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Article



# Techno-Economic Evaluation of Co-Removal of $NO_x$ and $SO_x$ Species from Flue Gases via Enhanced Oxidation of NO by $ClO_2$ —Case Studies of Implementation at a Pulp and Paper Mill, Waste-to-Heat Plant and a Cruise Ship

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**Abstract:** Co-absorption of NO<sub>2</sub> and SO<sub>2</sub> from flue gases, in combination with the enhanced oxidation of NO by ClO<sub>2</sub>(g), is studied for three different flue gas sources: a medium sized waste-to-heat plant; the kraft recovery boiler of a pulp and paper mill; and a cruise ship. Process modeling results are used to present the technical potential for each site together with cost estimation and optimization using a bottom-up approach. A process set-up is proposed for each site together with equipment sizing and resulting flows of process fluids. The simulation results, supported by experimental results, show that removal rates equal to or greater than current best available technologies are achievable with more than 90% of NO<sub>x</sub> and 99% of SO<sub>2</sub> removed from the flue gas. The resulting cost of removing both NO<sub>x</sub> and SO<sub>2</sub> from the flue gases is 2100 €/ton for the waste-to-heat plant, 800 €/ton for the cruise ship and 3900 €/ton for the recovery boiler. The cost estimation show that the consumption and cost of chemical additives will play a decisive role in the economic feasibility of the investigated concept, between 50% and 90% of the total cost per ton acid gas removed.

Keywords: flue gas cleaning; co-absorption; NOx; SO2; gas-phase oxidation; ClO2

# 1. Introduction

The relevance of control technologies for nitrogen and sulfur oxide (NO<sub>x</sub> and SO<sub>x</sub>) emissions remains high. Emission targets are continuously becoming increasingly strict. In the European Union, a national emission reduction commitment (NEC) directive is in force which specifies national emission reduction levels of NO<sub>x</sub> and SO<sub>x</sub> for each member state to comply with the 2020 and 2030 targets [1]. Currently 12 of the EU28 states need to decrease NO<sub>x</sub> emissions to reach the 2020 goal and every state needs to decrease NO<sub>x</sub> emissions to reach the 2020 goal and every state needs to decrease NO<sub>x</sub> emission targets, reduction across every sector is required. The more stringent requirements on sources with current emission control systems employed or implementation of NO<sub>x</sub> and SO<sub>x</sub> control on point sources without previous emission control systems will arguably require new technologies suited for these applications.

Over the last decade, multipollutant emission control systems to reduce operating and capital costs of flue gas cleaning have been given much attention as part of a possible solution to the problem [3–6]. The idea is to remove NO<sub>x</sub> and SO<sub>x</sub> concomitantly in one unit. The combined removal of NO<sub>x</sub> and SO<sub>x</sub> is generally proposed to take place in a wet scrubber preceded by an oxidation step where NO is oxidized to NO<sub>2</sub>. Several studies have shown that the co-absorption of SO<sub>x</sub> and NO<sub>x</sub> work for a variety of flue gas compositions and process conditions. Our research group studied the combined absorption of NO<sub>x</sub> and SO<sub>x</sub> from the flue gases of a waste incineration plant using gaseous chlorine dioxide (ClO<sub>2</sub>) to oxidize NO to NO<sub>2</sub>. It was shown that >90% of the SO<sub>x</sub> and >80% of the NO<sub>x</sub> can be removed using NaOH and Na<sub>2</sub>SO<sub>3</sub> as scrubber additives [7]. Meng et al. studied the



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**Copyright:** © 2021 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (https:// creativecommons.org/licenses/by/ 4.0/). removal of NO<sub>x</sub> and SO<sub>x</sub> from a simulated flue gas using ozone (O<sub>3</sub>) to oxidize NO. They used a steel slag slurry together with thiosulfate  $S_2O_3^{2-}$  to increase the absorption and reached 100% of SO<sub>x</sub> removal and >78% of NO<sub>x</sub> removal [8]. In another study Hutson et al. investigated the simultaneous removal of SO<sub>2</sub>, NO<sub>x</sub> and Hg from a simulated coal flue gas using aqueous sodium chlorite (NaClO<sub>2</sub>). They showed that complete removal of SO<sub>2</sub> and Hg can be achieved while removing about 60% of the NO<sub>2</sub> [9]. Furthermore, Si et al. performed experiments at pilot scale, 5000 Nm<sup>3</sup>/h, using a newly designed scrubber where SO<sub>2</sub> and NO<sub>x</sub> was removed using O<sub>3</sub> for oxidation of NO. They confirmed that increasing amount of SO<sub>2</sub> in the flue gas increases NO<sub>2</sub> absorption and they reached almost complete SO<sub>2</sub> absorption and >80% absorption of NO<sub>x</sub> [10].

However, despite the amount of research and technical readiness, no techno-economic evaluation of the simultaneous removal of  $SO_x$  and  $NO_x$  has been published to the best of the authors' knowledge. Therefore, this work aims to conceptualize the design and derive cost estimations for end users. In this study, reaction modeling of the co-absorption of  $NO_x$  and  $SO_x$  is used to make techno-economic performance estimates of the proposed concept to evaluate economic feasibility for three different cases: a waste-to-heat plant; the kraft recovery boiler of a pulp and paper mill; and a cruise ship.

# 2. Methods

Techno-economic evaluation of simultaneous removal of  $NO_x$  and  $SO_x$  from flue gases were performed for three applications: a medium-sized waste-to-heat plant (20 megawatt, MW); a medium-sized pulp and paper mill (500,000 tons pulp/year) recovery boiler; and a cruise ship (12 MW). The technical performance of each case was investigated through simulations based on previous experimental results [7]. Economic evaluations were then performed based on the technical study. Each application had important differences in operating conditions and in emission standards and that is why the proposed process design, technological and economic performance varied between cases.

#### 2.1. Case Studies

#### 2.1.1. Waste-to-Heat Plant

The waste-to-heat plant operates with an inhomogeneous fuel inherently creating variations in flue gas composition making NO<sub>x</sub> control more challenging. In Sweden, there is a charge on NO<sub>x</sub> of 4900 €/ton of NO<sub>x</sub> emitted together with an abatement system which returns 0.83 €/MWh (useful) to the emitter with the consequence that there is a cost for emitting more than 0.17 g NO<sub>x</sub>/kWh. If NO<sub>x</sub> can be prevented at a cost <4900 €/ton, profit can be made. The current best available technology (BAT) for NO<sub>x</sub> removal is selective catalytic reduction (SCR) with removal rates of ~90% and selective non catalytic reduction (SNCR) with removal rates of <50% [11]. For SO<sub>2</sub> removal, BAT consists of 5 different technologies, including wet flue gas desulfurization (WFGD) with removal of >99% [11]. Both NO<sub>x</sub> and SO<sub>x</sub> removal technologies are to be regarded as available and technically feasible.

