a foundation for a common language that could be used by the various disciplines that study or engineer shale. In the following sections we review the state-of-the-art methodologies used to address key subsets of coupled processes across the THCMB spectrum.

2. Methods

The selection of primary literature for this systematic review was based on whether the full range or any combination of the THCMB coupled processed is specifically addressed. Given the multi-scale multi-process nature of this review, we included both experimental, field, and theoretical studies. Only literature in English is included. We have identified the physical, chemical, mechanical, and geological properties of shale that distinguish it from other rock types. We reviewed the methodologies developed to address process coupling at various time and length scales, and synthesized the data to outline recommendations for future research on coupled processes and data integration.

3. Methodologies applied to coupled processes in shale

3.1. Post-depositional physical and chemical processes in shale (diagenesis)

Post-depositional changes cause further heterogeneity in fine-grained sediments and encompass a similar range of chemical and mechanical processes as are observed in the diagenesis of other sedimentary rocks. Compaction (Mondol et al., 2007, Schneider et al., 2011), cementation by carbonate, quartz, and other minerals (Milliken and Day-Stirrat, 2013), pressure solution (Evans, 1990), and grain replacement such
as albitization of feldspars (Milliken, 1994) are all observed in shales. The details of these processes in mudrocks are, however, still poorly understood compared to similar processes observed in sandstones and limestones.

The transformation of smectite layers in MLC with increasing temperature is the key diagenetic reaction in shale in sedimentary basins, including the northern Gulf of Mexico basin (Kharaka and Hanor, 2014, Lynch et al., 1997, Boles and Franks, 1979). Due to the large volumes of clay minerals, the water and solutes released and consumed by the MLC transformation are major factors shaping the hydrogeochemistry and petroleum resources of these basins (Lynch et al., 1997). Several incongruent reactions conserving aluminum (Al) or maintaining a constant total volume have been proposed for this transformation (Rask et al., 1997, Boles and Franks, 1979). The reaction of Eq. (1) (Kharaka and Hanor, 2014) conserving both Al and Mg, and precipitating chlorite, quartz, and illite is probably a closer approximation based on the composition of formation water observed in the northern Gulf of Mexico basin.

\[
\begin{align*}
10.8H^+ &+ 3.81K^+ \\
+1.69KNaCa2Mg4Fe4Al14Si36O100OH20 &+ 10H2O \leftrightarrow K5.5Mg2Fe1.5Al22Si35O100OH20 &+ 1.59Mg3Fe2AlSi3O10OH8 &+ 24.4SiO2s &+ 22.8H2O &+ 1.69Na^+ &+ 3.38Ca2^+ &+ 2.06Fe3^+ \\
\end{align*}
\]

Ferric iron (Fe\(^{3+}\)) in reaction (1) will be reduced by organic matter to Fe\(^{2+}\) and some may precipitate as pyrite or ankerite. The overall reaction consumes large amounts of potassium (K\(^{-}\)) and protons (H\(^{+}\)) and adds calcium (Ca\(^{2+}\)), sodium (Na\(^{+}\)) and some iron (Fe\(^{2+}\)) to the pore water.

Diagenetic illite and quartz are also the major pore-occluding cements in the petroleum reservoirs of North Sea (Bjørlykke et al., 1995) and Saudi Arabia (Franks and Zwingmann, 2010). Illite and quartz in these basins are formed from reactions of K-feldspar and kaolinite as depicted in reaction (2). Illitization of existing kaolinite is postulated to occur isochemically at a threshold temperature of ~ 140 °C (Bjørlykke et al., 1995).

\[
\begin{align*}
2KAlSi3O8 &+ Al2Si2O5OH4 \leftrightarrow KAl3Si3O10OH2 &+ 2SiO2s &+ H2O \\
\end{align*}
\]

As detailed in Kharaka et al. (2016), the salinity and chemical and isotopic compositions of pore waters from conventional and unconventional reservoirs from the same basin/sub-basin and pressure and temperature conditions appear comparable, indicating that diagenetic pathways experienced by fine-grained sediments are similar to the processes in coarser sediments (Haluszczak et al., 2013, Rowan et al., 2015).

Basin modeling and basin petroleum system modeling (BPSM) represent integrated approaches to recreate the diagenetic evolution of sedimentary basins, the latter focusing explicitly on hydrocarbon systems (Al-Hajeri et al., 2009). The traditional goal
of BPSM is to quantify the history of hydrocarbons in conventional reservoirs for exploration and resource extraction by reconstructing and/or incorporating the original hydrocarbon source location, generation, expulsion, migration pathways and preservation in hydrocarbon traps \cite{Al-Hajeri2009}. Original sediment deposition and subsequent diagenetic processes integrated by BPSM can include: sediment deposition, sediment types and structures, sedimentation rates, compaction, cementation, thermal history, dewatering and porosity evolution, kerogen maturation, hydrocarbon generation-migration-accumulation, multi-phase flow including relative permeability, capillary pressure, and assumptions on wettability, phase behavior of hydrocarbons and/or other fluids, pore pressure evolution, (effective) stress changes, and faulting. The BPSM approach can be applied spatially in 1, 2, or 3 dimensions. Major differences of BPSM from typical conventional reservoir modeling include large basin-scale domains (on the order of hundreds of kilometers with large grid block sizes in the numerical modeling) and geologic time scales (e.g., hundreds of millions of years) as opposed to meter or kilometer scale modeling over months to years \cite{Al-Hajeri2009}.

Application of BPSM to shale hydrocarbon plays is relatively new and still under development. Recent effort focuses on treating the shale play as both source and reservoir. \cite{Romero-Sarmiento2013} explain that “expelled [hydrocarbons] have been therefore used as a parameter to adjust for the assessment of conventional petroleum systems \cite{Romero-Sarmiento2013}. A thorough simulation of expulsion and retention mechanisms was not therefore necessary to define conventional oil and gas in place in the reservoirs.” \cite{Romero-Sarmiento2013} present approaches for basin-scale shale-play BPSM that include: source-rock kinetics, chemical transformations, and evolution of total organic carbon (TOC) and associated porosity, retention of hydrocarbon fluids (e.g., via sorption or as a free-phase in kerogen porosity), and assumptions of porosity evolution of kerogen-hosted pores versus mineral matrix pores (e.g. mineral-assemblage-hosted pores respond to stress state whereas organic pores respond mainly as a function of maturity i.e., thermal history) \cite{Rowan2015}. Recent and developing knowledge of the phase behavior of hydrocarbon fluids under nano-scale confinement, non-Darcy transport process (e.g., Knudsen transport), complex cementation/dissolution textures, and creep and other mechanical responses still need to be fully integrated in shale BPSM to enable better estimates of hydrocarbon resources in place and their exploitation potential.

3.2. Physical and chemical controls on fracturing in shale
As discussed above, the heterogeneous nature of organic-rich shales arises from textural complexity and variation in mineralogical composition. Fractures and coupled processes in shales cannot be properly understood without accounting for the role of these textural/compositional variations in controlling the mechanical properties, failure and the interactions among rock deformation, pore pressure, fluid flow and solid-liquid geochemistry (Gale et al., 2014).

The laminated nature and varying mineral/chemical composition of shale has been observed on multiple scales: from geologic observations at outcrops, in cores from organic-rich mudrock reservoirs, and from laboratory tests (with cm-resolution) that measure properties along the core length (Fig. 2, Fig. 3). Core measurements have shown that variation in mechanical properties of shale occur over intervals that range in scale from sub-centimeter to decimeters (Fig. 3). For example, unconfined strength in a single shale core can range from 69 MPa to 241 MPa (10,000 psi to 35,000 psi) among laminae, while Young Modulus has been observed to range from 7 GPa to 34 GPa (1 Mpsi to 5 Mpsi) on the same length scale (unpublished data). Equally large ranges of values of other mechanical properties are commonly observed (Suarez-Rivera, 2011).

