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# Effects of Impurities on CO<sub>2</sub> Transport, Injection and Storage

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#### Abstract

Effects of impurities in  $CO_2$  streams on geological storage of  $CO_2$  have been investigated. A number of key issues have been addressed, and several significant findings have been made. Highlighted among the findings are: (1) non-condensable impurities such as  $N_2$ ,  $O_2$  and Ar greatly reduce  $CO_2$  storage capacity of geological formations, and there is a maximum reduction of the storage capacity at a certain pressure under a given temperature. By contrast, impurities which are more condensable than  $CO_2$ , such as  $SO_2$ , can increase the storage capacity, and there is a maximum increase at a certain pressure under a given temperature; (2) change of density caused by non-condensable gas impurities results in lower injectivity of impure  $CO_2$  into geological formations. However, above a threshold pressure range the injectivity could reach the level of pure  $CO_2$  due to lowered viscosity; (3) non-condensable impurities increase the buoyancy of the  $CO_2$  plume. This would negatively affect the efficiency of solubility trapping and residual trapping of  $CO_2$ . (4) the effect of  $SO_2$  on reduction of rock porosity and hence  $CO_2$  injectivity would be much smaller than previously thought. A simple formula has been developed to enable quick determination of the effect.

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Keywords: CO<sub>2</sub>, impurities, storage capacity, injectivity

#### Effects of Impurities on Phase Behaviour of CO2

 $CO_2$  capture and storage (CCS) has been identified as a key potential technology in mitigating  $CO_2$  emissions. The capture process represents the major cost component of CCS.  $CO_2$  may be captured by different technologies, and various gaseous impurities exist in the  $CO_2$  stream, such as  $N_2$ ,  $O_2$ ,  $SO_2$  and  $H_2S$ . Separation of the impurities would drastically increase the cost of capture and it would be more cost effective to co-inject the impurities with  $CO_2$  for underground storage. Moreover, some technologies aim at co-capture and co-storage of multiple air pollutants together with  $CO_2$ .

Unfortunately, impurities in the  $CO_2$  stream can have negative effects on transport, injection and storage of  $CO_2$ . For instance, non-condensable impurities such as  $N_2$ ,  $O_2$  and Ar would increase the saturation pressure of liquid  $CO_2$  and decrease the critical temperature. As a result, lower temperature and additional overpressure is required to avoid two-phase flow in  $CO_2$  pipeline transport. Non-condensable impurities would also increase the pressure for injection

and reduce the capacity of the storage sites by decreasing the density of the  $CO_2$  stream. Moreover, acid impurities like  $SO_x$  and  $NO_x$  can react with formation and cap rocks and affect the injectivity and storage integrity. In addition, the fate of hazardous impurities is of special concern in the event of  $CO_2$  leakage, as they may be released to the environment.

Natural Resources Canada (NRCan) has been leading Canada's federal CCS programs. Along with a range of R&D projects for capturing CO<sub>2</sub> from coal-fired power plants and other sources, NRCan is also involved in and collaborating in research endeavours for CO<sub>2</sub> storage, including CO<sub>2</sub> injection, monitoring, measurement and verification, storage integrity assessment, and capacity estimation. The effects of impurities on transport, injection and storage of captured CO<sub>2</sub> are of vital importance to successful implementation of the CCS research strategies. In this context NRCan has undertaken a study supported by IEA GHG on the effects of the impurities, focusing on storage of CO<sub>2</sub> in saline formations – a promising target for underground CO<sub>2</sub> storage. As the literature on a number of critical impurity effects is scarce, such as the effects on storage capacity, storage integrity and injectivity, we have carried out extensive evaluations of our own. Part of the findings are presented here.

# Effects of Impurities on Phase Behaviour of CO2

Figure 1 shows phase envelopes calculated using the Peng-Robinson equation of state for the following cases:

- 1) CO<sub>2</sub> stream from oxyfuel combustion in a fluidized bed pilot plant combustor in CanmetENERGY, containing 5.2 vol % O<sub>2</sub>, 221 ppm CO, 1431 ppm SO<sub>2</sub> and 243 ppm NO [1];
- 2)  $CO_2$  stream from a zero-emissions process proposed by CanmetENERGY, containing 1.05% CO, 1.7%  $SO_2$ , 0.32%  $H_2$  and 690 ppm  $H_2S$  [2];
- 3) CO<sub>2</sub> stream from Cansolv® absorption system containing 2.9% SO<sub>2</sub>, studied in a previous IEA GHG report [3];
- 4) Expected  $CO_2$  stream from a pre-combustion removal plant, whose composition information is provided by IEA GHG, including 1 vol %  $H_2$ , 0.9 vol %  $N_2$ , 300 ppm Ar, 100 ppm  $H_2S + COS$  and other impurities in ppm level;
- 5) Expected  $CO_2$  stream from an oxyfuel combustion plant, whose composition information is also provided by IEA GHG, including 5.8 vol %  $N_2$ , 4.7 vol %  $O_2$ , 4.47 vol % Ar, 100 ppm  $NO_x$ , 50 ppm  $SO_2$ , 20 ppm  $SO_2$  and 50 ppm CO.

