Environmental Performances of Various CCU Options in the Framework of an Integrated Chemical Plant

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Abstract: Several carbon capture processes are investigated to separate a part of the CO\textsubscript{2} contained in the flue gas of a coal-fired power plant located in a chemical integrated plant, with the objective of using it as a raw material in a production process. The expected results are to reduce the impact on global warming potential (GWP) and to increase the productivity of the plant. The study is based on the modelling of the combination of systems in the plant using a process simulation software and using life cycle assessment to evaluate both technical feasibility and environmental aspects. Models for the power plant, the production processes, amine chemical absorption, membrane separation and adsorption on activated coal are developed and validated against industrial and literature data. The life cycle inventory is obtained from the mass and energy balances given by the systems model. A first set of calculations is launched with a high purity requirement for the CO\textsubscript{2} stream (95%) recycled into the process. Those calculations show a 12% increase in productivity for the chemical process considered, but result in no significant gain in terms of GWP. Conversely, scenarios with a lower CO\textsubscript{2} purity (40%) show a drop around 9% of the impacts on GWP using membrane separation and activated coal adsorption, while keeping the other impacts at about the same level.

Keywords: LCA; process modelling; carbon capture and utilization (CCU); adsorption; absorption; membranes

1. Introduction

Among the numerous environmental issues the chemical industry sector has to face, global warming will be one of the most challenging in coming next decades. Keeping productivity, product quality and production costs stable while reducing greenhouse gases (GHG) emissions is a major topic to tackle the global warming issue and anticipate the evolution of the carbon market in the next decades.

Many carbon capture (CC) technologies are now entering into their mature phase, offering new opportunities for cleaner production at the industrial scale. Nonetheless capturing CO\textsubscript{2} raises the question of the fate of the captured CO\textsubscript{2}. Storage, for example geological storage (CCS), is of course a possibility, but it is often associated with animated debates in the scientific community and in the civil society [1]. Moreover, when stored, CO\textsubscript{2} becomes another burden to be looked at. A more satisfying option, when it is possible, is to consider the captured CO\textsubscript{2} as a new raw material and to find ways to use it (Carbon Capture and Utilization—CCU) [2].

In the framework of the present study, we consider the particular case of an integrated chemical plant, which is responsible for large CO\textsubscript{2} emissions, mainly because of the presence of a coal-fired power plant onsite which paradoxically requires CO\textsubscript{2} as raw material in the chemical process itself. With such a configuration, CCU appears to be particularly well adapted to both reduce CO\textsubscript{2} emissions and valorize the captured CO\textsubscript{2} onsite. As we worked on an existing plant, post-combustion options, which are easier to implement in brownfield projects, were considered. Chemical absorption with solvents is a mature technology [3], which was first implemented at industrial scale in Denmark.
on a power station with monoethanolamine (MEA) [4]. Other options using solutions of
demixing amines were studied in [5–8] and showed a better energetic efficiency of the
process. Other solvent types were also investigated [9–11]. Membrane processes were
deeply investigated in [12–14] and a recent paper [15] showed that they could be more
advantageous than absorption processes in terms of environmental impacts. Adsorption
on activated coal, zeolites or grafted nanofibrous adsorbents may be operated with various
configurations [16–19]. Both membranes and adsorption processes compete with chemical
absorption in terms of performances. Among these processes we decided to consider one
process of each category to produce a CO₂ stream from the flue gas of the power plant
that could be eligible for direct utilization in the chemical process: chemical absorption
with MEA, which is the most mature process [2], polymer membrane separation, and
temperature swing adsorption (TSA) on activated coal.

Our work was then divided into three main tasks. First, a complete model of the plant
was built using the process simulator Aspen Plus and validated against industrial data [20].
This model includes the different part of the chemical process itself and the coal-fired
power plant as well. Models of the selected CC technologies were then developed using
Aspen Plus and Aspen adsorption [21] and incorporated in a global Aspen Plus flowsheet
representative of the whole plant. Such an approach ensures that all the mass and energy
balances are respected and that the CC facilities considered are well designed to fit all the
requirements and constraints of the considered plant. In the last part of the study, which is
the heart of the present paper, we used the results of the Aspen calculations as a basis to
build the life cycle inventory (LCI) of the different scenarios and we proceeded to a com-
parative life cycle assessment (LCA) of those scenarios. This hybrid methodology, which
associates systems modelling and LCA was already adopted several times [22–25] in other
contexts. It allows the design and the comparison of processes in terms of environmental
impacts with very limited use of generic commercial data bases, which may not be relevant
in such specific configurations. To our knowledge, it is the first time this methodology is
applied to compare chemical absorption, polymer membrane separation and temperature
swing adsorption in the framework of their potential integration on an industrial plant
for CCU.

