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An Outlook on Tar Abatement, Carbon Capture and its Utilization for a Clean Gasification Process

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Abstract: The generation of syngas via gasification is accompanied by greenhouse gas emissions and certain impurities like tar. This is the major problem that makes the technology unattractive for commercialisation. Tar content present in the syngas limits its application as it causes damage to engines, while the CO_2 that accompanies it adds to the climate issues when released to the atmosphere. In this short review, an overview of techniques used for tar abatement, CO_2 capturing, and potential areas of CO_2 utilisation were discussed.

Keywords: tar removal; carbon capture and utilisation; syngas; gasification

1. Introduction

Renewable energy, a clean environment, and a sustainable economy are the three major forces driving the global search for the alternative, replenishable, and lasting energy source 1, 2). The rapid growth of the global population, industrialisation and rapid urbanisation has raised the energy requirement in the last century. Fossil fuel utilisation to catch up with this energy demand continuously generates problems of atmospheric pollution, global warming, ozone layer depletion and climate change, among others 3, 4). Hence, it is noticed that increasing interest in the reduction of greenhouse gasses and fossil fuel usage has triggered research on renewable energy sources ^{5, 6)}. Among these energy sources, biomass energy conversion seems to be the best as it is capable of supplying energy in solid form (bio char), liquid form (bio-oil), and gas (producer/ syngas) which are potential feedstock for the production of chemicals like methanol and ethanol 7,8).

Biomass energy is that which is obtained from plants or plant materials. Biomass can be converted into fuel via thermochemical (combustion, pyrolysis, and gasification) and biochemical (fermentation, methanation and liquefaction) routes ^{9, 10)}. Thermochemical conversion is more preferable to biochemical due to rapid conversion rate and a higher quantity of feedstock ¹¹⁻¹²⁾. Biomass gasification simply means the thermochemical transformation of biomass to gaseous mixtures in the

presence of a gasifying medium, which may be air, oxygen, or steam. The gaseous mixture comprises of methane (CH₄), carbon monoxide (CO), hydrogen gas (H₂) and carbon dioxide (CO₂), sulphur oxides (SOx) and alkaline gases and complex hydrocarbons $^{13-15}$). The fuel can be utilised for producing chemicals and heat and power generation in internal combustion engines. Gasification gives the liberty of using individual feedstock or mixing different ones to achieve co-gasification 16).

Despite the wonderful products of biomass gasification (H₂, CO, and CH₄), the process is associated with some unwanted substances which have turned out to be archrivals of gasification 17). These unwanted substances are the complex hydrocarbons commonly referred to as tar and carbon dioxide (CO₂), which accompany the producer gas, as illustrated in Fig.1. The tar and the CO₂ contents limit the usage of the syngas obtained, and they have made gasification technology unattractive from a commercial perspective. Therefore, they are the significant problems of gasification that need to be addressed. The tar is formed at the pyrolysis stage during decomposition and therefore is released in the form of an aerosol, condensing to a sticky component. On the other hand, the CO₂ usually develops as a constituent of the gas stream (syngas) and therefore has to be captured. While the tar is responsible for various damages in the downstream usage like blockage and fouling in turbines and engines, cracking in filter-pores, low gasification efficiency and syngas calorific value, cocking, and

deactivation of catalyst 18), the CO₂ limits the efficiency of the syngas as it is a non-flammable gas. Both tar and CO₂ concurrently accompany the syngas during gasification, and therefore addressing the two issues simultaneously becomes imperative. Improved gasification systems incorporated with CO2 capture and sequestration mechanisms can fully store up to 90% of the CO₂ emitted from facilities that use biomass/fossil fuels to produce power, H₂ or syngas. Most importantly, such technologies are referred to as negative emissions technologies (NETs) and are essential contributors to curbing down CO₂ emissions from the environment ¹⁹⁾. New emerging technologies like the sorption enhanced gasification (SEG) process allow gasification and CO₂ capture simultaneously by applying a solid sorbent. SEG is therefore influenced mainly by sorbent properties (e.g. type and activity).

Depending on the gasifier type, the tar content ranges from 1 g/m^3 to $100 g/m^3$. The produced tar results to lower gas yield, high maintenance and operation challenges such as corrosion, blocking and clogging of fuel pipes, filters, nozzles, and turbines $^{20-22)}$. Sasujit $^{13)}$ argued that high tar

modification (separation of pyrolysis and reduction zones), optimised process conditions (temperature, gasification medium, pressure, additive/ catalyst, residence time and mass ratio between fuel and gasification medium). Hence this method limits or decreases the potential of tar formation during biomass gasification.

On the other hand, post-gasification abatement applies after the producer gas has been produced. Then, the tar is collected through physical processes involving cyclones, filters scrubbers, electrostatic precipitators or chemical treatment by catalytic and thermal processes and partial oxidation to purify the gas. Combining both methods is recommended for maximum tar removal with good quality product gas ²³⁻²⁵).

