BIOMASS CONVERSION TO FUELS AND CHEMICALS

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Introduction
Biomass has a significant potential to provide liquid, solid and gaseous fuels in conjunction with the co-production of food, feed, fiber and new bioproducts. The selection of a conversion technology will be a function of both the supply of locally available feedstock and the markets for products and will determine the degree to which various biological and chemical routes are deployed. This paper reviews the major technical principles and challenges in the evaluation of feedstocks and conversion systems for the energy markets of the future. The challenges of energy supply, reliability, environmental performance and lifecycle impacts will also be discussed for thermochemical biomass conversion via gasification.

Biomass Supply
The growing acceptance of biomass as the only renewable source of carbon for the future requires that a realistic appraisal of its potential be undertaken. While solar energy captured by wind, PV or high temperature concentrating solar systems, have very high energy supply potentials—far exceeding today’s total primary energy supply of approximately 417 EJ (IEA,TPES data for 2000)—it will be delivered in the form of electricity and potentially hydrogen. However, before a purely hydrogen and electrical economy can arrive, during the 21st century the world has to undergo a carbon usage transition from directly emitting fossil carbon dioxide into the atmosphere to one in which fossil carbon is captured and sequestered, and biomass becomes the sole form of the earth’s carbon resource that can be emitted directly to the atmosphere due to its short term photosynthetic cycle in the biosphere.

Unlike fossil fuels for which exploration, discovery and assessment of resources and reserves is a well known science, renewable energy accounting introduces a flow concept to describe the size and availability of the resource. Biomass is rather unusual; it exists somewhere between the intermittent, flux-only resources such as solar and wind, and the traditional fossil fuel stock perspective. This is because of its annual and perennial storage capability, allowing dispersed and captured solar energy to be concentrated and converted into useful, energy and products on a continuous basis at centralized facilities according to normal demand patterns. Thus, the photosynthesis process of solar collection, crop harvest, and transport of biomass fuels will define the availability of biomass for energy and materials purposes.

As with fossil resources, the energy conversion technologies play a huge role in enabling the solar resources by dictating the efficiency, and the utility of renewables with respect to society and the environment—in sum, their sustainability. However, this presentation will concentrate on the biomass system only up to the conversion plant gate.

Biomass resources in the USA
The renewable energy cycle for biomass starts with photosynthesis, which captures the fraction of light between 400 nm and 700 nm in the solar spectrum to provide the energy to fix carbon dioxide and split water molecules, producing raw photosynthate according to the equation:

\[
\text{CO}_2 + \text{H}_2\text{O} + \text{sunlight} \rightarrow \text{CH}_2\text{O} + \text{O}_2
\]

The photosynthate formula, CH2O, is essentially the same as sugar, starch, or cellulose, the latter being the most common natural polymer on earth. Energy is obtained from biomass by means of combustion with oxygen as in a fire, or through metabolic processes, e.g., respiration to provide chemical energy for muscles and other processes:

\[
\text{CH}_2\text{O (Biomass)} + \text{O}_2 \rightarrow \text{CO}_2 + \text{H}_2\text{O}
\]

Either way, the net result is a closed cycle in which carbon dioxide and water vapor are returned to the atmosphere. This solar-energy powered closed-loop supports all life on earth, providing food for humanity, feed for animals, fibers for clothing, and materials for buildings and consumer products, in addition to fuel.

A theoretical estimate of the efficiency of photosynthesis is about 5.5%. The energy received from the sun at any given location varies with latitude and season; and the annual photosynthetic capture is a function of the water and nutrient availability and the seasonal climate variations. As a result actual crops achieve much less than this. The variation is very large ranging from the annual accumulation of less than 1 t ha\(^{-1}\) y\(^{-1}\) of dry biomass in a mature boreal forest, to approximately 50 t ha\(^{-1}\) y\(^{-1}\) of total biomass for a record corn yield (in Iowa, in 1999) of 408.2 bushels of corn per acre. This is calculated by accounting for the stalk and other above ground biomass components by means of the harvest index, which has a value of 0.5 applied to the 25.6 t ha\(^{-1}\) y\(^{-1}\) of corn kernels.

Sources of bioenergy
The primary biomass sources are natural forests, planted trees and shrubs, and agricultural crops. However, much of today’s bioenergy is derived from secondary, tertiary, and post-consumer residues derived from these primary resources. Secondary residues are mainly harvesting residues and include: forest slash; crop residues such as straw and stover from cereal production; as well as minor sources such as orchard, rights of way, and vineyard prunings. Tertiary sources are usually process residues and include: black liquor produced in kraft pulping; sawdust and bark from solid wood processing; and sugar cane bagasse. Food processing tertiary residues include: orange and potato peelings; and large quantities of water containing various amounts of proteins, starches, and sugars. Animal operations that produce dairy products, eggs, and meat generate a large stream of residues (at ever increasing scales) from large feedlot operations. These are becoming regulated as concentrated animal feed operations (CAFOs). These tertiary residues are usually high moisture and most often are treated by a process of anaerobic digestion to produce biogas, a mixture of methane and carbon dioxide, as fuel.

Much of the post consumer residues are in fact biomass: paper, packaging, crates, pallets, demolition lumber, food scraps, and sewage treatment residues. These are more and more often entering the bioenergy, biofuels, and bioproducts stream as society practices resource conservation and use. The Environmental Protection Agency’s (EPA) preferred method is source reduction, which includes reuse, followed by recycling and composting, and, lastly, disposal in combustion facilities and landfills.

These primary, secondary, tertiary, and post consumer pathways are significant when compared with the US TPES of about 100 EJ. The total forest harvest is on the order of 203 M tonnes of air-dried...
round wood equivalents per year, which corresponds to > 3.5 EJ of primary energy equivalent. In recent years the annual corn harvest of almost 10 billion bushels (254 Mt) corresponds to over 4 EJ of primary energy equivalent produced on 3,000 km$^2$ (about 75 million acres, or about half of the arable crop area of the USA). Though corn stover is partly required to remain on the land to maintain fertility and reduce soil erosion, the quantity available is similar to the amount of crop and represents the largest agricultural residue potential.

**Effect of Transportation on Cost**

The key determinant of the feasibility of biomass is the cost of the biomass that contains the stored “free” solar energy. As the cost varies with the transportation distance from the process plant to the field or forest, as well as the opportunity cost of the feedstock, the cost is usually obtained from a supply curve. There are many factors that go into deriving the supply curve for the resource. Due to the bulk of biomass, its transportation over long distances by road is not economical, though with low cost transportation by rail, the distance can be increased. While an 80 km collection is often cited, this is only an order of magnitude. In each case, the actual collection radius will be a function of the biomass density per unit area, the nature of the terrain, road network density, and the condition and type of transportation available.

**Density Relationship**

One of the highest biomass densities is, in fact, the post consumer residue production of the urban areas. Cities draw upon the resources of vast areas and concentrate them in a small area resulting in a “footprint” that is much greater than the physical urban area. A useful number is a per capita urban residue energy generation rate of about 22 MJ caput$^{-1}$ d$^{-1}$ (1.86 kg d$^{-1}$ of residue with 11.6 MJ kg$^{-1}$ heat of combustion, 1990). Major metropolitan areas have population densities of greater than 2000 person km$^{-2}$. For this population density, the thermal energy equivalent in their residues is about 4.4 GWh km$^{-2}$. If a short rotation woody crop was grown at yields of 15 tonne ha$^{-1}$ y$^{-1}$, it would produce about 7.8 GWh km$^{-2}$.

**Conclusion**

The predicted 2020 potential of biomass costing less than 4 $ GJ$^{-1}$ is about 8 EJ. The sources and constraints on the biomass resources contributing to this total will be discussed.
BIO-OIL FROM RICE STRAW BY PYROLYSIS USING FLUIDIZED BED AND CHAR REMOVAL SYSTEM
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Introduction
Rice straw is one of abundant biomass whose production sums up to about 7 milion ton annually in Korea. It is usually utilized as a compost or forage in Korea. Nowadays intensive attention has been paid to rice straw from the viewpoint of a potential renewable energy source, due to high price of petroleum. Thermochimical conversion of biomass is one of the most common routes for conversion into energy. Among thermochimical conversion processes, pyrolysis of biomass is a promising tool to provide bio-oil which can be used as an alternative fuel oil or chemical feedstocks. In the last two decades, extensive researches have been carried out to understand the complexity of pyrolysis, to obtain optimal conditions for producing bio-oil, and to analyze the composition of bio-oil quantitatively and qualitatively. A great portion of past researches have focused on the pyrolysis of wood or wood wastes. Pyrolysis of rice straw, however, has been conducted not so intensively and once it is performed, only in small laboratory scale plants. This paper reports the results of pyrolysis of rice straw in a bench scale plant equipped mainly with a fluidized bed, a char separating system and a quench system. We investigated the influence of reaction temperature on the production of bio-oil, the efficiency of a char-separating system and some physical and chemical analyses of bio-oil, product gas and char.

Experimental
Rice straw. Rice straw obtained from a province in Korea was cut into rectangular form with a size of about 5 mm. Including elemental analysis, some analyses were performed. Table 1 shows characteristics of rice straw fed. The amount of cellulose, hemicellulose and lignin is based on volatile portion of rice straw.

Table 1. Analysis of rice straw.

<table>
<thead>
<tr>
<th>Composition</th>
<th>wt%</th>
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<tbody>
<tr>
<td>cellulose</td>
<td>54.67</td>
</tr>
<tr>
<td>hemicellulose</td>
<td>32.53</td>
</tr>
<tr>
<td>lignin</td>
<td>12.83</td>
</tr>
<tr>
<td>Water, ash, volatile</td>
<td>wt%</td>
</tr>
<tr>
<td>water</td>
<td>6.48</td>
</tr>
<tr>
<td>volatile</td>
<td>82.30</td>
</tr>
<tr>
<td>ash</td>
<td>11.22</td>
</tr>
<tr>
<td>Elemental analysis</td>
<td>wt%</td>
</tr>
<tr>
<td>C</td>
<td>39.2</td>
</tr>
<tr>
<td>H</td>
<td>4.84</td>
</tr>
<tr>
<td>N</td>
<td>1.60</td>
</tr>
<tr>
<td>S</td>
<td>0.67</td>
</tr>
<tr>
<td>O</td>
<td>53.69</td>
</tr>
<tr>
<td>Alkali metal Concentration (ppm)</td>
<td></td>
</tr>
<tr>
<td>Na</td>
<td>250</td>
</tr>
<tr>
<td>Mg</td>
<td>930</td>
</tr>
<tr>
<td>Ca</td>
<td>2100</td>
</tr>
<tr>
<td>K</td>
<td>12000</td>
</tr>
</tbody>
</table>

Bench scale pyrolysis plant. Experiments were carried out using a bench scale plant with a capacity of up to 3 kg/h. The fluidized bed reactor made of sus-304 has a free diameter of 154mm and a length of 616 mm. It is heated indirectly by electric power. Rice straw enters the reactor through two screw conveyors and then is pyrolyzed. A cyclone and a hot filter serve as char separator. These separators are heated up to 400 °C to prevent from the condensation of product gas. A series of quenching system is applied to cool product gas efficiently and quickly. Product gas is circulated into the fluidized bed using a compressor.

Figure 1. Fluidized bed reactor system.

Reaction conditions. Reaction conditions are shown in Table 2. In every experiment, particle size of feed was about 5 mm and product gas was served as a fluidizing medium. The residence time of product gas in the reactor lay ca. 2 or 3 seconds in each experiment and this makes it possible to compare the influence of reaction temperature on the product spectrum relatively clearly. Particle size of the sand used in the fluidized bed was ca. 0.4 mm.

Table 2. Parameters of experiments.

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>Run1</th>
<th>Run2</th>
<th>Run3</th>
<th>Run4</th>
<th>Run5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amount of input (g)</td>
<td>1700</td>
<td>1500</td>
<td>1560</td>
<td>1500</td>
<td>1500</td>
</tr>
<tr>
<td>Duration (min)</td>
<td>118</td>
<td>87</td>
<td>67</td>
<td>80</td>
<td>80</td>
</tr>
<tr>
<td>Feed rate (g/h)</td>
<td>864</td>
<td>1034</td>
<td>1397</td>
<td>1125</td>
<td>1125</td>
</tr>
</tbody>
</table>

Analysis. Qualitative and quantitative analysis of pyrolysis gas and oil has been performed by GC and GC-MS system. Using ICP the content of alkali metals in products was also analyzed. Solid content in bio-oil was measured using acetone as a solvent. In addition to chemical analysis, physical characteristics of bio-oil was also examined, such as density, viscosity etc.

Results and Discussion
Mass balance The behavior of biomass pyrolysis is dependent on many parameters, such as reaction temperature, residence time of gas in reactor, particle size of feed material and feed rate. This paper focuses on the influence of reaction temperature. Mass balance of each experiment is shown in Figure 2. As reaction temperature rises, the amount of gas increases. It is also shown that there is an optimum temperature to produce more bio-oil. In the reaction temperature range between 412 and 516 °C, the yield of oil was relative constant to give a maximum yield of 50 wt% and then decreases distinctly with increasing temperature.

This trend is typical for the pyrolysis of biomass, which results from the secondary reaction of volatiles[1]. The yield of bio-oil is lower than in the case of pyrolysis of wood, which is largely due to lower content of organic portion in rice straw than in wood and a larger content of ash in rice straw, some metallic components in which lower the yield of bio-oil by catalytic decomposition of oil portion. The use of hot filter can reduce the yield of bio-oil in which thermal cracking of pyrolysis vapor takes place. In the experiments, we used rice straw particle whose diameter lies about 5 mm for easy transporting it into the reactor. This relative larger size in comparison with typical pyrolysis of biomass can have an influence on the reduction of bio-oil yield [2].

**Product gas.** Figure 3 shows the development of main components in product gas with temperature. Methane, CO and CO₂ consist mainly of product gas. Ethene and ethane was other important components in the product gas. Hydrogen content was under 1 wt% in every experiment. It is shown in the Figure 3 that higher temperature leads to high yield of methane and CO, which results mainly from the secondary cracking of volatiles[3]. This tendency including higher content of other hydrocarbon gases at elevated reaction temperature gives high heating value to product gas.

**Bio-oil.** Obtained bio-oil shows a phase separation. Large portion of tar was deposited in bio-oil containers. Therefore, it was very difficult to sample it accurately. We conducted sampling according to the procedure at VTT and various chemical and physical characterizations of bio-oil were carried out [4]. Some distinct features of bio-oil are listed in Table 3. Water content in bio-oil was very variable according to samples but lay in the range of 50-60wt%. Product oil showed nearly metal-free character, which simplifies the treatment of bio-oil during up-grading and reduces catalytic reactions during storage. Reduction of density and increase of pH value in comparison with typical bio-oil from wood is caused mainly by high content of water in bio-oil. Solid content in bio-oil lay around 0.03 wt%. The value is somewhat high in comparison with mineral oil whose value is about 0.01 wt%. Applying of a hot filter, however, reduced solid content distinctly, compared with other processes without it. Solid seems to consist of hydrocarbons implied by the ash content in bio-oil (0.007 wt%).

**Table 3. Characteristics of bio-oil.**

<table>
<thead>
<tr>
<th>Analysis</th>
<th>Run5 (598°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alkali Metals (g/L)</td>
<td>Na</td>
</tr>
<tr>
<td></td>
<td>Ca</td>
</tr>
<tr>
<td></td>
<td>Mg</td>
</tr>
<tr>
<td></td>
<td>K</td>
</tr>
<tr>
<td>Solid content (wt%)</td>
<td>0.03</td>
</tr>
<tr>
<td>pH</td>
<td>4.07</td>
</tr>
<tr>
<td>Ash (wt%)</td>
<td>0.007</td>
</tr>
<tr>
<td>Density (15°C, g/cm³)</td>
<td>1.0089</td>
</tr>
<tr>
<td>Flash point (°C)</td>
<td>69</td>
</tr>
<tr>
<td>Pour point (°C)</td>
<td>-10</td>
</tr>
<tr>
<td>Viscosity (50 ℃, cSt)</td>
<td>71</td>
</tr>
</tbody>
</table>

Bio-oil was analyzed by GC and GC-MS system. Using acetone as a solvent, qualitative analysis was conducted. Some typical components, such as acetic acid, phenol and alkylated phenols, furan derivatives, furfural and anhydro sugars were identified.

**Char.** Char formed during pyrolysis serves as vapor cracking catalysts to reduce the yield of bio-oil. It can also raise bio-oil viscosity through catalytic reaction during storage, and is likely to be detrimental to most applications. Therefore efficient removal of char is necessary for the production of bio-oil of high quality. We used a hot filter and a cyclone for removal of char. In cyclone, particles with sizes of 10-100 µm were captured. In contrast, the hot filter could catch particle size around 0.1µm. Most of alkali metals in feed was found in char and the C/H ratio of char was in the range of 4 and 5. Figure 4 shows particle distributions of char obtained in hot filter.

**Figure 4.** Particle size distribution of char from hot filter.

**Conclusion**

High quality bio-oil was obtained from rice straw using a fluidized reactor and a char removal system. Optimum reaction temperature for the high yield of bio-oil lay between 410 and 510 °C. Bio-oils obtained in experiments showed very low content of alkali metals, and solid content in bio-oil was about 0.03wt%. In the future, some more works are needed for more yield of bio-oil with smaller size of particle to feed and less water content in feed by drying.

