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Technoeconomic Analysis of a Waste Tire to Liquefied Synthetic Natural Gas (SNG) energy system

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Abstract

Thermochemical conversion of solid wastes through gasification offers the dual benefit of production of high-value fuels and environmentally friendly waste disposal. In this paper, we propose a novel process for production of liquefied synthetic natural gas (SNG) from waste tires via a rotary kiln gasification process. We use a combination of experimental data available in the open literature, first principles mathematical models and empirical models to study three design cases (without CCS, with precombustion CCS and with pre- and postcombustion CCS) in two locations (USA and Norway). The thermodynamic, economic and environmental performance of the concept is studied. The results show that minimum selling prices of 16.7, 17.5 and 19.9 $\$/GJ_{LHV,SNG}$ are required for USA and 20.9, 21.8 and 24.9 $\$/GJ_{LHV,SNG}$ for Norway. We note that these prices may become competitive under certain regulatory conditions (such as recent public policy movement in British Columbia, Canada requiring public utilities to purchase natural gas made from renewables at prices up to 30 $\$/GJ_{LHV,SNG}$). The minimum selling price reduces substantially with process scale and with levying tipping fees. The design situated in Norway with both pre- and post-combustion CCS has near zero direct and indirect CO₂ emissions.

Keywords: Waste-to-Energy, Waste tire, Gasification, CO₂ capture, Synthetic Natural Gas (SNG), Rubber

1. Introduction

The growing global demand for energy is motivating research efforts both to utilize alternative feedstocks such as solid wastes (waste tires, plastics and municipal solid waste) as well as to develop efficient and environmentally sustainable conversion processes. Concurrently, increased population growth creates large quantities of wastes that require appropriate management. Thermochemical conversion processes, such as gasification, are promising options that offer the dual benefit of both recovery of high-value products from these solid wastes as well as their environmentally friendly disposal. Belgiorno et al. suggest that homogeneous high-energy density wastes such as waste tires are particularly suitable for gasification as a result of their high volatile matter content ($\sim 67\%$), low ash content ($\sim 7\%$) and high calorific value (LHV of ~ 33.96 MJ/kg, higher than coal) [1]. In the developed world, approximately 1 waste tire per person per year is produced resulting in approximately 1 billion discarded tires annually [2]. In addition, there are currently an estimated 4 billion waste tires in landfills and stockpiles worldwide highlighting the extent of the disposal problem. Previous waste tire management strategies have aimed at either material recovery or energy recovery as reviewed in [3]. The use of gasification as an alternative conversion process allows for both material and energy recovery through the production of higher value products through the syngas route.

However, research into waste tire gasification has been limited and mainly focussed on bench scale experimental studies as reviewed by Labaki and Jeguirim [4] and Oboirien and North [5]. Several gasifier types have been studied including fixed bed, fluidized bed and rotary kiln gasifiers. Fixed bed reactors may be suitable for small-scale processes [6] and their use has been investigated for producing hydrogen-rich syngas ([7], [8], [9]). Fluidized beds may be suitable for large-scale processes but are generally fairly expensive and require expensive external tar cracking and particulate removal [6]. In addition, experimental results on syngas yields and composition from fluidized bed gasifiers have been inconsistent [10], [11], [12], [13].

Rotary kiln gasifiers are a mature technology used commercially in several applications (such as cement production [14]). They consist of a downward-sloping rotating drum into which the waste and the oxidizing agent are fed. Heat is provided by either direct firing, when combustion takes place within the drum, or indirect firing when the drum is housed within a furnace. Amodeo et al. suggest that rotary kilns are particularly suited for

solid wastes because they allow a wider range of feed particle sizes, compositions and densities and are less sensitive to feed moisture content (allowing up to 50 %) [15]. In addition, rotary kilns have minimal maintenance requirements and sufficient turndown capacity, produce syngas with consistent quality and low tar content, and can handle difficult wastes such as waste tires. Robust experimental results were obtained by researchers at the Italian National Agency for New Technologies (ENEA) who conducted several studies on steam gasification in a rotary kiln at temperatures between 850 - 1000 °C. The syngas produced contained mostly H₂ (50 - 65 %v/v) and significant CH₄ (10 - 30 %v/v), CO (8 - 21 %v/v), C₂H₄ (0 - 10 %v/v) and CO₂ (2 - 8%v/v) with the overall gas yield increasing with temperature [16], [17], [18], [19]. However, all experiments were run at atmospheric pressure which may not be favorable for large-scale processes as expensive syngas compression may be required prior to downstream synthesis [20]. In addition, significant amounts of tar and particulate matter were reported in the syngas thus requiring an expensive external tar cracker and ceramic filter. Despite these drawbacks, we use rotary kiln gasification in this work because of the advantages mentioned and the availability of robust experimental data.

The different gasifier configurations result in different syngas qualities with a range of H₂:CO ratios. Depending on the ratio obtained, syngas could be used for production of fuels (such as diesel, gasoline, synthetic natural gas (SNG) and hydrogen), chemicals (such as methanol, dimethyl ether (DME), ethanol, and olefins), or electricity. The syngas could also produce multiple of these products as part of a polygeneration scheme [21]. In this work, considering that tire-derived syngas in rotary kilns has a high H₂:CO ratio of 3.77, we investigate the production of SNG. Haldor Topsøe's TREMP methanation process is used to convert syngas to SNG [22]. This process does not require an exact H₂:CO ratio of 3.0; in order to compensate for the excess H₂ and maximize conversion to SNG, a small amount of CO₂ is left in the syngas as explained in Section 2.2.2. This eliminates the need for an inefficient reverse water gas shift step. The decision to produce SNG was also influenced by the Norwegian context in which natural gas exports are expected to have an increasing share of revenue earned. Liquefaction of SNG enables the product to be sold on the fast-growing global LNG market as well as to utilize existing Norwegian LNG supply chain infrastructure. Furthermore, the highest demand growth for natural gas is expected to arise from China and South-East Asia [23]. Thus, transportation of liquefied SNG from Norway is typically the more suitable option compared to building pipelines. We

use the Poly-Refrigerated Integrated Cycle Operations (PRICO) (or Single Mixed Refrigerant (SMR)) process for liquefaction because it is the simplest and most researched commercial process [24].

Promising studies are available in the literature for production of SNG from biomass. For instance, Batidzirai et al. investigated the thermodynamic, economic, and environmental performance of the entire biomass-based SNG supply chain [25]. Van der Meijden et al. studied wood to SNG systems using different types of gasifiers [26]. Zwart et al. analyzed the potential of bioSNG production in a integrated system with the aid of a bench scale set-up [27]. Gassner and Maréchal studied a variety of thermo-conversion options for biomass to SNG including indirect and direct-heated gasification [28]. Further, the optimal design strategies for polygeneration of SNG, power and heat were investigated [29].

However, to the authors' knowledge, there have been no research efforts analyzing waste tire to liquefied SNG processes. Therefore the objective of this paper is to evaluate the thermodynamic, economic and environmental performance of the proposed waste tire to liquefied SNG concept from the systems perspective at industrially relevant scales. Three designs (without CCS, with precombustion CCS and with pre- and postcombustion CCS) are studied as illustrated in Figure 1. Two plant locations are considered: USA and Norway. Furthermore, the influence of the process scale, tipping fees and CO₂ tax rates on economic performance is determined by means of a sensitivity analysis.