High greenhouse gas emissions such as CO₂ that accompanies the producer gas stream are harmful to the ecosystem ^{26, 27)}. By implementing carbon capture, it can be handled as an essential raw material for the manufacture of several value-added fuels and chemicals, thus curbing emission problems, energy supply challenges, and multiplying the biomass fuel conversion yield. The idea of carbon capture, utilisation and sequestration

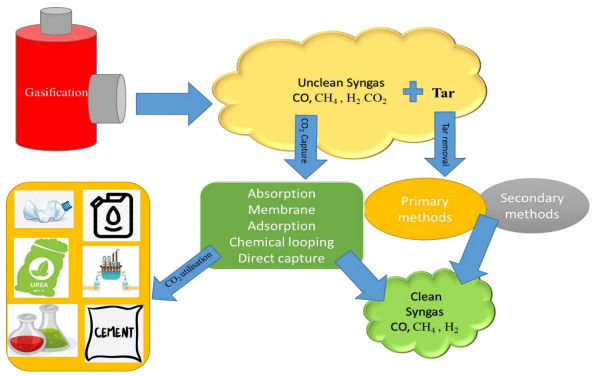


Figure 1. Schematic representation of CO₂ and tar removal during gasification

content discourages the implementation of the technology in high-efficiency IC engines and fuel cells where utmost priority is given to an extremely clean gas stream. The tar is a combination of different chemical compounds ¹¹⁾ as demonstrated in Fig. 2. There are numerous methods of tar abatement which are broadly classified into two: (1) Primary (in-situ) methods and (2) secondary (postgasification) methods. In-situ reduction avoids tar formation, and is attained either by gasifier design

(CCUS) are concepts of CO₂ regulation and climate crisis. Carbon capture and storage (CCS) deals with the mechanism of selective ejection of CO₂ from producer gas streams, compressing it to a supercritical condition, after which it finally get transported and sequestrated in exhausted oil and gas reservoirs and geologic formations ^{28,29)}. The challenging fact with CCS is the high cost associated with capture and compression, which consumes about 75% of the entire cost of the CCS process,

and has highly impacted its large scale implementation. As a more promising and appealing substitute, carbon capture and utilisation technologies (CCU) have lately dominated the scene for the credit of transforming the captured CO2 into a precious asset as a substitute for permanent sequestration. Styring et al. 30) argued that CCU manipulates captured CO₂ as a renewable energy resource to replace fossil fuel resources. Even though CCU is more advantageous than CCS, it is faced with the challenges of thermodynamic stability during the CO₂ conversion and utilisation in chemical reactions. CO2 capture technologies are classified into two groups: (1) postcombustion which refers to direct CO₂ dismissal from flue streams and (2) pre-combustion which refers to using improved low carbon comprehensive combustion systems which involves an integrated gasification combined cycle (IGCC) and oxyfuel gasification (addition of uncontaminated oxygen to lower the CO₂ concentration of power generation) ^{28, 31)}.

This paper provides a short review of the most recent developments in carbon capture and tar abatements as methods that will purify producer gas stream obtained after biomass gasification. The concept has been demonstrated in Fig.1. The ultimate goal of producing syngas is to use it as fuel for power generation or feedstock for chemical synthesis. This may only be achieved through proper cleaning of the gas to remove the tar and capture the CO₂ ^{32, 33)}. The paper gives an overview of the different pathways and the mechanisms involved in removing both tar and CO₂ according to past literature and further provides the potential application of the captured carbon. The paper is organised in the following order; section 1 introduction, section 2 biomass gasification, section 3 tar definition, formation, classification and reduction methods. Section 4 carbon capture separation techniques and utilisation of the captured CO2, and lastly section 5 conclusion.

2. Biomass gasification

The term "gasification" refers to the thermochemical transformation of organic materials/solid fuels within a vessel referred to as the gasifier. This results in gaseous products commonly called producer gas. The producer gas is a mixture of gases such as carbon monoxide (CO), hydrogen (H_2) , methane (CH_4) and carbon dioxide (CO_2) , volatile organic matter, water vapour, hydrogen sulphide (H₂S), tar, char, and ash ^{25, 34)}. The reaction takes place within temperature ranges of 800-100 °C, within a medium which may be air, oxygen, steam and flue gases ^{35, 36)}. Details of gasification reactions are given in Table 1. The average calorific value of the producer gas obtained from biomass gasification ranges from 4-10MJ/m³, while the carbon conversion efficiency ranges from 50-70% ³⁷⁾. The gasification process involves four stages as summarised below:

> Drying/Vaporisation stage: The earliest stage in gasification where excess water is removed

from the feedstock, at temperature ranges of $70\text{-}200~^{\circ}\text{C}~^{38)}$. However, no thermal decomposition takes place in this zone as a result of low temperature $^{39)}$.

Drying: Moist feedstock + Heat \rightarrow Dry feedstock + water

- Devolatilization/pyrolysis: This stage occurs at a temperature of about 300-500 °C, and involves the thermal breakdown of materials in an inert condition, which results in the emergence of liquid tar, solid char and gases, such as CO, CO₂, H₂, CH₄, H₂O, etc. Pyrolysis: Dry feedstock + Heat → Char + Volatiles
- Combustion/ partial oxidation: Due to the presence of oxygen in this zone, the combustion reaction occurs with the volatile products and char obtained during pyrolysis. The reactions in this stage are highly exothermic in this zone and occur at a temperature greater than 500 °C. Varying reactions occur between solid carbonized fuel and oxygen, resulting in carbon dioxide and a considerable amount of heat. Hydrogen also combines with oxygen to produce water vapours.

$$C + O_2 \rightarrow CO_2 + 406 \text{ kJ/g mole}$$
 (1)
 $2H_2 + O_2 \rightarrow 2H_2O + 242 \text{ MJ/kg mole}$ (2)

• Reduction/gasification: The char is converted into gases by reacting with hot gases coming from the oxidation zone at about 800 to 1000 °C, resulting in the transformation of the sensible heat of the solid char and gases into the chemical energy of the producer gas. During this process, some combustible products are generated such as, CO, CH₄, and H₂. In addition, the following chemical reactions occur in the reduction zone ³⁹⁾:

$$\begin{array}{cccc} C & + & CO_2 & \rightarrow & 2CO-172.6 & kJ/g & mole \\ (Boudourd) & & (3) \\ C & + & H_2O & \rightarrow & CO & + & H_2 & - & 131.4 & kJ/g & mole \\ (steam reaction) & & (4) \\ CO & + & H_2O & \rightarrow & CO_2 & + & H_2 & + & 42.3 & kJ/g & mole \\ (Water gas shift & & (5) \\ C & + & 2H_2 & \rightarrow & CH_4 & + & 75 & kJ/g & mole \\ (Methanation reaction) & & (6) \end{array}$$