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**References**

VALUABLE AND STABLE CARBON CO-PRODUCT FROM FOSSIL FUEL EXHAUST SCRUBBING

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Introduction

A Sustainable Carbon Sink With Hydrogen Production. The increasing anthropogenic CO2 emissions and possible global warming have challenged the United States and other countries to find new and better ways to meet the world’s increasing needs for energy while reducing greenhouse gas emissions. The need for sustainable energy with little greenhouse gas emissions has lead to demonstration work in the production of hydrogen from biomass through steam reforming of pyrolysis gas and pyrolysis liquids. Our research to date has demonstrated the ability to produce hydrogen from biomass under stable conditions. Future large-scale renewable hydrogen production using non-oxidative technologies will generate co-products in the form of a solid sequestered carbon. This char and carbon (“C”) material represent a form of sequestered C that decomposes extremely slow and retains the bio-capture CO2 for centuries. The limitation of the use of this form of carbon is a profit centric use. It was apparent that additional value needed to be added to this material that would justify large-scale handling and usage. In 1990’s, C in the form of CO2, accumulated at rates ranging from 1.9 to 6.0 Pg C/yr and increasing CO2 levels by 0.9 to 2.8 ppm/yr. The volume of waste and unused biomass economically available in the United States is over 314 gigatons per year. Sequestering a small percentage as valued added carbon could significantly reduce the atmospheric loading of CO2 while simultaneously producing hydrogen. Normally hydrogen is referred to as a zero emissions fuel, however from life cycle perspective it can be viewed as a negative emissions fuel. In order to accomplish this economically, the sequestered C must have a very large and beneficial application such as a soil amendment and/or fertilizer.

The concept of utilizing charcoal as a soil amendment is not new. Man-made sites of charcoal rich soils intermingled with pottery shards and human artifacts have been identified covering 50,000 hectares of the Central Amazon rainforest each averaging 20ha and the largest at 350ha. The radiocarbon dating of the sites have shown ages dating back 740–2,460 years BP and 30% of the soil organic matter is made up of pyrolytic black carbon which is 35 times higher than the adjacent poorer quality soils (Oxisols). Most terra preta sites are identified covering 50,000 hectares of the Brazilian savanna and give evidence to the man-made techniques (slow burning fields and/or fertilizer). In this novel system a portion of the charcoal is sequestered C. This community-based solution operates as a closed loop process, integrating C sequestration, pollutant removal, fertilizer production, increased crop productivity and restoration of topsoil through the return of carbon and trace minerals. The benefits of producing a value added sequestering co-product from coal fired power plants and other fossil energy producing operations, can help bridge the transitions to clean energy systems that are in harmony with the earth’s ecosystem. An important benefit of this approach to the power industry is that it does not require compressors or prior separation of the CO2. The use of biomass in combination with fossil energy production, can allow agriculture, and the agrochemical industry infrastructure to assume a more holistic relationship of mutual support in helping each meet Kyoto greenhouse gas reduction targets.

The char component of the material acts to provide the same benefits as in terra preta sites reducing the leaching of soluble nutrients. This increases plant growth, nutrient uptake and reduces nitrogen runoff. One experimental goal of the project was to identify process parameters that would produce a carbon material optimized for agricultural use. Since it has been shown that the that charcoal addition from 2000 years ago is still providing significant soil fertility benefits and farmers report up to three times crop yields over immediately adjacent non-terra preta soils. Recent research, conducted by Steiner, on the addition of charcoal to non terra preta soils showed significant crop yield increases.

Charcoal has been found to support microbial communities even greater than activated carbon (which is charcoal processed at higher temperatures with steam). Prior researchers assumed that the porous structure provided safe haven but we propose that it may be the availability of microbial nutrients. The processes of pyrolysis create an intrapore deposition of organic polycondensates. The deposition of these materials may increase microbial activity.

Experimental Set-up and Results

SEM Investigation of a Low Temperature Char. The low temperature char particles are hydrophobic in nature and grind easily. The internal gases that escape from the material during the charring help develop charcoals natural porosity. The evolution of this adsorbent material provides a porous internal structure as well. We selected 400 °C as the target temperature for the char before being discharged from the pyrolysis reactor. The resulting char was cooled for 24 hours then fed through a two-roll crushe and then sieved with a mechanized screen through 30 mesh and 45 mesh screens. The resulting fraction remaining above the smaller screen was chosen as our starting material.

SEM Investigation of an Enriched Carbon, Organic Slow-release Sequestering (ECOSS) Fertilizer. Bench scale demonstrations by Oak Ridge National Laboratory recently demonstrated the removal of flue-gas CO2 via formation of solid NH4HCO3 through ammonia carbonation in the gas phase. [23] The results indicated that it is possible to use NH4+ H2O+CO2 solidsifying in process in gas phase to remove greenhouse-gas emissions from industrial facilities such as a coal-fired power plant. A study of agriculturally optimized charcoal produced by Eprida were combined with the above process created via the sequestration of a CO2 stream. It was proposed that the char could act as a catalyst (providing more effective nucleation sites) to speed up the formation of solid NH4HCO3 particles and enhance the efficiency of the gas phase process. A pilot demonstration was constructed to evaluate production and material characteristics of the NH4HCO3-char product. The process also showed promise that it could remove SOx and NOx, enhance sequestration of carbon into soils; providing an ideal “Enriched Carbon, Organic Slow-release Sequestering” (“ECOSS”) fertilizer and nutrient carrier for plant growth. The value would be enhanced if the production of NH4HCO3 could be developed inside the porous carbon media.

To test the production of a coal-based NH4HCO3 fertilizer, we used a mechanical fluidized cyclone, easily adaptable to any gas stream and injected CO2, and hydrated ammonia. A 250g charge of 30-45 mesh 400 °C char was fed in at regular intervals varying from 15-30 minutes. A higher rotor speed increased the fluidization and suspended the particles until they became too heavy from the deposition of NH4HCO3 to be supported by fluidized gas flows.

SEM Investigation of the Interior of an ECOSS-15 Char Particle. The material produced was evaluated by scanning electron microscopy. The examinations revealed The very small molecules of NH3.H2O (hydrated ammonia) are adsorbed into the char fractures and internal cavities. As CO2 enters, it converts the NH3.H2O into the solid NH4HCO3, trapping it inside the microporous material. The SEM’s of the original char and the resulting product clearly evidenced the intra-pore development of the fibrous NH4HCO3 inside the carbon-charcoal framework. The material accumulated as internal flat-top volcano like structures. This demonstration of the process showed that we can deposit nutrients inside the porous media using a low cost gas phase application. We analyzed the required amounts of hydrogen, reformed from biomass, which would be required as ammonia and calculated that 31.6% of the H2 would be necessary for ECOSS production leaving (using all charcoal available from the process) leaving 68.4% of the hydrogen for use as a fuel.

The amount of C, directly converted from exhaust CO2 is equal to 15.2% of the total sequestered carbon as a ammonium bicarbonate and for each 100kg of biomass, we will produce a total of 28.3kg of utilized carbon. In acid soils, this part of the carbon will convert to CO2 but in alkaline soils, (pH=8) it will mineralize and remain stable. According to USDA reports, 60-70% of worldwide farmland is alkaline, so conservatively allowing for 50% of the bicarbonate to convert to CO2, this will leave us with approximately 25 kg of stable carbon in our soils for each 100kg of biomass processed. This carbon represents 91.5 kg of CO2 of which 88% is stored as a very beneficial and stable charcoal. A different way to look at this is to compare the amount of energy produced and the resulting CO2 impact. With 6.78 kg hydrogen extra produced per 100 kg of biomass, then 25/6.78 = 3.69 kg C / kg H or 3.66X3.69 = 13.5 kg CO2 / kg of hydrogen produced and used for energy. From a power perspective, that is 13.5kg CO2 / 120,000 KJ of hydrogen consumed as a renewable energy or 112 kg/GJ of utilized and stored CO2.

Global Potential

The large majority of increases in CO2 will come from developing countries and a sustainable technology needs to be able to scale to meet the growing population needs. The second point is that The energy from biomass systems point of view could create a viable pathway to carbon negative energy as detailed in the IJASA focus on Bioenergy Utilization with CO2 Capture and Sequestration (BECS) [20]. The effects (i.e. providing 112kg of CO2 removal for each GJ of energy used) could allow major manufacturers to offset their carbon costs. For a quick test of reasonableness, if we take the atmospheric rise of 6.19T and divide by 112kg/GJ = 54.5EJ. This number falls amazingly along the 5EJ estimate of the current amount of biomass that is used for energy in the world today. [23] While the potential reaches many times this for the future utilization of biomass, this shows that there is a chance that we can be proactive in our approach. Economic projections of a study based on the ORNL process were compared to this process. Equivalent 10% CO2 reductions and credits needed to return a 13% ROI was estimated to cost $46 million for a 700MW facility utilizing purchased ammonia. However, if the market for nitrogen was an upper limit, and renewable hydrogen were used for producing the worlds ammonia, and all the world's N fertilizer requirements were met from NH4HCO3 scrubbed from power plant exhaust, then the total carbon capture at (1999 N levels) then coal combustion CO2 could be reduced by ~16%. The factors of increased biomass growth with the addition of charcoal as found by Mann [29], Hoshi [29], Glaser [30], Nishio [31], and Ogawa [32] show increase biomass growth from 17% to as 280% with non-optimized char. The direct utilization of an optimized char plus slow release nitrogen/nutrients may allow the increase biomass growth targets worldwide. A portion of this increased biomass growth will be converted to soil organic matter, further increasing C capture (especially if no till management practices are adopted).

The ability to slow down the release of ammonia in the soil will allow plants to increase their uptake of nitrogen. This will lead to a reduction in NOx atmospheric release. For each ton of nitrogen produced, 0.32 tons of C are released, and the 80.95 million tons of nitrogen utilized would represent 26 million tons of C. This is a small a small number in relative terms to the amounts released by combustion of coal (2427 million tons) [33] however if we assume 1.25% of our nitrogen fertilizer escapes into the atmosphere as N2O, then 1.923 million tons of N2O are released, with a CO2 equivalence of 595.9 million tons or 162.5 million tons of CO equivalent.

The economics of hydrogen from biomass has been addressed in the 2001 report by Spath [35]. Our analysis shows that inside plant use of renewable hydrogen would no longer be 2.4-2.8 times the costs from methane, but is approaching 1.6-1.9 times with the increase in natural gas prices. Since market price of nitrogen increases with natural gas prices and this process shows intra plant usage of renewable hydrogen (i.e. no storage or transport expense) becomes significantly more competitive at our current natural gas prices. A review of traditional ammonia processing, shows that due to unfavorable equilibrium conditions inherent in NH3 conversion, only 20-30% of the hydrogen is converted in a single pass. We determined that the ECOSS process could only utilize 31.6% of the hydrogen as we were limited by the total amount of char produced and the target 10% nitrogen loading. This means that a single pass NH3 converter could be used and the expense of separating and recycling unconverted hydrogen is eliminated. The 68.4% hydrogen is then available for sale or use by the power company/fertilizer partnership. This last bracket shows that the ECOSS process thus favors the inefficiencies of ammonia production and reduces costs inherent in trying to achieve high conversion rates of hydrogen.

Conclusions

This concept of biomass energy production with agricultural charcoal utilization may open the door to millions of tons of CO2 being removed from industrial emissions while utilizing captured C to restore valuable soil carbon content. This process simultaneously produces a zero emission fuel that can be used to operate farm machinery and provide electricity for rural users, agricultural irrigation pumps, and rural industrial parks. The use of value added carbon while producing hydrogen (or energy) from biomass can lead to energy with an associated carbon credit (i.e negative carbon energy). With this development non-renewable of carbon dioxide producers

can work with agricultural communities to play a significant part in reducing greenhouse gas emissions while building sustainable economic development programs for agricultural areas in the industrialized and economically developing societies.

Acknowledgements. Kim Magrini, Stefan Czernik, Bob Evans and Calvin Fiek and Don Reicosky and James Lee who have been seeking solutions to our CO2 buildup. Thanks to Nigel Smith, Bruno Glaser, Christoph Steiner, Johannes Lehman, Bill Woods, Bill Denevan and Susanna Hecht who has taught us with the history of terra preta and finally Michael Obersteiner for his encouragement.

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PRODUCTION OF HYDROGEN FROM VEGETABLE OIL

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Introduction

The production of hydrogen from biomass and bio-oils represents a realistic renewable source. The development of the bio-diesel infrastructure could enable hydrogen to be produced from vegetable oil derived materials by either decentralised stationary reformers or on-board fuel processors.

Unmixed steam reforming first proposed by Kumar, Lyon and Cole [1], is an emerging concept for enhancing the steam reforming process. It encompasses the concepts of unmixed combustion, regenerative sorption assisted steam reforming, and is potentially autothermal. It can be operated on a smaller scale to conventional steam reforming at lower pressures. The process operates in two steps forming a cycle. Firstly, an oxygen mass transfer catalyst is oxidised under air flow. In this study, the catalyst is Ni-based and its oxidation results in the formation of NiO compounds. The reaction is exothermic and heats up evenly the reactor bed. During this step, the reactor also contains a carbonate, here a Spanish dolomite (50/50% CaCO3/MgCO3). Some of the heat evolved during the Ni oxidation is used to decompose the carbonate, releasing the CO2. Secondly, the air feed is discontinued and is switched to the fuel/steam mix. Upon contact with the hot catalyst bed, the steam reforming reaction progresses, and CH4, CO, CO2, and H2 are evolved. Under this step, the NiO reduces back to Ni while there is CO2 capture on the adsorbent. The latter shifts the equilibrium of the water gas reaction toward increased hydrogen production. Hydrogen production is therefore intermittent, and for a continuous hydrogen yield, at least two identical reactor beds would need to run in parallel, each operating out of step. The cycle of reactions involved in the unmixed reforming process of an oxygenated hydrocarbon fuel such as vegetable oil is postulated below:

- Under air feed:
  \[ Ni + 0.5 O_2 \rightarrow NiO \quad (\Delta H<0) \]
  \[ MO + CO_2 \rightarrow MO-CO_2 \quad (\Delta H>0) \]
  where M is either Mg or Ca.
  The gas product from this half cycle is an O2-depleted, CO2-rich air stream.

- Under fuel/steam feed:
  \[ (MO+NiO) + fuel_{\text{vij}} + H_2O_{\text{vij}} \rightarrow (MO_3+Ni) + \text{little (CO, CO}_2) + \text{ca.80% H}_2 + H_2O \quad (\Delta H>0) \]
  For sunflower oil fuel, the elemental composition is taken to be \( C_{18}H_{34}O_2 \).

The benefits of this process include an evenly heated reactor bed across its whole cross section, two product streams being either H2 or CO2 rich, low sensitivity to coking, low sensitivity to sulphur poisoning, autothermal behaviour, and, when running on a renewable fuel, no net-CO2 emissions.

Experimental

Following a period of catalyst and CO2-adsorbent screening using a micro-reactor [2], the bench scale demonstration unit of unmixed reforming described in Figure 1 was designed to run on either methane or vegetable oil. The unit operates at pressures between 1-2 bar. The reactor temperature is maintained within the region of 600-800°C, initially by means of an external heating coil, which can be phased out in the event of autothermal operation setting-in.

The liquid flow rates of oil or water are maintained within the range 0-3.2 ml/min depending on conditions. The two feed flows are preheated at 400°C, the temperature at which vegetable oil is completely vapourised. For a more detailed description of the reactor and its operation we refer the reader to [3].

During the fuel step a flow of either nitrogen diluted methane or nitrogen and vapourised vegetable oil mixture pass through the reactor together, this is then switched to air during the oxidative step. When using CH4 and N2 as the fuel feed, the inlet flows are 200 and 600 sccm respectively. When using sunflower oil, the liquid flow is 0.57 ml/min. Various steam to carbon ratios were studied.

The product gases are cooled using a water cooled condenser before entering a water trap followed by a chemical water trap (silica gel). The dry gaseous products are fed through a series of analysers. The gas volume concentrations of CO2, CO, CH4, H2 and O2 were monitored online using analysers from ABB. The gas reaching the analysers was dry and at room temperature.

Figure 1. Schematic diagram of bench scale unmixed reforming reactor.

Results and Discussion

H2 production under fuel feed

![Figure 2. Selectivity to H2 (%) with varying steam/carbon ratio during the methane feed step experiments when there is no CO2 adsorbent in the reactor. Cycle 1 has no steam, cycles 2, 3, and 4 have steam/C of 1.65, 3.3 and 4.95 respectively. Water produced in the reactor makes for the balance to 100 %](image-url)

Because the peak dry H\textsubscript{2} concentrations corresponding to Figure 2 were quite low (10\%, 16\%, 25\%, 27\% respectively), due to the presence of the diluent N\textsubscript{2}, we choose to show how the selectivity to H\textsubscript{2} improved with the presence of steam in the feed. In Figure 3, the effect of reactor temperature for the non-diluted methane feed in the reactor containing the CO\textsubscript{2} adsorbent can be seen on the dry H\textsubscript{2} concentration.

