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Design and development of manufacturing methods for manufacturing of PEM fuel cell MEA's

Nikhil Pramod Kulkarni

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DESIGN AND DEVELOPMENT OF MANUFACTURING METHODS FOR
MANUFACTURING OF PEM FUEL CELL MEA’S

by

NIKHIL PRAMOD KULKARNI

A THESIS

Presented to the Faculty of the Graduate School of the

MISSOURI UNIVERSITY OF SCIENCE AND TECHNOLOGY

In Partial Fulfillment of the Requirements for the Degree

MASTER OF SCIENCE IN MECHANICAL ENGINEERING

2011

Approved by

Frank W. Liou
Joseph W. Newkirk
K. Chandrashekhara
This thesis is presented in publication format and is divided into two separate papers. Pages 2 through 34 are published at the 2009 International Solid Freeform Fabrication Conference for publication in the symposium proceedings under the title, “Fuel Cell Development using Additive Manufacturing Technologies - A Review”. Pages 35 through 66 are to be submitted for publication in the Journal of Sustainable manufacturing and renewable energy, under the title, “Comparison of Direct Deposition Process and Electro-write Process for Proton Exchange Membrane Fuel cell MEA manufacturing”.

ABSTRACT

This thesis focuses on design and manufacturing of Fuel cell components using Additive Manufacturing techniques and then in the later part on design and manufacturing of Membrane Electrode Assembly (MEA) which is a very important component in fuel cells. Additive manufacturing methods are fast and efficient manufacturing methods which are additive building up components layer by layer instead of conventional subtractive manufacturing techniques. This ensures low cost and faster manufacturing. Additive manufacturing is important for fuel cell component manufacturing since it is important in fuel cells to minimize wastage and reduce the cost. MEA is the basis of the cost factor in Proton Exchange Membrane (PEM) fuel cells. It contributes for more than 50% of the cost in a fuel cell. In order to reduce the cost of a fuel cell/kW, it is necessary to achieve the maximum performance of the fuel cell using least amount of the platinum catalyst. The best way to achieve that is to achieve a uniform loading of the catalyst through the entire area of the MEA. Along with this, it is important to have an efficient and at the same time a fast manufacturing method for MEA’s. This thesis discusses two methods, namely Direct Deposition Process (DDP) and Electro-Write Process (EWP) and compares the efficiencies of the two using a novel way employing Scanning Electron Microscopy (SEM) and Energy Dispersive X-ray Spectroscopy (EDS) techniques. This thesis also focuses on the importance of cost model and efficiency measurement techniques for monitoring a manufacturing method in order to know the impact of every manufacturing method of individual components on the total cost of the product.
ACKNOWLEDGMENTS

I am greatly indebted to my advisor, Dr. F.W. Liou, whose encouragement, guidance, and support helped me throughout my research and writing of my thesis. He provided expertise, inspiration and support for this research without which my studies would not have taken place.

I would like to thank Dr. J.W. Newkirk for giving me the required guidance in my research, suggesting me novel techniques to carry out tasks and Dr. K. Chandrashekhara for serving as committee members and examining the thesis. I would sincerely like to thank all the members of the LAMP lab, especially Todd Sparks for helping me during the experiments and providing me with valuable suggestions which have been very critical in completing my research work. My sincere appreciation to Miss Shruti Mahadik from the Chemistry department for the help she extended to me for nanoparticle measurements. I would also like to thank my friends here in Rolla, who have made the past two years memorable.

The financial assistance provided to me in the form of Graduate Research Assistantship and Graduate Teaching Assistantship through the Intelligent Systems Center and Department of Mechanical and Aerospace Engineering are gratefully acknowledged.

Finally, I would like to thank my family, most especially, my parents Gauri and Pramod Kulkarni, my brother Aditya Kulkarni for their help, encouragement and motivation throughout my life without which I would not have accomplished this task. I would like to dedicate this thesis to my parents.
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1. INTRODUCTION

The development of Proton Exchange Membrane Fuel cells has been taking place since many decades but it has not been able to generate significant interest for portable fuel cell applications due to various reasons such as cost, ease of manufacturing etc.

Additive manufacturing is an exciting new arena in the field of manufacturing which intends to reduce time of manufacturing and cost due to its fundamental principle which is to build equipments by addition of materials in place of conventional subtractive manufacturing which leads to a lot of material wastage and hence higher costs. The number of materials which can be employed with Additive manufacturing is also increasing by the day, hence opening up a wide number of applications for it.

The first section presents a study of additive manufacturing processes which can be used for fuel cell manufacturing. The second section is a comparison of two such processes, namely Direct Deposition process and Electro-write process followed by the in-depth study of the process which is more efficient. This is very important since, in fuel cell manufacturing it is very important to reduce the manufacturing cost to reduce the cost of the product.

The second section also gives a very simple method of determining the efficiency of the processes using the SEM and the EDS techniques.
Fuel cells are being perceived as the future clean energy source by many
developed countries in the world. The key today for clean power is the reliance of fuel
cells not only to power automobiles but also for residential, small commercial, backup
power etc. which calls for production on a large scale. Additive manufacturing is
perceived as a way to develop cost effective fuel cells. It imparts flexibility to design
different kinds of fuel cells along with reduction in material wastage. This paper deals
with the review of additive manufacturing processes for research and development of fuel
cell components, such as synthesizing and prototyping new materials for fuel cell
components, fuel cell system design and prototyping, designing well sealed fuel cells,
bridging from fuel cell design to manufacturing tooling, etc.
1. INTRODUCTION

Fuel cells are electrochemical devices similar to batteries which convert energy from chemical state to electricity. There is an anode side and a cathode side in it. Fuel enters the cell from the anode side and oxidant flows into it from the cathode side. The reactants react inside the cell and the reaction products or the waste products flow out of it. The basic difference between batteries and a fuel cell is that fuel cell is only an energy conversion device and not energy storage device. Fuel cells consume reactant (fuel) from an external source which must be replenished. Hence, fuel cells represent a thermodynamically open system. However, batteries are both energy storage and conversion devices and hence they represent a thermodynamically closed system. The advantage of separating the storage and conversion functions is that power and energy capacity can be sized independently of each other. Also, many different fuels can be used as the primary energy source of the fuel cell setup depending on the types of fuels compatible with the type of fuel cell being employed [Spiegel 2006].

This paper primarily discusses Proton Exchange Membrane (PEM) fuel cells because most of the research regarding fuel cells has been undertaken with regards to PEM fuel cells, due to its many advantages such as versatility. PEM fuel cells can be employed for various uses starting from portable power to automotive power to stationary residential power. The by-product of a PEM fuel cell is water, which is not only non-polluting but can be used as a potable water supply.
In the PEM fuel cell, hydrogen is the fuel which enters the fuel cell through the anode end and oxygen through the cathode end. The following reactions take place at the cathode and the anode:

Anode: \( \text{H}_2 (g) \rightarrow 2\text{H}^+ (\text{aq}) + 2e^- \)

Cathode: \( \frac{1}{2} \text{O}_2 (g) + 2\text{H}^+ (\text{aq}) + 2e^- \rightarrow \text{H}_2\text{O} (l) \)

Overall Reaction: \( \text{H}_2 (g) + \frac{1}{2} \text{O}_2 (g) \rightarrow \text{H}_2\text{O} (l) \)

The components in a fuel cell are:

1. Bipolar Plates.
2. Membrane Electrolyte Assemblies (MEA’s).
3. Gas diffusion electrode layers.

Apart from this there are various auxiliary components such as gas flow pipes, the gaskets (seals), the connectors, end plates and cooling plates (required in fuel cell stack). The cost of these auxiliary components is relatively insignificant as compared to the cost of the major components. The percentage cost of components is illustrated in Table 1.