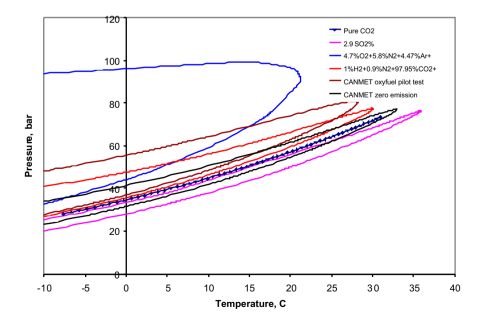


Figure 1 Calculated phase evelopess for CO<sub>2</sub> and CO<sub>2</sub> mixtures.

From the results it can be seen that the critical temperature and pressure of the mixtures are quite different from that of pure  $CO_2$ .  $N_2$ ,  $O_2$ , Ar and  $H_2$  show the greatest effect of increasing the saturation pressure of the liquid and decreasing the critical temperature. One extreme case is the mixture from oxyfuel combustion containing 5.8%  $N_2$ , 4.7%  $O_2$ , 4.47% Ar and other impurities at ppm level. The critical temperature decreases by about 10°C in comparison with that of pure  $CO_2$ , and the liquefaction pressure increases by over 50 bar. On the other hand,  $SO_2$  results in a decrease in the saturation pressure and an increase of the critical temperature, as is expected from the high critical temperature of pure  $SO_2$  (157.6°C). It can also be seen that low-concentration impurities, such as CO and  $NO_x$  would not significantly affect the phase behaviour of  $CO_2$ .

### **Effects of Impurities on Storage Capacity**

Impurities will affect the volumetric properties of injected CO<sub>2</sub>. Figure 2 shows calculated density of supercritical CO<sub>2</sub> with non-condensable impurities for a number of cases at 330 K, which is in the typical temperature range for CO<sub>2</sub> storage evaluations in western Canada. It can be seen that non-condensable impurities such as O<sub>2</sub>, Ar, N<sub>2</sub> and H<sub>2</sub> significantly reduce the density of the supercritical CO<sub>2</sub> stream. The reduced density is largely related to increased volume, except for H<sub>2</sub>, where the effect of smaller molecular weight is also significant. All these impurity components in CO<sub>2</sub> would cause a volume increase greater than their molar or volume fractions at standard temperature and pressure (STP) For example, 5% vol (STP) N<sub>2</sub> will result in a volume increase greater than 5 percent in the temperature range not high above the critical temperature of CO<sub>2</sub>. This can be understood from the fact that non-condensable impurities are less dense than CO<sub>2</sub> and hence take greater volumes. If they have the same molar volume as CO<sub>2</sub> there would be no volume increase, provided the interactions between unlike molecules are negligible.

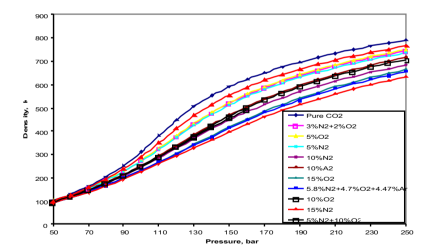


Figure 2 Calculated density for CO<sub>2</sub> and CO<sub>2</sub> mixtures as a function of pressure at 330 K.

The decrease of  $CO_2$  storage capacity due to the contained impurities is thus not only caused by the lower volume fraction of  $CO_2$ , but also by the additional volume of less dense impurities. To produce a simple relation regarding the effect of the impurities on the storage capacity for  $CO_2$  for given storage volume, we propose the following expression for the storage capacity (expressed in mass) as a function of density of the  $CO_2$  stream:

$$\frac{M}{M_0} = \frac{\overline{\rho}}{\rho_0 (1 + \sum_{i} m_i / m_{CO2})}$$
 (1)