Pervasive mm- to cm- to decimeter-scale layering with sharply contrasting properties is abundant (Fig. 2). Large scale layering, on the order of meters, is also common and is caused by the presence of carbonate benches, or other mineral concentrations, and intercalated shale units (Fig. 3). The sharply changing properties between thin rock units and the stacked rock architecture of organic-rich shales also results in the presence and distribution of weak interfaces with specific orientations (often bed parallel) which also affect fluid mobility and fracture propagation. Some of these interfaces were activated, parted, and/or mineralized, during basin development. Other interfaces are susceptible to slip when stresses and/or pore pressure change during drilling, hydraulic fracturing, production, as well as from associated regional changes in stress and deformation. For example, stresses and fluid pressure can vary widely in regions with a high density of wells where large volume of fluids are pumped in during hydraulic fracturing, and withdrawn during production. The presence of layering and weak interfaces results in preferential directions of deformation, failure, and fluid flow. Often these observed preferential directions do not align as predicted by simple homogeneous isotropic models.
Fig. 2. Intense layering with contrasting properties in shales. Outcrops photos (top), and scanning electron microscope (SEM) images (bottom).
Fig. 3. Elemental composition of shale measured by x-ray fluorescence (XRF) with centimeter-resolution. The presence and distribution of calcite-rich layers intermixed with mudstone units are observed over an 85 ft. (25 m) section. These challenges have been clear to the mining and civil engineering industries, and their workflows and models include orientations and modes of rock failure that are controlled primarily by the presence, distributions, and properties of fault sets and discontinuities (i.e., planes of weakness). The role of the intrinsic rock properties on failure is proposed to be of secondary importance. Block Theory (Goodman and Shi, 1985) is an example of a methodology developed to understand and predict rock failure and excavation stability based on a detailed mapping of the orientation and distribution of planes of weakness in the regional rock system. Admittedly the mining and civil engineering industries focus predominantly on “stronger” rocks, where the intrinsic rock strength is larger than the shear strength of faults or other planes of weakness; and the confining stresses may be lower because of the shallower nature of their applications. Nevertheless, in contrast, the method and models the oil industry uses for rock deformation and failure (e.g., wellbore stability, hydraulic fracturing, depletion-induced compaction, and others), are primarily dependent on the intrinsic, homogenized, rock properties. These are typically obtained or predicted at well-log resolution (averaged over 2 ft. (60 cm), Table 2) and, as a consequence, this minimizes or removes the
representation of existing thin layering and interfaces – e.g. micron-scale fractures (Fig. 4). Such a method may be appropriate and successful for conventional reservoirs, but is limiting for unconventional, over pressured, heterogeneous, anisotropic reservoirs. Regarding hydraulic fracturing, for example, it is now accepted that the layered nature of mudrocks and the ubiquitous presence of planes of weakness in them (often bed parallel) give rise to complex hydraulic fracturing geometries, the presence of multiple branches of fracture propagation, fish-bone structures of fracturing and leak off, step overs, and other geometric effects that do not occur in homogeneous isotropic materials and cannot be explained using homogeneous models (Fig. 5) (Suarez-Rivera et al., 2013).

Table 2. Types of data available for studies on shale, their spatial and temporal resolution, and their process couplings.

<table>
<thead>
<tr>
<th>Data type</th>
<th>Spatial resolution or size of testable sample</th>
<th>Temporal resolution</th>
<th>Couples with‘ or strongly affects</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Texture (grain size, shape, orientation of individual grain, overall sorting)</td>
<td>Mudrock has &gt; 50% of particles &lt; 62.5 μm; fine mud (clay and very fine silt) is &lt; 8 μm</td>
<td>Time-scale of depositional setting (see Bedding); up to geological time for mechanical diagenesis and recrystallization</td>
<td>Porosity, permeability, capillary pressure characteristics; strength, Young's modulus</td>
<td>Informs sediment provenance, water column energy level, and geologic controls on rock properties including porosity and permeability (Lazar et al., 2015)</td>
</tr>
<tr>
<td>Pore network imaging, pore types and distribution, porosity (FIB-SEM, USAN/SANS, core plug measurements) NMR</td>
<td>nm to 100s of μm (imaging); 0.5 μm to 100s of μm (EDS); 1 nm to 10s of μm (USANS/SANS) cm's (core plug)</td>
<td>Time-scale of transport in pore networks: up to seconds</td>
<td>Porosity, permeability, chemical composition /wettability</td>
<td>Direct imaging and indirect measurements of porosity and connectivity—useful for modeling of pore-scale transport (including wettability) and potential mechanical behavior</td>
</tr>
<tr>
<td>Bedding</td>
<td>Lamina: fraction of mm to mms; Laminaset: mm to cms; Beds: typically mms to 10s of cms (do not have minimum or maximum absolute thickness); laterally meters to kms</td>
<td>Laminae: forms in seconds to one or more years Beds: minutes to “many moments of geological time” (Campbell, 1967)</td>
<td>“Larger”-scale flow or mechanical units; natural fracture spacing may correlate with larger-scale stratal units</td>
<td>Relevant for interpreting: sediment input, accumulation; energy of deposition; degree of bioturbation (Lazar et al., 2015); curved, wavy planar (parallel and nonparallel) affect heterogeneity and other transport/mechanical properties</td>
</tr>
<tr>
<td>Solid composition</td>
<td>Can vary from</td>
<td>Time-scale controlling</td>
<td>Brine chemistry;</td>
<td>Reflects primary</td>
</tr>
<tr>
<td>Data type</td>
<td>Spatial resolution or size of testable sample</td>
<td>Temporal resolution</td>
<td>Couples with or strongly affects</td>
<td>Comments</td>
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<td>--------------------------------------------------------------------------</td>
<td>---------------------------------------------</td>
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<td>----------------------------------------------------------------------------------------------------</td>
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</tr>
<tr>
<td>(mineralogy, chemical composition, kerogen type; total organic carbon)</td>
<td>individual particles (or cements) to large-scale systematic variation up the bedset or larger-scale stratal units (such as parasequences)</td>
<td>composition from seconds (depositional setting) to oil and gas extraction and CO₂ storage (10s to 1000s of years) to geological time (diagenesis)</td>
<td>pore-lining phases and geometry of pore networks; (mixed) wettability</td>
<td>deposition and diagenetic conditions; pore-lining phases affect wettability and chemical reactivity</td>
</tr>
<tr>
<td>Fluid chemistry (brine, fracturing fluids, hydrocarbon liquids and gases, possibly injected CO₂)</td>
<td>Single pores to regional scale (10s to 100s of km)</td>
<td>Minutes/hours/weeks/years for engineered (injection); geologic time scales for natural systems</td>
<td>Mineralogy; permeability, porosity (coupled through reactive transport) and nano-scale confinement effects and phase behavior; sub-critical fracture growth</td>
<td>Fluid chemistry can strongly couple with many processes (transport, mechanical behavior, heat flow, and microbiology)</td>
</tr>
<tr>
<td>Absolute permeability (and relative permeability)</td>
<td>10s of microns (FIB-SEM and modeling) to cm (core plug)</td>
<td>Permeability evolution of natural systems: geological time scale for diagenesis; permeability evolution can be over hours, weeks, days, years for engineered systems</td>
<td>Porosity; pore pressure distribution; drained versus undrained behavior; texture, chemistry</td>
<td>Continuum concept; REV may vary for shales and is not very well documented yet and may vary for different types of pore structure (e.g., matrix vs fractures) Relative permeability is not routinely measured in mudrocks</td>
</tr>
<tr>
<td>Capillary pressure curves and pore-throat size distributions</td>
<td>cm (core plug); possibility down to 100’s of microns if using FIB-SEM data and a modeling method (e.g., LB)</td>
<td>Probably up to seconds of transport (in FIB-SEM models)</td>
<td>Porosity, permeability, textural analysis; activity of water or fluids in nano-scale confined pore networks</td>
<td>Capillary imbibition strongly coupled with texture</td>
</tr>
<tr>
<td>Effective diffusion coefficient</td>
<td>cm (core plug); possibility down to 100s of microns if using FIB-SEM data</td>
<td>Geologic time scales for natural transport processes; large surface area of induced fractures may reduce time scales to that of reservoir production</td>
<td>Texture, composition, fluid compositions, permeability</td>
<td>Diffusion processes may range from Ordinary Fickian to Knudsen diffusion</td>
</tr>
<tr>
<td>Sorption/adsorption isotherms</td>
<td>mm to cm (core plug or crushed samples)</td>
<td>Seconds; may depend on reservoir pressure and thus time scales of field operations (years)</td>
<td>Texture, composition, fluid compositions, permeability</td>
<td>Consider sorption/adsorption for fine-grained minerals and organic material</td>
</tr>
<tr>
<td>Data type</td>
<td>Spatial resolution or size of testable sample</td>
<td>Temporal resolution</td>
<td>Couples with or strongly affects</td>
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<tr>
<td>Geomechanics and constitutive models; Young Modulus and Poisson ratio (isotropic); Transversely isotropic (five parameters); critical state mechanics; general plasticity; failure models and parameters (e.g., Mohr-Coulomb)</td>
<td>From micron (nanoindentation) to cm (core plug); core plugs may need to be taken at different angles for estimates of transversely or fully anisotropic parameters</td>
<td>Seconds, hours, weeks (engineered) to geological timescales (natural)</td>
<td>Porosity, pore network properties, texture, composition, diageneric textures</td>
<td>Some parameters may be dynamically estimated from seismic wave velocities; sophistication of geomechanical properties estimated can vary greatly</td>
</tr>
<tr>
<td>Unconfined compressive strength</td>
<td>micron to mm to cm (micron indenter on FIB-ed columns; calibrated scratch test; core plugs)</td>
<td>Seconds, hours, weeks (engineered) to geological timescales (natural)</td>
<td>Porosity, pore network properties, texture, composition, fluid composition contacting fracture process zone, chemistry-related</td>
<td>In situ stress state, including magnitudes, directions, and pore pressure distribution should be taken into account in predicting subsurface behavior</td>
</tr>
<tr>
<td>Fracture toughness</td>
<td>From micron (nanoindentation) to cm (double-torsion)</td>
<td>Seconds, hours, weeks (engineered) to geological timescales (natural)</td>
<td>Porosity, pore network properties, texture, composition, fluid composition contacting fracture process zone</td>
<td>In situ stress state and fluid composition should be taken into account</td>
</tr>
<tr>
<td>Tracer logs or spinner surveys; distributed fiber optic sensing</td>
<td>10's of cm, up to length of completion of wellbore</td>
<td>Hours, weeks, years (for permanently installed fiber optic arrays)</td>
<td>Porosity, permeability, fracture network characteristics</td>
<td>Tracer logging and spinner surveys may reflect aspects of completions and the reservoir itself</td>
</tr>
<tr>
<td>Production decline analysis; pressure or rate transient analysis; yields estimates of a variety of reservoir parameters</td>
<td>Length scale of fracture spacing; provides information on the entire length of completion of wellbore</td>
<td>Months to decades; the progression of flow regimes in shale reservoirs may take decades to develop</td>
<td>Integrates permeability (potentially temporally varying), porosity, multi-phase flow; may include double porosity</td>
<td>Attempts to invert for reservoir parameters and reverses or oil/gas in place, and forecasts production</td>
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<tr>
<td>Wireline measurements (neutron porosity, gamma ray, well resistivity, NMR, in situ fluid saturations, etc.)</td>
<td>Resolution 1–2 ft (30–60 cm)</td>
<td>Wireline measurements on order or hours</td>
<td>Porosity, permeability, and fluid composition</td>
<td>Important tools for measurements of reservoir properties over length of completed zones</td>
</tr>
<tr>
<td>Microseismic</td>
<td>10s of meters</td>
<td>Probably hours for measurements</td>
<td>Porosity, permeability,</td>
<td>Important for mapping fracture complexity at</td>
</tr>
<tr>
<td>Data type</td>
<td>Spatial resolution or size of testable sample</td>
<td>Temporal resolution</td>
<td>Couples with or strongly affects</td>
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<td>geomechanical properties</td>
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Notes: * identifies where methodologies for integrating multi-disciplinary data are well established.