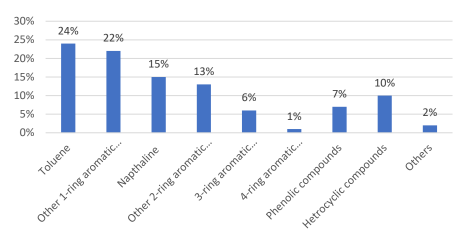


Figure 2. Typical composition of biomass tars $^{11)}$

Table 1. The gasification process reaction of ordinary biomass 40-42)

Process	Type of reaction	
Volatile matter = $CH_4 + C$	Partially exothermic	Char oxidation
$C + \frac{1}{2}O_2 \to CO$	$\Delta H^* = -110.60 \text{ kJ/mol}$	
$CO + O_2 \rightarrow CO_2$	$\Delta H^* = -393.60 \text{kJ/mol}$	
$CO + \frac{1}{2}O_2 \rightarrow CO_2$	$\Delta H^* = -283 \text{ kJ/mol}$	Oxidation of volatiles
$H_2 + \frac{1}{2}O_2 \rightarrow H_2O$	$\Delta H^* = -241.90 \text{kJ/mol}$	
$C + CO_2 \rightarrow 2CO$	$\Delta H^* = 172.50 \text{k J/mol}$	Boudourd
$C + H_2 O \leftrightarrow CO + H_2$	$\Delta H^* = 131.30 \text{ kJ/mol}$	Primary water gas
$C + 2H_2O \leftrightarrow CO_2 + 2H_2$	$\Delta H^* = 90.20 k J/mol$	Secondary water gas
$C + 2H_2 \leftrightarrow CH_4$	$\Delta H^* = -74.90 \text{ kJ/mol}$	Methane formation
$CO_2 + 4H_2 \leftrightarrow CH_4 + 2H_2O$	$\Delta H^* = -165 \text{kJ/mol}$	
$C + 3H_2 \leftrightarrow CH_4 + H_2O$	$\Delta H^* = -206 \mathrm{kJ/mol}$	
$CO + H_2O \leftrightarrow CO_2 + H_2$	$\Delta H^* = -41.2 \mathrm{kJ/mol}$	Water gas shift

Gasifiers are broadly grouped into three, namely, entrained flow, fixed bed, and fluidised bed gasifiers, subtypes exist and are given in Table 2. The classification is based on the arrangement of fuel and produced gas flow and the contact mode of fuel and gasification agent. In the fixed bed gasifier, there is intimate mixing between fuel particles and oxidising agents. The fuel travels naturally along the four stages (drying, pyrolysis, combustion, gasification) and makes a bed of solid fuel, over which the gasification agent moves freely. The fluidised bed

fluidises the fuel with low-pressure gas or air, allowing more contact between the fuel and oxidiser for the reactions. The particle contact is low in entrained bed type, as the fuel particles are made smaller and fed through an atomiser to ensure maximum surface contact with the oxidisation agent. Further classification may be also due to the source of heating which may be allo-thermal that is externally sourced heating or auto-thermal if the source of heating is as a result of partial combustion of the fuel ²⁴).

Table 2. Gasifier Types ^{24, 37, 43)}

	Gasifier	Gasifier sub-	Gasification	Suitable	Tar	Feed stock
	types	types	temperature	scale		preparation
			°C			
	Fixed/	Downdraft	700-1200	Small-	Low	Very critical
Gasification	Moving bed			medium		
technologies		Updraft	700-900	Small-	High	critical
				medium		
		Crossdraft	-	-	-	-
	Fluidised bed	Bubbling	< 900	Small-large	Medium	Less critical
		Circulating	1450	Medium-	Medium	Less critical
				Large		
		Twin bed	-	-	-	-
	Entrained	Coaxial down-	1450	Medium-	Very low	Only fine
	Flow	flow		large		particles
		Opposed jet	-	-		-

3. Tar

In the spring of 1998, a meeting was held in Brussels by the Energy Department of United States, the Directorate General of Energy of the European Commission (DG XVII) and the International Energy Agency (IEA) about tar quantification protocol, and was agreed that tar refers to all hydrocarbons whose molecular weight was greater than benzene (C_6H_6) ⁴⁴. Devi et al. ⁴⁵ defined tar as a combination of condensable hydrocarbons, including aromatic compounds with up to five rings that may be oxygenated and Polycyclic Aromatic Hydrocarbons (PAHs).

3.1 Tar Formation, content and categorisation

Tar formation is initiated at the pyrolysis zone when the biomass fuel decomposes into char, liquid (primary) tar and volatiles at a temperature range between 200 to 500°C. Generally tar produced during producer gas condensation may be classified as primary, secondary and tertiary compounds. Primary tar is formed at temperatures below 500 °C, while at higher temperatures in the oxidation zone, the primary tar reforms to miniature molecules, noncondensable gases (CO, H₂O, CO₂) and larger molecules (secondary tar). At a later stage with a higher temperature about of 800 °C, the secondary tar decomposes completely and tertiary tar is produced ²⁵⁾. Another classification of tar was developed by the Applied Scientific research center of Netherlands, Energy research center of Netherlands (ECN) and the University of Twente