2. Materials and Methods

2.1. Plant Overview

The chemical plant considered is composed of several processes that can be schemati-
cally gathered in four main parts: the coaled-fired power plant, a pre-process that produces
an intermediate product (Prod I), and two main processes (A and B) producing two different
products (Prod A and Prod B respectively), as shown in Figure 1.

Figure 1. Schematic representation of the considered plant.
The production of Prod A, which is a high-quality grade product for pharmaceutical applications, requires CO$_2$. This CO$_2$ is provided by the pre-process (co-product) and is a limiting reactant in the chemical reactions involved in process A, which means that amount of Prod A produced is bounded by the amount of CO$_2$ collected at the pre-process. Prod B is of lower quality and does not require CO$_2$ for its production. The actual plant (without CC) produces 3.15 tons of Prod B for 1 ton of Prod A. The whole process is supplied in energy by the coal-fired power plant onsite.

In this paper, we consider the possible addition of a CC unit: a limited amount of CO$_2$ from the exhaust gas of the power plant would be recovered to supply process A with extra CO$_2$ and increase Prod A yield (marked in red in Figure 1). At the same time, a reduction in GWP impacts could be expected. The CC facilities would be supplied in energy (electricity and steam) by the power plant.

To be eligible for plant integration, the CC facilities should have a minimal CO$_2$ production capacity of 1 ton per hour with a minimal CO$_2$ recovery of 75%. In addition, the minimal CO$_2$ purity should be 95 mol%, according to specific requirements of Prod A. Given those requirements, around 13 tons of exhaust gas from the power plant would be processed to produce 1 tCO$_2$/h. The detail of the chemical process and its modelling are not described here for confidentiality reasons.

2.2. Power Plant

The coal-fired power plant was modelled on Aspen Plus. The model is schematically divided into two sub models—the boiler and the steam cycle—which are connected.

Coal is considered a ‘non-conventional solid’ described by its chemical composition, diameter (0 to 50 mm), humidity (10%) and lower heating value (28,100 kJ/kg). In the model, the coal is dried before entering a ‘R-Yield’ reactor where it is decomposed into chemical elements. Combustion is then performed into a ‘R-Gibbs’ reactor. The Redlich Kwong-Soave thermodynamic state equation is used to describe the gas phase into the boiler. The hot combustion gas is sent to heat exchangers where its energy is transferred to the steam cycle. The steam cycle is modelled by a seven-stage (two high pressure, two medium pressure and three low pressure) Hirn cycle with overheated steam delivering 20 MWhe. Isentropic efficiencies are respectively 90, 92 and 85% for high, medium and low-pressure turbines, respectively. Electromechanical efficiency is set to 95%. The vapour mass fraction at the end of the cycle is 0.78 at 30 $^\circ$C and 50 mbar. At each stage water is condensed and collected before starting a new cycle.

The exhaust gas is cleaned from dust through filters before being sent to the de-NOx and de-SOx processes. As NOx and SOx are considered to be very harmful to the considered CC facilities, we decided to enhance the de-NOx and de-SOx processes of the actual plant to be able to reach a negligible concentration of NOx and SOx in the flue gas. De-NOx is performed through Selective Non-Catalytic Reduction (SNCR) with ammonia solution (25 wt-%). The flow rate of the ammonia solution was increased by 18% in comparison to the reference process. De-SOx on the plant is achieved using dry slaked lime produced onsite (intermediate product in the pre-process). This process was completed by an additional treatment in a column with a soda solution (20 wt-%). Beforehand the gas was cooled down to 50 $^\circ$C in a cooler. This additional treatment requires 4.75 GJ of steam, 1120 kg of demineralized water and 280 kg of NaOH to produce 1 ton of CO$_2$ per hour; those values are given in the case of membrane separation and slightly change depending on the calculated performances of the considered process.

