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Biomass

Top Value Added Chemicals from Biomass Volume I—Results of Screening for Potential Candidates from Sugars and Synthesis Gas

Produced by the Staff at
Pacific Northwest National Laboratory (PNNL)
National Renewable Energy Laboratory (NREL)
Office of Biomass Program (EERE)
For the Office of the Biomass Program
T. Werpy and G. Petersen, Editors

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Produced by Staff at
the Pacific Northwest National Laboratory (PNNL) and
the National Renewable Energy Laboratory (NREL)

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Executive Summary

This report identifies twelve building block chemicals that can be produced from sugars via biological or chemical conversions. The twelve building blocks can be subsequently converted to a number of high-value bio-based chemicals or materials. Building block chemicals, as considered for this analysis, are molecules with multiple functional groups that possess the potential to be transformed into new families of useful molecules. The twelve sugar-based building blocks are 1,4-diacids (succinic, fumaric and malic), 2,5-furan dicarboxylic acid, 3-hydroxy propionic acid, aspartic acid, glucaric acid, glutamic acid, itaconic acid, levulinic acid, 3-hydroxybutyrolactone, glycerol, sorbitol, and xylitol/arabinitol.

Building Blocks
1,4 succinic, fumaric and malic acids
2,5 furan dicarboxylic acid
3 hydroxy propionic acid
aspartic acid
glucaric acid
glutamic acid
itaconic acid
levulinic acid
3-hydroxybutyrolactone
glycerol
sorbitol
xylitol/arabinitol

The synthesis for each of the top building blocks and their derivatives was examined as a two-part pathway. The first part is the transformation of sugars to the building blocks. The second part is the conversion of the building blocks to secondary chemicals or families of derivatives. Biological transformations account for the majority of routes from plant feedstocks to building blocks, but chemical transformations predominate in the conversion of building blocks to molecular derivatives and intermediates. The challenges and complexity of these pathways, as they relate to the use of biomass derived sugars and chemicals, were briefly examined in order to highlight R&D needs that could help improve the economics of producing these building blocks and derivatives. Not surprisingly, many of the transformations and barriers revealed in this analysis are common to the existing biological and chemical processing of sugars.

The final selection of 12 building blocks began with a list of more than 300 candidates. The shorter list of 30 potential candidates was selected using an iterative review process based on the petrochemical model of building blocks, chemical data, known market data, properties, performance of the potential candidates and the prior industry experience of the team at PNNL and NREL. This list of 30 was ultimately reduced to 12 by examining the potential markets for the building blocks and their derivatives and the technical complexity of

the synthesis pathways. A second-tier group of building blocks was also identified as viable candidates. These include gluconic acid, lactic acid, malonic acid, propionic acid, the triacids, citric and aconitic; xylonic acid, acetoin, furfural, levoglucosan, lysine, serine and threonine. Recommendations for moving forward include examining top value products from biomass components such as aromatics, polysaccharides, and oils; evaluating technical challenges in more detail related to chemical and biological conversions; and increasing the suites of potential pathways to these candidates.

1 Background

America is fortunate to possess abundant and diverse agricultural and forest resources, unused cropland and favorable climates. Together with a remarkable talent to develop new technologies, we have a tremendous opportunity to use domestic, sustainable resources from plants and plant-derived resources to augment our domestic energy supply.

The Biomass Program, in the Energy Efficiency and Renewable Energy Office in the Department of Energy directly supports the goals of The President's National Energy Policy, the Biomass R&D Act of 2000 and the Farm Security and Rural Investment Act of 2002. To accomplish these goals, the Program supports the integrated biorefinery, a processing facility that extracts carbohydrates, oils, lignin, and other materials from biomass, converts them into multiple products including fuels and high value chemicals and materials. Already today, corn wet and dry mills, and pulp and paper mills are examples of biorefinery facilities that produce some combination of food, feed, power and industrial and consumer products.

This report, the first of several envisioned to examine value-added products from all biomass components, identifies a group of promising sugar-derived chemicals and materials that could serve as an economic driver for a biorefinery. By integrating the production of higher value bioproducts into the biorefinery's fuel and power output, the overall profitability and productivity of all energy related products will be improved. Increased profitability makes it more attractive for new biobased companies to contribute to our domestic fuel and power supply by reinvesting in new biorefineries. Increased productivity and efficiency can also be achieved through operations that lower the overall energy intensity of the biorefinery's unit operations, maximize the use of all feedstock components, byproducts and waste streams, and use economies of scale, common processing operations, materials, and equipment to drive down all production costs.

2 Objective

In 2002, The US DOE Office of Energy Efficiency and Renewable Energy reorganized to combine previously separate biofuels, biopower, and biobased products programs into a single Biomass Program. Promotion of biorefineries producing multiple products, including higher-value chemicals as well as fuels and power, is a main objective of the consolidated program. The Office of the Biomass Program asked research staff at the National Renewable Energy Laboratory (NREL) and Pacific Northwest National Laboratory (PNNL) to identify the top ten opportunities for the production of value-added chemicals from biomass that would economically and technically support the production of fuels and power in an integrated biorefinery and identify the common challenges and barriers of associated production technologies. This report is a companion study to ongoing program planning reports for the Biomass Office including a Multiyear Program Plan, a Multiyear Technical Plan, an Analysis Plan, a Communications Plan, and an Annual Operating Plan.

3 Overall Approach

The separate steps in the overall consideration for this analysis are depicted in the following flow diagram (Figure 1).