Netherlands, based on solubility and condensability of different tar compounds. The two classifications, however, complement each other and are both critical 25). The classification is presented in Table 3. Several researchers claimed lignin as the primary antecedent of tar as a result of its aromatic nature with the main components vanillin (4-hydroxy-3pyrolysis as methoxybenzaldehyde, $C_8H_8O_3$), guaicol (1,2methoxyphenol, $C_7H_8O_2$), catechol dihydroxybenzene, C₆H₆O₂) and anisol (metoxibenzeno, C₇H₈O) ⁴⁶⁻⁵¹⁾. However, Sasujit et al. claimed that tar composition depends on the temperature of the reaction, type of the reactor and the feedstock. For biomass tar, the main components are naphthalene, ethylbenzene, and acenaphthylene 13). Feng et al. 52) used Gas Chromatography Mass Spectrometry (GC-MS) analysis to examine the evolution of tar during homogeneous conversion at temperature ranges between 500 and

900 °C . It was discovered that at temperatures below 600 °C, the highest proportion of the tar is composed of components based on the primary biomass tar consisting of oxygen and compounds like levoglucosan and dimethoxymethane. At temperatures greater than 700 °C, the transformation of the primary pyrolyzed tar takes place, and the composition becomes mainly made of aromatic compounds like naphthalene. Temperature addition led to the reduction or total tar removal containing branched compounds and the subsequent formation of PAHs. Some tar components break down into more minor molecular gases and C1~C5 hydrocarbons. As the temperature reaches 900 °C, PAH constituents like naphthalene, phenanthrene, and anthracene increased, leading to the evolution of more stable aromatic hydrocarbons.

Table 3. Classification of tar compound 53)

Tar class	Class name	Property	Representative compounds		
1	GC-undectable tar	Extremely large tar	Obtained by subtracting the gas		
		compounds, not visible	chromatography-detectable tar fraction		
		under a GC-FID or GC-MS	from the total gravimetric tar		
		equipped with a non-polar			
		capillary column			
2	Heterocyclic	Tars containing heteroatoms;	Pyridine, phenol, cresols, quinolone,		
		highly water-soluble	dibenzophenol		
		compounds			
3	Light aromatic (1	Usually light hydrocarbons	Toluene, ethylbenzene,xylenes,styrene		
	ring)	with a single ring; do not			
		pose a problem regarding			
		condensability and solubility			
4	Light PAHs	2 and 3 ring compounds;	Indene, naphthalene, methylnaphalene,		
	compounds (2-3)	condense at low temperature	biphenyl, acenaphthalene, fluorine,		
	rings	even at very low	phenanthrene, anthracene		
		concentration			
5	Heavy PAHs	Compound larger than 3	Fluoranthene, perylene, pyrene,		
	compound (4-7	rings, these components	coronene, chrysene		
	rings)	condense at high			
		temperatures			

3.2 Tar reduction

Effective use of producer gas in different applications

such as gas turbines, internal combustion engines, fuel cells, methanol, and ammonia synthesis depends on its tar level. Therefore, the acceptable limit of tar level is recommended to be between 50-100 mg/Nm³. Tar limits for various applications is given in Table 4. Hence it becomes imperative to bring down the level of tar within producer gas to enable its use for multiple applications. As mentioned earlier, there are two methods of tar reduction: the in-situ (primary) method and the post-gasification (secondary) method.

3.2.1 In- situ (primary) methods

In-situ methods refer to precautions implemented during the gasification stage to completely avoid tar formation or break down the tar formed in the gasifier. This is possible through the adoption of proper operating conditions and gasifier design.

3.2.1.1 Operating conditions

Important operating conditions such as temperature, pressure, residence time, gasification medium, catalyst/ additives and mass ratio between fuel and gasification medium play a vital role in the resulting gasification quality. These factors affect the syngas quality, tar formation and carbon conversion efficiency 45). The gasification temperature and pressure play a significant role in the reduction of tar as it enhances cracking. Several studies have depicted that with an increase in temperature, the levels of tar in producer gas decrease. Chan et al. gasified sawdust in air and increased the temperature from 780-857 °C, and the tar produced dropped from 2.0-0.3 g/Nm³ 54). The same scenario is also valid on the effect of pressure, as the more the gasification pressure, the less the tar. Tuomi et al. 55 demonstrated the impact of pressure on tar decomposition activity of different catalysts. As the pressure was increased from 1-10 bar, the catalytic activity for olivine and sand increased thereby increasing tar conversion. The gasification medium, which may be air, steam or oxygen, affects the production of tar. Rios et al. ²⁵⁾ reported that more tar is produced when steam is used while less is obtained when air is used. This may be due to the lowering of the temperature by steam injection. Air fuel ratio also influences tar production. When air is used as the gasification medium, the parameter is referred to as the equivalent ratio. It is reported that with increased ER, tar content decreases as a result of more oxygen availability to break the volatile compounds in the pyrolysis zone. In terms of residence time, Rios et al. ²⁵⁾ further reported that with longer residence time, 1 and 2 aromatic ring compounds decreased significantly, resulting from long exposure of tar compounds to oxidising medium, resulting in a 75% of tar reduction. Catalyst also plays a significant role in tar removal and have proven to be effective. Catalytic cracking can transform tar to low molecular weight hydrocarbons and valuable gas products (H₂ and CO). According to Islam ¹⁸⁾, tar abatement is best achieved through the use of catalyst, and these catalysts have been characterised with the following properties;

• It is necessary that the catalyst must be efficient

for tar reduction in a gas stream having maximum amounts of H_2 , CO, CO_2 , and H_2O within temperature limits of $600~^{\circ}C-900~^{\circ}C$.