The main results of the simulation of the power plant are reported and compared to onsite-measured data in Table 1.
Table 1. Results of simulations for 1 MWhe.

| Compounds                  | Model        | Plant Data   |
|----------------------------|--------------|--------------|
| Coal (kg)                  | 1059.8       | 1058.9       |
| Natural gas (kg)           | 23.3         | 21.8         |
| Ammonia (kg)               | 25.3         | 21.4         |
| Demineralized water (m³)  | 7426.1       | 7426.5       |
| Process water (m³)         | 2443.1       | 2445.0       |
| Condensate (m³)            | 787.0        | 803.5        |
| Electricity produced (MWhe)| 1            | 1            |
| Steam produced (GJ)        | 24.5         | 25.0         |
| Ash (kg)                   | 109.7        | 125.6        |
| Flue gas temperature (°C) | 50           | 50           |
| Flue gas pressure (atm)   | 1            | 1            |
| Flue gas composition (%m) |              |              |
| CO₂                        | 8.5          | 8.5          |
| N₂                         | 74.1         | 74           |
| H₂O                        | 7.1          | 7            |
| O₂                         | 10.3         | 10.5         |
| SOx                        | 0            | 0.015        |
| NOx                        | 0            | 0.007        |

The calculated results are in good agreement with measurements. As mentioned above, SOx and NOx concentrations are negligible in our model because of the enhanced de-NOx and de-SOx treatments considered. The extra ammonia required for the enhanced de-NOx treatment explains the difference of ammonia consumption between the model and the plant. The noticeable difference between the calculated and measured ash production should be attributed to the variability of the coal composition on the plant, which changes with the supply, whereas it was fixed in the model. The concentration of CO₂ of the flue gas is relatively low, around 8.5 wt-%. This model was connected to the global flowsheet of the plant on Aspen Plus, and automatically adapts its energy production to supply the various facilities of the plant.

2.3. CC Technologies Considered

- Membrane separation

  The membrane separation process was modelled using MEMSIC, a third-party module [26], which was integrated in the Aspen Plus model. Different configurations were tested and a two-stage arrangement of two different polymer membranes was selected to achieve the specifications [21].

  The polymer membranes were chosen on the basis of their performances, their well-known physicochemical properties, and their availability: polyethylene oxide (PEO), which is highly selective, and Pebax 1657, which is highly permeable to CO₂ [27]. These membranes are associated to a compressor and a vacuum pump as shown in Figure 2. The pressure of the gas fed at the membrane entrance is 1.5 bar and the pressure of permeate is 0.15 bar. The properties of the two membranes, reported in the figure, were calculated from the model to reach the desired performances.

- Chemical absorption with MEA

  The chemical absorption with MEA was modelled on Aspen Plus as well. A rate-based kinetic approach was employed and heat and mass transfers inside each column were considered. Details of the model can be found in [21] and main characteristics of the process are reported in Table 2.
Figure 2. Schematic representation of the selected membrane process.

Table 2. Main characteristics of the chemical absorption with MEA.

| Column Properties | Absorber | Stripper |
|-------------------|----------|----------|
| Column type       | FLEXIPAC, 250Y, Koch, Metal | FLEXIPAC, 250Y, Koch, Metal |
| Volume (m$^3$)    | 58.9     | 45.2     |
| Head losses (bar) | 0.1      | 0.2      |
| Number of stages *| 20       | 20       |

| Operating Parameters |
|----------------------|
| Stripper pressure (bar)| 2.0 |
| MEA conc. (%mass)     | 30.0 |
| Solvent flowrate (m$^3$/h) | 38.0 |
| Extra MEA adding (kg/h) | 1.41 |
| MEA purge (kg/tCO$_2$) | 0.38 |

* including two stages of scrubbing.

- Temperature swing adsorption

The third CO$_2$ capture process, TSA on activated carbon, was modelled [21] with Aspen Adsorption, through a dynamic approach, using cycles for simulating the successive adsorption and desorption steps. For a given column, the cycle consists of four steps: adsorption, regeneration (the bed is heated indirectly by steam circulating in pipes inside the bed), a hot gas purge with CO$_2$ to avoid CO$_2$ dilution and N$_2$ cooling. The process is made of three columns to allow a continuous operation. The main characteristics of the columns and the process operating parameters are indicated in Table 3.