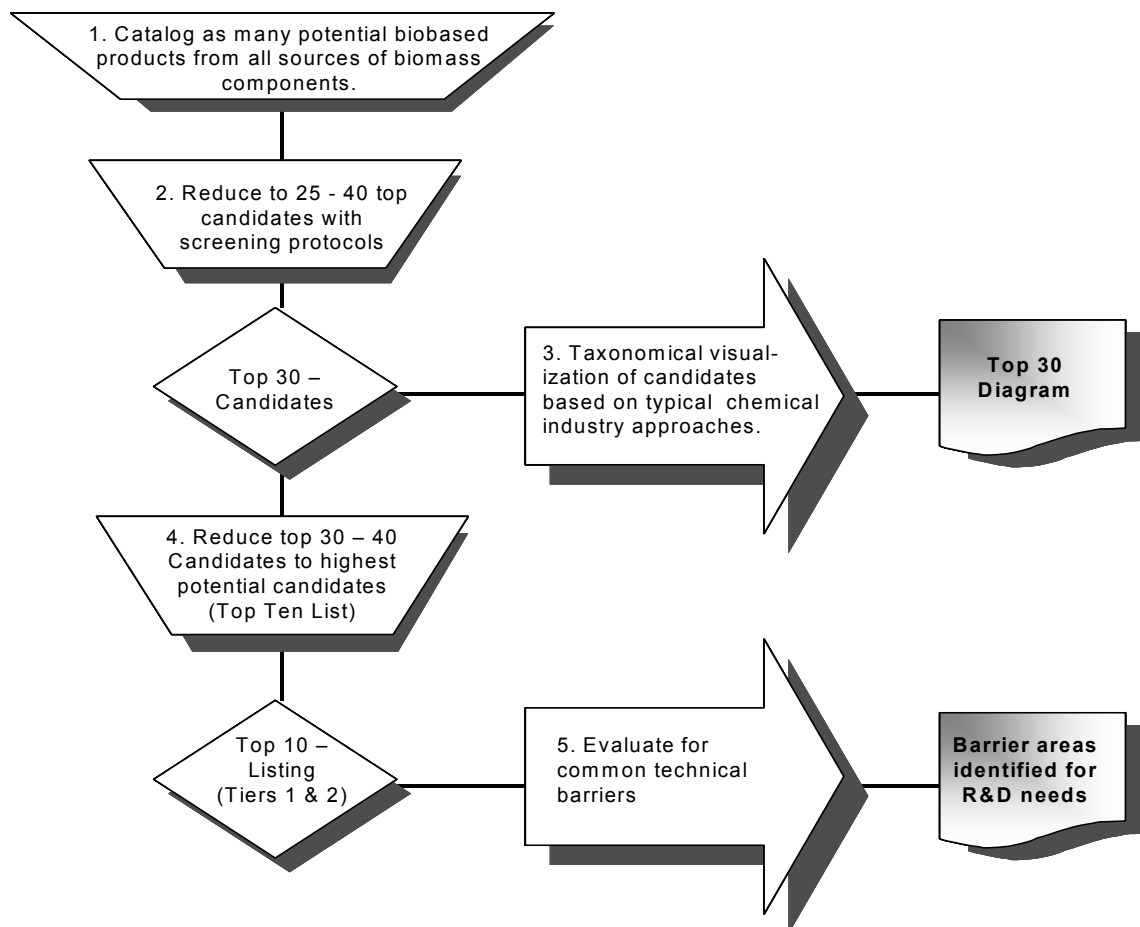


Figure 1 - Visual Representation of Overall Selection Strategy

A group of over 300 possible building block chemicals was assembled from a variety of resources and compiled in an Access database. The source materials included previous DOE and National Laboratory reports and industry and academic studies listed in the Bibliography. The database includes a chemical name, structure, sources for the biomass feedstock, the current and potential production processes, a designation as a commodity, specialty, polymer or food/ag chemical, and pertinent citation information. The initial screening criteria included the cost of feedstock, estimated processing costs, current market volumes and prices, and relevance to current or future biorefinery operations. Interestingly, this first criteria set did not provide sufficient differentiation among the sugar based candidates within the database to produce the smaller number of candidates desired in step 2 of Figure 1. A different approach was needed and developed as described in the next section.

4 Initial Screening to the Top 30

A more effective screening tool was found using the concepts employed in traditional petrochemical industry flow-charts as shown in the representative example in Figure 2. All of the products from the petrochemical industry are derived from 8-9 foundation chemicals. An iterative review process was established which used chemical and market production data, estimates of the material and performance properties of the potential candidates and over 75 years of cumulative industry experience of the research team as the basis for the down selection. Figure 3 gives a graphical representation of the top 30 building blocks analogous to the example of the petrochemical industry flow chart shown in Figure 2. This figure depicts the value chain approach used in the downselection process.

From the initial list of over 300, the team systematically down selected to a smaller list using factors that are important components of the strategic criteria shown in Table 1. The screening criteria for this first round included the raw material and estimated processing costs, estimated selling price, the technical complexity associated with the best available processing pathway and the market potential for each of the candidate building blocks.

Table 1 - Biorefinery Strategic Fit Criteria

	Direct Product Replacement	Novel Products	Building Block Intermediates
Characteristic	Competes directly against existing products and chemicals derived from petroleum	Possesses new and improved properties for replacement of existing functionality or new applications	Provide basis of a diverse portfolio of products from a single intermediate
Examples	Acrylic acid obtained from either propylene or lactic acid	Poly(lactic acid) (glucose via lactic acid is sole viable source)	Succinic, levulinic, glutamic acids, glycerol, syngas
Upside	Markets already exist Understand cost structures and growth potential Substantial reduction in market risk	Novel products with unique properties hence cost issues less important No competitive petrochemical routes Differentiation usually based on desired performance New market opportunities Most effective use of properties inherent in biomass	Product swing strategies can be employed to reduce market risks Market potential is expanded Capital investments can be spread across wider number of unit operations Incorporates advantages of both replacement and novel products
Downside	Strictly competing on cost Competing against depreciated capital	Market not clearly defined Capital risk is high Time to commercialization	Identifying where to focus R&D

	Direct Product Replacement	Novel Products	Building Block Intermediates
	Limited (green label) “market differentiation” for biobased vs. petrochemical based sources	may be long	

Almost 50 potential building block candidates resulted from this initial screening.

Continuing to use the strategic fit criteria (direct replacement, novel properties, and potential utility as a building block intermediate) shown in Table 1 above, the team organized the 50 candidates using a carbon number classification framework of one to six carbon compounds (C1 to C6).

Next the team reviewed the candidate group for chemical functionality and potential use. Chemical functionality can be based on the number of potential derivatives that can be synthesized in chemical and biological transformations. Simply, a candidate with one functional group will have a limited potential for derivatives where candidate molecules with multiple functional groups will have a much larger potential for derivatives and new families of useful molecules.

Each candidate molecule was then classified for its current utility to serve as a simple intermediate in traditional chemical processing, as a reagent molecule for adding functionality to hydrocarbons, or as byproducts from petrochemical syntheses. Examples of candidates that fell into this category included acetic acid, acetic anhydride, or acetone.

The team then reviewed the candidate group for potential status as a super commodity chemical. Super commodity chemicals are derived from building block chemicals or are co-products in petrochemical refining. Although the ability of biomass to serve as a source of these compounds is real, the economic hurdles of large capital investments and low market price competitors would be difficult to overcome. Table 2 shows the results of this first screen classified by the carbon number taxonomy C1-C6.

Table 2 - Top Candidates from the First Screen

C #	Name	Projected or Known Use (Building block, reagent, intermediate)	Selected for top 30	Rationale
1	Formic Acid	Reagent	N	Very limited BB, use mostly for adding C1
1	Methanol	BB- limited	N	Super commodity from syngas
1	Carbon Monoxide (+ H ₂ gives syngas)	BB	Y	
1	Carbon dioxide	Reagent	N	Thermodynamics barrier
2	Acetaldehyde	Intermediate	N	V. limited BB.
2	Acetic acid & anhydride	Reagents and Intermediates	N	Limited BB, large commodity scale today from syngas. Adds C2
2	Ethanol	Fuel	N	Major use envisioned as fuel. Limited BB. Will become supercommodity
2	Glycine	Reagent	N	V. limited BB. Few uses envisioned
2	Oxalic acid	Reagent	N	Used primarily as chelator and reagent
2	Ethylene glycol	BB & Product	N	Super commodity
2	Ethylene oxide	BB & Reagent	N	Super commodity
3	Alanine	Intermediate	N	V. limited BB. Few uses envisioned
3	Glycerol	BB	Y	
3	3-Hydroxypropionic acid	BB	Y	
3	Lactic acid	BB	Y	
3	Malonic acid	BB & reagent	Y	
3	Serine	BB	Y	
3	Propionic acid	BB & reagent	Y	
3	Acetone	Intermediate	N	Super commodity, by-product from cumene to phenol synthesis
4	Acetoin	BB	Y	
4	Aspartic acid	BB	Y	
4	Butanol	Intermediate	N	Large commodity chemical, Not a good BB, but large intermediates market. No competitive advantage from biomass
4	Fumaric acid	BB	Y	

C #	Name	Projected or Known Use (Building block, reagent, intermediate)	Selected for top 30	Rationale
4	3-Hydroxybutyrolactone	BB	Y	
4	Malic acid	BB	Y	
4	Succinic acid	BB	Y	
4	Threonine	BB	Y	
5	Arabinitol	BB	Y	
5	Furfural	BB	Y	
5	Glutamic acid	BB	Y	
5	Glutaric acid	BB	N	V. limited market. Indeterminate set of derivatives
5	Itaconic acid	BB	Y	
5	Levulinic acid	BB	Y	
5	Proline	BB	N	Limited market. Indeterminate set of derivatives
5	Xylitol	BB	Y	
5	Xylonic acid	BB	Y	
6	Aconitic acid	BB	Y	
6	Adipic acid	Intermediate	N	Super commodity. Examined previously by DOE/industry with little success
6	Ascorbic acid	BB	N	Limited market. Indeterminate set of derivatives. Subject of successful NIST ATP work
6	Citric acid	BB	Y	
6	Fructose	BB	N	Other routes to the derivatives would be easier than from fructose
6	2,5 Furan dicarboxylic acid	BB	Y	
6	Glucaric acid	BB	Y	
6	Gluconic acid	BB	Y	
6	Kojic & Comeric acid	BB	N	V. limited market. Indeterminate set of derivatives
6	Lysine	BB	Y	
6	Sorbitol	BB	Y	