#### 2.1.2. Kraft Recovery Boiler

The emissions from the kraft recovery boiler are regulated in the permit to operate where a maximum allowed limit of kg NO<sub>2</sub> emitted per produced ton of pulp produced is set. The limit is set by the local authorities and varies depending on age and location of plant. There are ongoing investigations on enforcing national levels, which will require exhaust gas treatment systems. Increasing emission standards can also be seen globally, where some pulp and paper mills in China are being required to install NO<sub>x</sub> control on kraft recovery boilers [12]. Kraft recovery boilers have low SO<sub>2</sub> emissions which is not favorable for the proposed technology, but they have processes and facilities in place and long experience of handling the proposed process chemicals. There is also the opportunity for waste treatment in the plants' existing sulfur recirculation, which makes the proposed technology more interesting from a business case perspective. The BAT for SO<sub>2</sub> control from kraft recovery boilers is WFGD with recirculation of the sulfur, achieving >90% removal of SO<sub>2</sub> [13]. Current BAT for NO<sub>x</sub> control is SNCR with >50% removal of NO<sub>x</sub>. SCR is discussed as a high removal rate alternative (>80% removal) but without current implementations in Europe [13]. The high dust content in the recovery boiler provides a difficult environment for SCR operation and requires additional dust removal units.

# 2.1.3. Cruiser

The cruiser differs from the two other cases in that the requirements will differ depending on geographic positioning. The international maritime organization (IMO) has standards that are detailed in annex VI for both  $SO_2$  and  $NO_x$  emissions from shipping detailed in Regulations 14 and 13, respectively [14,15]. In addition to the agreed global limits that are binding for ratifying states, specific emission control areas (ECAs) also exist where the emission limits are considerably lower compared to the global standards. In annex VI, regulation 14 the  $SO_2$  emissions are to either be controlled through the use of low sulfur-containing fuels or by the usage of exhaust gas treatment systems, generally scrubbers. The emission limits are set equivalent to a fuel sulfur content of 0.1% m/m in ECAs and 0.5% m/m globally. In the IMO annex VI, regulation 13 the NO<sub>x</sub> emission standards are divided into three tiers related to engine speed, where tiers 2 and 3 are currently in place and tier 1 is outdated. Tier 2 (rpm >2000) is the globally enforced limit at 7.7 g NO<sub>x</sub>/kWh and tier 3 (rpm >2000), which limits emissions of NO<sub>x</sub> to 2 g/kWh is only enforced in ECAs. From an economic perspective this makes a flexible exhaust gas treatment system attractive since the stricter emission limits can be met when required and expenses can be spared outside of ECAs. The current BAT for NO<sub>x</sub> control on cruisers is SCR with current installations in operation.

#### 2.2. Choice of Oxidizing Agent

It is vital for the absorption of NO<sub>x</sub> that NO is oxidized to NO<sub>2</sub> to increase its solubility. Different chemicals have been tested for NO oxidation, including ClO<sub>2</sub>, H<sub>2</sub>O<sub>2</sub> and O<sub>3</sub> [8,16,17]. The process integration of the oxidation step varies depending on the chemical used. For gas phase oxidation, H<sub>2</sub>O<sub>2</sub> requires high temperatures, above 300 °C, for the oxidation to take place [18]. ClO<sub>2</sub> and O<sub>3</sub> can operate at lower temperatures, 70–160 °C while maintaining efficiency [16,19]. H<sub>2</sub>O<sub>2</sub> and O<sub>3</sub> have an advantage compared to ClO<sub>2</sub> in that preferable reaction products are formed, H<sub>2</sub>O and O<sub>2</sub>, compared to the HCl formed from ClO<sub>2</sub>, but selectivity and reactivity with respect to NO to NO<sub>2</sub> oxidation is still below that of ClO<sub>2</sub> [20]. The high oxidation potential of O<sub>3</sub> has been shown to oxidize NO to higher oxidation states beyond NO<sub>2</sub> if the added O<sub>3</sub> exceeds a molar ratio with NO of 1.2 [21].

In this study  $ClO_2$  was used in the oxidation step due to the previous research on  $ClO_2$  by our group that shows high efficiency in a lower temperature range and simple process integration of the oxidation [7]. However, it should be noted that all aforementioned oxidation chemicals appear to be viable options and depending on regional legislations, site facilities and flue gas temperatures, one oxidizing agent might be favored above the others. In Table 1 the three discussed oxidizing agents are compared based on efficiency. While both NaClO<sub>2</sub>(aq) and H<sub>2</sub>O<sub>2</sub>(aq) are commonly used in experiments only gaseous oxidation is included since the consumption, translating into efficiency, of aqueous oxidizing agents is seldom reported in the literature. Often concentration is used to determine the efficiency of NO oxidation but without specifying the amount of oxidizing agent continuously added to maintain this concentration.

Method	NO Oxidation Rate	Main Active Species	Oxidizing Agent Efficiency (OA/NO)	Reference
ClO <sub>2</sub>	99%	ClO, ClO <sub>2</sub>	0.5	[7]
O3	95%	O3	1	[8]
O <sub>3</sub>	95%	O <sub>3</sub>	1.25	[22]
$H_2O_2$	95%	$H_2O_2$	2.7	[17]

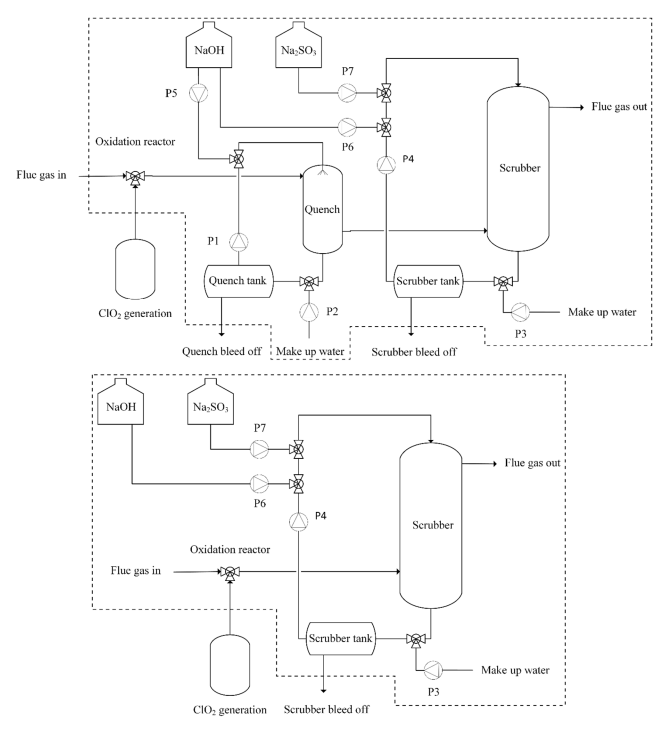
**Table 1.** List of oxidizing agents to be considered and associated technical results. OA stands for oxidizing agent and the "OA/NO" is the molar ratio between OA and NO in the flue gas.