Table 3. Column properties and operating parameters for the TSA process.

| Column Properties |
|-------------------|
| Particle diameter | 3.8 mm |
| Particle porosity | 0.46   |
| Particle tortuosity | 2.2 |
| Columns height    | 1.2 m  |

| Operating Parameters |
|----------------------|
| Adsorption |
| Stop criteria | $x_{CO2} \geq 2$ mol % in the output gas |
2.4. Life Cycle Assessment

The goal of the present study is to evaluate from an environmental point of view the whole plant in its standard operating conditions (without CC, reference scenario) and to assess the potential benefit of the integration of the different CC options onsite. A LCA following the ISO14040 and 14044 standards [28,29] was carried out in both cases using the commercial software GABI 8. The life cycle inventory was mainly established on the basis of the computer simulations conducted on Aspen Plus software described above. These data were completed for standard processes by the commercial database ecoinvent 3.5 integrated to GABI 8.

The functional unit is “production of 1 ton of product A and 3.15 tons of product B”, which is representative of the standard operation of the plant. Calculations showed that with the implementation of CC facilities, the rate of production of product A would be increased of 12% without any change regarding the production of Prod B. This means that with CO2 capture and valorisation, for 1 ton of Prod A, 2.8 tons of Prod B is produced. To stick to the functional unit, an additional production of 0.35 ton of Prod B onsite is considered (boundaries extension).

Four scenarios were considered: reference case without CCU, plant with chemical absorption with MEA, plant with membrane separation and plant with adsorption on activated carbon (TSA).

The other LCA assumptions are as follows. The construction and end of life of the infrastructure of the chemical process and of the power plant are not considered. The coal comes from Russia and is transported to the plant by train and trucks. The collected ashes are buried. The life duration of the membranes is 5 years. The degraded membranes are not recycled but directly incinerated. The activated carbon comes from Russia and has to be changed every 5 years. The degraded MEA from purges is collected and incinerated.

The impact assessment was performed with the method ReCiPe v1.08 midpoint indicators. Considering the nature of the processes at study and the associated emissions, we focused on the following categories of impacts: global warming potential (GWP100), eutrophication potential (EP), freshwater eutrophication (FEP), terrestrial acidification (TAP), particulate matter formation potential (PMFP), photochemical oxidant formation potential (POFP), ozone depletion potential (ODP), human toxicity potential (HTP), freshwater ecotoxicity potential (FETP) and terrestrial ecotoxicity potential (TETP).

3. Results

3.1. LCI

The inventory was calculated on the basis of the results of the simulations conducted with our models. The main flows of materials and energy calculated are listed below, for the different units of the plant.

- Power plant

The principal results of the inventory (per functional unit, FU) for the power plant in the reference scenario are given in Table 4 for the functional unit. Depending on the scenario considered, the production of electricity is of course subject to variations, which are accounted for in the following.

### Table 3. Cont.

| Regeneration |  |
|--------------|---|
| Steam temperature | 180 °C |
| Bed temperature | 150 °C |

| Purge with Hot CO2 |  |
|-------------------|---|
| Duration | 120 s |
| CO2 temperature | 150 °C |

| Cooling with N2 |  |
|-----------------|---|
| Stop criteria | Bed temp. ≤ 30 °C |
| | Gas temperature | 30 °C |
Table 4. Calculated inventory (main results) for the power plant (reference scenario).

| Inputs/FU | Power | Outputs/FU | Energy |
|-----------|-------|------------|--------|
| Materials (t) | Power | Materials (t) | Energy |
| Coal | 1.68 | Electricity from the grid-French mix (GWhₑ) | 6.28 × 10⁻³ BO₂ | 4.54 |
| Demineralized water | 11.83 | Electric (GWhₑ) | 1.41 × 10⁻³ |
| Cooling water | 3.89 | Electricity self-consumed (GWhₑ) | 0.34 |
| Ammonia | 0.005 | | |

- Production processes

The main results for the inventory (per functional unit, FU) of the production processes under the reference scenario are reported in Table 5.