BB = Building Block

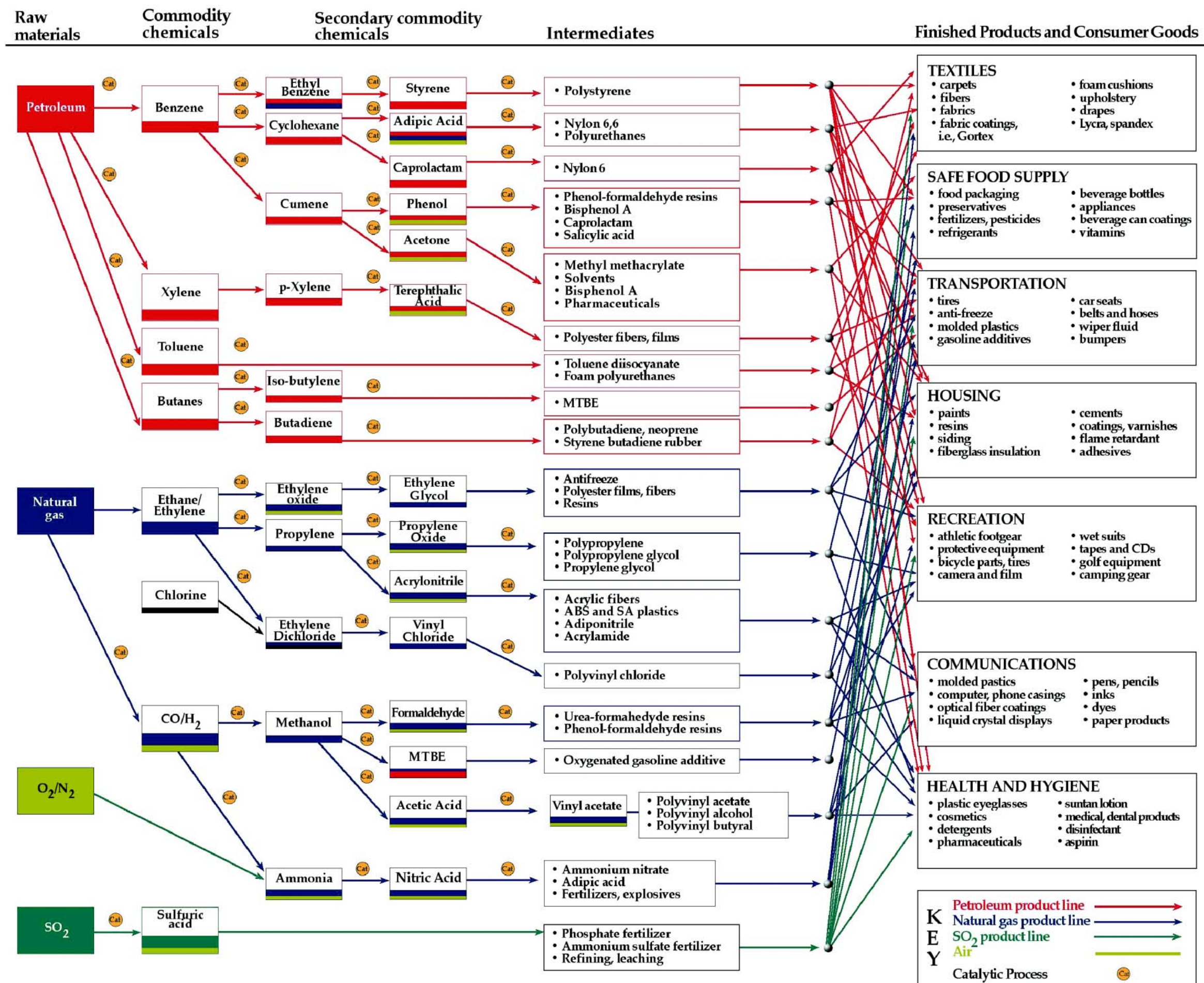


Figure 2 – An Example of a Flow-Chart for Products from Petroleum-based Feedstocks

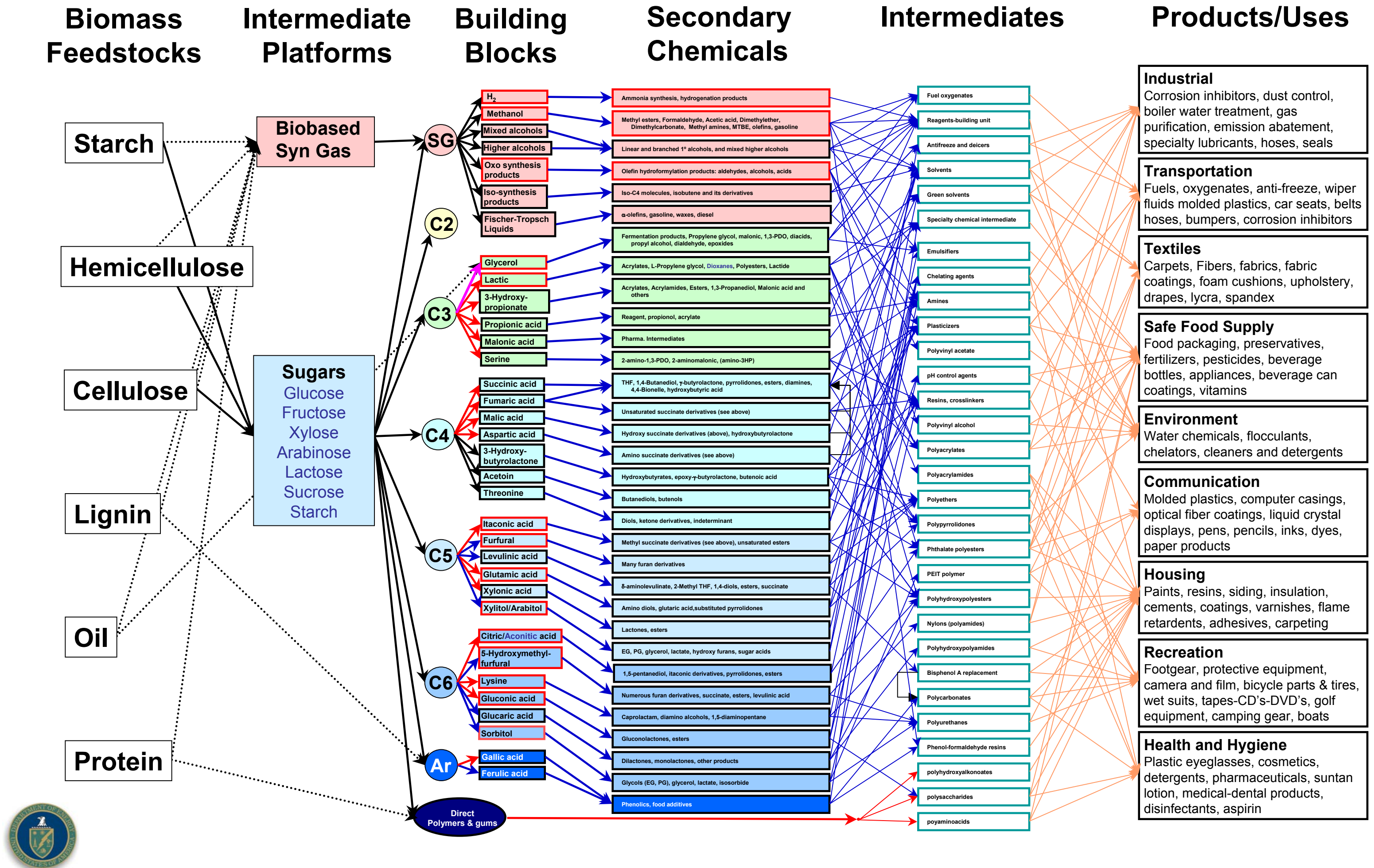


Figure 3 – Analogous Model of a Biobased Product Flow-chart for Biomass Feedstocks

By eliminating those that did not meet the criteria, a list of top 30 building block candidates was produced that 1) exhibited multiple functionalities suitable for further conversion as derivatives or molecular families, 2) could be produced from both lignocellulosics and starch, 3) were C1-C6 monomers, 4) were not aromatics derived from lignin, and 5) were not already supercommodity chemicals. These are shown in Table 3.

Table 3 - Down Selection – Top 30 Results

Carbon Number	Potential Top 30 candidates
1	Carbon monoxide & hydrogen (syngas)
2	None
3	Glycerol, 3 hydroxypropionic acid, lactic acid, malonic acid, propionic acid, serine
4	Acetoin, aspartic acid, fumaric acid, 3-hydroxybutyrolactone, malic acid, succinic acid, threonine
5	Arabinitol, furfural, glutamic acid, itaconic acid, levulinic acid, proline, xylitol, xylonic acid
6	Aconitic acid, citric acid, 2,5 furan dicarboxylic acid, glucaric acid, lysine, levoglucosan, sorbitol

Of note, C2 compounds such as acetic acid and acetic anhydride, were considered to have lower potential and C3 compounds such as acetone which is already a petrochemical byproduct were not included.