#### 2.3. Process Designs and Contingencies

Figure 1 (upper) shows a conceptual design of a combined  $NO_x$  and  $SO_x$  control system for a waste-to-heat plant and a recovery boiler where  $ClO_2$  is used to enhance NO to NO2 oxidation. ClO2 is injected to the main flue gas path, preferably at a point where the flue gas temperature is  $150 \pm 10$  °C in order to maximize efficiency and minimize the risk of corrosion. In the conceptual design the oxidation reactor is constructed of Stainless steel 304 L. The reactor should be designed to increase mixing with minimal pressure drop. It is noteworthy that results obtained in previous studies indicate that the oxidation of NO to NO<sub>2</sub> via ClO<sub>2</sub> is rapid and not dependent on an advanced mixing reactor design [16,23]. The flue gas continues to a quench where temperature decrease, and the gas becomes saturated. The quench is maintained at an acidic pH of 1.5 and is of sufficient height to absorb the vast majority of chloride present in the flue gas. A residence time of 5 s is used in this study for design of equipment. The quench is constructed using Stainless steel 6MO/SMO to withstand the harsh environment. The scrubber is designed to achieve 90% absorption of incoming NO<sub>2</sub>. Sodium sulfite (Na<sub>2</sub>SO<sub>3</sub>) and sodium hydroxide (NaOH) is supplied directly to the liquid feed and not into the scrubber tank in order to minimize reactions with the liquid bulk. In this design, the scrubber is constructed of stainless steel SS2205 to ensure the lifetime of the equipment. The quench and scrubber tank is given dimensions such that it has a volume equal to 3 min of liquid flow [24]. Both tanks are constructed of SS2205. The chemical storage units for NaOH and Na<sub>2</sub>SO<sub>3</sub> are designed to hold a volume equal to 1 week of operation. All pumps are centrifugal and sized to fit.

In the case of the cruise ship, the conceptual design is different and seen in Figure 1 (lower).  $SO_x$  emission control on ships usually allows both for closed and open loop operation, where the wastewater is stored for treatment while in closed loop and rejected to the surrounding water in open configuration. While the release of wastewater is allowed outside of ECAs, the system will need to be able to operate in such a way to be economically feasible. It is also desirable for the system to be able to run without scrubber operation so the system must allow for high temperatures. A quench is not deemed necessary to separate chloride since the cruiser uses sea water as scrubber solution.

For a complete list of equipment and sizing for each case see the Supplementary Materials. The sizing and installation of the chlorine dioxide production unit is not included in the study.



**Figure 1.** Principal process diagram of the implementation of co-removal of  $NO_x$  and  $SO_x$  from flue gases via enhanced oxidation of NO by  $CIO_2$ . The upper scheme illustrating the design at the waste-to-heat plant and the pulp and paper mill. The lower design illustrates that of a cruise ship where no quench is present. Dashed lines indicate system boundary for economic analysis where the cost associated to wastewater treatment is not included, additionally only the production cost of  $CIO_2$  and not the production unit is included. For a complete list of sizing and materials, see Supplementary Materials.

# 2.4. Process Modeling

For each case, the sizing of the equipment and evaluation of the performance of the gas treatment process, the simulation software Aspen Plus V.11 was used. The simulations are based on the design criteria and flue gas concentrations available in Table 2. The removal efficiencies of SO<sub>2</sub> and NO<sub>2</sub> are set to >99% and >90%, respectively. The target of >90%

 $NO_2$  removal is set as to match the performance of a SCR system and the  $SO_2$  removal target is set close to 100% since this has been shown experimentally to follow with the  $NO_2$  absorption. The flue gas concentrations used in the simulations are based on typical values for each application.

**Table 2.** Comparison of the parameters applied in the three different applications. HCl concentration corresponding to the amount of ClO<sub>2</sub> needed for NO to NO<sub>2</sub> oxidation.

Parameter	Waste to Heat Plant	Kraft Recovery Boiler	Cruise Ship			
SO <sub>2</sub> removal efficiency (%)	>99	>99	>99			
$NO_2$ removal efficiency (%)	>90	>90	>90			
Raw Flue Gas Flow (Nm <sup>3</sup> /h)	57,500	357,000	61,600			
Flue gas concentrations after oxidation						
N <sub>2</sub> (%)	65	68	76			
CO <sub>2</sub> (%)	8	12	3.2			
H <sub>2</sub> O (%)	16	16	9.2			
$O_2(\%)$	11	3	11.3			
SO <sub>2</sub> (ppm wet)	150	20	900			
NO <sub>2</sub> (ppm wet)	150	150	600			
HCl (ppm wet)	60	60	220			

The process model considers the reactions in the tanks, in addition to the mass transfer and reaction kinetics in the scrubber and quench. The quench is modeled as a packed tower with a high void fraction of 0.96 to minimize pressure drop. Both the quench and the scrubber tower have diameters designed based on an 80% approach to flooding to maximize the efficiency. The chemical reactions and eventual steady state composition of the bleed stream is not considered in this work. Each case is modeled.