where M and  $M_0$  denote the mass of  $CO_2$  in the mixture and in the pure stream, respectively, which occupy the same volume.  $\overline{\rho}$  and  $\rho_0$  are the density of the mixture and the pure stream, and  $m_i/m_{CO2}$  is the ratio of the mass of impurity i to the mass of  $CO_2$  in the mixture. The ratio  $M/M_0$  on the left hand side of this relation, which can also be viewed as the ratio of the mass of  $CO_2$  per unit volume in the mixture to that in the pure state, represents a normalized storage capacity for  $CO_2$  in its supercritical phase, i.e., the capacity for structural trapping of  $CO_2$ . In the case of pure  $CO_2$  (zero impurity effect) the ratio equals unity. The right hand side is a function of temperature, pressure and mixture composition and can be calculated from equations of state. Accordingly, the normalized storage capacity can be determined for given temperature and pressure conditions. Calculated results for a number of mixtures are shown in Figure 3.

The existence of the minimum has been verified through both theoretical analysis and experimental data (details will be shown in a new IEA GHG report).

For impurities which have higher critical temperature than that of  $CO_2$ , no storage capacity minimum would occur in the corresponding  $CO_2$  mixture. Rather, a maximum can appear based on an analysis similar to that for the non-condensable impurities. This has been verified with calculated results for the  $CO_2$  mixture with 2.9 vol %  $SO_2$  (the  $Cansolv^{\oplus}$  absorption system case presented in the previous IEA GHG report [3]). The result shows a maximum at about 110 bar, where the storage capacity is increased by over 5% (figure not shown). It is interesting to see that in the maximum region,  $SO_2$  can create space for  $CO_2$ . This can be rationalized from the consideration that  $SO_2$  decreases average distance between the molecules of the mixture – an opposite effect of that of the non-condensable gases.

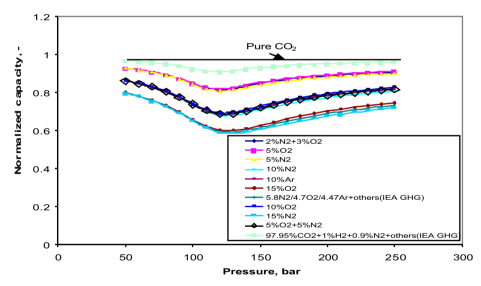


Figure 3 Normalized CO<sub>2</sub> storage capacity at 330 K in terms of Equation 1.

# Impact of the Minimum CO<sub>2</sub> Storage Capacity

The minimum caused by non-condensable impurities is of particular concern, because these impurities are typical for CO<sub>2</sub> streams from oxyfuel combustion. Besides, the pressure for this minimum falls in the pressure range for CO<sub>2</sub> storage. Calculated pressure dependence of the normalized CO<sub>2</sub> storage capacity for several temperatures show that with temperature increasing, the minimum shifts to higher pressure and the magnitude of the minimum decreases (Figure 4). In subsurface formations, the pressure and temperature are interrelated as both of them increase with the depth from the surface. At certain depths, the minimums which correspond to the largest decreases of storage capacities may occur due to pressure and temperature conditions. By assuming a hydrostatic pressure gradient of 10 MPa/km, and geothermal gradients from 20 to 33 °C/km for the subsurface and a ground temperature of 15 °C, the impact of the minimum on the CO<sub>2</sub> storage capacity for the high-impurity stream are illustrated in Table 1. As can be understood, the storage capacity can decrease to as low as 54.3% at 320 K (47°C) and 56.4% at

330 K (57°C) in the case of 33°C/km geothermal gradient. At higher temperature, 350 K (77°C), the decrease in the storage capacity is smaller and at still higher temperatures the decrease would be smaller still.

As structural trapping is deemed the most important trapping mechanism for  $CO_2$  storage at least in the injection phase of a CCS operation, the pressure effect should be taken into account in estimation of capacity for underground formations. Moreover, in storage operations, the minimum would cause reduced storage efficiency. At a given depth, increasing the storage pressure well beyond the level for the minimum would increase storage efficiency. This may be attained for closed formations within the allowable overpressure range. For open formations where overpressure may not be attainable, one possibility to increase the storage efficiency is to increase the depth of injection and storage. With increasing depth the temperature and pressure will increase so that the capacity decrease can be alleviated.

The above discussion should also be applicable to residual trapping of  $CO_2$ , where  $CO_2$  is trapped in rock pores as an immobile phase. Here the changes of interfacial tensions of the  $CO_2$ /water/pore wall system due to the impurities are neglected. The decrease of the trapping capacity due to the impurities can be greater than their molar fractions or STP volume fractions. Moreover, at lower temperatures a minimum may occur at a certain pressure, causing a greater drop in the storage capacity. Furthermore, injection at deeper levels can increase the storage efficiency of impure  $CO_2$  streams.

