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The energy penalty of post-combustion CO₂ capture & storage and its implications for retrofitting the U.S. installed base

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A review of the literature has found a factor of 4 spread in the estimated values of the energy penalty for post-combustion capture and storage of CO₂ from pulverized-coal (PC) fired power plants. We elucidate the cause of that spread by deriving an analytic relationship for the energy penalty from thermodynamic principles and by identifying which variables are most difficult to constrain. We define the energy penalty for CCS to be the fraction of fuel that must be dedicated to CCS for a fixed quantity of work output. That penalty can manifest itself as either the additional fuel required to maintain a power plant’s output or the loss of output for a constant fuel input. Of the 17 parameters that constitute the energy penalty, only the fraction of available waste heat that is recovered for use and the 2nd-law separation efficiency are poorly constrained. We provide an absolute lower bound for the energy penalty of ~11%, and we demonstrate to what degree increasing the fraction of available-waste-heat recovery can reduce the energy penalty from the higher values reported. It is further argued that an energy penalty of ~40% will be easily achieved while one of ~29% represents a decent target value. Furthermore, we analyze the distribution of PC plants in the U.S. and calculate a distribution for the additional fuel required to operate all these plants with CO₂ capture and storage (CCS).

Introduction

Global carbon dioxide (CO₂) emissions have accelerated from 1.1%/yr in the 1990’s to over 3%/yr since 2000. Those continued growth rates would result in global CO₂ emissions of ~40 GtCO₂/yr and ~100 GtCO₂/yr by 2050, respectively. Stabilizing atmospheric CO₂ concentration below 550 ppm, however, requires emissions to essentially stay flat for the next 42 years. CO₂ capture and storage (CCS) is a promising technology that has the potential to address the ~40% of emission emanating from large-point sources such as power plants. CCS for existing plants involves separating the CO₂ from the plant’s flue gas, compressing the CO₂ for pipeline transport, and injecting the CO₂ into a geologic formation where it is intended to remain for millennia.

The U.S. has 1493 coal-fired power plants that constitute 336 gigawatts (GW) of rated power generation capacity. Nearly all of these plants involve pulverized-coal (PC) combustion, where the coal is pulverized such that over 98% of it is less than 300 μm, and then it is combusted in air at atmospheric pressure. In 2006, these plants composed about 70% of U.S. fossil-fuel derived electricity and about 50% of total electricity production. To produce that electricity, the plants burned ~930 million tonnes of coal and produced ~1.9 gigatonnes (Gt) of CO₂, about 1/3 of total U.S. emissions. In addition, PC power constitutes well over 90% of coal-fired power in the world. The dominance of PC power plants makes significant reduction in national or global

Broader context

We derive an analytic relationship for the energy penalty from thermodynamic principles, and we apply that relationship to the installed base of U.S. coal-fired power plants to determine the energetic requirements of retrofitting that base for CCS. Pulverized-coal (PC) facilities compose over 95% of the CO₂ produced by U.S. coal-fired power plants. It is unlikely that either national or global CO₂ emissions can be substantially reduced without either shutting down or retrofitting these plants for CCS. The economics of CCS from PC power plants depend, to a large degree, on the thermodynamic work required to capture and store the CO₂. The data demonstrate that—under reasonable assumptions—retrofitting the most efficient 10% of existing plants will offset 30% more CO₂ per unit of additional fuel than retrofitting the least efficient 10% of plants. Indeed, CCS on the least-efficient plants may not make economic sense compared with building new capacity and shutting down the least efficient plants. Finally, we show that a reduction in electrical power demand by between 15% and 20%, combined with retrofitting existing plants for CCS, would lower CO₂ emissions from electricity production by ~65% if the newly liberated power were used for CCS.
CO₂ emissions from power-plants dependent on either shutting down a substantial fraction of the existing PC plants or retrofitting those plants for post-combustion capture and storage.

Each of the CCS steps—separation, compression, transport, and storage—requires work. To perform that work, a fraction of the fuel input must be dedicated to CCS. That fuel requirement constitutes the CCS energy penalty. The energy penalty can be realized as either additional fuel input to maintain the baseline power output or as reduced power output for a constant fuel input. Several studies have estimated the energy penalty for PC plants, but these estimates differ between studies by nearly a factor of 4 (Fig. 1).²⁻¹⁸

In this paper, we calculate the thermodynamic work required for the various steps of CCS. We elucidate the cause of the spread in previously estimated energy penalties by deriving an analytic relationship for the energy penalty from first principles and by identifying which of the variables are most difficult to constrain. We show that the energy penalty associated with capturing and storing all the CO₂ generated by U.S. PC plants will require either burning an additional ~400–600 million tonnes of coal per year or building an additional ~100 GW of CO₂-free baseload power. In the latter scenario, the additional baseload power would be required to make up for the reduced power output of the retrofitted PC plants. It should be noted that separate end-use efficiency improvements could serve to offset the CCS energy penalty. In the discussion, we apply our analysis to the actual U.S. fleet of PC power plants. Since the energy penalty is a function of the power-plant’s baseline efficiency, then we derive the expected distribution of energy penalties that would result from retrofitting the U.S. installed base of PC plants.

Minimum work required to sequester CO₂

We first derive the lower bound for the work required to capture and store CO₂ from a PC plant. Throughout this section, we neglect frictional losses in both pipeline transport and resistance.

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Our analysis follows carbon through the process of oxidation to CO₂ in a power plant to storage in a geologic formation. Those losses are addressed later. For the calculation of this ideal limit, the work to store CO₂ is described in three steps (Fig. 2); (1) The work required to separate the CO₂ from the mixture of gases in power-plant flue gas; (2) the work to compress the CO₂ for transport and injection at hydrostatic pressure (pipeline pressure is typically ~14 MPa and hydrostatic pressure in reservoirs is roughly 1 MPa per 100 meters of depth);¹⁹ (3) the work required to emplace the CO₂ at depth, displacing denser groundwater upward. Each step is required to overcome particular physical barriers: separation overcomes entropy; compression to pipeline and hydrostatic pressure overcomes pressure; and emplacing beneath the groundwater overcomes gravity and surface tension.

