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<td><strong>Author(s)</strong></td>
<td>Nagle, Lorraine C.; Rohan, James F.</td>
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<td><strong>Publication date</strong></td>
<td>2011-08</td>
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| **Original citation** | Nagle, Lorraine C., Rohan, James F. (2011)  
  'Nanoporous gold anode catalyst for direct borohydride fuel cell'.  
| **Type of publication** | Article (peer-reviewed) |
| **Link to publisher's version** |  
  [http://dx.doi.org/10.1016/j.ijhydene.2010.09.077](http://dx.doi.org/10.1016/j.ijhydene.2010.09.077) |
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  A definitive version was subsequently published in International Journal of Hydrogen Energy, [Volume 36, Issue 16, August 2011]  
  DOI: [http://dx.doi.org/10.1016/j.ijhydene.2010.09.077](http://dx.doi.org/10.1016/j.ijhydene.2010.09.077) |
| **Item downloaded from** | [http://hdl.handle.net/10468/1072](http://hdl.handle.net/10468/1072) |

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Nanoporous gold anode catalyst for direct borohydride fuel cell

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Abstract
Nanoporous gold (NPG) electrodes were fabricated in film and wire array formats by selectively dealloying Ag from Au$_{0.18}$Ag$_{0.82}$. Borohydride oxidation reaction (BOR) was studied by cyclic voltammetry at the NPG electrodes. The onset potential for the oxidation at a NPG wire array shifted to more negative potentials than observed at a Au disc and higher currents were realised. An onset potential of -1.07 V vs. Ag/AgCl which is 0.207 V lower than that at a Au disc was recorded. The oxidation current for 20 mM borohydride in 1 M NaOH increased to 73.6 mA cm$^{-2}$ from 3.17 at a Au disc. A value of 7.49 electrons was determined for n out of a possible 8 for borohydride oxidation. NPG presents an attractive alternative to gold nanoparticle-based catalysts for use in direct borohydride fuel cells. NPG can establish intimate contact with an electrical substrate and eliminates the need for a carbon support.

Keywords
Borohydride
Gold
Nanoporous gold
Direct Borohydride fuel cell

1. Introduction
Direct Borohydride Fuel Cells (DBFCs) are currently under investigation in an effort to develop sustainable power sources. Their target specification of 600–1000 Wh kg$^{-1}$ is 5-6 times the energy density of lithium-ion batteries. The DBFC system was first proposed in the 1960s: Indig and Snyder reported a practical demonstration of direct electricity generation from borohydride ions [1]. Research on DBFC technology stagnated then until the late 1990s. Amendola et al. reported on the performance of a DBFC system using Au–Pt alloy electroplated on carbon cloth as the anode [2]. Following further impressive research efforts over the next 5 years the first demonstration of a DBFC system for laptop computers was presented in 2005 by the Materials and Energy Research Institute (MERIT) who succeeded in increasing the DBFC system output from 10 to 400 W [3]. Recent vigorous research on DBFCs has greatly raised the expectation for the realization of the various benefits promised by this fuel cell system for portable applications [4]. Establishing the technology requires an understanding of the mechanism of borohydride oxidation and its associated catalytic species. The DBFC decomposes and oxidizes the fuel directly,

Cathode: $2\text{O}_2 + 4\text{H}_2\text{O} + 8\text{e}^- \rightarrow 8\text{OH}^- \quad E^0 = 0.40 \text{ V}$ (1)

Anode: $\text{NaBH}_4 + 8\text{OH}^- \rightarrow \text{NaBO}_2 + 6\text{H}_2\text{O} + 8\text{e}^- \quad E^0 = 1.24 \text{ V}$ (2)

The equilibrium voltage, $E^0 = 1.64 \text{ V}$, compares favourably with DMFC and H$_2$/O$_2$ fuel cells that have equilibrium voltages of 1.21 V and 1.24 V, respectively. Borohydride is
a promising fuel given its high theoretical specific energy, high specific capacity and good anode performance. Careful selection of the anode catalyst is crucial in minimising competing borohydride hydrolysis given by Eqns. 3 and 4 which can result in heterogeneous, non-faradaic hydrogen evolution that decreases fuel utilisation and lowers cell performance,

\[
\begin{align*}
BH_4^- + H_2O & \rightarrow BO_2^- + H_2 \quad (3) \\
BH_3OH^- + H_2O & \rightarrow BO_2^- + 3H_2 \quad (4)
\end{align*}
\]