2. Methodology

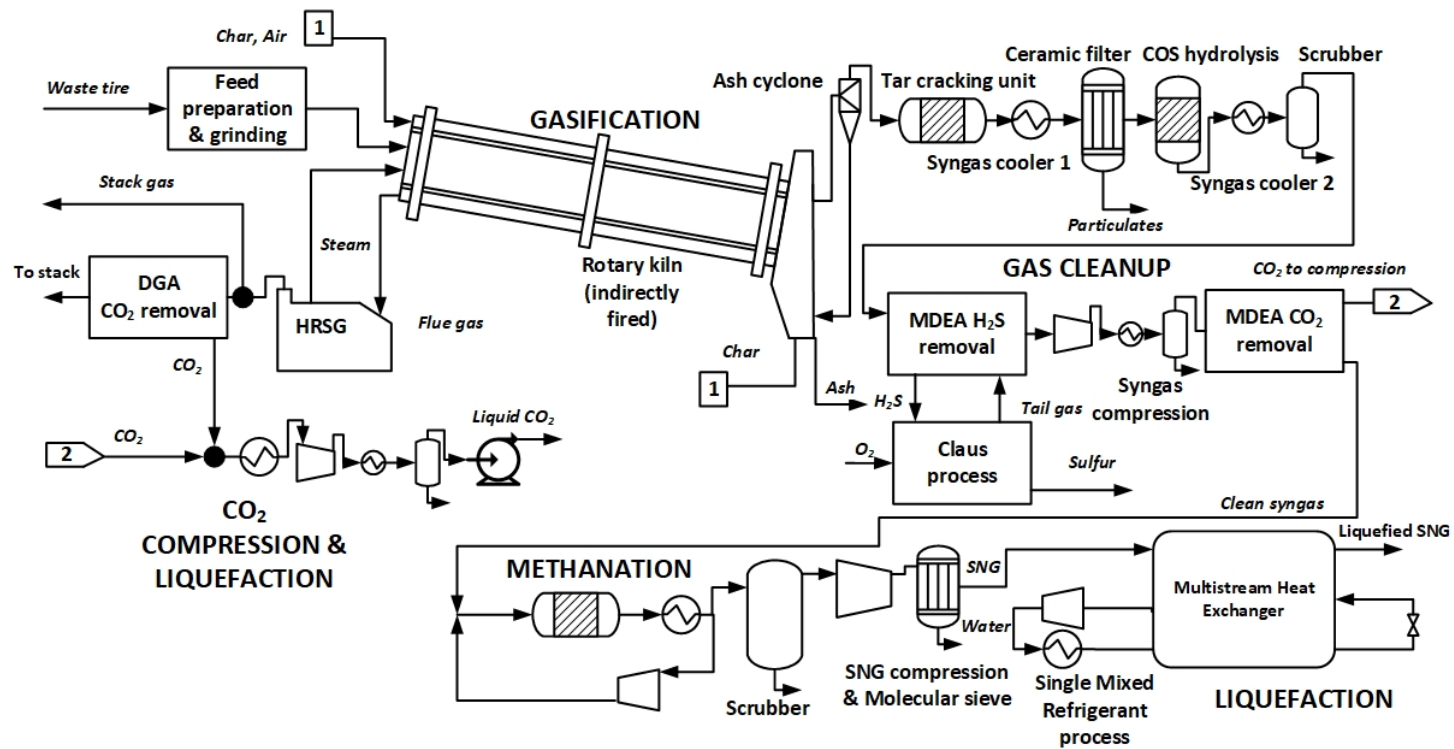


Figure 1: Detailed process flow diagram for the waste tire to liquefied synthetic natural gas (SNG) process

2.1. Simulation basis

Figure 1 illustrates the major sections of the proposed waste tire to liquefied synthetic natural gas process. Three design cases are studied. For the case without CO₂ capture, the waste tire is gasified with steam in a rotary kiln to generate syngas that undergoes cleaning using a two-stage MDEA process to remove H₂S and CO₂ before heading to the downstream synthesis section. The captured H₂S stream is converted to elemental sulfur in a Claus process, while the CO₂ is emitted. Heat for the gasification is provided by housing the kiln in a furnace in which combustion of the unreacted char occurs, with the resulting flue gas emitted. The cleaned syngas is then converted to methane in the methanation section. The methane is then compressed before heading to the liquefaction section where it exits as liquefied SNG. The second case with pre-combustion CO₂ capture features essentially the same units, except that the CO₂ captured in the MDEA process is compressed and liquefied for sequestration. Similarly, the case with both pre and post-combustion CCS features all the above units except that the flue gas stream passes through a cleaning process with Diglycol Amine (DGA) to capture CO₂, which then heads to the CO₂ compression and liquefaction section.

These three concepts are analyzed using a combination of commercial process simulation software and experimental data available from the literature. Mass and energy balances for most of the unit operations are developed using Aspen Plus v10, except for the MDEA-based H₂S and CO₂ removal sections that are modeled using BR&E's ProMax v4. For physical property calculations, the Peng-Robinson equation of state with the Boston-Mathias modification (PR-BM) was used for the gasification, Claus, CO₂ compression, and SNG liquefaction sections in order to be consistent with previous work [30], [31], [32]. The Redlich-Kwong-Soave equation of state with modified Huron-Vidal mixing rules is used for the methanation section as validated in [33]. The Amine package in ProMax was used for the H₂S and CO₂ removal sections consistent with previous work [31]. The ElecNRTL package in Aspen Plus is used for the DGA-based CO₂ capture process [34]. For the complex gasification process, experimental results published by Portofino et al. [19] are used to specify the syngas yield and composition.

The waste tire feedstock is assumed to be identical to that studied in [19] with respect to proximate and ultimate composition, as illustrated in Table 1. The liquefied SNG produced contains at least 99.5 mol% CH₄. The Aspen Plus simulation contained the following conventional components: CO, CO₂, H₂, H₂O, N₂, AR, CH₄, H₂S, O₂, NO, NO₂, SO₂, SO₃, COS, CH₃OH, C₂H₆,

C_2H_4 , C_3H_8 , $i-C_4H_{10}$, $n-C_4H_{10}$, $i-C_5H_{12}$, and $n-C_5H_{12}$, S_2 , S_3 , S_4 , S_5 , S_6 , S_7 , S_8 , DGA , DGA^+ , H_3O^+ , $DGACOO^-$, OH^- , HCO_3^- , CO_3^{2-} . Solids: C and S, and Nonconventional solids: Waste tire, Char, Tar, Soot and Ash. Table 2 presents a summary of the operating parameters and assumptions of the simulation. The three concepts are scaled such that the thermal input of the waste tire feedstock is $893 MW_{LHV}$ (equivalent to approximately 26.3 kg/s of waste tire) in order to be consistent with previous studies [20].

