

## Article

# Simulation and Techno-Economic Analysis of a Power-to-Hydrogen Process for Oxyfuel Glass Melting

Sebastian Gärtner <sup>1,2,\*</sup> , Daniel Rank <sup>1</sup> , Michael Heberl <sup>1</sup> , Matthias Gaderer <sup>2</sup> , Belal Dawoud <sup>3</sup> , Anton Haumer <sup>4</sup> and Michael Sterner <sup>1</sup> 

- <sup>1</sup> Research Center on Energy Transmission and Storage (FENES), Faculty of Electrical and Information Technology, University of Applied Sciences (OTH) Regensburg, Seybothstrasse 2, D-93053 Regensburg, Germany; daniel.rank@oth-regensburg.de (D.R.); michael.heberl@oth-regensburg.de (M.H.); michael.sterner@oth-regensburg.de (M.S.)
- <sup>2</sup> Chair of Regenerative Energy Systems (RES), Campus Straubing for Biotechnology and Sustainability, Technical University Munich, Schulgasse 16, D-94315 Straubing, Germany; gaderer@tum.de
- <sup>3</sup> Laboratory of Sorption Processes (LSP), Faculty of Mechanical Engineering, Technical University of Applied Sciences (OTH) Regensburg, Galgenbergstraße 30, D-93053 Regensburg, Germany; belal.dawoud@oth-regensburg.de
- <sup>4</sup> Faculty of Electric and Information Technology, Technical University of Applied Sciences (OTH) Regensburg, Seybothstrasse 2, D-93053 Regensburg, Germany; anton.haumer@oth-regensburg.de
- \* Correspondence: s.gaertner@tum.de; Tel.: +49-941-943-9808

**Abstract:** As an energy-intensive industry sector, the glass industry is strongly affected by the increasingly stringent climate protection targets. As established combustion-based production systems ensure high process stability and glass quality, an immediate switch to low greenhouse gas emission processes is difficult. To approach these challenges, this work investigates a step-by-step integration of a Power-to-Hydrogen concept into established oxyfuel glass melting processes using a simulation approach. This is complemented by a case study for economic analysis on a selected German glass industry site by simulating the power production of a nearby renewable energy park and subsequent optimization of the power-to-hydrogen plant performance and capacities. The results of this study indicate, that the proposed system can reduce specific carbon dioxide emissions by up to 60%, while increasing specific energy demand by a maximum of 25%. Investigations of the impact of altered combustion and furnace properties like adiabatic flame temperature (+25 °C), temperature efficiency ( $\Delta\zeta = -0.003$ ) and heat capacity flow ratio ( $\Delta z_{HL} = -0.009$ ) indicate that pure hydrogen-oxygen combustion has less impact on melting properties than assumed so far. Within the case study, high CO<sub>2</sub> abatement costs of 295 €/t CO<sub>2</sub>-eq. were determined. This is mainly due to the insufficient performance of renewable energy sources. The correlations between process scaling and economic parameters presented in this study show promising potential for further economic optimization of the proposed energy system in the future.

**Keywords:** Power-to-Gas; hydrogen; electrolysis; oxyfuel; glass industry; decarbonization; carbon dioxide emissions; renewable energies



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## 1. Introduction

Climate change is an urgent global challenge that affects human living conditions through weather extremes, droughts, and floods. To prevent the increasing risks of climate change, a rapid and significant reduction in anthropogenic carbon dioxide (CO<sub>2</sub>) emissions is crucial. These CO<sub>2</sub> emissions are mainly responsible for climate change, and are primarily caused by leading industrial nations. Therefore, steps towards a significant reduction of CO<sub>2</sub> emissions in energy generation, the building sector, and transport have been continuously discussed and established in highly industrialized nations. However, energy-intensive industries such as steel, paper, and chemicals production are also of increasing interest in these considerations. Although often neglected at present, the glass industry is

one of these energy-intensive industries. This work focuses particularly on the German glass industry, which accounts for 20% of European glass production. In Germany, about 56,000 people are employed within 390 companies, generating a total revenue of 9.8 billion€ per year. Since most of the products are intended for European and global export, the German glass industry is of considerable economic and social importance [1].

The German glass industry is facing increasing financial and social pressure to reduce CO<sub>2</sub> emissions and energy consumption. The main reason for its high energy consumption is the melting process, accounting for 50–80% of the total energy input (in 2015: approx. 18.53 TWh) [2–4]. In addition, more than 75% of the glass industry's total energy consumption is covered by so far low-cost natural gas, which in turn has a negative impact on CO<sub>2</sub> emissions. Further energy sources electricity (approx. 17%) and mineral oil (approx. 7%). Over the years, extensive know-how has been accumulated on fossil fuel-based melting processes, which in turn guarantees consistently high glass quality. If the German glass industry is to remain economically competitive, access to a cost-effective energy supply for glass manufacturers must be ensured in the future, also with a view to future climate protection regulations. A major challenge in this context is the transition of the energy supply to an environmentally friendly, low-CO<sub>2</sub> emission system.

### *1.1. CO<sub>2</sub> Emissions of the German Glass Industry*

The CO<sub>2</sub> emissions caused by the German glass industry have stagnated at a relatively constant level of ≈4 million t CO<sub>2</sub>-eq. per year since 2005 [5]. The industry sector is listed as exposed to carbon leakage and has therefore received free allocated emission allowances. As a result, the established European Union emission trading system (EU-ETS) has not been able to implement any financial incentives to reduce CO<sub>2</sub> emissions. However, since the beginning of the third trading period of the EU-ETS (2013) and the associated reduction in the amount of free allocated emission allowances, the actual CO<sub>2</sub> emissions of the German glass industry have exceeded the amount of these free allowances [5]. An expected further reduction in the amount of free allocated emission allowances in the fourth trading period of the EU-ETS, and the associated price dynamics in the European Energy Exchange (EEX) trading system, will exert significant financial pressure on the glass industry to reduce CO<sub>2</sub> emissions in the future. This pressure is further intensified by the introduction of taxation for CO<sub>2</sub> emissions, currently being at about 30–40 €/t CO<sub>2</sub>-eq. with a subsequent fast linear increase up to 180 €/t CO<sub>2</sub>-eq. by 2045 [6].

### *1.2. Structure and Scope of This Work*

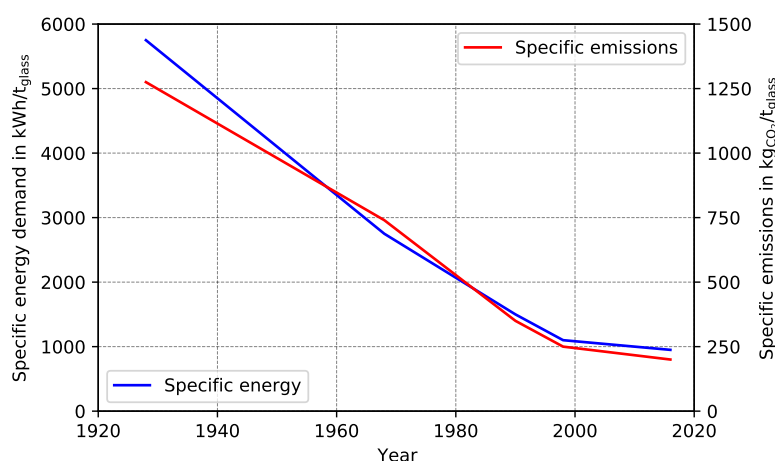
In recent years, various research approaches have been pursued to develop energy concepts for reducing CO<sub>2</sub> emissions while simultaneously integrating fluctuating renewable energy sources into the glass industry. So far no comprehensive overview of the current state of the art in glass production is available in the literature. Therefore, a brief overview of the current state of the art of glass production is provided and current approaches for reducing CO<sub>2</sub> emissions are discussed. Since these approaches have drawbacks for large-scale application, a new, innovative approach to integrating a power-to-hydrogen (PtH<sub>2</sub>) concept into glass melting processes is described. This concept enables a step-wise conversion of combustion processes to hydrogen-based melting processes, which could overcome the limitations of current approaches in terms of CO<sub>2</sub> reduction and integration of renewable energy sources in the glass industry. The main focus of this study is the numerical evaluation of this concept, which is depicted and evaluated by means of simulations based on mass and energy balances. The predictions of these models enable a subsequent energy efficiency analysis as well as the evaluation of positive effects on CO<sub>2</sub> emission reduction of the integration of PtH<sub>2</sub> into oxyfuel-based glass melting processes. In a final step, the presented models are incorporated within a case study for the integration of PtH<sub>2</sub> into the German container glass industry, thus enabling an initial techno-economic analysis.

## 2. Literature Review

CO<sub>2</sub> emissions during glass melting are depending on the specific energy demand as well as the efficiency of the furnace. Therefore, these two parameters will be briefly addressed, before various established and innovative melting tank designs are described. Some of these more advanced processes also focus on reducing CO<sub>2</sub> emissions.

### 2.1. Specific Energy Demand

The specific energy demand is highly dependent on the glass type being produced, its batch and glass composition, the melting technology, and the furnace capacity. Improvements in energy efficiency and material utilization enabled the reduction of the specific energy demand from 4000 kWh/t glass in the 1930s to under 1000 kWh/t glass in 2016 (see Figure 1). Associated with this, the specific CO<sub>2</sub> emissions per ton of molten glass have been reduced from approximately 1300 kg CO<sub>2</sub>-eq./t glass in the 1930s to an average value of 250 kg CO<sub>2</sub>-eq./t glass in 2016.

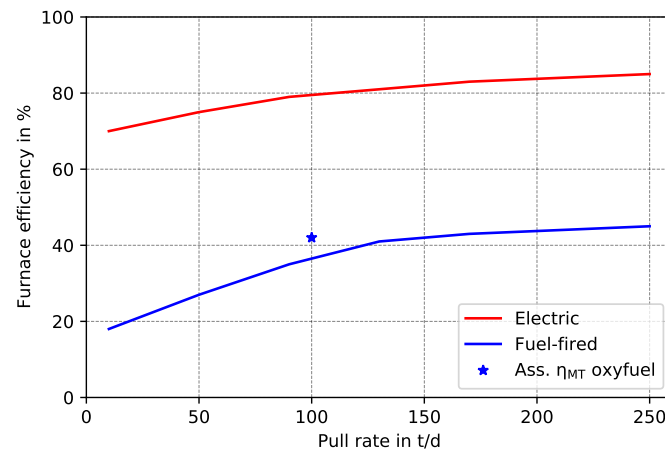


**Figure 1.** Historical development of specific energy demand and specific CO<sub>2</sub> emissions based on data from the glass industry in the EU across the overall glass sectors (e.g., glass fibers, container glass, special glass). Adapted with the permission of Ref. [7]. Copyright 2017 M.Lindig.