In addition, commercial scale conversion of syngas to hydrogen, ammonia, methanol, alcohols and aldehydes, (oxosyntheses), and Fischer-Tropsch products already exist.

5 Selected Sugar-derived Chemicals

The second round of down selection from the top 30 building block candidates identified twelve sugar derived building blocks shown in Table 4.

Table 4 - The Top Sugar-derived Building Blocks

Building Blocks
1,4 diacids (succinic, fumaric and malic)
2,5 furan dicarboxylic acid
3 hydroxy propionic acid
aspartic acid
glucaric acid
glutamic acid
itaconic acid
levulinic acid
3-hydroxybutyrolactone
glycerol
sorbitol
xylitol/arabinitol

In some cases molecules have been grouped together because of the potential synergy related to their structures. These molecules can be 1) isomers, 2) interconverted to afford the same molecule, and/or 3) the derivatives pathway leading to essentially the same family of products. These candidate groups are xylitol/arabinitol and the 1,4-dicarboxylic acids, succinic, malic and fumaric acid.

Summary information about each building block is presented in two different formats. First, a table, or “bio” of qualitative properties was compiled characterizing each building block and derivative. For example, Table 5 shows the *bio* for the 3-HPA building block. Tables 6 and 7 show families of 3-HPA transformations that in these cases involve reduction and dehydration to the corresponding derivatives. The remaining building block and derivative pathway *bios* and further discussion appear in Section 9.

Table 5 - Sugar Transformation to 3-HPA

Building Block Pathway	Technical Barriers	Direct Uses of Building Block
Chemical – Unknown or multistep, costly process		
Biotransformation-Fermentation	Being done by industry. Fermentation pathway not known General needs in fermentation Improving microbial biocatalyst to 1) reduce other acid coproducts, 2) increase yields and productivities Lower costs of recovery process to reduce unwanted salts Scale-up and system integration issues	None

Table 6 - Reductive Transformation – 3HP to 1,3 PDO via catalytic dehydrogenation

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
1, 3-propanediol	Direct reduction of carboxylic acid with high selectivity Reduction at mild conditions – moderate hydrogen pressure, low T Tolerance to inhibitory elements or components of biomass based feedstocks Robust catalysts and catalyst lifetimes	Sorona fiber (new material)

Table 7 - Dehydrative Transformation – 3-HPA to acrylic acid via catalytic dehydration

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
Acrylate family	Selective dehydration without side reactions (high value need for biomass) New heterogeneous catalysts (i.e. solid acid catalysts) to replace liquid catalysts and to improve existing catalyst based systems	Contact lenses, diapers (Super Absorbent Polymers SAPs)

In addition to each *bio* summary a visual representation, or “star” diagram, was created for each building block. These diagrams use the star’s projections to represent the different pathways for converting building blocks to derivatives or secondary chemicals. The number and nature of each derivative pathway represents each building block’s potential value as a starting material for a variety of end uses. Unlike the building blocks, the derivatives can be categorized into two types. One type of derivative could be used as current replacements for industrial petrochemicals or biochemicals. The second type of derivative could serve as novel material with new performance characteristics that would allow new applications or create a new market segment.

Each transformation pathway was assessed for 4 characteristics including 1) current industrial use, 2) a transformation analogous to a currently known technology, 3) moderate process development requirement and 4) significant process development requirement.

The 3-hydroxypropionic acid star diagram is shown below in 2. The building block star diagrams for all the top 12 appear in Section 9. Derivatives are grouped by families as a function of chemical transformations.

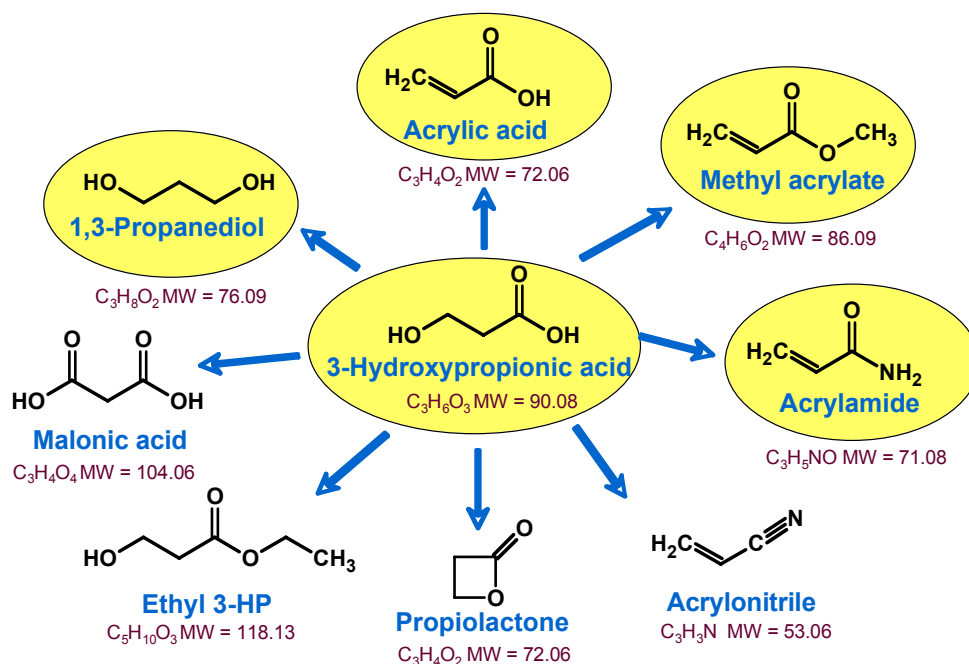


Figure 4 - Star Diagram of 3-Hydroxypropionic Acid

The circled derivatives are those in commercial use and produced in commodity-scale volume today. A dashed line indicates a lack of knowledge about how to undertake the proposed pathway. The team attempted to identify most of the derivative pathways that could be replacements for petrochemically-derived compounds and for novel compounds that have growth potential. Taken together the number of potential pathways and associated technical barriers for each *star* are an indication of the value of the candidate as a building block.

As part of the screening criteria the team used standard reference documents (listed in the Bibliography) and industrial chemistry experience to identify the potential transformation pathways. Four criteria were used to rank the potential building block candidates: 1) strategic fit for lignocellulosic and starch biomass within the biorefinery, 2) value of the building block and its derivatives as a replacement or novel chemical, 3) the technical complexity of each part of the pathway transformation (sugars to building blocks and building blocks to derivatives), and 4) the building block's potential to produce families or groups of similar derivatives. Each building block candidate was given a consensus score. From a statistical analysis 12 candidates ranked above average, three were at the mean (lactic, levoglucosan and lysine) with the remaining falling below the average. Tier one building blocks or those whose score was above average are listed in Table 4 above. Tier two candidates, gluconic acid, lactic acid, malonic acid, propionic acid, the triacids (citric and aconitic) xylonic acid, acetoin, furfural, levoglucosan, lysine, serine and threonine were judged to have somewhat lower potential.

6 Syngas Results – Top Products

For the purposes of this study hydrogen and methanol comprise the best near-term prospects for biobased commodity chemical production because obtaining simple alcohols, aldehydes, mixed alcohols and Fischer-Tropsch liquids from biomass are not economically viable and require additional development. Therefore no further down select from syngas-derived products was undertaken. This determination was based on a review of the literature and a progress review of the OBP Thermochemical Platform R&D at NREL in August 2003. The review identified gas cleanliness as a key barrier to economic production of syngas from biomass. A comprehensive report including economic analysis, technical challenges and energy impacts of syngas to liquid processes is available.¹

¹P. Spath and D. Dayton Preliminary Screening – Technical and Economic Assessment of Synthesis Gas to Fuels and Chemicals with Emphasis on the Potential for Biomass-derived Syngas, NREL Technical Report NREL/TP-510-34929 , December 2003)

7 Pathways and Challenges

Before common technical barriers can be listed and described, the team identified the most viable biological and chemical transformation pathways from sugars to building blocks. A large number of sugar to building block transformations can be done by aerobic fermentation employing fungi, yeast or bacteria. Chemical and enzymatic transformations are also important process options. It should be noted however, that pathways with more challenges and barriers are less likely be considered as viable industrial processes. Currently known sugars to building blocks that are commercially available are listed in Table 8.