Ultimate Analysis	wt %	Proximate Analysis	wt %
C	77.3	Volatile Matter	67.7
H	6.2	Fixed Carbon	25.5
N	0.6	Ash	6.8
S	1.8		
O	7.3		
LHV [MJ/kg]	33.96		

Table 1: Characterization of waste tire

Unit	Parameters	Reference
Waste tire preparation	Crumb tire size = 6.0 mm	[19],[35]
Rotary kiln gasifier	Rotary kiln steam gasification. 66.7 wt% steam/33.3wt% waste tire T = 1000 °C, P = 1.013 bar	[19]
Furnace	T = 1050 °C, P = 1.013 bar	Assumed
COS hydrolysis	T = 200 °C, P = 1.013 bar	
H ₂ S removal	Solvent composition: 50.0 wt% MDEA: 50.0 wt% H ₂ O T = 45 °C, P = 1.013 bar, Removal: 96.2 %	[34]
CO ₂ removal	Solvent composition: 45.0 wt% MDEA: 5.0 wt% piperazine: 50.0 wt% H ₂ O, T = 45 °C, P = 23 bar, Removal: 88.2 %	[34] -
Claus process	Two-stage sulfur conversion, Furnace: T = 950 °C	[36]
Methanation	Four-stage conversion, Inlet T = 300 °C, Inlet P = 23 bar. ΔP = 3 bar Recycle rate = 75 %	[37]
SNG compression & purification	Outlet pressure = 55 bar, Molecular sieve removes 98.5% CO ₂ and all H ₂ O	[36]
SNG liquefaction	SNG flowrate = 9.7 kg/s, P = 55 bar, Inlet T = 22 °C, Outlet T = -157 °C SNG Mole composition: CH ₄ : 99.4 %, H ₂ : 0.5 %, CO ₂ : 0.03 % MSHE UA _{max} = 25.0 MW/K, Pressure ratio = 6.47 Refrigerant: Mole composition: N ₂ : 8.32, CH ₄ : 24.02, C ₂ H ₆ : 36.88, C ₃ H ₈ : 0.00, n-C ₄ H ₁₀ : 30.77 Low P = 2.79 bar, high P = 18.04 bar, Δ T _{min} = 0.95 K, Flowrate = 58.5 kg/s	[24]
CO ₂ compression	Multistage compressors CO ₂ purity = 99.1 %, Outlet T = 25 °C, P = 153 bar	[38]
Postcombustion CO ₂ capture	Solvent composition: 72.3 wt% DGA: 27.3 wt% H ₂ O T = 70 °C, P = 1.0 bar, CO ₂ Removal = 95.0 %	[34]
Compressors	Isentropic efficiency = 80 %, maximum pressure ratio = 5	[30]
Pumps	Isentropic Efficiency = 80 %	[30]
Heat Exchanger Network	ΔT _{min} = 5 °C, Maximum 10 branch splits per stream Utilities: Cooling water: T _{in} = 10 °C, T _{out} = 16 °C, Cost = 5.15e-9 \$ ₂₀₁₆ /kJ LP steam generation: T = 125 °C, P = 1.2 bar °C, Cost = 1.89e-6 \$ ₂₀₁₆ /kJ MP steam : T = 300 °C, P = 12 bar °C, Cost = 2.20e-6 \$ ₂₀₁₆ /kJ HP steam generation: T = 480 °C, P = 50 bar °C, Cost = 2.49e-6 \$ ₂₀₁₆ /kJ	[31], AEA AEA Assumed [31] [31]

Table 2: Operating parameters and specifications of the simulation

2.2. Case 1: Without CCS

2.2.1. Rotary kiln gasification

Waste tires received at the plant gate are shredded and ground to a maximum size of 6 mm to meet the specifications suggested by Portofino et al. [19], [35]. The resulting crumb tire together with steam is fed to the rotary kiln gasifier. The gasification phenomena include pyrolysis, devolatilization, char gasification, sulfur reaction, and a complex series of chemical reactions to give a gaseous product (consisting primarily of H₂, CO, CH₄ and CO₂), tar and soot, and unreacted char as shown in Table 3 [19], [18]. The gasifier is operated at 1000 °C and 1.013 bar because experimental results at these operating conditions produced the highest gas yield as shown in Table 3 [19].

Product	Yield [kg/kg _{feed}]	Composition
Raw syngas	2.62	Mole fractions: H ₂ O: 0.416, CH ₄ : 0.048 H ₂ : 0.309, CO ₂ : 0.136 CO: 0.082, COS: 0.003 C ₂ H ₄ : 0.002, C ₂ H ₆ : 0.002
Char	0.33	Proximate Analysis: Volatile Matter: 5.65 Fixed Carbon 71.99 Ash: 21.92

Table 3: Yield and composition of products of steam gasification at 1000 °C and 1.013 bar obtained from [19]

The experimental results for the syngas composition are incorporated in the Aspen Plus simulation by specifying the yields in an RYIELD reactor. The solid residue consisting of unreacted char and ash rolls down the length of the sloped kiln and is discharged at the reactor outlet while the raw gases are first directed to a cyclone to remove any entrained ash or unconverted char, and then to a tar cracking unit to eliminate tars from the generated syngas [20]. The syngas is then cooled to 200 °C and passed through a ceramic barrier filter for removal of particulate matter before being directed to the acid gas removal section.

Similar to the experimental setup in [19], the gasifier is indirectly fired by housing the kiln in a furnace operating at 1050 °C as illustrated in [39]. The entire fuel requirement for the furnace is provided by the unreacted char

from the rotary kiln which is mixed with preheated air for combustion. In order to minimize heat loss and protect the shell of the furnace, a refractory lining is used on the inner side of the wall between the furnace and the ambient air. The wall between the furnace and the gasifier is made out of a temperature-resistant alloy in order to maximize heat transfer to the kiln [39]. The pyrolysis (breakdown) of the solid char to elemental species is first modeled as a decomposition process occurring at 500 °C in an RYIELD block together with a calculator block that specifies the component yields according to the ultimate composition of the char. Then the combustion reaction is modeled using the RGIBBS reactor assuming unrestricted chemical equilibrium to give flue gas and ash. The air flowrate is adjusted such that the heat requirement of the gasifier is satisfied. Heat loss to the environment of 1% of the gasifier duty is assumed and included in the energy balance calculations.

2.2.2. Acid Gas Removal

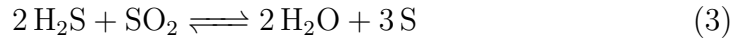
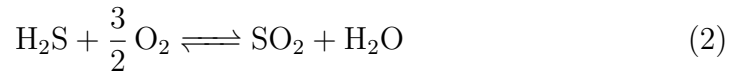
The gaseous product contains 0.03 mol% COS that needs to be removed to sub-ppm levels to prevent poisoning of downstream catalysts. The COS hydrolysis process is used to convert COS to H₂S. This process is modeled with the REQUIL reactor in Aspen Plus assuming chemical equilibrium is achieved. In order to achieve 99.5% COS conversion, an activated alumina-based catalyst is used [40]. The syngas is then cooled and flashed before heading to the acid gas removal section. Physical absorption is used for removal of H₂S and CO₂ using MDEA as a solvent. The process detailed in [34] is used. The low pressure absorber is used to remove H₂S. The sweetened gas is then compressed to 23 bar for CO₂ removal and the captured sour gas stream is sent to the Claus section. Piperazine is added to the solvent in the CO₂ removal section to enhance absorption. It is essential to control the amount of CO₂ removed in the MDEA section because the conversion of syngas in the downstream methanation process is sensitive to the relative ratios of CO, CO₂, and H₂ [41]. The contribution of the CO₂ feed fraction to the methanation process through the Sabatier reaction is accounted for by using the “feed gas module”, M, (Equation 1) which is set to 3 to maximize methane production [41]. This specification resulted in 87.8 wt % CO₂ removal from syngas.

$$M = \frac{X_{\text{H}_2, \text{feed}} - X_{\text{CO}_2, \text{feed}}}{X_{\text{CO}, \text{feed}} + X_{\text{CO}_2, \text{feed}}} \quad (1)$$

Simulation of the H₂S and CO₂ removal processes is done using BR&E's ProMax software as detailed in [34].