The chemistry in the scrubber and quench, except for one reaction, is based on a previously developed mechanism and is discussed in detail elsewhere [25]. The reaction included in this analysis, and not in our prior work, is the oxidation of S(IV) to S(VI) by O<sub>2</sub>. The oxidation of S(IV) to S(VI) in a combined NO<sub>2</sub>–SO<sub>2</sub> removal system is discussed thoroughly in literature but the rate of the reaction is still debated. In our previous experiments we have seen that the oxidation of S(IV) is rapid without any addition of other chemicals. With addition of  $Na_2S_2O_3$ , the oxidation rate of sulfite is decreased.  $Na_2S_2O_3$ is also an effective chemical for NO<sub>2</sub> absorption and we have in our experiments reached 90% NO<sub>2</sub> removal with Na<sub>2</sub>S<sub>2</sub>O<sub>3</sub> addition to the scrubber. However, there are no kinetics for a  $Na_2S_2O_3/Na_2SO_3$  system for NO<sub>2</sub> absorption why only  $Na_2SO_3$  is used in these simulations. To better match the rapid oxidation of S(IV) that has been observed, a worstcase scenario has been adopted when running the simulations. The reactions in which bisulfite (R1) and sulfite (R2) react with dissolved oxygen to form bisulfate and sulfate respectively were added to the scrubber chemistry as first order reactions in regards to S(IV) and O<sub>2</sub> with an activation energy of 29 kJ/mol and rate constant of  $6E6 \text{ s}^{-1}$ , which is the highest reported reaction rate that the authors have found in literature [26]. The reactions are defined as follows,

$$\mathrm{HSO}_{3}^{-} + 0.5\mathrm{O}_{2} \to \mathrm{HSO}_{4}^{-} \tag{R1}$$

and

$$SO_3^{2-} + 0.5O_2 \to SO_4^{2-}.$$
 (R2)

The added reaction will have a significant impact on the overall chemistry and increase the amount of S(IV) needed to reach the NO<sub>2</sub> absorption criteria. With the added reaction, results are more in line with what has been observed during experiments using Na<sub>2</sub>SO<sub>3</sub>. However, it is assumed that all, or major parts of the Na<sub>2</sub>SO<sub>3</sub> mentioned in this work will be substituted to Na<sub>2</sub>S<sub>2</sub>O<sub>3</sub> in future full-scale applications. The added amount of absorption reagent (Na<sub>2</sub>SO<sub>3</sub> and Na<sub>2</sub>S<sub>2</sub>O<sub>3</sub>) should, therefore, be expected to drop compared to the results presented here.

# 2.5. Cost Estimation

The cost of the acid gas (NO<sub>2</sub>, SO<sub>2</sub>) removal system, as shown in Figure 1 (dashed lines) is estimated using Aspen Process Economic Analyzer V11 based on the process model described using typical values [27]. The cost basis is for a nth of a kind plant and before taxation. Total direct cost (TDC) is the equipment cost together with piping, instrumentation, electrical etc., together with process contingencies. Total indirect cost includes construction and engineering together with project contingencies. The total direct cost together with indirect cost and owner's cost is equal to the total plant cost (TPC). The main assumptions in the economic analysis are presented in Table 3. In addition to TDC and TPC, total specific cost, which consists of annualized TDC plus variable operating cost, is defined to evaluate the cost of every ton of removed NO<sub>2</sub> and SO<sub>2</sub>.

Cost of equipment is largely dependent on material choice. For the specification of the reviewed process, high grade materials have been used due to the corrosive nature of the solutions. The material and component specification for each case study is available as Supplementary Materials. Fixed operating and maintenance (O&M) cost, including all labor and maintenance, are estimated as percentages of TPC with estimates based on the values in Rao et al. [28]. Variable operation cost, including chemical reagents, water and electricity price is estimated based on available literature but might be subject to large deviation depending on local availability and scale of operation.

It should be noted that all assumptions are made for land-based installation and production. The cost variation based on marine application is not taken into account. The cruiser is not bound to be produced in the EU or to operate in the EU and, therefore, costs might change on the production and installation. The harsh environment at sea can increase the maintenance cost of the unit and operating hours will vary depending on the designated route. The chemical cost should remain similar as for land based installation and while electricity generation at sea will be based on the usage of fuel, prices per kWh are similar [29]. In addition to the uncertainties of the values in Table 3, the limited space available on cruisers will pose a challenge for storage/production of chemicals to be used in the process.

**Project Region Base** EU € (2020) Project currency Conversion rate €/US. \$ 0.85 Lifetime of Unit 25 years Tax rate 0% Inflation rate 0% Annual operating hours 8000 10% [27] Process contingencies Construction time 1 year 14% of TDC [30] Indirect costs Project contingencies 15% of TDC [27] Owner's cost 7% of TDC [27] Number of full-time employees required 0.5 2.5% of TPC [28] Total Maintenance cost NaOH 100 wt% 400 €/ton <sup>1</sup> 500 €/ton <sup>2</sup> Na2SO3 100 wt% 1200 €/ton [31] ClO<sub>2</sub> 100% 68.4€/MWh Electricity 6.65 €/m<sup>3</sup> [30] Make-up water

Table 3. Main assumptions for the economic analysis.

<sup>1</sup> Estimated average. Values range between ~200–600 €/ton. <sup>2</sup> Estimated average. Values range between ~400–1000 €/ton.

# 3. Results and Discussion

This section is presented in two parts, technical performance, and economic performance. In the technical performance, process design and absorption rates are investigated. The economic performance presents capital and operating cost for each application. However, it should be noted that the design is derived from in an iterative procedure between process and economic performance.

#### 3.1. Technical Performance

The design of the proposed units of each case is based on the criteria in Table 2 and the design specification is available in Supplementary Materials. In Table 4 the resulting flows of water, liquid bleed concentration and supporting chemicals to reach 90%  $NO_x$  absorption, with scrubber dimensions specified in Supplementary Materials, are displayed.

The flows for each case are directly proportional to the size of the unit and the ratio and amount of SO<sub>x</sub> and NO<sub>x</sub>. The scrubber liquid flow is adjusted to maintain a constant liquid to gas ratio (L/G) at 10. The NaOH flow corresponds to the amount of acid gases absorbed and the Na<sub>2</sub>SO<sub>3</sub> flow corresponds to the SO<sub>x</sub>–NO<sub>x</sub> ratio in the flue gas. The relation between the variables is evident with the highest NaOH flow and lowest Na<sub>2</sub>SO<sub>3</sub> flow for the cruiser, which has the highest SO<sub>x</sub>–NO<sub>x</sub> ratio and highest total amount of acid gases removed. The amount of ClO<sub>2</sub> is fixed at 0.6 kg ClO<sub>2</sub> per kg NO<sub>2</sub> in the flue gas.

