Advances in Geo-Energy Research⁻

Original article

The influence of adsorption layer thickness and pore geometry on gas production from tight compressible shales

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Keywords:

Adsorption model adsorbed layer thickness compressible shale shale gas production

Cited as:

Abolghasemi, E., Andersen, P. Ø. The influence of adsorption layer thickness and pore geometry on gas production from tight compressible shales. Advances in Geo-Energy Research, 2021, 6(1): x-x. https://doi.org/10.46690/ager.2022.01.02

Abstract:

In tight shales, gas is stored in both free and adsorbed form. A one-dimensional model is derived for shale gas production by pressure depletion where the adsorbed layer thickness is of similar magnitude as the pore radius and can affect flow performance. The adsorbed layer thickness is a function of pressure. Different pore geometries are assumed varying continuously between spherical pores to more fracture shaped pores. The shale is assumed compressible and its porosity and pore radius reduce with pressure depletion. The effective pore radius (pore radius minus adsorption layer thickness) controls intrinsic and apparent permeability. The impact of the adsorption layer, compressibility and geometry are investigated. A given adsorbed layer thickness fills more of the pores when they are more spherical and concentrates more of the volumetric flow to the effective pore boundaries giving lower permeability for a given effective radius. Increasing the adsorbed thickness increases the adsorbed fraction initial gas in place. A high volume fraction adsorbed gas reduces apparent permeability and delays production. Pressure depletion causes both pore radius and adsorbed layer to be reduced. The change in adsorbed layer thickness is low at high pressure and greater at low pressure, while pore radius changes more linearly and more with higher compressibility. The free gas saturation and slip increases with pressure depletion for low compressible cases, but if matrix compression dominates there can be a net reduced permeability. Recovery was linear with the square root of time for all cases. Adsorbed gas is less effectively produced by pressure depletion than free gas and higher adsorbed content by more spherical pores or higher layer thickness reduces end recovery. Higher compressibility reduces permeability and delays recovery but increases end recovery.

1. Introduction

Shale gas reservoirs are organic-rich shale formations which act as both reservoir and source rock. These formations are characterized by ultra-low permeability and can contain a significant quantity of adsorbed gas in addition to free gas. The amount of adsorbed gas can vary between 20-85% of the total gas-in-place volumes (Curtis, 2002; Montgomery et al., 2005; Wu et al., 2014; Gou and Xu, 2019). The adsorbed gas is densely packed on the surface of the shale matrix and in the organic matter known as kerogen while the free gas mainly

resides in nanopores and naturally occurring fractures (Yu and Sepehrnoori, 2014; Chen et al., 2019; Sheng et al., 2020). The adsorption capacity of a shale depends on factors such as total organic content, clay content, the specific surface area of nanopores and pore-size distribution (Strapoc et al., 2010; Wang et al., 2017; Klewiah et al., 2020). CO_2 can adsorb preferentially over CH₄ and with greater capacity, hence CO_2 injection in shales can be a means to sequester CO_2 , mitigate climate changes and enhance shale gas production (Berawala and Andersen, 2020a; Klewiah et al., 2020). Gas adsorption

Yandy Scientific Press

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2207-9963 © The Author(s) 2021. Received November 11, 2021; revised November 29, 2021; accepted November 30, 2021; available online December 3, 2021. in shales is commonly modeled using a Langmuir isotherm (Langmuir, 1916) with the characteristic feature that substantial desorption does not occur until a significant pressure drop is obtained. Yu and Sepehrnoori (2014) history matched Marcellus and Barnett shales and found that desorption for this reason was most significant during late time production.

Technological advancements in horizontal drilling with hydraulic fracturing has made the extraction of gas from shale reservoirs economically viable. The overall recovery, which is still very low, 10-30% of the gas-in-place, however, depends on effective stimulation (Wu et al., 2014). Strong capillary or osmotic forces in shale can draw fracturing water into the matrix where it can be trapped and block gas flow (Andersen, 2021b). Typical gas production rates decline steeply after an initial peak, which affects the economics of many shale gas projects (Valko and Lee, 2010; Patzek et al., 2014). The tight nature of shales means that mainly the stimulated reservoir volume around a single well is produced effectively. Commonly, shale gas production can be studied by means of the behavior around representative individual hydraulic fractures (Berawala et al., 2019; Berawala and Andersen, 2020b).

In shales, gas transport takes place on spatial scales covering several orders of magnitude. The transport processes can effectively be divided into matrix and fractures. The flow in the matrix, which occurs on nanometer scale, is characterized by slip flow, transition flow or free molecular flow regimes based on the Knudsen number. The mean free path of gas molecules in the matrix is usually of the same order of magnitude or larger than the size of pore throat. This may cause acceleration of gas molecules leading to an increase in the apparent permeability (Tang et al., 2017). The adsorbed gas is mostly produced in the latter stages of production when a considerable amount of free gas has been depleted (Mengal and Wattenbarger, 2011). In addition to matrix flow, fracture flow is critically important in unconventional reservoirs (Zhao and Du, 2019). A challenge is to open and maintain the fractures for effective gas transport.

Significant efforts have been made to effectively model the shale gas production system. A rarefaction coefficient was introduced by Beskok and Karniadakis (1999) to account for the reduced intermolecular collisions in the transition and freemolecular regimes. Javadpour (2009) introduced an apparent permeability term that includes the complexity of flow in nanopores, for application in the Darcy equation so it easily can be implemented in reservoir simulations. Civan (2010) introduced a methodology to provide accurate and meaningful correlations of data including intrinsic permeability, porosity, tortuosity effect on apparent gas permeability, rarefaction coefficient, and Klinkenberg gas slippage factor. Different modelling approaches such as molecular dynamics (Bird and Brady, 1994), direct Monte Carlo simulation (Karniadakis et al., 2006), Burnett equation (Karniadakis et al., 2006) and Lattice-Boltzmann (Hornyak et al., 2008) methods have also been implemented to model gas flow in nanopores. Characteristics like pore pressure, reservoir temperature, surface roughness, and phase change have also been studied for shale (Higashi et al., 1963; Zarragoicoechea and Kuz, 2004; Chen et al., 2008). Clarkson et al. (2012) introduced a dynamicslippage model to study production dynamics. Knudsen diffusion was extended into the slippage factor by Civan et al. (2011) to explain gas transport. Pore transport of free and adsorbed gas was modeled by Sheng et al. (2020) and generalized to account for pore size distribution.

In addition to gas storage, adsorption can also affect gas flow relations. Xiong et al. (2012) assumed the adsorbed layer to have a thickness of one molecular diameter and calculated that a 2 nm pore could lose half its porosity and 75% of its permeability by adsorption. Li et al. (2016) modeled shale gas flow at pore scale with the Lattice-Boltzmann approach incorporating pressure dependent layer thickness. They found that permeability was reduced when the adsorbed layer filled more of the pores. Cao et al. (2016) included adsorption as a volumetric strain in a poroelastic model for shale gas. Pang et al. (2017) measured adsorption and compressibility in shale. Based on volumetric calculations the experimental effective porosity reduced when a nonadsorbing gas was present during pressure depletion, but when an adsorbing gas was present the effective porosity increased due to desorption. Similar findings were made by Memon et al. (2020). Jiang and Yang (2018) coupled a geomechanical and fluid flow model. They observed higher gas rate when the adsorbed layer was not included. Sheng and Javadpour (2018) accounted for compressibility and adsorbed volume in the organic pores and presented apparent porosity and apparent permeability formulations for use in a dual-continua reservoir model.

This paper presents a one-dimensional (1D) dynamic model for shale gas production from the matrix with nspherical pore geometry. Adsorbed gas occupies portions of the volume as a finite layer and has implications on flow, storage and production dynamics. The shale is assumed compressible and its porosity and pore radius reduce with pressure depletion. Consistent relations for porosity and permeability capturing these effects are presented. The gas transport equation incorporates pore geometry as a free parameter n, apparent permeability, compressibility, adsorption layer thickness and free gas saturation. The system is solved numerically using operator splitting and experimental data from the literature are used as input with significant adsorption and compressibility effects. The proposed model addresses some questions of practical importance:

- 1) How does adsorption affect flow performance?
- 2) What is the impact of varying adsorbed layer thickness, pore geometry and compressibility on porosity, permeability, free gas saturation and gas recovery?
- 3) Which parameters control final recovery?

These questions are answered with analytical relations for recovery and other output; sensitivity analyses on these relations as function of pressure for different pore geometry, compressibility and adsorption layer thickness; and investigating the evolution of shale gas production for such variations.