<table>
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<th>Cell Stack</th>
<th>Contribution to the Entire Cost of the Fuel Cells [DOE 2005]</th>
</tr>
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<tr>
<td>Membrane</td>
<td>35~40 %</td>
</tr>
<tr>
<td>Catalyst</td>
<td>15~20 %</td>
</tr>
<tr>
<td>Bipolar plates</td>
<td>10~15 %</td>
</tr>
<tr>
<td>MEA’s</td>
<td>30~35 %</td>
</tr>
</tbody>
</table>

MEA is the heart of the fuel cell; rather it is the distinguishing criteria for different types of fuel cells. An MEA, as the name suggests, is the assembly of the
membrane and two electrodes on either side of the membrane. An electrode is a carbon cloth which is fabricated in a particular pattern depending on the mesh size required. Also, it needs to have specific properties to facilitate proper water management throughout the cell.

As for the membrane, it is the electrolyte which is being employed for that particular fuel cell. The most common electrolyte used for a PEM fuel cell has been Nafion®. Nafion® is a generic brand name given by its developer DuPont. Its chemical name is sulfonated Polytetrafluoroethylene (PTFE). Although Nafion® is the most common polymer membrane employed in PEMFC, extensive research is being carried out to find a cost effective alternative which is as mechanically and chemically stable as Nafion® [Payne 2009].

The most common catalyst used for the PEMFC is Platinum due to its stability in highly corrosive atmospheres as well as its performance characteristics. The methods used for applying the catalyst are screen printing and hand painting. However, the uniformity of the catalyst deposited is not easily controlled. Also, these processes are time consuming, and require iterations of painting, drying and massing to achieve the required loading of the catalyst. The reproducibility of these methods is poor. There is considerable amount of catalyst wasted in the feed lines due to clogging which results in an increase in the production cost [Taylor 2007].

The aforementioned commercial methods of producing major components of fuel cells are not in accordance with the economic threshold value as required by the US Department of Energy. These processes combine costly materials and processes that result in increased costs of fabrication of fuel cells.
To achieve the target of production cost of $30/kw by 2015 as set by the US DOE [DOE 2005], there is a need to achieve low cost fabrication of fuel cells and use alternate cheaper materials in the manufacturing processes. Based on the Results of the Workshop on Manufacturing R&D for the Hydrogen Economy, several challenges confront the transformation of the U.S. manufacturing sector to support the hydrogen energy economy such as:

- Develop innovative, low-cost manufacturing technologies for new materials and material applications.
- Adapt laboratory fabrication methods to low-cost, high-volume production.

Rapid manufacturing is an innovative manufacturing technique which can be used for the fabrication of fuel cells which goes hand-in-hand with the aim of US DOE. Rapid prototyping is defined as a machine technology which is used to fabricate 3-dimensional models and prototype parts from a numerical description (typically a CAD model) using an additive approach to form physical models. That is why Rapid prototyping is also referred to as ‘Additive Manufacturing’. Additive Manufacturing (AM), as the name suggests is the process of fabrication of physical models or prototypes by addition of materials. This addition takes place layer by layer incrementally. By this process, the problems of form generation and material composition are addressed. The smaller the incremental volume of material better is the accuracy of the form generated and also the control over system parameters. AM doesn’t require any external tooling for the manufacturing of 3D freeform objects.
There are various kinds of Additive Manufacturing techniques such as:

1. Selective Laser Sintering (SLS)
2. Fused Deposition Modeling (FDM)
3. Stereolithography (SLA)
4. 3D printing
5. Laminated object manufacturing (LOM)
6. Electron Beam Melting (EBM)

Variations of these processes also exist but it is not important to be listed above since a small variation of some system parameters might lead to an entirely different process. For the manufacturing of PEM fuel cells, more importance has been given to a few processes such as Fused Deposition Modeling (FDM), 3D Printing and Selective Laser Sintering (SLS) which are discussed in detail in this paper.

The advantages of AM itself make it an attractive way to build fuel cells. With additive manufacturing technologies, you have the flexibility to change the design of the fuel cells without the need to change the entire setup as would be required with regards to conventional manufacturing technologies. In this paper, there is an example of a planar array fuel cell with a mono polar plate design. It gives a good proof of the flexibility of additive manufacturing technology. This feature of AM enhances the prospects of further cost reduction. Inkjet printing aids the process of precision manufacturing since we can deposit materials with micrometer precision thereby again reducing material waste. Impressive results from the three processes as described in this paper maximize the scope of AM for building fuel cells. It might happen that, under a single roof, we see multiple AM techniques used to build an entire fuel cell.
2. FDM (FUSED DEPOSITION MODELING)

Fused Deposition Modeling is an additive fabrication technology which constructs superior rapid prototypes from 3D CAD data where in a thermo plastic material is extruded in the form of beads layer by layer using a temperature controlled head which is actually controlled by Computer Aided Manufacturing (CAM) software [Zhong 2000].

A plastic filament supplies material to an extrusion nozzle which is heated so as to melt the material and deposit the required amount of material in horizontal and vertical directions (i.e., wherever it is necessary). The material hardens as soon as it is extruded from the nozzle.

The thermoplastic materials used in the FDM process have good stability and durability of the mechanical properties over time; they have high heat resistance and also produce smooth parts with all the finest details intact. The commonly used materials with this process are Acrylonitrile Butadiene Styrene (ABS) polymer, elastomers, investment casting wax and some of the water soluble materials are used in this process which acts as support structures during the manufacturing process [Masood 2004]. In Figure 1, the schematic of the FDM process is illustrated.
FDM has been used for the fabrication of miniature fuel cell stack in a planar array form [Chen 2008]. For the development of miniature fuel cells, it is required to have pin-point precision, since the aim of this type of fuel cell is to have high power density in a small stack. A study has been made to develop a 10 cell planar array air breathing fuel cell using FDM as the RP (Rapid Prototyping) process [Chen 2008]. Figure 2 shows the construction of the stack using components manufactured by FDM process and Figure 3 shows the layout of the 10 cells for the analysis of the configurations in series and parallel.
Figure 2 Construction of the stack [Chen 2008]

Figure 3 Layout of the stack [Chen 2008]
The alternative processes for the development of such miniature fuel cell are the Micro-Electro-Mechanical processes (MEMS) and the conventional CNC machining processes. The flow field plates were the parts which were fabricated using RP, more specifically Fused Deposition Modeling (FDM). Acrylonitrile-butadiene-styrene (ABS) was the material used for the fabrication of flow field plates since it has high mechanical strength, low cost and easy to fabricate by RP. Table 2 gives a comparison of the manufacturing time required for the geometry of flow field plates by different processes [Chen 2008].

Table 2 Comparison of different methods for fabricating the flow field geometry [Chen 2008]

<table>
<thead>
<tr>
<th>Process</th>
<th>Time (approx)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rapid Prototyping</td>
<td>1 hour</td>
</tr>
<tr>
<td>CNC</td>
<td>2 hours</td>
</tr>
<tr>
<td>MEM’s</td>
<td>12-36 hours</td>
</tr>
</tbody>
</table>

It is noted that RP is faster than the rest of the methods. Also, if the flow field plate is designed more and more complicated, CNC machining may not be possible at all. This kind of miniature PEM planar array FC stack is a first try in both academic as well as industrial world.

The FDM process used for planar array fuel cell fabrication is described next with respect to the design and performance of the PEMFC.
2.1 DESIGN AND CHARACTERISTICS OF A PLANAR ARRAY PEMFC

- There are 10-segments in the PEM. Hence the total reactive area is 17cm$^2$
- The cell operating temperature is 70°C internal and ambient temperature is 25°C.
- Two configurations have been tried: Parallel and series with natural and forced convection
- The anode is on the same side of the membrane whereas the cathode is on the opposite end or the ventilated end. Hence, it is called as a mono-polar stack design.

In Figures 4 and 5, the performance curves of the cell stack in parallel and series connection is illustrated, respectively, and Tables 3 & 4 state the performance statistics.