### 2.2. Furnace Efficiency

The furnace efficiency of a melting tank is defined as the ratio of the input energy (e.g., fuel, electrical power) to the energy transferred into the molten glass (see Equation (7)). Further, furnace efficiency of a melting system is a function of its pull rate (Figure 2). The pull rate indicates the sizing factor of a melting system and determines how much glass can be produced within a day of operation. An increasing pull rate yields an improved area-to-volume ratio and thus, reduced heat losses.

So far, reducing energy costs has been the main motivation for the glass industry to enhance both material and energy utilization efficiencies. In terms of enhanced material utilization, the addition of cullets to the glass batch has been the most important step in recent decades. The addition of only 10% of cullets to the batch composition already reduces the average energy consumption of the melting process by 2–3% [8,9]. The addition of cullets further reduces the required amount of carbonates in the batch. As the decomposition reactions of carbonates cause CO<sub>2</sub> emissions, the addition of cullets also causes a reduction of CO<sub>2</sub> emissions. However, cullets from production residues are used in most cases. The extensive use of waste glass cullets is limited, due to its impurities, various glass colors and expensive logistic costs [9].



**Figure 2.** Efficiencies of fuel-fired and electric furnaces as a function of pull rate, adapted from Ref. [10]. In this work, a melting tank efficiency  $\eta_{MT}$  of 42% is assumed for a 100 t/d oxyfuel melting system (see Section 4.4.1).

### 2.3. Melting Tank Design

The design of a melting tank is decisive in terms of energy consumption and related CO<sub>2</sub> emissions. The most common tank designs are (i) regenerative, (ii) recuperative, (iii) oxyfuel and (iv) all electric melting. A comparison of key values for this established designs is shown in Table 1.

Regenerative furnaces can be considered as the current standard technology for glass melting and are established in all industry sectors. This concept was invented by Friedrich and Hans Siemens in 1862 [11]. Because of their long use, regenerative furnaces have recently achieved only modest improvements in energy efficiency [12].

**Table 1.** Comparison of key values for established melting tank designs. As efficiency depends on several factors (e.g., glass type, melting capacity, fuel) only a qualitatively assessment can be provided. An increasing number of “+” indicates higher efficiency [12–18].

	Regenerative	Recuperative	Oxyfuel	All-Electric
Heat-exchanger system	direct contact ceramic lattice structure	indirect contact dual shell, shell-and-tube	none	none
Burner arrangement	U-flame, cross fired	U-flame, cross fired	cross-fired	none
Industry sector	Flat-, container-, fiber-, special-glass	Container- and special-glass	special glass	special glass
Efficiency	++	+	+++	++++

Within recuperative melting tanks, combustion air is preheated by indirect contact with the exhaust gas through a dual shell or shell-and-tube heat exchangers. Thus, the efficiency of heat transfer between exhaust gas and ambient is lower, while a cleaner furnace atmosphere is possible [13]. Besides regenerative or recuperative preheating of combustion air, the preheating of cullets and/or batch raw materials is established. Therefore, indirect heat exchanger systems are primarily used, as direct contact between batch and exhaust gas can cause contamination of raw materials [14].

An alternative approach is oxyfuel melting. Within oxyfuel-fired melting, natural gas is combusted in an atmosphere of nearly pure oxygen (O<sub>2</sub>), rather than ambient air. This results in increased furnace efficiency, due to much higher adiabatic flame temperatures (2775 °C vs. 2440 °C for common air-gas) and increased heat capacity flow [15]. Thus, the efficiency of oxyfuel furnaces is increased by about 5–10% compared to a similar regenerative

furnace, and about 25–40% compared to a similar recuperative furnace, respectively [13]. Moreover, the associated reduction of nitrogen ( $N_2$ ) fractions in the furnace atmosphere enable a significant reduction of the nitrogen oxide ( $NO_x$ ) emissions of oxyfuel melting furnaces making them more attractive concerning stricter environmental regulations [16]. The application of oxyfuel furnaces is carefully considered in each case (high costs for  $O_2$ -infrastructure, high-quality ceramics for brick-work, etc.) and is often omitted due to insufficient returns on investment. Currently, oxyfuel melting is widely used in the special glass industry, where glass compositions are more energy-intensive [17]. The full potential of oxyfuel furnaces has not been fully exploited yet, as they have largely been constructed as simplified regenerative furnaces, without further special adaptation of the furnace geometry [18].

A further improvement in terms of efficiency of the oxyfuel melting process can be achieved by thermochemical reformation (TCR) of natural gas (e.g., Optimelt process by Praxair). This enhances both the recovery efficiency by means of natural gas reformation to carbon monoxide (CO) and Hydrogen ( $H_2$ ) and the inferior calorific value of the fuel gas mixture from approx. 10 kWh/m<sup>3</sup> to 12.6 kWh/m<sup>3</sup> [19].

As a combination of regenerative and oxyfuel concepts, the so-called HeatOx process has recently been applied on an industrial scale. It involves preheating of  $O_2$  and natural gas for oxyfuel combustion. This enables an additional increase in the efficiency of the furnace, and a further reduction in pollutant emissions [20].

All-electric melting has several advantages over combustion-based melting, such as higher overall efficiency (see Figure 2), higher specific melting capacity (t/m<sup>2</sup>), and lower emissions of dust and toxic gases [21]. Despite these promising properties, all-electric melting is currently only used in small-scale applications of glass production (up to 50 t/d). Whether an extension and upscaling of this technology without extensive changes in the production processes is possible is still the subject of controversial debate [22]. Nevertheless, there have been successful pilot plants of up to 240 t/d recently. However, these plants require very complex technical solutions such as rotating melting areas, an extensive heat control strategy and careful positioning of electrodes [7]. In addition, currently established all-electric melting systems do not allow flexible integration of electricity from renewable energies. A strictly constant input of heat is crucial for a stable batch-to-melt conversion without excessive bubbling and therefore a constant pull rate of glass [13].

The advantages of electric melting are also used in combustion-based melting processes. Therefore, electrodes are locally installed within the molten glass, to achieve a specific improvement of glass quality and melting capacity [7]. This concept of so-called electrical boosting thus improves the overall efficiency of a melting tank [12]. However, electrical boosting usually covers only up to 10% of the total energy demand [23].

#### 2.4. Innovative Melting Concepts and Options for Low $CO_2$ -Emission Glass Production

There are numerous suggestions for innovative melting processes that may be considered more disruptive and have not significantly progressed beyond an experimental scale yet. Some of these are:

1. complete  $H_2$ - $O_2$  combustion [24],
2. stirred melters (RAMAR melter) [25],
3. rotary kilns [26] and the related P10 melter concept [25],
4. plasma arc melting [25,27] or
5. the submerged combustion melter (SCM) [28].

While some of these concepts would result in significant reductions of  $CO_2$ -emissions (1.), not all of them have been developed with a particular focus on this objective. However, research and development processes are focusing on low  $CO_2$ -emission glass production. One of them is the extension of electrical boosting to so-called horizontal hybrid melting [29]. This involves operating the furnace 100% electrically when electric energy prices are lower, e.g., due to an oversupply of renewable energy sources, whereas when electric energy prices are high, fossil fuel-fired melting is used. Nevertheless, horizontal hybrid

melting requires a high level of control engineering of a melting tank. Therefore, major changes in glass temperature and combustion area temperatures, but also in flow patterns and fining performance, are expected for hybrid melting tank [7]. In addition, research projects investigate flexible all-electric melting. However, they are focusing on a small scale and only a short-term reduction of energy input, for example at nighttime [30]. A large-scale all-electric melting system, powered only by renewables such as photovoltaics and wind power would thus require large and expensive electricity storage capacities.

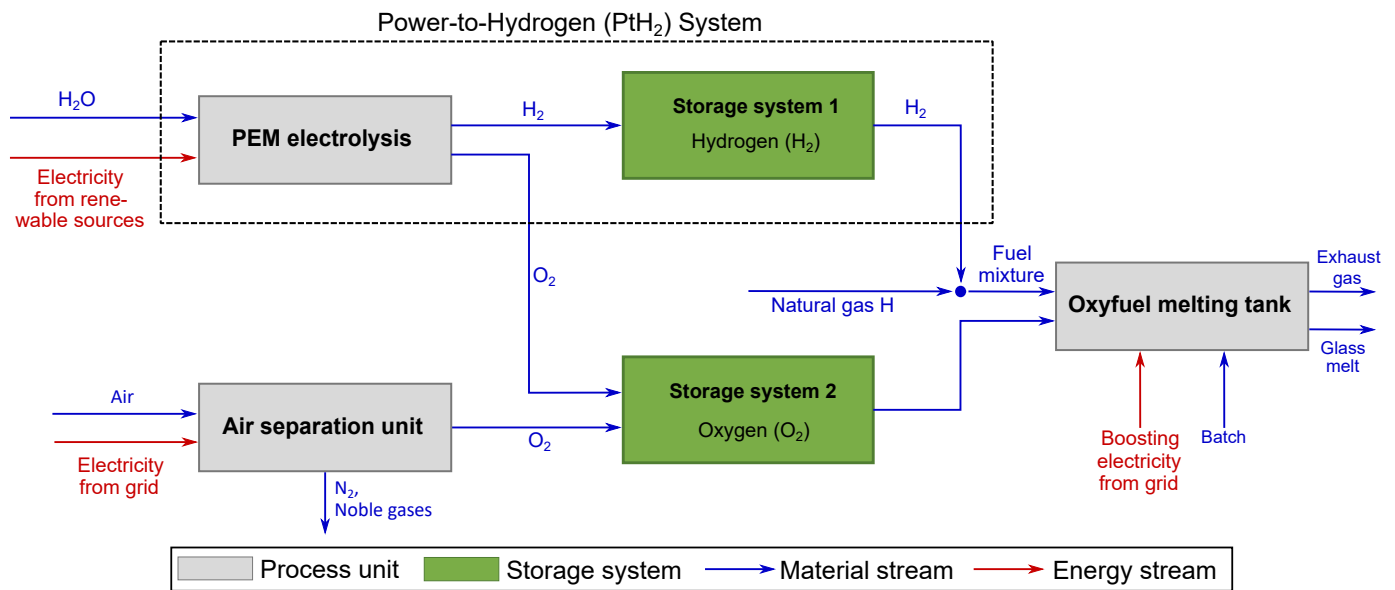
It is evident that the glass industry has implemented numerous measures to increase furnace efficiency and reduce CO<sub>2</sub> emissions. However, all of these measures show weaknesses in terms of their full application, especially on a large scale. It can be concluded that the current state of the art in glass production is not sufficient to meet future challenges in terms of CO<sub>2</sub> emission reduction and integration of fluctuating renewable energy sources. Thus, there is an urgent need for low carbon and flexible energy concepts, which can be implemented fast and with minimal impact on established processes. This work presents an alternative option to achieve this by integrating a Power-to-Hydrogen (PtH<sub>2</sub>) system into oxyfuel glass melting processes. This step-wise integration concept is completely new and has not been considered so far.