Our analysis follows carbon through the process of oxidation to CO₂ in a power plant to storage in a geologic formation. As such, we have defined 6 different chemical and physical states during this process:

State 0: Reduced carbon in the plant’s combustor
State 1: Dilute CO₂ mixed with N₂ and H₂O in the flue gas
State 2: Concentrated N₂ stream at low pressure that has been separated from the flue gas
State 3: Concentrated CO₂ stream at low absolute pressure exiting the separation process

Fig. 1 (A) Published values for the additional fuel required to maintain constant electric with CCS and the cost of CO₂ avoided in 2007 US dollars²⁻¹⁸ for post-combustion capture and storage from pulverized coal plants. The blue diamonds are for new construction projects, the red squares are for retrofits, and the black triangles are for retrofits with boiler upgrades. The horizontal intercept—labeled N/A—is for three studies that estimated the energy penalty but not the cost of the CO₂ avoided. (B) Values from the same studies of the CCS energy penalty and the levelized cost of electricity in constant 2007 dollars from the CCS power plant. Note that while the cost of CO₂ avoided is much higher for PC retrofits than for new projects, the levelized cost electricity is essentially the same in both cases.
State 4: Concentrated CO₂ stream compressed for injection at the surface
State 5: Concentrated CO₂ stream emplaced beneath the pore water in the geologic formation

The process of sequestering CO₂ requires the input of work to transfer the system from state 1 to states 2 and 3, from state 3 to state 4, and from state 4 to state 5. We have labeled the work required for these three transitions: $W_a$, $W_b$, and $W_c$.

### $W_a$ (from state 1 to states 2 and 3)

**Separate the CO₂ from the flue gas.** Separating the CO₂ from flue gas is justified because separating and venting gases other than CO₂ back to the atmosphere reduces the net sequestration work by lowering the compression and injection costs. In the discussion, we calculate the optimal separation fraction that minimizes the total primary energy requirement for a range of efficiencies.

Flue gas emitted from typical coal-burning power plants contain $\sim 78\%$ N₂ from the atmosphere, $\sim 15\%$ CO₂ from the oxidation of the carbon in the hydrocarbon, and $\sim 7\%$ water from both the oxidation of hydrogen in the coal and the vaporization of water that was adsorbed on the coal. Removing H₂O from the flue gas is—in principle—thermodynamically favorable because H₂O condenses at surface conditions indicating the enthalpy change of separation is not zero. In the thermodynamic limit, the minimum work to separate the flue gas into one concentrated CO₂ stream and one concentrated N₂ stream is the difference in the thermodynamic availability before and after separation. For an isothermal and isobaric process, the work equals the change in free energy before and after separation:

$$W_{\text{min}} = -dG$$  \hspace{1cm} (1)

where $G$ is the Gibbs free energy.

For the separation, we employ the ideal gas assumption as the pressure is near atmospheric and N₂ and CO₂ do not chemically interact. The mole fraction of N₂ and CO₂ in the fully mixed state is $X_{N1}$ and $X_{C1}$. After the separation, the mole fraction in state 2 (low CO₂) are $X_{N2}$ and $X_{C2}$, and the mole fractions for state 3 (high CO₂) are $X_{N3}$ and $X_{C3}$.

The partial molar Gibbs energy for each gas in an ideal mixture is given by:

$$\frac{\partial G}{\partial n_i} = \frac{G_i^0 + RT \ln \left( \frac{P_i}{P} \right)}{n_i}$$

where $P_i$ is the partial pressure of the $i$th gas and $P$ is the total pressure. Thus, the total free energy of an ideal gas mixture is:

$$G_{\text{tot}} = \sum_i n_i \frac{\partial G}{\partial n_i}$$  \hspace{1cm} (3)

If we assume that none of the states are completely pure (i.e., $X_{ij} > 0$ for all $i$ and $j$), then the previous equation will determine the water table is independent of the size of the domain and the geometry of the injected CO₂ plume. If CO₂ were injected beneath twice the land area, then the change in potential energy would not change because a greater quantity of water would be lifted a corresponding smaller distance.
the free energy of each state. The minimum work to change a system’s state is given by the change in free energy between those states:

\[ W = \Delta G_{\text{sep}} = (G_2 + G_3) - G_1 \]  

(4)

To calculate the minimum work required to transfer the system from state 1 into the distinct states 2 and 3, we calculate the free energy of each state:

\[
G_1 = n_3G_0^2 + n_{N_1}G_0^N + RT(n_{C_1}\ln(X_{C_1}) + n_{N_1}\ln(X_{N_1}))
\]

\[
G_2 = n_2G_0^2 + n_{N_2}G_0^N + RT(n_{C_2}\ln(X_{C_2}) + n_{N_2}\ln(X_{N_2}))
\]

\[
G_3 = n_3G_0^2 + n_{N_3}G_0^N + RT(n_{C_3}\ln(X_{C_3}) + n_{N_3}\ln(X_{N_3}))
\]

(5)

So, the minimum required work is:

\[
W = (RT(n_{C_2}\ln(X_{C_2}) + n_{N_2}\ln(X_{N_2})) + RT(n_{C_3}\ln(X_{C_3}) + n_{N_3}\ln(X_{N_3}))) - RT(n_{C_1}\ln(X_{C_1}) + n_{N_1}\ln(X_{N_1}))
\]

(6)

And the work per mole of CO₂ in the flue gas is:

\[
W_a = \frac{1}{n_{C_1}}(RT(n_{C_2}\ln(X_{C_2}) + n_{N_2}\ln(X_{N_2})) + RT(n_{C_3}\ln(X_{C_3}) + n_{N_3}\ln(X_{N_3}))) - RT(n_{C_1}\ln(X_{C_1}) + n_{N_1}\ln(X_{N_1}))
\]

(7)

The number of parameters can be reduced by substituting the definition of the mole fraction into eqn (6), and the minimum work required to isothermally separate an ideal gas mixture into two ideal gas mixtures of different concentrations (\(W_a\)) is given by:

\[
W_a = RT\left[\frac{n_{C_2}\ln\left(\frac{n_{C_2}}{n_{C_2} + n_{N_2}}\right) + n_{N_2}\ln\left(\frac{n_{N_2}}{n_{C_2} + n_{N_2}}\right)}{n_{C_1} - n_{C_2} + n_{N_1} - n_{N_2}}\right]
\]

\[
+ (n_{C_1} - n_{C_2})\ln\left(\frac{n_{C_1} - n_{C_2}}{n_{C_1} - n_{C_2} + n_{N_1} - n_{N_2}}\right)
\]

\[
+ (n_{N_1} - n_{N_2})\ln\left(\frac{n_{N_1} - n_{N_2}}{n_{C_1} - n_{C_2} + n_{N_1} - n_{N_2}}\right)
\]

\[
- \left(n_{C_1}\ln\left(\frac{n_{C_1}}{n_{C_1} + n_{N_1}}\right) + n_{N_1}\ln\left(\frac{n_{N_1}}{n_{C_1} + n_{N_1}}\right)\right)
\]

(8)

where \(n\) is the number of moles of either N₂ or CO₂, subscripted N or C, in either the original mixture, the concentrated CO₂ stream, or the concentrated N₂ stream, subscripted 1, 2, and 3 respectively. For typical values, \(W_a\) equals \(\sim 9\) kJ/(mol CO₂).