- If syngas is the target output, then the catalyst should be capable of methane reforming
- The catalysts should provide a suitable syngas ratio (H₂/CO) for the intended process.
- The catalysts should withstand deactivation in the event of carbon fouling, sintering, and poisoning.
 - It should be simple to regenerate the catalyst.
 - The catalysts should be attrition resistant.
- The catalysts should be affordable and easily accessible.
 - It should not be environmentally hazardous.
- The catalyst should be resistant to sulfur poisoning.

Table 4. Permissible tar levels in producer gas for different applications ²⁵⁾

Application	Tar Limit (mg/Nm³)	
Direct combustion	No defined limit	
Internal combustion engine	<100	
Gas turbine	<5.0	
Synthesis of methanol	<0.1	
Compressors	50-500	
Fuel cells	<1.0	
Fischer Tropsch synthesis	<1.0 μL/L (class 2,BTX)	

3.2.1.2 Gasifier design

The type of gasifier and its design affects the level of tar in a producer gas ⁵⁶⁾. This might be a result of the gasifier configuration. In the updraft gasifier, the produced gas with a mixture of tar escapes from the top of the gasifier. And as a result of heat exchange with cooler surfaces of the gasifier, conversion of the primary tar to secondary tar and gases becomes minimal. This results in producer gas with a high tar content of about 50 g/Nm³. In downdraft gasifiers, the produced gas passes through the bottom of the gasifier along with built tar. In so doing, the primary tar passes through the combustion zone. The high temperature and sufficient oxygen present in that zone enable tar to crack into condensable gases. This results in producer gas with a less tar content <20 g/Nm³. When fluidised bed gasifiers are used, lower tar content is recorded due to the high level of mixing between the bed material and fuel. This results in high biomass conversion to gases with minimum tar. For bubbling and circulating fluidised bed, tar content ranges from 5-20 g/Nm³ and 1-5 g/Nm³, respectively. In an entrained flow gasifier, the tar level is very low as the produced tar passes through a very high-temperature region > 1000 °C. The separation of the

gasifier zones results in a multistage process that is different zones of drying, pyrolysis, reduction and gasification. This leads to increased efficiency of the process with qualitative producer gas 25). An effective reduction in tar level of about 10° mg/Nm³ has been reported by Cao et al. 57), and this was achieved by developing a lab-scale combined 2-region reactor gasifier. Galindo et al. 58) also conducted a similar study where tar level was brought down by introducing air in 2-stage gasification, which triggered the temperature within the reactor. Hence this influenced thermal tar cracking, thereby lowering 87% of tar content in the syngas. Some gasification systems feature separate zones of drying, pyrolysis, gasification and combustion, and the multiple stage process has been found to reduce the amount of tar. Xiao et al. 59) constructed a dual-chamber Internally Circulating Fluidized Bed Gasifier (ICFG) having separate functions for gasification and combustion with bed material circulating between two chambers. The resulting syngas had minimum tar. A similar low tar result was also reported by Burhenne et al.600 where a fixed bed gasifier with four different zones was used for syngas production

3.2.2 Post-gasification (secondary method)

Post-gasification methods are applied after the completion of the gasification process, and product gas has been collected. The process of removing tar from the gas may be a through physical process or partial oxidation, thermal and plasma cracking.

3.2.2.1 Physical/ mechanical process

The physical cleaning methods collect the tar in a condensed form and are divided into two depending on the temperature of the gas. The dry gas cleaning method is employed at a temperature range of 200-500 °C or 600-800 °C. However, it is advisable to cool the gas when using fabric filters as elevated temperatures affect the resistance of bag material fabric 61). Equipment used for dry cleaning includes cyclonic separators, electrostatic precipitators, ceramic filters, bag filters, and catalytic filters. The wet gas cleaning is conducted at a temperature range of 20-60 °C when the gas temperature cools down. Equipment used for wet cleaning includes the spray towers scrubbers, impingement, wet cyclone, venturi scrubbers and wet electrostatic precipitators. A combination of dry and wet methods has also been developed, referred to as the OLGA method 62). The OLGA technique removes and reuses valuable tar components with minimum operational cost. Its operating temperature lies in the range of 60- 450 °C. Oil is utilised as the absorbent medium instead of water. Like water scrubbing, tars classified as 1, 4, and 5 are regenerated by condensation as the temperature is lowered below the tar dew point. Lighter tar compounds classified as 2 and 3 are subsequently removed via absorption into a second liquid scrub. Compared to conventional dry and wet procedures, the OLGA process brings numerous benefits. In addition to eliminating waste treatment, this strategy overcomes the technical and financial challenges of catalytic and high-temperature tar removal. Highly poisonous PAHs are always an issue; however, these tars are usually easier to extract from the water because of their volatility and limited water solubility. The main problem is highly soluble tar and phenol in particular. They easily dissolve in water hence challenging to remove. The removal of these tars using oil prevents costly wastewater treatment. The oil may then be regenerated simply or utilized as a feedstock ²⁵⁾.

3.2.2.2 Thermal oxidation plasma and thermal cracking

The outlined methods are only applied to reactors with high-temperature zone 900-1300 °C. Partial oxidation is achieved through the addition of air or O2 in oxidation reactors. It proves to be an effective way of tar removal when it is conducted at a very high temperature. In a study by Zheng et al., partial oxidation was performed between 600 to 1400 °C, and the results indicated tar reduction with temperature increase 63). Lately, plasma, the fourth state of matter consisting of excited atoms and molecules, a mixture of ions, radicals, neutral particles, is now used for tar reduction 13). It is classified into two; thermal and non-thermal plasma. For thermal plasma-like gliding arc discharge and arc discharge, the gas temperature exceeds 1730 °C. In the case of non-thermal plasma, such as corona discharge/ and dielectric discharge, the electrons temperature may get up to 104-105 °C while the gas temperature may be the same as that of the surrounding. High tar abatement efficiency could be achieved by implementing plasma technique despite some drawbacks like high cost, having a limited lifetime for pulse power supplies and a high energy requirement from the overall process. When a catalyst is incorporated in a plasma reactor, there seems to be a synergy that leads to increased tar conversion into valuable energy and syngas products. This is because catalysts can enhance water gas shift and steam reforming reactions that stimulate the production of H2 and CO. Hence, a synergistic effect is likely to result through a combination of non-thermal plasma with a catalyst ²⁵⁾.

