Ir(I)-NHC-phosphine complex catalyzed hydrogen generation and storage in aqueous formate/bicarbonate solutions with use of a flow reactor dynamic response to changes in hydrogen demand[¶]

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[¶]Dedicated to Professor Gábor Laurenczy on the occasion of his 65th birthday in recognition of his numerous achievements in coordination chemistry and homogeneous catalysis with particular emphasis on chemical storage and delivery of molecular hydrogen.

Abstract

Hydrogen gas generation with dynamic response to changing application demands was achieved with use of a hydrogen battery based on aqueous cesium bicarbonate hydrogenation/ formate dehydrogenation, homogeneously catalyzed by an Ir(I)-N-heterocyclic carbene complex. In this device the storage solution was circulated through a small volume tubular reactor heated to the required high temperature to allow fast hydrogen evolution while the high volume reservoir was kept at ambient temperature at which no H_2 was generated. By simple control of the reactor temperature it was possible to regulate the rate of hydrogen evolution as required. The results also demonstrate the applicability of homogeneous catalysis for hydrogen generation in flow systems.

Keywords

Bicarbonate hydrogenation; Formate dehydrogenation; Hydrogen battery; Iridium; N-Heterocyclic carbene

1. Introduction

Harnessing the Sun by photovoltaic devices, wind turbines or by other means leads to fluctuating energy production which in turn highlights the need for temporary storage of energy.[1] Hydrogen is considered a suitable energy carrier for use in industry, transportation and in everyday life ("hydrogen economy"), and its production using renewable energy could mitigate the fluctuations in available energy. In this case, the original problem of energy storage translates to the question of reversible hydrogen storage.[2] Among other possibilities, this can be achieved by reversible hydrogenation/dehydrogenation of suitable chemicals; processes which usually need catalysts.[3,4] Devices, in which the hydrogenation and dehydrogenation cycles (H₂ loading or delivery) are achieved simply by changing the hydrogen pressure can be termed hydrogen batteries.[5-7]

A widely studied way of hydrogen storage is the reversible formation and decomposition of formic acid (FA) and several highly active catalysts [8], both homogeneous [5-7, 9-18] and heterogeneous [19-21], have been discovered for decomposition of FA. In contrast, the catalytic hydrogenation of gaseous CO_2 to yield liquid FA is thermodynamically disfavoured.[22] Reaction of dissolved H₂ and CO₂ leads to high conversions to FA only in the presence of basic additives, [23-27] such as various amines, although a few processes are known leading to low concentrations of dissolved FA.[28-30] For this reason hydrogen storage devices based on reversible CO_2 hydrogenation/FA dehydrogenation can be constructed only with use of bases, of which the various ethanolamines, widely applied for CO_2 -scrubbing from gases have received special attention.[31]

Although our early attempts [32] to achieve catalytic hydrogenation of CO₂ in aqueous media remained inconclusive, in later studies [33] bicarbonate was succesfully hydrogenated to formate in aqueous solutions. We also demonstrated that the processes of catalytic hydrogenation of bicarbonate and dehydrogenation of formate (as alkali metal salts) can be coupled, Eq. (1), to give working hydrogen batteries. The water-soluble complexes [{RuCl₂(*m*tppms-Na)₂}₂] [5] and [Ir(cod)(emim)(*m*tppms)] [6,34] in the presence of *m*tppms-Na and *m*tppts-Na₃, respectively (*m*tppms-Na = sodium-diphenylphosphinobenzene-3-sulfonate, *m*tppts-Na₃ = tris(meta-sulfonatophenyl)phosphine trisodium salt, cod = 1,5-cyclooctadiene, emim = 1-ethyl-3-methylimidazol-2-ylidene) were identified as active catalysts for both hydrogenation and dehydrogenation. Consequently, loading and delivery of H₂ (the charge-discharge process of the battery) could be simply regulated by the hydrogen pressure. An important feature is that in addition to the catalyst the aqueous reaction mixture

contains only inorganic salts with no other components, so the long-term operation of the battery depends only on the chemical stability of the catalyst.

In homogeneous catalysis the substrates and the catalysts reside in the same phase. In chemical hydrogen storage this can lead to unwanted H₂-generation when the battery is not in use. Furthermore, in case the reactions (either hydrogenation or dehydrogenation, or both) require elevated temperatures it is impractical to heat the entire storage solution (possibly of large volume). In the following we present a flow reaction system for reversible hydrogen storage/delivery based on the aqueous bicarbonate/formate equilibrium, Eq. (1) in which all the difficulties mentioned above are circumvented.