2.2 PERFORMANCE TEST IN PARALLEL CONNECTION

![Performance Test in Parallel Connection](image)

Figure 4. Comparison of forced and natural convection in a parallel connection [Chen 2008]
Table 3 Performance characteristics in a parallel connection [Chen 2008]

<table>
<thead>
<tr>
<th></th>
<th>Voltage</th>
<th>Current Density</th>
<th>Power Density</th>
</tr>
</thead>
<tbody>
<tr>
<td>Free convection</td>
<td>0.425 V</td>
<td>233 mA/cm²</td>
<td>99 mW/ cm²</td>
</tr>
<tr>
<td>Forced Convection</td>
<td>0.425</td>
<td>289 mA/cm²</td>
<td>123 mW/ cm²</td>
</tr>
</tbody>
</table>

2.3 PERFORMANCE TEST IN SERIES CONNECTION

Figure 5 Comparison of forced and natural convection in a series connection [Chen 2008]
Table 4 Performance characteristics in a series connection [Chen 2008]

<table>
<thead>
<tr>
<th></th>
<th>Voltage</th>
<th>Current Density</th>
<th>Power Density</th>
</tr>
</thead>
<tbody>
<tr>
<td>Free convection</td>
<td>4.25 V</td>
<td>216 mA/cm²</td>
<td>92 mW/cm²</td>
</tr>
<tr>
<td>Forced Convection</td>
<td>5.25 V</td>
<td>200 mA/cm²</td>
<td>105 mW/cm²</td>
</tr>
</tbody>
</table>

The parallel connection stack had higher power density than the serial connected stack since some cells performing badly will affect serial connection where parallel connection won’t be affected a lot. [Chen 2008]. The performance of the stack reached power density of the state of the art planar array fuel cells (100-120 mW/cm²) [Chen 2008]. Clearly, fuel cell components made by RP (FDM) instead of conventional CNC machining or more costly MEM processes do deliver performances as required which does speak about the reliability of the process. So, we infer that RP is a successful procedure in prototyping the components. In the future, we might even see RP being applied for larger scale production.
3. 3D INKJET PRINTING

3D Inkjet printing (3D IJP) is yet another form of Additive Manufacturing. What differentiates 3D printing from other forms of additive manufacturing is that it is much more affordable than other processes existing till date. Inkjet printers are plug and play devices that require little setup, training or maintenance.

Inkjet printing utilizes drop-on-demand technology to deposit various materials in a colloidal ink form. Also, there is no contact between the printer head and the substrate on which it is going to be deposited. There are two types of inkjet printers- one which use piezoelectric transducers and one which use thermal resistors to expel droplets through the nozzles. Development of inkjet printers will result in smaller nozzle sizes and hence ink droplets, which will result in higher resolution (dots per inch) as well as in printing intricate features, patterns which is advantageous in the development of fuel cell components.

Inkjet printing can be employed in printing different MEA’s since the composition is not very different from each other. Inkjet printing can be considered as an efficient method used for the deposition of catalyst layers because of the performance it gives in terms of controlled catalyst deposition for ultra loadings of Platinum which results in a better utilization of Pt as compared to conventional catalyst deposition methods like Screen Printing and Hand Painting. Inkjet printing will also help in optimizing the Pt loading which will result in reduction of costs. The reproducibility produced in the catalyst printing is incredible and this will in turn lead to lesser cell failure rates. Figure 6 shows some of the shapes of catalyst layers printed with 3D IJP.
3.1 3D INKJET PRINTING AS COMPARED TO OTHER PRINTING TECHNIQUES

IJP as compared to other printing techniques proves to be more advantageous as it allows for a uniform distribution of catalyst material onto the surface of GDL and provides picolitre precision and control of the deposition of each print and thus paves a way for ultra low loadings. IJP is also found to be reproducible due to the elimination of some of the intermediate steps of drying and massing which are two important steps in Hand painting and screen printing [Taylor 2007]. The comparison of the Hand painting method which is the conventional method for catalyst layer printing and that of 3D Inkjet Printing is shown in Figure 7.
Catalyst inks should be similar to the OEM inks as specified by the manufacturer so that the printing can be executed smoothly. These properties are specified in table 5.

Table 5 Usual properties for home/office printer inks [Towne 2007]

<table>
<thead>
<tr>
<th>Property</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Viscosity</td>
<td>1-4 cP</td>
</tr>
<tr>
<td>Surface tension</td>
<td>30-35 mN/m</td>
</tr>
<tr>
<td>Average Particle size</td>
<td>0.2 μm</td>
</tr>
</tbody>
</table>
3.2 SETUP USED FOR THE EXPERIMENT

Catalyst formation takes place by thoroughly mixing a carbon supported catalyst with Nafion® solution and de-ionized water. Water, ethylene glycol and isopropanol are added to achieve the required properties of surface tension and viscosity. The Nafion® membrane is prepared by washing in 3% wt H₂O₂ for 1 hour, rinsed and boiled in de-ionized water for 1 hour and stored in Milli-Q grade de-ionized water. The printers considered for this experiment were simply off the shelf printers whose cartridges were cleaned of the original ink and replaced with the catalyst ink with the help of a syringe. Illustrator software is used for making different size and shape electrodes and for different amounts of platinum loading by changing the hue, saturation and luminescence [Towne 2007].

3.3 RESULTS AND DISCUSSION

The characteristics of the ink prepared by the process stated earlier are tabulated in Table 6.

<table>
<thead>
<tr>
<th>Property</th>
<th>Result</th>
<th>Within range</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Particle size</td>
<td>&lt; 2µm</td>
<td>No</td>
<td>Not within range but the ink is maintaining colloidal stability.</td>
</tr>
<tr>
<td>Surface Tension</td>
<td>35.5 mN/m</td>
<td>No</td>
<td>Just a tad out of range.</td>
</tr>
<tr>
<td>Viscosity</td>
<td>3.35 cP</td>
<td>Yes</td>
<td></td>
</tr>
</tbody>
</table>
There were two attempts to print the catalyst layers. One was on a cellulose acetate substrate and the other was on Nafion® substrate. These two attempts are shown in Figures 8 & 9.

In Figure 8, the catalyst layer is seen between the epoxy layer and the substrate and its thickness is 1.02 μm thick since below the thickness of 580 nm, the resistivity of the catalyst increases tremendously which is detrimental to our interests. In Figure 9, a layer thickness up to 3.2 μm was measured. We can observe here that the thickness of the catalyst layer is not uniform and is very uneven. This happens due to swelling of the Nafion® membrane.

As expected, the Nafion® substrate swells due to the water and alcohol in the catalyst ink. Water alone can lead to swelling of the membrane by 32%. This results in
uneven printing. The cross section of the single layer catalyst section as seen in Figure 9 clearly shows the swelling of the membrane.

The next analysis presented in the paper is about the control of the deposition of the catalyst ink onto the GDL using illumination characteristics such as brightness and tint. The Figure 10 shows the optical micrographs of the same.

![Optical micrographs of printed layers taken at 15x magnification](image)

Figure 10 Optical micrographs of printed layers taken at 15x magnification; these images show evidence of banding in three samples of different thickness [Towne 2007]

The three figures above show 3 single layer inkjet printed catalysts. They have different amounts of thicknesses (drop amounts). The darker the layer, the thicker it is and hence well connected. This leads to better conductivity. It is evident from these images that inkjet printing allows excellent control over the individual layer thicknesses. Hence, many layers and ultimately thicker electrodes can be deposited.

The earlier images show that it is difficult to print the catalyst layer on a Nafion membrane, however, with some post processing, the catalyst layer can be made uniform as well as well mechanically adhered to the membrane. The usual post processing steps are hot pressing and water extraction.
Hot pressing leads to removal of ethylene glycol and also it leads to more uniform catalyst formation on the membrane. The Transmission Electron Microscopy (TEM) images are of the catalyst layers before and after processing are shown in Figures 11 & 12.

In Figure 11, the arrow represents carbon particles and depicts the discrete nature of platinum and carbon particles. This definitely affects the interconnectivity and thus the conductivity. In Figure 12, you cannot make out the separate layers of carbon and platinum particles, thus showing the continuity. Hot pressing was done at 2045 psi, 125°C for 5 min.
3.4 PERFORMANCE OF SINGLE CELLS

The testing of these catalyst printed membranes was carried out by making a single cell out of it. MEA’s had printed layers on both anode and cathode. Only 2.75% H₂ was used for initial studies. The specifications of the MEA are shown in Table 7.