\(W_b\) (from state 3 to state 4)

**Compress the concentrated stream at the surface.** To inject the concentrated CO₂ stream into a geologic formation, it must be compressed such that at the bottom of the well, its pressure equals the reservoir pore-pressure:

\[
P_a = P_3 - g \left[\int_0^{L_3} \rho_s(z)dz + \rho_w g L_4 - g \int_0^{L_4} \rho_s(z)dz\right]
\]

(9)

\(P_a\) is the pressure at the top of the well, where subscript 4 indicates the state of concentrated and compressed CO₂ at the surface. \(P_3\) is the pore-pressure at the bottom of the well, which is assumed to initially equal the product of the density of water (\(\rho_w\)), the gravitational constant (g), and the depth of injection (\(L_4\)). The integral in the second term accounts for gravitational compression and thermal expansion of the concentrated CO₂ stream within the borehole.

The minimum work required to compress the concentrated CO₂ stream from state 3 to state 4 (\(W_b\)) is given by the reversible isothermal compression:

\[
W_b = -\int_{v_1}^{v_4} (P(v, T) - P_a)dv
\]

(10)

where \(v\) is the molar volume of the concentrated CO₂ stream, \(P_3\) and \(T_3\) are the conditions at which the plant supplies the highly concentrated stream, \(P_4\) and \(T_4\) are the post compression conditions, and \(P_a\) is the atmospheric pressure, which assists in the compression. For typical values, \(W_b\) equals \(\sim 13\) kJ/(mol CO₂).

\(W_c\) (from state 4 to state 5)

**Push the compressed CO₂ into the formation.** Once the CO₂ at the surface is compressed to the pressure at the bottom of the borehole (\(P_a\)), it must be pushed out of the compressor and into the well, which causes CO₂ at the well screen to flow into the formation and vertically displace the ground-water. In the limit of zero friction, pushing the CO₂ into the formation requires work to vertically displace the groundwater (\(W_{c1}\)) and work to create the interfacial surface between the CO₂ and the pore water (\(W_{c2}\)).

The work required to vertically displace the ground water is equal to the volume of ground water displaced times the well-head pressure:

\[
W_{c1} = (P_4 - P_a)v_4
\]

(11)

In the limiting case of zero viscous drag, the minimum value of \(W_{c1}\) occurs when \(P_4\) equals the hydrostatic pressure of the ground water at the well-head. For typical values, \(W_{c1}\) equals \(\sim 1-2\) kJ/(mol CO₂).

In addition, work is required to create the interfacial surface between the CO₂ and the pore-water because, on the injection time-scale, the CO₂ acts primarily as an immiscible phase. The interfacial surface tension between supercritical CO₂ and water at the relevant conditions is \(\sim 0.02\) J/m², and the work required to increase the surface area goes as the interfacial surface tension (\(\gamma\)) and the change in surface area of the interface:

\[
W_{c2} = \gamma dA = \Delta V \Delta P_{\text{cap}}
\]

(12)

\(W_{c2}\) is the work required to overcome the capillary force; where \(\Delta V\) is the total swept out pore volume, and \(\Delta P_{\text{cap}}\) is the capillary pressure:

\[
\Delta P_{\text{cap}} = \gamma \left(\frac{1}{R_1} + \frac{1}{R_2}\right)
\]

(13)

\(R_1\) and \(R_2\) are the radii of curvature of the surface, and for a typical sand-reservoir, the pore sizes are on the order of
The molar volume of CO$_2$ at reservoir conditions is $10^{-4}$–$10^{-5}$ m$^3$. Therefore, the capillary pressure is on the order of 2000 Pa, and $W_{c2}$ is on the order of 0.2 J/(mol CO$_2$). Since $W_{c1}$, on the other hand, is on the order of $10^3$ J/(mol CO$_2$), then $W_{c2}$ can be safely ignored.

The total compression work is the sum of $W_b$ and $W_c$, where $W_b$ is the sum of $W_{c1}$ and $W_{c2}$. By the reverse integration by parts:

$$W_{bc} = W_b + W_c = \frac{1}{\gamma-1} (P_v(T) - P_a)v_4 + (P_4 - P_a)v_4$$

$$= \int_{P=P_3}^{P_4} v(P, T)\,dp$$

(14)

$W_{bc}$ is minimized under isothermal conditions, but regardless of the final temperature, the minimum work input required to achieve this lower bound injection pressure equals the change in thermodynamic availability of the concentrated CO$_2$ stream during compression:

$$W_{bc} = n_C((h_{C4} - h_{C3}) - T_0(s_{C4} - s_{C3})) + n_N((h_{N4} - h_{N3}) - T_0(s_{N4} - s_{N3}))$$

(15)

where $h$ and $s$ are the molar enthalpies and entropies, respectively. The subscripts, 3 and 4 indicate the surface conditions before and after compression, and $T_0$ is the ground state temperature. Integration of eqn (14) from $P_3$ to $P_4$ will always equal the state change of the thermodynamic availability (eqn (15)) as shown in Fig. 3.

A different approach to ascertain the work required to emplace the CO$_2$ below the pore water is to calculate the change in gravitational potential energy between state 4 and state 5. The total turbine shaft work is then the sum of $W_b$ and the change in gravitational potential energy:

$$W_{bc} = W_b + \Delta U = W_2 + gL_d \rho_0 \left( \frac{m_C p_C}{\rho_C} + \frac{m_N p_N}{\rho_N} \right)$$

(16)

where $m_C$ and $m_N$ are the molar masses of CO$_2$ and N$_2$ respectively, $\rho_1$ are densities of H$_2$O, CO$_2$, and N$_2$ in state 5, and $\Delta U$ is the change in gravitational potential energy of the system from state 4 to state 5.

Eqn (14), (15), and (16) all yield the same value for $W_{bc}$, the minimum possible work. It should be noted that this lower limit depends on the depth of injection, but it is independent of the reservoir geometry and area [see Fig. 2(c)]. That is to say that creating sufficient pore-space for a given volume of CO$_2$ in a reservoir with smaller area requires greater vertical displacement of less pore-water while emplacing the CO$_2$ into a reservoir with larger area requires less vertical displacement of more pore-water. The displaced volume and the required work are the same in both cases.