2. Materials and methods

[Ir(cod)(emim)(*m*tppms)] [35], monosulfonated and trisulfonated triphenylphosphine, *m*tppms-Na [36], and *m*tppts-Na₃ [37], respectively, were synthetized by published procedures. CsHCO₃ and CsHCO₂ were purchased from Alfa Aesar GmbH. Ion-exchanged water ($S \le 1 \mu S$) was used throughout. All other reagents were high purity commercial products.

Both bicarbonate hydrogenations and formate dehydrogenations were carried out by using a H-Cube microfluidic hydrogenation flow reactor [38] (ThalesNano Nanotechnology Inc, Budapest, Hungary). Aqueous reaction mixtures containing the dissolved catalyst were pumped through the empty reactor heated to the appropriate reaction temperature. Volumes of the generated H₂ were measured with the use of a thermostated gas-burette kept at the appropriate $T \pm 0.1$ °C temperature by using a Haake K10 circulator.

Formate concentrations in the reaction mixtures were determined by HPLC (AGILENT 1220 INFINITY, Supelcogel 610H column, sample volume 20 μ l, eluent 0.1% H₃PO₄ in water, flow rate 1 mL/min). Formate was detected at λ =210 nm and its concentration was determined by integration of peak areas using a calibration curve.

General procedure for homogeneous catalytic *dehydrogenation of aqueous Csformate.* Aqueous reaction mixtures containing the catalyst ([Ir(cod)(emim)(*m*tppms)]) together with excess phosphine (*m*tppts-Na₃) and CsHCO₂ were prepared under argon in a Schlenk-tube (the reservoir) and pumped through the heated empty tubular reactor (CatCart, dimensions 30 (l) × 4 mm (i.d.)) of the H-Cube device. No H₂ was mixed into the flow. Flow rate was varied between 0.2-2.0 mL×min⁻¹. The reaction was studied in two ways. According to Method **A** (single pass) the reaction mixture was sampled and analyzed directly after leaving the reactor. According to Method **B** (cumulative), the reaction mixture leaving the CatCart was continuously fed back to the stirred reservoir and the formate concentration of the resulting solution in the reservoir was then determined over time by HPLC. Volumes of the evolved hydrogen were measured with use of a thermostated gas-burette connected to the reservoir (see Figure 2).

General procedure for homogeneous catalytic *hydrogenation of Cs-bicarbonate*. Aqueous reaction mixtures containing the catalyst ([Ir(cod)(emim)(*m*tppms)]) together with excess phosphine (*m*tppts-Na₃) and Cs-bicarbonate were prepared in a Schlenk-tube (the reservoir) and pumped through the heated empty tubular reactor (CatCart, dimensions $30 (1) \times 4 \text{ mm} (i.d.)$). Hydrogen, generated by the device electrolytically, was mixed into the liquid flow before entering the tubular reactor, where the hydrogen pressure was regulated in the range of 1-90 bar, and the temperature within 25-100 °C. Flow rate was varied between 0.5-2.0 mL×min⁻¹. Hydrogenation of CsHCO₃ was investigated using both Method **A** (single pass) and Method **B** (cumulative) – see previous paragraph. Composition of the reaction mixtures (formate concentration) were followed by HPLC.

3. Results and discussion

Hydrogenation of cesium bicarbonate and dehydrogenation of cesium formate (Equation 1) were studied in a microfluidic hydrogenation reactor (H-Cube [38]). [Ir(cod)(emim)(*m*tppms)] + *m*tppts-Na₃ was applied as catalyst.[6,34] Cesium salts were chosen due to their high aqueous solubilities [39]: CsHCO₃ – 209 g/100 g H₂O (15°C), CsHCO₂ – 450 g/100 g H₂O (20°C) and outstanding reactivities.[40-41] The catalyst was dissolved in the aqueous solutions of CsHCO₃ or CsHCO₂ and the reaction mixture was pumped through a tubular reactor held at a controlled temperature. The reactions could be conveniently followed in the 60-100°C temperature range.