Table 7 Specifications of the MEA [Towne 2007]

<table>
<thead>
<tr>
<th>Specification</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Active area of the MEA</td>
<td>2.25 cm²</td>
</tr>
<tr>
<td>Platinum loading</td>
<td>0.094 mg/cm²</td>
</tr>
<tr>
<td>Drop size</td>
<td>3 pL</td>
</tr>
</tbody>
</table>

In Figure 13, the comparison of unprocessed MEA with MEA’s processed at different pressures at 125°C is shown.

Figure 13 Graph comparing printed MEA with processed MEA’s [Towne 2007]
The maximum performance is achieved at 2045 psi giving maximum power density of 31.5 mW/cm². It was presumed that hot pressing did not remove the ethylene glycol completely. Hence water soaking was carried out on the printed MEA to remove the rest of the ethylene glycol. Water soaking led to maximum current output of 106mA/cm² at 0.401V. Thus the density of power is 42.4mW/cm².

After this initial testing, 100% H₂ was used for better comparison with the commercial MEA’s. Figure 14 describes the comparison of different MEA’s with 100% H₂ and with different treatments after the printing.

Figure 14 Power curves comparing a commercial MEA with printed MEA’s

[Towne 2007]
In Figure 14, it is clear the commercial MEA’s outperformed those inkjet printed MEA’s are having the catalyst loading of 0.094 mg Pt/cm². However, when the catalyst loading was 0.2 mg Pt/cm², it was a comparable performance as compared to commercial MEA’s, which is tabulated in Table 8.

Table 8 Comparison of the improved MEA with the commercial MEA [Towne 2007]

<table>
<thead>
<tr>
<th>Type of MEA</th>
<th>Platinum loading ( mg Pt cm²⁻² )</th>
<th>Peak power density ( mWcm²⁻² )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inkjet printed</td>
<td>0.2</td>
<td>155</td>
</tr>
<tr>
<td>Commercial</td>
<td>0.3</td>
<td>167</td>
</tr>
</tbody>
</table>

Hence, with a 33% lower catalyst loading, only 7% lower power density was obtained. This result proves that inkjet printed MEA’s can compete with the commercial ones.

Thus, it is evident how efficient can inkjet printing method for fabricating MEA’s is as compared to commercial MEA’s. The efficiency of the catalyst usage or loading can be further enhanced by grading the amount of platinum loading in every layer. Previous literatures suggest that the graded catalysts were found to perform better than the uniformly loaded catalyst in every layer [Xie 2005, Wang 2004].

Paganin et al. clearly suggests that platinum is better utilized when it is more concentrated near either the GDE layer or the electrolyte membrane layer [Paganin 1996]. Inkjet printing makes it possible to grade the platinum loading print after print. Previous research carried out by Taylor et al. demonstrated that a graded catalyst of Pt wt% 10-50 on carbon black outperformed the uniform catalyst structure of 20 % wt Pt on
carbon black at nearly the same amount of overall platinum loading [Taylor 2007]. Figure 15 shows the possible grading scheme for better performance and Figure 16 shows the performance comparison of a standard catalyst to that of a graded catalyst.
4. SELECTIVE LASER SINTERING

The functions of bipolar plate are:

- To provide electrical contact between two adjacent MEA’s.
- Uniformly distribute hydrogen gas and oxygen gas/air to the anode and the cathode side of the MEA respectively.
- To serve as a platform to support the soft MEA.
- To act as an outlet medium for heat and water vapor generated from the net reaction.

Hence, the requirements for a bipolar plate accordingly are:

- High electrical conductivity
- Plate material electrically compatible with the electrode
- Very low gas permeability for reactant gases.
- High thermal conductivity to make use of the waste heat.
- Chemical stability i.e. corrosion resistant.
- Low density plate material to keep the stack weight low.
- Inexpensive plate material.

4.1 MATERIAL SELECTION FOR BIPOLAR PLATE

There are three types of materials identified for the manufacture of fuel cell bipolar plates which are: pure graphite, metallic materials, and carbon-polymer composites.
Pure graphite, with peak conductivity of $1.44 \times 10^3$ S/cm is suitable for Bipolar Plates because they need to be highly conductive. Graphite is very difficult to machine when it comes to the machining of the flow field channels because of its flaky microstructure and irregular geometry. This also reduces its mechanical strength [Chen 2006].

Metals such as stainless steel, titanium, gold, aluminum, have good machining characteristics as compared to graphite. However, gold and titanium are very costly. Aluminum can be used with a gold coating. However, there is large difference in coefficient of thermal expansion which leads to micro-cracks in the coating. Stainless steel has corrosion issues [Maeda 2004, Chen 2006].

Composite materials suitable for the application of bipolar plates are a combination of porous graphite along with polycarbonate plastic. Graphite is an allotrope of carbon and a semimetal. The carbon based materials suitable are resins such as polyethylene, phenolic, Vinyl ester etc. with filler materials like carbon black and carbon/graphite powders. These composite systems provide electrical conductivity as well as corrosion resistance and mechanical strength [Chen 2006].

4.2 SETUP AND PROCEDURE FOR THE EXPERIMENT

There are two kinds of SLS procedures namely, Direct and Indirect. Direct SLS means parts are produced by just laser sintering of the powder without any post processing measures. Indirect SLS involved production of a porous green part held together by a certain polymer binder followed by some post processing measures.
According to the research conducted by Chen, indirect SLS of carbon based composite material accommodates the material and procedure selection criteria for the PEM fuel cell bipolar plate fabrication [Chen 2006]. This Indirect SLS proceeded in 3 stages to meet all the plate requirements:

1. SLS of bipolar plates
2. Carbonization of the binder
3. Epoxy infiltration

In Table 9, the parameters for the first stage which is the SLS process are shown.

<table>
<thead>
<tr>
<th>Powder constituents</th>
<th>Graphite (GrafTech GS150E) and Phenolic resin (Georgia Pacific GP5546)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Composition:</td>
<td>70w% graphite and 30 w% phenolic resin</td>
</tr>
<tr>
<td>Average particle size</td>
<td>Graphite: 80 μm  Phenolic resin:11μm</td>
</tr>
<tr>
<td>SLS machine</td>
<td>DTM Sinterstation 2000</td>
</tr>
<tr>
<td>CO₂ laser power</td>
<td>10~20 W</td>
</tr>
<tr>
<td>Laser scan speed</td>
<td>60 in/s</td>
</tr>
<tr>
<td>Powder layer thickness</td>
<td>0.004 in</td>
</tr>
<tr>
<td>Powder bed preheating Temp</td>
<td>60°C</td>
</tr>
<tr>
<td>Purging gas</td>
<td>Nitrogen</td>
</tr>
</tbody>
</table>

After the SLS process, post processing of the bipolar plates was further carried out. The post processing basically consisted of two steps of binder carbonization and epoxy infiltration.
Carbonization process

A vacuum furnace was used for this purpose. The maximum heating capacity of the vacuum furnace being 2000°C. Argon gas was filled into the chamber to prevent oxidation of carbonized phenolic resins which reduce the glassy carbon yields. The temperature rise and the ramp rates for this process are tabulated in Table 10.

<table>
<thead>
<tr>
<th></th>
<th>Temperature</th>
<th>Ramp Rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initial Profile</td>
<td>Room Temp-200°C</td>
<td>60°C/hr</td>
</tr>
<tr>
<td>Intermediate</td>
<td>200°C-600°C</td>
<td>30°C/hr</td>
</tr>
<tr>
<td>Final Profile</td>
<td>At 800°C</td>
<td>0</td>
</tr>
</tbody>
</table>

At 800°C, the dwell time was 1 hour. During this process the phenolic binders are burned off and a part of it was converted into glassy carbon. This resulted in good interconnected pores which increased the electrical conductivity.

Epoxy infiltration for final sealing

After the carbonization process, the structure was still found to be porous. Epoxy resin was chosen as the infiltrant to seal these pores because of its good mechanical strength, chemical stability and ability to wet most substrate materials. Clear coat resin which is a mixture of more than 70% diglycidyl ether of bisphenol A and less than 30% alkylglycidyl ether was used for this purpose. The resin was initially cured with the help of a hardener and then diluted with solvents like toluene, xylene etc., this was done in order to reduce the viscosity of the resin so that it can easily penetrate through the cured
pore structure. The epoxy, hardener and the solvent should be mixed in proper ratios (2:1:1) to avoid the formation of un-reacted epoxy and hardener which affect the final part properties. So as to form a gas tight plate structure the brown part were immersed in the infiltrant at least twice. The parts were then oven dried at 60°C for several hours to remove residual moisture. The electrical conductivity of the infiltrated parts was found to be better than the brown parts. The final properties of the bipolar plates are shown in Table 11.