2. Mathematical model

2.1 Geometry

Shale gas production from a shale matrix 1D system (e.g.,

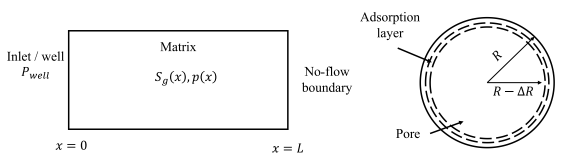


Fig. 1. Geometrical representation of the system. Left: The matrix. Right: Representation of a pore in the matrix.

a core sample or shale connected to a hydraulic fracture) is considered with length L, as shown in Fig. 1. Production takes place at x = 0 with a fixed pressure P_{well} , while there is a noflow boundary at x = L representing the length of the system or half the fracture spacing. The matrix initially contains free and adsorbed gas, with same composition (assumed to be methane). The free gas is compressible, modeled as a real gas, while the adsorbed gas has a constant, relatively high density. No other phases are assumed present. Pores have the shape of an *n*-sphere. This means, for n = 1 the pores are planar, for n = 2 cylindrical and for n = 3 spherical, while intermediate values indicate geometrical configurations in between. In accordance with standard reservoir modeling assumptions, compression is accounted for via changes in the pore volume and a fixed bulk volume is considered.

2.2 Transport equation

Performing a mass balance on the system, the change in mass of free and adsorbed gas in a bulk volume Adx is:

$$\begin{aligned} \left[\phi(\rho_g S_g + \rho_a S_a) dx \right]_{t+\Delta t} &- \left[\phi(\rho_g S_g + \rho_a S_a) dx \right]_t \\ &= (\rho_g u)_x \Delta t - (\rho_g u)_{x+\Delta x} \Delta t \end{aligned}$$
(1)

where ρ_g and ρ_a are free and adsorbed gas densities respectively (in mass per phase volume), ϕ is prosity, S_g is free phase saturation, S_a is adsorbed phase saturation, *t* is time, *u* is Darcy free gas velocity and *A* is the constant cross section area (omitted from the equation). Rearranging and letting Δx , $\Delta t \rightarrow 0$ gives the following mass conservation equation:

$$\partial_t \left\{ \phi \left[\rho_g S_g + \rho_a \left(1 - S_g \right) \right] \right\} = -\partial_x (\rho_g u) \tag{2}$$

Above, it is used that the two saturations sum to 1 and the adsorbed phase saturation was eliminated. ρ_a will be assumed constant (similar to a liquid) while ρ_g behaves according to a real gas:

$$\rho_g = \frac{pM}{zR_gT} \tag{3}$$

where p is gas pressure, R_g is the gas constant, T is absolute temperature, M is gas molar weight and z is compressibility factor. The Peng and Robinson (1976) equation of state is used to calculate z, see Appendix A. In Eq. (2) the gas flux u is described using Darcy's law:

$$u = -\frac{K_a}{\mu} \partial_x p \tag{4}$$

in which K_a is pressure-dependent apparent permeability and μ is pressure-dependent gas viscosity. $\mu(p)$ is calculated using the Lee et al. (1964) correlation, see Appendix B. The above relations lead to the following form of the transport equation:

$$\partial_t \left[\phi \left(\frac{p}{z} S_g + R_g T \rho_a \frac{1 - S_g}{M} \right) \right] = \partial_x \left(\frac{p}{z} \frac{K_a}{\mu} \partial_x p \right) \tag{5}$$

2.3 Porosity model accounting for pore geometry and adsorption

The *n*-volume V_n of an *n*-sphere pore with radius *R* is (Evans, 1947):

$$V_n = \frac{\pi^{\frac{n}{2}}}{\Gamma\left(\frac{n}{2}+1\right)} R^n \tag{6}$$

The Gamma function (Γ) above is defined as a semi-infinite integral:

$$\Gamma(m) = \int_{x=0}^{\infty} x^{m-1} e^{-x} dx \tag{7}$$

which is corresponding to the factorial function as below for integer input values:

$$\Gamma(m) = (m-1)! \tag{8}$$

By considering an *n*-sphere, the *n*-volume will be obtained. To turn this into a three-dimensional volume a factor *C* (volume per *n*-volume) was used. For n = 1, *C* equals the cross section of a fracture, for n = 2 it is the length of a cylinder, and for n = 3 it equals 1. Let *N* be the total number of pores per unit bulk volume. The total pore volume V_p is then:

$$V_p = NCV_n \tag{9}$$

Porosity follows as:

$$\phi = \frac{V_p}{V_b} = \frac{NC}{V_b} V_n \tag{10}$$

Assuming the porosity is known at initial time ϕ_0 , then:

$$\phi_0 = \frac{NC}{V_b} V_n^0 \tag{11}$$

where V_b is bulk volume, V_n^0 is the *n*-volume of a pore at initial state. The ratio NC/V_b may be determined from Eq. (11) and substituted into Eq. (10) to yield porosity written as:

$$\phi = \phi_0 \frac{V_n}{V_n^0} = \phi_0 \left(\frac{R}{R_0}\right)^n \tag{12}$$

where R is pore radius, R_0 is initial pore radius. These relations can then be used to equivalently express pore radius as function of porosity given the initial radius and porosity:

$$R = R_0 \left(\frac{\phi}{\phi_0}\right)^{\frac{1}{n}} \tag{13}$$

Porosity may be taken as a function of pressure and (pore) compressibility c_k :

$$\phi = \phi_0 \exp[c_k (p - p_0)] \tag{14}$$

where p_0 is the initial pressure, corresponding to the initial porosity ϕ_0 and radius R_0 . The amount of adsorbed gas in shale is commonly described by the Langmuir isotherm which assumes gas attaches to the surface as a single layer (a monolayer). Regardless of pore geometry, an adsorbed layer in the pores with thickness ΔR following a Langmuir relation with pressure may be defined as (Xiong et al., 2012; Jiang and Yang, 2018):

$$\Delta R = \Delta R_{\max} \frac{p}{p_L + p} \tag{15}$$

where p_L is the Langmuir pressure, ΔR_{max} is the maximum thickness at theoretically infinite pressure the thickness is half that value at the Langmuir pressure p_L . By representing the adsorbed layer by an isotherm it is implicitly assumed that the adsorption process occurs instantaneously. Adsorption can occur at a finite rate where a source term is better suited to represent the process. In that case, the thickness ΔR would be a function of both pressure and time and reach the equilibrium state Eq. (15) only when sufficient time had passed.

For a given layer thickness ΔR in a pore, using Eqs. (6) and (9), the volume of free gas V_g in the pore volume is:

$$V_g = NC \frac{\pi^{\frac{n}{2}}}{\Gamma\left(\frac{n}{2}+1\right)} R_e^n \tag{16}$$

$$R_e = R - \Delta R \tag{17}$$

where R_e is the effective radius. The free gas saturation S_g is defined as the fraction of pore volume filled by free gas. The remaining fraction is adsorbed gas.

$$S_g = \frac{V_g}{V_p} = \left(\frac{R_e}{R}\right)^n \tag{18}$$

This can also be used to define an effective porosity (ϕ_e), i.e., the volume fraction of the bulk volume occupied by free gas:

$$\phi_e = \phi S_g = \phi_0 \left(\frac{R}{R_0}\right)^n \left(\frac{R_e}{R}\right)^n = \phi_0 \left(\frac{R_e}{R_0}\right)^n \qquad (19)$$

At initial state it holds that:

$$\phi_{e0} = \phi_0 \left(\frac{R_{e0}}{R_0}\right)^n \tag{20}$$

Combining the two Eqs. (19) and (20) results in:

$$\phi_e = \phi_{e0} \left(\frac{R_e}{R_{e0}}\right)^n \tag{21}$$

2.4 Apparent permeability model accounting for pore geometry and adsorption

Apparent permeability is typically separated into two factors, where the first is the intrinsic permeability k_{∞} and the second is the slip correction factor f, typically modeled as dependent on the Knudsen number K_n :

$$K_a = k_{\infty} f(K_n) \tag{22}$$

2.4.1 Intrinsic permeability model

A similar derivation will be followed as Zimmermann and Bodvarsson (1996) did for a plane fracture (n = 1). Consider flow through a pore with geometry characterized by the effective radius R_e and parameter n. The interstitial velocity field in the pore is denoted $\mathbf{v}(r,x,t) = (v_r(r,x,t), v_x(r,x,t))$ where it is assumed that the field can depend on the radial distance r from the pore center, the position along the general flow axis x and time t. r is normal to x. The evolution of the velocity field is given by Navier-Stokes equation (Batchelor and Batchelor, 2000):

$$\partial_t \mathbf{v} + (\mathbf{v} \cdot \nabla) \mathbf{v} = \mathbf{F} - \frac{1}{\rho} \nabla p + \frac{\mu}{\rho} \nabla^2 \mathbf{v}$$
(23)

where **F** is a body force (gravity) and p(x, w, t) is the pressure field. Steady state is assumed and gravity ignored such that $\partial_t u = 0$ and **F** = 0 which yields:

$$\mu \nabla^2 \mathbf{v} - \rho (\mathbf{v} \cdot \nabla) \mathbf{v} = \nabla p \tag{24}$$