### 3. Integration of Power-to-Hydrogen in Oxyfuel Glass Melting Processes

PtH<sub>2</sub> is originally intended as a long-term energy storage system for fluctuating renewable energy sources, using H<sub>2</sub> as an energy carrier. For this purpose, water is converted into H<sub>2</sub> and O<sub>2</sub> in an electrolysis process. Subsequently, H<sub>2</sub> is temporarily stored in underground caverns, pressure storage tanks, or, up to a certain amount, in the already established natural gas infrastructure [31]. Additional approaches include the use of H<sub>2</sub> as a basic material for energy applications, such as methane (CH<sub>4</sub>) and higher chained hydrocarbon synthesis, or as a raw material for the chemical industry [32]. Meanwhile, this concept has been implemented and tested in many demonstration plants and is considered robust and applicable on a large scale [33].

PtH<sub>2</sub> is of particular interest to the glass industry, as it can provide O<sub>2</sub> for oxyfuel combustion, and H<sub>2</sub> as a substitute for hydrocarbon-based fuels. Figure 3 shows a basic flow sheet of the integration of a PtH<sub>2</sub> system into an oxyfuel glass melting process. The actual PtH<sub>2</sub> system consists of a proton exchange membrane (PEM) electrolysis and an H<sub>2</sub> storage system. PEM electrolysis is commonly considered, due to its highly adjustable partial load behavior, which allows an adaption to fluctuations in renewable energy sources like wind power (WP) and photovoltaic (PV) [34]. After generation, H<sub>2</sub> is intermediately stored in an overground pressure storage system to ensure a constant H<sub>2</sub> volume fraction in the fuel mixture for constant combustion properties. This is crucial, as unexpected and sudden fluctuations in the fuel composition have previously been identified to be critical on the constant heat input into the molten glass and consequently on glass quality [35].

The O<sub>2</sub> produced from the electrolysis is supplied into the O<sub>2</sub> storage system and is used for oxyfuel combustion. In most cases, glass manufacturers that already use oxyfuel furnaces have O<sub>2</sub> storage systems in place. These storage systems can be adopted from existing glass manufacturer infrastructure, which commonly use O<sub>2</sub> storage systems to provide backup for the ASU or to ensure permanent availability of delivered liquid O<sub>2</sub>. By using O<sub>2</sub> from electrolysis, the required ASU power can be reduced.



**Figure 3.** Basic flow sheet for the integration of Power-to-Hydrogen (PtH<sub>2</sub>) into oxyfuel glass melting processes.

Current concepts for pure H<sub>2</sub>-O<sub>2</sub> combustion have not proved to be suitable for quick implementation in existing oxyfuel melting systems, as major changes in furnace atmosphere composition are expected. The influence of these changes on the glass chemistry has not yet been adequately investigated [24]. To maintain a high level of process stability, it is, therefore, appropriate to add step-wise fractions of H<sub>2</sub> to the fuel mixture. By this method, established burner technology can still be used, while a reduction in CO<sub>2</sub> emissions can be achieved. In this work, five scenarios of H<sub>2</sub> volume fractions in the fuel mixture are investigated. In the first scenario, 10 vol% of H<sub>2</sub> are examined, which is equivalent to the mid-term H<sub>2</sub> volume fraction in the German natural gas grid [36]. In further scenarios 25, 50, and 75 vol% of H<sub>2</sub> in the fuel mixture are considered, respectively. Finally, a scenario with pure H<sub>2</sub>-O<sub>2</sub> combustion is investigated provide an outlook for future concepts. Table 2 gives an overview of the fuel mixture compositions.

In addition, a conventional scenario is defined to provide a basis for the evaluation of the changes caused by the PtH<sub>2</sub> process introduced. The conventional scenario is defined as a steady-state operating oxyfuel glass melting furnace with a pull rate of 100 t/d and additional electrical boosting, covers a constant share of 10 % of the required heat demand. A soda-lime glass, which is typical for the container glass industry, is melted. As a fuel mixture, the composition given in the “Conventional” column of Table 2 is used as natural gas H from Russia [37]. The O<sub>2</sub> supply for the oxyfuel combustion is completely provided by an ASU, which uses electricity from the German grid.

**Table 2.** Composition of fuel mixtures in the analyzed scenarios. Each composition is given in mole fractions  $x_i$  (mol/mol fuel). Mole fractions are equivalent to volume fractions  $\varphi_i$  at standard conditions (STP, 0 °C and 1.013 bar). The initial composition of natural gas H from Russia for the conventional scenario is adapted from [37].

Component	Conventional 0 vol% H <sub>2</sub> mol/mol	Scenario 10 vol% H <sub>2</sub> mol/mol	Scenario 25 vol% H <sub>2</sub> mol/mol	Scenario 50 vol% H <sub>2</sub> mol/mol	Scenario 75 vol% H <sub>2</sub> mol/mol	Scenario 100 vol% H <sub>2</sub> mol/mol
Methane (CH <sub>4</sub> )	0.9642	0.8678	0.7238	0.4822	0.2410	0.0000
Ethane (C <sub>2</sub> H <sub>6</sub> )	0.0258	0.0232	0.0193	0.0129	0.0065	0.0000
Propane (C <sub>3</sub> H <sub>8</sub> )	0.0017	0.0015	0.0013	0.0008	0.0004	0.0000
Carbon dioxide (CO <sub>2</sub> )	0.0033	0.0030	0.0025	0.0016	0.0008	0.0000
Nitrogen (N <sub>2</sub> )	0.0050	0.0045	0.0037	0.0026	0.0013	0.0000
Hydrogen (H <sub>2</sub> )	0.0000	0.1000	0.2500	0.5000	0.7500	1.0000

## 4. Materials and Methods

Most of the simulation models in this work are developed using the hyperphysical modeling language Modelica, using Dymola as a development environment. Modelica is characterized by its acausal modeling approach, which allows the description of transient physical systems [38]. Due to the open-source approach of Modelica, a large number of libraries are available. Wherever possible, existing Modelica modeling approaches are used to describe the process shown in Figure 3. In a first step, the usage of established libraries for modeling renewable energy sources is described. Since a detailed consideration of the thermodynamic states of the hydrogen storage process is not meaningful at this point, the storage model contained in the open energy modeling framework (OEMOF) was used [39]. Thereupon, a more detailed description of the custom-developed oxyfuel melting system model is provided.

### 4.1. Modelling of Renewable Energy Sources

The renewable energy sources wind power (WP) and photovoltaic (PV) are modeled based on existing Modelica libraries. The required boundary conditions, such as ambient temperature, solar irradiance, and wind speed, are extracted from the test reference year (TRY) data sets of the German meteorological service (DWD) [40].

#### 4.1.1. Photovoltaic Power Plants

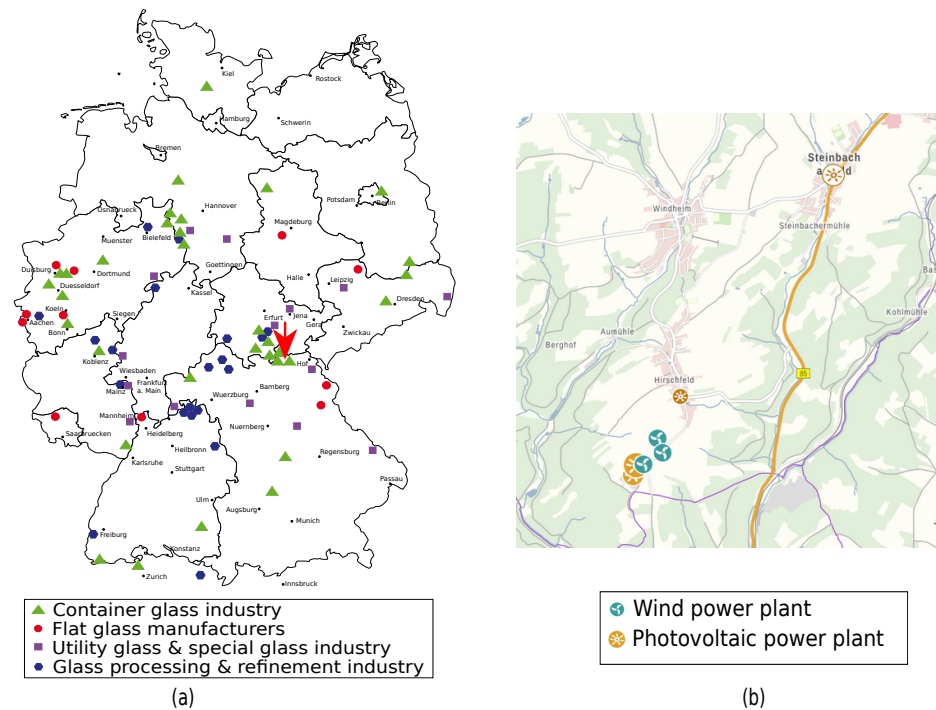
The models by Brkic et al. [41] are used to calculate the sun angle and sun azimuth at a given location. To determine the irradiance on an inclined surface, the approach of Liu et al. [42] is applied. To calculate the alternating current (AC) and direct current (DC) output of the PV plant, the modeling approach of the TransiEnt library is used [43,44]. Thus, the DC power generation of arbitrary PV modules is determined based on irradiation and temperature-dependent power curves. The AC power output is calculated based on a power-dependent efficiency curve of specific inverters. For the later evaluation of CO<sub>2</sub> emissions from power generation by PV plants, specific emissions of  $s_{CO_2e,PV} = 67 \text{ g CO}_2\text{-eq./kWh}$ , are assumed [45].