Similarly, the team examined the most common transformations involved in converting building blocks to derivatives. Chemical reduction, oxidation, dehydration, bond cleavage, and direct polymerization predominated. Here enzymatic biotransformations comprised the largest group of biological conversions. Additionally some biological conversions can be accomplished without the need for an intermediate building block. 1,3-propanediol is a case in point where a set of successive biological processes convert sugar directly to an end product.

Each pathway has its own set of advantages and disadvantages. Biological conversions, of course, can be tailored to result in a specific molecular structure but the operating conditions must be relatively mild. Chemical transformations can operate at high throughput but less conversion specificity is achieved.

Prioritizing the technical barrier categories for chemicals is less clear. For example, biomass is already highly oxidized (contains significant fractions of oxygen) so the numbers of transformations requiring oxidation are relatively low. On the other hand those requiring hydrogenation and dehydration (adding hydrogen or removing oxygen) were much higher. This belies the fact that oxidation is still a difficult thing to do and possesses significant benefits not only for sugars but also oils, lignin, and other biomass components.

Table 8 – Pathways to Building Blocks from Sugars

Identification of Actual and Potential Pathways to Building Blocks from Sugars						
Building Blocks	Yeast or Fungal	Bacterial	Yeast or Fungal	Bacterial	Chemical & Catalytic Processes	Biotransformation
	AEROBIC FERMENTATIONS		ANAEROBIC FERMENTATIONS		CHEM-Enzyme TRANSFORMATIONS	
3 Carbon	Commercial Product - C	Commercial Product - C	Commercial Product - C	Commercial Product - C	Commercial Product - C	Commercial Product - C
3-Hydroxy propionic acid	X	X				
Glycerol	X	X	X	X	C	
Lactic acid	X		X	C		
Malonic acid	X				X	
Propionic acid				X		
Serine	X	C				C
4 Carbon						
3-Hydroxy butyrolactone					X	
Acetoin	X	X		X		
Aspartic Acid	X	X				X
Fumaric Acid	X	X			X	
Malic acid	X	X				
Succinic acid	X	X		X		X
Threonine	X	C				
5 Carbon						
Arabitol	X		X		C	X
Furfural					C	
Glutamic	X	C				
Itaconic Acid	C					
Levulinic acid					X	
Xylitol	X		X	X		C
6 Carbon						
2,5 Furan dicarboxylic acid					X	
Aconitic acid	X					
Citric acid	C					
Glucaric acid	X	X			X	
Gluconic acid	C	X				X
Levogluconan					X	
Lysine	X	C				
Sorbitol	X	X			C	X
Number in each Pathway category*	21	14	4	6	11	7
Commercial processes	3	4	0	1	4	2

* All of the top 30 were used in the evaluation but only those involved in the final downselect are shown here, hence total pathway in each category numbers may not add up on this specific chart.

8 Moving Forward

In reality, the first screen choices should not be viewed as an elimination but as generic guidance using criteria that allowed the selection of a top twelve list. Clearly, the sugar focus of this analysis limited the number of opportunities for value-added chemicals beyond C6 compounds. For example, aromatics comprise a very large commodity market for polymers and surfactants. Polysaccharides are a growing market segment because of their potentially valuable properties in various applications such as enhanced oil recovery and paper/metal finishing. Oils, although produced in an established industry, have a broad range of opportunities for diverse market applications. Lignin can afford the entire family of aromatic compounds that are difficult to produce via sugars or oils. It would be worthwhile to assess the potential value of products derived from both oils and lignin as has been done here for sugars and syngas.

We also know that new knowledge and better technologies are needed in dealing with chemical transformations that involve milder oxidations, selective reductions and dehydrations, better control of bond cleavage, and improvements to direct polymerization of multifunctional monomers. For biological transformations, we need better pathway engineering of industrial hosts, better understanding of metabolic pathways and cell biology, lower downstream recovery costs, increased utility of mixed sugar streams, improved molecular thermal stability; and better understanding of enzyme functionality. While it is possible to prepare a very large number of molecular structures from the top building blocks, we have almost no publicly available information about these molecules' behavior, material performance or industrial processing properties. Hence, a comprehensive database on biopolymer performance characteristics would prove extremely useful to both the public and private sector.

It is highly likely we will need to expand the suites of potential pathways and increase our understanding of all the technical barriers beyond the ones summarized here. This knowledge would also lead to a better definition of which biobased feedstock materials hold the most promise as economic drivers for an integrated biorefinery.

9 Top 12 Candidate Summary Bios

9.1 **Four Carbon 1,4-Diacids (Succinic, Fumaric, and Malic)**

9.1.1 Pathways to Building Block From Sugars

Table 9 – Pathways to Building Block From Sugars [Four Carbon 1,4 Diacids (Succinic, Fumaric, and Malic)]

Type of pathway	Technical Barriers	Direct Uses of Building Block
Chemical – None		
Biotransformation - Fermentation to overproduce C4 diacids from Krebs cycle pathways	Improving microbial biocatalyst to 1) reduce acetic acid coproducts, 2) increase yields and productivities Lower costs of recovery process to reduce unwanted salts Scale-up and system integration issues	

9.1.2 Primary Transformation Pathway(s) to Derivatives

Table 10 – Family 1: Reductions [Primary Transformation Pathway(s) to Derivatives Four Carbon 1,4-Diacids (Succinic, Fumaric, and Malic)]

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
THF, BDO, GBL Family	Selective reductions: controlling reduction of acids to alcohols, lactones and furans Operation at mild conditions (pressure, T, etc.) Catalyst tolerance to inhibitory compounds and catalyst lifetime	Solvents, fibers such as lycra

Table 11 – Family 2: Reductive Aminations [Primary Transformation Pathway(s) to Derivatives - Four Carbon 1,4-Diacids (Succinic, Fumaric, and Malic)]

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
Pyrrolidinone Family	Selective reductions of acid salts Operation at mild conditions (pressure, T, etc.) Catalyst tolerance to inhibitory compounds Catalyst lifetime in continuous processes	Green solvents, water soluble polymers (water treatment)

Table 12 – Family 3: Direct Polymerization [Primary Transformation Pathway(s) to Derivatives - Four Carbon 1,4-Diacids (Succinic, Fumaric, and Malic)]

Derivative or Derivative Family	Technical barriers	Potential use of derivative product
Straight chain polymers	Commercial polymer processes	Fibers (lycra, others)
Branched polymers	Selective esterifications to control branching Control of molecular weight & properties	TBD

Building Block: Four Carbon Diacids (Succinic, Fumaric, and Malic)

Primary Derivatives:

Family 1: Reductions

Family 2: Reductive Aminations

Family 3: Direct Polymerization

9.1.3 Building Block Considerations

The family of 4-carbon diacids is best grouped together, since their production arises from very similar biochemical paths. For the purposes of this summary, succinic acid will be used as a prototypical example, but the concepts described herein apply equally well to fumaric and malic acid. Star diagrams for fumaric and malic acid are found elsewhere in this report. Four carbon dicarboxylic acids have the potential to be a key building blocks for deriving both commodity and specialty chemicals. The basic chemistry of succinic acid is similar to that of the petrochemically derived maleic acid/anhydride. Succinic acid is produced biochemically from glucose using an engineered form of the organism *A. succiniciproducens* and most recently via an engineered *Eschericia coli* strain developed by DOE laboratories and licensed to a small business. The chemistry of succinic acid to the primary families of derivatives is shown in Figure 5.