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Fig. 4. Healed bedding plane fracture in Eau Claire Formation, a major seal for the Mt. Simon Sandstone in the mid-continent of the U.S.
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Fig. 5. (a) Borehole breakouts result along preferential direction under a uniform stress field, because the presence of planes of weakness in the rock; (b) Hydraulic fracture grows by overcoming one layer at a time, despite the imposed uniform stress field; (c and d) Thin mineralized interfaces and weak bedding control the propagation of fractures during fracture toughness experiments; (e) Fish-bone structures of fracture.
propagation and fluid leak off are common on laboratory hydraulic fracturing experiments on laminated shales. Elastic or seismic wave characterization is often used to determine the mechanical properties of rocks in the subsurface from measurements of wave attenuation and velocity. However, interpretation of geophysical signals is complicated for shale because of the potential for competing sources of anisotropy: textural versus structural. Textural anisotropy arises from laminae, thin parallel layers of alternating composition and moduli (e.g. carbonaceous, silty, dolomitic, clayey, or lithoclastic) that range in scale from micrometers to centimeters, to decimeters (Fig. 1, Fig. 2, Fig. 3), and with orientations that depend on the original depositional environment and post-depositional tectonic processes. Structural anisotropy arises from micro-cracks, fractures, joints, and the aforementioned interfaces that range in scale from micrometers to meters, are sensitive to stress, and have orientations and spacings that may or may not align with the textural features based on the diagenetic and tectonic history or any activities that perturb the subsurface system. These competing sources of anisotropy can mask either the presence of fractures and/or the matrix/fabric anisotropy depending on the state of stress (i.e., stress magnitude and orientation).

The effect of competing anisotropy has been clearly demonstrated in the laboratory measurements on manufactured anisotropic medium. The acoustic wavefronts (Fig. 6) were propagated through this anisotropic medium with fractures perpendicular to subwavelength layering (Shao, 2015). The unique symmetry axis for the fracture set is vertical while that for the matrix is horizontal. In Fig. 6, the spatial distribution of energy is shown at a fixed arrival time. At low or high stress, the measured anisotropy is controlled by either the fracture orientation or the matrix texture, respectively. However, at an intermediate stress, the fractured anisotropic medium appears isotropic as indicated by the circular shape of the wavefronts, i.e. the energy spreads out nearly uniformly in all directions. Interpreting rheological properties from velocities measured under the condition of the intermediate stress state would, incorrectly, yield isotropic moduli.
matrix-dominated

Increasing Stress

Fracture-dominated
Acoustic wavefronts propagated through garolite (fiberglass-epoxy laminate) sample with a set of parallel fractures. Stress was applied perpendicular to the fractures (solid black lines) or parallel to the layering (dashed gray lines). Thus a question arises whether competing sources of anisotropy in fractured shale can be delineated using seismic or elastic wave techniques. Fractures and other mechanical discontinuities often give rise to converted-, guided- and scattered modes that could be used to interpret fracture and matrix properties. For example, the velocity of waves guided between or along fractures has shown promise as potential tool to extract both fracture and matrix properties from elastic waves. These guided modes depend on the orientation of a fracture relative to layering, the matrix and fracture mechanical properties, layer/fracture spacing and signal frequency (Shao and Pyrak-Nolte, 2013, Shao et al., 2015). Heterogeneity leads to scaling complexities that are the undeniable challenge for evaluating and modeling failure, fracture, and fluid mobility behavior of organic-rich mudrocks, for understanding their coupled effects, and for defining the type of measurements that will be relevant. Additional research is needed to assess the contributions to the scattered wave field not only from fractures but also from fracture intersections, fracture sets, stress gradients and fluids in order to characterize dynamically evolving fractured shale systems. This future research will determine whether competing sources of anisotropy are separable, if the dominant symmetry axis depends on stress and fluid conditions, and if the dominant symmetry axis also indicates fluid flow anisotropy.