#### 4.1.2. Wind Power Plants

The simulation models for wind turbines are also based on the TransiEnt library [43,44]. The wind speed at 10 m above ground ( $v_{10}$ ), given in the TRY data set, is extrapolated to the wind speed at hub height ( $v_{Hub}$ ) of a given wind turbine following the approach given by Hau [46]:

$$v_{Hub} = v_{10} \cdot \frac{\ln \frac{h_{Hub}}{z_0}}{\ln \frac{h_{Ref}}{z_0}} \quad (1)$$

$h_{Ref}$  is the reference height of the wind speed,  $h_{Hub}$  the hub height of the wind turbine, and  $z_0$  is the roughness length of the surrounding terrain surface. In this work, a roughness length representative for forest,  $z_0 = 0.8$  is assumed, based on the local conditions of the exemplary WP plants investigated in this study (see Figure 4). The extrapolated wind speed profile ( $v_{Hub}$ ) is used to determine the electrical power output of arbitrary wind power plants from their power curves [47]. A hysteresis controller of the Modelica Standard Library (MSL) is used to model minimal startup wind speed for power generation and safety shutdown if wind speed exceeds a critical value. The specific CO<sub>2</sub> emissions reported in [45] for onshore wind power generation (11 g CO<sub>2</sub>-eq./kWh) are used.



**Figure 4.** (a) Distribution of the glass industry in Germany, highlighting the selected location (red arrow, Steinberg am Wald) for the case study of the exemplary evaluation of CO<sub>2</sub> abatement costs [1]. (b) Emergence of renewable energy in the vicinity of the selected case study site (i.e., Steinberg am Wald, latitude: 50.404387°, longitude: 11.332333°, WGS84 coordinates) [48].

#### 4.2. Electrolysis

PEM electrolysis is modeled based on mass and energy balance, following the approach of [49]. Reaction Equation (2) is used as the basis for the mass balance. An efficiency of 72 % [49] based on the lower heating value of H<sub>2</sub> is assumed to calculate the H<sub>2</sub> mass flow. The energy demand of the electrolysis is assumed to be covered by the renewable energy sources PV and WP.



#### 4.3. Air Separation Unit

The ASU model is based on the specific primary energy demand  $e_{ASU}$  to produce the mass flow of oxygen  $\dot{m}_{\text{O}_2,req}$  required for the combustion. The resulting total energy demand  $E_{ASU}$  is calculated as:

$$\frac{dE_{ASU}}{dt} = (\dot{m}_{\text{O}_2,req} - \dot{m}_{\text{O}_2,El}) \cdot e_{ASU} = P_{ASU} \quad (3)$$

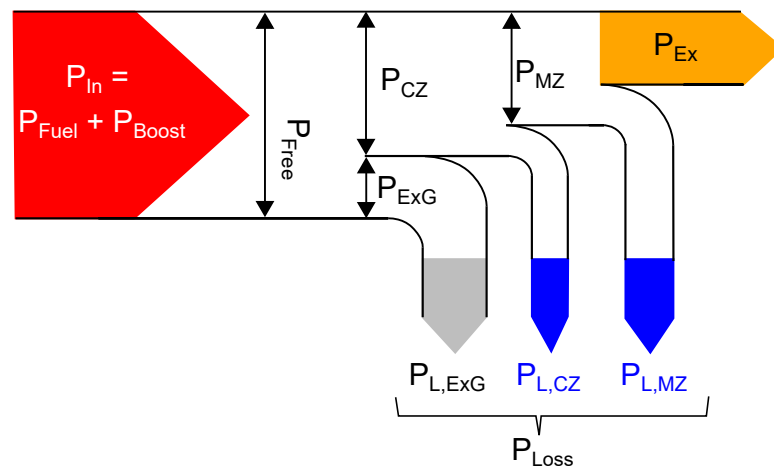
By incorporating O<sub>2</sub> from electrolysis,  $\dot{m}_{\text{O}_2,El}$  can be reduced and thus the required ASU power is reduced.  $e_{ASU}$  is set to 0.32 kWh/kg O<sub>2</sub> [50].  $E_{ASU}$  allows to calculate the CO<sub>2</sub> emissions resulting from electricity consumption from the German electric energy mix (in this work: 427 g CO<sub>2</sub>-eq./kWh [51]). The output fluid stream from the ASU is referred to as the Oxygen carrier fluid (OCF) in the following. As ASUs can produce oxygen purities of >99.0 vol% [50], the OCF is assumed to be a mixture of oxygen and nitrogen ( $x_{\text{O}_2,OCF} = 0.99 \text{ mol/mol}$  and  $x_{\text{N}_2,OCF} = 0.01 \text{ mol/mol}$ ). This is legitimate since the nitrogen component both represents the unavoidable false air component in oxyfuel glass melting systems, and allows a comparison with conventional air combustion.

#### 4.4. Oxyfuel Glass Melting Tank

So far, no detailed modeling approach for a continuously operating oxyfuel melting tank with additional electrical boosting is described in literature. Therefore, a novel model is developed. The model of a continuously operated oxyfuel melting tank is composed of (i) a heat balance and (ii) a combustion submodel.

##### 4.4.1. Heat Balance Submodel

The heat balance submodel is adapted from the heat balances for regenerative furnaces, presented in [15,23,52,53]. The recirculated heat from the exhaust gas heat exchanger is removed, as shown in Figure 5. Therefore, this heat balance represents a simplified version of a regenerative melting tank, which is justified as indicated in Section 2 [18].



**Figure 5.** Efficiencies of fuel-fired and electric furnaces as a function of pull rate, adapted from Ref. [10]. In this work, a melting tank efficiency  $\eta_{MT}$  of 42% is assumed for a 100 t/d oxyfuel melting system (see Section 4.4.1).

Thus, the heat balance in terms of the total power released in the furnace  $P_{Free}$  is calculated as:

$$P_{Free} = P_{In} = P_{Fuel} + P_{Boost} = P_{Fuel} + \gamma_B \cdot P_{In} \quad (4)$$

$P_{In}$  is the external power input of the melting system, defined as the sum of the heating power generated by fuel combustion  $P_{Fuel}$  and additional electrical boosting  $P_{Boost}$ .  $P_{Boost}$  is calculated as a fraction  $\gamma_B = 0.09$  of  $P_{In}$  [23,53]. Power losses  $P_{Loss}$  are composed of the remaining power in the exhaust gas  $P_{ExG}$  and losses in the combustion  $P_{L,CZ}$  and melting zone  $P_{L,MZ}$ . As no comprehensive data on wall losses, etc., is available, no specific calculation of these parameters is included. Rather, the most important parameter of the melting tank, i.e., the total power input  $P_{In}$  and the exploited power  $P_{Ex}$ , are calculated separately following the approach provided in [15]:

$$P_{Ex} = \frac{d(H_{Ex} \cdot z_r)}{dt} \quad (5)$$

$H_{Ex}$  denotes the enthalpy in the molten glass, leaving the tank for post-processing and  $z_r$  the pull rate of the melting tank. Within this work, a glass melting tank with a constant pull rate of  $z_r = 100$  t/d is assumed. However, the pull rate is in- or decreased within certain limits depending on economic pressure. This can result in considerably faster furnace abrasion [15]. The remaining variable  $H_{Ex}$  in Equation (6) is calculated as:

$$H_{Ex} = (1 - \gamma_c) \cdot \Delta H_{chem}^0 + \Delta H(T_{Ex}) \quad (6)$$

whereby  $\gamma_c$  is the cullet fraction,  $\Delta H_{chem}^0$  the standard enthalpy demand of batch-to-melt conversion (referring to the thermodynamic standard states given in [15] of 25 °C and 1.00 bar) and  $\Delta H(T_{Ex})$  the enthalpy of the melt at exit temperature  $T_{Ex}$ . The exact determination of  $\Delta H_{chem}^0$  and  $\Delta H(T_{Ex})$  requires detailed information about the glass composition as well as the properties of raw materials used for the batch [53]. Thus, in this work,  $H_{Ex}$  is assumed to be 550 kWh/t glass, which is representative for soda-lime glass, with  $\gamma_c = 50\%$  and  $T_{Ex} = 1350$  C [52]. These parameters can be considered as standard glass composition used in the container glass industry [53]. Based on  $P_{Ex}$  and the energy efficiency  $\eta_{MT}$  of the melting tank,  $P_{In}$  is calculated as:

$$\eta_{MT} = \frac{P_{Ex}}{P_{In}} \Rightarrow P_{In} = \frac{P_{Ex}}{\eta_{MT}} \quad (7)$$

As already indicated in Section 2, the efficiency of fuel-fired tanks is a function of its pull rate (Figure 2) [10]. At the assumed pull rate of  $z_r = 100$  t/d, fuel-fired (e.g., regenerative) melting tanks are expected to have an efficiency of about 37% [10] (Figure 2). However, oxyfuel furnaces are associated with a 5–10% increase in efficiency over a comparable regenerative furnace [13]. This increased efficiency can be attributed to the higher adiabatic flame temperature and improved heat capacity flow [15]. Thus, in this work, and energy efficiency of  $\eta_{MT} = 0.42$  is assumed for the oxyfuel melting tank.

To further characterize the melting tank model, the dimensionless key indicators temperature efficiency  $\zeta$  and heat capacity flow ratio  $z_{HL}$  are introduced. The temperature efficiency  $\zeta$  is defined as:

$$\zeta = \frac{T_{ex} - T_0}{T_{ad} - T_0} \quad (8)$$

$T_{ex}$  is the exit temperature of the glass melt,  $T_{ad}$  the adiabatic flame temperature and  $T_0$  the ambient temperature, also representing the three most relevant temperature levels of a melting tank. Common values for  $T_{Ex}$  and  $T_0$  are 1300 °C and 25 °C, respectively [15,52]. The modeling approach used to calculate  $T_{ad}$  is described in Section 4.4.3.

The heat capacity flow ratio  $z_{HL}$ , is calculated as:

$$z_{HL} = \zeta \cdot \frac{P_{In}}{P_{Ex}} \cdot \left(1 - \frac{P_{Boost}}{P_{In}}\right) = \frac{\dot{m}_{hot} \cdot c_{p,hot}}{\dot{m}_{cold} \cdot c_{p,cold}} \quad (9)$$

$\dot{m}_{hot} \cdot c_{p,hot}$  is the heat capacity flow into the furnace by the hot stream (combustion gases) and  $\dot{m}_{cold} \cdot c_{p,cold}$  the heat capacity flow from the cold stream (batch and molten glass).  $z_{HL}$  is the ratio of the heat input into the furnace by the hot stream (combustion gases,  $\dot{m}_{hot} \cdot c_{p,hot}$ ) to the cold stream (batch and molten glass,  $\dot{m}_{cold} \cdot c_{p,cold}$ ). Thus,  $z_{HL}$  evaluates this efficiency at an idealized infinitely large heat exchange surface or infinitely large heat transfer rate. Consequently,  $z_{HL}$  would reach its optimum at  $z_{HL} \rightarrow 1.00$  [15].