4.3 RESULTS AND DISCUSSION

Table 11 Properties of the SLS bipolar plates [Chen 2006]

<table>
<thead>
<tr>
<th>PROPERTY</th>
<th>TEST METHOD</th>
<th>VALUE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Flexural strength</td>
<td>Two point bend test</td>
<td>1730psi</td>
</tr>
<tr>
<td>Electrical Conductivity</td>
<td>Four point probe test</td>
<td>80 S/cm</td>
</tr>
<tr>
<td>Specific weight</td>
<td>Archimedes principle of fluid displacement</td>
<td>Avg. density=1.27g/cm³</td>
</tr>
<tr>
<td>Corrosion Rate</td>
<td>Tafel extrapolation method</td>
<td>6µA/Cm²</td>
</tr>
<tr>
<td>Gas Permeability</td>
<td>Mass spectrometer leak detector</td>
<td>5x10^-6 Cm³/Cm².s</td>
</tr>
<tr>
<td>Interfacial contact resistance</td>
<td></td>
<td>&lt;200mΩ.cm² / 1.6MPa</td>
</tr>
</tbody>
</table>

All the above properties were found to be satisfactory but the electrical conductivity of these bipolar plates could be enhanced to meet the target set by the DOE [DOE 2005].
The following methods were followed to improve the electrical conductivity [Chen 2006]:

- Infiltration of brown parts with conductive polymer
- Addition of a liquid phenolic infiltration/re-curing step prior to final sealing
- Reduction of glassy carbon resistivity by curing process parameter control

These processes showed results which are quantified as below in Table 12 and the improvements in electrical conductivity are illustrated in Figure 17.

<table>
<thead>
<tr>
<th>First Infiltration/Recurring step</th>
<th>~108 S/cm (35% boost in the conductivity)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Second Infiltration/Recurring step</td>
<td>~117 S/cm (8.3% further enhancement)</td>
</tr>
</tbody>
</table>

Figure 17 Improvement in electrical conductivity with each step [Chen 2006]
5. CONCLUSIONS

The paper reviews three Additive Manufacturing (AM) processes. Each process is suitable for building specific fuel cell components. Performance characteristics of these components fabricated using AM processes prove that they give performance equal or better than the components fabricated by conventional techniques. Inkjet printing, amongst all AM processes is the process where most of the research has been undertaken with regards to building fuel cell components. The reason for that is it is easiest to commercialize as compared to rest of the methods since off the shelf printers have been demonstrated to produce components competitive with the commercial ones. FDM is convenient for planar array fuel cells as compared to MEMS and CNC manufacturing. Indirect SLS of graphite composite bipolar plate demonstrates fabrication of plates having superior characteristics which also meet DOE specifications. Development of these processes on a commercial basis is still under a lot of investigation.

6. ACKNOWLEDGEMENTS

This research was supported by a grant from the U.S. Air Force Research Laboratory contract #FA4819-09-C-0018. Support from the Missouri S&T Intelligent Systems Center is also greatly appreciated.
7. REFERENCES


II. COMPARISON OF DIRECT DEPOSITION PROCESS AND ELECTRO-WRITE PROCESS FOR PROTON EXCHANGE MEMBRANE FUEL CELL MEA MANUFACTURING

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\textsuperscript{b}Department of Material Science and Engineering, Missouri University of Science and Technology, Rolla, MO 65409 USA

ABSTRACT

Fuel cells are an important source of power for the future. Being in an energy demanding era, we are in dire need of new efficient power sources. However, there are issues regarding fuel cell manufacturing which need urgent attention. The paper discusses the manufacturing of the Proton Exchange Membrane (PEM) fuel cell MEA’s by two methods, namely Direct Deposition Process (DDP) and Electro-write Deposition Process (EWP). The comparison is carried out to provide us with the knowhow of the most suitable method for MEA manufacturing. The paper discusses the impact on the cost of the fuel cell by means of comparison of the two processes, the DDP and EDP in terms of their efficiency by a unique method. This paper is an introductory work for forming a basis of comparison and more detailed works will follow.
1. INTRODUCTION

In most countries around the world, the current energy supply system is considered as not being sustainable, in particular because of climate change impacts and the consumption of non-renewable energy resources [Krewitt 2006]. It is clear that transition from conventional fuels to clean and non-exhaustible ones is unavoidable [Asghari 2010]. Among the various renewable energy sources, fuel cell technology has received considerable attention as an alternative to the conventional power units due to its higher efficiency, clean operation and cost-effective supply of power demanded by the consumers [Erdinc 2010]. For small portable applications, fuel cells are the closest possible alternative to batteries since batteries do not provide the expected power density. Amongst all the different types of fuel cells, the Proton Exchange Membrane (PEM) fuel cells have received the highest attention due to low fuel permeability, high proton conductivity, high efficiency and good thermal stability [Peighambardoust 2010]. The main shortcomings for fuel cell development, however, are the cost and non-feasibility of mass production. The two factors however are interlinked with each other. If fuel cells could be mass produced in the near future, the cost will go down substantially. The other factors which can reduce the cost of PEM fuel cells are the high utilization of catalysts, low cost manufacturing process, use of different catalysts, heat and water management.

A fuel cell consists of many components such as Bipolar Plates, Gas Diffusion Layers (GDL), Membrane Electrode Assemblies (MEA’s) etc. The GDL allows direct and uniform access of the fuel and oxidant to the catalyst layer, which stimulates each half reaction [Mehta 2003].
The cost of a fuel cell is highly dependent on the utilization of the catalysts. Hence, to minimize the wastage of catalysts, research has been carried out to spray the catalyst onto the GDL with maximum efficiency. However, there is no sure way to estimate how efficient the process is. The paper discusses the efficiency of a manufacturing process in detail with regards to two catalyst spraying processes, the Direct Deposition Process (DDP) and the Electro-Write Process (EWP)
2. EXPERIMENTAL WORK

MEA manufacturing is the most complicated manufacturing process of all the components in a fuel cell. It’s mainly because MEA is not a single component but a series of components which need to be bonded precisely. For this reason, the manufacture of MEA proceeds in multiple steps. Hence, initially a cost model was estimated to analyze the costs incurred in an MEA manufacturing process. After it was calculated, the next important step is to analyze which is the most critical step in the entire manufacturing process. For this critical step, it is important to know what the most efficient manufacturing process is for it. This was carried out using the surface characterization techniques such as Scanning Electron Microscopy (SEM) and Energy-dispersive X-ray spectroscopy (EDS). SEM is a high resolution imaging technique that helps in analyzing the surface characteristics of the sample and EDS helps to identify the elemental nature of the surface. As described earlier, MEA manufacturing proceeds in multiple steps such as Catalyst Ink preparation followed by Catalyst spraying, which is spraying the catalyst ink onto the GDL using an XYZ platform and syringe disposing gear. After the catalyst spraying process, the GDL becomes a GDE (Gas Diffusion Electrode) since now the GDL contains the catalyst and thus becomes an electrode of the fuel cell. This is followed by hot pressing, in which the proton exchange membrane is pressed between two GDE’s to form an MEA. The flowchart for MEA manufacturing is presented in the following sections of this paper.
2.1 CATALYST INK PREPARATION

The catalyst ink was prepared using catalyst particles and Nafion LQ-1105 5% by weight NAFION®, 1100 EW. From the literature survey, it is known that Nafion® forms solutions having dielectric constants i.e. ‘ε’ more than 10, colloidal solution for ε between 3 and 10, while precipitate for ε less than 3 [Shin 2002]. Isopropyl alcohol has ε of 18.3 and hence it was used. Isopropyl alcohol and a dispersant were added to ensure uniform dispersion. The quantity of these ingredients was chosen in a way so that the solution has a required range of viscosity and surface tension. The range of viscosity chosen was between 2-5 cP and the range of surface tension chosen was between 35-40 mN/m. This range for both the viscosity and the surface tension was chosen so that the solution does not form lumps on the surface of the substrate or it does not remain in the syringe disposer for a long time either. After adding the ingredients, the ink was kept for stirring for approximately 36 hours using Fisher Scientific Isotemp magnetic stirrer. The stirring was carried out to make sure the solution has a uniform dispersion. In general, the nano scaled catalyst particles should come in touch with other components uniformly which is why the stirring is carried out [Zhang 2008]. The mixing was initially carried out in small steps by adding the ingredients slowly and simultaneously checking for its viscosity and surface tension to ensure that they stay in the desired range.