Friction. The energetic calculations are now extended to include friction associated with transport and storage. As the CO$_2$ is transported from the injection zone to the storage site, viscous drag within the pipeline decreases the fluid pressure. The pressure drop in a pipeline is given by:

$$\Delta P = \rho\beta \left( \frac{LV^2}{2D} \right)$$

(17)

where $\rho$ is the fluid density, $L$ is pipeline length, $V$ is the fluid velocity, $D$ is the pipeline diameter, and $\beta$ is the friction factor, which depend on the Reynolds number of the fluid and inner roughness of the pipeline. For typical values ($i.e.$, $\rho = 900$ kg/m$^3$, $D = 0.5$ m, $V = 2$ m/s) $\beta \approx 10^{-2}$ and $\Delta P \approx 50$ (kg/(m$^2$s$^2$)). Therefore, a typical pressure drop is $\sim 0.5$ bars per km or 25 bars per 50 km. Eqn (14) indicates that—in the thermodynamic limit—the work required to re-compress the CO$_2$ from 90 bars to 115 is $\sim 0.4$ kJ/(mol) per 50 km.

Friction is encountered again when CO$_2$ is injected into the formation and work is required to compress the pore-fluid and expand the pore-space. That work dissipates into friction, which is manifest in elevated injection pressure requiring additional compression work ($W_b$ and $W_c$). The frictional loss is larger for high rates of flow—so the compression work is greater for faster injection. But, storage within the reservoir reduces work by slowing down flow.

Induced fracturing, however, limits reservoir pressures to the formation’s least compressive stress. The fracture threshold is predominately determined by the least compressive stress. If the vertical stress, which is often near lithostatic pressure, is the weakest stress, then horizontal fractures will form and accommodate the injected CO$_2$. It is more likely, however, that one of the horizontal stresses is the weakest indicating that vertical fractures are most likely.

In any case, the pressure and the required work are bounded by pressure-induced fracturing. Prior to having reached fracture...
pressure, however, additional compression work is required to maintain the constant flow of CO₂ into the formation. Over the lifetime of the injection, the reservoir pressures can increase from the hydrostatic pressure of the formation to the fracture pressure, which is ∼1.4–2.0 times the hydrostatic pressure.²⁷

The additional compression work required (i.e., the increase in \( W_b \)) to accommodate the increased reservoir pressure is small compared with the initial compression as the CO₂ pressure–volume curve (Fig. 3) flattens out significantly above 100 bars. On the other hand, the additional work required to emplace the CO₂ beneath the pore-water by pushing the fluid into the reservoir (i.e., the increase in \( W_c \)) is on the same order as the initial work \( W_c \). That is because eqn (11) is linear in pressure, and the fracture pressure is about twice the initial hydrostatic pressure. Specifically, \( W_b \) changes from ∼12 kJ/(mol CO₂) initially to 13 kJ/(mol CO₂) once fracture pressure is reached, while \( W_c \) changes from 1 kJ/mol to ∼2 kJ/mol. Thus, the friction related work increases \( W_{bc} \) by ∼15%.

### Primary energy required and waste heat recovery

The minimum total work required to separate, compress, transport, and store a unit of CO₂ produced from a power plant is:

\[
W_{\text{tot}} = W_a + W_b + W_c
\]

Performing work \( W_a \), \( W_b \), and \( W_c \), requires primary energy \( E_a \), \( E_b \), and \( E_c \). Calculating \( E_S \)—the sum of \( E_a \), \( E_b \), and \( E_c \)—requires determining the theoretical limit for each conversion of primary energy to work. Subsequently, we extend our calculations to consider reasonable upper bounds on efficiencies, and hence reasonable lower bounds for the energy penalty.

\( E_a \)

A well developed technology for post-combustion separation of CO₂ from flue gas is the temperature-swing system with the solvent monothenolamine (MEA).²⁹ \( E_a \) is not set by the enthalpy of desorption because nearly the same quantity of heat is generated during the exothermic absorption as the endothermic desorption. Rather, \( E_a \) is set by the 2nd-law of thermodynamics, which limits the efficiency of any heat-to-work conversion.²⁹ Therefore, \( E_a \) is given by \( W_a \) over the product of the ideal heat-to-work conversion efficiency for separation (\( \eta_{\text{hsad}} \)) and the 2nd-law efficiency of the actual separation process (\( \eta_{\text{hsad}} \)):

\[
E_a = \frac{W_a}{\eta_{\text{hsad}} \eta_{\text{h2ad}}}
\]

\( \eta_{\text{hsad}} \) for a temperature-swing separation—such as monoethanolamine (MEA)—is given by the efficiency of a Carnot engine running between its highest and lowest temperatures. The lowest temperature (\( T_a \sim 310 \) K) of an MEA system is the temperature of the absorber unit that binds lean MEA to CO₂, while the highest temperature (\( T_S \sim 390 \) K) is the temperature of the of the stripper unit that desorbs the rich MEA (Fig. 2):²⁹

\[
\eta_{\text{hsad}} = 1 - \frac{T_S}{T_a}
\]

Temperature-swing separation systems—like MEA—often work at low temperatures.²⁹ For that reason, harnessing low-temperature waste heat for this type of separation is practical, and as such, the additional primary energy required for separation, \( E_{\text{sep}} \), is lower than \( E_a \). If the quantity of waste heat that can—in principle—be used for separation is \( E_w \), then and the primary energy required from additional combustion of fuel is:

\[
E_{\text{sep}} = \frac{W_a}{\eta_{\text{hsad}} \eta_{\text{h2ad}}} - E_w
\]

Or,

\[
E_{\text{sep}} = \frac{W_a}{(1 - \frac{T_S}{T_a}) \eta_{\text{h2ad}}} - E_w
\]

\( E_w \)—the available waste heat—is calculated by considering the Carnot efficiency of the power plant:

\[
\eta_c = 1 - \frac{T_L}{T_H} = 1 - \frac{E_L}{E_H}
\]

where \( T_L \) is the minimum temperature of the working fluid (i.e., the condenser), \( T_H \) is maximum temperature of the working fluid (i.e., modern-ultra-critical steam cycles run at ∼1000 K²), \( E_L \) is the minimum quantity of heat transferred to the environment, and \( E_H \) is the primary energy content of the fuel, which for a high rank coal is ∼400 kJ/(mol CO₂).³⁰

The actual power plant efficiency (\( \eta_{pp} \)) is given by:

\[
\eta_{pp} = 1 - \frac{E'_1}{E_H}
\]

where \( E'_1 \) is the actual quantity of heat dumped to the environment, which is equal to \( E_L \) plus the total waste heat produced by irreversible processes (or inefficiencies) within the plant (\( E'_w \)). Thus,

\[
\eta_{pp} = 1 - \frac{E_L + E'_w}{E_H}
\]

Solving for \( E'_w \) with the above 3 equations yields:

\[
E'_w = E_H(\eta_c - \eta_{pp})
\]