In practice the actual anodic reaction depends on the ratio of hydroxide to borohydrate ion and is more realistically represented by Eqn. 5

\[
BH_4^- + nOH^- \rightarrow BO_2^- + (n-2)H_2O + ne + (4-0.5n)H_2 \quad (5)
\]

Earlier it was believed that the platinum group metals show poor faradaic efficiency for borohydride oxidation while gold was believed to be an effective catalyst for borohydride oxidation and non-catalytic with regard to its hydrolysis regarded. The oxidation of borohydride and its hydrolysis product, \(BH_3OH^-\) has been studied extensively at a range Au electrodes \([5\text{-}adjust]\). It was shown recently that gold does in fact show catalytic activity for hydrolysis of \(BH_4^-\) and \(BH_3OH^-\) \([9, 10]\).

The faradaic efficiency of gold for borohydride oxidation has been shown to be lower than that assumed previously \([11\text{ and above new ref}]\). Gold does in fact catalyse \(BH_4^-\) and \(BH_3OH^-\) hydrolysis to produce \(H_2\). Chatenet \([above new ref]\) showed that by coupling online MS with electrochemistry evidence of \(H_2\) formation during oxidation of \(BH_4^-\) on Au could be obtained. This originated from the partial hydrolysis of \(BH_4^-\) into \(BH_3OH^-\). They observed the onset of faradaic \(BH_4^-\) oxidation coincided with \(H_2\) evolution and this increases monotonically with electrode potential up to 0.8 V vs RHE (ca. -0.2 V SCE) until it reaches ca. 6.5–7.4 electrons per \(BH_4^-\) species for 0.1 M NaBH4 and 0.015 or 0.003 M NaBH4, respectively in the high potential region. Their proposed pathway assumes the generation of \(BH_3OH^-\) and \(H_2\) at low potential, followed by the oxidation of \(BH_3OH^-\) when these species can remain long enough at the vicinity of the electrode surface. At higher potential, the direct and complete BOR becomes more likely, yielding to a rather high faradaic efficiency. In a RDE analysis of \(BH_4^-\) oxidation Finkelstein \([11]\) showed that the electron count, \(n\), for the reaction is potential dependent. They identified two regions of mass transport limited current from -0.45 to -0.35 V and -0.20 to +0.28 V vs Ag/AgCl. Levich analysis showed that the higher potential wave yields 7 e per molecule while the lower potential wave yields 4.5 e. In our study a \(n\) value of 7.49 at the oxidation plateau centered at 0.0 V vs SCE was determined which is in agreement with the findings of Chatenet and Finkelstein \([11\text{ and above new ref}]\).
It was just recently shown that the platinum can show high catalytic activity for borohydride oxidation and low activity for its competing hydrolysis under certain conditions and may yield nearly 8 electrons per BH₄⁻ species. For smooth Pt Finkelstein et al [11] demonstrated that the direct oxidation of borohydride at low potentials (-0.8 to -0.25 V vs Ag/AgCl) yields 7 electrons and requires the presence of Pt surface hydrides. Molina Concha et Chatenet [ref 2 from Reviewer 2] showed that for carbon-supported platinum catalysts a n value of almost 8 has been obtained for a certain active catalyst layer thickness. This was explained in terms of an increased residence time of H₂ formed by hydrolysis and BOR intermediates within the layer to favour the complete BOR and/or hydrogen oxidation.


NPG is an interconnected, bicontinuous ligament network containing regions of both negative and positive curvature which topologically requires a high step density. NPG can be formed by a chemical etching process called dealloying which involves selective metal dissolution. Forty has shown that depletion gilding of a less-noble metal from Au alloys results in an open, continuous nanoporous structure composed of Au [16]. As the Ag atoms in an AgₓAuᵧ alloy are dissolved in acid, remaining Au atoms gather together in clusters that create a rough surface causing Au to evolve into a porous material. The sponge-like 3D structure is a system of interconnecting pores/tunnels in a skeleton of filaments of the metal. The filament size can range from 5-50 nm with surface areas as high as 20 m² g⁻¹ and a porosity of 70 % or higher possible. Biener et al. [17] indicated that NPG brings together two seemingly conflicting properties: high strength and high porosity. They characterized the size-dependent mechanical properties of NPG using a combination of nanoindentation, column microcompression, and molecular dynamics simulations. It was shown that NPG can be as strong as bulk Au, despite being a highly porous material and that its ligaments approach the theoretical yield strength of Au. Structurally it bears a resemblance to naturally occurring zeolites (filament size 1-2 nm and surface areas 100 m² g⁻¹) [18].