2.2.3. Claus process

The Claus process serves the dual purpose of treating the captured H₂S to prevent emissions to the outside air and producing elemental sulfur as a valuable by-product. The two key reactions that occur are the partial oxidation of H₂S to produce SO₂ (Equation 2), and the reaction of H₂S with the produced SO₂ to produce elemental sulfur (Equation 3).

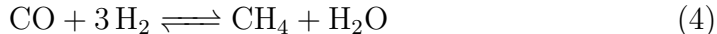


The relatively small quantity of oxygen required is assumed to be available on-site as a utility. Two stages of conversion to sulfur take place with intermediate sulfur condensation and removal in order to drive the forward reaction resulting in a sulfur recovery of 96% [36]. The unreacted H₂S after the second stage is recycled back to the MDEA H₂S removal process. The produced sulfur is cooled and stored for sale. A rigorous model is developed in Aspen Plus for the Claus process using the approach detailed in [42].

2.2.4. Methanation

The TREMP methanation process by Haldor-Topsøe was used, which consists of four catalytic fixed bed reactors each operating at a temperature of 300 °C with intermediate cooling [36],[37], [41]. Hydrogenation of CO and CO₂ occurs over a nickel-based catalyst according to the methanation and Sabatier reactions, which are presented in Equations 4 and 5 respectively. Due to the highly exothermic nature of these two reactions, it is necessary to implement a temperature control strategy, such as recycling a substantial fraction of the product from the first reactor in order to dilute the feed, in addition to using high temperature tolerant catalysts. The outlet stream from the third reactor is cooled to 150 °C and condensed water removed so as to enhance the conversion to methane in the final reactor. The SNG product stream is passed through a molecular sieve in order to remove 98.5 vol% of CO₂ and any remaining water to sub-ppm levels. Molecular sieves are designed to separate molecules based on differences in polarity and molecular

size as detailed in [43]. The SNG product is then compressed to 55 bar in order to satisfy specifications for the natural gas liquefaction process [36].



2.2.5. Liquefaction of SNG

The pressurized SNG stream is desuperheated, liquefied, and subcooled in a multistream plate-and-fin type heat exchanger [24] before exiting as liquefied SNG at -157°C . The cooling duty is provided by a refrigeration cycle. The refrigerant stream operates at two pressure levels, as shown in Figure 1: High pressure refrigerant is partially-liquefied in a condenser and then enters the multistream heat exchanger as a hot stream where it is also cooled to -157°C before undergoing adiabatic expansion through a throttle valve. The expansion slightly lowers the stream temperature after which the low pressure refrigerant re-enters the multistream heat exchanger as the only cold stream. Vaporization of the low pressure refrigerant provides the entire refrigeration duty for the system. The evaporated refrigerant then undergoes two-stage compression with intercooling to complete the cycle.

Determining the optimal operating conditions for the liquefaction process is a subject of active research [44]. In this work, we used the operating conditions suggested by Watson et al. [24] (presented in Table 2) which show good agreement with the results of Austbø and Gundersen [45]. The results from the case with maximum heat exchanger conductance (UA_{max}) set to 25 MW/K were used.

Simulation of the natural gas liquefaction process is challenging particularly as a result of the complex multistream heat exchanger block. Simulation in Aspen Plus with the MHEATX block commonly fail to converge to a physical solution especially if done in isolation [46]. This is because the MHEATX block only solves an energy balance of the multistream heat exchanger but does not have constraints to prevent temperature crossovers, thus making it necessary to either know in advance which operating conditions and heat exchanger parameter values avoid crossovers or perform a “guess-and-check” procedure. To avoid these issues, in this work the values of all pressures and compositions (except for the outlet temperature of the low pressure refrigerant) obtained from an equation-based simulation done in a previous study [46],[24] are passed on to the MHEATX block. The MHEATX block

is discretized into 10 zones with the option to add extra zones for phase change, and stream entry/exit enabled. The MHEATX block then solves for the low pressure refrigerant outlet temperature, and provides results of the zone profiles, duties, UA value and log mean temperature difference. Thus, the Aspen Plus simulation is used to validate the physical feasibility of the operating conditions suggested in [24].

2.2.6. Heat Integration and Steam Generation

In order to minimize utility costs and increase energy efficiency, a heat integration strategy is implemented on a plantwide scale. Relevant stream data (supply and target temperatures, heat capacities, flowrates) are exported from the Aspen Plus and ProMax simulations to the commercial Aspen Energy Analyzer (AEA) software. Data on the utilities available are also specified in AEA; Cold utilities considered are Cooling water, LP, MP, and HP steam generation while hot utilities considered are LP, MP, and HP steam as detailed in Table 2.

AEA determines (near) optimal designs of the heat exchanger network (HEN) using a methodology detailed by Shethna et al. [47]. Default cost correlations are used for the heat exchangers. The Grand Composite Curve utility allocation method is used as a heuristic to maximize the use of the utilities closest to the pinch (the hottest cold utilities and the coldest hot utilities i.e. the cheapest utilities) first. Default values for the unit costs of utilities in Aspen Plus are used as detailed in Table 2, with steam generation incurring a negative cost. The cooling water system consists of an evaporative mechanical draft multicell cooling tower, circulating water pumps, and a make-up water system as detailed in [40]. AEA does not guarantee a globally optimal HEN design. Thus, in this work, the best design in terms of lowest TAC out of 100 proposed near-optimal designs is chosen. A visual representation of the HEN design is presented in the Excel sheets in the Supplementary Materials.

2.3. Case 2: With precombustion CCS

The case with precombustion CO_2 capture has essentially the same units as Case 1 except that the CO_2 captured in the MDEA unit goes through a two stage compression process with intercooling and water removal to supercritical conditions. The CO_2 is then pumped to 153 bar.

2.4. Case 3: With pre and postcombustion CCS

The case with postcombustion CO₂ capture has essentially the same units as Case 2 except that an additional DGA-based unit is included to remove CO₂ from the flue gas exiting the furnace. The DGA process is similar to the MDEA process and is detailed in [34]. The captured CO₂ stream is directed to the CO₂ compression and liquefaction section prior to sequestration.

2.5. Economic Analysis

The three waste to liquefied SNG design strategies are compared using a profitability analysis for the two plant locations. The net present value (NPV) is calculated using the discounted cash flow rate of return approach. The approach follows the work of Seider et al. [48]. The base scale used for the economic analysis is 893 MW_{LHV,input} (consistent with the analysis of Larson et al. [20]) equivalent to 26.3 kg/s of waste tire. All cases also have the same SNG output of 483.8 MW_{LHV}.