Eqn (26) gives the total waste heat produced by irreversible processes (or inefficiencies) within the plant by a power plant. \( E'_w \), however, is greater than \( E_w \)—the available waste—because temperature-swing separation systems have a minimum temperature at which heat can be used. The stripper accepts heat at or above \( T_S \) while the absorber dumps heat to the environment (or the condenser) at or above \( T_L \) (Fig. 2). As such, waste heat can only be utilized if it is greater than \( T_S \). The fraction of available waste heat (\( \eta_w \)) approaches zero as \( T_S \) approaches \( T_H \). Likewise, \( \eta_w \) approaches 1 as \( T_S \) approaches \( T_L \). There is limited published data on the temperature distribution of power-plant waste heat. For that reason, a linear temperature distribution between \( T_L \) and \( T_H \) was assumed. Such a waste–heat distribution yields:

\[
\eta_w = \frac{T_H - T_S}{T_H - T_L}
\]
Careful research is needed to more accurately access the waste–heat distributions. Such work would be very valuable, but also very difficult as it would require detailed engineering studies of existing power-plants. The goal of this study is to constrain what is physically possible given actual and theoretical power plant efficiencies.

Given our assumed waste–heat distribution, the quantity of available waste heat is:

\[
E_w = E_H (\eta_c - \eta_{pp}) \frac{T_H - T_S}{T_H - T_L} \tag{28}
\]

Or,

\[
E_w = E_H \left(1 - \frac{T_L}{T_H}\right) - \eta_{pp} \frac{T_H - T_S}{T_H - T_L} \tag{29}
\]

Only a fraction of the available waste heat, however, will actually be harnessed. Therefore, we introduce a new variable \(E_{w2nd}\), which is the 2nd-law efficiency (i.e., the fraction of maximum possible) of the waste heat recovery, and therefore, the total quantity of waste heat that is actually recovered for productive use is:

\[
E_{w2nd} = E_H \left(1 - \frac{T_L}{T_H}\right) - \eta_{pp} \frac{T_H - T_S}{T_H - T_L} \eta_{w2nd} \tag{30}
\]

Combining eqn (21) and (30) yields the additional primary energy required to perform \(W_a\) for a temperature-swing separation process:

\[
E_{sep} = E_1 - E_{w2nd} = \frac{W_a}{\left(1 - \frac{T_S}{T_H}\right) \eta_{sep}} - E_H \left(1 - \frac{T_L}{T_H}\right) - \eta_{pp} \frac{T_H - T_S}{T_H - T_L} \eta_{w2nd} \tag{31}
\]

\(E_b\) and \(E_c\)

The primary energy required for compression depends on the efficiency of the turbine to produce shaft work and the 2nd-law efficiency of the compressors themselves. No waste heat is allocated to compression as that would require building a more efficient power plant. Therefore:

\[
E_b + E_c = \frac{W_b + W_c}{\eta_{pp} \eta_{com}} \tag{32}
\]

where \(\eta_{pp}\) is the power plant thermal efficiency and \(\eta_{com}\) is the isothermal compressor efficiency. Available compressors report isothermal efficiencies of between ~57% and 66%.

The re-compression necessary to overcome pipeline and reservoir friction will be powered by either natural-gas from a parallel pipeline or electricity from the grid. Calculating the associated \(CO_2\) emissions requires either the efficiency of natural gas compressors or the collective efficiency of grid power with respect to \(CO_2\) emissions (i.e., all the electric power produced over all the thermal power from \(CO_2\) production). Assuming the \(CO_2\) is not transported more than a few hundred kilometers, the primary energy calculated from such values increases \(E_{bc}\) by less than 10%.

Adding eqn (31) and (32), we arrive at the lower limit of the total primary energy used by a temperature-swing separation system for \(CO_2\) sequestration:

\[
E_S = \frac{W_a}{\left(1 - \frac{T_S}{T_H}\right) \eta_{sep}} - E_H \left(1 - \frac{T_L}{T_H}\right) - \eta_{pp} \frac{T_H - T_S}{T_H - T_L} \eta_{w2nd} + \frac{W_b + W_c}{\eta_{pp} \eta_{com}} \tag{33}
\]

For non-temperature-swing separation systems—such as membranes or pressure-swing absorption systems—waste heat is not useful because the separation work is a parasitic load on the power-plant turbine. As such, \(E_S\) is:

\[
E_S = \frac{W_a}{\eta_{sep} \eta_{pp}} + \frac{W_b + W_c}{\eta_{com} \eta_{pp}} \tag{34}
\]

The energy penalty

The energy penalty, \(f_1\), is the fraction of the fuel that must be dedicated to CCS activities for a given quantity of fuel input \((E_H)\):

\[
f_1 = \frac{E_S}{E_H} \tag{35}
\]

If the quantity of fuel is fixed, then the energy penalty is manifest in the reduction of the plant’s power output. The new power output per unit fuel is:

\[
W' = \eta_{pp} E_H \left(1 - \frac{E_S}{E_H}\right) = \eta_{pp} E_H (1 - f_1) \tag{36}
\]

If, on the other hand, the power output is fixed, then the energy penalty is manifest as the increase in fuel necessary to maintain that constant power output. This additional fuel requirement is expressed as the ratio of the fuel for CCS \((E_S)\) to the fuel that produces power output \((E_H - E_S)\). Thus, the fraction of additional fuel required to maintain the constant power output associated with \(E_H\) is:

\[
f_2 = \frac{E_S}{E_H - E_S} = \frac{E_S}{E_H - E_S - E_S/E_H} = \frac{f_1}{1 - f_1} \tag{37}
\]

This additional fuel requirement can also derived as the geometric sum of the energy penalty. The additional fuel required to sequester the \(CO_2\) produced for a unit of power generation is \(f_1\), but in burning this fuel, more \(CO_2\) is produced that requires \(f_1^2\) additional fuel to sequester, which in turn requires more fuel, ad infinitum, to give the sum of the infinite geometric series:

\[
f_2 = f_1 + f_1^2 + f_1^3 + ... = \frac{f_1}{1 - f_1} \tag{38}
\]

By combining eqn (33) with eqn (35), the energy penalty \((f_1)\) and the additional fuel requirement \((f_2)\) for a temperature-swing separation system can be written in terms of basic system parameters as well as the minimum work required for separation and compression:
For a temperature-swing separation process, eqn (39) provides insight into that spread as well as into the thermodynamic limit for sequestration with a temperature-swing separation system that implies an energy penalty of approximately 45 kJ/(mol CO₂), implying an f₁ of ~11% and an f₂ of ~13%.