![Figure 3](image-url)

**Figure 3.** Typical dry gas composition of gaseous products during the fuel step for:
7a. methane at 650\ºC, Steam/C of 4.
7b. methane at 600\ºC, Steam/C of 4.

Figure 3 a-b show that CO\textsubscript{2} and CH\textsubscript{4} dry gas concentrations are significantly reduced to the benefit of that of H\textsubscript{2} when the average reactor temperature of the fuel step is lowered from 650\ºC to 600\ºC. This indicates increased adsorption of CO\textsubscript{2} and the expected effect of shift in the water gas reaction. The extent to which CO\textsubscript{2} adsorption is maintained on a continuous basis is dependent upon the extent of regeneration of the adsorbent achieved during the previous oxidation cycle.

![Figure 4](image-url)

**Figure 4.** shows the effect of steam carbon ratio on the peak dry H\textsubscript{2} concentration when using a nitrogen (558 sccm) and sunflower oil (0.57 ml/min) mix in a reactor containing just the Ni catalyst, in the absence of CO\textsubscript{2} adsorbent.

The points shown in Figure 4 are the results of experiments carried out on three different days on the same catalyst. In spite of this, a clear trend can be picked up whereby the maximum dry H\textsubscript{2} concentration is obtained for steam carbon ratios between 2 and 4, as was found for methane fuel. However, the selectivity to H\textsubscript{2} when using sunflower oil was of the order of few percent, hardly comparable to the methane experiments. This was concurrent with a production of water which was of the same order as the amount as the water reactant.

**Catalyst redox cycles under air and fuel feed.** Before we could attempt to operate the process autothermally, a major hurdle had to be overcome, namely the ability of the catalyst to reduce under the fuel feed and oxidize at similar rates under the air feed. Our first experiments with methane fuel indicated a rate of reduction two orders of magnitude slower than the oxidation rate, hence the subsequent experiments with sunflower oil were operated with an equivalent fuel molar rate two orders of magnitude larger than the methane. This had several effects. The reduction and oxidation rates were now both of the order of 10\textsuperscript{4} mol/s, but for a small adjustment. In counterpart, using a much larger fuel flow without changing the air feed had the effect of increased carbon formation by two orders of magnitude as well (order 10\textsuperscript{4} mol/s). This increased carbon formation was an additional oxygen sink to the nickel catalyst during the air feed which tended to produce CO and CO\textsubscript{2} in similar concentrations while it burned at a rate ca. 10\textsuperscript{5} mol/s. Under the previous lower methane flow, hardly any CO had evolved during the air feed, while all products of oxidation were NiO and CO\textsubscript{2}. Clearly, further optimization of the flows need to be carried out before the timing of the flows can be adjusted for autotherm operation.

**Conclusions**

During the methane feed step, the main dry gas product was hydrogen. However, solid carbon is the main carbon product throughout the whole fuel feed step, while water is the principal hydrogen containing product during an initial phase. A much improved selectivity to H\textsubscript{2} product (up to 80\%) is achieved when introducing steam in the feed. During the air feed step, the oxidation of Ni to NiO is slowed down by the carbon deposited during the previous fuel step. When operating with an adsorbent, the production of hydrogen is increased although the adsorption of CO\textsubscript{2} is sensitive to the bed temperature.

When using vegetable oil at a rate two orders of magnitude larger than in the methane experiments, chosen in order to achieve the desired NiO reduction rate, carbon formation is increased in proportion. The dry gas H\textsubscript{2} concentration is larger than with methane but the selectivity to H\textsubscript{2} much lower. The NiO reduction rate under fuel feed and the Ni oxidation rate under air feed are of similar magnitude, suggesting both feeds need little adjustment in the next optimisation phase of the process. Future work will focus on achieving autothermal operation in the reactor while maximizing the H\textsubscript{2} yield and the separation of the CO\textsubscript{2} from the H\textsubscript{2} product stream.

**Acknowledgement.** The work presented here was financially supported by the Engineering and Physical Science Research Council (EPSRC). The authors are also grateful to Mr Gavin Rickett and Dr Phil Ingram at Johnson Matthey.

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SUSTAINABLE HYDROGEN FROM BIO-OIL - CATALYTIC STEAM REFORMING OF ACETIC ACID AS A MODEL OXYGENATE

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Introduction

Hydrogen as a fuel has recently been an important issue due to developments in fuel cell technology. Hydrogen production from sustainable sources, e.g. biomass, is gaining attention for a CO2 neutral energy supply. Recent developments in flash pyrolysis technologies make it possible to convert lignocellulosic biomass efficiently to a bio-oil, which is easier for handling and transport. The bio-oil so generated contains a variety of aliphatic and aromatic oxygenates (aldehydes, ketones, acids, alcohols).

Combination of catalytic steam reforming and water gas shift reaction is the most promising option to generate hydrogen since these reaction steps maximize hydrogen yield. During steam reforming of bio-oil, deactivation due to coke/oligomer deposition on the catalysts is a severe problem. It has been reported that deactivation is unavoidable even in the presence of excess steam (H2O/C > 5). Economically, lower steam/carbon (S/C) ratio (< 2) is preferable. In this study, Pt/ZrO2 has been investigated since earlier work has shown excellent activity and stability for Pt/ZrO2 under conditions favorable for coke formation, e.g., even in the absence of steam, during CO2 reforming of natural gas.

Bio-oil is a mixture of oxygenates. In order to provide a knowledge base for the design of active and stable catalysts for steam reforming of bio-oil, studies have been initiated with model oxygenates. Acetic acid (HAc) has been chosen as a model compound in this study because it is a major component of bio-oil (up to 32 wt.%). It is known that steam reforming and water gas shift reaction (CO + H2O → CO2 + H2) occur simultaneously and the overall stoichiometry is represented (Equation 1),

\[
CH_3COOH + 2H_2O → CO_2 + 4H_2 \quad \ldots(1)
\]

This study describes attempts to understand the reaction and deactivation mechanisms in order to establish guidelines for the design of efficient catalysts for steam reforming of bio-oil.

Experimental

Catalyst preparation. Pt/ZrO2 was prepared by wet impregnation technique. Pt loading was set to 0.5 wt%. The catalyst was calcined at 925 K for 15 h.

Catalytic measurements. The amounts of 50 mg of catalysts were loaded in a fixed bed reactor and held by quartz wool plugs. The catalyst was first reduced in 5% H2 in N2 at 925 K for 1 hr. An aqueous solution of HAc giving a steam to carbon molar ratio (S/C) of 5 was introduced by using a microfeder (KD Scientific) and a syringe (Hamilton co.). Gas mixture, HAc/H2O/N2/Ar, was fed to give a ratio of 3/30/5/82 ml min\(^{-1}\) (N2 as an internal standard). For analysis, a flame ionization detector (FID) and a Hayesep Q column were used for HAc, acetone, ketene, and hydrocarbons; a thermal conductivity detector (TCD) and a Carbosieve column were used for H2, N2, CO, CH4, and CO2.

Pulse experiments were performed with 20 mg of Pt/ZrO2. After reduction at 925 K, the system was purged in Ar at a flow rate of 37.5 ml min\(^{-1}\). Pulses of HAc aqueous solution (S/C = 5, 3.5 µmol of HAc) at 293 K were introduced.

The hydrogen yields were calculated based on Equation 1. For carbon containing compounds, the yields were calculated based on C1 equivalent values.

Catalyst characterization. Thermogravimetry (TG) (Mettler Toledo TGA/SDTA 851E) was used to determine the amount of deposits in the catalysts.

IR spectra were recorded in situ under vacuum in transmission mode (Bruker, Vector 22, MCT detector). The catalyst reduced ex situ at 925 K was mounted in the IR cell and firstly reduced in situ by introducing 0.1 mbar of H2 at 725 K. After evacuation, oxygenates and steam were introduced into the system with an S/C molar ratio of 5 at a total pressure of 0.1 mbar at 725 K. Each spectrum consists of 32 scans taken at 4 cm\(^{-1}\) resolution.

Results and Discussions

Figure 1 shows HAc conversion and yields of products during HAc/H2O reaction over Pt/ZrO2. Conversion was close to 100% and constant for 3 h, however, yields of products changed with time. In the beginning (5 min), H2 and CO2 were the main products, CH4 and CO were observed in small quantities. H2 yield (47%) was close to equilibrium value (56%). The product yields and pattern indicates that the catalyst was active for steam reforming. However, after 25 minutes time on stream, H2 and CO2 yields decreased drastically, and an increase in acetone was observed.

![Figure 1](https://example.com/figure1.png)

**Figure 1.** Conversion and yields for steam reforming of HAc over Pt/ZrO2 catalyst. (725 K, S/C = 5, GHSV = 160000 h\(^{-1}\)).

Figure 2 shows HAc conversion and yields of products during HAc/H2O reaction over ZrO2 (without Pt). HAc conversion was close to 90%. The conversion of HAc and yields of the products were constant for 3 h. However, no steam reforming activity (H2 and CO) was observed, and only acetone and CO2 were observed as products. HAc is known to convert to acetone easily and the reaction is catalyzed by a variety of oxides. ZrO2 is also known as such an oxide. The stoichiometry of the products observed (acetone and CO2, 1:1) correspond to the ketonization reaction (Equation 2).

\[
2CH_3COOH → CH_3COCH_3 + CO_2 + H_2O \quad \ldots(2)
\]

Both Pt/ZrO2 and ZrO2 were very active for HAc conversion. However, H2 and CO, i.e., steam reforming products were produced only over Pt/ZrO2 and not over ZrO2. This implies that presence of Pt is essential for steam reforming. ZrO2 shows acetone yields similar to those observed over Pt/ZrO2 after 25 minutes time on stream (Figure 1), indicating that even after the active sites for steam reforming were deactivated, the accessible ZrO2 sites still provided the activity to form acetone and CO2.

For Pt/ZrO2 and ZrO2, TG experiments showed approximately 1.2 and 0.8 wt% deposits, respectively, after 3 hour test. Regeneration in air restored catalytic activity, indicating that blocking active sites by deposits caused deactivation of the Pt/ZrO2 catalyst for the steam reforming.

Presence of acetone in the product mixture and formation of deposits on \( \text{ZrO}_2 \) indicates a role for acetone in catalyst deactivation. Acetone is known to undergo aldol type condensation and give products such as diacette alcohol, mesityl oxide (MO) and MO is known to form oligomers easily (\( \text{CH}_3\text{COCH}_3 \rightarrow (\text{CH}_3)_2\text{C(OH)}\text{CH}_2\text{COCH}_3 \rightarrow (\text{CH}_3)_2\text{C}=\text{CHCOCH}_3 \rightarrow \text{oligomers} \)). In order to check if these oligomers play a role in deactivation, kinetic and \textit{in situ} IR measurements were carried out with \( \text{Pt/ZrO}_2 \) pre-exposed to MO.

Table 1 shows comparison between HAc conversion and \( \text{H}_2 \) yield with or without MO treatment during pulse experiment. It is seen that \( \text{Pt/ZrO}_2 \) catalyst deactivated for \( \text{H}_2 \) formation (steam reforming) more rapidly when contacted with MO in comparison to HAc.

**Table 1. Comparison of HAc Conversion and \( \text{H}_2 \) Yield between Fresh and MO Treated Pt/\( \text{ZrO}_2 \)**

<table>
<thead>
<tr>
<th>Catalyst (Pt/( \text{ZrO}_2 ))</th>
<th>HAc conv. (%)</th>
<th>( \text{H}_2 ) yield (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fresh</td>
<td>98.2</td>
<td>60.0</td>
</tr>
<tr>
<td>MO treated(^a)</td>
<td>82.4</td>
<td>29.1</td>
</tr>
</tbody>
</table>

\(^a\)MO/\( \text{H}_2\text{O} \) pulse (0.32 \( \mu \)mol-MO, \( \text{S/C} = 5 \)) was introduced before a HAc/\( \text{H}_2\text{O} \) pulse.

\textit{In situ} IR spectra recorded over Pt/\( \text{ZrO}_2 \) are shown in Figure 3. In the presence of HAc/\( \text{H}_2\text{O} \) (Figure 3a) characteristic peaks for linear CO adsorbed on Pt at 2060 cm\(^{-1} \) were observed. This confirms the production of CO in the \textit{in situ} experiment. Additionally bands due to oxygenates (acetate, carbonate) derived from HAc (1541, 1426 and 1405 cm\(^{-1} \)) were also seen. Exposure-evacuation cycles with HAc/\( \text{H}_2\text{O} \) were repeated three times (not shown) and no significant difference in IR spectra were observed.

After pre-exposure to MO, \textit{in situ} IR spectra were recorded with HAc/\( \text{H}_2\text{O} \) (Figure 3b). The spectra showed similar oxygenate peaks on the support but the CO peak was very weak. This implies the activity towards CO formation (steam reforming) was deactivated as expected from kinetic experiments. However, when CO was introduced in the gas phase, the intensity of the peak for CO significantly increased (Figure 3c). This indicates that the Pt surface was not blocked completely even though the activity for steam reforming was lost. This further implies that only part of the surface Pt atoms are responsible for steam reforming. Since oligomerization reactions of MO occur on the support, and the fact that Pt sites were deactivated with MO, we conclude that the Pt sites active for steam reforming are at the close vicinity of the support. This is in agreement with a bi-functional mechanism proposed by us for the steam or CO\(_2\) reforming of methane, where methane is activated on the metal and steam or \( \text{CO}_2 \) on the support, the reaction occurring at the metal support boundary.\(^,\)\(^5\) Thus, \( \text{ZrO}_2 \) may have a key role in the mechanism for steam reforming in the activation of water. Once MO, or the oligomers derived from it, is formed on the support even in small amounts, they could block the boundary sites and deactivate the Pt/\( \text{ZrO}_2 \) catalyst.

**Figure 3. In situ IR spectra over Pt/\( \text{ZrO}_2 \) at 725 K (Steam/Carbon = 5, 0.1 mbar). (a) HAc/\( \text{H}_2\text{O} \). (b) HAc/\( \text{H}_2\text{O} \) after MO/\( \text{H}_2\text{O} \) treatment and evacuation. (c) \( \text{CO} \) treatment after (b) without evacuation.**

The acid-base properties of the support oxides play a key role in catalyzing the condensation reactions.\(^5\) \( \text{ZrO}_2 \) has an important role in steam reforming \textit{via} steam activation. However, condensation reaction could also happen over \( \text{ZrO}_2 \). Thus, design of a stable catalyst should be aimed at improvement of the support, minimizing the condensation reactions as well as enhancing the steam activation capacity.

**Conclusions**

The Pt/\( \text{ZrO}_2 \) catalysts showed high activities at initial time on stream, but lost its activity for steam reforming (\( \text{H}_2 \) production) rapidly.

Results obtained in the study point to a bifunctional mechanism for steam reforming of HAc on Pt/\( \text{ZrO}_2 \), i.e., HAc activation takes place on Pt and steam activation on \( \text{ZrO}_2 \). The reaction occurs at the Pt periphery in close proximity of \( \text{ZrO}_2 \). The catalysts deactivate due to blocking of active sites by oligomers derived from HAc/acetone on \( \text{ZrO}_2 \). In order to develop a durable catalyst, the oligomerization/condensation reactions which take place on the support need to be minimized.

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**References**

DETERMINATION OF THE OXIDATION CHARACTERISTICS OF FIR (Abies bornmuelleraiana) WOOD

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Introduction

About 90% of the energy used by humankind is based on combustion – mainly combustion of fossil fuels (oil, coal, and natural gas), with a relatively small contribution from the combustion of biomass (wood). The main exceptions to combustion are nuclear fission and hydro for the generation of electricity. Energy conversion by combustion has a significant impact on the planet’s atmosphere by means of ground-level air pollution (photochemical smog, acid rain, global climate change, and stratosphere particulate matter, SO2, CO, toxic organic gases and vapors and toxic metals), acid rain, global climate change, and stratosphere ozone depletion.

Although energy production from biomass residues is limited due to availability and high cost compared to fossil fuels, biomass provides a renewable source of energy on a scale sufficient to play a significant role in the development of sustainable energy programmes, so vital if we are to create a greener, more ecologically sensitive society.

By optimal utilization of biomass resources and development of new technical advances, biomass will become more efficient and competitive as an energy source. In order to consider the optimum conditions for biomass utilization one should start investigating the combustion characteristics of the biomass. In this study we aimed to find the combustion characteristics of wood samples as the starting point for the evaluation of to use as an energy source.

Experimental

The wood sample used in this work was a bark-free fir (Abies bornmuelleraiana) sawdust sample obtained from Bolu forests (northwest Anatolia) in Turkey. The proximate and elemental analyses of the wood sample were done at the Instrumental Analysis Laboratory of the Scientific and Technical Research Council of Turkey, Ankara, is given in Table 1. The sawdust was ground and sieved to below 175 μm (80 mesh) size.