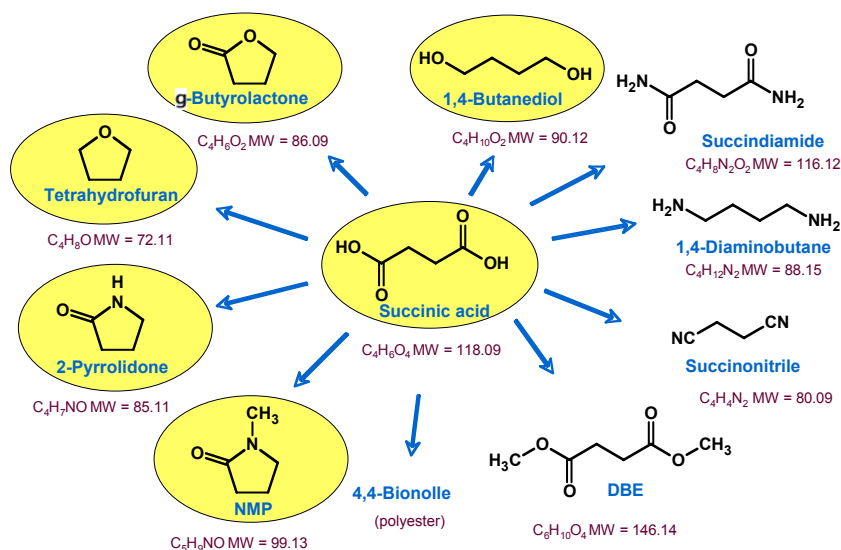


Figure 5 - Succinic Acid Chemistry to Derivatives

The major technical hurdles for the development of succinic acid as a building block include the development of very low cost fermentation routes. There are currently two organisms under active development for the fermentation of sugars (both C6 and C5) to produce succinic acid. Based on the available information in the literature regarding these two organisms significant improvement in the fermentation is still required to be competitive with petrochemical routes. Figure 6 depicts a simplified PFD of glucose fermentation to succinic acid.

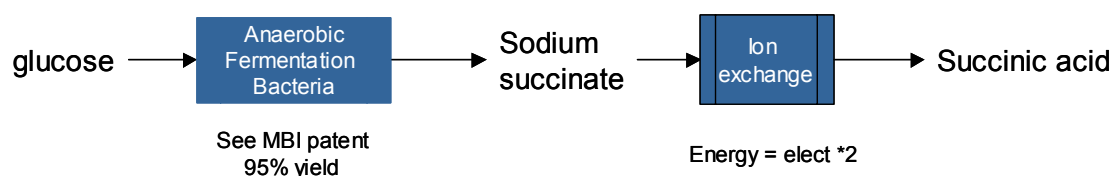


Figure 6 - Simplified PFD of Glucose Fermentation to Succinic Acid

The major elements of improvement in the fermentation include the following:

Productivity: Productivity improvements are required to reduce the capital and operating costs of the fermentation. A minimum productivity of 2.5 g/L/hr needs to be achieved in order for the process to economically competitive.

Nutrient Requirements: It is essential for commercial fermentations to be run using minimal nutrients. Expensive nutrient components such as yeast extract and biotin must be eliminated. The nutrient requirements should be limited to the use of corn steep liquor or equivalent.

Final Titer: Final titer is also important when considering overall process costs. This is not a showstopper but a high final titer will reduce overall separation and concentrating costs.

pH Considerations: In an ideal situation the fermentation would be run at low pH, most preferably without requiring any neutralization. The cost of neutralization is not necessarily cost prohibitive, but the conversion of the salt to the free acid does add significant costs. If derivatives such as BDO, THF and GBL are going to be competitive from a cost perspective then low pH fermentation will be essential.

9.1.4 Derivative Considerations

A primary technology for use of succinic acid is selective reduction to give the well-known butanediol (BDO), tetrahydrofuran (THF) and gamma-butyrolactone (GBL) family. The hydrogenation/reduction chemistry for the conversion of succinic acid to BDO, THF and BGL is well known and is similar to the conversion of maleic anhydride to the same family of compounds. The only real technical consideration here is the development of catalysts that would not be affected by impurities in the fermentation. This is a significant challenge but would not necessarily be a high priority research item until the costs of the fermentation are substantially reduced.

Pyrrolidinones are materials that can be derived from GBL, and address a large solvent market. Reaction of GBL with various amines leads to the production of materials such as pyrrolidinone and N-methylpyrrolidinone. Succinic acid can also be converted more directly to pyrrolidinones through the fermentative production of diammonium succinate. One advantage with the fermentation derived succinic acid is that the conversion of diammonium succinate to the pyrrolidones directly could offer a significant cost advantage. This would eliminate the need for low a low pH fermentation for the direct production of succinic acid.

Similar chemistry can be applied to transformations of malic and fumaric acid. Of particular interest is the ability to use selective reduction for the conversion of fumaric acid to succinic, and the use of malic acid in the production of substituted THF or NMP derivatives.

9.1.5 Overall Outlook

There is a significant market opportunity for the development of biobased products from the C4 building block diacids. The major challenges are primarily associated with reducing the overall cost of the fermentation. In order to competitive with petrochemicals derived products the fermentation cost needs to be at or below \$0.25/pound. This is a significant technical challenge and should be undertaken with a long-term perspective. When considered in aggregate, the diacid family offers access to a wide range of products that address a number of high volume chemical markets.

9.2 2,5-Furan dicarboxylic acid (FDCA)

9.2.1 Pathways to Building Block from Sugars

Table 13 –Pathways to Building Block From Sugars [2,5-Furan dicarboxylic Acid (FDCA)]

Type of pathway	Technical Barriers	Direct Uses of Building Block
Chemical – Oxidative dehydration of C6 sugars	Dehydration Selective dehydrations without side reactions Dehydration steps to anhydrides or lactones New heterogeneous catalyst systems (solid acid catalyst) to replace liquid catalysts Oxidations Alcohols (ROH) to acids (RCOOH) Avoiding exotic oxidants in favor of air, oxygen, dilute hydrogen peroxide Tolerance to inhibitory components of biomass processing streams Oxidation of aldehydes to acids and alcohols to aldehydes	PET analogs with potentially new properties (bottles, films, containers)
Biotransformation - Possibly enzymatic conversions	Unknown	

9.2.2 Primary Transformation Pathway(s) to Derivatives

Table 14 – Family 1: Reduction [Primary Transformation Pathway(s) to Derivatives: 2,5-Furan dicarboxylic Acid (FDCA)]

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
Diols and Aminations	Selective reduction of acids in presence of alkenes. Direct reduction of carboxylic acids to alcohols. Knowledge of properties of polymer derivatives	New polyesters and nylons with new properties likely for fiber applications
Levulinic and Succinic Acids	Selective catalytic tools	All uses of succinic and levulinic

Table 15 – Family 2: Direct Polymerization [Primary Transformation Pathway(s) to Derivatives: 2,5-Furan dicarboxylic Acid (FDCA)]

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
Polyethylene terephthalate analogs.	Reactivity of monomer. Controlling rates Selective esterifications to control branching Control of molecular weight & properties	Furanoic polyesters for bottles, containers, films,
Furanoic Polyamines	Reactivity of monomer. Controlling rates Selective esterifications to control branching Control of molecular weight & properties	Polyamide market for use in new nylons

Building block: 2,5 – Furan dicarboxylic acid (FDCA)

Family 1 – Reduction

Family 2 – Direct Polymerization

9.2.3 Building Block Considerations

Dehydration of the sugars available within the biorefinery can lead to a family of products, including dehydrosugars, furans, and levulinic acid. FDCA is a member of the furan family, and is formed by an oxidative dehydration of glucose. The process has been reported to proceed using oxygen, or electrochemistry. The conversion can also be carried out by oxidation of 5-hydroxymethylfurfural, which is an intermediate in the conversion of 6-carbon sugars into levulinic acid, another member of the top 10. Figure 7 describes some of the potential utility of FDCA.