Table 1 reveals the end-member cases for post-combustion capture and storage between the thermodynamic lower-bound and upper-bound values being reported for current technology. From these values, it is clear that the energy penalties achieved from temperature-swing separation systems are more uncertain, but also that waste-heat recovery offers a significant opportunity to decrease the energy penalty.

The thermodynamic limit for sequestration with a temperature-swing separation system indicates that capturing and storing all the CO₂ generated from current U.S. PC plants while delivering the same power output would require—at the very least—consuming an additional ~120 million tonnes of coal annually. If, on the other hand, the energy penalty were incurred by decreasing the electrical work output—rather than increasing the fuel consumption—then the electrical work output of the U.S. coal fleet will drop by—at the very least—~37 GW. That means that either an additional 37 GW of base-load CO₂-free power have to be built, or national electricity use would have to be reduced by 37 GW.

The energy penalty for post-combustion capture and storage of PC power-plant CO₂ has been estimated by several different studies. A review of that literature demonstrates a relationship between the energy penalty and the economics of CCS (Fig. 1); it also reveals a significant spread between the various published estimates of the energy penalty. The reviewed studies of PC retrofits include only small amounts of waste-heat recovery, and the associated f₂ values are between 43% and 77%, which indicates that waste-heat recovery offers a significant opportunity to decrease the energy penalty.

Eqn (40) provides insight into that spread as well as into future CCS development. Eqn (40) depends on 17 parameters, yet in practice only two of them—the fraction of available waste heat that is actually recovered (ηₜₚₚ) and the 2nd-law separation efficiency (ηₛₛ)—are poorly constrained (ηₛₛ varies significantly, but it is well constrained). Fig. 4 reveals how f₂ depends on these poorly constrained parameters.

New construction projects have two distinct advantages; first, the power plants themselves—having supercritical or ultracritical steam cycles—are more efficient which results in a lower energy penalty since the primary energy required for compression is function of ηₛₛ; and second, new construction projects can more easily be designed to utilize low-grade waste heat for CO₂ separation. The absolute value of the contour slopes in Fig. 4 are mostly less than 1, which indicates that f₂ is generally more sensitive to changes of ηₛₛ than to changes of ηₜₚₚ. Since current systems already achieve ηₛₛ values in the range of ~40%–60%, ηₜₚₚ represents the most potential for...
improvement as the contour slopes are significantly steeper for $h_{s2nd}$ values above 50%.

The energy penalty—as derived in this paper—can be used to calculate the optimal values for various independent variables. For instance, $W_a$ and $W_{bc}$ depend on the degree of separation in opposite directions. Zero separation minimizes $W_a$, but it maximizes $W_{bc}$. The same is true—naturally—for $E_a$ and $E_{bc}$.

![Fig. 4](image) The additional fuel requirement ($f_2$) for coal-fired power plants employing a MEA separation system. The horizontal axis is the fraction of available available-waste heat employed for separation, and the vertical axis is the 2nd-law efficiency of the separation process ($\eta_{s2nd}$). For these calculations, the stripper temperature ($T_S$) = 390 K, the absorber temperature ($T_A$) = 310 K, the power plant efficiency = 33% ($\eta_{pp}$), the isothermal compressor efficiency = 65% ($\eta_{com}$), the highest turbine temperature ($T_H$) = 1000 K, and the environmental temperature ($T_L$) = 300 K.

Fig. 5 shows the optimal separation as a function of the fraction of flue-gas CO$_2$ that is emitted to the atmosphere.

The stripper temperature is another parameter for optimization as $T_S$ affects $f_1$ in two different directions. The ideal separation efficiency increases with $T_S$ (eqn (20)), but the quantity of available waste heat decreases with $T_S$ (eqn (27)). MEA strippers operate at $\sim$390 K, and research is ongoing to identify new absorption materials—such as ionic liquids$^{32}$—that can operate at higher temperatures with the goal of increasing the heat to work conversion efficiency. Due to the decrease in available waste heat, however, these efforts might be limited in their potential. For typical efficiency and waste–heat recovery values,
increasing the $T_s$ beyond $\sim$500 K might not be helpful because the loss of available waste heat compensates for the increase in separation efficiency [Fig. 6(a)]. Indeed, given the likelihood that the waste–heat temperature distribution is more skewed toward lower temperatures than the linear distribution assumed here, it is probable that the optimal $T_s$ is below 500 K, suggesting that current systems are operating near the optimal $T_s$. That assumes, however, that effective engineering can harness the available waste heat.

Sensitivity analysis on the total energy penalty is performed by varying the power-plant efficiency. The additional fuel requirement ($f_2$) depends on the power-plant efficiency in two ways: $E_{bc}$ decreases as $\eta_{pp}$ increases, but $E_s$ can actually decrease as $\eta_{pp}$ increases because the available waste heat decreases as $\eta_{pp}$ increases. Fig. 6(b) shows $f_2$ as a function of $\eta_{pp}$, and that figure demonstrates that $f_2$ is minimized for particular power-plant efficiencies. If available waste heat recovery rates can exceed 30%, then it may not be beneficial to target inefficient plants for CCS retrofits.

The reviewed studies also indicate significant differences in the energy penalty between new construction projects and retrofits. Those differences are primarily driven by 3 factors that are made clear from our analysis of the energy penalty: (1) the degree of available-waste-heat recovery ($\eta_{w2nd}$), (2) the baseline power plant efficiency ($\eta_{pp}$), and (3) the 2nd-law separation efficiency ($\eta_{s2nd}$). All the studies of new construction projects involve either supercritical or ultra-supercritical cycles whose superior plant efficiencies result in lower energy penalties than subcritical cycles. In addition, waste–heat recovery for separation is easier to implement in new construction projects than in retrofits.

The U.S. installed base of PC plants has a total thermal efficiency ($\eta_{pp}$) of 33%. Fig. 7(a) shows the distribution of thermal efficiencies for the installed base of PC plants.\textsuperscript{13}

In 2007, the most efficient plant recorded a thermal efficiency of 46.4% while the least efficient plant recorded a value 18.7%. The energy penalties ($f_1$) to capture and store the CO\textsubscript{2} from those two plants with a modern temperature-swing separation system are 34% and 52%, respectively. From the distribution of thermal efficiencies, $f_1$ and the additional fuel requirement ($f_2$) associated with converting all or some of the U.S. coal-fleet to CCS can be calculated.

Fig. 7(b) shows the distribution of $f_2$ for the U.S. installed base with 20% available-waste-heat recovery. That distribution yields a spread in $E_{bc}$ between 78 and 96 kJ/mol because less efficient plants have more available waste heat. Fig. 7(b) shows the corresponding $f_2$ distribution, which spreads from 0.57 to 1.01 with a mean value of 0.66 and standard deviation of 0.05. Converting the entire PC installed base to CCS while keeping its electrical work output constant would require an additional $\sim$460 million tonnes of coal annually (assuming an average energy content of 25 GJ/(tonne coal)). Alternatively, the energy penalty could be manifest in a decreased plant output. In that case, the power output of the U.S. coal fleet would drop by $\sim$78 GW.