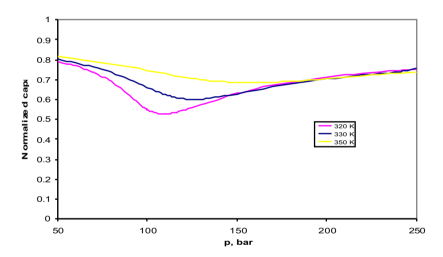


Figure 4 Normalized  $CO_2$  storage capacity for the high-impurity oxyfuel combustion stream (5.8 vol %  $N_2$ , 4.7 vol %  $O_2$ , 4.47 vol % Ar and other impurities at ppm level) at different temperatures.

Table 1 Normalized CO<sub>2</sub> storage capacity for the high impurity CO<sub>2</sub> stream at several subsurface temperatures

T gradient	20°C/km		25°C/km		33°C/km	
T (K)	p (bar)	Capacity (-)	p (bar)	Capacity (-)	p (bar)	Capacity (-)
320	160	0.653	130	0.574	100	0.543
330	210	0.723	170	0.672	130	0.564
350	310	0.755	250	0.734	190	0.698

#### **Permeation Flux**

The presence of impurities would also affect CO<sub>2</sub> injection. Here we discuss their physical effects (no chemical reactions involved) first. For the single phase flow of injected CO<sub>2</sub>, the effects may be analyzed in terms of an expression based on Darcy's Law, for the permeation flux of injected CO<sub>2</sub>:

$$\dot{M} = \frac{-\rho k}{\mu} \nabla p \tag{2}$$

where M is the mass flow per unit area,  $\rho$  is the density of injected stream, k is rock permeability,  $\nabla$  is the gradient operator, p is the pressure, and  $\mu$  is the viscosity of the fluid. As the impurities lower the density of the  $CO_2$  stream, the mass flux will decrease for the same pressure drop. However, the impurities also affect the viscosity of the injected fluid. When the viscosity of the impure  $CO_2$  stream is lower than the viscosity of pure  $CO_2$ , the  $CO_2$  flux increases, hence the decrease in density may be compensated by a corresponding decrease in viscosity. The density and viscosity are functions of temperature and pressure. The permeability and pressure gradient vary case by case. However, for an estimation of the impurity effects under the same permeability and pressure drop conditions, one may use the following relation which is a consequence of Equation 2.15

$$\frac{\dot{M}_{CO2}}{\dot{M}_0} = \frac{\rho(\mu_0/\mu)}{\rho_0(1 + \sum_i m_i/m_0)}$$
(3)

This expression represents a normalized permeation flux, and should be able to provide a measure of the relative injectivity of the impure  $CO_2$  stream. As non-condensable impurities reduce the density of the  $CO_2$  stream, and thus reduce the permeation flux. However, the impurities would also reduce the viscosity of the stream, and thus compensate the effect of reduced density. Calculated result for the high impurity  $CO_2$  stream (with about 15%  $N_2/O_2/Ar$  and other minor-quantity impurities) shows that the permeation flux is lower than that of pure  $CO_2$  by more than 15 percent at lower pressures, but reaches the same level as pure  $CO_2$  above a transition pressure (Figure not shown).

# **Buoyancy and Rising Velocity**

The light-impurity species will result in greater buoyancy for the CO<sub>2</sub> plume, due to the lower density. When a mass of the plume in a unit volume is in contact with the formation water, the buoyant force can be expressed as

$$F = (\rho_{H2O} - \rho_m)g \tag{4}$$

where  $\rho_{H2O}$  and  $\rho_m$  are the density of the water and the plume, respectively. The effect of impurities on this force with reference to pure  $CO_2$  may be given as

$$\frac{F}{F_0} = \frac{\rho_{H2O} - \rho_m}{\rho_{H2O} - \rho_{CO2}} \tag{5}$$

where F and  $F_0$  are the buoyancy force for the  $CO_2$  mixture and pure  $CO_2$ , respectively. The greater the difference between the densities of the mixture and  $CO_2$ , the greater the change in the buoyancy would be. Calculated result shows that with about 15 vol% non-condensable impurities (the high-impurity stream from oxyfuel combustion) the buoyancy of the  $CO_2$  plume can increase by over 50%. This would significantly increase the rising velocity of the plume. The velocity of the impure  $CO_2$  stream ( $\nu$ ) can be related to that of the pure  $CO_2$  stream ( $\nu$ ) by