4. Carbon capture

A sufficient amount of carbon dioxide is generated during thermochemical processes like gasification and combustion of natural gas, which is either released to the atmosphere or utilised in manufacturing plants or food processing industries ⁶⁴). Likewise, in the case of thermochemical processes, especially biomass gasification, a considerable amount of CO₂ (about 20%) is released, and most of it goes back to the atmosphere. Quite a number of methods, either chemical or physical, have been applied to control CO₂ emission in gasification

and combustion processes ⁶⁵⁾. Amongst these methods, two chemical techniques have been proven to be the best and most conventional in terms of gasification and combustion processes owing to their high efficiency. These are the calcium looping method that results in a low CO₂ gas stream ⁶⁶⁾ and chemical absorption via monoethanolamine (MEA), widely used as a commercial method on an industrial scale ⁶⁷⁾. In addition to capturing the CO₂ for storage in geological formations, its utilisation will influence the commercialisation of the biomass thermochemical conversion processes ⁶⁸⁾. This can be achieved by compressing the CO₂ to a dense fluid (liquid state) and transporting it via pipelines, leading to an underground storage facility (geological formations).

4.1 Carbon capture separation techniques

4.1.1 Absorption

Absorption based CO2 removal is performed by the application of physical or chemical solvents. Chemical solvents like aqueous ammonia, amine-based solvents such as monoethanolamine (MEA), diethanolamine (DEA), and N-methyldiethanolamine (MDEA) and alkaline solvents such as Ca(OH)2 and NaOH, are used in post-combustion capture ²⁸⁾. This post-combustion capture is achieved via the stripping process, where CO2 and the chemical solvent (saturated) are introduced to heat, producing a new solvent and releasing the CO2 at the tip of the stripping chamber ⁶⁴⁾. In the physical methods for CO₂ capture, the gas is made to come in contact with the solvent stream, and the CO2 is being captured by the solvent physically. The physical methods have been wellestablished not long ago and include Selexol, Rectixol, Purisol, and Fluor. Lately, ionic liquids have been recognised as potential physical solvents due to their properties, including elevated thermal stability at high temperatures, low volatility, low vapour pressure, and low energy requirements. However, the main hindrance to their utilisation is their low working capacity. The challenge facing the absorption technology is that high energy is required for solvent regeneration. Other challenges include corrosion and a large volume of water requirement. It is recommended that solvents with a high heat of absorption (>60 kJ mol⁻¹) may reduce energy consumption while utilisation of thermally stable solvents with low regeneration energy requirements can significantly improve the thermodynamic efficiency of the separation process. Additionally, impurities in the gas streams quickly deteriorate the chemical solvents; therefore, high tolerant solvents should be considered in such applications ²⁸⁾.

4.1.2 Membrane

Membrane separation is established on the Knudsen diffusion principle and occurs under steady state conditions with pressure difference across the membrane driving the permeation process. The separation performance of a membrane is influenced by its structure,

morphology, make-up and operating conditions. Membrane separation is more suitable for high-pressure pre-combustion techniques such as IGCC, and less ideal for low-pressure post-combustion applications. Different membrane types exist, including the porous inorganic membrane composed of zeolites, metal organic frameworks (MOFs), carbon molecular sieves (CMS), ceramics, and a few oxides. Inorganic membranes can endure elevated temperatures and have high mechanical stability, but their high cost lowers the chances of commercialisation. Their main limiting factor to their industrial scale usage is their long term stability, reliability and fabrication route, which is very expensive. There is also the polymeric membrane which has a high potential for industrial usage with a high packing density greater than that of inorganic membranes. However, some of their disadvantages in CO2 capture include low separation performance, low CO₂/N₂ selectivity and permeability for post-combustion processes, a limited imbalance between permeability and selectivity, swelling, deterioration, reactive to impurities, and mechanical stability, especially high-pressure operations. Facilitated-transport membranes (liquid membrane, ion exchange membrane) are also recommended for CO₂ capture. However, their drawbacks include contamination by SOx and NOx and long term stability. The mixed matrix membrane originates from dispersed highly selective molecular-sieve particles such as zeolites and carbon nanotubes in a polymer matrix. They are currently at the initial stage of development and are very expensive and complex to manufacture 69).

4.1.3 Adsorption

Adsorbents are porous solid materials used to remove CO₂ from gas streams and are classified into high temperature and low temperature materials. The high temperature materials are all chemisorbents, including hydrotalcites, alkali/alkaline earth oxides like calcium oxides, alkali silicates, and double salts. Low temperature adsorbents are physisorbents like carbon based materials such as graphene, carbon nanotubes, carbon nanofibres, porous polymer networks, molecular sieves, and covalent organic frameworks. It is desirable that adsorbents are cheap, have a high working capacity, low regeneration requirement, stability and rapid kinetics. Also, for optimum efficiency of the process, factors like cycle time, pressure, temperature and bed quantity, cycle configuration and a number of steps and number of the bed are among the important parameters to be into account for effective capture ^{28, 70, 71)}. One of the recent upcoming technologies as mentioned earlier is the SEG in which the use of limestone especially CaCO₃ has been applied due its ability to capture carbon in gas stream during gasification. Optimisation of the CO2 capture in the gasifier and the release of CO2 in the combustor, requires operation of the reactors at particular temperature settings as defined by the CaO/CaCO₃ equilibrium. Major chemical reactions that take place during SEG process are enumerated ¹⁹⁾:

