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SOLAR HYDROGEN PRODUCTION BY A TWO-STEP CYCLE BASED ON MIXED IRON OXIDES

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ABSTRACT

A very promising method for the conversion and storage of solar energy into a fuel is the dissociation of water to oxygen and hydrogen, carried out via a two-step process using metal oxide redox systems such as mixed iron oxides, coated upon multi-channeled honeycomb ceramic supports capable of absorbing solar irradiation, in a configuration similar to that encountered in automobile exhaust catalytic converters. With this configuration, the whole process can be carried out in a single solar energy converter, the process temperature can be significantly lowered compared to other thermo-chemical cycles and the re-combination of oxygen and hydrogen is prevented by fixing the oxygen in the metal oxide. For the realization of the integrated concept, research work proceeded in three parallel directions: synthesis of active redox systems, manufacture of ceramic honeycomb supports and manufacture, testing and optimization of operating conditions of a thermochemical solar receiver-reactor. The receiver-reactor has been developed and installed in the solar furnace in Cologne, Germany. It was proven that solar hydrogen production is feasible by this process demonstrating that multi cycling of the process was possible in principle.

Keywords: Hydrogen, Redox System, honeycomb reactors, solar, Thermochemical Cycle

NOMENCLATURE

ReSiC - Recrystallized Silicon Carbide

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SiSiC - Siliconized Silicon Carbide SSS - solid-state synthesis SHS –self-propagating high-temperature synthesis

INTRODUCTION

By far the most attractive reaction for the production of hydrogen is the thermal decomposition of water yielding directly pure hydrogen [1]. However because of unfavorable thermodynamics, interesting yields can only be achieved at very high temperatures imposing therefore technological difficulties to any ideas trying to couple solar energy as the driving energy for the reaction [2]. The current state of the art of solar chemistry for water splitting and hydrogen production is basically focused on processes for water splitting at lower temperatures [3]. From all technological possibilities the most interesting is the thermochemical two-step water splitting process using redox systems. According to this idea, during the first step (water splitting) the reduced and therefore activated material (usually a metal oxide) is oxidized by taking oxygen from water and producing hydrogen according to the reaction:

(1)
$$MO_{reduced} + H_2O \rightarrow MO_{oxidized} + H_2$$

During the second step (re-generation) the material is reduced to be used again, delivering some of the oxygen of its lattice according to the reaction:

(2) $MQ_{oxidized} \rightarrow MQ_{educed} + O_2$

The advantage is the production of pure hydrogen without separation need of hydrogen and oxygen. The challenge is a two-step process consisting of a splitting and a regeneration step. The redox materials that have been evaluated consist of oxide pairs of multivalent metals (Fe₃O₄/FeO, Mn₃O₄/MnO[4]) or systems of metal oxide/metal (e.g. ZnO/Zn [5]) the most representative of which can be considered the Fe₃O₄/FeO system [6]. Water splitting is taking place at temperatures below 900K while the reduction of the metal oxide, i.e. the regeneration, takes place at much higher temperatures. The concept has been proven experimentally [7, 8], however the regeneration temperatures are still high. The high temperatures of the reduction of a two-step water splitting process and a concentrating solar system.

A crucial factor for the applicability of the reaction in technical processes is the stability of the redox system. Ehrensberger et al. showed that non stoichiometric iron oxides and mixed manganese oxides form stable crystals that loose their activity over several consecutive cycles [9]. A major part of the work is to develop materials that does not show this behavior. The investigations of the materials (by x-ray diffraction e.g.) is in progress and will be reported in a following paper.

The aim of the work is to exploit solar energy for the dissociation of water and the production of hydrogen. The basic idea is to combine a support structure capable of achieving high temperatures when heated by concentrated solar radiation, with a redox pair system suitable for the performance of water dissociation and at the same time suitable for regeneration at these temperatures, so that complete operation of the whole process (water splitting and regeneration of the metal oxide) can be achieved by a single solar energy converter. A highly compact reactor structure is that of a honeycomb monolith consisting of a plurality of channels on the surface of which active compounds can be coated. Such reactors made of advanced ceramics (Cordierite, SiC) find extensive application in automotive emission control [10,11].SiC-based ceramics in particular have shown superior thermal properties [12 13] and can be coated with a variety of traditional and novel techniques [10,14]. Based on this experience, SiC has been used in an advanced solar volumetric receiver [15,16] with very good performance and it was selected to be the material for the solar water splitting reactor. Therefore the four main tasks are:

- Development of novel and efficient oxide materials to carry out the water dissociation reaction, able to be reversibly oxidized and reduced within the temperature interval of 800-1200°C. The mainstream materials of choice are based on the ferrite spinel family [17]. It is aimed to utilize the material activity caused by thermodynamic instabilities in the defect chemistry, which will be introduced on purpose into the crystal structures.
- Development of ceramic monolithic structures able to absorb efficiently concentrated solar radiation up to 1.5MW/m².
- Integration of the redox material-coated monoliths on the solar reactor/receiver and development of a complete hydrogen fuel production unit based on solar energy.
- Evaluation of technical and economic feasibility of the whole process

TESTING OF A SOLAR REACTOR WITH RESPECT TO WATER SPLITTING EFFICIENCY AND REDOX MATERIAL REGENERATION CAPABILITY

To achieve these aims a solar test reactor was designed and built at the German Aerospace Center DLR, Cologne, Germany [18].

The reactor (Fig. 1) contains a ceramic support structure that is able to host different coated structures in order to compare different redox pair systems. The ceramic honeycombs tested were made from two kinds of Silicon Carbide: re-crystallized and siliconized (Re-SiC and SiSiC respectively). The housing is made of stainless steel and the reactor has a quartz window to allow a volumetric heating of a space closed to the environment.

The reactor is designed to withstand temperatures up to 1400 °C. It is connected to an exhaust treatment system to analyze and dispose the gases.

In a first step the reactor was qualified in the solar furnace of DLR in Cologne (Fig.2) as a test bed for different metal oxide-support assemblies.

a) Preparations

To enable the quantitative testing of coated honeycombs the following actions were carried out:

- Implementation and qualification of reactor and peripherals
- Determination of operational limits with regards to temperatures, mass flows, solar input power, heating ramps and detection limits
- Elaboration of detailed experimental plan for the systematic performance of the tests
- Measurement of flux densities and flux distributions allowing thermal balancing and giving input for reactor simulation



Figure 1: Solar test reactor DLR Cologne Germany.



Figure 2: The solar furnace at DLR Cologne, Germany

b) Performance of testing

Testing of uncoated monoliths

The first measurements done in the solar furnace concerned the calibration of a mass spectrometer for the quantitative analysis of the produced gases. Therefore compound gases in accurately defined amounts, with the same quantitative gas composition as expected in the product gas were measured. Systematic series of measurements regarding mass flow or concentration with the gases $N_2 + O_2$ and $N_2 + H_2$ were accomplished to determine the lower detection limit of the spectrometer for O₂ and H₂ and to provide calibration functions for the identification of the concentrations of these gases during operation with the coated honeycombs. The subsequent experiments concerned the effects of water vapor in the gas stream on the H₂O- signal and on the H₂-signal. Firstly water vapor only was fed to the reactor and afterwards mixtures of steam and H₂. The resulting difference provided the contribution of pure H₂ to the relevant signal of the mass spectrometer by giving a correction function of the contributions from the splitting of water in the spectrometer. Finally pure nitrogen was fed in to generate baselines for the relevant species H₂ and O₂.

A strong fluctuation of O_2 concentration in the exhaust (product) gas (Fig. 3) was observed when using SiSiC monoliths independently of whether the monoliths were coated or not. The fluctuation started at the time when the structure reached 800°C and lasted for approximately one hour, decreasing and gradually vanishing after about two hours. The fluctuation varied in intensity from monolith to monolith (there are SiSiC monoliths with only minor grade of fluctuation). This phenomenon was not observed when testing ReSiC monoliths (Fig. 4).



Figure 3: Progress of O_2 signal provided by a mass spectrometer using SiSiC monoliths.



Figure 4: Progress of O_2 signal provided by a mass spectrometer using a ReSiC monolith.

These fluctuations have been interpreted as alternating oxygen absorption and desorption processes. They shall occur when residues of carbon or silicon are present on the surface of the monoliths after fabrication. The consequence of this observation is to pre-treat the monoliths under oxygencontaining atmosphere to artificially age their surfaces and thereby render them inert.