The flowchart is Figure 1 describes the flowchart for the MEA manufacturing process.
2.2 FLOWCHART FOR PREPARING THE MEA

Ink preparation
- Taking optimum quantity of Nafion solution, Isopropyl Alcohol, Catalyst
- Mixing the components and stirring using magnetic stirrers for 24-48 hours

Spraying
- Taking the prepared catalyst ink and loading into the dispensing syringe
- Using the process to spray the catalyst ink onto the Carbon Paper substrate

Hot Pressing
- Drying the Substrate containing the sprayed ink
- Hot pressing the 2 sprayed substrates onto two sides of the proton exchange membrane at 1250 psi, 110°C for 3 minutes

Figure 1 Flowchart for MEA manufacturing
3. ELECTROSPRAY PROCESS MODEL

In this paper, unlike the usual performance analysis, the efficiency of a process is compared. Before the comparison, it is necessary to understand which the most critical process is in the entire MEA manufacturing process. For this, an experiment is carried out using a horizontal electrospraying apparatus to spray the catalyst ink onto the GDL which is the Toray Paper TGP H-090.

The electrospraying process was taken into consideration initially for the manufacture of GDE. A 5cm×5cm Toray cloth was the GDL for the process. The Iridium oxide catalyst content was 25 mg/10ml of ink. Hence, for achieving a loading of 0.4mg/cm² which is considered as a standard loading, we prepared only 4 ml of ink. To achieve uniform dispersion, it was stirred additionally for 1 hour before it was loaded in the syringe.

The optimum parameters for the electrospraying process were fixed by experimentation. In this, one parameter which was the droplet ejaculation rate was fixed and the other two parameters which are the voltage and the distance of the needle from the GDL were varied. The droplet ejaculation rate was also found out by a similar method in which the other two factors were fixed and only the droplet ejaculation rate was varied to find out the optimum rate at which the droplet is absorbed into the GDL without formation of the drop on the surface of the GDL. In this way, the following parameters were found are best for the electrospraying process to manufacture a GDE which are listed in Table 1.
Table 1 Optimum parameters for electrospraying process

<table>
<thead>
<tr>
<th>Voltage</th>
<th>Droplet Ejaculation Rate</th>
<th>Distance from the GDL</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>75-80 µL/min</td>
<td>0.25 cm</td>
</tr>
</tbody>
</table>

After this task, the electrospraying apparatus is mounted on a XYZ table and automatic electrospraying of the catalyst ink onto the GDL is carried out. From this study, the optimum traversing speed for the electrospraying process was found out to be 2 in/min for the entire quantity of ink to spread uniformly on the surface of the GDL. This optimum speed is found on the basis of visual inspection of non-formation of any droplet on the surface of the GDL. The setup of the process is shown in Figure 2.

Figure 2 Horizontal electrospray apparatus
The pattern in which the ink is sprayed on the GDL is shown below. The spraying was started from the left bottom end of the GDL and then the progress is as shown in the figure. Totally, it took 25 minutes for the entire GDL to be sprayed with the optimum amount of ink which is 4 ml. The pattern is shown in Figure 3

![Pattern of ink spraying on GDL](image)

Figure 3 Tool path for horizontal electrospray process

### 3.1 ELECTROSPRAYING PROCESS COST MODEL

The cost model for the electrospraying process was calculated considering the cost of the process per liter of catalyst ink. The catalyst loading considered for the MEA is 0.2 mg/cm², a standard loading according to previous literature review.

As every process has certain wastage associated with it, a wastage rate of 10% was considered for each of the process in the final cost. Tables 2-7 list the processes and the component costs in the entire MEA manufacturing process.
Table 2 Catalyst ink component cost

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>for 10 ml</th>
<th>for 1 liter</th>
<th>Cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nafion liquid</td>
<td>1ml</td>
<td>100 ml</td>
<td>$105</td>
</tr>
<tr>
<td>Catalyst particles</td>
<td>10mg</td>
<td>1 g</td>
<td>$110</td>
</tr>
<tr>
<td>Isopropyl alcohol</td>
<td>8ml</td>
<td>800 ml</td>
<td>$5</td>
</tr>
<tr>
<td>Dispersing agent</td>
<td>1 ml</td>
<td>100 ml</td>
<td>$5</td>
</tr>
<tr>
<td><strong>Total cost</strong></td>
<td></td>
<td></td>
<td><strong>$225</strong></td>
</tr>
</tbody>
</table>

**Final Cost (10% wastage considered)**  
$247.5

Table 3 Catalyst ink preparation cost

<table>
<thead>
<tr>
<th>Description</th>
<th>Cost/Quantity</th>
</tr>
</thead>
<tbody>
<tr>
<td>No of stirrers</td>
<td>2 units</td>
</tr>
<tr>
<td>Stirrer cost ($)/liter</td>
<td>~ $0.71</td>
</tr>
<tr>
<td>power consumed</td>
<td>$1</td>
</tr>
<tr>
<td>1 liter beaker cost for 2 units</td>
<td>$18</td>
</tr>
<tr>
<td><strong>Total operational cost</strong></td>
<td><strong>$19.71</strong></td>
</tr>
</tbody>
</table>

Number of MEA’s using loading of 0.2 mg/cm² using 1 liter catalyst ink is 100.

For making 100 MEA’s number of GDL’s required will be 200 and number of membranes required will be 100. The following table gives the material costs.

Table 4 MEA component cost

<table>
<thead>
<tr>
<th>Component</th>
<th>Units required</th>
<th>Cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>GDL</td>
<td>200</td>
<td>$235</td>
</tr>
<tr>
<td>Membranes</td>
<td>100</td>
<td>$750</td>
</tr>
<tr>
<td><strong>Total material cost</strong></td>
<td></td>
<td><strong>$985</strong></td>
</tr>
</tbody>
</table>

**Final Cost (10% wastage considered)**  
$1083.5
Table 5 Electrospray apparatus cost

<table>
<thead>
<tr>
<th>Activity</th>
<th>Time/Cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>Catalyst spraying time with electrospray apparatus</td>
<td>30 minutes</td>
</tr>
<tr>
<td>Considering there are 10 machines in the shop</td>
<td>300 minutes</td>
</tr>
<tr>
<td>Setup time for 1 run</td>
<td>10 minutes</td>
</tr>
<tr>
<td>Labor cost ($20/hr, 4 employees)</td>
<td>$480</td>
</tr>
<tr>
<td>Equipment cost for 10 syringe dispensers</td>
<td>3200/240</td>
</tr>
<tr>
<td>Battery and power supply cost for 10 dispensers</td>
<td>2220/240</td>
</tr>
<tr>
<td>Syringe costs for 1 liter ink</td>
<td>20 syringes/liter</td>
</tr>
<tr>
<td>Total operational cost</td>
<td>$512.58</td>
</tr>
<tr>
<td><strong>Final cost (10% wastage considered)</strong></td>
<td><strong>$563.84</strong></td>
</tr>
</tbody>
</table>

In Table 5, the equipment cost for the syringes dispensers is considered as per day cost. The total cost of 10 syringe dispensers for a day is stated in the last column and the same holds true for the battery and power supply equipments too. The number of working days in an year is considered to be 240 days.