Since permeability is an intrinsic property, flow conditions can be based on an incompressible fluid, which has the property:

$$\nabla \cdot \mathbf{v} = 0 \tag{25}$$

Gas compressibility and slip will be accounted for as separate mechanisms. Three equations then result for the two velocity components v_r , v_x and the pressure p. The flow is considered to be in one direction x, such that the radial velocity component is zero, $v_r = 0$ while the velocity component in x-direction v_x will vary with r only: $v_x = v_x(r)$. The incompressibility condition is then obeyed:

$$\nabla \cdot \mathbf{v} = (\partial_r, \partial_x) \cdot (v_r, v_x) = \partial_r v_r + \partial_x v_x = 0$$
(26)

Also, the pressure gradient components are considered a constant in *x*-direction and zero in *r*-direction, respectively:

$$\nabla p = (\partial_r p, \partial_x p) = (0, \partial_x p) \tag{27}$$

The differential terms in Eq. (24) turn into:

$$(\mathbf{v} \cdot \nabla) \mathbf{v} = (v_r \partial_r + v_x \partial_x) (v_r, v_x) = (v_r \partial_r v_r + v_x \partial_x v_r, v_r \partial_r v_x + v_x \partial_x v_x) = (0,0) \nabla^2 \mathbf{v} = \left\{ 0, \frac{1}{r^{n-1}} \partial_r \left[r^{n-1} \partial_r v_x (r) \right] + \partial_{xx} v_x (r) \right\} = \left\{ 0, \frac{1}{r^{n-1}} \partial_r \left[r^{n-1} \partial_r u_x (r) \right] \right\}$$
(28)

The following equation is then obtained for the velocity component $v_x(r)$:

$$\mu \frac{1}{r^{n-1}} \partial_r \left[r^{n-1} \partial_r v_x(r) \right] = \partial_x p \tag{30}$$

Integrating the equation once gives:

$$\partial_r v_x(r) = \frac{1}{r^{n-1}} \left(\frac{1}{n} \frac{\partial_x p}{\mu} r^n + \Upsilon \right)$$
(31)

where Υ is an integration constant. It is assumed the profile is symmetric such that $\partial_r v_x|_{r=0} = 0$:

$$\partial_r v_x|_{r=0} = \Upsilon r^{-(n-1)}|_{r=0} = 0$$
 (32)

Since $n \ge 1$ the above equality can only hold if $\Upsilon = 0$. Eq. (31) thus simplifies to:

$$\partial_r v_x(r) = \frac{1}{n} \frac{\partial_x p}{\mu} r \tag{33}$$

Integrating again results in:

$$v_x(r) = -\frac{1}{n} \frac{\partial_x p}{\mu} \frac{1}{2} \left(R_e^2 - r^2 \right) \tag{34}$$

During the integration a no-flow condition was applied at $r = R_e$, the outermost radius accessible to gas flow, i.e., $v_x(r = R_e) = 0$. Implicitly the adsorbed phase is assumed immobile. As a remark, note that surface diffusion, the movement of adsorbed phase due to a gradient in adsorbed phase concentration, has been considered important in some works (Yang et al., 2016; Ren et al., 2017; Yin et al., 2017), but is not accounted for here.

Assume that the pore has an n-1-area A_{n-1} (with unit m^{n-1}) when the pore has *n*-volume V_n (with unit m^n) which depend on *n* and the effective radius R_e as follows:

$$A_{n-1}(R_e) = \frac{2\pi^{\frac{n}{2}}}{\Gamma\left(\frac{n}{2}\right)} R_e^{n-1}$$

$$V_n(R_e) = \frac{\pi^{\frac{n}{2}}}{\Gamma\left(\frac{n}{2}+1\right)} R_e^n$$
(35)

The total flow rate Q can be calculated by considering the contributions of flow through all *n*-volume elements dV_n in the pore. The incremental change in V_n given a change in radius for an arbitrary value r is related as:

$$dV_n(r) = A_{n-1}(r)dr \tag{36}$$

The cumulative flow is found by integrating the flow through every element of the entire cross section:

$$Q = \int_{r=0}^{R_e} v_x(r) C_f A_{n-1} dr$$

= $-\frac{C_f}{2n} \frac{\partial_x p}{\mu} \frac{2\pi^{\frac{n}{2}}}{\Gamma\left(\frac{n}{2}\right)} \int_{r=0}^{R_e} \left(R_e^2 r^{n-1} - r^{n+1}\right) dr$
= $-C_f \frac{\partial_x p}{\mu} \frac{2\pi^{\frac{n}{2}}}{\Gamma\left(\frac{n}{2}\right)} \frac{R_e^{n+2}}{n^2(n+2)}$ (37)

where the expressions for the flow profile $v_x(r)$ and the n-1-area A_{n-1} from Eqs. (34) and (35) have been inserted, respectively. The effective cross section open for gas flow is:

$$A\phi_e = \int_{r=0}^{R_e} C_f \frac{dV_n}{dr} dr = C_f V_n(R_e)$$

= $C_f \frac{\pi^{\frac{n}{2}}}{\Gamma\left(\frac{n}{2}+1\right)} R_e^n$ (38)

The average interstitial velocity v in x-direction is found by dividing the total flow rate by the porous cross section and the tortuosity τ (accounting for the fact that flow may not be along a straight line).

$$v = \frac{Q}{A\phi_e \tau} = -\frac{C_f \frac{\partial_x p}{\mu} \frac{2\pi^{\frac{n}{2}}}{\Gamma\left(\frac{n}{2}\right)} \frac{R_e^{n+2}}{n^2(n+2)}}{C_f \frac{\pi^{\frac{n}{2}}}{\Gamma\left(\frac{n}{2}+1\right)} R_e^n \tau}$$
(39)
$$= -\frac{\partial_x p}{\tau \mu} \frac{R_e^2}{n(n+2)}$$

The Darcy velocity equals the product of interstitial velocity and effective porosity:

$$u = \phi_e v = -\frac{\phi_e}{\tau \mu} \frac{R_e^2}{n(n+2)} \partial_x p \tag{40}$$

This can be compared with Darcy's law which states (without consideration of slip):

$$u = -\frac{k_{\infty}}{\mu} \partial_x p \tag{41}$$

This implies that the intrinsic permeability k_{∞} for the different geometries can be expressed as:

$$k_{\infty} = \frac{\phi_e}{\tau} \frac{R_e^2}{n(n+2)} \tag{42}$$

Tortuosity is often correlated with porosity, as reviewed by Shen and Chen (2007). A tortuosity relation with effective porosity is applied using the correlation by Bruggemann (1935):

$$\tau = \phi_e^{-0.25} \tag{43}$$

As special cases of the derived model for intrinsic permeability k_{∞} , note that n = 1 gives planar fracture pores, and n = 2 gives cylindrical pores. The corresponding *n*-volumes and n - 1-areas from Eq. (35) are then:

$$A_0(R_e) = 2, \quad V_1(R_e) = 2R_e, \quad (n = 1)$$
 (44)

$$A_1(R_e) = 2\pi R_e, \quad V_2(R_e) = \pi R_e^2, \quad (n=2)$$
 (45)

For n = 2, V_2 already corresponds to the cross section of the pore, while for n = 1 the factor C_f should equal the height h of the fracture to give the cross section. The permeability expressions become:

$$k_{\infty} = \frac{\phi_e}{\tau} \frac{R_e^2}{3} = \frac{\phi_e}{\tau} \frac{w_e^2}{12}, \quad (n = 1)$$
(46)

$$k_{\infty} = \frac{\phi_e}{\tau} \frac{R_e^2}{8}, \quad (n=2)$$
 (47)

where $w_e = 2R_e$ is the width of the fracture. Both these expressions are consistent with the literature (Carman, 1956; Zimmermann and Bodvarsson, 1996; Civan, 2007), although they typically do not account for the influence of adsorption and thus use the full porosity ϕ and radius *R*. Finally, by using the relations between effective radius and effective porosity from Eq. (21) and between tortuosity and effective porosity from Eq. (43), the relation Eq. (42) can be written as:

$$\frac{k_{\infty}}{k_{\infty 0}} = \left(\frac{R_e}{R_{e0}}\right)^{2+1.25n} = \left(\frac{\phi_e}{\phi_{e0}}\right)^{1.25+\frac{2}{n}}$$
(48)