Testing of coated monoliths

Monoliths coated with redox pair systems were tested subsequently – each in a separate experiment (Fig. 5). The replacement of a tested monolith was only possible after cooling of the whole reactor overnight. At the end of each test day, an abbreviated O_2 and H_2 calibration was carried out. Seven coated honeycombs were tested, four of them loaded with SHS synthesized Zn,Mn-doped iron oxides and the other three with SSS activated iron oxide [19,20] (with different values of oxide loading).

The intention of the test programme was the quantitative evaluation of capability for hydrogen production and metal oxide reduction by monitoring the progression of the two reaction steps – splitting of water vapor and regeneration of the metal oxide – by mass spectrometry. The basic procedure was to heat up the reactor to 800° C and add water vapor for 20 minutes. Then the temperature was raised to 1200° C and maintained for 20 minutes that turned out as an appropriate time for both steps of the process. This procedure was repeated at least twice and for as long as the coating was regenerated and capable of splitting water.

The following procedure was employed: regeneration temperature was mostly 1200°C (sometimes higher if necessary). During the splitting step the temperature was

gradually raised from 800°C to 1200°C (1. cycle: 800°C, 2. cycle: 900°C, 3. cycle: 1000°C, 4. cycle: 1100°C, 5. cycle: 1200°C). The amount of water vapor was varied in the splitting step.



Figure 5: Honeycomb coated with activated iron oxide in the centre of the reactor, ready for testing.

c) Results

The first solar hydrogen was successfully produced by irradiating Zn,Mn-doped iron oxide coated on a ReSiC monolith at 800 °C in a mixture of steam and nitrogen (Fig. 6). The samples were coated with the reduced material and shipped under inert conditions to Cologne, therefore it was not necessary to activate them. The amount of hydrogen produced (8,23 mmol H₂/g oxide, the loading of one sample was 3,25 g oxide) corresponded very well to that produced in laboratory experiments performed with the metal oxide in the form of powders. After completion of the water splitting step the temperature was raised to 1200 °C under flushing by pure nitrogen. This procedure initiated a release of 3,0 mmol of oxygen per g oxide. Due to sintering of the oxide coating only one cycle could be operated.



Figure 6: Mass flow of hydrogen during splitting of water on Zn,Mn-doped iron oxide on ReSiC



Figure 7: Mass flow during release of oxygen (Zn,Mn-doped iron oxide on ReSiC).

Multi-cycling was possible when applying a coating of activated Fe₂O₃ on a SiSiC monolith. The coated monolith could perform six cycles at 800 °C with respect to solar water splitting and at 1200 °C to be partly re-activated (Fig. 8). The last two cycles in Fig. 8 (cycle 7 and 8) suggested initially that the coating is significantly more active with respect to water splitting when increasing the splitting step reaction temperature (here 1200 °C). However, further investigations showed that the hydrogen production at this elevated temperature is due to participation and thereby destruction of the support material and not desirable. The splitting temperature should not exceed 1000 °C. Thus a maximum of six cycles can be stated for the set of coated monoliths tested so far. Although the kinetics of the reactions are relatively slow this is favorable to the technical process. In the experiments the reduction step was completed after 10 - 15 minutes. This time is sufficient for a reliable operation even in a large scale reactor.

Table 1 summarizes the main results of the test campaign. Except the first monolith tested all coated samples were able to split water at least once. The monoliths coated by activated iron oxide are capable of multi-cycling (up to 6 cycles).

In conclusion it can be stated that solar thermo-chemical splitting of water and hydrogen production was successfully demonstrated several times. The release of oxygen from the redox coating was also realized by using solar energy. Multi cycling of the process is possible in principle. The amounts of hydrogen produced correspond very well to laboratory results on the redox systems in the form of powders indicating that the iron oxide as coating is almost as active as the powder in laboratory tests. ReSiC was proven more suitable in these tests compared to SiSiC, due to its resistance towards higher temperatures, lower tendency of absorption/desorption of oxygen, lower tendency of reactions with the coating and easier fabrication of monoliths with higher number of channels per square inch.



Figure 8: Mass flow of hydrogen produced during subsequent splitting steps (activated Fe₂O₃ on SiSiC).