The capital costs of the different pieces of equipment are estimated from data available for similar processing units in established literature sources as presented in the Supplementary Materials. For sections where cost data is not available, we make conservative estimates based on the most similar unit operations available in the literature. For instance, the capital costs of the waste tire shredding plant is estimated from [35] with the conservative assumption that the cost to produce crumb tire of size 6 mm is one third of that to produce 0.18 mm. Similarly, for the rotary kiln gasifier, we effectively double the capital cost estimate (by using a process contingency of 100%) presented in [49] for the related use of rotary kilns in cement manufacture. For the indirect firing, we include the additional cost of a furnace using an estimate made using Aspen Process Economic Analyzer (APEA). Finally, the multistream heat exchanger in the liquefaction section is costed using the method outlined by Fu and Gundersen [50] using Equation 6

$$C_{HE} = f_m r_p C_{volume} V_{MSHE} \quad (6)$$

where C_{volume} [\$₂₀₁₈/m³] is the MSHE cost per unit volume and V_{MSHE} [m³] is the volume of the MSHE. The factors f_m and r_p take on values of 2.0 and 1.1 and are used to account for the installation costs and MSHE operation at pressures higher than 25 bar respectively. The volumetric heat transfer coefficient method is used to determine C_{volume} following the methodology outlined in [51]. The MSHE is divided into zones, and the volume corresponding to each zone (V_z) is calculated using Equation 7. A local volumetric heat

transfer coefficient (β_z) of 100 kW/m³K [52] is assumed while the zone duties (\dot{Q}_z) and zone LMTDs ($\Delta T_{m,z}$) are obtained using the MHEATX simulation in Aspen Plus.

$$V_z = \frac{\dot{Q}_z / \Delta T_{m,z}}{\beta_z} \quad (7)$$

The overall MSHE volume is determined using Equation 8 with an additional 15% allowance for headers.

$$V_{MSHE} = 1.15 \sum_{i=1}^n V_z \quad (8)$$

A C_{volume} of 20,800 [\$₁₉₉₇/m³] is assumed with the costs scaled to \$₂₀₁₈ using the CEPCI index method [50].

Variable operating costs including raw material, catalyst and solvent, and waste disposal costs are estimated from closely related sections in published reports as detailed in the Supplementary Materials. For the base case, no cost or tipping fees are assumed for the waste tires and no CO₂ taxes are levied; the influence of these two factors on plant profitability is determined using a sensitivity analysis. Utility costs are estimated from AEA. The electricity requirement for preparation and shredding of waste tire (El_{tire}) [kWh/ton] is determined using the correlation with final crumb tire size (S_{tire} [inch]) suggested in [35] (Equation 9).

$$El_{tire} = 97.91 S_{tire}^{-0.222} \quad (9)$$

A capacity factor of 85 % is assumed. All costs are scaled to \$₂₀₁₈ assuming a yearly inflation rate of 2.75 %. The fixed operating costs including labor, maintenance, operating overhead, and property insurance and tax costs are estimated based on the methodology of Seider et al. [48] as detailed in Supplementary Materials. A purchasing power parity (PPP) for Norway of 10.142 NOK/USD and an average exchange rate of 8.133 NOK/USD is used. The minimum selling price (MSP) of liquefied SNG is calculated as the value that results in a NPV of zero. For the base case, the assumed parameters for the discounted cash flow rate of return analysis to calculate the NPV are presented in the Supplementary Materials.

3. Results and Discussion

3.1. Simulation results

The Aspen Plus and ProMax simulation files are made open-source and are available for download at the LAPSE digital archive at: <http://psecommunity.org/LAPSE:2019.1261> [53].

In addition, the Excel sheets containing relevant stream conditions, CAPEX and OPEX calculations as well as the DCFROR analysis are made available.

3.1.1. Thermodynamic analysis

A summary of the power consumed by the various sections as well as plantwide utility requirements are presented in Table 4. The fuel efficiency (Equation 10) only considers the proportion of the feedstock that is converted to the final product and so neglects the energetic input in utilities and electricity. The contribution of utilities and electricity is accounted for in the overall energy efficiency (Equation 11).

$$\eta_{fuel} = \frac{SNG_{LHV}}{Tire_{LHV}} \quad (10)$$

$$\eta_{energy} = \frac{SNG_{LHV}}{Tire_{LHV} + Electricity\ consumed + Utilities\ consumed} \quad (11)$$

Only a small energy penalty is incurred by incorporating precombustion CCS (Case 2) because CO₂ has to be separated anyway in Case 1. However, including postcombustion CO₂ capture results in a substantially lower energy efficiency because the parasitic load of the DGA process is supplied by additional LP steam. Table 4 shows that it is possible to recover 54.2% of the energy content of waste tires as liquefied SNG. In addition, for each kg of waste tire converted, 0.37 kg of liquefied SNG is obtained thus illustrating the potential value of the concept.

No comparable research efforts exist for the production of SNG from waste tires. Instead, we provide a comparison with studies on biomass-based SNG production. Van der Meijden et al. reported overall energy efficiencies of 54%_{LHV}, 58%_{LHV} and 67 %_{LHV} for entrained flow, circulating fluidized bed and indirect-fired gasification respectively [26]. Zwart et al. estimated an energy efficiency of 70% while Gassner and Maréchal reported overall efficiencies of 65-76 %_{LHV} [28], [29]. The lower efficiency of the proposed

concept can be partly explained by the fact that biomass typically has a very low sulfur content (less than 0.1% [54]) thus separate H₂S removal using the energetically expensive MDEA and Claus processes is not necessary. Instead, the small quantities of H₂S can be co-removed with CO₂ and oxidized to SO₂ and released to the atmosphere at levels substantially below emission limits [20]. In addition, inefficiency arises from running the gasifier at atmospheric pressure which necessitates expensive syngas compression prior to CO₂ compression. Thus, there is substantial scope for optimization with different gasifier configurations, and these will be explored in future work. For instance, using entrained flow gasification (with essentially the same gasifier design used commercially for coal gasification) at higher temperatures and pressures may result in higher syngas yields and thus efficiencies as well as allow the use of a different acid gas removal strategies like the selexol process. In addition, using entrained flow gasification with hybrid feedstocks has been shown to increase efficiency as a result of tighter heat integration of the high temperature syngas cooling section with endothermic processes such as natural gas reforming [30], [31], [55].

3.1.2. CO₂ emissions

The carbon efficiency of the three design configurations which denotes the percentage of carbon atoms in the waste tires that end up in the liquefied SNG is calculated using Equation 12.

$$\eta_{carbon} = \frac{SNG_{carbon}}{Tire_{carbon}} \quad (12)$$

The direct and indirect emissions associated with the waste tire to liquefied SNG concept for the three design cases and two locations are presented in Table 5. The direct emissions include all the GHG emissions (converted to equivalent CO₂ emissions) from processes occurring within the plant gates while the indirect emissions include all the supply chain emissions associated with all the process inputs such as electricity and utilities which vary with location. The GHG emissions computed followed the International Panel on Climate Change (IPCC) 100-year metric. However, indirect emissions involved in the manufacture of process equipment are not included; performing a full life cycle analysis of the process is left as future work. Electricity emissions are estimated based on statistics for the average electricity mix in 2017 of 687.5 kg_{CO₂,eq}/MWh [56] and 16.4 kg_{CO₂,eq}/MWh [57] for the US and Norway respectively. The indirect emission factors of utilities are determined

from data from the EPA [58]: Steam and heat, and cooling water have emissions of 0.128 kg_{CO_{2,eq}}/MJ and 0.016 kg_{CO_{2,eq}}/MJ respectively [59]. Indirect emissions for waste tires are assumed to be zero; emissions associated with tire manufacturing are attributed to the automobile sector. The emissions that may arise out of stockpiling tires are not considered as well.