After the electrospraying operation, the next step requires some post processing in order to bond the GDE’s with the membranes. The best parameters identified for the hot pressing are 1000 psi, 100°C and 2 minutes [Therdthianwong 2007]. Also, the equipment depreciation has to be taken into consideration while computing the process cost. The life time of the hot pressing apparatus is considered to be 7 years. Thus the hot pressing apparatus cost can be calculated using the following equation

$$\text{Hot pressing cost} = \frac{N \times A}{Y \times D}$$ ................................................................. (1)

where,

N= Number of machines required
A= Cost of 1 machine

Y= Life of the equipment in number of years

D= Number of operational days in an year

Thus, the hot pressing cost will be \[ \frac{N \times A}{Y \times D} = \frac{2 \times 25000}{7 \times 240} = 29.7619 \]

Considering the hourly labor charges as $20, 5 hours would be required to hot press 100 MEA’s. Accordingly, for 1 employee, the labor cost comes to $100.

Table 6 Hot pressing cost

<table>
<thead>
<tr>
<th>Activity</th>
<th>Cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hot pressing apparatus cost</td>
<td>$29.7619</td>
</tr>
<tr>
<td>Labor cost ($20/hr, 1 employee)</td>
<td>$100</td>
</tr>
<tr>
<td>Total operational cost</td>
<td>$129.76</td>
</tr>
<tr>
<td><strong>Final cost</strong></td>
<td><strong>$129.76</strong></td>
</tr>
</tbody>
</table>

Table 7 Total cost of MEA manufacturing

<table>
<thead>
<tr>
<th>Operation</th>
<th>Cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>Components for catalyst ink</td>
<td>$247.5</td>
</tr>
<tr>
<td>Catalyst ink preparation</td>
<td>$19.71</td>
</tr>
<tr>
<td>MEA components</td>
<td>$1083.5</td>
</tr>
<tr>
<td>Electrospraying</td>
<td>$563.84</td>
</tr>
<tr>
<td>Hot pressing</td>
<td>$129.76</td>
</tr>
<tr>
<td><strong>Total cost</strong></td>
<td><strong>$2044.31</strong></td>
</tr>
</tbody>
</table>
3.2 EXPERIMENT CONCLUSION

Thus the total cost of preparation of 100 MEA’s comes out to be $2044.31. Hence the cost per piece is $20.44, which is considerably less expensive as compared to the commercially available MEA’s. However, it is understood from the overall study that the cost depends primarily on the process of GDE preparation because the other costs incurred are mainly raw material costs. Hence, if the process of GDE preparation is optimized to achieve best results in shortest time, a lot of cost reduction can be achieved.

Optimizing the GDE preparation process means that the speed of the process should be increased so that it does not act as a bottleneck. If it takes 25 minutes to spray the ink over 1 MEA, it will certainly act as a bottleneck to the entire MEA manufacturing assembly. Hence, to accelerate the process, DDP (Direct Deposition Process) and EWP (Electro-write process) are chosen to be compared since they are the faster than conventional catalyst ink spraying processes such has hand painting and screen printing processes [Taylor 2007]. DDP consists of a vertical syringe mounted on an XYZ platform and which can be programmed to deposit over a tool path. Hence, the Z axis doesn’t move as might be expected from an XYZ platform. EWP is essentially the same process as DDP except the fact that the electrospraying circuit is added to the DDP process to accelerate it by increasing the rate of deposition. The rate of deposition increases because of the external electric field wherein the syringe needle acts as the anode and the substrate as the cathode thereby attracting the catalyst ink from the syringe needle.
4. ANALYSIS OF EXPERIMENTS

For the analysis, an X-Y-Z platform was constructed using the Fab@home model 2 apparatus. This apparatus is equipped with printing head capable of moving along the X &Y axes whereas the loading base moves vertically to give the 6 degrees of freedom. For the EWP, a power supply and a battery had to be used. During the EWP, the loading base was treated as cathode whereas the syringe needle was the anode. Other than the power supply and the battery, rest of the apparatus was common for both the DDP and the EWP. Figure 4 shows the setup of the apparatus.

Figure 4 XYZ platform used for both the DDP and EWP
4.1 EFFICIENCY CALCULATION FLOWCHART

The flowchart explains the steps required to calculate the efficiency of the catalyst ink spraying process. Next, the paper describes the efficiency calculation of the DDP in detail. To compare the two processes, it is necessary to set some common parameters. For the experiment described in this paper, the common parameters were:

- Quantity of the ink sprayed onto the GDL
- Translational speed of the XYZ platform
- Tool Path
- Time taken by the machine to complete the tool path
4.2 DDP EFFICIENCY CALCULATION

After the catalyst spraying process, the carbon substrate was analyzed by SEM and checked for the Iridium oxide particles. For the efficiency calculation, Iridium oxide particles were used because it is one of the novel catalysts in the field of PEM fuel cells and Iridium particles are easier to identify by SEM. 5 ml of Iridium oxide catalyst ink containing Nafion solution, Isopropyl alcohol, dispersant and Iridium oxide catalyst particles. A total of 6.25 mg of Iridium oxide was loaded into the ink. The ink was sprayed onto 25 cm² of carbon paper giving it a loading of 0.25 mg/cm². After the spraying operation, small sections of the GDE from the entire area of the GDE were cut to be analyzed by SEM techniques. This was carried out in order to collect information from the entire GDE and hence random sections were chosen. To identify the elemental nature of the surface characteristics, Energy Dispersive X-ray Spectroscopy (EDS) technique was used.

During the SEM analysis, every possible particle in the image was analyzed by Electro-Discharge Spectroscopy using the EDAX Genesis software. Each particle was analyzed and was checked for its elemental nature. Many such trials were carried out to know if it’s an Iridium particle or any other. After analyzing around 50 such particles, the rest of the particle count was carried out using just simple judgment since an Iridium particle stands out having a very high brightness as compared to the otherwise dark background. The procedure of calculation of efficiency in described next. Flowchart in Figure 6 demonstrates the steps required for calculating the efficiency.
The above flowchart describes efficiency calculation procedure for the processes. This procedure is common for both the processes. In the second step, while calculating the diameter, it is assumed the catalyst particles are spherical. Considering they are finely ground particles, it is a fair assumption.
Iridium oxide catalyst ink was deposited using the XYZ platform apparatus onto the Toray cloth. After deposition, the deposited Gas Diffusion Electrodes (GDE) were allowed to dry for a full day so that all the volatile ingredients from the ink evaporate. After drying it for a day, small samples of the GDE’s were cut from different areas of the GDE to be analyzed under the Scanning Electron Microscope (SEM). The high resolution images were captured using the SEM and then analyzed using the ImageJ software. The images were analyzed using the SEM Hitachi S-4700.

To identify if the particles are catalyst particles, EDS techniques were used. During EDS analysis, high energy beam of charged particles is focused onto the sample which leads to an emission of characteristic X-rays which are specific to individual elements. This is how the elements present in the sample were identified, which in this case are the catalyst particles.

Thus the average number of particles ranged from 11-23 in each image. After spotting the particles, the average particle diameter was analyzed using the ImageJ software and was found out to be 1.69 µm ± 0.17 µm. The average number of particles was found to be 14.125. Figure 7 shows the sample image of the GDE manufactured using the DDP and the arrow shows the catalyst particle. In this figure, it can be observed that the particle spread is less. The spread of the catalyst particles was similar in all the images captured.
After the analysis of the images and calculating volumes, the following are the results

- For 25 cm², the total volume of the catalyst particles was 0.082 mm³
- The density of catalyst is 22.42 mg/mm³ and hence, the mass is 1.837 mg
- Considering the initial loading as 12.5 mg for 25 cm² area, the efficiency will be

\[
\text{Efficiency} = \frac{\text{Calculated Loading}}{\text{Initial Loding}} = 15\% \\
\]

Thus, the efficiency of the DDP comes out to be 15% according the calculation procedure adopted here, which is common for both the DDP and the EWP.
The calculation of the EWP process is calculated in the similar way as for the DDP. As explained earlier the EWP is similar to DDP except the following addition of the Electrospray apparatus in which the substrate acts as a Cathode and the syringe needle acts as an Anode. The perceived advantages of this process are:

- The ink flow rate is more than in DDP because of the extra electric field.
- The ink stream is more uniform and very linear.