The ability to detect and monitor the dynamic evolution of fractured shale systems using geophysical methods requires a link between a remotely-measured geophysical response and a characteristic property (or properties) of a fracture. For over two decades, several researchers have demonstrated that fracture-specific stiffness can be estimated from seismic wave attenuation and velocity (Choi et al., 2014, Far et al., 2014, Hobday and Worthington, 2012, Verdon and Wüstefeld, 2013, Lubbe and Worthington, 2006, Lubbe et al., 2008, Majer et al., 1988, Pyrak-Nolte et al., 1990a, Pyrak-Nolte et al., 1990b, Sayers et al., 2009). The dependence of specific stiffness on the spatial and probability distribution of regions of contact between two fracture surfaces creates an implicit link to the hydraulic properties of a fracture through the fracture geometry (Pyrak-Nolte and Morris, 2000, Cook, 1992). Recently, Petrovitch et al., 2013, Petrovitch, 2013b determined the existence of a scaling relationship between fracture specific stiffness and fluid flow for single fractures (Petrovitch et al.,...
The numerical flow-stiffness data, simulated at multiple length scales, collapsed to a single scaling function because fracture specific stiffness captures the deformed fracture void topology that includes both changes in contact area and aperture caused by stress as well as by transport-dominated chemical erosion (Pyrak-Nolte and Nolte, 2016). The resulting hydro-mechanical scaling function potentially provides a link between fluid flow and the seismic response of a fracture, because fracture-specific stiffness affects seismic wave attenuation and velocity. However, several outstanding questions related to the deformation of fractures in shale and the scattered wave field must be addressed before extending these concepts from single fractures to fracture networks in subsurface shale. For example: does a viscoelastic matrix affect deformation of fracture void geometry in a manner that differs from pure elastic conditions? Will the flow-stiffness relationship hold for partially-mineralized fractures that are often found in shale? Can seismic data differentiate or unravel chemical, fluid and stress alteration of fractures? Can scattered wave fields delineate the effects of stress gradients that lead to non-uniform fracture topology and fluid distributions that mask or promote additional scattering? And how do proppants or geochemically induced reaction halos affect fracture deformation and in turn fracture specific stiffness?

The importance of the chemical effects on subsurface fracture behavior, or chemo-mechanical coupling, has been recognized for several decades. Chemically assisted subcritical fracture growth, also referred to as stress corrosion cracking, is proposed to control the time and deformation-rate-dependent failure of rocks (e.g., (Anderson and Grew, 1977, Swanson, 1984, Holder et al., 2001) and references therein). Resistance to subcritical fracturing depends on microstructural heterogeneities: micro-fractures, grain boundary cohesion and orientation, and mismatches in elastic properties between phases. Development of large fracture populations and fracture architecture is controlled by chemical processes in the micro-scale near-tip fracture regions (Hu and Hueckel, 2013, Schultz, 2000, Gale et al., 2004). In aqueous fluids, mineral reactions have been shown to play a key role in subcritical fracture development at both bulk and microscopic scales. A technique traditionally used for assessing reaction kinetic effects on subcritical fracture propagation is double-torsion geomechanical testing, which allows measuring fracture propagation rate and subcritical index in shale under controlled fluid composition and temperature (Holder et al., 2001). A novel approach for interrogating coupled chemical-mechanical fracture processes is X-ray computed tomography (CT) for imaging fractures while controlling stress conditions using a triaxial cell. This
technique enables characterization of fracture formation and permeability changes at in situ temperature, pressure and stress conditions (Carey et al., 2015). The CT data can be used as model input (Carey et al., 2015, Lei et al., 2014) enabling prediction of fracture behavior as a function of fluid pressure and changes of in situ stress. The accuracy of these models depends on the proper inclusion of interfaces and fluid flow processes.

Further development of coupled models is needed in order to predict coupled fracture, multi-phase flow, and multi-component reactive transport in the subsurface, and to capture the effect of geochemical reactions on fracture permeability. These new models require that the solid solvers are coupled to the multi-phase reactive flow and reactive transport codes. One difficulty in modeling chemical-mechanical processes in shale, is that a model must incorporate changes in volume that correspond to changes in chemical concentration which in turn are coupled to the diffusion processes. In addition, laboratory data to support or verify such models is sparse.

3.3. Fluid flow in multi-porosity systems

Fluid flow in shales takes place in a variety of void or pore structures. Salient features that influence flow and transport include: the occurrence of discrete pore networks within kerogen and inorganic components (e.g., clay and silt), that have different wettabilities and characteristic sizes (Akkutlu et al., 2015, Heath et al., 2011); a range of pore sizes that result in various transport modes including viscous flow, Knudsen flow, transition flow (viscous and Knudsen flow), and diffusive fluxes encompassing ordinary Fickian diffusion including effect of mass (Graham's Law), and adsorption-desorption processes on the surface of pores of the minerals or organic matter (Gensterblum et al., 2015); a variety of pore body and throat shapes, sizes, and topologies that can strongly affect multi-phase flow processes such as imbibition-drainage (Fig. 7), especially relevant for liquid and gas hydrocarbon systems (Sakhaee-Pour and Bryant, 2012); a vast range of natural or induced fracture types that reflect paleo and recent fluid flow, and precipitation-dissolution that may have modified the porous matrix or fracture permeability (Fig. 4). Different flow mechanisms can occur for different scales of the pore structures (Mehmani et al., 2013). The orientation of fractures (or faults) relative to the current regional and local stress regime can affect whether the fractures-faults are critically-stressed and “hydraulically active.”
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Fig. 7. (a) Dual beam focused ion beam (FIB)/SEM 3-D volume, (b) 3-D rendering of kerogen and (c) 3-D rendering of the pores in this volume of the Pt. Pleasant formation, Wood Co. West Virginia, 9503 ft. depth. Re-printed from Arthur and Cole (2014).

Additional complexity is introduced by the large variability of the organic matter associated with shale (Milliken and Curtis, 2016, Eliyahu et al., 2015). For example, laboratory measurements of hydrocarbon fluid extraction and composition indicate measurable differences in hydrocarbon compositions and in the partition between light components, intermediate molecular weight components and heavy components, within short intervals in the same formations and at the same levels of thermal maturity (Freeman et al., 2011). This is most likely due to the heterogeneous distribution of macerals in the system and also to the effect of rock texture and composition on moderating the thermal processes of hydrocarbon generation and cracking. Thus fluid separation (oil and water) in relation to surface energies (mineral and organic surfaces), and changes in hydrocarbon composition in relation to changes in rock texture and composition are factors that also contribute to the heterogeneous distribution of pore pressures, and control the flow in the system.

The large variety of pore types (Milliken and Curtis, 2016), many of which can occur in a shale formation at a range of scales, makes the quantification and/or prediction of flow and transport difficult. Current techniques for understanding the transport in the multi-porosity shale system include a combination of methods to characterize the pores and model flow and transport. Several studies use fully three-dimensional (3D) imaging or serial sectioning to characterize the pores and then reconstruct digital models of the pore geometry and connectivity of solid components. Primary methods (Table 2) include dual beam focused ion beam–scanning electron microscopy (FIB-SEM) (Fig. 7), X-ray CT and micro- to nano-tomography, and neutron tomography, with resolutions from ~ 1 nm to 10s of microns or higher for medical X-ray CT systems (Saraji and Piri, 2015). A major concern of reconstructions for flow modeling is whether the small volumes representative of the digital reconstructions are representative of bulk volume properties of the shale. These volumes can range in size from ~ 653 μm³ (Saraji and Piri, 2015) to ~ 10⁶ μm³ (Trebotich and Graves, 2015). Several studies address issues of representative elementary volume (REV) to determine the length scale at which shale properties become statistically stable and suited for modeling by continuum methods (Saraji and Piri, 2015, Yoon and Dewers, 2013, Chen et al., 2013, Gelb et al., 2011). It appears porosity REVs can be obtained for some shales from FIB-SEM 3D reconstructions: e.g., members of the Bakken, (Saraji and Piri, 2015); however, it is not
possible to obtain an REV for permeability for the same study because the FIB-SEM volumes are too small. Rui and Akkutlu (2016), have recently presented a scaling up approach predicting kerogen REV using a nanopore-network modeling, and predicted an REV of 5 μm³ (Fig. 8). Establishing the ranges of sizes for REVs in kerogen, clay mineral-rich components, and mixtures of these with larger grains has not yet been fully investigated; however, and limited data is available to determine if REVs for certain shale lithotypes can be universally applicable to other shales (or even regions within the same shale). The unique depositional and diagenetic histories may make it difficult to determine which REV length scales are appropriate for different shale lithofacies. Concepts of representative elementary time (RET) also come into play for the given process under study, as the different pore types have different characteristic length scales that in turn affect the characteristic time scales of processes (Milliken and Curtis, 2016). Other studies address transport through laboratory measurements of permeability, porosity (connected and disconnected), capillary breakthrough pressure, wettability, and fracture hydraulic aperture and conductance (Schneider et al., 2011, Gensterblum et al., 2015). Laboratory petrophysical properties suffer from the limitations due to retrieval and handling of core, which may induce microcracks and fractures that are not present in the subsurface. Additionally, the variety of techniques can give disparate results, especially when compared to digital pore network reconstructions.