2.4.2 Slip correction factor

To classify flow in small pores, the dimensionless Knudsen number is commonly used. It is defined as the ratio of molecular mean free path λ (Loeb, 2004) to a characteristic pore dimension, in this work assumed to be the effective radius R_e :

$$K_n = \frac{\lambda}{R_e} = \frac{z\mu}{pR_e} \sqrt{\frac{\pi R_g T}{2M}}$$
(49)

Apparent permeability K_a is related to intrinsic permeability k_{∞} according to the Knudsen number relation derived by Beskok and Karniadakis (1999):

$$K_{a} = k_{\infty} [1 + \alpha(K_{n})K_{n}] \left(1 + \frac{4K_{n}}{1 + K_{n}}\right)$$

$$\alpha(K_{n}) = \frac{128}{15\pi^{2}} \tan^{-1} \left(4K_{n}^{0.4}\right)$$
(50)

2.5 Initial and boundary conditions

The system has a defined initial pressure p_0 which also defines the initial radius, porosity, adsorption layer thickness and saturation:

$$p(x,t=0) = p_0, \quad R(x,t=0) = R(p_0)$$
 (51)

$$\Delta R(x,t=0) = \Delta R(p_0), \quad S_g(x,t=0) = S_g(p_0)$$
(52)

The boundary at x = 0 is open with a fixed pressure p_{well} while a closed boundary is applied at x = L:

$$p(x=0,t) = p_{well}, \quad (\partial_x p)_{x=L} = 0$$
 (53)

2.6 Original and current gas in place and gas recovery

The conserved property M_T in the transport equation (Eq. (5)) consists of a free gas term M_1 and an adsorbed gas term M_2 , both in units of pressure due to scaling constant properties:

$$M_{1} = \phi \frac{pS_{g}}{z}, \quad M_{2} = \phi \frac{R_{g}T\rho_{a}}{M}(1 - S_{g}), \quad M_{T} = M_{1} + M_{2}$$
(54)

Gas currently in place (G_{CIP} with units of length times pressure) can be evaluated by integrating over the system. Similarly, gas initially in place G_{IIP} is G_{CIP} uniformly evaluated at the initial pressure p_0 :

$$G_{CIP} = \int_{x=0}^{L} (M_1 + M_2) dx$$

$$G_{IIP} = G_{CIP}(p_0) = L[M_1(p_0) + M_2(p_0)]$$
(55)

Gas recovery factor RF is the fraction of G_{IIP} that has been produced:

$$RF = 1 - \frac{G_{CIP}}{G_{IIP}} \tag{56}$$

It was shown by Andersen (2021a), for a similar system, that if the conserved property M_T is monotonous with pressure (which follows since higher pressure leads to storing more gas), the system can be expressed as a nonlinear diffusion equation with M_T as the independent variable. It was demonstrated theoretically that recovery *RF* against time then is proportional to the square root of time until the no-flow boundary is encountered.

If a given (well) pressure is allowed to become uniform, the highest recovery possible at that state is obtained, in terms of original gas in place O_{GIP} , called the ultimate recovery factor $RF_{\infty}(p)$:

$$RF_{\infty}(p) = 1 - \frac{G_{CIP}(p)}{O_{GIP}} = 1 - \frac{\phi(p) \left\{ \frac{pS_g(p)}{z(p)} + \frac{R_g T \rho_a}{M} [1 - S_g(p)] \right\}}{\phi_0 \left[\frac{p_0 S_{g0}}{z_0} + \frac{R_g T \rho_a}{M} (1 - S_{g0}) \right]}$$
(57)

Calculations of other average properties are shown in Appendix C. In the limit where the pressure goes to zero, the adsorbed layer thickness goes to zero, see Eq. (15), while pore radius stays finite, thus saturation goes to 1 and recovery goes to 1. For the special case where adsorption is negligible (*na*), i.e., $S_g = 1$, and gas is ideal z = 1, recovery is given by:

$$RF_{\infty}^{na}(p) = 1 - \frac{p}{p_0} \frac{\phi(p)}{\phi_0}$$
 (58)

If also pore compressibility is negligible (*na*,*nc*):

$$RF_{\infty}^{na,nc}(p) = 1 - \frac{p}{p_0} \tag{59}$$

demonstrating a linear relation between recovery and pressure.

2.7 Implementation

The final model consists of the transport Eq. (5), coupled with the relations for porosity (Eq. (14)), viscosity (Eq. (B-1)) gas compressibility factor (Eq. (A-4)), radius (Eq. (13)), effective radius (Eq. (17)), intrinsic permeability (Eq. (42)), and apparent permeability (Eq. (50)). The system was solved by operator splitting into a 'Free gas transport' system with fixed-in-time spatial distributions of porosity, saturation and permeability, and a 'Redistribution' system where these properties were updated locally to satisfy the constraint equations and preservation of mass. The simulation time T_{total} was divided into equal splitting time steps ΔT ; the time running 'Free gas transport' before a 'Redistribution'. Appendix D provides the details and discretization for this process. The numerical procedure was implemented in MATLAB. The xaxis was discretized into 50 equal cells and ΔT was selected to 12 hrs, which was sufficient for convergence, see Appendix E. Numerical validation against a commercial software is presented in Appendix E.

Constant parameters	Value	Units
*Well pressure, p _{well}	2.75×10^{6}	Ра
*Initial pressure, p_0	27.5×10^{6}	Ра
Gas constant, R_g	8.314	J/mol·K
Matrix length, L	10	m
Initial pore radius, R_0	4	nm
*Temperature, T	355.37	Κ
Initial tortuosity, τ_0	2.197	-
Gas molar weight, M	16.04×10^{-3}	kg/mol
*Initial porosity, ϕ_0	0.06	-
*Ads. gas density, ρ_a	343.8	kg/m ³
*Langmuir pressure, p_L	8.11×10^{6}	Pa
Varied parameters (base value)	Value	Units
*Pore compressibility, c_k	1.55×10^{-9}	1/Pa
Geometry parameter, n	2	-
Max ads. thickness, ΔR_{max}	0.8	nm
*Langmuir volume, V _L	0.2338	mmol/g
Numerical parameters	Value	Units
Grid Cells, N_x	50	-
Splitting step, ΔT	12	hrs.
Simulation time, T _{total}	30000	d

 Table 1. Input parameters used for reference case simulations.

* from Pang et al. (2017).

3. Results and discussion

3.1 Model input

The model input reference parameters are shown in Table 1. They are taken from Pang et al. (2017) who quantified the impact of pore compressibility and adsorption on porosity based on experiments on shale from the Yanchang formation in the Ordos basin. A fixed adsorbed phase density is assumed of 343.8 kg/m³, as reported by Pang et al. (2017). Perez and Devegowda (2017) obtained a similar value from molecular simulations, while Chen et al. (2019) reported a wider range $(150-660 \text{ kg/m}^3)$, but at more varying experimental conditions. Reference ΔR_{max} was selected as 0.8 nm, and low and high values were taken as 0.4 and 2 nm. The low to high values of $\Delta R_{\rm max}$ corresponded to initial mass fractions of adsorbed gas (M_2^0/M_T^0) of 0.27, 0.47 and 0.78, similar to observations in the literature; 20% to 85% (Montgomery et al., 2005; Ambrose et al., 2012). The ΔR_{max} range is also comparable to measured methane adsorption thicknesses between 0.25 and 1.5 nm reported by Chen et al. (2019) for Longmaxi shale and the molecular diameter of methane which is 0.38 nm, often applied as monolayer adsorption thickness. Reference n was set to 2 to indicate cylindrical pores. The match between our

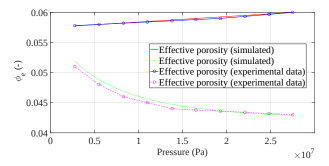


Fig. 2. Comparison of experimental effective porosity against pressure from Pang et al. (2017) and our model. The top curves show porosity reduction in presence of non-adsorbing gas, while the lower curves show porosity change when adsorption occupies a significant amount of the pore space.

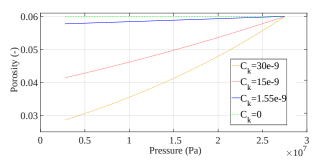


Fig. 3. Porosity as function of pressure for different compressibilities (in units of Pa^{-1}).

model and the data from Pang et al. (2017) is shown in Fig. 2, where the pore compressibility captures the decline in porosity on pressure depletion with non-adsorbing gas present, and adding the adsorption model captures how porosity actually increases from a low value on pressure depletion since desorption opens the pore space.