Table 1. Proximate and Elemental Analyses of the Fir Wood Sample

<table>
<thead>
<tr>
<th>Proximate analysis, % (as received)</th>
<th>85.5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Volatile matter</td>
<td>85.5</td>
</tr>
<tr>
<td>Fixed carbon</td>
<td>10.5</td>
</tr>
<tr>
<td>Moisture</td>
<td>3.7</td>
</tr>
<tr>
<td>Ash</td>
<td>0.3</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Elemental analysis, % (daf)</th>
<th>47.2</th>
</tr>
</thead>
<tbody>
<tr>
<td>C</td>
<td>47.2</td>
</tr>
<tr>
<td>H</td>
<td>6.1</td>
</tr>
<tr>
<td>N</td>
<td>0.3</td>
</tr>
<tr>
<td>O (by difference)</td>
<td>46.7</td>
</tr>
</tbody>
</table>

Characterization Techniques. Combustion and decomposition characteristics of the wood sample were analyzed by using a Netsch 449C thermogravimetric analyzer (TGA). TGA analysis was done in the following manner; the wood samples were heated up to a temperature range between 500°C and 1000°C with different heating rates of 5K/min, 10K/min, 20 K/min and 30 K/min. In addition to this, TGA analysis was combined with an Bruker Equinox 55 FTIR in order to monitor the released gases during combustion and decomposition. The ashes obtained from TGA analysis were examined with a Leo G34-Supra 35VP scanning electron microscope (SEM) and SEM images were compared with the images obtained from unburnt wood sample. Additionally the wood sample was objected to heat treatment at different temperatures between 100°C and 400°C. The mass loss according to the heat treatment was recorded and these samples were analyzed with an IKA-Calorimeter System-C4000-Adiabatic calorimeter, again included the comparison with the untreated sample.

Results and Discussion

The heating rate seemed to be very effective on the oxidation of the samples and different mass losses were observed. There were two main temperatures for mass losses; as the heating rate was increased the greatest mass loss was detected at higher temperatures.

Table 2. Percent Mass Loss and Change of Calorific Values

<table>
<thead>
<tr>
<th>Heat Treatment Temperature, °C</th>
<th>Mass Loss, %</th>
<th>Calorific Value, kJ/kg</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unheated</td>
<td>-</td>
<td>18746</td>
</tr>
<tr>
<td>100°C</td>
<td>5.9</td>
<td>19135</td>
</tr>
<tr>
<td>200°C</td>
<td>11.0</td>
<td>19521</td>
</tr>
<tr>
<td>300°C</td>
<td>32.0</td>
<td>3149</td>
</tr>
<tr>
<td>400°C</td>
<td>99.3</td>
<td>-</td>
</tr>
</tbody>
</table>

While heat treatment of the wood sample up to 200°C increased the calorific value of the wood due the removal of the low volatile compounds. As the heat treatment temperature was increased to 300°C and higher temperatures due to the pyrolytic losses of carbonaceous material from the structure of the wood the calorific values decreased sharply.

The FTIR spectra of the volatiles obtained during TGA measurements are presented in Figures 2-4. The peaks due to CO2 and H2O can be seen near 2335 cm⁻¹ and 3240 cm⁻¹, respectively and a complex peak near between 1500- and 1700 cm⁻¹ is due to H2O.

Figure 2. FTIR spectrum of the volatiles evolved at the 14th second during TGA measurement.

Figure 3. FTIR spectrum of the volatiles evolved at the 1506th second during TGA measurement.

Figure 4. FTIR spectrum of the volatiles evolved at the 2041st second during TGA measurement.

SEM micrographs of the wood dust before and after oxidation are presented in the Figure 5 and 6. Fibrous structure of the wood was destroyed and a structure with a relatively greater porosity seemed to form after the oxidation experiments.

Conclusions
The effect of heat treatment on the calorific values and structural changes of a wood sample before its oxidation was investigated by using a TGA system and scanning electron microscope. Heat treatment up to 200°C increased the calorific value of the wood. Structural changes during oxidation reactions observed indicated the loss of volatiles and production of porosity on the surface of the residue. The work is in progress and it is expected that the complete set of analysis of the material evolving during oxidation and changes in the morphology of the residual material of wood are being investigated in detail.

References
THE EFFECT OF ANTIOXIDANT ADDITION ON NO\textsubscript{x} EMISSIONS FROM BIODIESEL

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Introduction

Although the use of biodiesel offers many environmental advantages over petrodiesel, an increase in NO\textsubscript{x} emissions from biodiesel combustion, relative to levels observed from petrodiesel combustion, has been reported by several researchers. (1-3) This increase is of concern in areas that are subject to strict environmental regulations. For universal acceptance of biodiesel, it is desirable to reduce these NO\textsubscript{x} emissions at least to levels observed with petrodiesel combustion.

NO\textsubscript{x} can be formed by two major pathways during diesel fuel combustion: the Zeldovich (thermal) mechanism and the Fenimore (“prompt”) mechanism. The rate of the Zeldovich reactions is flame temperature-dependent, whereas the Fenimore pathway is more complex. In the Fenimore mechanism, free radicals formed from the fuel react with N\textsubscript{2} to eventually form NO\textsubscript{x}. This takes place very early in the combustion process and is partly dependent upon the fuel radical concentration and how it is established. (4)

Some data have indicated that the chemical makeup of biodiesel can influence the amount of NO\textsubscript{x} formed during its combustion. McCormick et al. (5) published data that correlated the properties of various biodiesels to NO\textsubscript{x} emissions. One notable result was a strong positive correlation between the iodine value of the fuel (a function of the number of double bonds) and NO\textsubscript{x}; as iodine value increased, so did the NO\textsubscript{x} emissions. These data suggest a role of double bonds in elevating NO\textsubscript{x} output. Sites of unsaturation are known to have a greater propensity for radical formation than do carbon-carbon single bonds. It is conceivable that this increase in free radical generation is at the heart of the increase in NO\textsubscript{x} output by some biodiesels. If so, a quenching of free radicals during combustion may reduce or eliminate this NO\textsubscript{x} elevation. Antioxidants are known to be free radical quenching agents.

Acting on this postulate, McCormick et al. (6) added a common antioxidant, tert-butyl hydroquinone (TBHQ), to biodiesel as a possible method of NO\textsubscript{x} reduction. They observed a slight reduction in NO\textsubscript{x} and suggested that the use of antioxidants as a NO\textsubscript{x} mitigation strategy deserved more study. Moreover, the addition of antioxidants to biodiesel is being investigated to increase its storage stability and it is important to understand the impact that such addition may have on NO\textsubscript{x} formation.

Experimental

Materials. Certified diesel fuel was obtained from Chevron Phillips Chemical Company (Houston, Texas). The fuel used was low sulfur (365 ppm), with a cetane number of 46.8. The biodiesel used was Soygold, a methyl soyate biodiesel purchased from Ag Environmental Products (Omaha, Nebraska) in one-gallon units to protect the fuel from extensive oxidation. 2-Ethylhexyl nitrate (2-EHN), 2,2′-methylenebis(6-tert-butyl-4-methylphenol), vitamin E, tert-butylhydroquinone, propyl gallate, diphenylamine, and butylated hydroxyanisole were purchased from Sigma-Aldrich Chemical Co. (St. Louis, Missouri). Structures of the antioxidants are shown in Figure 1.

The fuel used had a blend composition of 80% petrodiesel and 20% biodiesel (B20). Additives were tested at a level of 1000 ppm. 

Figure 1. Structures of selected additives. (I) 2-ethylhexyl nitrate. (II) 2,2′-methylenebis(6-tert-butyl-4-methylphenol) (III) Vitamin E (IV) TBHQ (V) Propyl gallate (VI) Diphenylamine (VII) BHA

Method. A Yanmar L100 single cylinder, four stroke, naturally aspirated, direct injection diesel engine from Bowers Power Systems (Kent, Washington) was installed and instrumented for NO\textsubscript{x} measurements. The engine had a compression ratio of 20:1, a displacement of 0.418 L, and a fuel injection timing of 17±0.5 degrees before top dead center (6TDC). The engine was connected to an electrical generator. Load was added to the engine by using the generator to power work lights. The results discussed here are obtained at a load of 5 kW (82% maximum load) and an engine speed of 3200 rpm. These conditions were chosen to generate NO\textsubscript{x} levels high enough to allow small changes in NO\textsubscript{x} emission levels to be detected.

NO\textsubscript{x} and NO\textsubscript{2} were measured using single gas monitors (model CA-CALC) manufactured by TSI Instruments ( Shoreview, Minnesota). These monitors have electrochemical sensors as opposed to the chemiluminescence methods normally employed for NO\textsubscript{x} detection. The monitors were chosen for simplicity and affordability, and were calibrated frequently using calibration gases. Exhaust temperature was monitored using a thermocouple from Omega Instruments (Stamford, Connecticut) inserted into the exhaust gas stream.

Since the engine was minimally instrumented with no provisions for the control of such variables as inlet air temperature and humidity, the baseline numbers were generally not repeatable on a daily basis. Therefore, for each test, NO\textsubscript{x} emission levels from petrodiesel, B20, and B20 with the additive were measured against those fuels without the additive. An experiment began by bringing the engine to steady state using certified diesel fuel. The speed of the engine was monitored using a tachometer. The engine was held at a constant speed and load for the duration of the test and the NO\textsubscript{x} emissions of the petrodiesel were measured. A minimum of triplicate measurements were made to ensure an accurate NO\textsubscript{x} measurement. The fuel was then switched to B20 containing 1000 ppm of the antioxidant additive again without disrupting engine operation. The engine was allowed to come to steady state and the NO\textsubscript{x} levels were measured again. Experiments were performed to ensure that when the fuel was switched back to B20, the same results as prior to running the additized blend were achieved. The NO\textsubscript{x} emission levels for the additized fuel were compared to the NO\textsubscript{x}
levels measured for B20 in that experiment only. The percent changes in NO\textsubscript{x} emissions from experiment to experiment were compared.

**Results and Discussion**

The engine system employed here proved effective and reliable as a screening method to determine if antioxidant fuel additives have an effect on NO\textsubscript{x} levels. Candidate additives that show promise with such a system will be tested at a rigorous engine test facility in the future.

It was necessary to confirm that the reported difference in NO\textsubscript{x} emissions between biodiesel and petrodiesel could be observed. As shown in Table 1, a 6.2% increase in NO\textsubscript{x} emissions was observed when B20 was used as opposed to petrodiesel. The increase observed here was fairly consistent throughout the tests and is in agreement with results reported in the literature. (1-3)

It was also of interest to establish that reduction in NO\textsubscript{x} by the use of accepted NO\textsubscript{x}-lowering treatments could be attained. To accomplish this 2-EHN was added to the fuel at a 1000 ppm level. 2-EHN is known to be an effective additive for reducing NO\textsubscript{x} emissions at this level in a variety of diesel engines. (7,8) This additive increases the cetane number of the fuel, which leads to shorter ignition times that result in decreased NO\textsubscript{x} emissions. Again, the results observed here corroborate those seen by others; the addition of 2-EHN at 1000 ppm reduced NO\textsubscript{x} by 4.5%. This observation suggests that this engine system responds typically to a known NO\textsubscript{x} reducing agent.

The antioxidants were chosen based on several considerations. Several are readily available and inexpensive since they are used in the food industry to prevent lipid oxidation. Additionally, Mittelbach and Schober (9) performed a study in which antioxidants were rated for their effectiveness in terms of fuel stability. Antioxidants that they found to be effective (such as n-propyl gallate and TBHQ) were tested in hopes that these additives would serve two functions, namely increasing storage stability and reducing NO\textsubscript{x}.

Table 1 lists the effect of the addition of each of six antioxidants on NO\textsubscript{x} emissions during B20 combustion. The antioxidants used had minimal effect on NO\textsubscript{x} emissions, but the results from three antioxidants were notable.

| Table 1. Effect of Additive Addition on NO\textsubscript{x} Emissions |
|-----------------------------|-------------------------|
| Fuel                        | Change in NO\textsubscript{x} from B20 Levels |
| Diesel Fuel (B0)            | -6.2%                   |
| B20 + 2-EHN                 | -4.5%                   |
| B20 + 2,2’-methylenebis(6-tort-buty1-4-methylphenol) | +0.2% |
| B20 + Vitamin E             | +0.3%                   |
| B20 + TBHQ                  | -0.9%                   |
| B20 + Propyl Gallate        | -0.4%                   |
| B20 + diphenylamine         | +0.7%                   |
| B20 + BHA                   | -4.4%                   |

TBHQ induced a 0.9% reduction in NO\textsubscript{x} emissions when added to B20 at 1000 ppm, but the data from this particular additive had a great deal of scatter. Reductions as high as 8.7% were observed, but other tests showed an increase as great as 5.3%. TBHQ was the only additive that exhibited such erratic behavior.

One possible explanation for this behavior may lie in the storage stability of the antioxidants. Although the antioxidants were stored in a cool, dry place and out of light, they were not stored under inert gas and may have reacted with oxygen, thus reducing their effectiveness. The possibility of pre-oxidation must be investigated further.

Diphenylamine was notable in that it consistently increased NO\textsubscript{x} emissions, though the increase was small (+0.7%). It is likely that this increase might have been due to the nitrogen present in the additive. Although this increase may not appear significant, it should be considered when selecting an antioxidant for fuel storage stability purposes.

Butylated hydroxyanisole (BHA) was the most effective of the antioxidants studied, giving a 4.4% decrease in measured NO\textsubscript{x} emissions. This decrease is approximately the same as was observed with an identical amount of 2-ethylhexyl nitrate. We conclude that the addition of BHA to the B20 does work to combat the “prompt” NO\textsubscript{x} that is formed during the combustion process.

BHA and TBHQ have similar structures, as shown in Figure 1. Decreases in NO\textsubscript{x} levels observed with TBHQ addition, however, were not repeatable. The similarity in structure of BHA and TBHQ suggests that structure may play a role in determining antioxidant activity and effectiveness in reducing NO\textsubscript{x} emissions.

**Conclusions**

The system described in this work was designed for screening additives that may potentially reduce NO\textsubscript{x} emissions from biodiesel. Despite minimal instrumentation, it was possible to observe changes in NO\textsubscript{x} levels with changes in fuel formulations. While more thorough engine testing will be required to confirm the results reported here, the system we used represents a relatively inexpensive approach to examining the effect of fuel additives on NO\textsubscript{x} production by biodiesel fuels.

The use of butylated hydroxyanisole (BHA) should be investigated further as this additive did produce a small but significant decrease in NO\textsubscript{x} emissions. TBHQ also should be investigated. Blends of additives should be investigated to look for synergistic effects.

**References**


Functionalization of nickel on 4-aminothiophenol modified glassy carbon electrodes and the enhanced electrochemical activity towards the oxidation of ethanol

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Introduction
Organized self-assembled monolayers (SAMs) on various solid supports have attracted much attention in recent years and are commonly used to impart desired chemical or physical properties to surfaces. SAM’s offer the possibility to control the film structure at the molecular and supramolecular level and therefore to realize the tailoring of the surface properties. Monolayers of organic thiols, especially those formed of alkanethiols, are the most studied self-assembled monolayer system since they offer stability and high degree of organization. Monolayers formed from aromatic thiols, although not that intensively studied, are of considerable interest since they offer together with the properties given by aliphatic thiols the additional advantage of having a high electrical conductivity. Their use as surface modification agents opens up new possibilities to arrange metal particles on modified electrode surfaces with supposedly enhanced electrochemical activity.

It is well-known from the literature that the thiol-containing group of any aliphatic and aromatic compound can be adsorbed onto Pt, Cu and Ag. This might be an interesting approach to the formation and arrangement of metal particles on aryl modified electrode surfaces. Nickel, showing a high affinity to the amine group in nickel amine complexes, is then expected to be functionalized onto the 4-aminothiophenol modified electrode. Nickel has been reported to be a good catalyst for the oxidation of various organic compounds. We have chosen in our studies ethanol as compound with the aim to investigate the electrochemical activity of nickel in this oxidation process.

Experimental
Glassy carbon electrodes (AIMCOR, Pittsburgh, grade GC-10) were prepared from 3 mm diameter rods embodied into epoxy resin (Hysol, 56C) and were used as working electrodes. Platinum gauze and a Ag/AgCl (saturated KCl) electrode were used as counter and reference electrodes, respectively. All potentials were reported versus the Ag/AgCl reference electrode. The glassy carbon electrode surface was cleaned by polishing with 1 µm and 0.05 µm alumina slurry. After polishing, the electrode was washed with water and ultrasonicated for 5 min in acetonitrile. Electrochemical modification of the glassy carbon electrode was carried out in acetonitrile containing 1 mM aminothiophenol and 0.1 M NBu3BF4. The applied potential was 2.0 V for the deposition of 4-aminothiophenol. After the modification, the electrode was thoroughly washed by rinsing in acetonitrile and then in deionized water. A solution containing 97.6 wt% methanol, 2 wt% acetic acid and 0.4 wt% nickel(II)chloride hexahydrate (reduced to Ni zero state by 0.4 M sodium borohydride) has been prepared in which the modified electrode has been soaked for 20 minutes at 45-50 °C to functionalize nickel. After rinsing with deionized water again, electrochemical measurements of the nickel prepared modified electrode were carried out at room temperature (18-23°C) in 1 M NaOH and in further experiments with the addition of ethanol using various concentrations. Before each experiment the solution was sparged for 10-15 minutes with a stream of highly purified nitrogen to displace dissolved oxygen. Electrochemical measurements were performed in a one-compartment cell using the three-electrode configuration. Cyclic voltammetry was performed using either a potentiotstat/galvanostat model M263A (EG&G Princeton Applied Research) or an electrochemical interface SI1287 (Solartron Instruments) interfaced with a PC and the electrochemical set-ups were both controlled with the model DC Corrware (Scribner Associates, version 1.2) software.