Table 5. Calculated inventory (main results) for the production processes.

| Input (/FU) | Energy (/FU) | Output (/FU) |
|------------|--------------|--------------|
| Pre-process |
| Brine (t) | 6.88 | Elec. (GWhₑ) | 6.9 × 10⁻⁴ |
| Limestone (t) | 7.04 | Steam (GJ) | 1.56 × 10¹ |
| Coke (t) | 1.89 | | |
| Ammonia (t) | 5.31 × 10⁻³ | CO₂ (t) | 9.11 × 10⁻¹ |
| Process water (t) | 1.13 × 10¹ | CO (t) | 1.16 × 10⁻¹ |
| Anthracite (GJ) | 1.19 × 10¹ | | |
| Process A |
| Prod I (t) | 9.11 × 10⁻¹ | Elec. (GWhₑ) | 8.46 × 10⁻⁵ |
| CO₂ (t) | 6.49 × 10⁻¹ | Steam (GJ) | 1.36 |
| Process water (t) | 1.09 | CO (t) | 2.89 × 10⁻² |
| Process B |
| Prod I (t) | 3.52 | Elec. (GWhₑ) | 2.07 × 10⁻⁴ |

- Carbon capture facilities

The flows of materials and energy associated to the use phase of the different CC facilities are given in Table 6.

Table 6. Main material and energy flows associated to the use phase (CO₂ separation) of the CC facilities (per functional unit).

| Membrane Separation | Chemical Absorption | Temp. Swing Adsorption |
|---------------------|---------------------|------------------------|
| PEO (kg) | 5.21 × 10⁻⁴ | MEA in loop (t) | 2.29 |
| Pebax 1657 (kg) | 9.93 × 10⁻⁵ | MEA losses (t) | 6.0 × 10⁻² |
| Process water (t) | 3.50 × 10⁻¹ | MEA purges (t) | 2.0 × 10⁻² |
| Energy (GJ) | 1.74 × 10⁻¹ | Energy (GJ) | 2.3 × 10⁻¹ |

| Membrane Separation | Chemical Absorption | Temp. Swing Adsorption |
|---------------------|---------------------|------------------------|
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| Pebax 1657 (kg) | 9.93 × 10⁻⁵ | MEA losses (t) | 6.0 × 10⁻² |
| Process water (t) | 3.50 × 10⁻¹ | MEA purges (t) | 2.0 × 10⁻² |
| Energy (GJ) | 1.74 × 10⁻¹ | Energy (GJ) | 2.3 × 10⁻¹ |
3.2. Impact Assessment

Figure 3 compares the four scenarios considered in terms of their impacts. In this graph, the reference scenario corresponding to the actual plant without CCU is set to 100% and the other scenarios are reported in terms of their relative environmental impacts.

![Graph showing comparison between CC scenarios and reference case](image)

Figure 3. Comparison between the different CC scenarios and the reference case (set to 100%) for the following impacts: global warming potential (GWP), eutrophication potential (EP), freshwater eutrophication (FEP), terrestrial acidification (TAP), particulate matter formation potential (PMFP), photochemical oxidant formation potential (POFP), ozone depletion potential (ODP), human toxicity potential (HTP), freshwater ecotoxicity potential (FETP), terrestrial eco-toxicity potential (TETP).

Globally, one observes that for all the selected categories of impacts, the results are very close to the reference case or worse; no significant improvement is observed in any category.

Concerning GWP, which is the main concern of the present study, the potential impacts for the four scenarios are in a nutshell. Chemical absorption, membranes and adsorption on activated coal reach 98.2, 97.2 and 97.8% of the reference case, respectively. This negligible and non-significant gain results from two factors: the energy required to run CC facilities, which erases the benefit of the CO$_2$ recycling, and the relatively low amount of CO$_2$ captured and reused onsite. Indeed, the CC facilities were designed to capture 1000 kg/h of CO$_2$, which hardly represents 2% of the standard emissions of the coal-fired power plant. At the same time, the production of Prod A is increased by the CO$_2$ recycling by around 12%, which means that a substantial improvement in term of GWP could be expected, after rescaling to the functional unit. However, it appears that the extra energy required to operate the CC facilities strongly mitigates the results. This is clearly illustrated on Figure 4, which represents the amount of CO$_2$ generated by the power plant to produce 1 kWh with (mean value) and without carbon capture. This graph shows that the implementation of CC facilities results in net CO$_2$ emissions.

Similar conclusions can be derived regarding the particulate matter formation potential (PMFP) category, which is directly linked to the emissions associated to the production of energy with the coal-fired power plant. The category photochemical oxidant formation potential (POFP) is also strongly linked to the energy production through the SOx emissions resulting from the combustion of coal, but one has to keep in mind that the absolute value of the indicator for this category is relatively weak due to the strong de-SOx purification process operated.
Figure 4. CO$_2$ emitted by the coal-fired power plant.