In the following, unless otherwise stated, all parameters take reference values. The initial porosity ϕ_0 and pore radius R_0 are set independently and the same in all example cases, but once set, they vary dependently. As indicated in the table, the main varied parameters are the pore compressibility c_k , the geometry parameter *n* and the max adsorption layer thickness ΔR_{max} . The initial pore volume will be the same in all cases, but it may vary how much is initially filled by adsorbed phase.

3.2 Static model behavior

3.2.1 Porosity relations

Fig. 3 shows the effect of compressibility on porosity. Higher compressibility results in greater porosity decline for a given pressure depletion. The reference compressibility of 1.55×10^{-9} Pa⁻¹ from Pang et al. (2017) reduces the porosity from 0.06 to 0.0575. For illustration this and higher values will be used to show how compressibility and adsorption interact, but it is noted that shales such as Marcellus, Barnett and Chinsui have lower compressibility around $0.1 - 0.8 \times 10^{-9}$ Pa⁻¹ (Dong et al., 2010; Yu and Sepehrnoori, 2014).

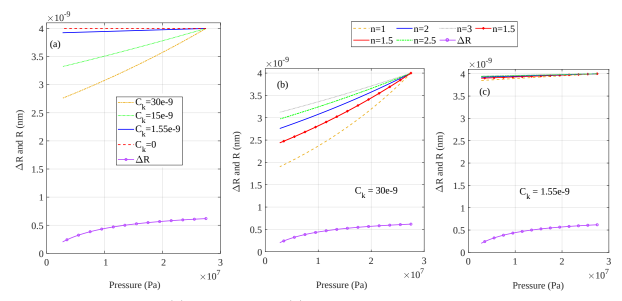


Fig. 4. Adsorbed layer thickness $\Delta R(p)$ and pore radius R(p) vs pressure for different values of compressibility (in Pa⁻¹) (a), and variations of geometry parameter *n* for a high compressibility (b) and low compressibility (c).

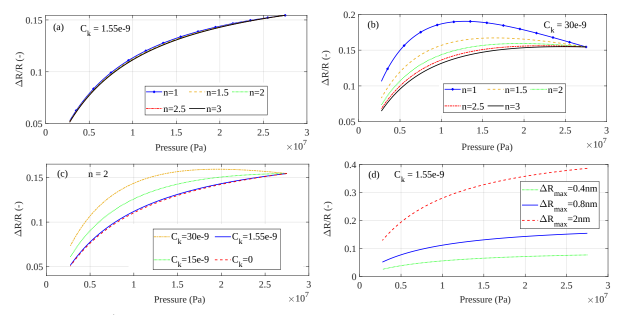


Fig. 5. The fraction $\Delta R/R$ vs pressure for variations of geometry parameter *n* for a low compressibility (a) and high compressibility (b), variation of compressibility (c) and ΔR_{max} (d). Note that the y-axis scales are not identical.

3.2.2 Pore radius and adsorption layer

In Fig. 4(a) it is observed that higher compressibility results in greater reduction of pore radius at a given pressure, which follows from Eqs. (13) and (14). In Fig. 4(b), *n* is varied using the base compressibility 1.55×10^{-9} Pa⁻¹. The small changes of *R* with *n* reflect that the radius (and porosity) stays near their original values. Changes from compressibility are then small irrespective of geometrical differences. When a high compressibility of 30×10^{-10} Pa⁻¹ is used in Fig. 4(c), the radius reduces and varies more with *n*. The radius reduces the most when *n* is low. In other words, the same reduction in porosity results in greater reduction of pore 'radius' when the pores are more planar (*n* close to 1) than spherical (*n* close to 3). The adsorption layer thickness is defined uniquely by pressure according to the Langmuir relation (Eq. (15)) and does not change with c_k or n. Radius appears to change close to linearly with pressure for different n and c_k , while layer thickness changes nonlinearly: Slowly at high pressures and more rapidly at low pressures.

The fraction $\Delta R/R$ is plotted vs pressure for variations of geometry parameter *n* with low and high compressibility (Figs. 5(a) and 5(b), respectively), variation of compressibility c_k (Fig. 5(c)), and ΔR_{max} (Fig. 5(d)). From the above considerations, for a given *n* (geometry) the layer fills a greater fraction $\Delta R/R$ with a higher compressibility than a low. And for a given compressibility, the layer fills a greater fraction $\Delta R/R$

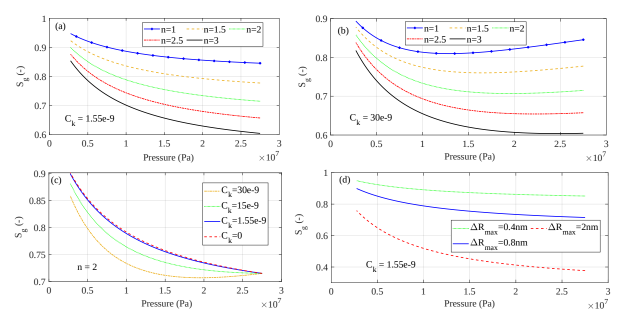


Fig. 6. Free gas saturation S_g vs pressure for variations of geometry parameter *n* for a low compressibility (a) and high compressibility (b), variation of compressibility (c) and ΔR_{max} (d). Note that the *y*-axis scales are not identical.

for a low n than a high n.

The fraction $\Delta R/R$ reduces towards low pressures for small changes in pore radius and greater reduction in adsorbed layer thickness. At zero compressibility (red curve in Fig. 5(c)) the radius is constant, while for more compressible cases, the radius changes linearly. As long as the radius changes little compared to the initial radius, the behavior is comparable to the incompressible case. For high compressibilities the pore radius can reduce more rapidly than the adsorbed layer at high pressures, but slower at low pressures. There is then a peak in how much the layer occupies of the pore radius at intermediate pressures (orange line in Fig. 5(c)). The effect of geometry comes more into play in highly compressible shale, see Fig. 5(b). High *n* maintains higher pore radius with pressure depletion, and thus the adsorbed layer fills a lower fraction of the radius at low pressure than low *n* cases. When maximum adsorption layer thickness is increased the fraction $\Delta R/R$ is generally increased, see Fig. 5(d).

3.2.3 Free gas saturation

Fig. 6 plots free gas saturation S_g as function of pressure using Eq. (18) for the aforementioned cases. The free gas saturation naturally has an opposite trend with pressure as the fraction $\Delta R/R$ but the relation is nonlinear according to the exponent *n*, see Eq. (18).

Pressure depletion reduces the fraction $\Delta R/R$ for low compressible cases which causes gas saturation to increase. Increased compressibility (Fig. 6(c)) increases the fraction $\Delta R/R$ and hence lowers the free gas saturation systematically. *n* not only affects the radius, but also influences how saturation is related to $\Delta R/R$. For example, in a round pore (high *n*) the same adsorbed layer thickness ΔR takes more of the volume than in a fracture-like pore (low *n*) for same porosity since the layer covers the radial interval taking most volume. Hence, for increased *n* the free gas saturation reduces quickly. Especially, the relations $\Delta R/R$ which were invariant with n at low compressibility (Fig. 5(a)) and varying little with high compressibility (Fig. 5(b)), are equivalent to saturation relations that vary significantly with n, see Figs. 6(a) and 6(b). The reduction of saturation with n includes the initial saturation which is natural since the radius and layer thickness are defined by the pressure, while the volume fraction depends on the pore geometry. Finally, increased ΔR_{max} reduces the free gas saturation (Fig. 6(d)).

3.2.4 Effective porosity

The effective porosity, which measures the volume fraction occupied by free gas, is shown in Fig. 7. In low compressible cases the desorption causes it to increase with pressure depletion, while in compressible dominated cases it can reduce. High n cases have the lowest effective porosity, with clear impact also for low compressibility cases.

3.2.5 Gas recovery

Recovery is shown vs pressure based on Eq. (57) in Fig. 8. Higher c_k (Fig. 8(c)) reduces the pore volume more at a given pressure and yields higher recovery. Although the adsorbed layer thickness is a unique function of pressure, the amount free gas is proportional to the remaining pore volume, which is reduced. Hence, recovery increases with compressibility.

Adsorbed gas is not produced significantly until low pressures. Recovery therefore generally decreases when the adsorbed fraction increases, e.g., when ΔR_{max} increases (Fig. 8(d)). More spherical pores (higher *n*) have a higher volume content of adsorbed gas (lower gas saturation, see Figs. 6(a) and 6(b)). The adsorbed mass fraction is even higher due to the density. Higher *n* therefore also results in lower recovery (Figs. 8(a) and 8(b)) at a given pressure.