Results and Discussion
Figure 1 shows the typical cyclic voltammetry in 1 M NaOH at 50 mV/s of nickel loaded onto 4-aminothiophenol corresponding to 1.8 x 10^-9 mol/cm². Initially the GC electrode surface (0.07 cm² electrode area) was modified with 4-aminothiophenol at 2.0 V vs Ag/AgCl for 240s. The voltammetry of nickel is characterized by reasonably well-understood surface reactions corresponding to the formation and stripping of hydroxide films at the metal surface. Starting on the anodic scan we can clearly note in figure 1 the presence of a well-established peak at 0.43 V attributable to the formation of NiOOH, and on the reverse scan the corresponding peak due to the reduction of the anodically formed NiOOH occurs at 0.34 V. More anodically the oxygen evolution arises at the onset potential of 0.55 V. The voltammetric features shown in this figure are comparable to those reported in the literature for ordinary nickel electrodes and dispersed nickel on modified electrode surfaces in basic solution. With the characteristics outlined in this figure we can clearly confirm the presence of nickel functionalized on the modified GC electrode. One could argue that the deposition of nickel might occur onto the glassy carbon surface and this may then be responsible for the voltammetric features given in figure 1. This assumption is rather unlikely since recent studies showed complete blocking of the electrochemistry on the modified electrode when using redox-active species so that any deposition of nickel onto GC in the presence of 4-aminothiophenol grafted onto the electrode is more to be doubted. This should therefore give strong indication that the presence of nickel in our voltammetry is only due to the surface reactions on the nickel film functionalized on the modified electrode surface.

![Graph](attachment:image.png)

**Figure 1.** Cyclic voltammetry in 1 M NaOH at 50 mV/s of nickel functionalized onto the 4-aminothiophenol modified GC electrode surface (0.07 cm² electrode area). 4-Aminothiophenol as surface modification agent was grafted at 2.0 V vs Ag/AgCl for 240s onto the glassy carbon (GC) electrode.

To give further evidence that nickel is present on the modified GC electrode and to make these studies conclusive, figure 2 shows the XPS core level spectrum of Ni2p obtained after the functionalization of nickel onto aminothiophenol. We restrict our studies to the Ni2p3/2 region since this will provide us with sufficient information. Curve fitting of the Ni2p3/2 gives different Ni species with binding energies of 855.9±0.1 and 857.2±0.1 eV. On the basis of their binding energies these peak values were assigned to Ni(OH)2 and NiOOH, respectively, with the ratio in area underneath the peak of 0.92. The Ni2p3/2 spectrum shows at higher binding energies (862±0.2 eV) an intense satellite signal which may be attributed to a multielectron excitation (shake-up peaks). It is relevant to point out that at lower binding energies below 854 eV the XP signal for nickel metal and NiO does not appear in this spectrum. This fact might indicate that Ni is fully covered by an hydroxide layer after its functionalization onto the modified electrode. Further support for this was given by the O1s spectrum. The fact that nickel was shown to be functionalized onto the 4-aminothiophenol electrode surface opens up new possibilities for further investigations in which we were particularly interested in the oxidation of ethanol on nickel.

![Figure 2](image)

**Figure 2.** X-ray photoelectron Ni2p core level spectrum for nickel functionalised onto the 4-aminothiophenol modified electrode surface.

Directing our attention towards figure 3 we notice on following the anodic scan in voltammogram a) a peak at 0.53 V with its corresponding peak on the reverse scan at the same potential. This experimental finding is consistent to that reported in the literature about the ethanol oxidation process on nickel electrodes so that we found our experimental results attributable to this electrochemical process. On continuing the anodic scan we see the start of the oxygen evolution with the onset potential of 0.68 V. Comparing these voltammetric features to those shown in voltammetry b) and c) we hardly note any difference except the fact that higher current densities were obtained with increasing ethanol concentration. This issue is essential, and we certainly want to stress this fact that with increasing the ethanol concentration in figure 3 fast kinetic processes and negligible shift of about a few mV in peak potential for the ethanol oxidation process occur when comparing the voltammograms and thus a clear separation between this process and the oxygen evolution is still be seen even at high ethanol concentrations (5M). These are remarkable features and are clearly in contrast to the results shown in published material on ordinary nickel electrodes and nickel based alloys dispersed on a modified electrode. Here the electrochemical surface reactions are accompanied by an increase in solution resistance near the electrode surface leading then at high ethanol concentrations to an ohmic drop. Kim and Park showed this behaviour in their work when using Ru-Ni alloys describing that the peak for the ethanol oxidation at higher concentrations (>0.5 M) is rather indistinguishable from the oxygen evolution in the voltammetry. In addition, we want to emphasize the fact that we obtained reasonable current densities for the ethanol oxidation process with a low amount of nickel loaded onto the modified electrode. Comparable electrochemical activity was obtained when using almost 300 fold less quantity of nickel as catalyst material. Running the ethanol oxidation constantly at 0.5 V vs Ag/AgCl in 1 M NaOH using the 4-aminothiophenol modified electrode on which nickel was loaded revealed appreciable current densities (>5mA/cm²). With the focus on further improvement we might therefore consider this electrode as one alternative to the Pt based alloy electrodes for the anodic oxidation of ethanol in fuel cells.

![Figure 3](image)

**Figure 3.** Set of cyclic voltammmograms at 50 mV/s in 1 M NaOH with increasing concentration of ethanol from 0.5 M a), 2 M b) to 5 M c) of nickel functionalised onto the 4-aminothiophenol modified GC electrode surface. 4-Aminothiophenol as surface modification agent was grafted at 2.0 V vs Ag/AgCl for 240s onto the GC electrode. In this voltammetry the same nickel prepared electrode was used as in figure 1

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**References**

Kinetics, Characterization and Mechanism for the Selective Dehydration of Ethanol to Diethyl Ether over Solid Acid Catalysts

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Introduction

Ethanol is a clean burning alternative automotive fuel to gasoline\(^1\). However, one serious disadvantage of ethanol is that it has a lower vapor pressure (-3 psia at 38 °C) than winter grade gasoline (~11 psia at 38 °F)\(^2\). A time-honored standard in the fuel industry mandates that fuels have sufficient vapor pressure to cold-start an automobile at temperatures as low as ~30 °C. This is not possible with ethanol. One promising way to solve this problem is to convert a portion of the ethanol into a more volatile compound, such as diethyl ether, on board the vehicle prior to or during the cold-start operation.

The main objective of this study is to develop a catalyst material that exhibits acceptable activity and selectivity for the dehydration of ethanol to diethyl ether (DEE) under the severe operating conditions that exist on board an automobile. Aliphatic alcohols, with exception of methanol, have two modes of dehydration, bimolecular dehydration to produce ethers and intra molecular dehydration to olefins, see equations (1) and (2).

\[
\begin{align*}
2\text{CH}_3\text{CH}_2\text{OH} & \rightarrow \text{CH}_3\text{CH}_2\text{OCH}_2\text{CH}_3 + \text{H}_2\text{O} \\
\text{CH}_3\text{CH}_2\text{OH} & \rightarrow \text{CH}_3=\text{CH}_2 + \text{H}_2\text{O}
\end{align*}
\]

Both dehydration reactions are known to proceed over solid acid catalysts\(^3\). However, ethylene is undesirable, since it contributes to automotive pollution and catalyst fouling. Diethyl ether formation is thermodynamically favorable over a wide range of temperatures, including the 50-500 °C range commonly employed in catalytic processes. However, the formation of ethylene is also thermodynamically favored and predominates at temperatures above 100 °C. Operating a solid acid catalyst at temperatures below 100 °C results in the selective formation of diethyl ether, but the reaction is kinetically limited and gives rates too slow for an automotive application. In order to overcome this dilemma, a catalyst is needed that can selectively produce diethyl ether at temperatures in excess of 100 °C. In this study, such a catalyst is reported and a possible mechanism for its observed selectivity for diethyl ether production is discussed.

In order to design an on board catalytic dehydration reactor, it is necessary to determine the required fuel properties of a ternary mixture of ethanol, water, and diethyl ether for cold-starting a vehicle. This information is needed in order to determine the % ethanol conversion that is required from an on-board catalytic reactor. This also plays an important role in selecting the appropriate catalyst material and the optimum reaction conditions in our laboratory studies. A significant amount of information has been published regarding ethanol as a fuel\(^2\), but the literature is nearly void of information regarding the diethyl ether assisted cold-starting of a vehicle\(^5\). Vapor-liquid equilibrium phase diagrams, for ternary mixtures of ethanol, diethyl ether, and water, at various temperatures, have been reported by Kito-Borsa and Cowley\(^6\) using ASPEN Plus software. Their results suggest that ethanol conversions between 40 and 85% at 100 % diethyl ether selectivity is required to produce an acceptable fuel mixture that can cold-start a vehicle at temperatures down to ~30 °C without forming two immiscible liquid phases.

Highly acidic ion exchange resins catalyze the dehydration of ethanol almost exclusively to diethyl ether over a temperature range of 80-120 °C\(^7-8\). Although these materials are highly selective, the narrow temperature range and low reaction rates severely limit their viability for any cold-start application. At temperatures above 120 °C, the particles begin to fuse together causing a dramatic increase in pressure drop through the catalyst bed or a complete blockage of flow\(^9\). Karpuk and Cowley\(^10\) reported the dehydration of methanol to dimethyl ether (DME) and water occurs readily over fluorinated γ-alumina. Amorphous silica-alumina, γ-alumina, and alumina-phosphoaluminates (APA) also gave good yields. In a review of hydrocarbon formation, Chang\(^1\) discusses the possible mechanisms for DME formation. Depending on the catalyst type, Lewis acid sites, Bronsted acid sites, or both appear to be involved in the methanol dehydration reaction through a complex set of surface reactions. Since all of these catalysts are highly active for methanol dehydration, they may also be suitable for ethanol dehydration as well. The selectivity of these materials for ether versus ethylene production has not been investigated.

Experimental

The following experimental procedures were used in the evaluation of the catalyst composition and the catalyst performance.

**Catalyst Preparation.** The aluminophosphate-alumina (APA) catalysts were prepared using a conventional coprecipitation method. In general, the APA catalysts were prepared by combining an aqueous solution of aluminum nitrate and orthophosphoric acid with an ammonium hydroxide solution to form a solid hydrogel. The gel was washed with ammonium hydroxide solution, dried at 120 °C for 24 hours, and calcined at 500 °C for 16 hours. The calcined materials were sized to 20/40 mesh. An attempt was made to prepare P/AI ratios of 0.0, 0.1, 0.5, 0.8 and 1.0 by varying the amount of orthophosphoric acid added. The actual P/AI ratios were determined by ICP analysis. The PS-1, PS-2 and PS-3 catalysts were prepared by impregnating 20/40 mesh sized silica gel (Davison Chemical grade 57) with a solution of phosphoric acid. The PA-1, PA-2 and PA3 catalyst was prepared by impregnating 20/40 mesh sized γ-alumina (Norton SA-6273) with a solution of phosphoric acid. The silica-alumina was provided by Davison Chemical (Grade 980-13).

**Catalyst Evaluation.** The catalyst evaluations were carried out under a variety of temperatures in a continuous-flow, fixed bed catalytic reactor, using 1.27 cm o.d. stainless steel tube. Ethanol was injected via syringe pump, then vaporized at 120 °C and mixed with a He (99.99999%) carrier gas. The partial pressure of ethanol in the feed was 0.48, and the space velocity was 2080 cm\(^3\)/cat-hr. Approximately, 0.5 g of catalyst was mixed with 1.5 g of quartz chips to insure uniform flow and minimize temperature gradients in the catalyst bed. Each catalyst was tested at 200, 250, and 300 °C for 3 hours at each temperature. The product gas was analyzed by an on-line gas chromatograph (SRI 8610B) using a TC detector and fitted with a 1.5 mm column packed with Porapak Q.

**Catalyst Characterization.** The XRD patterns of the catalysts were collected with a LINT 2000 diffractometer by Rigaku using a Cu K\(_\alpha\) source at an applied voltage of 40 kV and a current of 150 mA. The radial distribution functions (RDF) of the amorphous APA catalysts were obtained by collecting the x-ray scattering data in the same instrument using a Mo K\(_\alpha\) source with an applied voltage of 40 kV and a current of 200 mA.

The \(^{27}\)Al MAS NMR spectra were measured at 130 MHz with a CMX Infinity 300 NMR spectrometer. The sample spin speed was

14 kHz. The spectra were recorded with π/4 excitation with a pulse width of 1.0 μsec, and a pulse delay of 0.1 second. All of the 27Al chemical shifts were referenced to a 1 M aqueous solution of aluminum nitrate.

The XPS studies were performed on a Kratos HSi instrument using Al Kα radiation and a band pass of 20 eV.

Temperature programmed desorption (TPD) results were obtained by placing approximately 0.3 g of the catalyst sample in a quartz cell. The sample was pretreated by heating it to 525 °C for 3 hours in a helium flow of 20 cm³/min. Ammonia was adsorbed onto the catalyst sample at 175 °C by flowing a 1:1 volume mixture of anhydrous ammonia and helium through the cell for 30 min. After adsorption was complete, the sample was heated at a rate of 20 °C/min. in flowing helium until a final temperature of 500 °C was reached. The desorbed ammonia was passed through an scrubber containing 20.0 mL of standard 0.05 N sulfuric acid solution. The total amount of ammonia desorbed was determined by titrating the acid solution with a 0.05 N standard solution of sodium hydroxide. Pyridine was introduced into the helium flow using a saturator maintained at 0 °C. The adsorption period for pyridine was 1 hour.

Results and Discussion

The catalysts used in this study were selected because they have varying surface densities of Lewis and Brønsted acid sites. Gamma-alumina (GA) has predominantly Lewis acidity, silica-alumina (SA) has both Lewis and strong Brønsted acidity, alumina-phosphatized-alumina (APA) has both Lewis and weak Brønsted acidity, phosphoric acid on silica (PS) has only Brønsted acidity, and phosphoric acid on γ-alumina (PA) has both Lewis and weak Brønsted acidity. The catalyst compositions are reported in Table 1.

Table 1. Bulk Composition of Catalysts

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<tr>
<th>Catalyst</th>
<th>Wt. % SiO₂</th>
<th>Wt. % Al₂O₃</th>
<th>Wt. % P₂O₅</th>
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* All of the P₂O₅ is present on the surface of the catalyst, and is not distributed throughout the bulk of the sample.
** As determined by ICP analysis.

Catalyst Evaluation. The activity and selectivity were calculated using the following equations. Where Xₐ, Xₐ, Xₐ, and Xₐ represent the mole fractions of ethanol in the feed, and ethanol, dimethyl ether, and ethylene in the product. The catalyst test results are presented in Table 2. A blank run was made using only quartz chips in the catalyst tube. The absence of any ethanol conversion showed that catalytic wall or thermal effects were absent. The conversions for all catalysts were typically less than 10% activity at 200 °C. At temperatures of 250 and 300 °C, only the APA catalysts (0.5-1.0) give activity and selectivity sufficient to cold start a vehicle. A kinetic study was made using the APA 0.5 catalyst. The following rate expression gave the best fit to the data.

\[ \text{Rate} = \frac{kP_E}{(1 + K_dP_E + K_dP_B)} \]  

The adsorption constant for water is 436 and that for ethanol is 26.7, which implies that a strong product inhibition by water exists. In summary, the APA catalysts exhibit the desired activity and selectivity for the ethanol dehydration reaction. In order to gain a better understanding of the surface chemistry responsible for this desired result, selected catalysts were analyzed in more detail in order to understand the possible cause of this selectivity.

Table 2. Catalyst Activity and Selectivity

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>250 °</th>
<th>300 °</th>
</tr>
</thead>
<tbody>
<tr>
<td>GA</td>
<td>52.5</td>
<td>93.0</td>
</tr>
<tr>
<td>SA</td>
<td>79.3</td>
<td>71.0</td>
</tr>
<tr>
<td>APA 0.0</td>
<td>61.6</td>
<td>93.5</td>
</tr>
<tr>
<td>APA 0.1</td>
<td>23.3</td>
<td>96.2</td>
</tr>
<tr>
<td>APA 0.5</td>
<td>54.8</td>
<td>98.0</td>
</tr>
<tr>
<td>APA 0.8</td>
<td>55.3</td>
<td>98.1</td>
</tr>
<tr>
<td>APA 1.0</td>
<td>51.5</td>
<td>97.7</td>
</tr>
<tr>
<td>PS-1</td>
<td>0</td>
<td>2.9</td>
</tr>
<tr>
<td>PS-2</td>
<td>2.0</td>
<td>100</td>
</tr>
<tr>
<td>PS-3</td>
<td>2.4</td>
<td>100</td>
</tr>
<tr>
<td>PA-1</td>
<td>70.1</td>
<td>92.7</td>
</tr>
<tr>
<td>PA-2</td>
<td>61.9</td>
<td>94.9</td>
</tr>
<tr>
<td>PA-3</td>
<td>51.4</td>
<td>97.1</td>
</tr>
</tbody>
</table>

Catalyst Characterization. Selected catalysts were analyzed by XRD, RDF, NMR, and TPD. Figure 1 shows the XRD patterns for the APA catalysts. The APA 0.0 catalyst contains no aluminophosphate and gives a diffraction pattern similar to the commercial γ-alumina sample (GA). This was the expected result. The addition of even a small amount of phosphate to the structure inhibits the crystallization of the γ-alumina phase. See the XRD pattern for the APA 0.1 catalyst. The addition of more phosphate results in a pattern similar to that of amorphous silica at around 23 degrees 20.