Absorption with MEA exhibits significantly higher results than the other processes regarding the eutrophication potential (EP), freshwater eutrophication potential (FETP), terrestrial eutrophication potential (TETP), ozone depletion potential (ODP) and terrestrial acidification potential (TAP). These results are in agreement with the literature [30]. ODP category is strongly affected by the ethylene oxide associated to the production of MEA, and by the degradation of MEA during CO$_2$ separation, which represents 22% and 78% of the impacts associated to the absorption process in this category, respectively. The degradation of MEA also strongly affects EP, TETP and TAP categories. The end-of-life treatment of the solvent rather impacts FETP and EP.

The membrane technology is slightly better than MEA absorption in most categories of impacts, which is consistent with the results published in [31]. The ozone depletion potential (ODP) is strongly impacted by emissions of chlorofluorocarbides and hydrofluorocarbons during the incineration of the membranes and to a lesser extent by the emissions associated de production of the membranes. These emissions represent 52% and 28% of the impacts associated to the membrane process in this category, respectively. The production of the membranes is also responsible for emissions of hydrogen fluoride, which affects the human toxicity potential category.

Adsorption on activated coal, which appears to be the most attractive alternative, presents impacts relatively close to the reference case in most categories. The differences mainly come from the extra energy required to operate the facility. To our knowledge no dedicated literature investigated the environmental assessment of this technology.

Those results are globally disappointing and especially in terms of global warming potential. The integration of the CC facilities improves the production of Prod A but offers no benefit in terms of CO$_2$ emissions and more or less significantly increases the impacts in all the other categories for the given functional unit.

3.3. Parametrical Study

The previous calculations were conducted strictly meeting the performances criteria of the CC facilities previously presented. As those criteria led to unsatisfying results regarding the goal of the study, we decided to conduct a parametrical study to try to find a compromise. For each process, we identified one key parameter, which could be modified without compromising the separation process. The principal results are presented below.

- **Absorption with MEA**
  
  A significant part of the impacts associated to the MEA process comes from MEA and its degradation products released during CO$_2$ separation. Direct releases of MEA originate from the purge implemented in the process to maintain a constant quality of the solvent. Consequently, reducing the purge rate has a positive impact regarding the MEA releases and the global MEA consumption of the process, but at the same time it decreases the efficiency of the process because of the accumulation of degraded products in the solvent. We conducted calculations for a reduction of 15% in the purge rate and correspondingly the
consumption of extra MEA was reduced by 5%, the CO₂ recovery and the CO₂ purity fell to 73% and 85%, respectively. Those values are still satisfying but unfortunately in terms of environmental impacts, the benefice does not exceed 5% of reduction for ozone depletion potential and freshwater ecotoxicity potential and is not significant for the other categories, especially regarding GWP.

- Membranes

  The impacts associated to the membrane process mainly come from the energy consumption during separation and from the materials (polymers) of the membranes. We investigated thinner membranes switching from 1.5 μm to 0.07 μm for the Pebax 1657 and from 1 to 0.03 μm for the PEO, which resulted in significant material savings. Reducing the membranes thickness maintain a good efficiency of the process but implies a drop of 11% in the CO₂ purity.

  Impacts are globally unchanged except for ozone depletion and human toxicity (canc.), which are reduced by about 5%.

- Adsorption on activated coal

  For this process, the impacts are mainly associated to the energy required to heat the bed and to a lesser extent to the activated coal itself. We tried to reduce the amount of activated coal from 10%, which implied a drop of 13% in the CO₂ recovery. As expected, this modification hardly reduces the environmental impacts.

- Membranes and activated coal with 40% CO₂ purity

  The previous results clearly showed that a significant decrease in the different categories of impacts, and especially in GWP, could not be reached with a simple tuning of processes operating parameters. Nevertheless, we further thought about the minimal performances criteria of the separation processes established at the beginning of the study. The minimal CO₂ production capacity and the minimal CO₂ recovery rate are imposed by technical constraints that cannot be modified. However, it appeared that the CO₂ purity, which was first set to 95% to ensure a standard high purity, could be reduced without damage to the process A. Process A actually requires a very clean CO₂ to ensure the high purity of Prod A required by its market. But this purity is disconnected from the concentration of CO₂ in the feed gas. On the actual plant, process A is fed with a CO₂ stream, which is purified from unwanted components (dust, SOx and NOx) but which is highly diluted: the CO₂ molar fraction does not exceed 40%. Consequently, as the tail gas collected on the power plant is highly purified prior to CO₂ separation, it could be eligible for process A with a molar fraction considerably lower than the 95% considered previously. Two additional scenarios, with membrane process and chemical adsorption on activated coal, were then considered with a CO₂ molar fraction of 40%. Chemical absorption with MEA, which appeared to be the less promising process in terms of environmental impacts, was not considered in the following.