$$\frac{v}{v_0} = \frac{F/(\rho\mu)}{F_0/(\rho_0\mu_0)} \tag{6}$$

According to calculations, the velocity of the high-impurity plume can reach nearly three times that of the pure  $CO_2$  plume. The high rising velocity would reduce the time for  $CO_2$  contact with water and formation of dissolved species. It would also reduce the formation of a residual phase, because it reduces the lateral spreading of the plume, leaving a smaller area for trapping  $CO_2$  in the pores of rocks. When the plume reaches the caprock it will spread laterally, but the contact with water is only at the interface. Besides, near the caprock the pressure is lower than in the deeper region, and hence the efficiency of  $CO_2$  dissolution and trapping in the pores as a residual phase would be lower. As a result, the potential for leakage increases if pathways are available.

# Impact of Reactive Impurities on Injectivity of CO<sub>2</sub>

The most common reactive impurities are  $H_2S$ ,  $SO_2$ ,  $NO_x$  and CO. CO is highly reductive and will be oxidized to  $CO_2$  by oxygen or mineral oxides. There have been several studies on co-injection of  $H_2S$  and/or  $SO_2$  with  $CO_2$  [4-8]. The general view is that  $H_2S$  is not an issue, and it has been co-injected with  $CO_2$  for 20 years in Canada in acid gas disposal. However,  $SO_2$  is believed to alter the geochemistry, causing increased dissolution and precipitation.  $SO_2$  can greatly lower the pH of the formation water and hence enhance the dissolution of rock minerals. When the species become oversaturated precipitation of sulphates would occur. Reduction of pore volume and hence the injectivity of  $CO_2$  is a concern. The effect of the impurities is difficult to quantify. Numerical simulation studies performed so far have either assumed that  $SO_2$  is injected as an aqueous solution or the injection zone is never dry. This situation would be different from the injection operations where  $SO_2$  migrates with  $CO_2$  in an immiscible plume which results in a desiccation or dry-out zone [9-11]. The important point is that  $SO_2$  is far less reactive when it is dry. Further, continuous dissolution of minerals as in the "wet cases" would not occur in the dry-out zone. The contact of  $SO_2$  with water only occurs on the front of the plume and the two-phase flow zone, and the time would be limited. Therefore, build-up and precipitation of minerals in the downstream which were dissolved in the upstream and carried by water flow would be much less, compared with the "wet cases". In the following we give an analysis for the impact of  $SO_2$  on the injectivity of  $CO_2$ .

 $SO_2$  will be highly reactive if it is dissolved and oxidized to form sulphuric acid. In  $CO_2$  streams from oxyfuel combustion and post-combustion capture plants  $O_2$  and  $NO_x$  will also be present.  $NO_x$  is known to catalyze  $SO_2$  oxidation and hence formation of sulphuric acid according to the lead chamber process. The rate of the oxidation depends on various factors such as concentrations of reactants and products, pressure and temperature. However, the maximum effect of  $NO_x$  is to oxidize all  $SO_2$ . Therefore, we assume that all  $SO_2$  can be converted to sulphuric acid and the concentration of  $SO_4^{\ 2^-}$  ions is equal to that of dissolved  $SO_2$ , when  $NO_x$  and  $O_2$  are present in sufficient amounts

Clearly, the rate of sulphate precipitation will not exceed the rate of SO<sub>2</sub> supply. As a limiting case, we consider that all SO<sub>2</sub> passing the reaction zone, *i.e.*, from the front of the two-phase zone to the front of the dry-out zone, is converted to sulphate precipitate. Taking the dissolution of calcite and precipitation of anhydrite as an example, 2.9 vol % SO<sub>2</sub> (the stream considered in the previous IEA GHG report PH4/32 [3]) at 330 K under 200 bar would result in a pore volume decrease 0.44%. This is much less significant compared with the "wet cases".

#### Other Effects of Impurities

Other effects of impurities, such as the effects on cap rock integrity, corrosion of injection well materials, and cost of  $CO_2$  storage, have also been evaluated. The fate of hazardous impurities, such as  $SO_x$ ,  $NO_x$ , CO and Hg, has also been assessed. The results will be included in the new IEA GHG report.

#### **Conclusions**

Impurities have important impacts on  $CO_2$  storage capacity, storage integrity, injectivity, etc. However, previously existing knowledge could not provide solutions to the problems that will be inevitably encountered. The results obtained from the present work are expected to have significant applications or implications to all  $CO_2$  storage operations.

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