This useful but relatively unstudied form of Au most likely contains an intrinsically high step density which makes it attractive for catalysis studies. It is even more attractive because it can be formed into thin, high-conductivity foils that are easily adapted to electrocatalytic measurements. In terms of catalytic applications, NPG has at least two advantages over other catalysts or Au nanoparticles. Firstly, NPG remains active at low temperature (room temperature or even lower), unlike Pt or Pd catalysts [21 and Topics in Catal below]. Research has suggested that gold nanoparticle based catalysts can display low temperature activity and selectivity under a range of conditions relevant to automotive pollution control. [Topics in Catal below].

New ref to add

Topics in Catalysis Vols. 30/31, July 2004

The potential for use of gold in automotive pollution control technologies: a short review

G. Pattricka, E. van der Lingena, C.W. Cortib, R.J. Holliday,b,* and D.T. Thompsonb
Platinum group metals (PGMs) are currently used for both anode and cathode catalysts in most commercial proton exchange membrane fuel cells (PEM). The tendency for poisoning of platinum by CO contaminant in the hydrogen feedstock gas remains a major technical hurdle. Gold was shown to have particularly good catalytic performance for the oxidation of carbon monoxide. However, compared to research focused on the use of PGMs for automotive pollution control, the science of gold catalysis is still in its infancy.

Stable low temperature CO oxidation over unsupported NPG was shown by Xu et al. [21]. A significant CO conversion rate at -30oC was sustained for longer times that that recorded at room temperature. The room temperature deactivation originated from an increase in ligament size which resulted in pore clogging and a decrease in active surface area. Secondly, NPG exhibits good thermal stability and resistance to oxidation [19] and thus can overcome the aggregation or sintering limitations which Au nanoparticles encounter at elevated temperatures or in an oxidative environment [20]. NPG in a foam type structure has been shown to exhibit high CO oxidation activity, however, the catalytic reactions only occur at the foam surface [22]. Zeis et al. [23] demonstrated that NPG is an effective catalyst for the reduction of hydrogen peroxide to water. The reaction efficiency is sufficiently high to allow use of the material as a cathode for oxygen reduction in hydrogen PEM fuel cells, although the overall efficiency in this context is still far less efficient than that of Pt. Their results are consistent with their overall hypothesis that the central difference between NPG and bulk Au is due to the increased density of step edges in NPG over bulk Au. In summary, using nanoporous metal electrodes for fuel cell electrodes opens exciting new avenues for catalyst design.

In this paper we report on the exploitation of nanoporous gold (NPG) with high specific surface area and electrocatalytic activity to develop an anode catalyst with high catalytic activity for oxidation of borohydride and low activity for its competing hydrolysis.

2. Materials and methods

2.1 Materials used
Sodium borohydride (minimum purity 98 %), sodium carbonate (minimum purity 99%) and sodium hydroxide, NaOH (minimum purity 99%) were purchased from Sigma Aldrich and used as received. KAg(CN)₂ (minimum purity 99%) and KAu(CN)₂ (minimum purity 99%) were purchased from Johnson Matthey and used as received. Anodisc alumina circular membranes (Anodisc® 25) 2.5cm in diameter, 60 micron thick, 200 nm pore size and 10⁹ pore openings per cm² of membrane were supplied by Whatman™. These show compatibility with a wide range of solvents and acids and can be dissolved in sodium hydroxide and ammonium hydroxide solutions. The maximum temperature at which they are stable is 400°C. Deionized water of resistivity 18 MΩ cm was used to prepare all solutions. The working electrodes was 5 mm Au disc (Princeton Applied Research) supplied by Advanced Measurement Technology, U.K. These were polished with 0.5µm alumina powder obtained from Struers on a Buehler polishing cloth for 2 minutes and rinsed in deionized water. A 1 mm diameter Au counter electrode of 50 mm length was used.

2.2 Electrochemical measurements
Cyclic voltammograms (CV) were recorded with respect to a standard calomel electrode. The potential of the working electrode was controlled using a CH Instruments potentiostat model 660B with picoamp booster. All solutions were purged with nitrogen for 20 min prior to experiments in order to remove oxygen, and the experiments were performed at room temperature.