  Reducing the purity requirements to 40% considerably simplify the membrane separation process, as only one stage of Pebax 1657, without recycling loop, is enough to reach this concentration. The total surface of membranes and the energy required to operate the process are drastically reduced (by about 70% and 60%, respectively). The separation process with activated coal is also deeply impacted: a smaller quantity of coal is required (66% less), which leads to a strong cut in the energy required to heat the bed and run the process (60% less). The main characteristics of the new processes are reported in Table 7.
The results for these two alternatives are reported in Figure 5. This graph clearly shows a significant drop of the results in all the considered categories of impacts, compared to the previous calculations with a CO$_2$ purity of 95%. With a CO$_2$ purity of 40%, both membranes and activated coal options lie within the same range of impacts as the reference case. Impacts on GWP are now significantly lower than the reference case, with a 9% reduction for the membranes and an 8% reduction for the activated coal. This drop is mainly attributed to the saving in the energy required during CO$_2$ separation. Consequently, a significant decrease in the particulate matter formation potential impacts, which are directly linked to the energy consumption, is observed as well. For all the other categories, the impacts remain higher than for the reference case, though they are drastically reduced in comparison to scenarios with a CO$_2$ purity of 95%.

For all the categories considered, except for GWP and particulate matter formation potential, the membranes have higher impacts than the activated coal.

4. Discussion

The present study aimed at reducing the impacts on GWP associated to the production of two commercial products (A and B) of a chemical plant, while increasing the production of the most valued product (Prod A) through a partial capture of CO$_2$ from a coal-fired power plant and its reuse as a raw material for the production of Prod A. The calculations
conducted with a high purity CO₂ (95%) showed a significant improvement, around 12%, of the yield of Prod A but the expected decrease of the potential impacts on GWP was unfortunately not achieved. This was explained by an extra requirement of energy to operate the different facilities considered for CO₂ separation, which almost totally erased the benefit of the gain of productivity of Prod A and of the capture of a portion of the CO₂ released into the atmosphere. Furthermore, the impacts in all the other categories (except particulate matter formation potential) were significantly increased, which led us to conclude that despite the higher yield of Prod A, which may have an economic interest for the long term, the investigated options were not satisfying from an environmental point of view.

The parametrical study conducted on a limited set of parameters of the three CC processes considered showed that no significant improvements could be expected without reconsidering the minimal requirements imposed at the beginning of the study. Consequently, a new set of calculations were conducted for the most promising options—membrane and activated coal—with a CO₂ purity of 40%, which could be eligible for process A. With no surprise, the characteristics of the new separation processes are highly simplified (only one stage without recycling for membrane and a smaller column with less coal for adsorption) and led to a drastic saving in the energy consumed for CO₂ separation. With this new configuration, which presents the same advantage as previously in terms of Product A yield, both membranes and activated coal led to a significant drop in terms of GWP, around 9% compared to the reference case without CCU. This result is quite satisfying as only 2% of the CO₂ from the power plant is captured. In addition, all the results for the other categories of impacts are only 5% to 18% higher than the reference case, which is quite less than in the case with a CO₂ purity of 95%.

5. Conclusions

Although there are small differences in the different categories of impacts for membrane separation and adsorption on activated coal, it is not possible to choose one of these processes above the other one. The differences in the results are very weak and the uncertainties both in our modelling and in the impact assessment method do not allow ranking these processes in terms of environmental impacts. Both options are promising and only technical or economic considerations could lead to a choice.

One important conclusion of this study, which is independent of the application, is that CCU should not be taken as an ideal solution to reduce the environmental load of processes. CC technologies have proven their efficiency for drastically reducing the impact on GWP of standard power plants, while increasing the impacts in other categories such as acidification and human toxicity, as highlighted in [30]. However, when the CC is done for CCU, the results are highly dependent on the local context. In our study, despite an actual gain in terms of impacts on GWP (minus 9% for the case with a 40% CO₂ purity), the very low amount of the processed flue gas (only 2% of the CO₂ is captured) drastically limited the potential of environmental gains.

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