For low n, a lower fraction gas is in adsorbed form (Figs. 6(a) and 6(b)). If porosity does not vary greatly and the

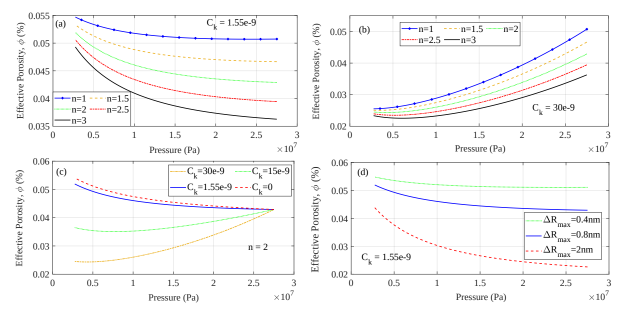


Fig. 7. Effective porosity ϕ_e vs pressure for variations of geometry parameter *n* for a low compressibility (a) and high compressibility (b), variation of compressibility (c) and ΔR_{max} (d). Note that the y-axis scales are not identical.

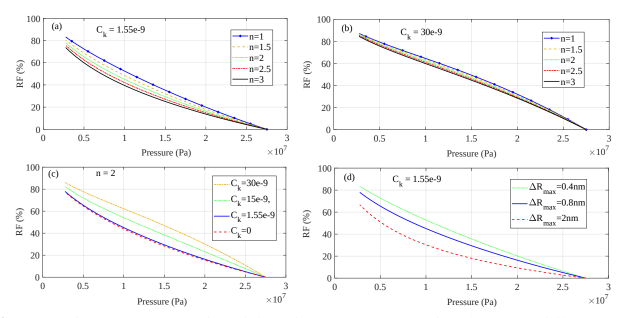


Fig. 8. Recovery factor RF vs pressure for variations of geometry parameter *n* for a low compressibility (a) and high compressibility (b), variation of compressibility (c) and ΔR_{max} (d).

adsorbed fraction is low, the relation between recovery and pressure becomes more linear in accordance with Eqs. (59) and (57), see Fig. 8(a).

3.2.6 Permeability

Intrinsic permeability is plotted against pressure in Fig. 9 for the presented cases. As noted from Eq. (42) it is a function of effective radius, effective porosity (where tortuosity is included) and geometry, as described by the parameters R_e , ϕ_e , *n*. Higher *n* shifts more of the volumetric flow towards the pore noflow boundary, meaning the intrinsic permeability is reduced (Figs. 9(a) and 9(b)). High compressibility reduces the pore radius at low pressures which reduces permeability (Fig. 9(c)). Desorption makes more of the pore space available for flow and increases permeability at low pressure, especially visible in less compressible cases (Fig. 9(d)).

The same plots are shown for apparent permeability in Fig. 10. Slip tends to increase permeability at low pressure and low effective radius which gives significant contributions in nanosize pores (Darabi et al., 2012). The apparent permeability is increasing with lowered pressure in all example cases, except in the high compressibility cases where reduction in intrinsic permeability is more dominating.

In supplementary material (SM) Fig. 1 apparent permeability is plotted against effective porosity for the considered pressures. In low compressibility cases (SM Figs. 1(a) and

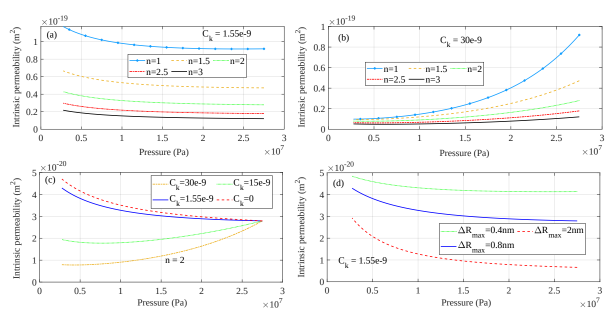


Fig. 9. Intrinsic permeability vs pressure for variations of geometry parameter *n* for a low compressibility (a) and high compressibility (b), variation of compressibility (c) and ΔR_{max} (d).

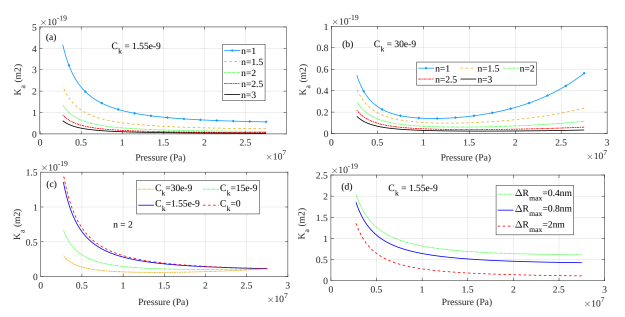


Fig. 10. Apparent permeability vs pressure for variations of geometry parameter *n* for a low compressibility (a) and high compressibility (b), variation of compressibility (c) and ΔR_{max} (d).

1(d)) the trend is monotonous and controlled by the increased effective radius R_e from desorption at constant pore radius R. Lower apparent permeability then correlates with lower effective porosity. For compressible cases however (SM Figs. 1(b) and 1(c)), the effective porosity first reduces and then increases with lowered pressure. The apparent permeability also reduces at high pressure and increases at low pressures, but the mechanism is also related to slip which results in non-unique porosity-permeability relations.

3.3 Dynamic simulation results

In this section, the dynamic interplay of all properties discussed in the previous section will be examined using transient simulations. The reference values are applied including an initial pore radius of 4 nm, initial porosity 0.06 and system length 10 m.

3.3.1 Introduction

Fig. 11 shows the effect of compressibility on gas production by illustrating the evolution of recovery factor (a), average porosity (b), average free gas saturation (c) and average apparent permeability (d). These parameters are plotted against the square root of time for 30000 d.

Higher compressibility reduces permeability (Fig. 11(d)) and delays recovery, but the end recovery is higher (Fig. 11(a)). Plotted against the square root of time, recovery is completely

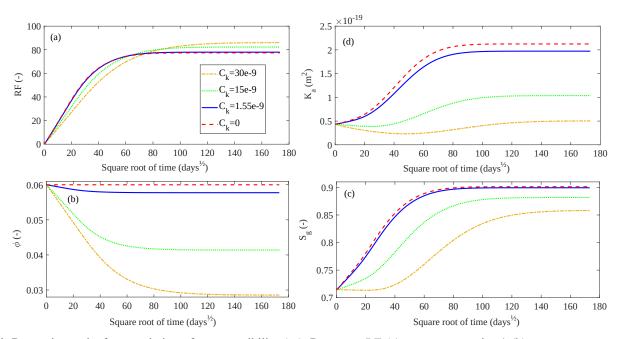


Fig. 11. Dynamic results from variation of compressibility (c_k) : Recovery *RF* (a), average porosity ϕ (b), average gas saturation S_g (c) and average apparent permeability (d).

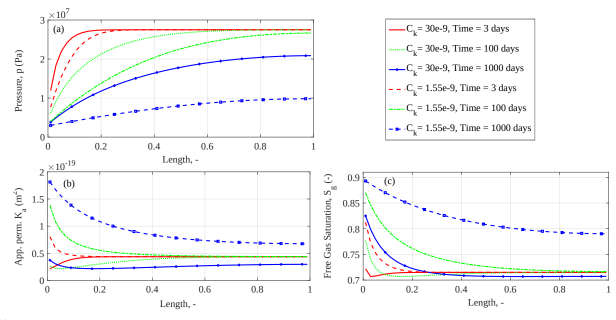


Fig. 12. Distributions of pressure p (a), apparent permeability K_a (b) and free gas saturation S_g (c) for a high compressibility (solid lines) and the low base case compressibility (dashed lines).

linear at early time as predicted in Andersen (2021a). Porosity declines more with higher compressibility (Fig. 11(b)). The higher compressibility gives smaller pore radius, but same adsorption layer thickness and hence generally lower free gas saturation (Fig. 11(c)).

Spatial distributions are presented of pressure (Fig. 12(a)), apparent permeability (Fig. 12(b)) and free gas saturation (Fig. 12(c)) at different times for a high and low (base) compressibility. The pressure profile has a steep gradient initially which reduces at later times (Fig. 12(a)). The cases with high compressibility are delayed as the pore space deformation acts as a buffer. As the well indicates a fixed pressure at x = 0, all the spatial profiles converge at x = 0 to properties determined by the well pressure (a finer grid would show this even clearer). For low compressible cases, the highest apparent permeability is found at the well with values decreasing monotonously into the system (dashed lines in Fig. 12(b)). The apparent permeability is increased by slip and increased gas saturation (dashed lines in Fig. 12(c)) at low pressures near the well. For high compressibility cases the net effect of reduced pressure is a lower apparent permeability due to pore radius reduction, despite that gas saturation and slip increase.