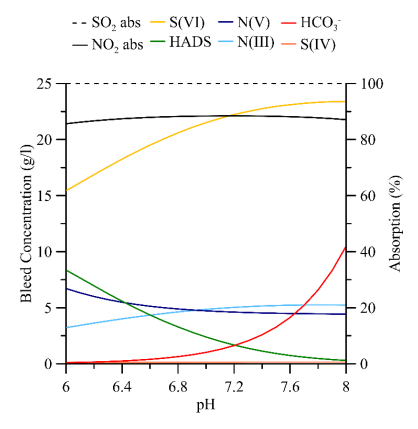
The scrubber bleed concentration profile for the waste-to-heat plant and the pulp and paper mill is similar due to the similarities in flue gas concentrations. The two major components present are S(VI) and N(III) followed by hydroxylamine disulfonic acid (HADS) and carbonate. Only a minor amount of S(IV) is present and almost no N(V). The S(IV)/NO<sub>2</sub> molar ratio, defined as the total amount of moles sulfur added to the scrubber in relation to the moles of NO<sub>2</sub> in gas phase entering the scrubber, is higher for the waste to heat plant compared to the recovery boiler. However, the resulting liquid concentration of S(IV) is lower for the waste to heat plant. This is due to the higher O<sub>2</sub> concentration in the flue gas for the waste to heat plant and the increased importance of the oxidation of S(IV) to S(VI) via O<sub>2</sub> compared to the pulp and paper mill. For the cruiser in closed mode, the share of nitrogen components to sulfur components is slightly higher. With open mode, all concentrations are low compared to the closed mode, except for the concentration of S(IV) which is almost as high as for the waste-to-heat plant.

Waste to Heat **Kraft Recovery** Cruise Ship, Parameter Plant Boiler Closed/Open Liquid Flow  $(m^3/h)$ 575 3570 616 L/G (kg liquid/kg gas) 10 10 10 NaOH (l/h) 58 75 350/280  $Na_2SO_3 (l/h)$ 60 460 10/150S(IV)/NO<sub>2</sub> ratio<sup>1</sup> 1.3 1.2 1.7/2.2 $ClO_2$  (kg/h) 8.5 52.7 33.7/33.7 SO<sub>2</sub> removed (kg/h) 20 20 130/130 50/50 $NO_2$  removed (kg/h) 13 80 Make up water  $(m^3/h)$ 1.344.166.26/616 1.38 Liquid bleed  $(m^3/h)$ 6.1 6.27/616 N(III) (g/L)8.8 10.5 7.9/0.1 N(V) (mg/L)9.4 1.98/0.67 36 S(IV) (mg/L) 60 140 1110/90S(VI) (g/L)25 25 25/0.11 HADS (g/L) 2.7 10 3.4/0.02.2 Carbonates (g/L) 1.8 0.67/0.0

**Table 4.** Resulting scrubber flows and species concentration in scrubber bleed to reach design criteria for each application.

 $^{1}$  S(IV)/NO<sub>2</sub> ratio is defined as the total amount of moles of sulfur added to the scrubber via Na<sub>2</sub>SO<sub>3</sub> and SO<sub>2</sub> divided by the total amount of moles NO<sub>2</sub> in the gas phase entering the scrubber.

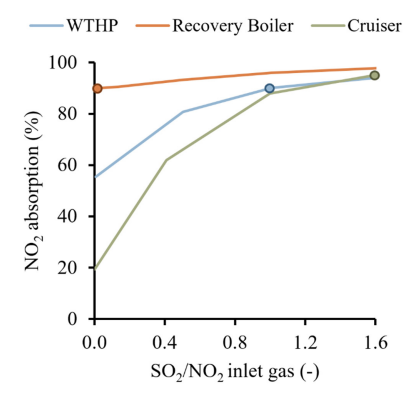
Figure 2 shows how the impact of scrubber pH on the liquid composition in the scrubber bleed. These simulations are for the waste-to-heat plant with operating conditions according to Table 2. The model predicts a maximal NO<sub>2</sub> absorption at a pH of 7.2. No differences are seen in SO<sub>2</sub> absorption in the pH-range from 6 to 8. The oxidation of absorbed SO<sub>2</sub> to S(VI) greatly increase the absorption capacity of SO<sub>2</sub> which is why the removal is always close to 100%. The increase in pH above 6 result in an exponentially increased amount of absorbed CO<sub>2</sub>. Absorbed CO<sub>2</sub> will influence process economics negatively since the amount of NaOH added changes in relation to this. The model predicts an exponential decrease of HADS when increasing pH from 6 to 8. HADS is formed from HSO<sub>3</sub> – and increased pH will shift the S(IV) equilibrium towards SO<sub>3</sub><sup>2–</sup>. Sulfite is never present in the bottom liquid at concentrations above 150 mg/L and the same is true for N(V) which requires a more acidic solution to be present. The increase in N(III) and S(VI) with increasing pH corresponds to the decreased concentration of HADS.



**Figure 2.** Liquid composition variations based on pH at scrubber bleed and resulting  $NO_2$  and  $SO_2$  absorption. Simulations are based on the waste-to-heat plant case.

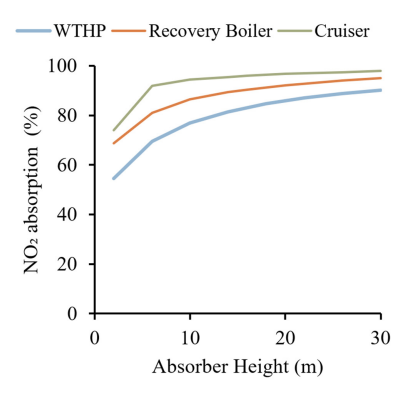
Figure 3 shows how the NO<sub>2</sub> absorption will increase with increasing total amount of S(IV) added to the system either from Na<sub>2</sub>SO<sub>3</sub> or SO<sub>2</sub> in the gas phase for the three cases. The molar ratio of added Na<sub>2</sub>SO<sub>3</sub> to NO<sub>2</sub> is constant at 0.3 for the waste-to-heat plant (WTHP in the figure), 1.1 for the recovery boiler and 0.04 for the cruiser. NaOH addition is adjusted to maintain the pH at 7.2 for each simulation. The results indicate that the required addition of  $S(IV)/NO_2$  (SO<sub>2</sub> and Na<sub>2</sub>SO<sub>3</sub>) is between 1.1 and 1.3 to reach 90% absorption of NO<sub>2</sub>. The operating point for each case is indicated with a circle on the respective line. The waste-to-heat plant is the most sensitive to changes up or down around the assumed SO<sub>2</sub> concentration (operating position) with respect to NO<sub>2</sub> absorption and will need to adjust Na<sub>2</sub>SO<sub>3</sub> addition to maintain constant absorption of NO<sub>2</sub>. The operating conditions and design for the recovery boiler and the cruiser is relatively insensitive to changes in SO<sub>2</sub> concentration. The recovery boiler relies almost solely on Na<sub>2</sub>SO<sub>3</sub> for NO<sub>2</sub> absorption and the amount is set at a level which reaches 90% NO<sub>2</sub> absorption. For the cruiser, the

sulfur content in the cruiser fuel could be decreased with 25% and still be enough for 90%  $NO_2$  absorption. Below a  $SO_2$ - $NO_2$  ratio of 1.1, the  $NO_2$  absorption drops rapidly for the cruiser.