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Fig. 8. Left: Three-dimensional pore network model consisting of nano-capillaries shown in two-dimensions for nanopore Pn. The model has been developed and used to investigate kerogen REV by Rui and Akkutlu (2016). Right: estimated permeability of the kerogen nanopore network as a function of the network volume for changing coordination number, Z. Methane transport is considered (including convection, diffusion and adsorption mechanisms) at average pore pressure of 2500 psi, and temperature 80 °C (353 K).

Modeling approaches for pore-scale flow and transport in shale range from theoretical to empirical—either idealized pore body and throat sizes and connectivity are assumed or digital reconstructions from direct imaging are used (Sakhaee-Pour and Bryant, 2012, Trebotich and Graves, 2015). Theoretical models typically capture laboratory-based behaviors such as drainage of a non-wetting phase and absolute permeability using network models such as bundle-of-tubes model; regular-lattice models; acyclic models; and multi-scale-multi-physics networks (Sakhaee-Pour and Bryant, 2012, Mehmami et al., 2013, Purcell, 1949, Washburn, 1921). Theoretical models have revealed that certain pore structures can capture multi-phase transport behaviors such as the non-plateau drainage in shale (Sakhaee-Pour and Bryant, 2012) and allow for estimation of fitting parameters that may be helpful for classification of different shales. Empirical modeling using digital reconstructions from shale samples involve realistic pore structures, geometries, and topologies. Approaches include level set methods, volume of fluid methods, gradient based computational fluid dynamics, and Lattice Boltzmann (LB) methods. LB methods in particular show much potential for shale as they capture pore geometries and can represent large differences in densities of two fluids occupying the pore space (Lei et al., 2014, Ho and Striolo, 2015). They also unfortunately suffer from the small REV problem discussed previously.

Wasaki and Akkutlu proposed a matrix permeability model for organic-rich shale and considered its coupling to a fracture during shale gas/oil production (Wasaki and Akkutlu, 2015). They argued that the shale permeability is not a petrophysical quantity in the classical sense reflecting the fluid transmitting ability of the pore network but instead it is an overall mass transfer coefficient that needs to be carefully tuned to honor the total mass flux of fluids draining into the fractures. They presented a conceptual transport model for the shale matrix with dual-porosity and single-permeability delineating the transport mechanisms at multiple-scales: (i) adsorption and diffusion mainly in the kerogen pores; and (ii) diffusion and convection in micro-cracks and other slit-shape inorganic pores.
Here, $\mathbf{I}$ is second order identity matrix. \textbf{Wasaki and Akkutlu (2015)} recognized that the observed anisotropy also exists in the flow field and inherently belongs to micro-cracks contribution ($\kappa_m$) which is sensitive to effective stress (Fig. 9). The authors gave analytical expressions for the micro-crack permeability components, $\{\kappa_m\}_{11}$, $\{\kappa_m\}_{12}$, and $\{\kappa_m\}_{22}$. Accordingly, if horizontal flow is considered, $\{\kappa_m\}_{11}$ and $\{\kappa_m\}_{12}$ are the elements that will affect the flow, and $\{\kappa_m\}_{22}$ can be important if pressure gradient in vertical direction exists. Note that only the natural gas flow has it non-Darcian effects, as shown in Eq. (3b), due to the presence of molecular transport mechanisms taking place in the kerogen. These mechanisms are pore diffusion and cluster diffusion of the adsorbed molecules and here represented by the isotropic diffusion coefficients, $D$, and $D_s$, respectively, following terminology presented by \textbf{Akkutlu and Fathi (2012)}.

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Fig. 9. Left: Schematic illustrating the uniform distribution of micro-cracks and their orientation angle $\theta$ in a cross section of organic-rich shale in vertical direction. The organic-rich shale is shown as gray background. Directions of vertical and horizontal minimum stresses are also shown. Right: Permeability tensor $\kappa_m$ elements for various microcrack orientation angle ($\theta$) with $\kappa_{rr} = 0$.

At the production scale, another type of modeling – pressure transient or rate transient analysis (PTA/RTA) – involves inversion of pressure and/or flow rate measurements at a production wellhead to estimate reservoir transport parameters and properties.
PTA/RTA methods can represent the pore space as dual, triple, or multi-continua in order to attempt to capture the multiple pore types and their interchange of fluids (Clarkson, 2013, Ezulike and Dehghanpour, 2016, Kuhlman et al., 2015). Reactive transport modeling (and other geochemical modeling) can be used for mm- to reservoir scale systems and is commonly applied to addresses mineral dissolution/precipitation, adsorption/desorption, and homogeneous chemical reactions (Bethke, 2008). Fluid flow in shale is often coupled to other physical and chemical processes, resulting in coupled phenomena. These can include mechanical behavior where the shale behaves as a partially-drained or undrained medium, thus affecting the mechanical constitutive behavior. Distinct pore networks in kerogen or the inorganic clay-silt components can exhibit different stiffness/compliance for different deviatoric stress and conditions of the pore fluids e.g., Biot and Skempton coefficients (Suarez-Rivera and Fjaer, 2013). Thus, the permeability of kerogen may be affected by flow and compaction or other mechanical processes at time-scales different from the pore networks of the inorganic components, which in turn affects the overall flow regimes in shale impacted by natural or engineering activities. The texture of a shale can strongly affect flow due to matric (combined effects of capillarity and adsorptive forces) and osmotic potentials—nano-scale confinement affects the phase behavior e.g., mean free path of a gas molecule and hence transport behaviors. Texture can cause flow mechanisms to be species-dependent as the mean free path may vary due to pressure changes (Knudsen number can vary during production of a reservoir). The ability to flow methane gas in a system with porosity dominated by kerogen-matrix is also a function of the percolation threshold of the connected kerogen volume in any given organic-rich shale system.

3.4. Nano-scale confinement, activity of water, and pore-scale coupled processes in shale

Due to fine-grained shale fabric and presence of nano-pores, fluids (gas, oil, brine) in shales are often present as nano-scale thin films, and occupy nano- and submicron-scale pores (Ambrose et al., 2010, Wang and Reed, 2009, Bennion et al., 2002, Al-Bazali et al., 2009, Wang, 2014). The resulting interfacial areas (fluid-fluid, and mineral-fluid) are relatively large for the pore volume and therefore control chemical and transport behavior in shale. As shown below, the combined effect of surface strain and fluid confinement results in a unique (and, largely unknown) set of thermodynamic parameters, different from those observed in the bulk phase (Firincioglu et al., 2012a, Wu, 2015, Teklu et al., 2014, Akkutlu and Rahmani, 2013). For
example, *molecular dynamic* (MD) simulations reported by Phan et al. (2014; 2015) have shown that methane solubility in confined water may far exceed that in bulk systems, and the release of methane from kerogen nano-pores is controlled by the geometry and connectivity of these pores (Ho et al., 2016). Methane solubility in confined water strongly depends on the confining material, with *silica* yielding the highest solubility followed by aluminum (Al₂O₃) and magnesium (MgO) *oxides* (Phan et al., 2014, Phan et al., 2016). Investigators looking into hydrocarbon behavior under confinement recorded confinement effects on phase behavior, and fluid properties (Akkutlu and Rahmani, 2013, Singh et al., 2009, Firincioglu et al., 2012b, Rahmani Didar and Akkutlu, 2013). Fig. 10 shows the suppression effect of nanopores on the phase envelop of pure methane and pure n-butane. The asymmetry in the suppression of the phase diagrams—the greater gap between the bulk and confined fluid in the left-hand-side compared to the right-hand-side on the phase diagram, indicates the greater impact of confinement on the vapor branch than on the liquid branch.
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Fig. 10. **Phase diagrams** of methane (top, left) and n-butane (top, right) in 4 and 8 nm size pores obtained from simulations and compared to bulk behavior. Phase diagram of binary methane-n-butane mixture (bottom). The bulk behavior is extracted from Peng-Robinson equation of state.