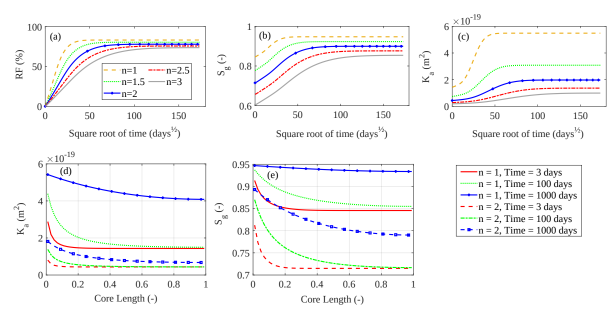


Fig. 13. Dynamic results from variation of geometry parameter *n*. Recovery *RF* (a), average gas saturation S_g (b) and average apparent permeability K_a (c) plotted against square root of time. Spatial profiles are given of apparent permeability (d) and free gas saturation (e) for the cases n = 1 (solid lines) and n = 2 (dashed lines).

3.3.2 The role of n-sphere geometry

The effect of the geometrical parameter n is investigated regarding the evolution of RF (Fig. 13(a)), S_g (b) and K_a (c) over 30000 days of production. The base compressibility 1.55×10^{-9} Pa⁻¹ was used, resulting in minor changes of porosity. The pore radius is then close to constant and as noted in Fig. 5(a) the fraction $\Delta R/R$, and hence the effective radius R_e , reduces towards lower pressure with little dependence on n. On the other hand, there is more gas adsorbed per volume and thus lower free gas saturation at high n, see Fig. 13(c). The intrinsic permeability is considered proportional to S_{g} in Eq. (42) through ϕ_e (note that R_e stays the same at a given pressure) and high n cases therefore have lower intrinsic permeability (Fig. 13(d)) and slower recovery (Fig. 13(a)) than low n cases. Higher n cases also reach lower end recovery as more of the gas is adsorbed and difficult to produce. Fig. 13 also illustrates distributions of apparent permeability (d) and gas saturation (e) for two cases where solid lines represent n = 1 and dashed lines represent the base case n = 2. Faster recovery in the low *n* case can be linked to the higher free gas saturations and apparent permeabilities.

3.3.3 Role of maximum adsorption layer thickness

The effect of maximum adsorption layer thickness is investigated for three cases $\Delta R_{\text{max}} = 0.4$, 0.8 and 2 nm corresponding to initial adsorbed gas fractions 0.27, 0.47 and 0.78. Fig. 14 shows the results in terms of RF (a), free gas saturation S_g (b) and apparent permeability (c). The base compressibility yields negligible porosity changes. Both higher end recovery and recovery at a given time are achieved for low ΔR_{max} , i.e., when more of the shale gas is in free form (green curve). With low ΔR_{max} the pores are more open for flow as R_e and S_g are larger, giving higher apparent permeability. Spatial distributions are plotted for two cases $\Delta R_{\text{max}} = 0.8$ and 2 nm. When ΔR_{max} is low, the free gas saturation (d) is higher and so is the apparent permeability (e). Accordingly, recovery goes faster for low ΔR_{max} . Note that the change in free saturation is higher when ΔR_{max} is high (from initially 0.38 to 0.75 vs from 0.72 to 0.9), The reason is that for low ΔR_{max} the desorption occurs in the outermost layers with most volume and hence most impact on saturation.

3.3.4 The importance of compressibility and adsorption layer

Fig. 15 illustrates the evolution of recovery factor for four different cases based on different influence of compressibility and desorption. All cases have same initial conditions of pore radius, porosity and adsorbed layer thickness as given by the reference parameters. Two cases use a zero compressibility (red and orange curves), while the other two (blue and green curves) use a high compressibility of 30×10^{-9} Pa⁻¹. Further, two cases (green and orange curves) consider desorption of the adsorbed layer with pressure depletion (as before), while the other two (blue and red curves) assume the layer is not desorbed but remains at its initial thickness regardless of pressure depletion, i.e., $\Delta R = \Delta R(p_0) = 0.8$ nm.

For these cases, the (reference) initial adsorbed mass fraction is 47%. The lowest end recovery (orange curve) is obtained when gas does not desorb and the pores are not compressed. If the pores are compressed but gas does not desorb (green curve) the recovery is slowed down by permeability reduction, but end recovery is increased.

The fastest recovery is seen when gas desorbs and the pores are not compressed (red curve). Then permeability is increased by reduction in the adsorbed layer thickness and the pore radius is maintained. However, the highest end recovery is seen when accounting for both desorption and pore compression

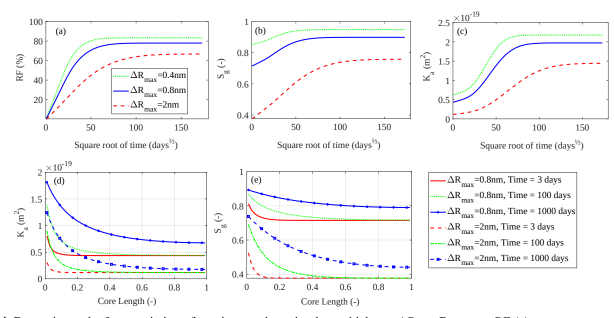


Fig. 14. Dynamic results from variation of maximum adsorption layer thickness ΔR_{max} : Recovery *RF* (a), average gas saturation S_g (b) and average apparent permeability (e) are plotted against the square root of time. Spatial distributions are given of apparent permeability K_a (d) and free gas saturation S_g (e) for the cases $\Delta R_{\text{max}} = 0.8$ nm (solid lines) and $\Delta R_{\text{max}} = 2$ nm (dashed lines).

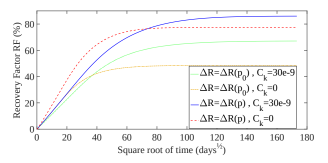


Fig. 15. Recovery factor RF vs time for four cases: Zero compressibility with desorption (red) and without desorption (orange) and high compressibility of 30×10^{-9} Pa⁻¹ with desorption (blue) and without desorption (green).

(blue curve). Then adsorbed gas is produced and pore volume is compacted taking out more of the free gas. However, the reduction of pore radius reduces permeability and slows down recovery compared to the incompressible case.

4. Conclusions

A mathematical model has been derived for shale gas production considering the interplay of free and adsorbed gas in *n*-spherical pores where the adsorbed phase layer has a thickness of comparable dimension as the pore radius. A higher *n* indicates more spherical pores, while a lower *n* means more fracture shaped pores. The role of geometric and intrinsic properties were investigated under static and dynamic conditions and a consistent model was presented for variation between pore radius, adsorption layer thickness, porosity and apparent permeability. The permeability model yields the established expressions for a cylinder or fracture as special cases. The main findings are:

· For the same porosity and adsorbed layer thickness (same

pressure), in more spherical shaped pores (higher n) a greater volume fraction of the pores is filled by the adsorbed gas since it is attached to the parts of the pores where a layer takes more volume.

- Pressure depletion causes a reduction in the adsorbed layer thickness, and if the shale is compressible, also the pore radius. The change in adsorbed layer thickness with pressure is low at high pressure and greater at low pressure. Pore radius changes more linearly with pressure. The free gas saturation can thus increase with pressure depletion (time) for low compressibility cases, or reach a minimum before increasing in high compressible cases.
- A larger fraction adsorbed gas results in reduced permeability by narrower pores, less linear relation between recovery and pressure and lower end recovery fraction. This is due to the low desorption taking place at high pressures compared to low pressures. Mainly free gas is produced at early times. Systems with higher adsorbed gas fraction see lower recovery at a given pressure.
- Apparent permeability increased monotonously towards the well where the pressure was lowest for incompressible cases. That is because desorption removed more of the adsorbed layer and because of higher slip factor with lower pressure. For compressible cases, the reduction in pore radius near the well could be more dominating and lead to a net reduction in permeability towards the well.
- Compressibility and desorption can both affect end recovery and recovery rate. Higher end recovery is seen when accounting for desorption and compressibility and vice versa. The fastest recovery is seen when accounting for desorption and not compressibility as the effective pore radius and permeability increases.