Figure 1. XRD patterns for APA 0.0, 0.1, 0.5, 0.8, and 1.0 catalyst samples.
Radial distribution function (RDF) analysis can be used to investigate amorphous materials. A comparison of silica glass, APA 0.1 and APA 1.0 are shown in figure 2. The Al-O (1.74 Å) and P-O (1.74 Å) bond distances are consistent with a tetrahedral configuration, similar to that observed for Si-O (1.61 Å). The variation in the Al-O and P-O bonds is likely responsible for the broader peak observed for the APA 1.0 sample. The APA 0.1 sample gives an Al-O (1.88 Å) bond distance. This value is essentially identical to the Al-O (1.9 Å) bond length reported for an aluminum octahedron. This result suggests that the addition of phosphate forces more aluminum atoms to adopt a tetrahedral configuration.

Figure 2. RDFs of Silica, APA 0.1 and APA 1.0.

The chemical shifts for the MAS NMR spectra for the $^{27}$Al atoms in samples of GA, APA 0.1, APA 1.0, PA-1, and PA-3 are given in table 3. For the GA sample, the $^{27}$Al atoms are present in 4-coordinate (tetrahedral) sites at 67.4 ppm and 6-coordinate (octahedral) sites at 91.1 ppm, see figure 3. The observed octahedral to tetrahedral ratio of 2.0 : 0.9 is typical for $\gamma$-alumina. As phosphate is added to the structure, a new 5-coordinate site appears at 34.0 ppm, apparently at the expense of tetrahedral sites. The APA 1.0 sample shows a dramatic change in the NMR spectra, with all three $^{27}$Al peaks being shifted to the right, and a significant increase in the 4-coordinate species. This is in agreement with the RDF data. It is likely that this new species plays a role in making the APA catalysts more selective for diethyl ether production. The PA-1 and PA-3 samples gave NMR spectra very similar in peak area and chemical shifts to that for $\gamma$-alumina. This is expected since the phosphate is only located at the surface and the preponderance of the NMR signal comes from the $\gamma$-alumina support.

Table 3. Chemical Shifts for NMR Spectra of $^{27}$Al.

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>4-coord.</th>
<th>5-coord.</th>
<th>6-coord.</th>
</tr>
</thead>
<tbody>
<tr>
<td>GA</td>
<td>67.4</td>
<td>0</td>
<td>9.1</td>
</tr>
<tr>
<td>APA 0.1</td>
<td>71.3</td>
<td>34.0</td>
<td>5.2</td>
</tr>
<tr>
<td>APA 1.0</td>
<td>44.1</td>
<td>17.6</td>
<td>-9.6</td>
</tr>
<tr>
<td>PA-1</td>
<td>67.4</td>
<td>0</td>
<td>8.3</td>
</tr>
<tr>
<td>PA-3</td>
<td>69.0</td>
<td>0</td>
<td>9.9</td>
</tr>
</tbody>
</table>

A decrease in the ratio of pyridine to ammonia desorption suggests a decrease in the population of Lewis acid sites. The pyridine to ammonia ratio decreases for the APA catalysts as the phosphate

Figure 3. $^{27}$Al MAS NMR Spectra of GA, APA 0.1, and APA 1.0 catalyst samples.

XPS analysis of the APA samples was done to determine if there was a direct correlation between the surface and bulk phosphorus to aluminum (P:Al) mole ratio. No enrichment of phosphorus on the surface of the APA samples was observed.

Ammonia and pyridine TPD studies provide information about the relative amounts of Lewis and Brønsted acid sites on the catalyst surfaces. Ammonia is known to adsorb onto both Lewis and Brønsted acid sites, while pyridine absorbs only onto Lewis acid sites. Since Brønsted sites of $\gamma$-alumina, Al-OH, are too weak to protonate pyridine, Lewis acidity is thought to be exclusively responsible for the observed acidity for $\gamma$-alumina. Silica is known to have very little acidity. Table 4 gives the TPD results for the desorption of ammonia and pyridine from our catalyst samples.

Table 4. Relative amounts of ammonia and pyridine desorption from catalyst samples.

<table>
<thead>
<tr>
<th>Catalyst Sample</th>
<th>Ammonia Desorbed</th>
<th>Pyridine Desorbed</th>
<th>Pyridine to Ammonia Des.Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>GA</td>
<td>0.89</td>
<td>1.69</td>
<td>1.91</td>
</tr>
<tr>
<td>SA</td>
<td>0.55</td>
<td>0.48</td>
<td>0.86</td>
</tr>
<tr>
<td>APA 0.0</td>
<td>0.76</td>
<td>1.27</td>
<td>1.67</td>
</tr>
<tr>
<td>APA 0.1</td>
<td>0.55</td>
<td>0.74</td>
<td>1.36</td>
</tr>
<tr>
<td>APA 0.5</td>
<td>2.12</td>
<td>1.10</td>
<td>0.52</td>
</tr>
<tr>
<td>APA 0.8</td>
<td>2.59</td>
<td>0.86</td>
<td>0.33</td>
</tr>
<tr>
<td>APA 1.0</td>
<td>3.04</td>
<td>0.74</td>
<td>0.24</td>
</tr>
<tr>
<td>Silica</td>
<td>0.04</td>
<td>0.03</td>
<td>0.57</td>
</tr>
<tr>
<td>PS-1</td>
<td>0.34</td>
<td>0.29</td>
<td>0.86</td>
</tr>
<tr>
<td>PS-3</td>
<td>2.04</td>
<td>0.74</td>
<td>0.36</td>
</tr>
<tr>
<td>PA-1</td>
<td>0.91</td>
<td>1.60</td>
<td>1.77</td>
</tr>
<tr>
<td>PA-2</td>
<td>0.88</td>
<td>1.52</td>
<td>1.73</td>
</tr>
<tr>
<td>PA-3</td>
<td>0.99</td>
<td>1.46</td>
<td>1.47</td>
</tr>
</tbody>
</table>

content increases. Considering these observations, it is apparent that the addition of phosphate decreases the surface concentration of Lewis acid sites. This has important implications with respect to the surface dehydration mechanism. It is proposed that the dehydration of ethanol to ethylene requires two adjacent Lewis acid sites or strong Brønsted acid sites. The APA catalysts dilute the surface concentration of Lewis acid sites without adding strong Brønsted acid sites. The dehydration of ethanol to diethyl ether requires only a single Lewis acid site.

Conclusions
The unique acidic properties of the aluminophosphate-alumina catalysts make them highly selective for the dehydration of ethanol to diethyl ether at elevated temperatures (200-300 °C).

Acknowledgement. This work was funded by DOE and NREL. Catalyst support samples were provided gratis of Grace (Davison Division) and Norton Chemical Companies.

References
EFFECTS OF NITROGEN CHEMISTRY ON CHAR FORMATION FROM BIOMASS

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Introduction

The fate of nitrogen during pyrolysis or combustion is an important consideration in the utilization of biomass. Nitrogen oxide emissions from biomass combustion are thought to arise primarily from fuel-bound nitrogen and an understanding of the chemical transformations involving nitrogen may lead to improvements in these emissions. An important source of nitrogen in biomass is proteins and amino acids, which can react catalytically with the carbohydrates by reactions generally characterized as Maillard chemistry. This chemistry produces complex mixtures of products and has been the subject of numerous studies. However, less is known about the role that this chemistry plays in the formation of char. The partitioning of nitrogen into char or gas phase products will be critical towards determining the routes for converting nitrogen from biopolymers into NOx emissions.

We have studied reactions of amino acids with carbohydrates and how these reactions affect the composition of the resulting char. We have used Molecular Beam Mass Spectrometry (MBMS) to characterize the gaseous emissions from the pyrolysis of model mixtures of amino acids with carbohydrates. The remaining char has been characterized using elemental analysis and reflectance FTIR spectroscopy.

Of particular interest in the char formation are effects due to the Maillard chemistry involving proline. We have shown that proline is more reactive in Maillard chemistry than other amino acids, which is consistent with literature reports of proline as a catalyst for asymmetric aldol condensations. In particular, we would like to investigate how proline affects the maturation of char.

Experimental Techniques

Experiments were conducted by sequentially heating samples of model compounds or biopolymers in flowing helium. Gas phase products were measured directly using an MBMS, and the remaining solid materials were retained for further analysis. Samples (100 mg to 400 mg) were placed in a quartz boat and slid into a heated tubular reactor. The furnace was set so that sliding the sample further into the reactor produces higher temperatures. Sequential heating experiments were conducted by sliding the sample into a fixed temperature region (as determined by a thermocouple attached to the boat) and holding for five minutes. A subsequent heating was conducted by sliding the material further into the reactor and holding it at that higher temperature for five minutes.

Results and Discussion

Figure 1 shows integrated mass spectra for staged heating experiments using proline mixed with a carbohydrate matrix composed of glucose, cellulose and pectin (Pro:Glu:Cell:Pect = 1:1:2:2). This mixture was used because the glucose will react vigorously with the glucose and the cellulose/pectin mixture produces a significant amount of char. In these experiments, the samples are heated to 550 °C directly (bottom spectrum) or sequentially (top two spectra). In the spectrum on the top, the sample was heated to 170 °C for 5 min, then to 350 °C for 5 min. before being heated to 550 °C. In the middle spectrum the sample was heated to 350 °C before being heated to 550 °C. The mass spectra in this figure are obtained by integrating over the pulse that occurs during the last temperature of each given sequence.

The intensity of the high m/z peaks are shown in the insets and the scales for these are on the right side of the plots. As can be seen, there are characteristic clusters of peaks in the high m/z portion of these spectra. For instance, there are clusters around m/z = 200, 214, 227 and 242. These spectral clusters are also observed for other amino acids but are not as prominent as for proline. Furthermore, the location of these clusters seems to be dependent upon the amino acid but not the type of carbohydrate used. These peaks appear to be associated with Maillard reactions between proline and the carbohydrates and probably result from secondary reactions. As can be seen from these spectra, the formation of these high m/z clusters of peaks is enhanced when the sample is not thermally pretreated. They are more intense in the bottom spectrum than in the middle spectrum and more intense in the middle spectrum than in the top spectrum. This suggests that low temperature reactions of the proline with the carbohydrates tend to decrease the formation of these secondary Maillard products.

Figure 1. Mass spectra from the staged heating of proline mixed with a matrix of glucose, cellulose and pectin. These spectra have been averaged over triplicate experiments.

The infrared spectra of the char also shows interesting changes in the solid that result from Maillard reactions. Fig. 2 shows a plot of
the FTIR spectra of the char from the pyrolysis of our carbohydrate matrix without proline. The lower spectrum is of the unheated matrix. The peaks at 3300 cm\(^{-1}\) (O-H stretch), 2890 cm\(^{-1}\) (C-H stretch), 1330 cm\(^{-1}\) (C-H bend) and 1050 cm\(^{-1}\) (C-O stretch) are characteristic of carbohydrates. At moderate pyrolysis temperatures (170°C and 350°C) carbonyl peaks are formed (1750 cm\(^{-1}\) and 1630 cm\(^{-1}\), C=O stretch). At 550°C peaks arising from aromatic groups are seen at 3040 cm\(^{-1}\) (C-H stretch), 1590 and 1436 cm\(^{-1}\) (C-C stretch), 1230 cm\(^{-1}\) (C-H in-plane bend) and 815 cm\(^{-1}\) (C-H out-of-plane bend).

Similar peaks are seen in the char FTIR spectra shown for proline (Fig. 3) mixed with the carbohydrate matrix. However, the presence of the proline leads to maturation of the char at lower temperatures. For neat matrix experiments there is still significant intensity for the carbohydrate peaks at 350°C, while in the presence of proline, these peaks are essentially gone at 350°C. For the mixture including proline the carbonyl peaks form at 170°C but disappear at 350°C, where they are replaced by aromatic peaks. This again shows the ability of proline to accelerate the char maturation process. Other amino acid can also accelerate the formation of aromatic groups in char but not to the extent that occurs with proline.

Measurement of char yields from the sequential pyrolysis experiments and elemental analysis of the remaining solid material allow a determination of the partitioning of nitrogen between the solid and gas phase. The plot in Fig. 4 shows this partitioning for sequentially heating a mixture of proline and the carbohydrate matrix to 170°C, 350°C and 550°C. This plot shows that the nitrogen has a tendency to stay in the char.

![Figure 4](image)

**Figure 4.** Partitioning of elements during sequential heating of proline mixed with the carbohydrate matrix (Glu+Cell+Pect). The composition in the lower left is for the starting material and the values along the bottom are for the char after heating. The values at the top are for the gas phase after heating.

**Conclusions.** The conclusions of this work are that proline can drastically change the rate of formation of mature char (highly aromatized) through its vigorous reaction with carbohydrates. Further, this Maillard chemistry appears to leave significant amounts of the nitrogen in the char.

**Acknowledgment.** We would like to acknowledge the support of Philip Morris USA, and stimulating discussions with Geoffrey Chan and Mohammad R. Hajirolig.

**References**
NMR STUDY OF THE TRANSFORMATIONS IN
PECTIN AND CELLULOSE (WITH 10% N-15
PROLINE) DURING PYROLYSIS

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Introduction
In preparation for the analysis of tars made from a tobacco
plant grown from N-15 enriched nutrients several chars made from
pectin and cellulose (with 10% by weight of N-15 enriched proline
added) have been studied by C-13 and N-15 NMR and ESR
methods. The carbon skeletal structure was analyzed in terms of the
previously described structural and lattice parameters. Only a
few of the highest temperature chars (550 and 600 °C) start to show
a small amount of graphite like structures. The spin lattice
relaxation time in both the laboratory and rotating frame were
measured for all samples and showed correlation with the
carbonization of the samples.

Experimental
All C-13 NMR spectra were run on a Chemagnetics CMX-100
spectrometer operating at a carbon frequency of 25.152 MHz and
using a 7.5 mm probe with a ceramic housing for reduced carbon
background. The proton decoupling power was 62.5 kHz. The
spinning speed was locked to 4.1 kHz. Chemical shift ranges used
in references 1 and 2 were modified to somewhat account for the
overlap of the aromatic region with the anomeric carbon in
carbohydrates in some of the samples. The contact time chosen for
the dipolar dephasing experiments was five times the longest TCH as
measured in variable contact time experiments.

Nitrogen-15 spectra were run on a Chemagnetics CMX-200
spectrometer also with a 7.5 mm probe. This field was chosen,
instead of the higher S/N spectrometer operating at 400 MHz, so
that the formation of pyridinic type nitrogens with large chemical
shift anisotropies could be more accurately followed.

All chars were prepared at Philip Morris and heated to
temperatures of 150 to 600 °C.

Results and Discussion
Carbon-13 CPMAS spectra of some pectin chars are shown in
Figure 1. The pectin resonances are the larger broader resonances at
about: 171 ppm for the carboxyl carbon, 101 ppm for the anomeric
carbon, 70 ppm for carbons next to one oxygen and 54 ppm for the
methoxy group. The five resonances from proline are the smaller
sharper lines. As can be seen in the spectrum from the sample
heated to 150 °C proline is the first to react. A small amount of
aromatic and carboxyl carbons have been produced where there
were none in the original mixture. Also the aliphatic part of the
spectra has become broad in the region where the sharp proline
resonances were before. At 250 °C most of the pectin has also
reacted and the spectrum looks like a lignite coal. A large amount
of aromatic carbons have been formed but the sample still retains a
wide range of oxygen functional groups: carboxyls, carboxyls,
phenolics and oxygenated aliphatic carbons. There may still be a
small amount of anomeric carbons under the aromatic band at
about 105 ppm. These may be able to be separated from the

![Figure 1. C-13 CPMAS spectra of the pectin and proline mixture heated to the listed temperatures.](Image 328x434 to 543x729)

aromatic carbon by a csa type filtering experiment. At 350 °C the
original material has mostly reacted and the oxygen functional
groups have greatly decreased along with some aliphatic material.
As the temperature is increased further the samples become more
and more aromatic. At 550 °C the aromaticity is 0.95. The average
aromatic cluster size is estimated to be about 9 in the 150 °C sample
where the aromatic material first starts to appear and progressively increases to about 19 carbons in the 550 °C sample.

Some important NMR relaxation parameters for the pectin/proline samples are presented in Table 1. One can see that
after some scatter in the starting sample and the 150 °C sample that
the T1 values decrease as the temperature that the sample was
heated increased. Others have seen this relaxation time go through a minimum and then increase when the electron spin exchange rate
changes and the relaxation is not proportional to the number of
static unpaired electrons. The spin-lattice relaxation in the rotating
frame, T1, also decreases as the temperature is increased, reaches
a minimum at 450 °C, and then starts to increase for the 550 °C
sample. This is a common occurrence that has been seen in other
pyrolysis series.