Indirect emissions are highest for Case 3 with postcombustion CO₂ capture as a result of higher steam requirements for the DGA process as well as higher electricity for CO₂ compression and liquefaction. Total GHG emissions are substantially lower for the cases with CO₂ capture. In order to quantify the economic penalty of implementing pre- or postcombustion CO₂ capture, we calculate the cost of CO₂ capture (CCC) metric defined according to Equation 13. The results (presented in Table 5) show that implementing only pre-combustion CCS results is substantially cheaper than implementing both pre- and postcombustion CCS since CO₂ has to be separated prior to methanation whether CCS is implemented or not.

$$CCC = \frac{MSP_{Design\ with\ CCS} - MSP_{Design\ without\ CCS}}{GHG_{Design\ without\ CCS} - GHG_{Design\ with\ CCS}} \quad (13)$$

In order to quantify the relative environmental performance of the proposed process, we compare the total emissions with those associated with the status quo (conventional natural gas). For this study, we used 20 kg_{CO_{2,eq}}/GJ_{LHV} as the value of total emissions associated with extraction and transportation of conventional natural gas [60]. Factoring this into account, only the case with both pre- and postcombustion CO₂ capture implemented in Norway has a net reduction in life cycle global warming potential compared to conventional natural gas. For this case, we calculate the cost of CO₂ avoided (CCA) using Equation 14 to be 1,313 \$/tonne_{CO_{2,eq}}. While this is expensive, similarly high CCA values are attained for the production of biological biobutanol (472 \$/tonne_{CO_{2,eq}}) [61], thermochemical biobutanol (136 \$/tonne_{CO_{2,eq}}) [62], biodiesel (400 \$/tonne_{CO_{2,eq}}) [63], or corn ethanol (potentially up to 750 \$/tonne_{CO_{2,eq}}) [31], [63]. We note that since the majority of emissions arise as indirect emissions from the high electricity requirement (primarily required for syngas compression prior to CO₂ removal), a potential option to improve environmental performance is to use a different gasifier configuration with higher operating pressure.

$$CCA = \frac{MSP_{SNG\ plant} - MSP_{conventional\ NG}}{GHG_{conventional\ NG} - GHG_{SNG\ plant}} \quad (14)$$

Case		Case 1	Case 2	Case 3
CCS enabled?		No	Only precombustion	Both pre- & postcombustion
Waste tire flow rate	kg/s	26.3	26.3	26.3
Thermal input	MW _{LHV}	893	893	893
Total electricity requirement	MW	59.62	61.71	63.84
Grinding to crumb tire	MW	14.1	14.1	14.1
Syngas compression	MW	32.9	32.9	32.9
H ₂ S stream compression	MW	0.1	0.1	0.1
Methanation recycle compression	MW	0.3	0.3	0.3
MDEA pump	MW	0.5	0.5	0.5
SNG compression	MW	2.1	2.1	2.1
Liquefaction compressor duty	MW	9.7	9.7	9.7
CO ₂ compressor duty	MW	0.0	2.1	4.2
Net hot utility requirement	MW	157.7	159.1	283.9
LP steam	MW	-16.1	-16.0	-16.0
MP steam	MW	173.8	175.1	299.9
HP steam	MW	<0.1	<0.1	<0.1
Total cooling water requirement	MW	535.2	540.7	622.5
SNG flowrate	kg/s	9.7	9.7	9.7
Sulfur flowrate	kg/s	0.3	0.3	0.3
Heating value of SNG produced	MW _{LHV}	483.8	483.8	483.8
Heating value of SNG produced	MW _{HHV}	537.5	537.5	537.5
Overall Fuel Efficiency (η_{fuel})	% _{LHV}	54.2	54.2	54.2
Overall Fuel Efficiency (η_{fuel})	% _{HHV}	55.7	55.7	55.7
Overall Energy Efficiency (η_{energy})	% _{LHV}	43.6	43.4	39.0
Overall Energy Efficiency (η_{energy})	% _{HHV}	45.4	45.3	40.9

Table 4: Energy balances for the three designs. These energy balances are independent of location.

Case		Case 1	Case 1	Case 2	Case 2	Case 3	Case 3
Location		USA	Norway	USA	Norway	USA	Norway
CCS enabled?		No	No	Only precombustion	Only precombustion	Both pre- & post combustion	Both pre- & postcombustion
Direct GHG emitted	Mt _{CO_{2,eq}} /year	1.19	1.19	0.62	0.62	0.03	0.03
CO ₂ sequestered	Mt/year	0.00	0.00	0.57	0.57	1.16	1.16
Indirect GHG emitted	Mt _{CO_{2,eq}} /year	0.57	0.01	0.59	0.01	0.81	0.02
Total GHG emitted	Mt _{CO_{2,eq}} /year	1.76	1.20	1.20	0.63	0.84	0.05
Direct GHG emitted (scaled)	kg _{CO_{2,eq}} /GJ _{SNG}	91.69	91.69	47.57	47.57	2.38	2.38
Indirect GHG emitted (scaled)	kg _{CO_{2,eq}} /GJ _{SNG}	44.07	1.05	45.07	1.08	62.17	1.48
Total GHG emitted (scaled)	kg _{CO_{2,eq}} /GJ _{SNG}	135.75	92.74	92.64	48.65	64.55	3.86
Carbon efficiency	%	35.7	35.7	35.7	35.7	35.7	35.7
CCS	\$/t _{CO_{2,eq}}	-	-	18.6	20.4	44.9	45.0
CCA	\$/t _{CO_{2,eq}}	-	-	-	-	-	1,313

Table 5: Environmental performance of three design strategies for the two locations.

Case Location		Case 1 USA	Case 1 Norway	Case 2 USA	Case 2 Norway	Case 3 USA	Case 3 Norway
CCS enabled?		No	No	Only precombustion	Only precombustion	Both pre- & post combustion	Both pre- & postcombustion
Total capital costs	M\$	746.3	930.7	751.6	937.3	825.2	1,029.1
Solids handling	M\$	48.8	60.8	48.8	60.8	48.8	60.8
Crumb rubber plant	M\$	30.3	37.8	30.3	37.8	30.3	37.8
Water systems	M\$	103.0	128.4	103.7	129.4	114.5	142.8
Ash cyclone, tar cracker & ceramic filter	M\$	84.9	105.9	84.9	105.9	84.9	105.9
Rotary kiln gasifier & Furnace	M\$	77.1	96.2	77.1	96.2	77.1	96.2
Gas cleaning	M\$	144.9	180.7	144.9	180.7	198.7	247.7
Syngas compression	M\$	58.8	73.4	58.8	73.4	58.8	73.4
Claus process	M\$	18.2	22.7	18.2	22.7	18.2	22.7
Methanation	M\$	33.1	41.2	33.1	41.2	33.1	41.2
SNG purification & compression	M\$	20.4	25.4	20.4	25.4	20.4	25.4
CO ₂ compression	M\$	0.0	0.0	3.9	4.8	7.8	9.7
SNG liquefaction	M\$	20.5	25.6	20.5	25.6	20.5	25.6
Heat Exchanger Network	M\$	3.1	4.0	3.5	4.4	3.8	4.7
Miscellaneous	M\$	52.6	65.6	52.6	65.6	52.6	65.6
Initial catalyst and solvent fill	M\$	1.7	2.1	1.7	2.1	1.9	2.3
Land	M\$	13.9	17.4	14.1	17.5	15.4	19.2
Working capital	M\$	34.9	43.5	35.1	43.8	38.6	48.1
Total annual costs (85% capacity)	M\$/yr	125.2	156.2	134.1	167.2	161.6	201.6
Raw materials, Solvents & Catalysts	M\$/yr	1.1	1.4	1.1	1.4	1.2	1.5
Waste disposal	M\$/yr	1.9	2.4	1.9	2.4	1.9	2.4
Utilities & HEN operating costs	M\$/yr	45.0	56.2	46.3	57.7	54.7	68.2
Total labor costs	M\$/yr	13.0	16.3	13.0	16.3	13.8	17.3
Maintenance	M\$/yr	44.9	56.0	45.2	56.4	49.7	61.9
Operating overhead	M\$/yr	8.2	10.3	8.3	10.3	9.0	11.3
Property insurance & tax	M\$/yr	11.1	13.9	11.2	14.0	12.4	15.4
Minimum Selling Price	\$/GJ _{LHV}	16.7	20.9	17.5	21.8	19.9	24.9