$$\begin{array}{ll} \textit{CaO} + \textit{CO}_2 \leftrightarrow \textit{CaCO}_3 & \Delta \textit{H} = -178 \; kJ \; mol^{-1} \\ \textit{CaCO}_3 \leftrightarrow \textit{CaO} + \; \textit{CO}_2 & \Delta \textit{H} = +178 \; kJ mol^{-1} \\ \textit{CO} + \textit{H}_2 \textit{O} \leftrightarrow \textit{CO}_2 + \; \textit{H}_2 \; \; \Delta \textit{H} = -41 \; kJ \; mol^{-1} \\ \end{array}$$

The effectiveness and cost of the adsorption process depend on adsorption processes like pressure/temperature swing adsorption, which are determined by the properties of the adsorbents, with process design and operation factors. For successful gas separation in industrial applications, adsorbents are required to be cheap, possess quick regeneration ability, adequate stability, capability and selectivity, low cost, low regeneration requirements, and high reactivity. Process parameters for maximum capture include cycle setting and duration, amount of steps and beds, and operation conditions (pressures and temperatures). Almamoori 28) related that adsorption-CO₂ capture could overcome most of the difficulties faced by absorption processes. However, the currently established technologies are not economically feasible at their present developmental stage. Industrial scale operation is yet to be fully implemented. In designing, developing and evaluating high-performance adsorbents, realistic performance and process considerations should be given priority.

4.1.4 Chemical looping

Chemical looping allows the separation of CO2 and H₂O from the gas stream. It also aids in minimising NOx emission. Chemical looping is combined with IGCC in the case of pre-combustion capture to produce syngas. The technique utilises a metal oxide as an oxygen driver between the air and fuel reactors. Hence in the case of an extensive application, the availability of suitable oxygen drivers with high reactivity in terms of oxidation and reduction, stability, resistance to agglomeration, and the high melting point seem to be challenges of the chemical looping process. Oxides of transition metal-like calcium (Ca), iron (Fe), copper (Cu), cobalt (Co), molybdenum (Mo), manganese (Mn), chromium (Cr), niobium (Nb), vanadium (V), cerium (Ce) and indium (In) are mainly employed for chemical looping. These transition metals used in this process are expected to withstand agglomeration and high temperatures, possess high oxidation/reduction activity, are mechanically stable, and should be economically and environmentally friendly. On the other hand, the challenge of this system is the highpressure requirement to attain maximum efficiency, and most of the chemical looping technologies to be applied for power generation are still at the lab scale under investigation 70).

4.1.5 Direct CO₂ capture from the atmosphere

When CO_2 is removed directly from the atmosphere or ambient air, it is called direct capture (DAC). It has

recently become an attractive technology as it avoids the need to transport captured CO2 from emitter site to sequestration sites. If widely adopted, it can reduce the overall CO2 level in the atmosphere. The technology is quite similar to adsorption-based capture but has the challenge of material requirement with strong binding affinities and high CO₂/N₂ selectivity. Several materials like NaOH, CaOH, potassium solutions and solid materials, including alkali and alkali-supported carbonates, anaionic-exchange resins, functionalised metal oxides and MOFs have been evaluated for DAC 72).

4.1.6 CO₂ captured by hybrid process

Hybrid CO₂ capture, as the name implies, combines two capture subsystems or more simultaneously. The hybrid technology combines two or more separation units in series or parallels to simultaneously optimise the separation efficiency and lower the cost of the separation. An example of such a hybrid system is that developed by American Air Liquide where the CO₂ capture is done through the hybrid membrane-cryogenic distillation process ^{28).} Also, hybrid concepts of membrane-pressure swing adsorption and membrane-distillation have been investigated. The hybrid capture may be considered to deliver novel methods that will be effective and economically viable. Researchers must focus on feasibility, economic viability, material availability, environmental concerns and process conditions⁷³⁾.

Most of the capture methods highlighted are still at the development stage and have not yet reached industrial scale. Thus engineers and material scientists have to work together on the material properties and process performance to develop feasible, cost-effective and functional CO₂ mechanisms. Some of the prospects and challenges of the capture techniques are highlighted in Table 5.

4.2 Captured CO₂ utilisation

The transformation of the captured carbon into valuable products is a very attractive way of embarking on the carbon capture culture, rather than releasing it freely into the atmosphere. To ensure sustainability, the process has to be eco-friendly safe and affordable. Table 6 highlights the opportunities and challenges rendered by the various ${\rm CO}_2$ utilisation technologies.

Table 5. Prospects and Challenges of CO_2 capture methods $^{28)}$

Capture Method	Prospects	Challenges		
Absorption	Ionic liquids and advanced amines	Equipment deterioration, amine degradation, increased		
	utilisation, improved commercially	energy consequences, effect on the environment, high		
	available technology for absorption	restoration energy		
Membrane	Mixed-matrix membranes, composite	Does not suit high temperature applications, high energy		
	hollow-fiber membranes, hybrid	required for post-combustion application, expensive to		
	membrane-cryogenic processes	fabricate novel membranes, low selectivity, trade-off between		
		purity and recovery		
Adsorption	Structured adsorbents, composite	Heat control, adsorbent wear and pressure drop, prolong		
	adsorbents, rapid swing cycles, hybrid compatibility to moisture and impurities			
	membrane- pressure swing adsorption			
Chemical	Changes to process design, composite	Reliable O ₂ conveyor materials, pressurised operation		
looping	oxides as O2 conveyors			
Direct capture	Structured adsorbents, direct air capture with	Extremely dilute CO ₂ , high energy requirement, manufacture		
	renewable energy resources	of robust materials		
Hybrid capture	Membrane distillation, pressure-	Not well researched, hybrid materials production,		
	temperature swing adsorption process,	improvement of synergy and process upgrade		
	membrane pressure swing adsorption			