Under the 20% available-waste-heat recovery assumption, the difference between retrofitting the most efficient plants for CCS and retrofitting the least efficient plants is significant [Fig. 7(b)]. If the 10 most efficient plants were retrofitted to capture and store 80% of their CO\textsubscript{2}, then an additional 6.5 million tonnes of coal would be required and 27 million tonnes of CO\textsubscript{2} emissions would be eliminated annually. Thus, the CO\textsubscript{2} abatement effectiveness (i.e., the mass ratio of CO\textsubscript{2} eliminated to additional coal required) for the top 10 plants is 4.1. On the other hand, retrofitting the 10 least efficient plants would require 2.4 million tonnes of additional coal and would only eliminate 6.8 million tonnes of CO\textsubscript{2} annually yielding a CO\textsubscript{2} abatement effectiveness of 2.8. Thus, retrofitting the 10 most efficient PC plants for CCS would eliminate 46% more CO\textsubscript{2} emissions per unit of additional coal than retrofitting the 10 least efficient plants.

These calculations were repeated for the most and least efficient 10\% and 25\% of current PC plants (Table 2). The columns in Table 2 reveal how the CO\textsubscript{2} abatement effectiveness depends on the fraction of available-waste heat that is recovered from retrofitting a particular ensemble of the most efficient plants versus the equivalent ensemble of least efficient plants.

---

**Fig. 7** (A) The thermal efficiency distribution of an ensemble of 420 large U.S. coal-fired power plants. These plants produced the equivalent of 218 GW of constant electric power in 2007 constituting 96% of all U.S. coal-fired power output. (B) The distribution of additional fuel requirements ($f_2$) is calculated from the power-plant efficiency distribution. From this distribution, the total additional fuel required is calculated to be 530 million tonnes of coal. These calculations assume, $\eta_{w2nd} = 40\%$, $\eta_{wout} = 20\%$, $\eta_{com} = 65\%$, and the national average coal heat content (25 GJ/(tonne)).
The CO₂ abatement effectiveness (i.e., the mass ratio of CO₂ eliminated to additional coal required) for 6 ensembles of U.S. coal plants: The ensembles are organized by each plant’s reported thermal efficiency. The first row labeled ‘Top 2%’ is for the most efficient 2% of U.S. coal plants. Each column assumes a different value for \( \eta_{w2nd} \), which is the fraction of available-waste heat that is harnessed for separation. In the \( \eta_{w2nd} = 0\% \) case, retrofitting the most efficient 10% of plants will eliminate nearly 30% more CO₂ per unit of additional coal than retrofitting the least 10% of plants. As the \( \eta_{w2nd} \) increases, then the gap in CO₂ abatement effectiveness decreases because less efficient plants have a greater amount of available waste heat. Table 2 reveals a narrowing of the CO₂ abatement effectiveness with increasing available-waste-heat recovery, but even in the high \( \eta_{w} \) scenario, retrofitting the most efficient PC plants is nevertheless measurably more effective than retrofitting the least efficient plants.

The financial costs of CCS are tightly related to the energy penalty (Fig. 1). Indeed, it is worth noting in Fig. 1 that while new PC construction appear superior to PC retrofits when measured by the common metric of dollars per tonne of CO₂ avoided (Fig. 1(a)), retrofits and new construction are about equal when measured by the more relevant metric of cost of electricity from a CCS power plant (Fig. 1(b)). That is partially the result of lower fixed costs associated with plants that have been fully or partially amortized.

The correlation between the costs of CCS and the energy penalty coupled with the variance in expected energy penalties (Fig. 7(b)) suggests an optimal CCS deployment strategy. The cheapest path to drastically reduce CO₂ emissions from electricity production will combine the selective retrofitting of the most efficient PC plants with the closing of the least efficient plants. Overall, our analysis strongly suggests that the supply curve for retrofitting PC plants for CCS is a function of the power-plant’s baseline thermal efficiency.

It should be noted, however, that the relationship between base-line efficiency and the energy penalty assumes that the power plant itself is providing the compression work. Other configurations are possible. For example, a dedicated natural-gas-fired compressor or even a wind turbine could provide the necessary compression work. In those scenarios, \( f_1 \) would manifest as the reduced power output from either natural gas or wind; \( f_2 \), however, would be lower because the CO₂ intensity of both gas and wind are lower than that of coal.

To demonstrate the potential value of available-waste-heat recovery, we calculate \( f_2 \) for the entire U.S. coal fleet with and without 1/3 available-waste-heat recovery. Retrofitting the entire U.S. coal fleet with zero available-waste-heat recovery would require additional ~600 million tonnes of coal annually. If, on the other hand, 1/3 of the available waste heat were productively used for separation, then the additional fuel requirement would drop from ~600 million to ~390 million tonnes of coal annually. Alternatively, if the energy penalty were manifest in a reduced power output, then with zero available-waste-heat recovery an additional ~92 GW of CO₂-free base-load power would be required to make up for the decrease in power output. With 1/3 available-waste-heat recovery, however, the additional power requirement would drop from ~91 GW to ~69 GW.

Improving end-use electrical efficiency is an additional path through which the CCS energy penalty could be offset. This path is intriguing because the total U.S. smoothed power output was ~472 GW in 2007,\(^4\) indicating that increasing end-use electrical efficiency of between 15% and 20% would be sufficient to make up for the decrease in power output after retrofitting the installed PC base for CCS. That would yield a ~65% reduction in CO₂ emissions from the power sector while not requiring any additional power-generation capacity to be built or any additional coal to be burned. The remaining 35% would come primarily from natural gas as well as a little from coal as we assumed 80% CO₂ capture. This approach may be feasible as California has been able to keep its per capita electricity use constant for the past 30 years, while average per capita electricity use in U.S. grew by nearly 50%.\(^8\)