2.3 Fabrication of NPG

Ag$_x$Au$_y$ films were deposited on a 200 nm Au film on pyrex at constant potential of -1.2 V from a solution of 100 mM KAg(CN)$_2$ and 20 mM KAu(CN)$_2$ in 250 mM Na$_2$CO$_3$, pH 13 for 5000 s. EDX confirmed that an alloy composition of Au$_{0.18}$Ag$_{0.82}$ was formed. It was demonstrated by Searson et. al [24] that the morphology and porous structure of NPG depends on the composition of the Ag$_x$Au$_y$ alloy for the range 0.18 ≤ x ≥ 0.32. The alloy composition Au$_{0.18}$Ag$_{0.82}$ gives the highest surface area NPG (6.9 m$^2$ g$^{-1}$) when dealloyed in nitric acid and the shortest ligament size of 20-30 nm. The CV response for the alloy in 1 M NaOH from -0.9 to 0.6 V vs. SCE at 10 mV s$^{-1}$ (not shown here) indicated the presence of Au and Ag from the metal oxide reduction peaks seen at -0.1 V and 0.4 V, respectively. Upon immersing the Au$_{0.18}$Ag$_{0.82}$ in 30 % nitric acid for 15 min a high surface area form of Au referred to as NPG is obtained. The thickness of NPG was measured as 2 micron using SEM. The NPG film was delaminated from the underlying Au film and sandwiched in a holey Cu folding TEM grid in order to obtain the TEM image shown in Fig. 1. The pore and ligament size are 20 nm and 30 nm, respectively.

The fabrication of NPG on a Au disc was achieved using the conditions given above for the fabrication of NPG on a planar Au film. The fabrication of a NPG in a 3D wire array format was achieved by dealloying Ag$_x$Au$_y$ nanowires that were deposited in Anodisc alumina membranes with 200 nm pore size using the conditions given above for the fabrication of NPG on a planar Au film. The alumina membrane had a conducting backing layer of 350 nm Au which was sputter deposited. The alumina was dissolved by soaking it in 0.5 M NaOH for 2 hours to release the NPG wires. The SEM image shown in Fig.2 was recorded for the resulting NPG wires.

The fabrication of a segmented NPG-Au wire array was demonstrated to add structural rigidity to the NPG wire array. A 3D Au wire array was deposited from a commercial Puramet bath in Anodisc alumina membranes with a 350 nm Au conducting backing layer which was sputter deposited. The Au wires were deposited to a length of 3 micron and 200 nm in diameter. The fabrication of NPG onto the Au wires was achieved by dealloying Ag$_x$Au$_y$ wires that were then deposited on the Au wires in the Anodisc alumina membranes using the conditions given above for the fabrication of NPG on a planar Au film. The alumina was dissolved by soaking it in 0.5 M NaOH for 2 hours to release the segmented Au-NPG wires. The SEM image shown in Fig. 3 was recorded for the resulting segmented NPG-Au 3D wire array.

2.4 Characterisation of NPG

TEM images were recorded using JEOL 2000FX at an accelerating voltage of 200 kV. SEM images were recorded using Nova Nanosem 630 at an accelerating voltage of 15 kV

3. Results and discussion
3.1 Cyclic voltammetry study

Cyclic voltammograms for a Au disc, NPG on the Au disc, NPG-Au segmented wire array and NPG wire array in 1 M NaOH are shown in Figure 4. They clearly illustrate the influence of the increased surface area of the electrodes. The oxidation response for 0.02 mM NaBH₄ in 1 M NaOH at a Au disc and NPG-coated Au disc under static conditions is shown in Fig. 5. The onset of oxidation at the Au disc is -0.80 V and oxidation is sustained at the gold surface until 0.05 V when gold oxides form which are inactive for BH₄⁻ oxidation. In the subsequent reverse sweep the oxidation recommences at 0.10 V upon gold oxide reduction. The onset of BH₄⁻ oxidation at the NPG-coated Au disc has shifted to -0.97 V. The peak a1 at -0.49 V has been reported previously in the literature [6,7,9] and has been assigned to the oxidation of a species in solution as shown by Chatenet at al [9] and can be attributed to a combination of H₂ oxidation and the low potential oxidation of BH₄⁻ [9, 11] The broad oxidation wave from -0.4 to 0.25 V may be ascribed to the high-potential oxidation of BH₄⁻ [9] and has been shown to correspond to the oxidation of adsorbed BH₄⁻ species. The oxidation of BH₂(OH)₂⁻ was noted [9] to occur at 0.30 V. The oxidation of BH₄⁻ is sustained into the region of gold oxide formation indicating higher activity for the reaction than at the clean Au disc over this potential region. The BH₄⁻ oxidation current at the plateau centered at 0.0 V has increased from 3.17 at the Au disc to 38 mA cm⁻² at the Au disc with NPG deposit.