The same procedure was carried out to study the images and to find the volume of the particles and then accordingly the efficiency. During the SEM analysis, many random images were captured from various different parts of the GDE to study the surface characteristics with regards to the spread and distribution of the catalyst particles over the area of the GDE. The particles in the images were analyzed using the EDS techniques for examining their elemental nature and verifying if they are the catalyst particles. The images were captured using the SEM Hitachi S-570 and the Revolution 4 pi software was used to use the EDS techniques for examination. After analyzing many such particles, the rest of the particle count was carried out using simple visual judgment since the catalyst particle stands out having a very high brightness as compared to the otherwise dark background.
Figure 8 shows a sample image of one of the areas of the GDE manufactured using the EWP. This image was taken with the SEM Hitachi S-570. In Figure 8, it is evident that the spread of catalyst particles is more than that observed in DDP.
After the analysis of the images and calculating volumes, the following are the results

- For 25 cm², the total volume of the catalyst particles was 0.025 mm³
- The density of catalyst is 22.42 mg/mm³ and hence, volume × density = mass.
- The mass thus calculated is 0.57 mg
- Considering the initial loading as 1.25 mg for 25 cm² area, the efficiency will be

\[
\text{Efficiency} = \frac{\text{Calculated Loading}}{\text{Initial Loading}} = 45.6\% \quad \text{............................................................. (3)}
\]

Thus, the efficiency of the EWP comes out to be 45.6% which is more than 3 times that of the DDP.

**4.4 PROCESS COMPARISON OBSERVATIONS**

One analysis which is important other than the efficiency is to know the uniformity of the ink which has been sprayed onto the GDL. The number of particles in the images can help in this regard.

For the DDP, 8 random images were taken with the same magnification from the various parts of the GDE and analyzed by the SEM. The number of particles ranged from 10 particles as the least number of particles to 19 particles as the most particles. The standard deviation for the DDP was 2.65.

For the EWP 6 images were taken with the same magnification from the various parts of the GDE and analyzed by the SEM and the number of particles ranged from 10 particles as the least number of particles to 33 particles as the most particles. The standard deviation for the EWP was 8.8.
This shows that there is a large variation in the number of particles found at different places on the GDE manufactured using the EWP. This can also affect the performance of the MEA.

This leads to a conclusion that deeper understanding of the EWP is needed to demonstrate its spraying uniformity. Hence, it was necessary to map the spread of the particles in various areas of the GDE to understand the variations in the amount of catalyst spread. It is necessary to do that in accordance with the tool path used for the EWP, which is similar to the DDP as already mentioned earlier. Hence, an analysis of the horizontal variation and the vertical variation in the density of the catalyst particles is carried out in the next part of the paper.

In this analysis, 2 small sections of the GDE manufactured by EWP, 5 mm in length are cut, one in horizontal direction, one in vertical direction. The horizontal section is the path where the ink has been deposited. Since, the ink spread is not 5 mm, the vertical section contains the path wherein some of the path might not have the ink spread which is what would be analyzed in this section. Figure 9 shows the same vertical and horizontal variation.
The Figure 9 shows the tool path on a 5cm×5cm carbon paper GDL. The horizontal variation and the vertical variation in the density of the catalyst sprayed onto the GDL have been analyzed in the next part of the paper. These variations have also been analyzed using the SEM and the EDS techniques.

**4.5 VERTICAL VARIATION IN EWP**

For studying the vertical variation, a 5 mm piece of the GDL where the top and the bottom boundaries are exactly the ink paths have been taken into consideration. The EDS was used to spot the Iridium particles and the particle count was taken in various places along the vertical length i.e. starting from top and ending at the bottom of the 5 mm piece. This variation is shown in Figure 10.
Figure 10 (a) Catalyst particles in top section, (b) middle section and (c) bottom section with EWP

The Figure 10 shows that quite a few particles can be spotted in the top section and the bottom section of the 5 mm piece. However, very few particles can be spotted in the middle section as seen from the image. However, there is need to quantify to know
the variation. Hence, finding the number of particles using EDS and ImageJ software is carried out in the next part of the paper. The variation of the count of particles as a function of the track width is shown in Figure 11

![Graph showing vertical variation of number of particles as a function of track width](image)

Figure 11 Vertical variation of number of particles as a function of trackwidth

The above figure shows the graph of the vertical variation in the number of catalyst particles as a function of distance from the top of the 5 mm GDE piece. In the Y axis scale, the ‘t’ stands for the track width point; for e.g. 0.6t means at 3/5\(^{th}\) the distance from the top. It is evident from the figure that there is a lot of variation in the vertical direction and it valleys in the middle section where there has been no spraying of catalyst.
4.6 HORIZONTAL VARIATION IN EWP

For studying the horizontal variation, a 5 mm rectangular piece of the GDE is taken into consideration which has been the ink path while spraying. The EDS technique was used to spot the Iridium particles and the particle count was taken in many places. The particle count was taken along the horizontal length. The Figure 12 shows the horizontal piece and Figure 13 shows the horizontal variation of the number of particles as analyzed by the SEM. Figure 14 graphically describes this variation.

![Ink path section for horizontal variation analysis](image)

Figure 12 Ink path section for horizontal variation analysis
Figure 13 (a) Catalyst particles in the left section, (b) middle section and (c) right section with EWP
The above figure shows the graph of the horizontal variation in the number of catalyst particles as a function of distance from the left of the 5 mm GDE piece. It is evident from the figure that there is not a high variation in the horizontal direction as much as it was in the vertical direction. This is primarily because of the high stirring in the ink which results in uniform spread of the catalyst particles in the entire volume of the ink. This proves that the uniformity in the catalyst layer on the GDE resulting in higher performance of the fuel cell in less cost.
From the above experiments, the most favorable conclusions which come out are that ink spread is substantially uniform in the horizontal directions of the tool path but not the vertical ones. It is hence imperative to spray the ink twice and the 2\textsuperscript{nd} tool path should be covering the areas which are left out from the first too path. Figure 15 illustrates the procedure.

![Figure 15 Additional path for catalyst ink deposition in the EWP](image)

Thus, from the above conclusions, it is understood that it is necessary to have the 2\textsuperscript{nd} tool path 2.5 mm away from the first tool path to achieve a uniform loading of the catalyst on the entire GDL.
5. CONCLUSIONS

This paper presents the unexplored part in fuel cell science such as catalyst spraying process efficiency. The cost model gives an idea of the most important process involved in the process of MEA manufacturing, which in turn needs to be optimized to achieve lower costs and efficiency. It is very important to focus on the efficiency of the process in order to reduce the cost and make fuel cells more efficient. From the study, it is evident that the electrospraying process is suitable for catalyst spraying operation as it is fast and efficient and along with a low cost XYZ translational platform can also be an inexpensive operation. The use of SEM and EDS techniques for calculating the efficiencies of the process gives accurate results. The EWP achieves more than 3 times the efficiency than that of DDP. In order to make the MEA more efficient, a second tool path to cover up the un-sprayed regions has been suggested.

6. ACKNOWLEDGEMENTS

This research was supported by the National Science Foundation grants DMI-9871185 and IIP-0637796, and a grant from the U.S. Air Force Research Laboratory contract. Support from Product Innovation and Engineering, LLC, MISSOURI S&T Intelligent Systems Center, is also greatly appreciated.
7. REFERENCES


SECTION

2. CONCLUSION

This paper gives a lot of details about the applications of additive manufacturing for manufacturing fuel cell components. Fuel cell components being small and detailed, additive manufacturing can open up wide number of areas for their manufacturing at low cost. Amongst various processes, SLS is the most suited process for bipolar plate manufacturing. 3D inkjet printing can be effectively used for the catalyst spraying process which is complicated and needs to be precise to spread the catalyst over the entire GDL uniformly. In the second part of the thesis, a method for comparison of the two processes used for manufacturing the MEA’s has been devised. The comparison and the efficiency calculation have been carried out using SEM and EDS techniques. This is very easy and cost-effective. Amongst the two processes compared for catalyst spraying operation, EWP came out topping the efficiency at 45.6% whereas the DDP was only 15% efficient. The EWP is further analyzed for further variation in its spraying. Further analysis of the EWP method led to a conclusion that because of the ink spread not being across the entire area of the GDL, an alternate ink path at 2.5 mm from the original ink path is warranted.
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