$$HCO_3^- + H_2 \rightleftharpoons HCO_2^- + H_2O \tag{1}$$

Hydrogenation of cesium bicarbonate and dehydrogenation of cesium formate were investigated at various temperatures, pressures, and flow rates, using solutions of various substrate concentrations (Figures A.1-A.7). In both reactions the conversions increased with increasing temperature (Figures A.1, A.3); remarkably, no formate dehydrogenation was observed at 25°C. The conversion showed inverse dependence on flow rate (Figures A.4, A.5). At low substrate concentrations conversions as high as approximately 20% could be observed in a single pass of the reaction mixture through the heated tubular reactor for both

hydrogenation of bicarbonate and dehydrogenation of formate (see e.g. Figure A.3). Note that the catalyst was able to handle even high substrate concentrations, yielding very high TO values (e.g. $TOF = 57800 h^{-1}$ for formate dehydrogenation at $[CsHCO_2]$:[Ir]=5000:1, Figure A.2). These observations are in agreement with our earlier findings obtained in batch reactions.[6] Furthermore, it was also found earlier [6] that the catalyst was stable under 100 bar H₂ pressure at room temperature for more than 71 days. Together with the results of the present measurements using a flow reactor, all the data show that aqueous CsHCO₃^{-/}CsHCO₂⁻ solutions and the [Ir(cod)(emim)(mtppms)] catalyst have sufficiently chemical stability and may serve as basis for reversible hydrogen storage.

Figure 1 shows the results of an experiment when the reaction mixture was continuously circulated through the heated hydrogenation reactor. The solution in the reservoir was sampled over time and analyzed by HPLC (Method **B**). Under the conditions of Figure 1, bicarbonate was gradually hydrogenated to formate and the conversion reached 85 % in 200 min. At this point the H₂ pressure was decreased from 90 to 1 bar and the solution was further circulated at the same temperature (100 °C) with unchanged flow rate. The pressure drop resulted in formate dehydrogenation according to Eq. (1) with approximately 50% conversion in 100 min.



Figure 1. Hydrogenation of HCO_3^- to formate (•; 0-200 min) and dehydrogenation of HCO_2^- to bicarbonate (•; 200-300 min) under flow conditions. [CsHCO₃]/[Ir] = 75 (•,•); [*m*tppts-Na₃]/[Ir] = 2; V(H₂O) = 25 mL; *P*(H₂) = 90 bar (0-200

 $[CsHCO_3]/[Ir] = 75 (\circ, \diamond); [mtppts-Na_3]/[Ir] = 2; V(H_2O) = 25 mL; P(H_2) = 90 bar (0-200 min) or 1 bar (200-300 min), <math>T = 100 \circ C; v = 1.0 mL \times min^{-1}.$

In other experiments the volume of the evolved H₂ was measured by using a gas-burette. The experimental setup is shown schematically on Figure 2 (in this case the burette and reservoir were kept at the same constant temperature). Note that the volume of the reservoir (at T = 25 °C) is about 66 times the volume of the heated flow reactor.



Figure 2. Schematic figure of the coupled flow reactor - gas-burette system



Figure 3. Hydrogen evolution in subsequent high temperature (100 °C; H) and low temperature (25 °C; L) intervals during temperature controlled dehydrogenation of aqueous Cs-formate.

 $[CsHCO_2] = 0.081 \text{ M}; [Ir-NHC] = 0.1 \text{ mM}; [mtppts-Na_3]/[Ir] = 2; V(H_2O) = 25 \text{ mL};$ v = 2.0 mL×min⁻¹, T = 100 °C (**H**) or 25 °C (**L**). Figure 3 shows the effect of temperature on the dehydrogenation of aqueous cesium formate. It can be seen that the reaction proceeds with high rate in the reactor set at 100 °C. However, hydrogen evolution stops immediately when the reactor temperature is switched to 25 °C. Apparently, the heat capacity of the entering cold reaction mixture is sufficient to cool the reactor in a few seconds. By switching the reactor temperature back and forth between 100 °C and 25 °C, generation of gaseous H₂ can be initiated and stopped at will. Such a possibility provided by flow systems may have high importance in the case of mobile devices (e.g. vehicles) where hydrogen supply must be dynamically adjusted to the feed requirements. It was also observed, that the temperature of the reaction mixture in the reservoir did not increase significantly even in the absence of temperature control; air cooling of the solution travelling through the capillary tube connection between the reactor and reservoir was efficient enough to keep the reservoir at room temperature.