Table 6: Summary of economic analysis of the three design strategies for the two locations

3.1.3. Economic Analysis

The results of the economic analysis for the different case studies are summarized in Table 6 with further details on the capital costs, annual operating costs and the discounted cash flow rate of return analysis available in the Supplementary Material. Cases 1, 2, and 3 have minimum selling prices for liquefied SNG of 16.7, 17.5 and 19.9 $\$/GJ_{LHV}$ in the US and 20.9, 21.8 and 24.9 $\$/GJ_{LHV}$ in Norway. All these prices are higher than the prices of conventional LNG in most locations which range from 2.90 $\$/GJ_{LHV}$ to 9.85 $\$/GJ_{LHV}$ [64]. This implies that the proposed concept is not profitable at current market conditions.

However, the minimum selling prices are comparable to those from SNG production from biomass. For instance, Batidzirai et al. report prices of 18.6 to 25.9 $\$/GJ_{LHV}$ for the 100 MW (thermal input) scale, and prices of 12.6 to 17.4 $\$/GJ_{LHV}$ at the 1000 MW (thermal input) scale although this value includes supply chain costs such as LNG transportation, regasification, compression and delivery to refuelling stations [25]. The minimum selling prices are lower than the range of 23.45 $\$/GJ_{LHV}$ (76 €/MWh) to 33.02 $\$/GJ_{LHV}$ (107 €/MWh) for the 20 MW (thermal input) scale and 18.20 $\$/GJ_{LHV}$ (59 €/MWh) to 29.93 $\$/GJ_{LHV}$ (97 €/MWh) for the 150 MW (thermal input) scale reported by Gassner and Marechal [28]. Zwart et al. report total SNG production costs of 34.22 $\$/GJ_{LHV}$ at the 10 MW (thermal input) scale and 17.00 $\$/GJ_{LHV}$ at the 100 MW (thermal input) scale [27].

We also note that the proposed process designs may become competitive under certain regulatory conditions. For instance, there is public policy movement in British Columbia (Canada) that requires public utilities to purchase natural gas made from renewables at prices up to 30 $\$/GJ_{LHV}$ [65]. In addition, the minimum selling prices are competitive with oil prices. A historical plot between 1984 and 2018 of the inflation-adjusted prices of natural gas and oil is available in our previous work [66]; oil prices ranged from 3 $\$/GJ$ in 1999 to 24 $\$/GJ$ during the energy crisis in 2008, with a recent price (June 2018) of 10.3 $\$/GJ$ [67]. The impact of plant scale, waste tire tipping fees and CO₂ taxes on minimum selling prices, and of liquefied SNG prices on net present value (NPV) are presented in the sensitivity analysis section.

3.2. Sensitivity Analysis

The results presented in the previous section are based on an assumed set of economic parameters which vary significantly over time and in different

locations worldwide. In order to determine the impact on economic performance as a result of different realizations of these uncertain parameters, a sensitivity analysis was performed. Figure 2 shows the impact of the Liquefied SNG price on NPV, with the minimum selling price resulting in an NPV of 0 \$.

Figure 3 shows the impact of the plant scale on the MSP of liquefied SNG. The results show that economies of scale have a substantial impact on lowering the MSP with large reductions in cost occurring for plant scales to 1000 MW (thermal input). However, we note that 100 MW of waste tire thermal input corresponds to approximately 10.5 million tires per year thus larger plant scales may necessitate either importing waste tires (for the Norwegian case) or situating the plant in a location with a high population density. One promising alternative that will be investigated in future work that retains the benefits of economies of scale is to utilize additional feedstocks such as waste plastics, municipal solid waste or natural gas.

Figure 4 shows that tipping fees of waste tires has a substantial impact on the MSP. Typical waste tire fees are in the range of 35 - 150 \$/tonne [35], with Sweden having a higher landfilling tax of 193 \$/tonne [68]. The results show that the proposed concept would become cost competitive with natural gas (at the upper price range of 9.85 \$/GJ_{LHV}) if tipping fees are increased to higher than 140 \$/tonne in the USA.

Figure 5 shows, in all cases, increases in CO₂ taxes increases the minimum selling price since the plant has positive direct emissions. However, the MSP for the cases with CO₂ capture increases very slightly with enabling pre-combustion CCS becoming cheaper at tax rates higher than 36 \$/tonne_{CO₂} and postcombustion CCS at tax rates higher than 54 \$/tonne_{CO₂} in both locations.