3.2 Determination of n

The influence of scan rate on the oxidation response for borohydride was studied at the NPG-coated Au disc. The positive shift of the peak potential for a1, Eₚ₂ᵃ₁, with increasing scan rate as shown in Fig. 6 is indicative of a sluggish reaction. The a1 peak current, Iₚᵃ₁ increases linearly with the square root of scan rate as shown in Figure 7, this suggests that a1 can be attributed to the oxidation of a species in solution and there is no interference from adsorption. Using the Randles-Sevcik Equation 6 and Equation 7 the number of electrons, n, associated with a1 was calculated as 4.26.

\[ I_p = 2.99 \times 10^5 \alpha^{1/2} n^{3/2} CD^{1/2} V^{1/2} \]  

(6)

\[ n\alpha = \frac{1.857 RT}{\zeta (E_p - E_{p,5})} \]  

(7)

The n value obtained indicates the direct oxidation of BH₄⁻ is incomplete at -0.49 V and its oxidation product is oxidised at more positive potentials.

The influence of rotation rate on the oxidation response NaBH₄ was studied at the NPG-coated Au disc, as is shown in Fig. 8. Levich analysis using equation 8 shown in Fig.10 of the oxidation plateau at 0.0 V revealed a diffusion-controlled reaction and the attainment of near-maximum coulombic efficiency for borohydride oxidation with n=7.49.

\[ I_{lim} = 0.62nFCD^{2/3} V^{-1/6} \Omega^{1/2} \]  

(8)

where the kinematic viscosity (ν) = 1.14×10⁻² cm² s⁻¹ and the diffusion coefficient (D) = 1.28×10⁻⁵ cm² s⁻¹.
The values used for D and ν were those recently determined by Chatenet et al (new ref to add below) for 10 mM NaBH4 in 1 M NaOH at 25°C.

New ref to add

The n values determined here support those recently determined in a RDE study of BH4− oxidation at Au by Finkelstein et al [11] where they reported that a 7-8 e− oxidation of BH4− occurs at -0.2 V vs. Ag/AgCl and that the oxidation peak at -0.52 V vs. Ag/AgCl yields 4.5 electrons per molecule with the remainder of the electrons possibly generating hydrogen.

3.3 Linear sweep voltammetry study
The linear sweep voltammograms for the oxidation of borohydride at the fabricated electrodes is compared in Fig. 10. The current for the oxidation of 20 mM borohydride at the Au disc, Au disc with NPG deposit and NPG wire array at the plateau centered at 0.0 V was measured as 3.17, 25, 38 and 73.6 mA cm⁻², respectively. The geometrical area exposed to the electrolyte was used to calculate the current density for each electrode. This equates to a 8-, 12- and 24-fold increase in current at the segmented Au-NPG wire array, Au disc with NPG deposit and NPG wire array, respectively over that recorded at the Au disc. The onset potential for borohydride oxidation shifts from -0.80 V at the Au disc to -0.87 V, -0.95 and -1.07 V at the segmented NPG-Au wire array, Au disc with NPG deposit and NPG wire array, respectively. The increased density of active sites at NPG over smooth gold may account for the lower onset potential for BOR. It was also shown by Chatenet et al that the onset for BOR is lower at carbon supported gold nanoparticles (-0.57 V vs NHE) than at bulk gold (-0.50V vs NHE).

New ref to add here:
Electrochim acta 2006
Kinetics of sodium borohydride direct oxidation and oxygen reduction in sodium hydroxide electrolyte Part I. BH4− electro-oxidation on Au and Ag catalysts
Marian Chatenet ∗, Fabrice Micoud, Ivan Roche, Eric Chainet

The electrochemically surface areas of the three NPG were estimated from the charge associated with the gold oxide reduction peak in 1 M NaOH assuming that the charge/area ratio is usually considered to be ca. 400 µC cm⁻² for a regular polycrystalline Au electrode. The real surface areas were calculated to be 610, 1009, 1240 cm² for the segmented NPG-Au 3D wire array, Au disc with NPG deposit and NPG wire array, respectively. The intrinsic electrode kinetic activity for the oxidation of 20 mM borohydride in 1M NaOH was calculated as 4.59, 7.52 and 72.1 µA cm⁻² for the segmented NPG-Au 3D wire array, Au disc with NPG deposit and NPG wire array, respectively. The intrinsic kinetic activity of the NPG electrodes cannot be compared with other gold-based electrodes from the literature as the electrochemically active surface areas for these are not given. The catalytic activity can only be compared to these electrodes based on the reported oxidation current densities based on the geometric electrode area. The reported oxidation current for 20 mM borohydride in 3M NaOH at 10 mVs⁻¹ at a gold foil, ETEK 10 wt% gold on Vulcan XC72R and gold nanoparticles supported on titanate nanotubes was 4.5,15
and 10 mA cm\(^{-2}\) [13]. The reported oxidation current for 100 mM NaBH\(_4\) in 1M NaOH at 10 mVs\(^{-1}\) at carbon-supported gold hollow nanospheres was 57 mA cm\(^{-2}\). [12].