Conclusions

A reversible hydrogen storage reaction system, based on hydrogenation of aqueous cesium bicarbonate and dehydrogenation of cesium formate both homogeneously catalyzed with the water-soluble [Ir(cod)(emim)(mtppms)] was analyzed in a flow reactor (H-Cube). Fast switches of the reactor temperature between high (e.g. 100 °C) and low (e.g. 25 °C) values allow precise and dynamic adjustment of the H₂ generation rate to application demand. Furthermore, depending on the relative volumes of storage solution and the flow reactor, only a fraction of the reaction mixture has to be heated avoiding undesired H₂ evolution. Until now, soluble complex catalysts were used only in heterogenized form on solid supports but mostly failed in the long run due to leaching of the anchored complexes. This work also demonstrates the practical feasibility of using homogeneous catalysis for hydrogen generation in flow systems provided it meets the following requirements: a) the catalytic H₂-storage reaction has sufficiently high activation energy to allow fast H₂-evolution at high temperatures with no or negligible H₂-formation at ambient temperature; b) all reaction partners of the storage reaction as well as the dissolved catalyst have high chemical stability, and c) the catalyst is able to catalyze the storage reaction (such as e.g. Eq. 1) in both directions.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at

Declarations of interest: none

Notes and references

[1] A. F. Dalebrook, W. Gan, M. Grasemann, S. Moret, G. Laurenczy, Chem. Commun.
2013, 49, 8735-8751; http://dx.doi.org/10.1039/C3CC43836H

Hydrogen storage: beyond conventional methods.

[2] P. Nikolaidis, A. Poullikkas, Renew. Sustain. Energy Rev. 2017, 67, 597-611; https://doi.org/10.1016/j.rser.2016.09.044

A comparative overview of hydrogen production processes.

[3] A. Boruane, M. Elanany, T. V. Pham, S. P. Katikaneni, Int. J. Hydrogen Energy 2016,
41, 23075-23091; https://doi.org/10.1016/j.ijhydene.2016.07.167

An overview of organic liquid phase hydrogen carriers.

[4] D. Teichmann, W. Arlt, P. Wasserscheid, R. Freymann, Energy Environ. Sci. 2011, 4, 2767-2773; http://dx.doi.org/10.1039/C1EE01454D

A future energy supply based on Liquid Organic Hydrogen Carriers (LOHC).

 [5] G. Papp, J. Csorba, G. Laurenczy, F. Joó, Angew. Chem. Int. Ed. 2011, 50, 10433-10435; https://doi.org/10.1002/anie.201104951

A Charge/Discharge Device for Chemical Hydrogen Storage and Generation.

[6] H. Horváth, G. Papp, R. Szabolcsi, A. Kathó, F. Joó, ChemSusChem 2015, 8, 3036 3038; https://doi.org/10.1002/cssc.201500808

Water-Soluble Iridium-NHC-Phosphine Complexes as Catalysts for Chemical Hydrogen Batteries Based on Formate.

[7] S.-F-Hsu, S. Rommel, P. Eversfield, K. Muller, E. Klemm, W. R. Thiel, B. Plietker, Angew. Chem. Int. Ed. 2014, 53, 7074-7078; https://doi.org/10.1002/anie.201310972

A Rechargeable Hydrogen Battery Based on Ru Catalysis.

[8] X. Wang, Q. Meng, L. Gao, Z. Jin, J. Ge, C. Liu, W. Xing, Int. J. Hydrogen Energy 2018, 43, 7055-7071; https://doi.org/10.1016/j.ijhydene.2018.02.146

Recent progress in hydrogen production from formic acid decomposition.

[9] N. Onishi, M. Iguchi, X. Yang, R. Kanega, H. Kawanami, Q. Xu, Y. Himeda, Adv. Energy Mater. 2018, *in press...*; <u>https://doi.org/10.1002/aenm.201801275</u>

Development of Effective Catalysts for Hydrogen Storage Technology Using Formic Acid.

[10] H. Kawanami, Y. Himeda, G. Laurenczy, Adv. Inorg. Chem. 2017, 70, 395-427; https://doi.org/10.1016/bs.adioch.2017.04.002

Formic Acid as a Hydrogen Carrier for Fuel Cells Toward a Sustainable Energy System.

[11] D. Mellmann, P. Sponholz, H. Junge, M. Beller, Chem. Soc. Rev. 2016, 45, 3954-3988; http://dx.doi.org/10.1039/C5CS00618J

Formic acid as a hydrogen storage material – development of homogeneous catalysts for selective hydrogen release.

[12] C. Fellay, P. J. Dyson, G. Laurenczy, Angew. Chem. Int. Ed. 2008, 47, 3966-3968; https://doi.org/10.1002/anie.200800320

A Viable Hydrogen-Storage System Based On Selective Formic Acid Decomposition with a Ruthenium Catalyst.