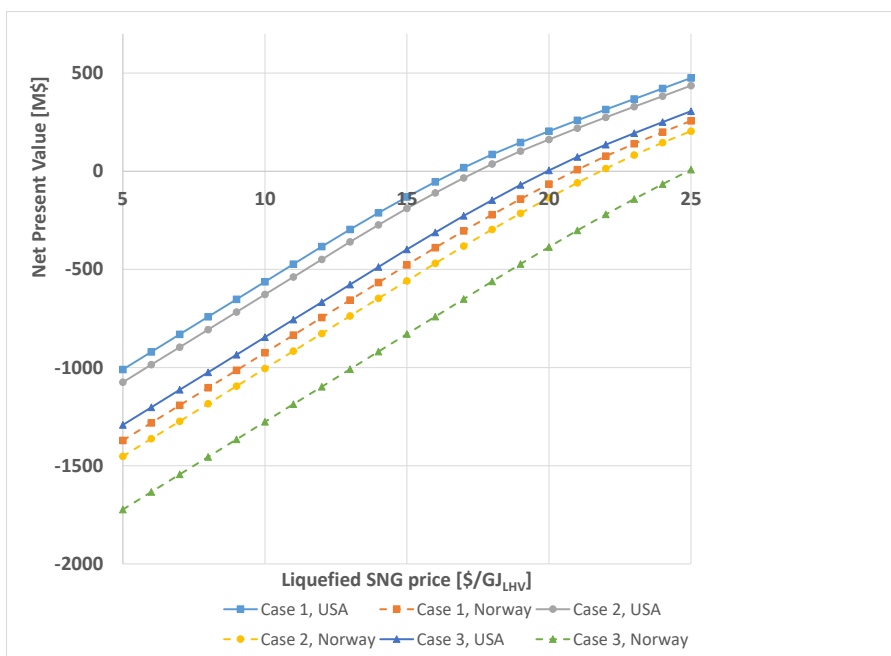


Figure 2: Impact of liquefied SNG prices on Net Present Value

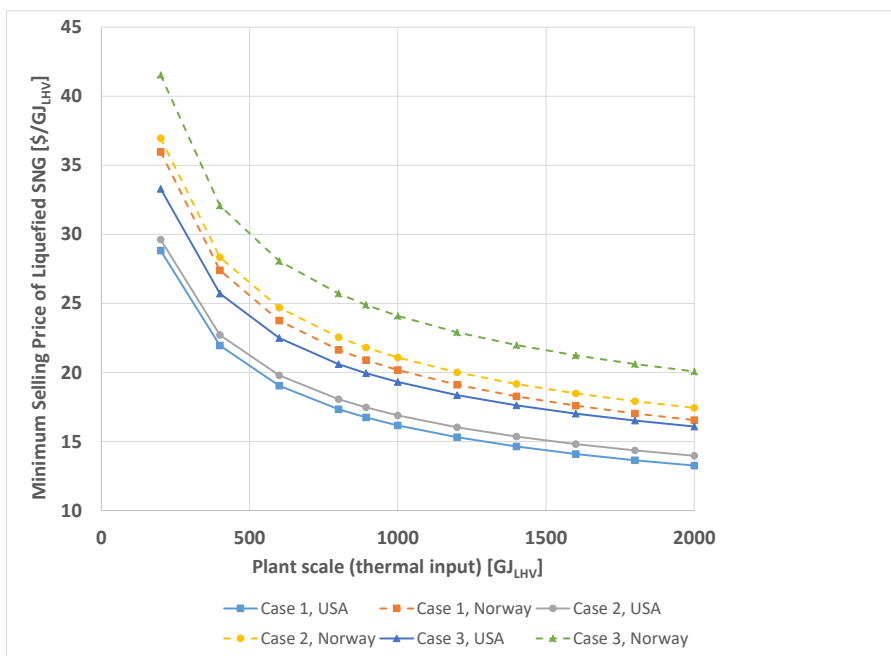


Figure 3: Impact of plant scale (thermal input) on the SNG minimum selling price

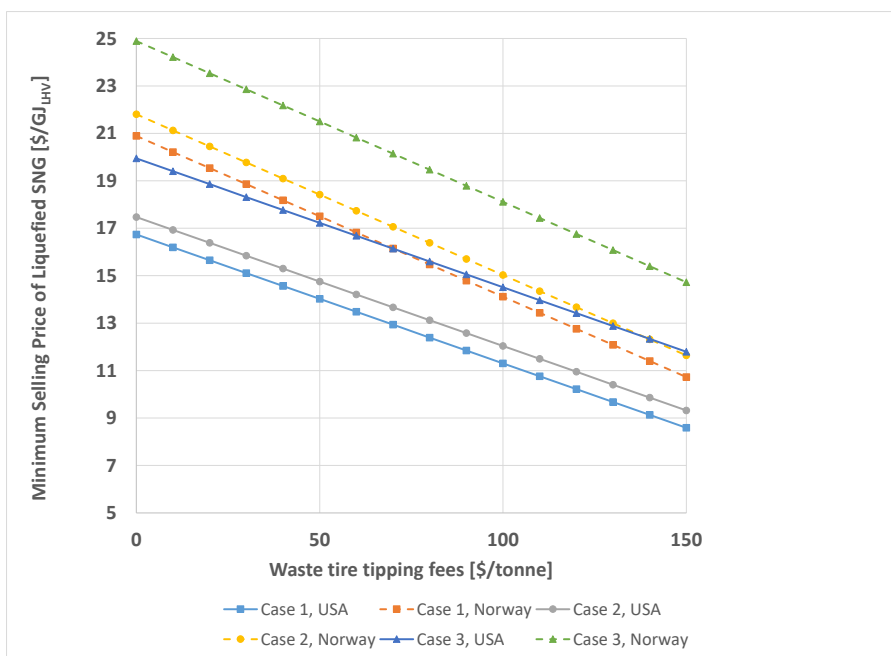


Figure 4: Impact of waste tire tipping fees on the SNG minimum selling price

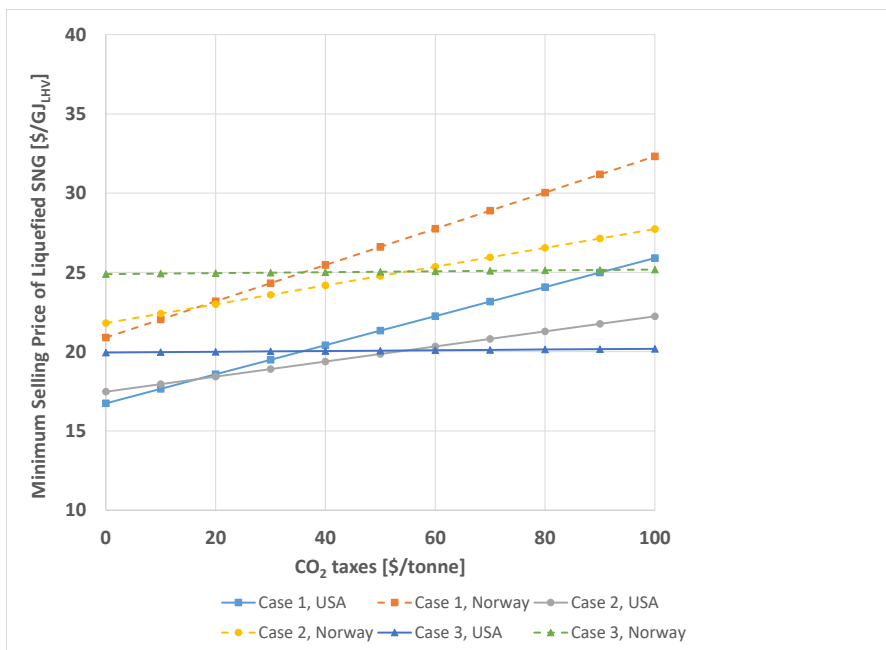


Figure 5: Impact of CO₂ taxes on the SNG minimum selling price

4. Conclusions and Future work

The proposed waste to liquefied SNG concept was found to be technologically feasible. Results of the energy analysis showed that the concept was less efficient than comparable biomass to SNG processes mostly because of the need for energetically expensive sulfur removal processes. The design situated in Norway for which both pre- and postcombustion CO₂ capture is implemented was found to be environmentally favorable to conventional natural gas extraction as it had near zero direct and indirect CO₂ emissions. Other designs may become competitive with conventional natural gas extraction under certain regulatory conditions. In addition, the minimum selling prices were found to lie within the range of oil prices in the recent decade. We also found that the minimum selling price reduces substantially with process scale and levying tipping fees.

Significant avenues for investigation and process improvement exist and these will be addressed in future work. First, different gasifier configurations such as high temperature and pressure entrained flow gasification with multiple feedstocks can be studied. In addition, different product portfolios can be investigated including production of multiple products as part of a poly-generation scheme. The design of flexible processes that are able to handle significant exogeneous uncertainties in market prices and CO₂ taxes can also be investigated.

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Conflicts of Interest

The authors declare no conflicts of interest.

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