**Figure 3.**  $NO_2$  absorption in relation to  $SO_2/NO_2$  ratio in the flue gas. The  $NO_2$  absorption is displayed for the three different cases as described in Table 2. The  $Na_2SO_3$  addition is kept constant for each case at 0.3 for the waste-to-heat plant (WTHP in figure), 1.1 for the recovery boiler and 0.04 for the cruiser. Each circle shows the operating position for each respective case. NaOH dosing is adjusted to maintain constant pH at 7.2.

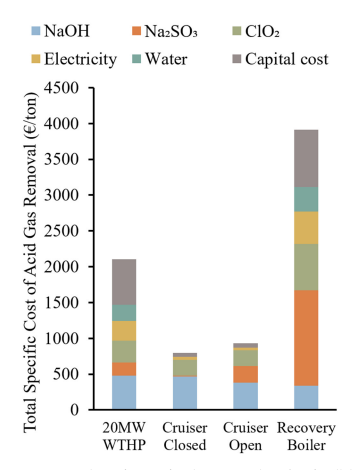
Figure 4 shows how the NO<sub>x</sub> absorption will increase with increasing scrubber height for the three different case designs. The corresponding gas residence time in the scrubber changes linearly between 2 and 30 m of height from 1 to 17 s for the waste-to-heat plant and the recovery boiler and from 1 to 12 s for the cruiser. Gas composition, flow and added Na<sub>2</sub>SO<sub>3</sub> remains constant while NaOH is changed to maintain pH constant at ~7.2. The increased absorption per meter added scrubber height is decreasing with each meter added, due to decreasing NO<sub>2</sub> partial pressure and S(IV) concentration. From the figure, the height of the scrubber for each final design case can be seen where the NO<sub>2</sub> absorption reaches 90% (except for the cruiser in closed mode which has 95% absorption). The difference between each case is due to the difference in S(IV)/NO<sub>2</sub> ratio in the scrubber where a higher S(IV)/NO<sub>2</sub> ratio will increase the reaction rate of NO<sub>2</sub> absorption. The cruiser has the highest S(IV)/NO<sub>2</sub> ratio and, therefore, requires the lowest height of the scrubber. The waste-to-heat plant design has the lowest S(IV)/NO<sub>2</sub> ratio and requires a much taller scrubber. Each S(IV)/NO<sub>2</sub> ratio is visible in Table 4.



**Figure 4.** NO<sub>x</sub> absorption in relation to scrubber height with all parameters retained according to Table 4. except for NaOH dosing to maintain pH constant at 7.2. Waste-to-heat plant abbreviated to "WTHP" in figure legend.

#### 3.2. Economic Performance

Figure 5 shows the total specific cost of acid gas removal (combined  $SO_2$  and  $NO_2$ ) in  $\mathcal{E}/(\text{ton SO}_x \text{ and NO}_x)$  removed for all the investigated cases. The results are based on simulations that reach the design criteria of 90%  $NO_x$  removal and >99%  $SO_2$  removal. It is apparent that chemical cost is a major factor when estimating the cost of the reviewed system. Approximately 50% of the total specific cost is related to the chemicals. For the waste-to-heat plant, the estimations give a cost of about  $2100 \notin$ /ton acid gases. The cruiser has a cost of about 800 €/ton acid gases in closed mode with a slightly higher cost of about 900  $\in$ /ton acid gases for the open mode. In open mode, the NO<sub>2</sub> absorption efficiency per added kilogram of absorption agent is decreased compared to closed mode due to small amounts of remaining S(IV) being rejected. The recovery boiler case is the most expensive due to the low concentrations of SO<sub>2</sub> in the flue gas at about 3900  $\notin$ /ton acid gases. Higher concentrations of  $SO_2$  in the flue gas decrease this result in two ways. Firstly, the removal of  $SO_2$  is cheaper, almost only increasing NaOH consumption and secondly it lowers the amount of Na<sub>2</sub>SO<sub>3</sub> needed for NO<sub>2</sub> absorption. There is an opportunity to lower chemical cost for the recovery boiler through the usage of white liquor as an absorption reagent. The availability on site could prove cost efficient. However, it will not be possible to use with the design as indicated in Figure 1 since the high pH of white liquor (>12) will result in a substantial absorption of CO<sub>2</sub>, as indicated by Figure 2. In a recent study, Hruška et al. investigated the usage of white liquor and NaOH to absorb H<sub>2</sub>S from black liquor gasification [32]. They concluded that the amount of  $CO_2$  absorbed by white liquor when used as an absorbent will increase the lime kiln load to a degree that will have significant negative impact on process economy.



**Figure 5.** Total specific cost of acid gas removal in €/ton for all the investigated cases as described in Table 2 divided by expense category. Capital cost is the annualized TPC.

Figure 6 shows the total specific cost of NO<sub>x</sub> removal in in  $\epsilon$ /ton removed. The result differs from that in Figure 5 by indirectly showing the amount of SO<sub>2</sub> that is removed by the process. The recovery boiler has the lowest SO<sub>2</sub> content in the gas while the cruise ship has the highest, shifting the total specific cost of NO<sub>2</sub> removal to increase to the highest level for the cruise ship and the lowest for the recovery boiler. The waste-to-heat plant has an estimated cost of ~5400  $\epsilon$ /ton, the cruise ship has ~2700 and 3300  $\epsilon$ /ton in closed and open modes, respectively, and the recovery boiler has an estimated cost at ~4700  $\epsilon$ /ton.