Adopted from Rahmani and Akkutlu (2015).
Cristancho et al. (2016) have recently quantified the impact of organic pore wall heterogeneities on the hydrocarbon fluid storage capacity of the pore using atomistic modeling and molecular simulations (Cristancho et al., 2016). For the purpose of creating heterogeneity, they have considered organic (carbon) walls with deficiencies and with non-hydrocarbon atoms, such as nitrogen. Their results, shown in Fig. 11, indicate that the measured excess amount due to confinement on the stored methane is most pronounced between 6.9 and 34.5 MPa (1000–5000 psi), which is the typical shale gas reservoir pressure range. The pore wall surface heterogeneities have the potential to impact storage depending on the type and level of heterogeneities. Among the investigated heterogeneities, nitrogen-doping at the pore walls is found to be the most influential. Both the adsorbed and excess amount decreased with the nitrogen-doped pore wall surfaces (Fig. 11).

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Fig. 11. Left: Organic pore wall model with surface heterogeneities. Carbon is shown in brown, hydrogen in yellow and nitrogen in blue. Right: Predicted storage of methane in the pore in adsorbed form (solid line) and excess (dashed line) methane. Predictions are based on grand canonical ensemble molecular simulations where the Lennard-Jones parameters (ε, σ) of methane-wall interactions have been estimated using quantum mechanical (DFT) calculations and ground-state energy optimization. Adopted from Cristancho et al. (2016)
The **dielectric constant** of water decreases with increasing nano-scale confinement (Marti et al., 2006, Senapati and Chandra, 2001), and this effect becomes more pronounced when the pore size approaches < 5 nm. Molecular dynamics simulations quantify the dielectric constant of water to be about one half of its unconfined value when water is contained in a 12 Å (1.2 nm) pore (Senapati and Chandra, 2001). Due to the decrease in the dielectric constant the equilibrium constant for the **mineral surface** protonation (pKa) change, ultimately changing the sorption behavior of the mineral surfaces. Bourg and Steefel (2012) calculated that the average pKa value of silanol surface sites in a 2 nm nanopore is 0.5 pH units higher compared to unconfined surfaces (Bourg and Steefel, 2012). Experiments indicate that in silica pores with < 5 nm pore size, both the density and the **surface tension** of water decrease with decreasing pore size (Takei et al., 2000). Other examples of emergent chemical behavior due to nano-scale confinement include the decrease in the **solvation** energy of metal **cations**, which promotes the formation of inner-sphere adsorption complexes over outer-sphere (Kalluri et al., 2011, Wang et al., 2003), enhanced solubility of gas in water (Diaz-Campos et al., 2009), enhanced adsorption (Wang et al., 2003, Nelson et al., 2014, Zimmerman et al., 2004), and modified redox properties (Jung et al., 2012, Patra et al., 2014a, Patra et al., 2014b). The nano-scale nature of the chemical environments in shale dictates the unique **chemical transport** and reactivity trends, in particular ion-selectivity and semi-permeable membrane behavior.

The nano-scale pore structure, mixed wettability, and multi-phase fluid conditions of shale have a strong effect on the disposition and movement of water, oil, and gas, which can all interfere with each other's movement. Quantitative measurements of shale samples by **nuclear magnetic resonance** (NMR) indicate that the amount of “free” water—water not structured by **electrostatic** forces—is largely variable for different shale samples, and does not correlate with the overall porosity (Dusseault, 2004). For example, only about 5% of total water content is “free” in smectite-rich shale of 10–20% porosity (Pierre II shale from Wyoming), while 50% of total interstitial water is “free” in the quartz-illite shale with the lower porosity of 6–8% (Queenston Shale from Ontario) (Dusseault, 2004). Surface-bound water is not likely to be oil wetting, thus oil migration through such pathways will require higher pressure. **NMR** measurements have also been used to demonstrate mixed wettability of source rock shale and for quantifying the water-wetting vs. oil-wetting porosity (Odusina et al., 2011). The resulting ratio of water (brine)-wetting to oil (dodecane)-wetting ranged from 0.34 to 2.93 (Odusina et al., 2011). This indicates that surfaces are heterogeneously wet - some shale surfaces are water-wetting, while others are hydrocarbon wetting. Interfacial contact lines will tend to pin at
locations where wettability changes, and additional pressure is required to cross these regions. Both adsorbed water and hydrocarbon will partially occlude the geometric area available for flow in nanopores affecting phase mobility (Sakhaee-Pour and Bryant, 2012, Hu et al., 2014).

Water in shale is subject to a variety of forces that affect its potential energy state (relative to a reference state) as expressed by the total potential. The total potential of water incorporates the matric potential, the osmotic (or solute) potential, the pressure potential, and the gravitational potential (Nitao and Bear, 1996). Pressure and gravitational potentials are invoked for understanding fluid flow in conventional reservoirs and aquifers. In tight shales, Darcian flow may not take place due to the limiting pore throat size. Because the majority of water may be “structured” – bound to the mineral surfaces – a threshold pressure gradient is required for advective transport to begin (Dusseault, 2004). High osmotic pressures (1–2 MPa) develop since hydrated ion flux is impeded (Dusseault, 2004). Fig. 12 shows the predicted osmotic pressure in the clay mineral pores as a function of the distance from the hydraulic fracture surface (Eveline et al., 2016). Hydraulic conductivity of smectite-rich shales depends on specific mineral surface characteristics (e.g., mineral surface charge, the ionic composition of the interstitial water, and temperature (“structured” water layer thickness decreases with increasing temperature, causing an increase in hydraulic conductivity). Therefore, hydraulic conductivity of shale is not a unique function of the pore and throat geometry. The following sections address the importance of the matric and osmotic potentials in shale with regard to the following: spontaneous imbibition-drainage, adsorptive fluid films and ion, and solute transport in shale.
Fig. 12. Osmotic pressure increase by the hydraulic fracture as a function of shut-in time at various locations (1, 5, 10, 50, and 100 cm) in the shale matrix near the fracture. The simulation involves flow of water in a rock (semipermeable membrane) driven by the pressure and chemical potential gradient. Membrane efficiency is 0.1 (top) and 1.0 (bottom). Osmotic pressure increase is defined as the difference between the initial pore pressure and pore pressure after fluid invasion. A constant pressure is applied on the left boundary. The objective of the simulation is to understand the effect of hydraulic fracturing water with low salinity (equivalent NaCl concentration of 10,000 ppm) on the adjacent shale matrix containing formation water with higher salinity (50,000 ppm). The initial pressure is 3000 psi and temperature is 50 °C. The rock has permeability of 200 nD and porosity of 10%. The further most left cell has a constant pressure of 3000 psi, which is representing shut-in pressure in the fracture, temperature of 50 °C. The matric potential expresses combined effects of capillarity and adsorptive forces of a porous medium (Hillel, 1982). Capillary forces arise due to a pressure difference.
related to the curvature of the interface between two immiscible fluids, as expressed by the Young-Laplace equation that incorporates the effects of the pore size, interfacial tension, and contact angle. The nanometer-scale pore sizes and water-wet or intermediate-wet conditions for certain pore networks in shale lead to high capillary pressure that can become significant in comparison to pressure potentials. The features of shale that control the matric potential include the pore size, shape, and wettability. Many studies have been performed to examine the pore structures in shales (Chalmers et al., 2012, Ambrose et al., 2010, Wang and Reed, 2009, Bustin et al., 2008, Curtis et al., 2011, Desbois et al., 2009, Dewers et al., 2012, Heath et al., 2012, Silin and Kneafsey, 2011), with Desbois et al. (2009) classifying pores. Pore geometry imposes the first order control on the mobility of water. Smaller, triangular pores or flat narrow pores (on the scale of 10 nm) have a tendency to imbibe water and spontaneously fill at a given water chemical potential compared to larger, circular pores (Or and Tuller, 1999) in part because adsorptive films take up a substantial portion of the pore space (Heath et al., 2014). “Corners” influence the curvature of the fluid-fluid interface and thus their geometry affects the amount of capillary-held water (Heath et al., 2014). The matric potential is currently being cited to explain the field observation when only a small portion of the water injected during hydraulic fracturing returns to the wellbore (e.g. ~ 30 ± 10% return, (Byrnes, 2011)). Gas shale systems typically have high thermal maturities with little to no smectite component left in the MLC (e.g. Marcellus and Haynesville shales). In the thermally mature systems only a fraction of injected water is recovered, due to the low chemical potential of water in these formations. In some cases, the observed return of injected water is higher: for examined wells in the Marcellus Shale, 10–50% of the fracturing fluid returns to the surface as produced water after a year of production (Rowan et al., 2015); while for Barnett Shale, large variability from < 20% to > 350% in the ratios of returned to injection water are observed after 4 years of production (Nicot et al., 2014). The return in excess of 100% indicates that wells produce local waters from adjacent formations. Extensive petrophysical evaluation of gas-bearing shale systems shows them to have very low water saturations and essentially no free mobile water phase. The limited water activity in organic-rich shale and how it is controlled by thermal maturity is an example of a coupled process that evolves over geological time scale.