Table 1. NMR Parameters from Pectin Char.

<table>
<thead>
<tr>
<th>Sample</th>
<th>$T_1^H$ (s)</th>
<th>$T_1^H$ (ms)</th>
<th>$T_1^L$ (ms)</th>
</tr>
</thead>
<tbody>
<tr>
<td>RT</td>
<td>0.188(76%)</td>
<td>4.97(24%)</td>
<td>--</td>
</tr>
<tr>
<td>150 °C</td>
<td>0.729(94%)</td>
<td>10.9(6%)</td>
<td>0.38</td>
</tr>
<tr>
<td>250 °C</td>
<td>0.271</td>
<td>5.4</td>
<td>0.67</td>
</tr>
<tr>
<td>350 °C</td>
<td>0.131</td>
<td>2.7</td>
<td>0.79</td>
</tr>
<tr>
<td>450 °C</td>
<td>0.027</td>
<td>2.1</td>
<td>1.3</td>
</tr>
<tr>
<td>550 °C</td>
<td>0.017</td>
<td>8.4</td>
<td>1.7</td>
</tr>
</tbody>
</table>

The last column in Table 1 is the Lorentzian (exponential) time constant from a dipolar dephasing experiment. This time constant represents dephasing from protons that are not directly attached to aromatic carbons. The dipolar dephasing experiment used here was sequence C from reference 7 that does not include dephasing from chemical shift anisotropy. As one can see as the temperature increases the time constant $T_1^L$ gets larger as there are fewer protons or the protons are farther away from the nonprotonated aromatic carbons.

Conclusions

This preprint shows that NMR can be used to quantify the changes in the structure in a pectin/proline mixture heated to various temperatures. Additional work is currently under way to study the nitrogen incorporation in the various chars and to measure the unpaired electron spin concentration and see how it correlates to the various NMR relaxation time constants. Additional samples heated between 150 °C and 250 °C are currently being examined because this is the temperature range where the largest amount of structural modification seems to occur.

Acknowledgement. This work was supported by Phillip Morris, USA.

References


PROBING THE PYROLYSIS OF METAL DOPED CELLULOSE CHARS AT DIFFERENT POINTS ALONG THE REACTION COORDINATE


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Golden, CO 80401-1887

Introduction

Cellulose is a major component of biomass, along with hemicellulose and lignin, and is a useful model for thermo-chemical conversion of biomass. The conversion of biomass via pyrolysis has important applications in the development of new renewable chemicals and bio-carbons, as well as combustion processes leading to combined heat and power, forest fires and cigarette smoking. The process of cellulose and cellulose char pyrolysis is still not fully understood despite rapid progress in the last couple of decades. The influence of metal salts on charring and subsequent pyrolysis has long been known to alter the chemistry of these processes. Metal cations occur naturally in biomass particularly in the hemicellulose, they may also be added to the growing plant in the form of fertilizer. In general inorganic materials in the form of ash are considered a problem that must be mitigated against in the pyrolysis and combustion of biomass. However, metal cations may be deliberately added to the biomass to alter its pyrolysis chemistry in a favorable way to reduce the formation of polycyclic hydrocarbons, PAH, or in the synthesis of new bio-carbons.

In order to more fully understand the effect of metal doping on cellulose pyrolysis chemistry we have doped Avicel cellulose with a series of mono- and di-valent cations. Both main group and transition metals were chosen in order to probe both the effects of promoting ionic pathways and possible catalytic pathways in the charring and pyrolysis chemistry. We chose to dope cellulose with metal acetates as not only is the acetate anion relatively inert but all of these metal acetates decompose below the charring temperature of 375 °C. The chars were characterized primarily by DRIFTS and NMR. The char structure and pyrolysis chemistry at very short times, several ms was studied by laser pyrolysis molecular beam mass spectroscopy, LP-MBMS. We also sampled the product gases and trapped the free radicals produced using cold and chemical trapping at slightly longer times, several 100 ms.

Experimental

Avicel cellulose (5 g) was doped at 1mol% with sodium, potassium, magnesium, calcium, palladium, copper and zinc acetates by the incipient wetness technique using aqueous solutions of the as received metal acetates. The doped cellulose samples were dried in an oven at 115 °C overnight pressed into a 1 in diameter pellet at 10,000 psi in a Carver press and charred under Ar(g) at 375 °C for 1/2 h.

For the DRIFTS experiments the chars were ground in air and placed in a Harrick Scientific, heatable variable atmosphere chamber in a Harrick Scientific Praying Mantis Diffuse Reflectance attachment. The chars were heated under dry He(g) for 10 mins at 400 °C and then cooled to room temperature at which point the DRIFTS was recorded using a Thermo-Nicolet Nexus 670 FT-IR spectrometer equipped with a N2(g) cooled MCT detector. The data was treated with a Kubelka-Munk transformation to enable its quantitative interpretation.

The details of the LP-MBMS experiment have been described previously. After cooling, the metal doped cellulose chars samples were transferred under inert gas to a He(g) filled chamber. LP-MBMS experiment performed at 6.5 W for 2 s. The first 0.25 s data corresponding to a temperature between 900 and 1000 °C is reported here.

For the cold-trapping experiments the charred pellet was mounted in the center of a N2(g) cooled brass collection cup, ca. 2 mm from the cold surface. The brass collection cup had been previously coated with Diphenylisulfide to which a known quantity of chlorobenzene had been added as an internal standard. The charred pellet was irradiated with the CO2 at 13 W whilst being spun through one revolution. After the experiment, the cold trap was washed with methylene chloride. The methylene chloride extract is filtered and concentrated by evaporation. The concentrated extract is then injected onto a capillary GC column and the species identified using a MSD. We cannot currently observe species smaller than toluene or benzothiophene, because of the elution of the large solvent peak.

Results and Discussion

The DRIFTS of the reheated and dried samples are shown in Figure 1. All of the samples show the same basic spectral features at the following approximate locations, a broad υ(OH) reaching a maximum at 3400 cm⁻¹, sharper aliphatic and aromatic υ(CH) at 2900 cm⁻¹, a υ(C=O) at 1750 cm⁻¹, an aromatic υ(C=CH) at 1650 cm⁻¹, a complex fingerprint region and two weaker aromatic υ(CH) bands at 850 cm⁻¹. Clearly metal doping does not influence the ratio of OH to carbonyl groups to hydrocarbon. The palladium doped char has by far the most OH remaining in the structure whereas sodium and potassium have the least. The copper doped char has the highest ratio of carbonyl to aromatic hydrocarbon.

Figure 1. DRIFTS of cellulose and 1 wt% doped cellulose 1/2 h 375 °C chars after heating to 400 °C under He(g).

Solid state MAS 13C NMR, not shown, also shows similar profound differences between the doped char samples. The zinc doped char appears similar to the undoped char, whereas most of the other chars appear very different or appear to have charred less.

The LP-MBMS experiment may be regarded as a snapshot of the initial reactive cloud of molecules evolved from the pyrolyzing char surface. The primary radical that we have proposed to evolve with these species is propargyl. When Avicel cellulose is doped by wt% of metal acetate and charred the amount of CO2 produced during pyrolysis is proportional to the molar amount of metal originally doped. This implies that one of the basic reactions of the metal in the char is to stoichiometrically oxidize the adjacent char. All metal doped chars produce slightly more toluene than the

undoped cellulose but palladium doped char produces a much larger quantity of this species than any other char during pyrolysis. The relative intensities of PAH species relative to m/z 202, pyrene are shown in Figure 2. It can be seen that whilst the initial ratios of the species produced are altered no distinct pattern emerges. All the chars produce more m/z 215, potassium doped more so. Universal suppression of PAH is only seen for the larger m/z 263 and 304. Most of the metals actually increase the initial amount of m/z 252, benzo-a-pyrene.

In the chemical/cold trap experiment radicals are trapped as their phenylsulfide adducts, Figure 3. No propargyl radical is observed in the cold trap experiment as presumably the reaction is being sampled further along the reaction coordinate. The dominant reactive species observed from cellulose char is the hydrogen radical, together with methyl, phenyl, and phenylacetylene (trapped as benzo(ghi)phenanthrene) radicals. Only calcium doping cause the amount of these trapped radicals to increase. Palladium and calcium create the observance of methyl naphthalene radical. All the other metals in general suppress the formation of radicals consistent with a more ionic char decomposition pathway during pyrolysis.

In addition to the chemically trapped radicals we also observe the PAH formed at this point of the reaction. The slate of aromatic and PAH species observed for the undoped cellulose char are shown in Figure 4. Toluene, naphthalene and acenaphthene are the dominant products. In addition to the hydrocarbon species we also observe a number of O-aromatic molecules. PAH as big as benzofluoranthene are easily observed. The ratio of these products changes with metal doping and further analysis of this data should provide insight to some interesting trends.

![Figure 2](image-url)  
**Figure 2.** Relative intensities vs. m/z 202, pyrene, of selective masses of cellulose and 1 wt% doped cellulose.

![Figure 3](image-url)  
**Figure 3.** Relative abundances of PhS trapped reactive intermediates from the 13 W laser pyrolysis of ½ h 375 °C Avicel cellulose char and 1 mol% metal acetate doped Avicel cellulose char.

**Conclusions**

Obviously the fact that the PAH are still in existence with radicals in the LP-MBMS and cold trap experiment suggest that we are not observing the final slate of products. We hope to be able to correlate the LP-MBMS data with the cold trapped data with conventional tube furnace data in order to build a complete picture of the reaction pathways involved in the pyrolysis evolved reacting volatile plume.

**Acknowledgement.** We would like to thank Philip Morris, USA, for support.

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QUANTUM MECHANICAL MODELING OF THE DECOMPOSITION OF SUGARS DURING ACID HYDROLYSIS OF BIOMASS

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Introduction
The treatment of biomass with hot acid is an intriguing option for preparing cellulose for enzymatic hydrolysis or for directly hydrolyzing cellulose. The goal of these two procedures is to convert the cellulose in biomass into sugars that can either be used as a fermentation substrate for ethanol, a transportation fuel, or as a feedstock for chemical production. A key issue during hydrolysis is the yield of the sugars. Ideally one would like to convert all of the hemicellulose and cellulose to monosaccharides. However, it is known that some of the sugars formed by hydrolysis are decomposed under acidic conditions. Further, it has been demonstrated that furanyl compounds are formed by this decomposition and these furanyl compounds can polymerize and may restrict access to unreacted polysaccharides.

In spite of the importance of sugar decomposition in hot acid, there is uncertainty concerning the mechanisms of these processes. Reaction (1) shows a proposed mechanism for conversion of xylose (1) to furfural (2) that involves a ring opening. Since xylan is an important constituent of corn stover, this process may be important in the full utilization of corn.

Ring Opening Mechanism:

\[
\begin{align*}
1 & \xrightarrow{\text{H}} 2 \\
2 & \xrightarrow{\text{H}} 3 \\
3 & \xrightarrow{\text{H}} 4 \\
4 & \xrightarrow{\text{H}} 5 \\
5 & \xrightarrow{\text{H}} 6 \\
6 & \xrightarrow{\text{H}} 7 \\
7 & \xrightarrow{\text{H}} 8 \\
8 & \xrightarrow{\text{H}} 9 \\
9 & \xrightarrow{\text{H}} 10 \\
10 & \xrightarrow{\text{H}} 11 \\
11 & \xrightarrow{\text{H}} 7 \\
\end{align*}
\]

In this mechanism, the sugar in the pyran form opens to the aldehyde (2), which then tautomerizes to an enol (3). Dehydration of this enol by a six centered transition state to form 4. Another tautomerization produces 5, which dehydrates again to form 6. This compound can cyclize and dehydrate to form furfural, 7.

An alternative mechanism for forming furfural from xylose involves direct formation of the furan after protonation of the pyran form (1). Xylose protonated at the 2 hydroxyl group can lose a water molecule and the resulting carbox cation is susceptible to attack by the ring ether forming a dihydroxy furan (9). This compound can then dehydrate to form furfural, 7.

Direct Mechanism:

\[
\begin{align*}
1 & \xrightarrow{\text{H}} 2 \\
2 & \xrightarrow{\text{H}} 3 \\
3 & \xrightarrow{\text{H}} 4 \\
4 & \xrightarrow{\text{H}} 5 \\
5 & \xrightarrow{\text{H}} 6 \\
6 & \xrightarrow{\text{H}} 7 \\
7 & \xrightarrow{\text{H}} 8 \\
8 & \xrightarrow{\text{H}} 9 \\
9 & \xrightarrow{\text{H}} 10 \\
10 & \xrightarrow{\text{H}} 11 \\
11 & \xrightarrow{\text{H}} 7 \\
\end{align*}
\]

We have studied the mechanisms of sugar decomposition shown in Rxns. (1) – (3) with quantum mechanical molecular modeling in an attempt to determine which mechanism is prevalent. Small molecular systems are used to model the reaction steps shown in this mechanism. This allows the use of accurate computational techniques to capture the essential chemistry of these mechanisms.

Computational Techniques
The calculations in this study were carried out using the Gaussian 03 suite of programs. Local minima and transitions states were found using B3LYP optimizations. Starting geometries for the transition state optimizations were often located using a potential energy scan, where one internal coordinate was systematically changed while all other coordinates were allowed to relax. The geometry at the calculated maximum of this scan was used as a starting geometry for a transition state (TS) optimization. Stable structures had only positive vibrational frequencies while transition states had one imaginary frequency. The transition states were confirmed by visual inspection of the imaginary frequency using the Molden visualization package.

Results and Discussion

Ring Opening Mechanism
The first step in this mechanism is the opening of the pyran ring, 1, to form the aldehyde, 2. This step involves a four-centered transition state as is shown for the simplified structure in Rxn. (4).

\[
\begin{align*}
1 & \xrightarrow{\text{H}} 2 \\
2 & \xrightarrow{\text{H}} 3 \\
3 & \xrightarrow{\text{H}} 4 \\
4 & \xrightarrow{\text{H}} 5 \\
5 & \xrightarrow{\text{H}} 6 \\
6 & \xrightarrow{\text{H}} 7 \\
7 & \xrightarrow{\text{H}} 8 \\
8 & \xrightarrow{\text{H}} 9 \\
9 & \xrightarrow{\text{H}} 10 \\
10 & \xrightarrow{\text{H}} 11 \\
11 & \xrightarrow{\text{H}} 7 \\
\end{align*}
\]

We have calculated the barrier for this reaction for several C5 and C4 sugars as well as hydroxymethyl methyl ether using density functional theory, B3LYP/6-31G(d). Values of 35 to 40 kcal mol\(^{-1}\) were obtained. Figure 1 shows a plot of the calculated results for hydroxymethyl methyl ether. The barriers for these reactions are greatly reduced with the addition of proton. For instance, with hydroxymethyl methyl ether the barrier is lowered from 40.4 kcal mol\(^{-1}\) to 14.3 kcal mol\(^{-1}\) with the addition of a proton. This suggests that protic solvents or added acid should accelerate this reaction. This lower barrier is also consistent with kinetic measurements made in water.

![Figure 1. A potential energy plot of the calculated relative energies for the unimolecular decomposition of hydroxymethyl methyl ether.](image)

The second step in the ring opening mechanism as show in Rxn. (1) is a keto-enol tautomerization. Unfortunately, the barriers for these reactions are typically high. Our calculations show that the tautomerization of acetaldehyde to vinyl alcohol has a barrier of 67.8 kcal mol\(^{-1}\). We find that protonation does not significantly lower this barrier. Since the mechanism in Rxn. (1) contains two tautomerization steps, this pathway appears unlikely at practical temperatures.
We propose an alternative mechanism for furfural formation whereby the opened sugar undergoes a double dehydration as shown in reaction (5). In this mechanism, a proton catalyzes the ring opening step. Protonation of the C3 hydroxyl group on the resulting sugar, 2, can lead to a double dehydration, forming 14. This compound can then cyclize and further dehydrate to give furfural, 7.

\[
\text{12} \quad \begin{array}{c}
\text{2} \\
\text{13} \\
\text{14} \\
\text{7}
\end{array}
\]

We have modeled this double dehydration using protonated glycerol as a model system. Figure 2 shows a plot of the calculated potential energy for this reaction. The barrier to the loss of the first water molecule from protonated glycerol is 22.4 kcal mol\(^{-1}\). The barrier to the loss of a second water is calculated to be 36.5 kcal mol\(^{-1}\). This second barrier is high because a primary carbocation is formed. In double dehydration of xylose, Rxn. (5), a secondary carbocation is formed. We calculated the second barrier when a methyl group is attached to glycerol to better simulate dehydration in xylose. In this case the barrier is 21.7 kcal mol\(^{-1}\) as shown in the figure. Thus the barriers for this double dehydration (22.4 and 21.7 kcal mol\(^{-1}\)) are low enough that this reaction could be fast at moderate temperatures.