**Conclusion**

Achieving substantial reductions in CO₂ emissions requires either shutting down a large fraction of the current installed base of coal-fired power plants or retrofitting those plants for CCS. Previous studies have estimated that the additional fuel required (\( f_2 \)) to maintain constant work output for a PC retrofit is between ~50% and 80%. An analysis of the thermodynamic limit indicates those values might be improved by harnessing more of the available waste heat and by improving the 2nd-law efficiency of temperature-swing separation systems. It appears difficult, however, to improve \( f_2 \) for post-combustion capture to below ~25% in practice. Our most likely efficiency scenario indicates that offsetting the energy penalty incurred from capturing and storing 80% of the U.S. coal fleet’s CO₂ emissions will require either an additional ~390–600 million tonnes of coal, an additional ~69–92 gigawatts of CO₂-free-base load power, or a 15%–20% reduction in overall electricity use.
Work:

\begin{align*}
W_a & \quad \text{The work required to separate the CO}_2 \text{ from the flue gas [thermodynamic limit} \sim 9 \text{ kJ/mol]} \\
W_b & \quad \text{The work required to compress the concentrated CO}_2 \text{ from atmospheric to reservoir pressure [thermodynamic limit} \sim 13 \text{ kJ/mol]} \\
W_{c1} & \quad \text{The work required to vertically displace groundwater [thermodynamic limit} \sim 1–2 \text{ kJ/mol]} \\
W_{c2} & \quad \text{The work required to generate a interface between CO}_2 \text{ and the pore-water [thermodynamic limit} <1 \text{ kJ/mol]} \\
W_c & \quad W_{c1} + W_{c2} [\text{thermodynamic limit} \sim 2 \text{ kJ/mol}] \\
W & \quad \text{Power plant work output after the addition of CCS} \\
W_{\text{tot}} & \quad W_a + W_b + W_c [\text{kJ/mol}] \\
W_{ab} & \quad W_a + W_b [\text{kJ/mol}] \\
W_{bc} & \quad W_b + W_c [\text{kJ/mol}] \\
\end{align*}

\begin{align*}
\text{Primary Energy:} & \\
E_a & \quad \text{The primary energy required to separate the CO}_2 \text{ from the flue gas [kJ]} \\
E_b & \quad \text{The primary energy required to compress the concentrated CO}_2 \text{ to reservoir pressure [kJ]} \\
E_c & \quad \text{The primary energy required to emplace compressed CO}_2 \text{ into the geologic formation [kJ]} \\
E_{\text{sep}} & \quad \text{The incremental primary energy required to separate CO}_2 \text{ from the flue gas [kJ]} \\
E_S & \quad \text{The total primary energy required for sequestration [kJ]} \\
E_w & \quad \text{The quantity of waste heat that can—in principle—be used in separation [kJ]} \\
E'_w & \quad \text{The total waste heat produced [kJ]} \\
E_L & \quad \text{The minimum quantity of heat transferred to the environment [kJ]} \\
E'_L & \quad \text{The actual quantity of heat transferred to the environment [kJ]} \\
E_H & \quad \text{The primary energy content of the fuel [kJ]} \\
\end{align*}

\begin{align*}
\text{Efficiencies:} & \\
\eta_{\text{sep}} & \quad \text{Ideal separation efficiency} (\eta_{\text{sep}} \ast \eta_{\text{com}} \text{ for pressure swing, } \sim 30\% \text{ for temperature swing)} \\
\eta_{\text{sep}} & \quad \text{2nd-law separation efficiency} (\sim 50\%) \\
\eta_c & \quad \text{The power-plant Carnot efficiency} (\sim 70\%) \\
\eta_{\text{pp}} & \quad \text{The power plant efficiency} (25\%–45\%) \\
\eta_{\text{com}} & \quad \text{Isothermal compression efficiency} (65\%) \\
\end{align*}

\begin{align*}
\text{Other Parameters:} & \\
X_{\text{N}_2} & \quad \text{The mole fraction of N}_2 \text{ in state } i \text{ (state 1: } \sim 80\%) \\
X_{\text{CO}_2} & \quad \text{The mole fraction of CO}_2 \text{ in state } i \text{ (state 1: } \sim 12\%) \\
\end{align*}

\begin{align*}
\eta_{\text{N}_2} & \quad \text{The number of moles of N}_2 \text{ in state } i \text{ (state 1: } \sim 0.80 \text{ moles N}_2 \text{ per mole of flue gas)} \\
\eta_{\text{CO}_2} & \quad \text{The number of moles of CO}_2 \text{ in state } i \text{ (state 1: } \sim 0.12 \text{ moles CO}_2 \text{ per mole of flue gas)} \\
P_f & \quad \text{The pressure of state } f [\text{Pa}] (\sim 10^6 \text{ Pa at the surface, } \sim 10^3 \text{ Pa in the reservoir}) \\
g & \quad \text{The gravitational acceleration [m/s}^2\text{]} \\
L_d & \quad \text{The depth of CO}_2 \text{ injection [} \sim 1000 \text{ m]} \\
L & \quad \text{Length of pipeline} \\
\rho_w & \quad \text{The density of H}_2\text{O} [\sim \text{kg/m}^3] (\sim 1000 \text{ kg/m}^3) \\
\rho_{\text{CO}_2} & \quad \text{The density of CO}_2 \text{ in state } i [\text{kg/m}^3] (\sim 2 \text{ kg/m}^3 \text{ at the surface, } \sim 400–600 \text{ kg/(m}^3\text{) in the reservoir}) \\
\rho_{\text{N}_2} & \quad \text{The density of N}_2 \text{ in state } i [\text{kg/m}^3] (\sim 1.2 \text{ kg/m}^3) \\
h_{\text{CO}_2} & \quad \text{The molar enthalpy of CO}_2 \text{ in state } i [\text{kJ/mol}] \\
h_{\text{N}_2} & \quad \text{The molar enthalpy of N}_2 \text{ in state } i [\text{kJ/mol}] \\
s_{\text{CO}_2} & \quad \text{The molar entropy of CO}_2 \text{ in state } i [\text{kJ/(K mol)}] \\
s_{\text{N}_2} & \quad \text{The molar entropy of N}_2 \text{ in state } i [\text{kJ/(K mol)}] \\
m_{\text{CO}_2} & \quad \text{The molar mass of CO}_2 [\text{kg/mol}] \\
m_{\text{N}_2} & \quad \text{The molar mass of N}_2 [\text{kg/mol}] \\
T_a & \quad \text{Temperature of the MEA absorber unit [} \sim 320–350 \text{ K]} \\
T_S & \quad \text{Temperature of the MEA stripper unit [} \sim 400 \text{ K]} \\
T_L & \quad \text{Temperature of the environment [} \sim 293 \text{ K]} \\
T_H & \quad \text{Temperature of the steam working fluid [} \sim 1000 \text{ K]} \\
f_1 & \quad \text{The energy penalty} \\
f_2 & \quad \text{The fraction of additional fuel required to maintain the constant power output} \\
V & \quad \text{Total swept out pore volume [m}^3\text{]} \\
v_i & \quad \text{Molar volume of state } i [\text{m}^3/\text{mol}] \\
G & \quad \text{The Gibbs free energy [kJ]} \\
\end{align*}

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