[13] K. Sordakis, C. Tang, L. K. Vogt, H. Junge, P. J. Dyson, M. Beller, G. Laurenczy, Chem. Rev. 2018, 118, 372-433; https://doi.org/10.1021/acs.chemrev.7b00182

Homogeneous Catalysis for Sustainable Hydrogen Storage in Formic Acid and Alcohols.

[14] J. F. Hull, Y. Himeda, W. H. Wang, B. Hashiguchi, R. Periana, D. J. Szalda, J. T. Muckerman, E. R. Fujita, Nat. Chem. 2012, 4, 383-388; https://doi.org/10.1038/nchem.1295

Reversible hydrogen storage using CO_2 and a proton-switchable iridium catalyst in aqueous media under mild temperatures and pressures.

[15] W. H. Wang, M. Z. Ertem, S. Xu, N. Onishi, Y. Manaka, Y. Suna, H. Kambayashi, J.
T. Muckerman, E. Fujita, Y. Himeda, ACS Catal. 2015, 5, 5496-5504; https://dx.doi.org/ 10.1021/acscatal.5b01090

Highly Robust Hydrogen Generation by Bioinspired Ir Complexes for Dehydrogenation of Formic Acid in Water: Experimental and Theoretical Mechanistic Investigations at Different pH.

[16] S.-M. Lu, Z. Wang, J. Wang, J. Li, C. Li, Green Chem. 2018, 20, 1835-1840; http://dx.doi.org/10.1039/C8GC00495A

Hydrogen generation from formic acid decomposition on a highly efficient iridium catalyst bearing a diaminoglyoxime ligand.

[17] G. A. Filonenko, R. van Putten, E. N. Schulpen, M. E. J. Hensen, E. A. Pidko, ChemCatChem 2014, 6, 1526-1530; https://doi.org/10.1002/cctc.201402119

Highly Efficient Reversible Hydrogenation of Carbon Dioxide to Formates Using a Ruthenium PNP-Pincer Catalyst.

[18] G. Papp, G. Ölveti, H. Horváth, Á. Kathó, F. Joó, Dalton Trans. 2016, 45, 1451614519; http://dx.doi.org/10.1039/C6DT01695B

Highly efficient dehydrogenation of formic acid in aqueous solution catalysed by an easily available water-soluble iridium(III) dihydride.

[19] X. Yang, P. Pachfule, Y. Chen, N. Tsumori, Q. Xu, Chem. Commun. 2016, 52, 4171 4174; http://dx.doi.org/10.1039/C5CC10311H

Highly efficient hydrogen generation from formic acid using a reduced graphene oxide-supported AuPd nanoparticle catalyst.

 [20] S. Wesselbaum, U. Hintermair, W. Leitner, Angew. Chem. Int. Ed. 2012, 51, 8585-8588; <u>https://doi.org/10.1002/anie.201203185</u>

Continuous-Flow Hydrogenation of Carbon Dioxide to Pure Formic Acid using an Integrated scCO₂ Process with Immobilized Catalyst and Base.

[21] C. Hu S-W. Ting, J. Tsui, K-Yu Chan, Int. J. Hydrogen Energy 2012, 37, 6372-6380;
 https://doi.org/10.1016/j.ijhydene.2012.01.062

Formic acid dehydrogenation over PtRuBiOx/C catalyst for generation of CO-free hydrogen in a continuous-flow reactor.

[22] W. Leitner, Angew. Chem. Int. Ed. 1995, 34, 2207-2221; https://doi.org/10.1002/anie.199522071

Carbon Dioxide as a Raw Material: The Synthesis of Formic Acid and Its Derivatives from CO₂.

[23] C. Guan, Y.Pan, E. P. L. Ang, J. Hu, C. Yao, M.-H. Huang, H. Li, Z. Lai, K.-W. Huang, Green Chem. 2018, 20, 4201-4205; doi: 10.1039/c8gc02186d

Conversion of CO_2 from air into formate using amines and phosphorus-nitrogen $PN^3P-Ru(II)$ pincer complexes.

[24] J. Klankermayer, S. Wesselbaum, K. Beydoun, W. Leitner, Angew. Chem. Int. Ed.
 2016, 55, 7296-7343; https://doi.org/10.1002/anie.201507458

Selective Catalytic Synthesis Using the Combination of Carbon Dioxide and Hydrogen: Catalytic Chess at the Interface of Energy and Chemistry.

[25] P. G. Jessop, T. Ikariya, R. Noyori, Chem. Rev. 1995, 95, 259-272; http://dx.doi.org/10.1021/cr00034a001

Homogeneous Hydrogenation of Carbon Dioxide.