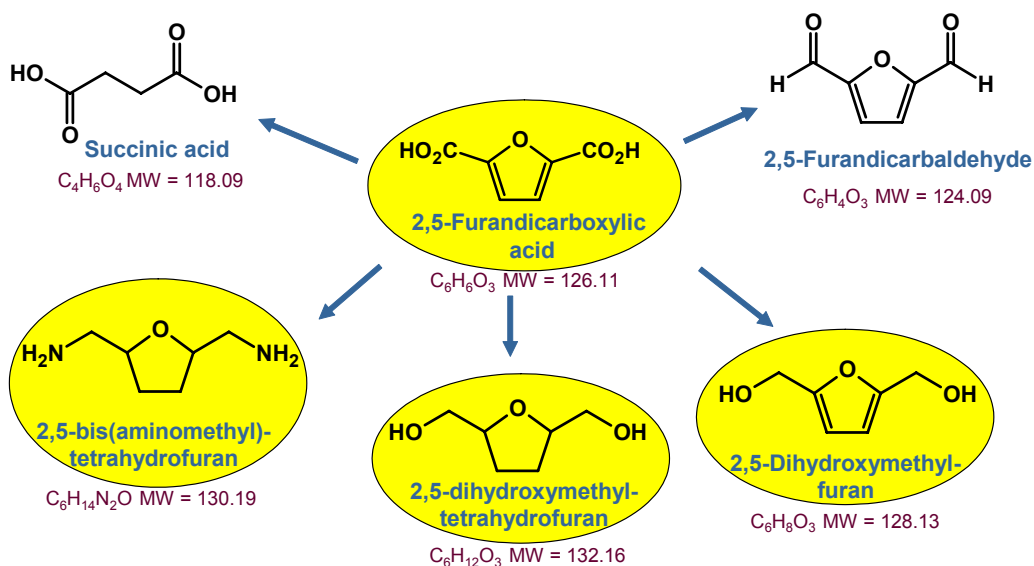


Figure 7 - Derivatives of FDCA

9.2.4 *Derivative Considerations*

FDCA has a large potential as a replacement for terephthalic acid, a widely used component in various polyesters, such as polyethylene terephthalate (PET) and polybutyleneterephthalate (PBT). PET has a market size approaching 4 billion lb/yr, and PBT is almost a billion lb/yr. The market value of PET polymers varies depending on the application, but is in the range of \$1.00 – 3.00/lb for uses as films and thermoplastic engineering polymers. The versatility of FDCA is also seen in the number of derivatives available via relatively simple chemical transformations. Selective reduction can lead to partially hydrogenated products, such as 2,5-dihydroxymethylfuran, and fully hydrogenated materials, such as 2,5-bis(hydroxymethyl)tetrahydrofuran. Both of these latter materials can serve as alcohol components in the production of new polyester, and their combination with FDCA would lead to a new family of completely biomass-derived products. Extension of these concepts to the production of new nylons, either through reaction of FDCA with diamines, or through the conversion of FDCA to 2,5-bis(aminomethyl)tetrahydrofuran could address a market of almost 9 billion lb/yr, with product values between \$0.85 and 2.20/lb, depending on the application. FDCA can also serve as a starting material for the production of succinic acid, whose utility is detailed elsewhere in this report.

The primary technical barriers to production and use of FDCA include development of effective and selective dehydration processes for sugars. The control of sugar dehydration could be a very powerful technology, leading to a wide range of additional, inexpensive building blocks, but it is not yet well understood. Currently, dehydration processes are generally nonselective, unless, immediately upon their formation, the unstable intermediate products can be transformed to more stable materials. Necessary R&D will include development of selective dehydration systems and catalysts. FDCA formation will require development of cost effective and industrially viable oxidation technology that can operate in concert with the necessary dehydration processes.

A number of technical barriers also exist with regard to the use of FDCA (and related compounds) in the production of new polymers. Development and control of esterification reactions, and control of the reactivity of the FDCA monomer will be of great importance. Understanding the link between the discrete chemistry occurring during polymer formation, and how this chemistry is reflected in the properties of the resulting polymer will provide useful information for industrial partners seeking to convert this technology into marketplace products.

9.2.5 *Overall Outlook*

The utility of FDCA as a PET/PBT analog offers an important opportunity to address a high volume, high value chemical market. To achieve this opportunity, R&D to develop selective oxidation and dehydration technology will need to be carried out. However, the return on investment might have applicability of interest to an important segment of the chemical industry.

9.3 3-Hydroxypropionic acid (3-HPA)

9.3.1 Pathways to Building Block from Sugars

Table 16 - Pathways to Building Block from Sugars (3-HPA)

Type of pathway	Technical Barriers	Direct Uses of Building Block
Chemical – Unknown or multistep, costly process		
Biotransformation-Fermentation	Being done by industry. Fermentation pathway not known General needs in fermentation Improving microbial biocatalyst to 1) reduce other acid coproducts, 2) increase yields and productivities Lower costs of recovery process to reduce unwanted salts Scale-up and system integration issues	None

9.3.2 Primary Transformation Pathway(s) to Derivatives

Table 17 – Family 1: Reductions [Primary Transformation Pathway(s) to Derivatives (3-HPA)]

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
1, 3 propane diol	Selective direct reduction of carboxylic acids Reduction at mild conditions – atmospheric pressure, low T Tolerance to inhibitory elements or components of biomass based feedstocks robust catalysts and catalyst lifetimes	Sorona fiber

Table 18 – Family 2: Dehydration [Primary Transformation Pathway(s) to Derivatives (3-HPA)]

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
Acrylate family	Selective dehydration without side reactions (high value need for biomass) New heterogeneous catalysts (i.e. solid acid catalysts) to replace liquid catalysts and to improve existing catalyst based systems	Contact lenses, diapers (Super Absorbent Polymers SAPs)

Building Block: 3-Hydroxypropionic acid (3-HPA)

Primary Derivatives:

Family 1: Reductions

Family 2: Dehydrations

3-Hydroxypropionic acid is a C3 building block and has the potential to be a key building block for deriving both commodity and specialty chemicals. The basic chemistry of 3-HPA is not represented by a current petrochemically derived technology. The chemistry of 3-HPA to the primary families of derivatives is shown in Figure 9.

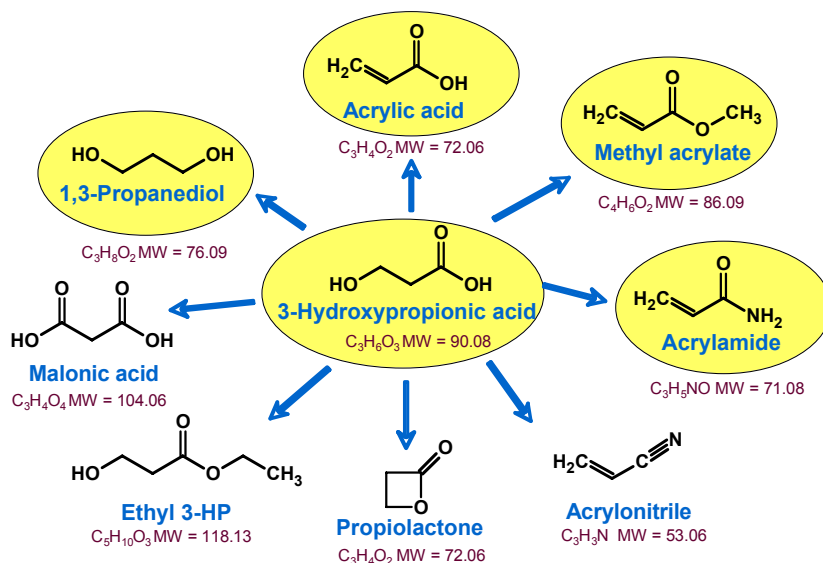


Figure 8 – Derivatives of 3-HPA

The major technical hurdles for the development of 3-HPA as a building block include the development of very low cost fermentation routes. Major technical considerations are the development of an organism with the appropriate pathways. In principle, the fermentation should be equivalent to lactic acid from a yield perspective. The major elements of improvement in the fermentation include the following considerations.

9.3.3 Building Block Considerations

Productivity: Productivity improvements are required to reduce the capital and operating costs of the fermentation. A minimum productivity of 2.5 g/Lhr. needs to be achieved in order to economically competitive.

Pathway Engineering: It will be necessary to engineer the appropriate pathway in an organism to produce 3-HPA. If successful the yield of the fermentation should be equivalent to lactic acid.

Nutrient Requirements: It is essential for commercial fermentations to be run using minimal nutrients. Expensive nutrient components such as yeast extract and biotin must be eliminated. The nutrient requirements should be limited to the use of corn steep liquor or equivalent if at all possible.

Final Titer: Final titer is also important when considering overall process costs. This is not a showstopper but a high final titer will reduce overall separation and concentrating costs.

pH Considerations: In an ideal situation the fermentation could be run at low pH, most preferably without requiring any neutralization. The cost of neutralization is not necessarily cost prohibitive, the conversion of the salt to the free acid does add significant costs. If derivatives such as BDO, THF and GBL are going to be competitive from a cost perspective then low pH fermentation will be essential.

9.3.4 Derivative Considerations

Family 1: 1,3-PDO

The hydrogenation/reduction chemistry for the conversion of 3-HPA to 1,3-PDO will require the development of new catalyst systems that are capable of the direct reduction of carboxylic acid groups to alcohols. A second option is to esterify the acid to an ester and reduce the ester. This may be technically easier but will add costs to the process. 1,3-PDO has been widely publicized by DuPont as a potential monomer for use in fibers for carpet. The new properties imparted by 1,3-PDO include better dye ability, and improved elasticity. Direct reduction of 3-HPA from a fermentation broth will require the development of robust catalysts that are not susceptible to fouling from impurities.