#### 4.4.2. Combustion Submodel

The combustion submodel aims to evaluate CO<sub>2</sub> emissions caused by natural gas combustion and their associated changes for different volume fractions of H<sub>2</sub> in the fuel mixture. This model is based on stoichiometrically complete combustion of (longer chained) hydrocarbons and H<sub>2</sub>, following the approaches given in [54,55]. The fuel mass flow  $\dot{m}_F$  required to cover  $P_{Fuel}$  is determined by considering the mass-related inferior calorific value  $H_{i,m}$ .

$$P_{Fuel} = \dot{m}_F \cdot H_{i,m} \quad (10)$$

The excess O<sub>2</sub> rate  $\lambda$  to reduce the formation of incomplete combustion products is commonly set to 1.02–1.08 in the glass industry [15]. In this work, a value of  $\lambda = 1.05$  is used. For the sake of model reduction, incomplete combustion products such as CO, NO<sub>x</sub> or SO<sub>x</sub> are omitted in the combustion submodel. Based on the determined fuel and O<sub>2</sub> demand, the composition and amount of exhaust gas can be calculated [54,55]. Therefore,

the fluid model “Simple flue gas for overstoichiometric O<sub>2</sub>-fuel ratios” implemented in the “Media” library of MSL is used. In this model, exhaust gas (EG) is described as a mixture of N<sub>2</sub>, O<sub>2</sub>, H<sub>2</sub>O and CO<sub>2</sub>.

To account for additional process-related CO<sub>2</sub> emissions  $\dot{m}_{\text{CO}_2, \text{batch}}$ , resulting from reactions during the batch-to-melt conversion, a constant value of 80 kg CO<sub>2</sub>-eq./t glass, representative for the container glass industry, is used [2].

#### 4.4.3. Adiabatic Flame Temperature

The adiabatic flame temperature  $T_{ad}$  is calculated within the open-source suite Cantera [56]. Thus, the GRI-MECH 3.0 mechanism is used to model advanced combustion phenomena [57]. This mechanism contains 325 reactions and 53 species, including higher chain hydrocarbons up to C<sub>3</sub>H<sub>8</sub> and various intermediates, e.g., like NO<sub>x</sub>, OH<sup>-</sup>, O<sup>+</sup>, H<sup>+</sup>. The built-in Cantera algorithm is used to equilibrate the fuel and oxidization fluid using the Gibbs minimization solver.

Inferior and gross calorific values of the various fuel mixtures are also calculated in Cantera. In this context, equilibrium analysis is omitted, following the established methods for calculating the calorific values according to industry standards [58].

#### 4.5. Economic Analysis and CO<sub>2</sub> Abatement Costs

For an economic evaluation of the presented PtH<sub>2</sub> concept, the established equivalent annual cost method is used to calculate the annual owning and operating costs for the H<sub>2</sub> storage system and the PEM electrolysis. The equivalent annual cost factor  $EAC$  is calculated by

$$EAC = \frac{(1+r)^n \cdot r}{(1+r)^a - 1} \quad (11)$$

with the annual interest rate  $r_a = 0.05$  and the number of years  $a = 20$  assumed for the components electrolysis and H<sub>2</sub> storage.

The CO<sub>2</sub>-abatement costs  $k_{\text{CO}_2}$  are calculated as

$$k_{\text{CO}_2} = \frac{k_M - k_{Ref}}{k_{e,Ref} - k_{e,M}} = \frac{\Delta k}{\Delta k_{e,M}} \quad (12)$$

with  $k_M$  being the specific energy costs in €/t glass of the CO<sub>2</sub> reduction measure and  $k_{Ref}$  the specific energy costs of the conventional scenario.  $k_e$ ,  $M$  and  $k_{e,Ref}$  are the specific CO<sub>2</sub> emissions of the reduction measure and the conventional scenario in kg CO<sub>2</sub>-eq./t glass, respectively [59].

#### 4.6. Energy System Optimization

The Open Energy Modeling Framework (OEMOF) is used for energy system optimization [39]. In order to optimize the system described in Figure 3, an OEMOF energy system is defined. This system consists of separate buses for electricity and H<sub>2</sub>. The electricity bus is fed by the intermittent renewable sources wind power and photovoltaic, whereas excess power is modeled as an energy sink. The electrolysis is connected to this bus as a transformer with the defined efficiency of 72%, and the given specific investment, where the peak power is defined as an optimization variable. The H<sub>2</sub> storage is integrated by a generic storage model, with an in- and output efficiency of 99%, no loss rate, and no initial storage capacity. The maximum storage capacity is defined as an optimization variable. A constraint for the H<sub>2</sub> storage system is the requirement to provide sufficient H<sub>2</sub> for each investigated mixing scenario. Figure 6 shows a block diagram of the OEMOF energy system.

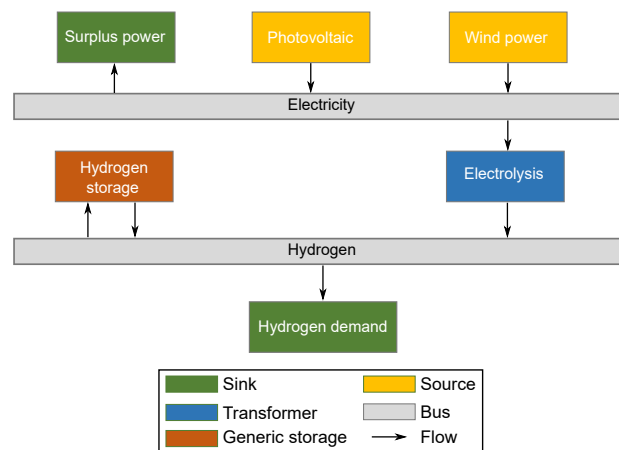


Figure 6. Block diagram of the OEMOF energy system.

## 5. Results and Discussion

To evaluate the impact of the PtH<sub>2</sub> process on oxyfuel glass melting, the changes in some characteristic values for fuel classification in the glass industry are analyzed, in particular, the inferior calorific (ICV) values and gross calorific values (GCV).

### 5.1. Changes of Heat Content in the Fuel Mixture

Figure 7 shows the changes in the mass- and volume-related heating value of the fuel mixture as a function of the volumetric hydrogen content. The gravimetric values increase with higher hydrogen contents, while the volumetric values decrease. This is due to the high gravimetric energy density of H<sub>2</sub>, in combination with its low density, and is further enhanced by the high density of the other fuel components (see Table 2). The volumetric values decrease to 50% at a volume fraction at approx. 75 vol% H<sub>2</sub> in the fuel mixture. This may result in more gas having to be transported through the infrastructure (e.g., via pipes, valves, compressors) to the burners to meet the energy demand. This can lead to process engineering challenges, but also considerable additional costs for glass manufacturers. In addition, H<sub>2</sub> embrittlement of the pipeline material has to be considered critical at higher H<sub>2</sub> vol% in the fuel mixture [60].

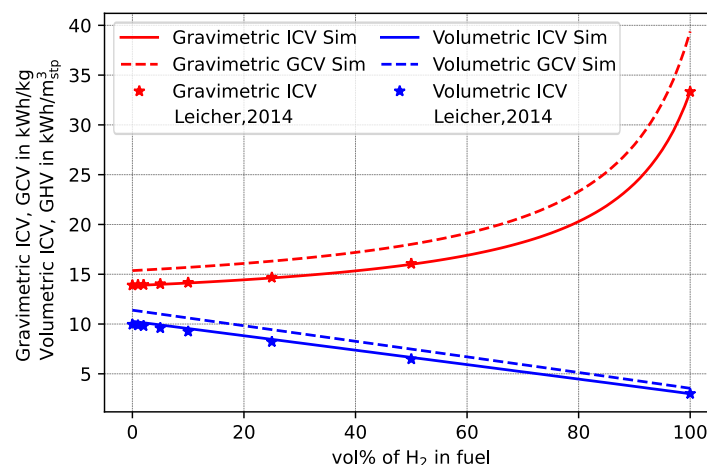
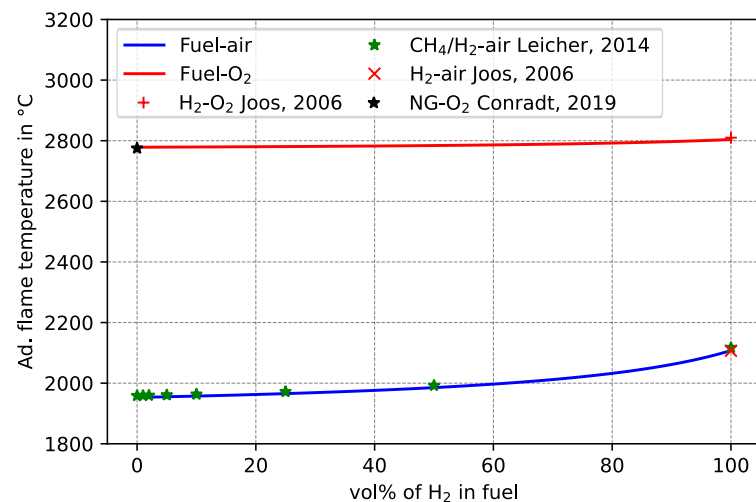


Figure 7. Predicted changes in mass- and volume-related gross calorific values (GCV) and inferior calorific (ICV) with increasing H<sub>2</sub> vol% in the fuel mixture, compared to literature data given in Leicher, 2014 [61].