No	Coating	Loading	Support	Cycles
		[g]	material	performed
1	Zn,Mn-doped	5.94	SiSiC	7
	FeO			
2	"	3.25	ReSiC	7
3	Act. Fe ₂ O ₃	3.20	SiSiC	3
4	"	5.32	SiSiC	10
5	"	3.08	SiSiC	3
6	Zn,Mn-doped	1.34	ReSiC	6
	FeO			
7	"	11.23	SiSiC	4
8	Act. Fe ₂ O ₃	3.18	SiSiC	3

 Table 1a:
 Quantitative evaluation of test campaign.

No	Number of	Quantity	Quantity	Quantity ratio
	cycles with	(H ₂)max	(H ₂) per	(H_2) of
	H ₂	per cycle	g(oxide)	subsequent
	formation	[mmol]	[mmol/g]	cycles
1	0	0	0	-
2	1	26,8	8,2	-
3	3	*	*	1:0,19:0,20
4	8	163	30,6	1:1,48:0,41:
				0,18:0,12:0,15
	(incl.1200°C)			(:0,37:0,32)*
5	3	29,8	9,7	1:0,45:0,09
6	5	*	*	0,08:1:0,05:
				0,01:0,1
7	2	23,4	2,1	1:0,1
8	3	*	*	-

Table 1b: Quantitative evaluation of test campaign.

* no absolute values available

These results are very encouraging with respect to the feasibility of the process investigated. The concept appears to get ready for scale-up after some further improvement of the stability of the coating. Currently, doped-iron-oxide-based redox pair materials synthesized via several routes (solid-state synthesis, combustion synthesis and aerosol spray pyrolysis) have been proven capable of splitting water at relatively low temperatures (800 °C) with water conversions up to 80 % and hydrogen being the only reaction product. This is the case for

the redox pair materials both as powders as well as coatings on a ceramic support.

Based on these results, the next efforts involve the execution and optimization of a test program for continuous water splitting and redox material regeneration in the solar reactor/receiver. The range of working conditions will be explored and the over-all reactor performance will be evaluated, before a final technical and economic evaluation of a solar plant for the production of solar Hydrogen.

TECHNICAL EVALUATION

A preliminary evaluation of technical and economic aspects of the process developed was done by a design study of a plant for solar water splitting in a commercial scale of hydrogen production by the 2-step "HYDROSOL" process.

Firstly a set-up of a plant concept was defined. A set of flow-sheets for the (chemical) part of the process was generated. From these point simulations were carried out to generation and selection of experimental and simulation data providing a basis for the design study

1) Plant concept

The set-up of a plant concept intensively corresponds with the design of a continuously operating reactor serving as a model reactor in commercial scale. The decision was taken in favor of reactor concept with modular set-up and with fixed position. Regarding commercial scale the solar heating of the process cannot be done by a solar furnace, which is a research instrument.

The temperature level of up to 1200°C in combination with a mass production of hydrogen can only be realized by using a solar tower system, a so called central receiver system consisting of a set of tracked mirrors (heliostat field), a tower, and a receiver. A sketch of the faceted receiver on the solar tower is shown in Fig. 9, and a simulation of the heliostat positions with respect to the tower in Fig. 10.



Figure 9: Design of a facetted receiver on a solar tower

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Figure 10: Simulation of heliostat field. Heliostat positions relating to tower (25 MW_{th})

2) Flow sheets

Two versions of the flow sheet have been generated and used for the lay-out of solar model plants. The first one is based on an equal temperature level for both process steps, the second one on alternating temperature levels of 800°C and 1200°C. Fig. 11 provides the basic flow sheet for the process applying equal temperature levels for both steps.



(1) condenser

(2) receiver reactor

(3) valve(4) heat exchanger (2 stages)

- (5) valve
- (6) steam generator

(7) heat exchanger (2 stages)

Figure 11: Basic flow-sheet of the water splitting process

3) Experimental data as input

The experimental campaigns have been evaluated not only regarding the feasibility of water splitting and oxide regeneration but also to generate a data base as a starting point for the modeling of a scale-up of the process. The following issues are the most important for that purpose.