Acknowledgement

Andersen acknowledges the Research Council of Norway and the industry partners, ConocoPhillips Skandinavia AS, Aker BP ASA, Vår Energi AS, Equinor ASA, Neptune Energy Norge AS, Lundin Norway AS, Halliburton AS, Schlumberger Norge AS, and Wintershall DEA, of The National IOR Centre of Norway for support.

Conflict of interest

The authors declare no competing interest.

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Appendix A: Calculation of real gas density using the Peng-Robinson equation of state

The Peng and Robinson (1976) equation of state is expressed below in terms of gas molar volume V_m :

$$p = \frac{R_g T}{V_m - b} - \frac{a(T)}{V_m (V_m + b) + b(V_m - b)}$$
(A-1)

$$a = \frac{R_g^2 T_c^2}{2.1870 p_c} \left[1 + k \left(1 - T_r^{0.5} \right) \right]^2, \quad b = \frac{R_g T_c}{12.8535 p_c}, \quad T_r = \frac{T}{T_c}$$
(A-2)

$$k = 0.37464 + 1.5422\omega - 0.26992\omega^2 \tag{A-3}$$

where p_c and T_c are critical pressure and critical temperature respectively, T_r is reduced temperature and ω is the dimensionless acentric factor. Eq. (A-1) can also be expressed with the compressibility factor z as a cubic equation:

$$z^3 + c_2 z^2 + c_1 z + c_0 = 0 \tag{A-4}$$

$$c_2 = -1 + B, \quad c_1 = A - 3B^2 - 2B, \quad c_0 = -AB + B^2 + B^3$$
 (A-5)

$$A = \frac{pT_c^2}{2.1870p_c T^2} \left[1 + k \left(1 - T_r^{0.5} \right) \right]^2, \quad B = \frac{pT_c}{12.8535p_c T}$$
(A-6)

By finding the roots of Eq. (A-4) the compressibility factor z can be calculated.

Appendix B: Calculation of gas viscosity

The gas viscosity correlation by Lee et al. (1964) was applied, which was modified from Starling and Ellington (1964) to represent mixtures and pure components:

$$\mu = 10^{-7} K \exp\left[X(T)\rho_g^{Y(T)}\right]$$
(B-1)

$$K = \frac{7.77 + 0.0063M) \left(\frac{5}{9}T\right)^{1.5}}{122.4 + 12.9M + \left(\frac{5}{9}T\right)}$$
(B-2)

$$X = 2.57 + \frac{1914.5}{\left(\frac{5}{9}T\right)} + 0.0095M, \quad Y = 1.11 + 0.04X$$
(B-3)

where T is absolute temperature in Kelvin and M is molecular weight.

Appendix C: Average properties

Average values of porosity, gas saturation and gas pressure are given as:

$$\overline{\phi} = \frac{1}{N_x} \sum_{i=1}^{N_x} \phi_i, \quad \overline{S}_g = \frac{\sum_{i=1}^{N_x} S_{gi} \phi_i}{\sum_{i=1}^{N_x} \phi_i}, \quad \overline{p} = \frac{\sum_{i=1}^{N_x} p_i \frac{S_{gi} \phi_i}{z_i}}{\sum_{i=1}^{N_x} \frac{S_{gi} \phi_i}{z_i}}$$
(C-1)

The average 'mass' of free and adsorbed gas at a given time are:

$$\overline{M}_{1} = \frac{1}{N_{x}} \sum_{i=1}^{N_{x}} \frac{p_{i} S_{gi}}{z_{i}} \phi_{i}, \quad \overline{M}_{2} = \frac{R_{g} T \rho_{a}}{M} \frac{1}{N_{x}} \sum_{i=1}^{N_{x}} \phi_{i} \left(1 - S_{gi}\right)$$
(C-2)

Appendix D: Operator splitting and discretization

The system in Section 2 is solved using an operator splitting approach (see Berawala and Andersen (2020a, 2020b) for examples of this approach in the shale gas context) by dividing the full system into more easily solved subsystems. The transport equation (Eq. (5)) is written with the notation:

$$\partial_t \left\{ \phi \left[\frac{p}{z} S_g + \hat{a} \left(1 - S_g \right) \right] \right\} = \partial_x \left(D \partial_x p \right), \quad \hat{a} = \frac{R_g T \rho_a}{M}, \quad D(p) = \frac{p K_a}{z \mu}$$
(D-1)

where \hat{a} is a constant and D is pressure and saturation dependent. This system is split into a 'Free gas transport' system where only free gas flow is considered and the adsorbed phase and porous medium are held undeformed; and a 'Redistribution' system where free gas, adsorbed gas and porosity are adjusted locally to meet equilibrium pressure constraints.

D.1: Free gas transport

For this system the spatial distributions of $S_g(x)$, $\phi(x)$, D(x), z(x) are kept fixed with time during the splitting step $(t, t + \Delta T)$. The following equation then needs to be solved:

$$\frac{S_g(x)\phi(x)}{z(x)}\partial_t(p) = \partial_x\left(D(x)\partial_x p\right) \tag{D-2}$$

The matrix is divided into N_x cells and an explicit discretization of Eq. (D-1) is used:

$$\left(\frac{S_g}{z}\phi\right)_i \frac{p_i^{j+1} - p_i^j}{\Delta t} = \frac{(D\partial_x p)_{i+1/2} - (D\partial_x p)_{i-1/2}}{\Delta x}, \quad (i = 1:N_x)$$
(D-3)

where i is the grid cell index and j the time step. For central boundaries the fluxes are defined by:

$$(D\partial_x p_g)_{i+1/2} = \frac{2}{\frac{1}{D_{i+1}} + \frac{1}{D_i}} \frac{p_{i+1}^J - p_i^J}{\Delta x}, \quad (D\partial_x p_g)_{i-1/2} = \frac{2}{\frac{1}{D_i} + \frac{1}{D_{i-1}}} \frac{p_i^J - p_{i-1}^J}{\Delta x}$$
(D-4)

while the fluxes at the outer boundaries are given as:

$$(D\partial_x p)_{1/2} = \frac{2}{\frac{1}{D_1} + \frac{1}{\infty}} \frac{p_1^J - p_{well}}{\frac{\Delta x}{2}}, \quad (D\partial_x p)_{N_x + 1/2} = 0$$
(D-5)

The time step Δt was selected sufficiently low to achieve stability of the solution.

D.2: Redistribution

While preserving the mass, gas is redistributed between free and adsorbed form and porosity is adjusted to the new pressure. No flow is assumed, giving:

$$\partial_t M_T = 0, \quad M_T = \phi \left[\frac{pS_g}{z} + \hat{a} \left(1 - S_g \right) \right]$$
 (D-6)

where M_T is the conserved property. Assume that the system was in equilibrium at an old state and that the transport solver was performed. Only the pressure in the first term (free gas) of M_T will then have changed, while the remaining terms remain evaluated at the old state. The local value of M_T is then:

$$M_{T} = \phi(p_{old}) \left\{ \frac{p_{new} S_g(p_{old})}{z(p_{old})} + \hat{a} [1 - S_g(p_{old})] \right\}$$
(D-7)

After redistribution, all the terms should be evaluated at the same adjusted pressure p_{adj} , while keeping the value of M_T (mass conservation). This pressure is found by solving:

$$f\left(p_{ad\,i}\right) = 0\tag{D-8}$$

where

$$f = \phi\left(p_{adj}\right) \left\{ \frac{p_{adj}S_g\left(p_{adj}\right)}{z\left(p_{adj}\right)} + \hat{a}\left[1 - S_g\left(p_{adj}\right)\right] \right\} - M_T$$
(D-9)

Appendix E: Convergence and numerical validation

The reference case was simulated using different ΔT and different N_x . Resulting recovery against the square root of time is shown in SM Figs. 2(a) and 2(b) where these parameters were varied, respectively. Keeping ΔT sufficiently low leads the solution to converge. The late time recovery does not depend much on ΔT , and only a minor impact can be noticed at early recovery for larger ΔT . Sufficient convergence was considered achieved at $\Delta T = 12$ hrs which was used as the reference value in our simulations. The model is also sufficiently converging for different N_x and only minor effects are noticeable during the simulation period.

The code was also validated numerically by running a case under identical conditions in both our code and the commercial reservoir simulation software Eclipse 100 (Geoquest, 2014). A relatively simple case was implemented using ideal gas, pore compressibility 1.55×10^{-9} Pa⁻¹, viscosity 0.02 cP. Adsorption and slip effects were ignored. The initial porosity was 0.06, and intrinsic permeability was set constant based on tortuosity $\tau = 2.197$, pore radius 4 nm and the initial porosity, resulting in 54.5 nD. As indicated in SM Fig. 3 our code is able to give same results as the commercial model under comparable conditions.