Osmotic, or solute potential, is defined as the potential of water molecules to move from a hypotonic solution to a hypertonic solution across a semi-permeable membrane (Hillel, 1982). Osmotic potential is a function of the gradient in solute concentration. Due to the nano-scale porosity and the permanent negative charge on the surfaces of clay
minerals, shale acts as an ion-selective semi-permeable membrane, with apparent ion mobility differing from the bulk-solutions (Lomba et al., 2000a). Therefore, shale can be conceptualized and modeled as an ion-exchange membrane with fixed charged sites along narrow pores (Lomba et al., 2000b). During Fickian ion transfer waters of hydration are transported with the ions. Flow occurs by osmotic pressure, governed by gradient in chemical potential. Shale exhibits a non-ideal, or “leaky” membrane behavior, due to the heterogeneity in pore size and pore throat geometry, with wider pores increasing the overall permeability to solutes. High concentrations of solutes have been observed in flowback water from hydraulic fracturing operations, indicating higher concentrations in the subsurface reservoir than in the injected water. The presence of this high-salinity brine and injection of low salinity brine is expected to drive an osmotic flow (Haluszczak et al., 2013, Arthur and Cole, 2014, Engelder et al., 2014). The effects of osmotic pressure have been observed during well completions. Water from introduced fluids is absorbed into shale due to the difference in osmotic potential between the pore waters and the drilling mud or fracturing fluid, causing sloughing of well walls (Al-Bazali et al., 2009, Chen et al., 2003, Schlemmer et al., 2003, van Oort et al., 1995). These processes have been incorporated in a coupled mechanical-thermal-physico-chemical model (Choi et al., 2004). To account for the driving force on fluid flow due to osmotic potential, osmotic pressure is explicitly included in the rock water potential. The gradient between the rock water potential and water potential in drilling mud (or, hydrofracturing fluid) is the driving force for pore fluid flow (Choi et al., 2004). Because shales may exist under low water saturations (for both liquid and gas hydrocarbons), the strong capillary and adsorption potential results in spontaneous imbibition and potentially counterflow of hydrocarbons from the shale matrix (Engelder, 2012, Engelder et al., 2014). Other researchers quantify water uptake in shales through imbibition experiments, while taking into account effects of rock texture, complex pore networks, interactions between hydrofracturing fluid and minerals, and changes in osmotic potential, with some studies accounting for how concurrent and countercurrent flow during water imbibition impacts liquid and gas hydrocarbon recovery (Ghanbari and Dehghanpour, 2015). Findings indicate that the connectivity of water-wet and oil-wet pathways may differ, which in turn affect water uptake and hydrocarbon expulsion (Engelder et al., 2014, Ghanbari and Dehghanpour, 2015). Thus, the matric potential is a major driving force for water (and other fluid) flow in the nano-scale matrix in shale.

3.5. Equilibrium-disequilibrium transitions in perturbed shale
The combination of heterogeneous rock texture and composition, heterogeneous surface forces acting on the solid and fluid (liquid and gas) constituents, and the potential separation of fluid types (water and hydrocarbons) based on their interactions with wetting and non-wetting surfaces suggests an added heterogeneity in the fluid distribution and the fluid pressure. The available models for coupled poro-elastic behavior are described by poro-elastic coefficients developed based on an assumption of homogeneous rock properties, homogeneous stresses, and hydraulic pressure equilibration. These models cannot be extrapolated to mudrocks, particularly due to the coarse – 2 ft. (60 cm) – volumetric averages typical for well-log measurements. The typical examples of perturbations include injection of CO₂ (into sandstone storage formations with shale caprock) and unconventional gas extraction by hydraulic fracturing of shale. In both cases, the initial state of shale formation is either at steady-state, or at equilibrium, before the drilling and introduction of large volumes of fluid disrupt the initial state. The response of the geologic system to this perturbation is non-linear in space and time, since for re-equilibration it has to reach a multi-component - equilibrium, not an independent equilibrium of each of the components (e.g., hydraulic equilibrium). For example, the overall fluid flow and pressure equilibration for fine-grained rocks is controlled by chemical effects (osmosis), surface effects (capillary), physical effects (electrical double layer), and thermal gradient effects (relative expansion between solids and fluids in relation to hydraulic diffusion and thermal diffusion). The deformation versus pore pressure equilibration of organic-rich laminated mudrocks has the same contributions as considered for other fine-grained rocks, complicated by complex distribution of texture and compositions. This problem cannot be resolved by measuring an equivalent homogenized Biot's coefficient; new models are required to define coupled behaviors in locally heterogeneous media.

The state of geochemical (dis)equilibrium is usually assessed by analyzing fluid samples. Detailed chemical and isotopic analyses of pore waters in shale and tight reservoirs have been reported for only a few recent case studies (Haluszczak et al., 2013, Rowan et al., 2015, Arthur and Cole, 2014, Council GWP, 2009). Chemical and isotopic data reported by oil companies from approximately 10,000 samples of ‘flowback’ and produced waters from these unconventional sources of petroleum have been recently compiled and added to the updated and expanded USGS Produced Waters Geochemical Database (Blondes et al., 2015). The reported salinity and chemical composition of water varies widely with time of sampling and carry large uncertainties, especially for the ‘flowback’ samples that are a variable mixture of pore formation water and the hydraulic fracturing fluids, which consist of large volumes
(~ 10,000–40,000 m³ per well) of generally fresh, local meteoric water, together with proppants (sand), and organic and inorganic chemicals additives (Kharaka et al., 2013, Bryndzia and Braunsdorf, 2014, Gallegos et al., 2015, Healy et al., 2015). During hydraulic fracturing, a significant fraction of the injected water can be imbibed into pores in some shales (e.g. Marcellus (Rowan et al., 2015, Engelder et al., 2014)), and the imbibition process may continue over a period of weeks to months (Byrnes, 2011, Nicot et al., 2014). Water salinities and chemical compositions obtained at steady chemical states, which may require a year or longer following production, vary greatly from basin to basin. Results show formation waters with relatively low salinities are present in Fayetteville Shale, AR (~ 15,000 mg L⁻¹) and in Monterey Formation, CA (~ 30,000 mg L⁻¹). Produced water salinities in Barnett Shale, TX, average at ~ 100,000 mg L⁻¹, but higher average salinities (~ 150,000 mg L⁻¹) are obtained in brines from the Marcellus Shale, PA, and Haynesville, TX; even higher average salinities (> 250,000 mg L⁻¹) are observed in brine in Bakken Shale. An important initial conclusion from these data is that the chemical and isotopic compositions of these samples are comparable with data from more than 150,000 samples currently listed in the same USGS Produced Waters Geochemical Database, but collected from conventional oil and gas wells (Blondes et al., 2015). It remains challenging, however, to use this geochemical data for predictive modeling.

4. Conclusions: approaches for coupled process studies and future research needs

4.1. Summary of applied approaches/methodologies

Table 2 indicates the various common data types relevant to shales, and their spatial and temporal resolution as well as well-established methods for integration of multi-disciplinary data.