The thermal behavior of the mini-plant including receiverreactor and all peripherals operated in the solar furnace in Cologne has been intensively analyzed by experiments. Beyond that, calculations have been applied for parameters not directly accessible by measurements. Important parameters for a technical and economic evaluation are thermal losses and net power available for the reactions. The major contribution to losses and power consumption are caused by the high temperature endothermic step of catalyst regeneration. Therefore the thermal balancing of the reactor is exemplarily shown for an absorber temperature of 1200°C and nitrogen as feed gas. The balancing was done at or near thermal equilibrium of the reactor. There are three contributions to the thermal losses of the mini-plant:

- Convective losses at the housing,
- Re-radiation losses trough the window,
- Heat transfer to the product/exhaust gas.

The third contribution will be recovered to a far extend in an optimized or commercial process. The experiments and calculations provided the results shown in Table 2.

1 able 2. Thermal balancing of mini-plant at 1 _{Absorber} – 1200	Table 2: Thermal bala	ncing of mini-pla	ant at $T_{Absorber} =$	1200 °	°C
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Contribution	Minimum [kW]	Maximum [kW]
Convection at reactor housing derived from surface temoperature distribution of the housing	0,7	1,0
Re-radiation measured by black- body absorber in front of the reactor aperture	2,0	2,5
Heat in exhaust/product gas =c _p (product gas)*T(product gas) - c _p (feed gas) * T(feed gas)	0,8	1,6
Overall losses without heat recovery	3,5	5,1
Overall losses with heat recovery from product gas (η = 70 %)	2,9	4,6
Solar input power from solar furnace	25,0	25,0
Max. available solar power for chemical reaction	20,1	21,4

Therefore, solar energy input beyond 3,5 and 5,1 kW respectively, is available to run the regeneration reaction in the reactor. Thus an upper limit of 80 % of the incoming radiation may be used for that reaction. Additionally 70 % solar field efficiency has to be considered and 0,35 kW (solar) are needed for the regeneration of 1 Nm³/h of nitrogen contaminated by oxygen This leads to a rough estimation of an overall efficiency of up to 40 % (ratio of higher heating value of produced hydrogen and solar input). The value has to be refined by determining potential receiver efficiencies by experiments, in

particular using a continuous operating reactor, by analyzing all unit operation represented by the process flow sheet, and finally by annual balancing of the operation of an optimized model plant.

ECONOMIC EVALUATION

Starting with these basic considerations a first calculation was performed with the following considerations for a solar plant (Fig. 12):

- solar only operation
- sited in Almería, Spain
- regeneration temperature: 1200 °C, splitting temperature 800 °C
- yield of hydrogen with respect to steam injected: 35 %
- 7-fold surplus of nitrogen needed for the regeneration
- size of heliostat field (90000 m² north field)
- 150 Euro per m² of mirror area installed
- losses of one percent of flushing gas during recycling
- annual production: 65,5 GWh of hydrogen (LHV)
- 1500 Euro per kg of metal oxide (coated)



Figure 12: Scheme of plant concept with integrated simple flow sheet.

Using these boundary conditions the investment costs and operating costs of such a facility have been calculated (Figs. 13, 14). The duration of depreciation is 10 years; the interest rate is 6 %. On this basis the production cost for hydrogen are 0,18 Euro/kWh (LHV), accordingly.



Figure 13: Investment costs

Sensitivity analyses were performed indicating that major reduction of the production costs is possible and likely due to the following parameters:

- reduction of price of the metal oxide
- enhancement of its reaction characteristics
- operation at temperatures lower than 1200 °C
- reduction of the necessary amount and purity of nitrogen
- potential co-generation of hydrogen and electric power



Figure 14: Operational costs

CONCLUSIONS

A solar reactor based on ceramic refractory honeycomb structures coated with active redox materials, was constructed. Its capability of achieving a uniform temperature profile of 1200 °C was demonstrated. Solar thermo-chemical splitting of water and hydrogen production was successfully demonstrated several times. The release of oxygen of the iron-oxide-based coating was realized by using solar energy. Multi cycling of the process is possible in principle. The amounts of hydrogen produced correspond very well to the laboratory experiments with the oxides in the form of powders. The redox pair materials as coatings are almost as active as the powders.

The analyses indicate that there is potential to reduce the production cost of hydrogen down to 10-12 Eurocent/kWh. Reduction of cost for the installation of the solar part of the plant which is expected to proceed with the ongoing

commercialization of solar thermal applications (in particular power plants) may provide even lower cost.

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