**Figure 2.** A potential energy plot of the calculated energies for the double dehydration of glycerol (CBS-QB3).

**Direct Mechanism**

We have calculated the barrier for a model reaction similar to reaction (2). The potential energy for the reaction of a protonated dihydroxy pyran is shown in Figure 3. We find that there is a concerted reaction for loss of water and rearrangement to the furan. The calculated barrier for this reaction is 13 kcal mol\(^{-1}\).

**Calculated Rate Constants**

The geometries and vibrational frequencies from these calculations can be used to estimate rate constants for the xylose decomposition reactions. These calculated values are used to estimate the entropy change in going from a reactant to a transition state (\(\Delta S^*\)), which is used to calculate the pre-exponential Arrhenius factor using Transition State Theory (TST).

\[
A_0 = \left( \frac{kT}{h} \right)^{1/2} \exp \left( \frac{\Delta S^*}{RT} \right)
\]

Rate constants calculated using Eqn. 6 are collected in Table 1. As can be seen, the rate constants are high at typical pretreatment conditions. The activation energies, \(A_{TST}\), for these mechanisms are consistent with measure global activation energies for xylose decomposition. The table is divided into rate constants for the ring opening mechanism, left, and the direct mechanism, right.

**Figure 3.** Plot of the potential energy for the direct conversion of a protonated pyran to a furan.

<table>
<thead>
<tr>
<th>Table 1 Calculated rate constants using TST</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Ring Opening Mech.</strong></td>
</tr>
<tr>
<td>****</td>
</tr>
<tr>
<td>Ea (kcal mol(^{-1}))</td>
</tr>
<tr>
<td>(A_{TST} (s^-1))</td>
</tr>
<tr>
<td>Temp. (°C)</td>
</tr>
<tr>
<td>Calculated TST Rate Constants</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
</tbody>
</table>

**Conclusions**

The rate constants that we obtain for the ring opening mechanism appear to be lower than for the direct mechanism. However, this does not include the dehydration steps such as are shown in Rxn. (3). These reactions should have barriers of about 20 kcal mol\(^{-1}\), which should make the two mechanisms more comparable.

**Acknowledgment**

We would like to acknowledge the Office of the Biomass Program of the U.S. Department of Energy, who funded portions of this work, and the National Center for Supercomputer Applications (NCSA Grant # CHE-980028n) for computer time.

**References**


MECHANISTIC INVESTIGATION INTO THE FORMATION OF POLYCYCLIC AROMATIC HYDROCARBONS FROM THE PYROLYSIS OF TERPENES

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Introduction

Currently, there is interest in utilizing renewable energy resources, such as wood and agriculture waste, for power generation, production of liquid fuels, and specialty chemicals as a consequence of the environmental benefits. However, pyrolysis, gasification, and combustion of biomass produces polycyclic aromatic hydrocarbons (PAHs), which are environmental pollutants and soot precursors. In pyrolysis, PAHs are proposed to be formed by pyrolysis, in which radicals undergo a series of bimolecular reactions with alkynes, alkynes, and aromatics to form larger ring structures. The yield of PAHs is found to increase with temperature and gas phase residence times. In gasification and combustion, it has been postulated that PAHs arise from pyrolysis reactions in oxygen-deficient regions. A better fundamental understanding of the initial pyrolysis reactions that occur in the thermochemical processing of biomass, and subsequent reactions of the vapor phase products could provide insight into reducing PAH emissions from biomass combustion, reducing tar formation from gasification, and enhancing production of specialty chemicals from the fast pyrolysis of biomass.

As a consequence of the chemical complexity of biomass and its pyrolysis products, it has been difficult to gain detailed information on the formation pathways of specific products. Thus, to gain fundamental kinetic and mechanistic information on product formation in biomass pyrolysis, efforts have focused on the pyrolysis of key chemical constituents found in biomass. In our previous investigations, we have focused on the pyrolysis of plant steroids as potential precursors to PAHs. It was shown that the native ring structure of the steroid was maintained to form PAHs, such as phenanthrene and chrysene, and the yield of PAHs was dependent on the steroid structure. PAH yield and ring size were also shown to increase with residence time, temperature, and concentration. In this investigation, the pyrolysis of plant terpenes was investigated at 600-800 °C and at short residence times (≤1 s). This study focused on the pyrolysis of solanesol, squalene, phytol, and β-carotene. These terpenes are predominately found in the foliage. The yield of PAHs from the pyrolysis of terpenes will be compared to those found from steroids to determine if the alkenes produced from the pyrolysis of terpenes, such as isoprene, enhance the formation of PAHs.

Experimental

The flow pyrolysis reactor used in this study has been previously described. After the pyrolysis, both cold traps were washed with CS2, and the washings were combined (total volume 10-20 mL). Solutions of hexatriacontane, phenanthrene-d10 and 12-benzanthracene-d12 were added as standards, and the reaction mixtures were analyzed by GC-MS and quantitated by GC-FID as previously described. Typical shot to shot reproducibility was ±2% with the exception of the smaller products (<0.1 mol%) in the presence of a large background in which the shot to shot reproducibility was typically ±5-10%. Products were quantitated by averaging the GC-FID output relative to the internal standards. Response factors were measured with authentic samples or estimated from measured response factors for structurally related compounds and based on carbon number relative to the internal standards (hexatriacontane, phenanthrene-d10 or 1.2-benzanthracene-d12). The limits of detection (LOD) for the PAHs depended upon the complexity of the reaction mixture, but the LOD was typically 5-20 μg g-1.

The residence time in the reaction chamber was calculated from the volume of the reactor (1.06 mL or 11.7 mL) in the hot zone, the helium flow rate at room temperature, and a correction for the gas flow rate at the reaction temperature. The reactor was also calibrated by the pyrolysis of phenethyl acetate, whose well-defined first order kinetics allows verification of the reactor temperature and residence time.

Results and Discussion

The flow pyrolysis of solanesol was investigated at high concentration at 700 and 800 °C with residence times of 0.13 and 1.0 s. The yields of selected PAHs are shown in Table 1. One surprising observation was the lack of substantial quantities of 3-ringed or larger PAHs at temperatures less than 800 °C. On the basis of Severson’s study, it was expected that solanesol would produce significant quantities of PAHs, since it was concluded that solanesol contributed as much as 40% of the benzopyrene produced from the pyrolysis of the petroleum ether extract of tobacco. Duplicate runs at 800 °C produced product yields that were ±20-30% of those reported in Table 1 confirming the low yields of PAHs from the pyrolysis of solanesol.

At 600 °C and 0.13 s residence time, dipentene (limonene, m/z 136) and isoprene (m/z 68) were the dominant products. Small amounts of toluene, n-xylene and isomeric products with m/z 136 (4 isomers) and 138 (1-methyl-4-(1-methyl)cyclohexene and isomer) were also found. At 700 °C with 0.13 s residence time, m-xylene, toluene, and isoprene were the dominant products, and only small amounts of dipentene and isomers were found. Small amounts of aromatic products, such as trimethylbenzene, methylisopropenylbenzene, methyldiene, and methylnaphthalene, were also formed. A similar product distribution was found in the low concentration (68 mg h⁻¹, 3x10⁴ M), short residence time (0.15 s), flow pyrolysis of solanesol at 700 °C, but the product yields were lower. As the temperature and residence times increased, larger PAHs, such as phenanthrene, were detected. However, four ringed aromatic compounds, such as pyrene and chrysene, were only found at 800 °C. Increasing the residence time from 1.0 s to 9.0 s at 700 °C significantly increased the yield of PAHs. It appears that solanesol only forms significant quantities of 3-5 ringed aromatics at high temperatures (800 °C) or at long residence times (≥1 sec).
It had been previously proposed that the terpene components of tobacco form PAHs by decomposing into isoprene and pyrosynthesis into larger PAHs. Isoprene is the major component of the unsaturated gaseous hydrocarbons present in tobacco smoke and pyrolysis of isoprene at 600–800 °C forms PAHs, including benzo[a]pyrene (2-3 s residence time). Larger PAHs can form from isoprene pyrolysis by concerted Diels–Alder or free radical reactions. Dimerization of isoprene, via Diels–Alder reaction, yields dipentene (also called p-metha-1,8-diene or limonene) and mixtures of dimethylvinylcyclohexenes (see below), but tobacco smoke contains only dipentene and 2,4-dimethyl-4-vinylcyclohexene.

This suggests that free radical pathways form the products in tobacco. The pyrolysis of solanesol has been studied previously (residence time unknown), and it was found that at temperatures up to 550 °C, the hydrocarbon fraction was mainly composed of monoterpenoids with only traces of aromatic hydrocarbons. For example, pyrolysis of solanesol at 500 °C under nitrogen produced pyrones, dimethyladiadiene, dimethylinylcyclohexene, \( \Delta^1 \)-methene, and dipentene. The major gas phase products were isoprene (64 parts), 2-methyl-2-butene (8 parts), and one part each of 2-methyl-1-butene and 3-methyl-1-butene. Running the pyrolysis in hydrogen altered the product yields confirming that the products were formed by free radical rather than a concerted pathway. At 600 °C, the monocyclic fraction (93 % by weight of the total sample) contained benzene (5 wt%), toluene (30 wt%), xylene (38 wt%) and \( C_9 \) alkylbenzenes (21 wt%). Similar results were found in our study at 600 °C. At temperatures above 650 °C, only aromatic compounds were formed and no terpenes were found.

The pyrolysis of dipentene (limonene) has been studied at 450 °C over copper pellets (residence time unknown). The dominant products were pyrones, other alkylcyclohexadienes, and a small amount (9%) of aromatic hydrocarbons. It was proposed that limonene decomposed by a biradical mechanism in which radicals from \( \text{C}_6 \text{H}_{10} \) homolysis of the bisallylic bond underwent disproportionation to form a triene that can isomerize (1,5-hydrogen shift) and undergo electrocyclic ring closure to form the pyrones and 1,3-dimethyl-1-ethyl-3,5-cyclohexadiene (not shown). Independent pyrolysis of the pyrones at 450 °C gave a similar distribution of aromatic hydrocarbons.

Thus, on the basis of the literature evidence, it is proposed that solanesol decomposes directly to isoprene and dipentene, as the major products, and alkylcyclohexenes by a free radical pathway. Depending on the reaction severity (residence time and temperature), these initial products can undergo additional free radical reactions to form alkyl-substituted monocyclic aromatic hydrocarbons, and larger PAHs. A potential mechanism for dipentene formation by a free radical pathway is shown in Figure 1a. The TG-MS of solanesol showed a significant loss of water at relatively low temperatures (i.e., 360 °C). Therefore, water can be eliminated by a concerted 1,4-elimination to form a terminal diene. Homolysis of the bisallylic \( \text{C}_8 \text{H}_{16} \) bond forms an isopropenyl radical (\( \text{C}_6 \text{H}_3 \)) that can hydrogen abstract from solanesol to give isoprene. The allylic radical can undergo an intramolecular 1,6-cyclization reaction competitively with \( \text{β} \)-scission. As previously reported, the ratio of intramolecular radical addition to \( \text{β} \)-scission for 1-hex-5-enyl radical is 124 at650 °C and 39 at 750 °C, which indicates that cyclization should dominate. After cyclization, \( \text{β} \)-scission of the tertiary radical forms dipentene and a radical that can continue the

Table 1. Product Yields From the Flow Pyrolysis of Solanesol at High Concentration.

<table>
<thead>
<tr>
<th>Product</th>
<th>Temperature (°C)</th>
<th>700</th>
<th>700</th>
<th>700</th>
<th>700</th>
<th>800</th>
</tr>
</thead>
<tbody>
<tr>
<td>Benzene</td>
<td>8.50</td>
<td>4.6</td>
<td>23.1</td>
<td>ND</td>
<td>67.6</td>
<td></td>
</tr>
<tr>
<td>Toluene</td>
<td>30.2</td>
<td>22.7</td>
<td>47.8</td>
<td>ND</td>
<td>61.3</td>
<td></td>
</tr>
<tr>
<td>Styrene</td>
<td>6.70</td>
<td>5.4</td>
<td>12.6</td>
<td>ND</td>
<td>21.5</td>
<td></td>
</tr>
<tr>
<td>Indene</td>
<td>2.03</td>
<td>0.16</td>
<td>5.45</td>
<td>ND</td>
<td>12.6</td>
<td></td>
</tr>
<tr>
<td>Naphthalene</td>
<td>0.91</td>
<td>0.53</td>
<td>3.67</td>
<td>ND</td>
<td>23.2</td>
<td></td>
</tr>
<tr>
<td>2-Methylnaphthalene</td>
<td>1.72</td>
<td>0.91</td>
<td>4.40</td>
<td>ND</td>
<td>13.4</td>
<td></td>
</tr>
<tr>
<td>1-Methylnaphthalene</td>
<td>1.07</td>
<td>0.55</td>
<td>2.94</td>
<td>10.8</td>
<td>8.28</td>
<td></td>
</tr>
<tr>
<td>Acenaphthylene</td>
<td>0.51</td>
<td>0.00</td>
<td>1.28</td>
<td>2.2</td>
<td>6.47</td>
<td></td>
</tr>
<tr>
<td>Fluorene</td>
<td>0.00</td>
<td>0.00</td>
<td>0.77</td>
<td>ND</td>
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<td></td>
</tr>
<tr>
<td>Phenanthrene</td>
<td>0.00</td>
<td>0.00</td>
<td>0.37</td>
<td>2.7</td>
<td>4.10</td>
<td></td>
</tr>
<tr>
<td>Anthracene</td>
<td>0.00</td>
<td>0.00</td>
<td>0.26</td>
<td>0.88</td>
<td>1.90</td>
<td></td>
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<tr>
<td>MethylPhen/An</td>
<td>0.00</td>
<td>0.00</td>
<td>1.30</td>
<td>5.4</td>
<td>6.71</td>
<td></td>
</tr>
<tr>
<td>Fluoranthene</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.89</td>
<td>0.71</td>
<td></td>
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<tr>
<td>Pyrene</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>1.5</td>
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<tr>
<td>Methylpyrenes</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>3.5</td>
<td>3.84</td>
<td></td>
</tr>
<tr>
<td>Benzo[a]anthracene</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.42</td>
<td>0.67</td>
<td></td>
</tr>
<tr>
<td>Chrysenne</td>
<td>0.00</td>
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<td>0.00</td>
<td>0.38</td>
<td>0.61</td>
<td></td>
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<tr>
<td>Benzo[a]pyrene</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.42</td>
<td>0.39</td>
<td></td>
</tr>
</tbody>
</table>

a Concentration determined by sublimation rate/gas flow rate at the reaction temperature. ND = Not Determined.

unzipping process. Hydrogen abstraction from solanesol at the allylic site between the double bonds also forms an intermediate that can undergo 1,6-cyclization to form dipentene. Hydrogen abstraction from the methyl group (Figure 1b) followed by 1,6-cyclizations and β–scission forms 2,4-dimethyl-4-vinylcyclohexene (which is found in tobacco smoke).

Larger PAHs must arise from reaction of the monocyclic aromatic hydrocarbons with the low molecular weight unsaturated products, such as isoprene. In the combustion of 1,3-butadiene in a sooting flame (i.e., fuel rich flame), it was concluded that monocyclic PAHs were made by free radical pathways involving C$_4$H$_8$• as the dominant species. The Diels–Alder reaction of butadiene was too slow and concentration of the reagents too low, to account for the PAHs formed in the flame. In the pyrolysis of solanesol, larger PAHs are most likely formed by a similar pathway with aromatic and benzyl radicals adding to unsaturated products, such as isoprene, in the hot zone of the furnace (i.e., pyrosynthesis). However, the specific reaction pathway from the primary products (i.e., isoprene and dipentene) to the large PAHs are only implied and are not specifically known. Solanesol and other terpenes could also enhance PAH yields from pyrolysis of other substrates by providing hydrocarbons (i.e., isoprene) for pyrosynthesis. For example, in the pyrolysis of cellulose, replacing the nitrogen carrier gas with propene increased the yield of benzo[a]pyrene 1.9 fold at 350-450 °C and 6.9 fold at 450-550 °C even though the propene did not produce benzo[a]pyrene at temperatures >550 °C.

Preliminary results on the yield of PAHs produced from the pyrolysis of solanesol, β-carotene, squalene, and stigmasterol (a plant steroid) at 800 °C with a residence time of 1.0 s is shown in Figure 2. In general, the steroid produces more PAHs than the terpenes, especially phenanthrene, anthracene, benz[a]anthracene, and chrysene which can be derived from dehydrogenation, dealkylation, and rearrangement of the steroid. Surprisingly, squalene produces fewer PAHs than solanesol. The pyrolys of β-carotene and squalene is currently being investigated at 700 and 600 °C with a residence time of 1.0 s and 0.13 s. These results will be discussed in the oral presentation.


(22) (a) Handford-Styring, S. M.; Walker, R. W. J. Chem. Soc., Faraday Trans. 1995, 91, 1431. The rate expression for β–scission to form ethylene and 1-but-3-enyl radical is log \( k(s') = 13.0 - 29.9/θ \), while intramolecular cyclization has a rate expression of \( logk(s') = 10.0 - 8.4/θ \). (b) For a cyclic system, the rate expression for the intramolecular cyclization of the 2-allylbenzyl radical is \( logk(s') = 11.14 - 16.28/θ \).