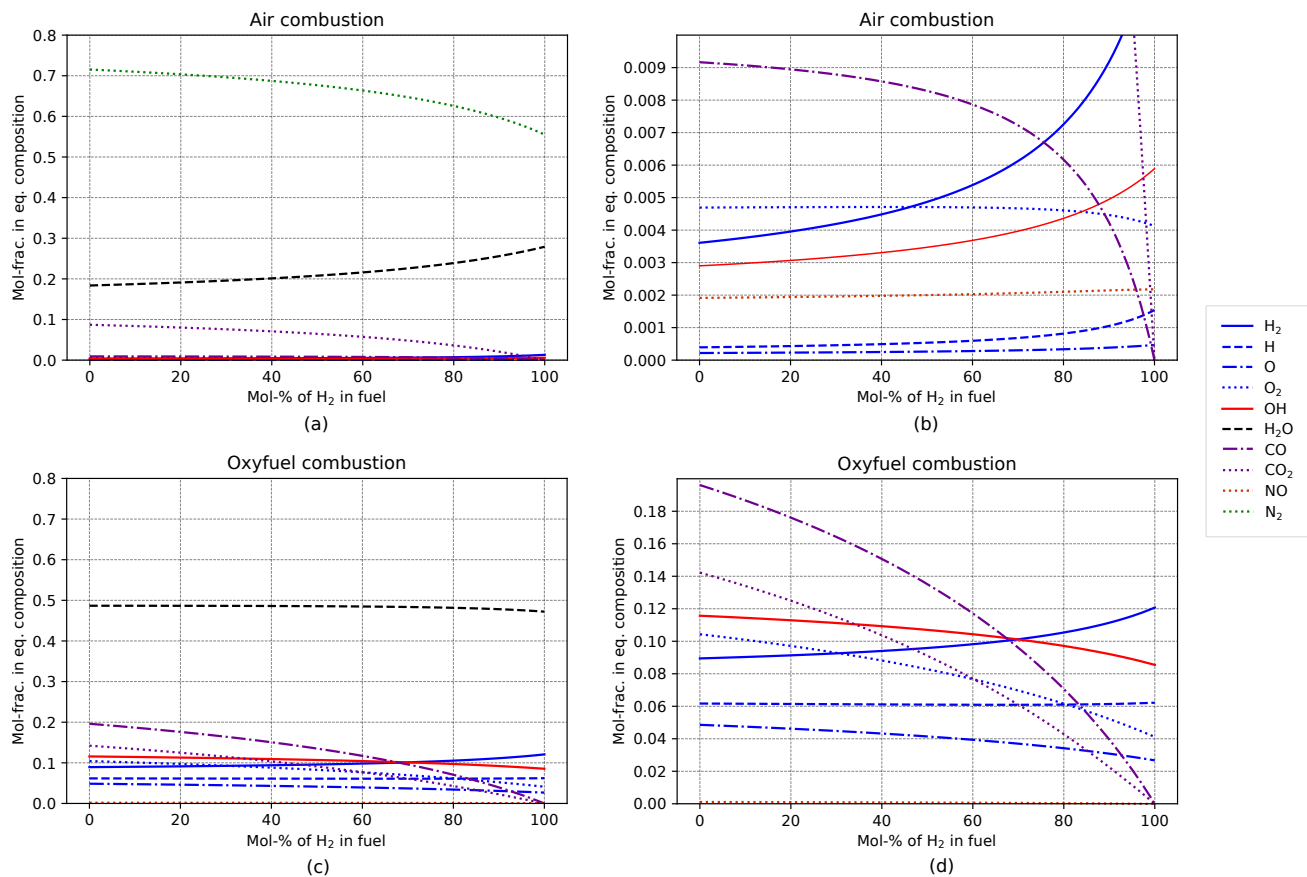
### 5.2. Changes of Adiabatic Flame Temperature

Besides ICV and GCV, adiabatic flame temperature ( $T_{ad}$ ) is used to evaluate fuels and the efficiency of glass melting furnaces. Figure 8 shows the comparison of the predicted  $T_{ad}$  with literature values. For combustion of the fuel mixtures in ambient air (21 vol% O<sub>2</sub>, 79 vol% N<sub>2</sub>)  $T_{ad}$  increases from approx. 1953 °C at 0 vol% H<sub>2</sub> to approx. 2108 °C for pure H<sub>2</sub>-air combustion. This is due to both the more efficient oxidation of H<sub>2</sub> compared to hydrocarbon compounds, as well as the decreasing hydrocarbon compound concentrations in the fuel mixture. This assumption is substituted by the consideration of the detailed composition of combustion gases (Figure 9a,b). Oxyfuel combustion (Figure 8, red solid line) only yields a slight increase, from approx. 2779 °C at 0 vol% H<sub>2</sub> to approx. 2804 °C for pure H<sub>2</sub>-O<sub>2</sub> combustion. Based on the product gas composition after adiabatic combustion (Figure 9c,d), it can be concluded that other combustion products play a more important role in oxyfuel combustion than in combustion with common air, e.g., in oxyfuel combustion significantly more CO<sub>2</sub> is formed than CO and no NO is present. Since CO<sub>2</sub> formation is more efficient than CO formation, the increased  $T_{ad}$  for oxyfuel combustion than for air combustion can be explained. Moreover, the almost complete absence of N<sub>2</sub> at oxyfuel combustion significantly reduces the total heat capacity of the combustion gas mixture and therefore also increases  $T_{ad}$ . Compared to air combustion, the exhaust gases of oxyfuel combustion are dominated by H compounds. Thus, the influences of higher H<sub>2</sub> vol% in the fuel mixture are less significant than for common air combustion.

It can be concluded that the change from conventional air to oxyfuel combustion has a more significant impact on  $T_{ad}$  than the addition of H<sub>2</sub> to the fuel mixtures. Nevertheless, the sole consideration of  $T_{ad}$  and the heating values are not sufficient to conclude the influences of high H<sub>2</sub> contents in fuel mixtures for oxyfuel glass melting tanks.



**Figure 8.** Comparison of the predicted adiabatic flame temperatures with increasing H<sub>2</sub> vol% in the fuel mixture with literature values for CH<sub>4</sub>/H<sub>2</sub>-air combustion (Leicher, 2014 [61]), pure H<sub>2</sub>-air combustion (Joos, 2006 [54]) and natural gas (NG)-air combustion (Conradt, 2019 [15]).



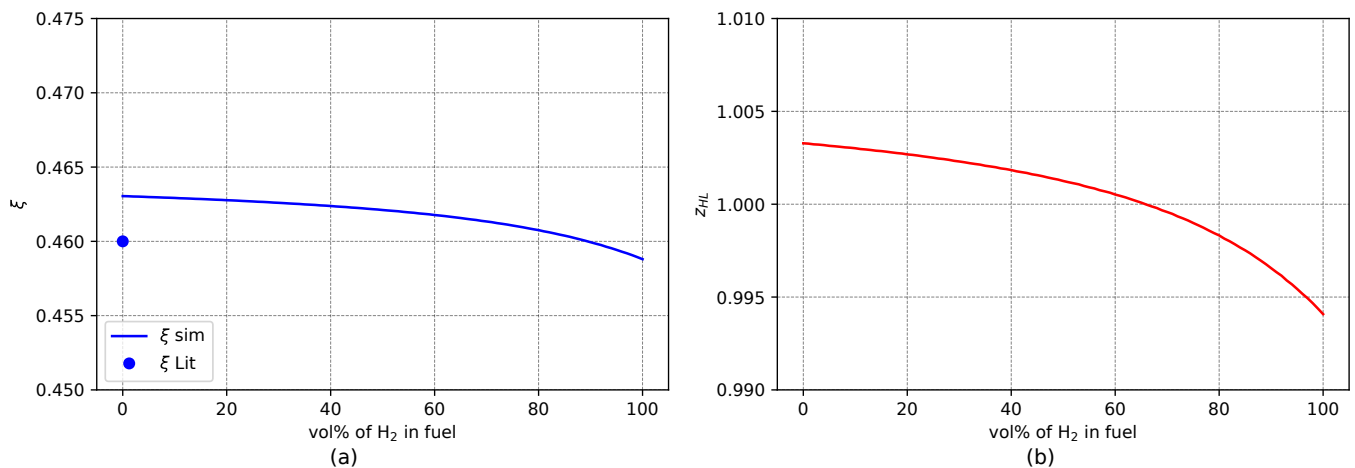
**Figure 9.** Predicted equilibrium composition of the gas mixtures after adiabatic combustion. (a) Main products of fuel-air combustion, with N<sub>2</sub>, H<sub>2</sub>O and CO<sub>2</sub> being the dominant products. (b) Highlight of the minor products of fuel-air combustion, i.e., CO, O<sub>2</sub>, H<sub>2</sub>, OH, NO, H and O. Despite their low mole fractions, CO and NO, in particular, have a significant impact on adiabatic flame temperatures. The other species can be considered negligible for adiabatic flame temperature investigations. (c) Main products of fuel-oxyfuel combustion with H<sub>2</sub>O, CO, CO<sub>2</sub>, OH and O<sub>2</sub> being dominating products. (d) Highlight of the minor products of fuel-oxyfuel combustion, i.e., H<sub>2</sub>, H and O.

### 5.3. Influences on Furnace Efficiency

The operating efficiency of a glass melting furnace is further characterized by the dimensionless key indicators temperature efficiency  $\zeta$  and heat capacity flow ratio  $z_{HL}$ , as defined by Equations (8) and (9) [15]. Usually,  $\zeta$  and  $z_{HL}$  are considered constant in previous studies for a given combustion mode. The novelty of this work is the evaluation of  $\zeta$  and  $z_{HL}$  as a function of H<sub>2</sub> vol% in the fuel rather than a constant (see Figure 10).

The calculated temperature efficiency of  $\zeta = 0.463$  (Figure 10a) shows a deviation of only 0.003 from literature data given in Ref. [15]. This is due to the fact that [15] assumed  $T_{ad} = 2780$  °C for oxyfuel combustion, whereas  $T_{ad}$  was explicitly calculated as 2779 °C in this work. Moreover, as the calculated  $T_{ad}$  varies only slightly ( $\Delta T_{ad} = +25$  C) for oxyfuel combustion,  $\zeta$  also only decreases moderately to 0.459 for higher H<sub>2</sub> contents in the fuel mixture. The heat capacity flow ratio  $z_{HL}$  (Figure 10b) deviates by 0.003 from the ideal value of 1.000 for pure NG-O<sub>2</sub> combustion and is, furthermore, within the typical range of 0.900–1.100 [15] for oxyfuel furnaces. At approx. 65 vol% H<sub>2</sub> in the fuel, the ideal value of  $z_{HL} = 1.000$  is reached. At higher vol% of H<sub>2</sub>,  $z_{HL}$  decreases to 0.994.

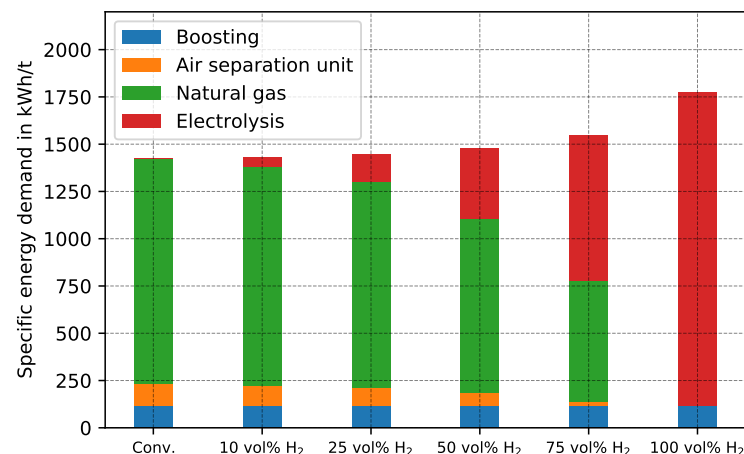
Based on the analysis of the key values  $\zeta$  and  $z_{HL}$ , it can be concluded that a higher H<sub>2</sub> vol% in the fuel has only a minor effect on the furnace efficiency. However, besides these energetic effects, chemical changes in the furnace atmosphere were found to have a significant effect on the melting process at pure H<sub>2</sub>-O<sub>2</sub> combustion [24]. As these chemical effects are not considered in this work, they should be investigated further in future studies.