The oxidation current for 20 mM borohydride in 1 M NaOH reported herein is 25, 38 and 73.6 mA cm\(^{-2}\) at a segmented NPG-Au wire array, Au disc with NPG deposit and NPG wire array, respectively. This comparison serves to highlight the superior activity of NPG over other gold-based electrodes for borohydride oxidation.

The primary reason that NPG is more catalytically active than bulk planar gold for BOR is that the catalytic sites are often crystal surface defects. At such sites gold atoms with lower bond coordination numbers exist which may aid adsorption of reactants via electronic or steric interactions. NPG intrinsically has a high step density and hence a greater number of active sites than bulk planar gold. Also, it is possible that the structure of the active layer at NPG promotes an increase in the residence time of BOR reaction intermediates at the surface leaving more time for complete oxidation to borate. Such behaviour was recently reported by Chatenet for Pt and Pt/C electrodes [new refs below] and by Schneider et al. [new ref below] for the (multi-step) oxygen reduction reaction at arrays of Pt nanostructures at planar glassy carbon electrodes.

New refs to insert

4. Conclusions
Borohydride oxidation was studied by cyclic voltammetry at NPG in a range of structures. The onset potential for borohydride oxidation shifted to more negative potentials than that observed at bulk Au. The onset potential shifts from -0.80 V at a Au disc to -0.87, -0.95 and -1.07 V vs. Ag/AgCl at a segmented NPG-Au wire array, Au disc with NPG deposit and NPG wire array, respectively. The oxidation current for 20 mM borohydride in 1 M NaOH increased from 3.17 at a Au disc to 25, 38 and 73.6 mA cm\(^{-2}\) at a segmented NPG-Au wire array, Au disc with NPG deposit and NPG wire array, respectively. A value of 7.49 electrons was determined for \(n\) out of a possible 8 for borohydride oxidation.

NPG presents an attractive alternative to gold nanoparticle-based catalysts for fuel cells as it does not require a carbon support thereby removing the stability issues associated with these. NPG as a thin foil may be incorporated as a porous catalyst electrode as it is shapeable and has mechanical, thermal and chemical stability coupled to high catalytic activity. It has a dual functionality in that it can act as a current collector and as a catalyst. It provides a solution to the sintering problems that plague nanoparticle based catalysts and will allow for establishing more intimate contact with an electrical substrate. The porous structure promotes mass transport of reactant to the active sites and release of gaseous by-products. The diffusion of an electroactive species to Au nanoparticles on a high surface area carbon support is limited by the low degree of porosity of the carbon support. The role of catalyst support is a topic of continuous debate and different mechanisms have been proposed. The controversy arises from the fact that the supported Au catalyst is a composite system and its catalytic activity results from an interplay of pretreatment conditions, particle size, preparation method and choice of support. By eliminating the support
effect in free-standing NPG a deeper insight into the origin of the exceptional catalytic behaviour of Au in nanostructured form can be gained.

NPG can be easily integrated into PEM fuel cells. NPG may be integrated into nafion-based MEAs in conventional PEM fuel cells. An example of a PEM fuel cell using Pt modified NPG was given in J. Power Sources 165 2007, p 65 Zeis et al. Another advantage from incorporation of NPG over platinum into fuel cells would be the useful enhancement in electrical conductivity that could be derived, due to the lower electrical resistivity of gold compared to platinum.

However, from a cost viewpoint NPG may not rival supported gold nanoparticle based catalysts due to the energy used in it’s production, the amount of silver lost in etching the precursor alloy and most importantly the significant amount of gold it constitutes.

Acknowledgement

This work is supported by EPA funding; it is part of a STRIVE fellowship entitled “Zero Carbon Emission Micro Fuel Cell Design”; Contract No. 2007-FS-ET-6-M5.

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Fig 1.
Fig. 3.
Fig. 5.
Fig. 6.
Fig. 7.

Fig. 8.
Fig. 9.
Fig. 10