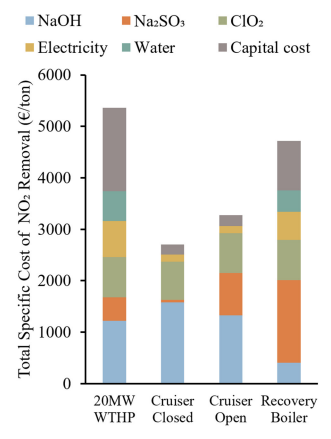
When relating the result to the aforementioned NO<sub>x</sub> charge in Sweden of 4900  $\notin$ /ton, the result indicates economic viability for installation of co-removal of NO<sub>x</sub> and SO<sub>x</sub> at a Swedish waste-to-heat plant. The total specific cost of NO<sub>x</sub> removal of 5300  $\notin$ /ton would net a 400  $\notin$ /ton loss when only NO<sub>2</sub> removal is considered. When also accounting for the SO<sub>2</sub> removal the relative cost looks competitive. In the waste-to-heat plant case, for each ton of NO<sub>2</sub> removed, 1.54 ton of SO<sub>2</sub> is also removed resulting in a cost of about 260  $\notin$ /ton SO<sub>2</sub>. In addition to this, a special case for the waste-to-heat plant is that the fuel is a source of income and lower-grade waste gives a higher income. This creates an incentive for being able to take in high sulfur containing wastes which will reduce the need for added Na<sub>2</sub>SO<sub>3</sub>.

The estimated specific cost of NO<sub>2</sub> removal for a recovery boiler unit is about 4700  $\notin$ /ton which nets a 200  $\notin$ /ton NO<sub>2</sub> profit, not even accounting for the SO<sub>2</sub> removal if the recovery boiler were to be included in the Swedish charge on NOx emission system. This application is the most sensitive to absorption reagent cost and effectiveness. Almost entirely relying on added sulfite as a source of active sulfur for NO<sub>2</sub> absorption the total specific cost of NO<sub>x</sub> removal would decrease with around 20% if the added sulfur was reduced by 50%, something that might be possible with the addition of Na<sub>2</sub>S<sub>2</sub>O<sub>3</sub>.

The target of 90% removal of  $NO_2$  at the recovery boiler might be higher than what is required at certain sites. If the required  $NO_2$  removal is lower than 90%, the cost of

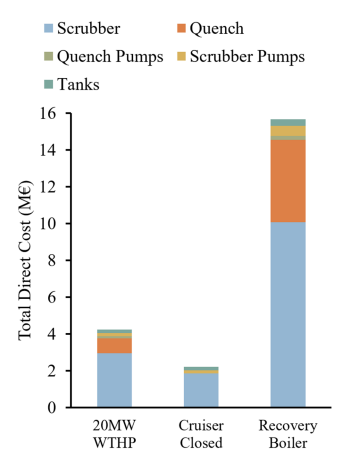
 $Na_2SO_3$  will drop significantly. However, if the required  $NO_2$  removal is lower than 50%, selective non-catalytic reduction will almost certainly be a cheaper alternative, even with the challenging implementation.

The cruiser has the lowest estimated total specific cost of  $NO_2$  removal due to the high amount of  $SO_2$  present in the flue gas and the relatively cold scrubber liquid (sea water). The cost might also be further decreased in retrofitting cases since it is assumed that the scrubber to be used is the same design as currently installed  $SO_2$  scrubbers. However, the economic feasibility for the ship application needs to be compared to other maritime alternatives. It is technically possible to reach the emission standards put in place by the IMO, but other techniques might be more affordable. The liquid treatment and storage when operating in closed mode should be regarded as an uncertainty in terms of cost.



**Figure 6.** Total specific cost of NO<sub>2</sub> removal in €/ton for all the investigated cases as described in Table 2 divided by expense category. Capital cost is the annualized TPC.

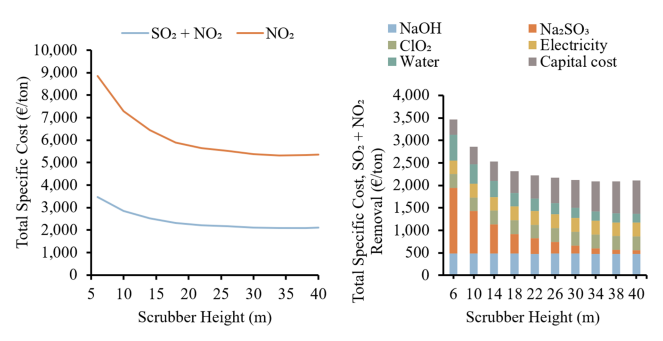
Figure 7 shows the estimated total direct cost for the three different cases. The main cost is related to the scrubber with over 60% of the total cost in all applications. The quench, when present, is estimated at about 25% of the total cost and tanks and pumps equates to roughly 15% of the total cost. The investigated process optimized for the currently employed design and cost assumptions have a large share of the total cost related to capital cost. If the consumption of sulfite can be decreased compared to what is simulated and/or if the cost of sulfite is lower than what is assumed in this work then a lower scrubber height than what has been used in this work will be a more cost-efficient option, significantly lowering the capital cost of the process. A summary of cost results, including TDC, TPC, variable and fixed cost can be found in Supplementary Materials.



**Figure 7.** Total direct cost in  $M \in$  divided between equipment category for all the investigated cases as described in Table 2. The recovery boiler has substantially higher total direct cost compared to the other two cases due to the much larger size of the plant, about six times larger.

Figure 8 visualize the impact on removal cost by increasing the scrubber height from 6 to 40 m while maintaining 90% NO<sub>2</sub> and 100% SO<sub>2</sub> removal for the waste-to-heat plant design. At a scrubber height lower than 6 m it is impossible to maintain NO<sub>2</sub> absorption above 90% while at the same time having a discharge concentration of sulfate at 25 g/L since a major part of the sulfur will leave the scrubber as  $SO_3^{2-}$ . Therefore, scrubber heights below 6 m are excluded from the figure. For all other cases, concentrations and efficiencies are constant. The cost for NaOH and ClO<sub>2</sub> remains almost unchanged with scrubber height since the total amount of acid gases absorbed remains constant. At the same time, water and Na<sub>2</sub>SO<sub>3</sub> cost decrease with increased scrubber height since less Na<sub>2</sub>SO<sub>3</sub> is needed to be added and capital cost increase with added height. The total specific cost of NO<sub>2</sub> and combined SO<sub>2</sub>/NO<sub>2</sub> removal is lowest, 5300 and 2100 €/ton respectively, at a scrubber height of 30–34 with the design conditions according to Table 2, "waste-to-heat plant". NaOH, includes the NaOH consumption for both the scrubber and the quench.