Table 6. Challenges and opportunities of CO₂ utilisation technologies ³⁰⁾

Utilisation	Chemical conversion	Enhanced oil/gas	Mineralisation	Desalination
technology		recovery		
Challenges	Complexity of reaction pathways	Transportation of CO ₂	Slow kinetics	Equipment corrosion
	High operation conditions	Large number of	High pressure and	Expensive operation
	Stability of catalyst to coke	parameters involved	high temperature	Large amount of brine
	formation	Fluctuations in oil	operation	
	Development of highly selective	price	Expensive to	
	catalyst		implement	
Opportunities	Dry reforming of methane	Water alternating gas	Indirect carbonation	Providing potable water
	Catalytic reduction to formic acid	system (WAG)	Utilisation of	to residential and
	and its derivatives	Compensated	inorganic wastes	municipal customers
	Noble-metal-doped transition-	Neutron log (CNL)		Possible
	metal catalyst			implementation in
	Biological pathways to synthetic			various regions
	fuels			Modified Solvay
	Oxidative dehydrogenation			process

Al mamoori et al $^{28)}$, enumerates the primary applications of the captured CO_2 as in:

- Enhanced oil and gas recovery
- Chemical conversion
- Mineralisation
- Desalination

Also, the US Department of Energy categorised captured carbon utilisation techniques into four main areas that the CCS program supports:

- Cement industry
- Polycarbonate plastics
- Mineralisation
- Enhanced hydrocarbon recovery

Broadly, CO_2 is used as a raw material in food industries for preservation, urea production, water treatment, enhanced oil and gas recovery, in chemical production and polymer synthesis with a current global usage of 232MT-1.

Recently, less than 1% of released CO2 into the atmosphere is being utilised in the mentioned industries ⁷⁴⁾. In the enhanced oil and gas recovery (EOG/EOR), CO₂ is introduced into an oil reservoir where it combines with trapped oil in the rock formation. The CO₂ together with the oil is then pumped out of the rock where the former is separated and re-injected back to repeat the process, and the oil is collected. Melza et al. 75) claims this process to be more efficient than the conventional oil recovery technique, as the oil, on mixing with CO₂, expands and becomes lighter and easier to recover. CO2 has the potential of partially replacing fossil fuels as its conversion into fuels seems to be the best way in CO2 utilisation. CO2 can be used in the synthesis of fuels such as methanol, methane, syngas, and alkanes, and such fuels may be used in power plants, transportation and fuel cells. Furthermore, CO₂ is used as feedstock in chemical production, most notably in the production of urea (160 Mt year-1), polyurethane (18 Mt year-1), inorganic carbonates (60 Mt year-1), acrylic acid and acrylates (10 Mt year-1), polycarbonates (4 Mt year-1), and alkene carbonates (a few kt year-1) ^{28, 76)}. Urea, a major fertilizer, possesses the highest market for CO2 utilisation and is also used for fine chemicals, polymer synthesis, pharmaceuticals, and inorganic chemicals 77). In the area of mineralisation, CO2 is treated with metal oxides like magnesium and calcium oxides that naturally exist as mineral silicates. This is done by increasing the temperature and injecting fluids that have high a concentration of CO₂, thereby enhancing the carbonation kinetics ⁷⁸⁾. Captured CO₂ is also utilised in removing total dissolved solids (TDS), thereby converting brine to pure water. Though the technique is efficient, but the main barrier towards its implementation is a cost implication. Also, CO₂ if exposed to sea water mixed with ammonia forms weak bonds, resulting in the removal of ions from the water phase. These compounds are formed NA₂CO₂ and NH₄Cl, and due to their weight, they settle at the bottom of the tank ⁷⁹⁾, while the fresh water is collected for other uses.

4.2 Conclusion

Tar removal and carbon capture techniques have been reviewed and shown to be effective in purifying gas streams. Even though some of the tar removal techniques have been applied to biomass gasification process, most of the CO₂ capture and utilisation techniques are yet to be applied. This may be due to the CCU techniques are still being at the laboratory stage of development. Combined CO₂ capture and utilisation will lead to smaller, cleaner and more energy efficient technologies. This will ensure the production of clean gas while mitigating the release of CO₂ into the atmosphere, as the captured carbon will be utilised for value-added products. Despite the promising prospect of this approach, challenges like that of cost implication, thermodynamic instability of the captured CO₂, technology scale-up, technology efficiency, and implementation risk currently limits its practical use. Full commercialisation of biomass gasification may be attractive to investors if a gasification system that incorporates both tar abatement and carbon capture systems may be developed. However, such studies are scarce in biomass gasification, even though some modelling studies on the aspect of carbon capture have been reported. Future research should focus on the design of systems, with integrated tar and CO2 capture units, as part of the producer gas conditioning (cooling and cleaning) system that will be financially viable. According to the review, efficient methods recommended can be incorporated, that is insitu and post gasification methods for tar abatement and calcium looping and chemical absorption via monoethanolamine for carbon capture. Biomass gasification has a lot of prospects in the near future where green fuels will take over the fossil fuel market. Hence there is need for developing a system that produces clean fuel free of tar and CO2 gas that could be used directly. This calls for collaboration between engineers and material scientists to come up with viable techniques through the use of sustainable materials and catalyst. This will go a long way in achieving cleaner syngas and commercialising the gasification process.

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