**Figure 10.** (a) Temperature efficiency  $\zeta$  with increasing H<sub>2</sub> shares in fuel, compared to a literature value for oxyfuel combustion given in Ref. [15]. (b) Heat capacity flow ratio  $z_{HL}$  with increasing vol% of H<sub>2</sub> in fuel for a normalized pull of  $\phi = 1.0$ . Ideal values are reached for  $z_{HL} \rightarrow 1.00$ . Literature values of  $z_{HL}$  for oxyfuel fired furnaces range between 0.85 and 1.15 [15].

#### 5.4. Specific Energy Demand

The change in specific energy demand resulting from the presented PtH<sub>2</sub> concept is a major point of interest. To validate the models presented in this paper in terms of their ability to predict specific energy demand, the conventional scenario (Figure 11, first bar) is compared with literature data. The predicted total specific energy demand of 1423 kWh/t glass is in accordance with [62], which reported an average value of 1431 kWh/t glass for 300 container-glass furnaces, for predominantly conventional air combustion. The slightly lower specific energy demand of 8 kWh/t can be considered as a key indicator for the low prevalence of oxyfuel melting processes in the container glass industry. The low energy savings and comparatively high investments costs in oxygen infrastructure, as well as more expensive ceramics for the furnace brickwork, prevent a lucrative return of investment.



**Figure 11.** Comparison of the proportional specific energy demand per ton of molten glass caused by boosting, air separation unit (ASU), natural gas and electrolysis between the integration of PtH<sub>2</sub> in oxyfuel-based glass melting processes with various H<sub>2</sub> vol% in the fuel mixture and the conventional scenario (i.e., 0 vol% H<sub>2</sub> in the fuel mixture).

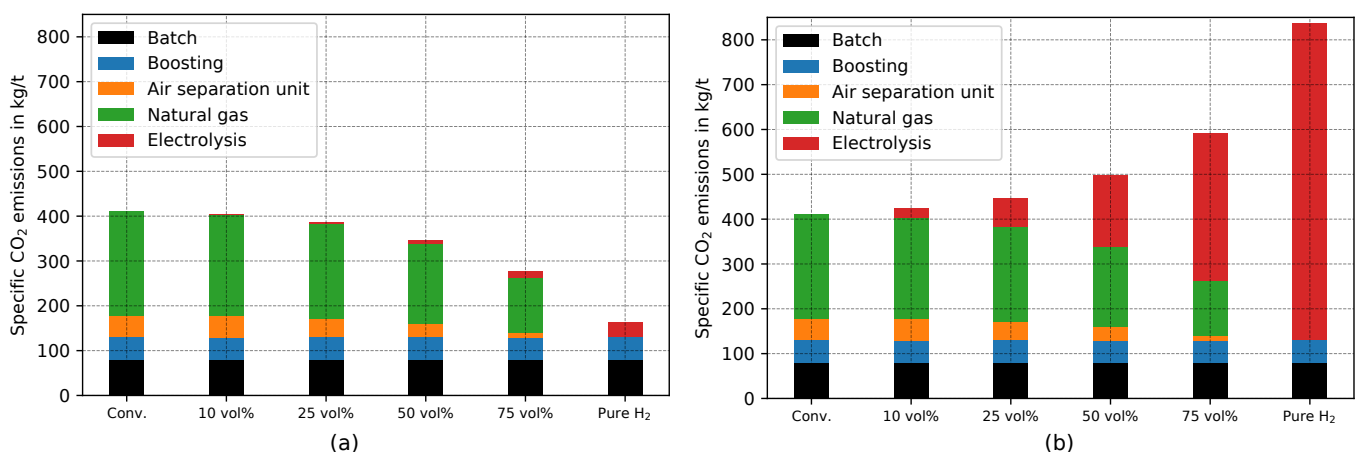
Figure 11 shows the changes of specific energy demand for the scenarios of 10, 25, 50, 75 and 100 vol% of H<sub>2</sub> in the fuel mixture. The energy demand for electrical boosting remains constant in all scenarios (118 kWh/t glass), as it is modeled as a constant fraction of the total heat demand of melting. With increasing H<sub>2</sub> vol% in the fuel mixture, the

demand for natural gas decreases (1192 kWh/t glass to 0 kWh/t), while the energy demand of electrolysis increases (0 to 1656 kWh/t glass). Moreover, the energy demand of the ASU declines (113 kWh/t glass to approx. 0 kWh/t glass), since oxygen is obtained from electrolysis and less carbon has to be oxidized during combustion.

### 5.5. Specific CO<sub>2</sub> Emissions

To validate the models presented in this paper in terms of their ability to predict the CO<sub>2</sub> emissions, the conventional scenario (Figure 12, first bar) is compared with literature data [2]. In [2], energy-related CO<sub>2</sub> emissions of 360 kg CO<sub>2</sub>-eq./t glass and process-related CO<sub>2</sub> emissions of 80 kg CO<sub>2</sub>-eq./t glass were reported as average values for the container glass industry in Germany. Energy-related emissions include natural gas combustion, electrical boosting, and the ASU, whereas process-related emissions are caused by glass batch due to carbonate reactions. For the conventional scenario, the presented modeling approach predicts energy-related CO<sub>2</sub> emissions of 331 kg CO<sub>2</sub>-eq./t (boosting: 50 kg CO<sub>2</sub>-eq./t glass, ASU: 48 kg CO<sub>2</sub>-eq./t glass, natural gas: 233 kg CO<sub>2</sub>-eq./t glass) and process-related emissions of 80 kg CO<sub>2</sub>-eq./t glass. The slightly decreased energy-related CO<sub>2</sub> emissions are associated with the lower specific CO<sub>2</sub> emissions for electricity assumed in Ref. [2] (512 g CO<sub>2</sub>-eq./kWh) compared to the present work (427 g CO<sub>2</sub>-eq./kWh). In addition, the slightly reduced energy demand of oxyfuel melting is a contributing factor. Moreover, as the deviation is still less than 10 % the validity of the simulation model is justified. It is not surprising that the predicted process-related CO<sub>2</sub> emissions match literature data perfectly, given that these were set to a constant value of 80 kg CO<sub>2</sub>-eq./t glass throughout the entire simulations.

Moreover, the specific CO<sub>2</sub> emissions were evaluated with respect to the electrical energy sources used. For renewable energy sources specific CO<sub>2</sub> emissions of 19 g CO<sub>2</sub>-eq./kWh of electricity were assumed. By using renewable energy sources, specific emissions can be reduced from 411 kg CO<sub>2</sub>-eq./t glass to 163 kg CO<sub>2</sub>-eq./t glass for pure H<sub>2</sub>-O<sub>2</sub> combustion, i.e., by 60% (Figure 12b). In contrast, if the energy demand of electrolysis is covered by electricity from the current German energy mix, the specific CO<sub>2</sub> emissions even increases (Figure 12b). In this case, the scenario for pure H<sub>2</sub>-O<sub>2</sub> combustion causes 837 kg CO<sub>2</sub>-eq./t glass, which is an increase of 104% compared to the conventional scenario. Thus, from an ecological perspective, the integration of PtH<sub>2</sub> is only reasonable if renewable energy sources are used.



**Figure 12.** (a) Specific CO<sub>2</sub> emissions with the energy demand for electrolysis being covered by wind and PV (19 g CO<sub>2</sub>-eq./kWh). (b) Specific CO<sub>2</sub> emissions with the energy demand for electrolysis energy demand being covered by the current German energy mix (427 g CO<sub>2</sub>-eq./kWh [51]).

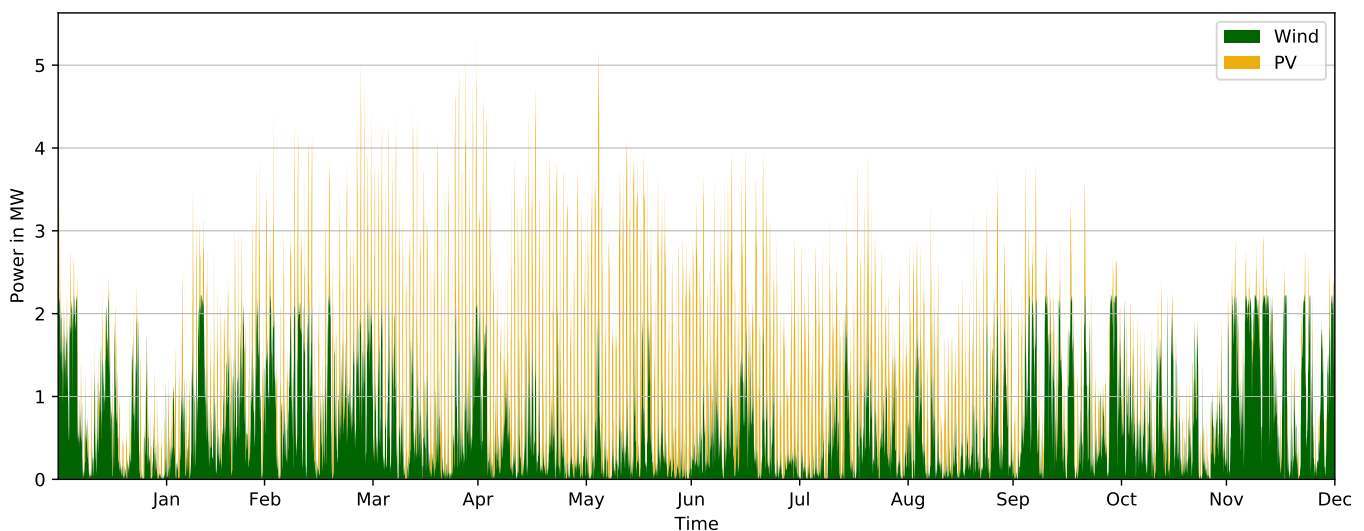
### 5.6. CO<sub>2</sub> Abatement Costs

In a case study, CO<sub>2</sub> abatement costs are calculated for Steinberg am Wald, county Kronach, Germany (latitude: 50.404387°, longitude: 11.332333°, WGS84 coordinates). This location can be considered as one of the main centers of the German container glass industry, as many companies are located here (Figure 4a). It is assumed that the energy demand for electrolysis to integrate PtH<sub>2</sub> into oxyfuel glass melting is met by a nearby renewable energy park, i.e., three wind turbines and two PV systems (Figure 4b). The technical data from this is shown in Table 3.