The absence of other techno-economic studies on the combined removal of  $NO_x$  and  $SO_x$  makes comparisons difficult. However, a comparison can still be made to other economic evaluations performed for only  $NO_x$  or  $SO_x$  removal. In 2017, Yaramenka et al. published a report in which they estimated the cost benefit of implementing SCR to cruise ships in the Baltic sea [33]. They estimated that a SCR system for  $NO_x$  control with 90% reduction would operate at a total specific cost ranging from 2000 to 5500  $\notin$ /ton  $NO_x$  depending on the lifetime of the equipment, from 5 to 15 years, and interest rates. Figure 6 shows the corresponding values found in this study to 2700 and 3300  $\notin$ /ton  $NO_x$  for closed and open scrubber configurations, respectively. The cost estimates for  $NO_x$  control on a cruiser in this study is close to those estimated for a SCR system with the same performance and operating conditions. However, the cost in this study also includes the removal of SO<sub>2</sub>.



**Figure 8.** Impact of scrubber height on total specific cost of NO<sub>2</sub> and acid gas removal with constant NO<sub>2</sub> removal of 90% for the waste-to-heat plant case. To the left, the absolute total specific cost of NO<sub>2</sub> and combined SO<sub>2</sub>/NO<sub>2</sub> removal in  $\notin$ /ton and to the right the total specific cost of combined SO<sub>2</sub>/NO<sub>2</sub> removal divided by category in  $\notin$ /ton.

In 2015, Eelco den Boer and Maarten Hoen published an assessment report where they brought together different available investment cost figures for the installation of a SO<sub>x</sub> scrubber on a medium-sized cruiser [34]. The capital expenditures of a scrubber for a 12 MW engine ranged from 1.2 to 2 M€ for an open loop scrubber and between 2.4 and 4.4 M€ for a closed loop scrubber. In Figure 7, we calculate the capital cost of a scrubber to be 2.2 M€. In this study, no cost is included for a waste treatment and storage solution which is part of the related increased cost for a closed loop scrubber in the compared values. The calculated cost in this study is close to the available data on installed marine scrubbers. Since the scrubber used for modelling in this study is based on an actual marine scrubber design, the numbers should be close to each other.

The operation and maintenance cost of a SO<sub>x</sub> scrubber for marine application reported by Eelco den Boer and Maarten Hoen ranged from 0.3 to  $2.5 \notin$ /MWh [34]. In this study the corresponding values can be derived from Figure 5 to be  $11.3 \notin$ /MWh. When removing the NO<sub>2</sub> related chemicals of ClO<sub>2</sub> and Na<sub>2</sub>SO<sub>3</sub> and excluding electricity price as this is not included in the values by Boer and Hoen, the cost drops to  $7.15 \notin$ /MWh. The resulting operating and maintenance cost in this study is high compared to the values found by Eelco den Boer and Maarten Hoen. This is not unreasonable and can be explained by the cost related to co-absorption of NO<sub>2</sub>. pH is kept higher and additional NaOH is needed to co-absorb the NO<sub>2</sub>. It is also possible that the cost of NaOH used in this study of 400  $\notin$ /ton is high compared to what shipping companies pay. In the report by Boer and Hoen, a price of 200  $\notin$ /ton NaOH was assumed.

The treatment of a liquid containing both nitrogen and sulfur compounds at the volumes present in the investigated technology is not studied nor currently employed at any plant to the best of the authors' knowledge. For the time being, the maritime sector is allowed to release the wastewater while outside of ECAs why no incentives are present to introduce any cost at the moment. For the two other applications there are a couple of studies with suggestions on how to best treat the liquid where the ultimate solution would be to turn the waste into products, for instance sulfuric acid and sodium nitrate. For the case of a more conservative outlook on the eventual cost coupled to waste treatment, a more ready solution would be to inject the scrubber effluent back into the boiler. Both waste-to-heat plants and recovery boilers are subject to high particle and sulfur loads as is and should therefore be relatively insensitive to the added sulfur load. Preliminary tests

performed have shown that  $SO_4^{2-}$  will remain as  $SO_4^{2-}$  when injected to the boiler in a region where the temperature is about 1000 °C and  $NO_2^-$  will be reduced to  $N_2$ . In a best-case scenario, this will only incur a minor cost from increased soot formation and depending on whether there is a flue gas condenser present or not is also an efficiency loss.

### 4. Conclusions

We evaluate the technical and economic performance of a co-absorption system for  $NO_x$  and  $SO_2$  emissions control. Three cases: a small sized waste-to-heat plant, a cruise ship and a kraft recovery boiler of a medium-sized pulp and paper mill, are identified as potential targets for the technology where either process conditions or existing facilities make for an interesting business case. Based on previous experiments, a simple process setup is established for each investigated case [7]. Each case process setup is simulated using a detailed reaction mechanism. The economic performance was investigated with investment cost estimated using the Aspen Process Economic Analyzer V11 and operational cost estimated using available data on chemical cost and simulation results. The key findings are summarized below:

- 1. The removal efficiencies of the co-absorption system are able to reach the level of what is currently viewed as best available technologies, with >90% and >99% removal for  $NO_x$  and  $SO_2$ , respectively.
- 2. The waste-to-heat plant had an estimated total specific cost of NO<sub>2</sub> removal of 5300 €/ton NO<sub>2</sub>, the recovery boiler 4700 €/ton NO<sub>2</sub> which is relatively close to the Swedish charge on NO<sub>2</sub> at 4900 €/ton NO<sub>2</sub>. The cruiser had an estimated total specific cost of NO<sub>2</sub> removal at 3300 €/ton NO<sub>2</sub>.
- 3. The consumption of absorption chemicals (Na<sub>2</sub>SO<sub>3</sub>) should be significantly reduced when either an inhibitor or another absorption chemical is used, for instance Na<sub>2</sub>S<sub>2</sub>O<sub>3</sub>.
- 4. The investigated technology is well suited for emission control of sources with a high share of SO<sub>2</sub> in the flue gas and/or sites with current availability of oxidizing chemicals and absorption chemicals.
- 5. The treatment of wastewater was identified as an unknown parameter in the cost evaluation.

**Supplementary Materials:** The following are available online at https://www.mdpi.com/article/10.3390/en14248512/s1, Table S1: Equipment list for the waste-to-heat plant, Table S2: Equipment list for the kraft recovery boiler, Table S3: Equipment list for the cruiser, Table S4: General operating conditions and design parameters for simulations and cost estimations, Table S5: Summary of Total direct cost for each investigated case together with all other resulting plant costs.

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