[26] P. G. Jessop, F. Joó, C.-C. Tai, Coord. Chem. Rev. 2004, 248, 2425-2442; https://doi.org/10.1016/j.ccr.2004.05.019

Recent advances in the homogeneous hydrogenation of carbon dioxide.

[27] W.-H. Wang, Y. Himeda, J. T. Muckerman, G. F. Manbeck, E. Fujita, Chem. Rev.
2015, 115, 12936-12973; http://dx.doi.org/10.1021/acs.chemrev.5b00197

 CO_2 Hydrogenation to Formate and Methanol as an Alternative to Photo- and Electrochemical CO_2 Reduction.

[28] S. Moret, P. J. Dyson, G. Laurenczy, Nat. Commun. 2014, 5, 4017-4024; https://doi.org/10.1038/ncomms5017

Direct synthesis of formic acid from carbon dioxide by hydrogenation in acidic media.

[29] K. Rohmann, J. Kothe, M. W. Haenel, U. Englert, M. Hölscher, W. Leitner, Angew. Chem. Int. Ed. 2016, 55, 8966-8969; https://doi.org/10.1002/anie.201603878

Hydrogenation of CO₂ to Formic Acid with a Highly Active Ruthenium Acriphos Complex in DMSO and DMSO/Water.

[30] G. Zhao, F. Joó, Catal. Commun. 2011, 14, 74–76; doi:10.1016/j.catcom.2011.07.017Free formic acid by hydrogenation of carbon dioxide in sodium formate solution.

[31] M. Scott, B. B. Molinos, C. Westhues, G. Franciò, W. Leitner, ChemSusChem 2017,

10, 1085-1093; https://doi.org/10.1002/cssc.201601814

Aqueous Biphasic Systems for the Synthesis of Formates by Catalytic CO2 Hydrogenation: Integrated Reaction and Catalyst Separation for CO₂-Scrubbing Solutions.

[32] F. Joó, M. T. Beck, React. Kinet. Catal. Lett. 1975, 2, 257-263; https://doi.org/10.1007/BF02068199

Formation and catalytic properties of water-soluble phosphine complexes.

[33] F. Joó, G. Laurenczy, L. Nádasdi, J. Elek, Chem. Commun. 1999, 971-972; http://dx.doi.org/10.1039/A902368B

Homogeneous hydrogenation of aqueous hydrogen carbonate to formate under exceedingly mild conditions - a novel possibility of carbon dioxide activation.

[34] H. Horváth, G. Papp, Á. Kathó, F. Joó, Hung. Pat. Appl. HU1300539; WO 2015/040440A2, 2013.

[35] H. Horváth, Á. Kathó, A. Udvardy, G. Papp, D. Szikszai, F. Joó, Organometallics2014, 33, 6330-6340; http://dx.doi.org/10.1021/om5006148

New Water-Soluble Iridium(I)–N-Heterocyclic Carbene–Tertiary Phosphine Mixed-Ligand Complexes as Catalysts of Hydrogenation and Redox Isomerization.

[36] F. Joó, J. Kovács, Á. Kathó, A. C. Bényei, T. Decuir, D. J. Darensbourg, A. Miedaner,D. L. Dubois, Inorg. Synth. 1998, 32, 1-8; https://doi.org/10.1002/9780470132630.ch1

(Meta-Sulfonatophenyl)Diphenylphosphine, Sodium Salt and its Complexes with Rhodium(I), Ruthenium(II), Iridium(I).

[37] W. A. Herrmann, C. W. Kohlpaintner, B. E. Hanson, X. Kang, Inorg. Synth. 1998, 32,
8-25; https://doi.org/10.1002/9780470132630.ch2

Syntheses of Water-Soluble Phosphines and their Transition Metal Complexes.

[38] www.thalesnano.com

- [39] R. C. Weast, Handbook of Chemistry and Physics, CRC, Boca Raton, FL, 1980
- [40] G. Papp, M. Purgel, F. Joó, 12th Int. Conf. CO₂ Utilization (ICCDU-12), June 23-27,
- 2013, Alexandria, VA, USA, Book of Abstracts P143. Storage of H₂ as formate in aqueous solutions.

[41] K. Sordakis, A. F. Dalebrook, G. Laurenczy, ChemCatChem 2015, 7, 2332-2339; https://doi.org/10.1002/cctc.201500359

A Viable Hydrogen Storage and Release System Based on Cesium Formate and Bicarbonate Salts: Mechanistic Insights into the Hydrogen Release Step.