Promising approaches for addressing coupled processes so far included dynamic consolidation problems with elastoplastic deformation and finite element modeling (FEM) (Lewis and Schrefler, 1987, De Borst et al., 1993, Armero, 1999, Kim, 2000, Chen et al., 2015a), quasi-static discrete element models (DEM) coupled with conjugate lattice network flow (Huang and Mattson, 2014), basin petroleum system modeling (Romero-Sarmiento et al., 2013), incorporating Knudsen diffusion and gas slippage (in addition to Darcy flow) into reservoir models for shale (Swami and Settari, 2012), and lattice Boltzmann (LB) approaches for coupled multi-component reactive flow and transport with the feedback between pore structure changes and flow
processes (Chen et al., 2015b, Chen et al., 2015c). The typical measurement resolution and modeling methods used for single and coupled processes in shale are shown in Fig. 13.
Electron and optical microscopy
SEM  BSE  Optical
TEM
FIB/SEM
Neutron methods
SANS/USANS  (lab) active/passive
EM techniques
SAXS/USAXS  micro to larger X-ray CT
Porosimetry
Nanotechnology
Contrast agents
Well logs
Resistivity/sonic dipole/FMI/etc
Seismic
Cross-well
Surface survey
Microseismic
Pressure/rate transient analysis
Fiber optic sensing
Ab-initio
LBM
Continuum
Single/double/multi FEM/XFEM
Pore network
Reactive transport
DEM
Discrete
Fracture
Network
The physics-based hydraulic fracturing simulator (Huang and Mattson, 2014) couples a quasi-static discrete element model for deformation and fracturing with conjugate lattice network flow model for fluid flow in both fractures and porous matrix. This two-dimensional model with coupled deformation and flow reproduces growth patterns of hydraulic fractures. The model accounts for in situ stress, fluid viscosity, heterogeneity of rock mechanical properties and injection rate. The modeling of a multistage horizontal wellbore confirms the strong coupling between observed complex fracture patterns and fluid pressure, small length scale heterogeneities, and elastic interactions among multiple propagating fractures (Huang and Mattson, 2014).

The lattice Boltzmann method is used for modeling pore-scale reactive transport and allows accounting for complex biogeochemical processes - mineral dissolution-precipitation and biofilm dynamics, and their feedback to transport (e.g. Yoon et al., 2015). LB models conceptualize flow as a collective behavior of pseudo-particles described by a discrete Boltzmann equation (Yoon et al., 2015). In shale applications, LB modeling is capable of predicting permeability and effective Knudsen diffusivity of the shale samples characterized by FIB-SEM (Chen et al., 2015b). LB models have been applied to multi-phase flow with phase transition (Shan and Chen, 1993) and development of preferential flow paths in porous media (Szymczak and Ladd, 2006). Basin petroleum system modeling is relatively new for unconventional hydrocarbon resources and is still under development and refinement. Basin-scale shale-play modeling accounts for source-rock kinetics and chemical transformations, including the evolution of TOC and associated porosity and adsorption of hydrocarbons to the mineral and organic components (Romero-Sarmiento et al., 2013). However, this modeling approach does not account for the chemical effects due to nano-scale confinement, non-Darcy transport process (e.g., Knudsen transport), and complex cementation/dissolution textures. The retention capacity in BPSM is also a coupled process between burial, uplift and geomechanical rock properties since the in situ fluid pressures are ultimately determined by the fracture gradient that a shale can sustain as a result of overburden loss during uplift. One of the most important challenges for resource assessment in unconventional plays is the need to quantitatively model the retention capacity of such organic-rich source rocks.
4.2. Coupled processes in shale: future research needs

The development of quantitative predictive models capturing process coupling and emergent phenomena at the necessary length and temporal scales requires a systematic approach involving a broad range of multi-disciplinary techniques. Current research is hindered due to sparse data from shale formations at *in situ* pressure and temperature conditions, limited spatial resolution of the well-logging/interpretation techniques, and methodologies for merging multi-disciplinary datasets at different length and time scales. Therefore, on the practical side, it is critical to define the type of relevant measurements, improve the resolution and link *in situ* well resistivity logging to chemistry, and develop better approaches for representative or standardized sampling and sample preservation. In particular, addressing heterogeneity and anisotropy, which leads to scaling complexity, is a major challenge. Below, we identify future research needs critical for fundamental and applied shale science, considering both single- and multi-disciplinary approaches.

It is necessary to develop unambiguous classification schemes for *mudrocks*, and develop further understanding of the material transport and cementation mechanisms during shale deposition and *diagenesis*. We also need to establish the range of sizes for REVs, which are different for *kerogen*, clay mineral-rich components, and individual *lithofacies*, and are controlled by the unique depositional and diagenetic history. As shown in our review, the complex cementation and dissolution textures—both solid components and pores in shale—control the fluid transport, chemical behavior, and mechanical properties. Understanding fundamental geological/chemical/physical controls on the formation of these textures, and proposing a comprehensive classification scheme, could lay the foundation for the development of predictive methods to understand physical flow, chemical behavior and reactive transport, and mechanical behavior of shale in native state and in engineered systems.

Another key research area is developing robust methods for integrating rock anisotropy into geomechanical analysis, especially proper characterization and modeling of mechanical interfaces, and developing new constitutive laws describing stress-strain relationships for shale. *Seismic methods* show promise in characterizing fractures; however, research is needed to understand the contributions to the scattered wave field from fracture intersections, fracture sets, stress gradients and fluids in the dynamically evolving fractured shale systems. To further refine the interpretation of *seismic data*, and to characterize dynamically evolving fracture populations, seismic data needs to be integrated with high resolution imaging. A potential bridge here is the integration of high
resolution image logs (centimeter scale) with petrological studies (micron scale) that may be upcaled to the seismic scale. The upscaling issue dominates the use of seismic methods for quantitative assessment of reservoir rock properties at all scales. Future research should focus on separating competing sources of anisotropy in rock and fluid flow, and how seismic signals change as a function of stress and fluid conditions. Future research needs to answer the fundamental question - can seismic data differentiate or unravel chemical, fluid and stress alteration of fractures? The geochemistry of shale is unique, characterized by high salinity, complex compositions of brine and solids, water-limited, and nano-scale confined chemical environments, resulting in ion-selectivity and semi-permeable membrane behavior. In order to interpret and predict the chemical behavior in these environments, we need to develop new thermodynamic databases, as well as a systematic approach for predicting shifts in chemical kinetics under these conditions. Fundamental science of nangeochemistry is still in its infancy.

As shown in our review, process coupling exerts major controls on the physical, mechanical and chemical behavior of shale. Further development of coupled models is crucial for predicting coupled fracture, multi-phase flow, and multi-component reactive transport in the subsurface. New models are required to address coupled behavior in locally heterogeneous shale media. For the development of these models, the solid solvers need to be coupled in a 2-way manner to the multi-phase reactive flow and transport codes, and incorporate changes in volume and chemical concentrations, and their feedback to the mechanical properties and permeability. Due to complex pore geometries and the large variety of pore types and their control on the flow processes in shale, the development of a porosity-permeability relationship is a challenge. Since fluid flow in shale is coupled to the mechanical behavior, a method is required for computing the effect of flow on the mechanical constitutive behavior of shale (as partially-drained or undrained medium). These models should also incorporate multi-porosity system behavior, as distinct pore networks in organic and mineral components can exhibit different stiffness/compliance as a function of stress, resulting in different time scales of permeability evolution. Additional laboratory data for calibration and verification of these coupled models is necessary.

Further development of data integration approaches is another critical research need, due to the primary controls that the nano-scale processes exert on the macro-scale behavior. In an ideal case, we should be able to merge data all the way from the mineral-water interface (sub-nanometer), to rock microstructure observations.
(micron), to the core-scale samples (centimeter-meter), well-log scale (meters), and linking to seismic scale (many meters to kilometers) (Table 2). For developing fully coupled multi-physics multi-scale models for shale, we need a fundamental understanding of the interplay between the physical and chemical processes, their rates and resulting emergent behavior. We need to understand and quantify the evolution of thermal, hydrologic, chemical, mechanical, and biological (dis)equilibrium during perturbations (withdrawal or emplacement of materials into subsurface), and identify the relevant representative elementary volumes, as well as representative elementary time scales. Further development in constitutive laws (or, equations of state) is needed to incorporate evolving stress-pressure and system transitions. This requires advanced modeling linking molecular-, to pore-, to macroscopic-scale processes and the formalization of heterogeneity and spatial and temporal scales. We need to improve and merge conceptual models and develop a common language for the multi-disciplinary research on coupled THCMB processes in shale.

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