**Table 3.** Renewable energy sources at the selected location (i.e., Steinberg am Wald, latitude: 50.404387°, longitude: 11.332333°) [48].

	Plant	Nominal Power in kW	Hub Height in m	Year of Construction
Wind power	AN Bonus 600	600	58	2001
	AN Bonus 600	600	58	2001
	AN Bonus 1000	1000	70	2001
Total:		2200		
Photovoltaic, open field	Plant 1	2997		2017
	Plant 2	538		2007
Total:		3535		

Based on the Modelica models described in Section 4.1, the energy output of these renewable energy parks during a TRY data set provided by DWD was determined for the selected location of the case study (Figure 13). The three wind power plants produce approx. 6100 MWh and the photovoltaic plants approx. 3900 MWh of electricity during a TRY.



**Figure 13.** Calculated generated power of wind power and photovoltaic plants during a test reference year (TRY) at the selected location of the case study (Steinberg am Wald, latitude: 50.404387° longitude: 11.332333° (WGS84)).

Subsequently, these data sets were used as input variables for the optimization of the energy system presented in Figure 3. Due to the different years of construction and the, in part, lengthy operating time of these renewable energy sources, an approach based on the investment costs of the plants is highly error-prone. Therefore, an approach based on the current literature on the electricity production costs for renewable energies in Germany is chosen instead [63]. A price range of 0.035–0.063 €/kWh for electricity from open-field PV plants and 0.04–0.08 €/kWh for WP was reported [63]. Based values, as well as rather less

favorable local factors for the renewable energy sources at the selected location (i.e., low solar irradiation of 1001–1020 W/m<sup>2</sup>, low wind speeds at 70 m hub height of about 8 m/s on average over TRY), electricity costs of 0.05 €/kWh from the described energy sources are assumed in the case study. Capital expenditures (CAPEX) and operational expenditures (OPEX) for electrolysis ( $\approx$ 1200 €/kW, [33]) and H<sub>2</sub> (storage (1100 €/kWh, [64])) are adopted from current literature. These cost parameters (i.e., electricity costs, CAPEX, OPEX) and the local energy profile (Figure 13), were subsequently used as input data in the OEMOF model.

Exemplary optimizations are performed for the 10 vol% H<sub>2</sub> and the 25 vol% H<sub>2</sub> scenario, respectively. Depending on the scenario, H<sub>2</sub> storage capacity and electrolysis power changes resulting in different prices for the generated H<sub>2</sub> (Table 4). The optimization based on the given TRY data set and the given high feed-in of renewable energy results in very high full load hours (FLH) of the electrolysis for both scenarios. Therefore, H<sub>2</sub> costs are quite low in these scenarios.

**Table 4.** Parameters of the optimized PtH<sub>2</sub> integration into oxyfuel glass-melting processes for the boundary conditions (location, renewable energy sources, TRY) with respect to cost minimization.

	Unit	Scenario 10 vol% H <sub>2</sub>	Scenario 25 vol% H <sub>2</sub>
PEM input power	kW	272	966
PEM electrolysis FLH	h	6979	5544
Used energy of total supply	%	19	53
H <sub>2</sub> Storage capacity	MWh	4.65	500.00
H <sub>2</sub> Costs	€/kWh	0.092	0.224
	€/m <sup>3</sup> STP	0.28	0.67
	€/kg H <sub>2</sub>	3.07	7.48

To determine the energy costs of the initial scenario (i.e., oxyfuel glass melting with no integration of PtH<sub>2</sub>), assumptions had to be made regarding the energy costs of glass manufacturers. The energy demand of the glass industry is currently mainly covered by natural gas (see Section 1). Therefore, German glass companies are not commonly considered energy-intensive in terms of electricity demand, and hardly benefit from exemptions from German energy taxes. As a result, a value of 0.12 €/kWh is assumed for electricity costs. No taxes, levies, or charges are considered for the electricity production costs from renewable energy sources. Further, 0.03 €/kWh for natural gas and 0.10 €/m<sup>3</sup> (STP) of O<sub>2</sub>, generated by ASU are assumed. The results of this specific cost distribution for the different estimated scenarios are shown in Table 5.

**Table 5.** Comparison of the various and total specific energy costs and CO<sub>2</sub> abatement costs of the conventional scenario (i.e., merely oxyfuel glass melting) with the scenarios including the integration of a PtH<sub>2</sub> into oxyfuel glass melting with 10 vol% H<sub>2</sub> and 25 vol% H<sub>2</sub> in the fuel mixture, respectively.

	Conventional Scenario		10 vol% H <sub>2</sub> Scenario		25 vol% H <sub>2</sub> Scenario	
	Demand	Specific Costs €/t Glass	Demand	Specific Costs €/t Glass	Demand	Specific Costs €/t Glass
Boosting	118 kWh/t glass	14.16	118 kWh/t glass	14.16	118 kWh/t glass	14.16
Natural gas	1192 kWh/t glass	35.76	1154 kWh/t glass	34.62	1086 kWh/t glass	32.58
O <sub>2</sub> from ASU	243 m <sup>3</sup> /t glass	24.30	240 m <sup>3</sup> /t glass	24.00	220 m <sup>3</sup> /t glass	22.00
H <sub>2</sub>	0 kWh/t glass	0.00	156 kWh/t glass	3.50	106 m <sup>3</sup> /t glass	23.78
<b>Total</b>		<b>74.22</b>		<b>76.28</b>		<b>92.52</b>
CO <sub>2</sub> em.	411 kg CO <sub>2</sub> -eq./t glass	—	404 kg CO <sub>2</sub> -eq./t glass	—	386 kg CO <sub>2</sub> -eq./t glass	—
CO <sub>2</sub> ab. costs			295 €/t CO <sub>2</sub>		732 €/t CO <sub>2</sub>	

The integration of PtH<sub>2</sub> into oxyfuel melting with 10 vol% H<sub>2</sub> and 25 vol% H<sub>2</sub> yields high CO<sub>2</sub> abatement costs of 295 €/t CO<sub>2</sub> and 732 €/t CO<sub>2</sub> in the case study, respectively. In the 10 vol% H<sub>2</sub> scenario, this is due to the only slight reduction in CO<sub>2</sub> emissions in combination with high costs for the PtH<sub>2</sub> infrastructure. The further significant increase in CO<sub>2</sub> abatement costs for the 25 vol% H<sub>2</sub> scenario is associated with a significant increase in the required H<sub>2</sub> storage capacity (factor of 100) and associated H<sub>2</sub> generation costs (factor

of 2.5) compared to the 10 vol% H<sub>2</sub> scenario (see Table 4). However, it should be noted that due to the low electrolysis power required, comparatively high FLH of the electrolysis are achieved, e.g. favoring the H<sub>2</sub> generation costs in the 10 vol% H<sub>2</sub> scenario. Moreover, storage capacity could be reduced for the 25 vol% scenario by repowering of the small-scale wind turbines.

In this work, the assessment of the CO<sub>2</sub> abatement costs associated with the integration of PtH<sub>2</sub> in oxyfuel glass production is based on the evaluation of a case study for a single location. However, this location was found not to offer the optimal conditions for this concept, e.g., with regard to the availability of renewable energy. At the same time, this enabled a neutral evaluation of the concept without artificial fine-tuning. Therefore, further factors and methods for increasing the profitability of integrating PtH<sub>2</sub> into oxyfuel glass production are presented below:

- A battery-storage concept for emission reduction of electrical boosting and ASU power supply should be considered. Through the associated use of renewable electricity, the remaining CO<sub>2</sub> emissions of both ASU and electrical boosting can be significantly reduced.
- The use of so-called green electricity plans that exhibit lower specific CO<sub>2</sub> emissions for electricity supplied from the grid will allow a further reduction in CO<sub>2</sub> emissions. In this context, electricity market-based control and optimization of the operating strategy for the PtH<sub>2</sub> can provide further cost benefits.
- Developments in electrolysis production technology through more intensive use of PtH<sub>2</sub> in other sectors may result in a further reduction in CAPEX.
- Political funding programs and changing situations of CO<sub>2</sub> taxation should be taken into account.

Therefore, it is obvious that the concept presented in this work on integrating PtH<sub>2</sub> into oxyfuel melting has a high techno-economic potential for improvement. Nevertheless, this process concept could pave the way for CO<sub>2</sub> reduction in the glass industry or, ideally, even for completely CO<sub>2</sub>-neutral glass production.

## 6. Conclusions

In this work, a process concept for the step-wise integration of PtH<sub>2</sub> processes into oxyfuel glass melting is presented, based on simulations. In this way, changes in specific energy demand and associated specific CO<sub>2</sub> emissions can be evaluated as a function of the H<sub>2</sub> content in the fuel mixture, the fuel composition, the combustion itself, and certain furnace parameters. The following conclusions can be drawn:

The addition of H<sub>2</sub> to the fuel mixture has a less significant effect on the adiabatic flame temperature of oxyfuel combustion than on conventional fuel-air combustion. As the H<sub>2</sub> content in the fuel increases, the specific energy demand caused by the PtH<sub>2</sub> concept is expected to increase by a maximum of 25% compared to conventional oxyfuel glass melting scenarios. Using the PtH<sub>2</sub> concept, CO<sub>2</sub> emissions may decrease by up to 60% if required electricity is covered by renewable energy sources. However, when using the conventional German energy mix, CO<sub>2</sub> emissions are twice as high as for conventional oxyfuel melting process. The integration of PtH<sub>2</sub> concept in the glass industry still shows high ideal techno-economic development potential, e.g., with regard to large-scale renewable energy sources, intelligent storage concepts and optimized energy purchasing.

The findings of this study motivate future numerical and experimental studies on the influences of high H<sub>2</sub> amounts in the fuel mixture on glass-industry-specific combustion systems.

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## Abbreviations

The following abbreviations are used in this manuscript:

AC	Alternating current
ASU	Air separation unit
CAPEX	Capital expenditures
DC	Direct current
DWD	German metrological service
EAC	Equivalent annual cost factor
EEX	European energy exchange
EU-ETS	European Union emission trading system
FLH	Full load hours
GCV	Gross calorific values
ICV	Inferior calorific value
MSL	Modelica standard library
OEMOF	Open energy modeling framework
OPEX	Operational expenditures
PEM	Proton exchange membrane
PtH <sub>2</sub>	Power-to-Hydrogen
PV	Photovoltaic
SCM	Submerged combustion melter
STP	Standard temperature and pressure conditions (0 °C and 1.013 bar)
TCR	Thermochemical reformation
TRY	Test reference year
WP	Wind power

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