Family 2: Acrylates

The dehydration of 3-HPA to the family of acrylates including acrylic acid and acrylamide will require the development of new acid catalyst systems that afford high selectivity. In addition, there is the potential for polymerization and this must be avoided during the dehydration. One advantage for the production of acrylamide is that the starting material could be ammonium 3-HPA. Starting with ammonium 3-HPA would eliminate the need for low pH fermentation.

9.3.5 Overall Outlook

There is a significant market opportunity for the development of biobased products from the C3 building block 3-HPA. The major challenges include the development of a low cost fermentation and a family of catalysts for the conversion of 3-HPA to desired products. In order to be competitive with petrochemical routes to acrylates both the fermentation and catalysis must afford high yields.

9.4 Aspartic acid

9.4.1 Pathways to Building Block

Table 19 – Pathways to Building Block - Aspartic Acid

Type of pathway	Technical Barriers	Direct Uses of Building Block
Chemical – Amination of fumaric acid with ammonia	Asymmetric aminations	Salts for chelating agent. Sweeteners

Type of pathway	Technical Barriers	Direct Uses of Building Block
Biotransformation - Conversion of oxaloacetate in the Krebs cycle can yield aspartic acid either fermentatively or via enzymatic conversion	Pathway engineering of biocatalytic organisms to overproduce oxaloacetate without compromising viability of organism Managing operating environment Enzymatic oxidation of oxaloacetate Need for low cost recovery Low cost sugars	

9.4.2 Primary Transformation Pathway(s) to Derivatives

Table 20 - Family 1: Reductions [Primary Transformation Pathway(s) to Derivatives - Aspartic Acid]

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
Amine butanediol, amine tetrahydrofuran, amine (-butyrolactone)	Analogous to those for succinic, malic, and fumaric transformations: Selective reductions Operation at mild conditions (atmospheric pressure, low T, etc.) Catalyst tolerance to inhibitory compounds acceptable catalyst lifetimes	Amino analogs of C4 1,4 dicarboxylic acids

Table 21 –Family 2: Dehydration - [Primary Transformation Pathway(s) to Derivatives - Aspartic Acid]

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
Aspartic anhydride	Selective dehydrations without side reactions New heterogeneous catalyst systems (solid acid catalyst) to replace liquid catalyst systems	New area

Table 22 – Family 3: Direct Polymerization [Primary Transformation Pathway(s) to Derivatives - Aspartic Acid]

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
Polyaspartic	Selective esterifications to control branching Control of molecular weight & properties	New area

Building Block: Aspartic Acid

Primary Derivatives:

Family 1: Selective Reductions

Family 2: Dehydration to anhydrides

Family 3: Direct Polymerizations

Aspartic acid is a 4-carbon amino acid that is an essential part of metabolism among many species, including humans, for protein production. There are several configurations of aspartic acid produced; however, L-aspartic is by far the most common. L-aspartic acid is primarily used to produce aspartame, a synthetic sweetener. The chemistry of aspartic acid to the primary families of derivatives is shown in Figure 9.

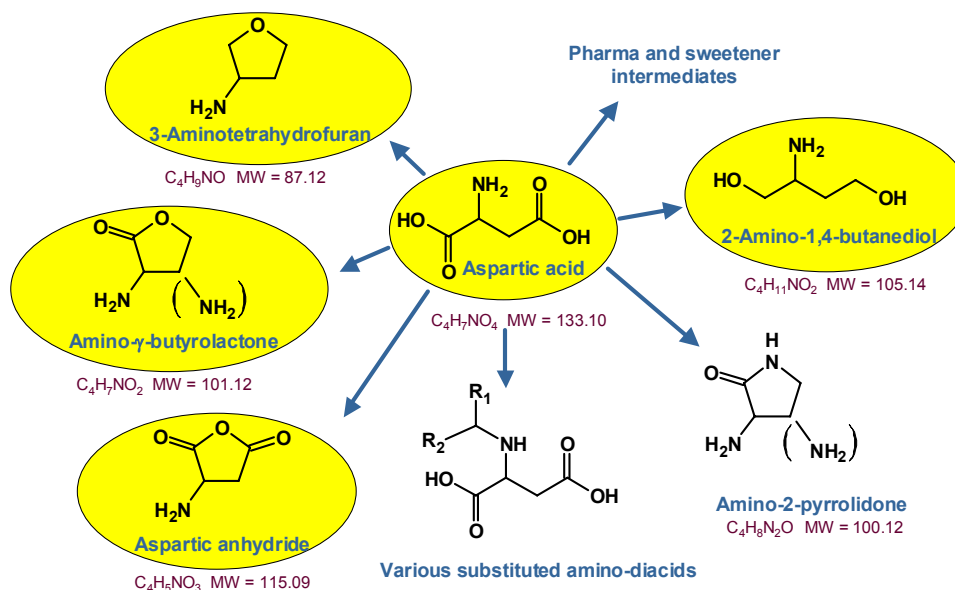


Figure 9 - Aspartic Acid Chemistry to Derivatives

There are 4 primary routes to producing L-aspartic acid: 1) chemical synthesis, 2) protein extraction, 3) fermentation, and 4) enzymatic conversion. The preferred method currently is the enzymatic route, reacting ammonia with fumaric acid, catalyzed by a lyase enzyme. The advantages to this pathway include high product concentration, high productivity, fewer byproducts, and ease of separation (crystallization).

The major technical hurdles for the development of aspartic acid as a building block involve developing a direct fermentation route (using sugar substrate) that is cost-competitive with the existing enzymatic conversion process. Direct fermentation routes are not cost-competitive yet, but the use of biotechnology holds promise to overcome this obstacle. A second strategy for reducing the cost of aspartic acid is to make improvements to the current technology. The primary focus of this effort would be to reduce the cost of fumaric acid, which is currently used as the feedstock for producing aspartic acid.

9.4.3 Building Block Considerations

High fermentation yields and product recovery are the two primary technical goals to strive for. A direct fermentation using sugar substrate could potentially be cheaper than fumaric

acid and ammonia feedstocks if the production technical performance can be improved. Bayer AG also has a competing route using maleic anhydride that may be potentially cheaper. Producing fumaric acid at a lower cost could have a near term impact on reducing the cost of aspartic acid. This strategy would have the advantage of using existing capital and infrastructure.

Productivity: Productivity improvements are required to reduce the capital and operating costs of the fermentation. The existing enzyme route through fumaric acid achieves productivities which are satisfactory for specialty applications. But for commodity-scale applications further improvements in productivity will be required.

Separation/recovery costs: The cost of separating aspartic acid from the fermentation broth could be a potential showstopper. The competing enzymatic route with ammonia and fumaric has high product concentration and uses crystallization to easily separate the final product. However, crystallization can be an expensive processing step and fermentation broth separation techniques may be able to compete through research.

Final Titer: Final titer is very important when considering overall process costs. A high final titer will reduce overall separation and concentrating costs.

Nutrient Requirements: If low-cost nutrients can be used, the production economics of aspartic acid can be significantly reduced.

9.4.4 Derivative Considerations

Family 1: Amino analogs of C4-dicarboxylic acids

Selective reduction of the carboxylic acids of aspartic acid would produce analogs to current high volume chemicals such as 1,4-butanediol, tetrahydrofuran and gamma-butyrolactone. These analogs have the potential for large market polymer and solvent applications. The specific technology drivers are developing the ability to selectively reduce the carboxylic acids in the presence of amine groups. The ability to do this in high selectivity and under mild conditions could make the derivatives competitive with the analog C4 compounds.

Family 2: Anhydrides

The selective dehydration to form the anhydrides is generally considered to be a thermal process in the presence of an acid-based catalyst. Development of new catalysts that allow for selective dehydration without side reactions will be critical for low cost anhydride formation.

Family 3: Direct polymerization to new polymers

Synthesis of biodegradable specialty polymers - polyaspartic acid and polyaspartates (PAA) - would be substitutes for polyacrylic acid and polycarboxylates. This synthesis is not expected to be difficult, but has not been undertaken. The polymerization would be analogous to polyglutamic acid that is a commercial process.

9.4.5 Overall Outlook

Improved fermentation for the direct production of aspartic acid or improved fermentation for reducing the costs of fumaric acid will depend on the utilization of both genetic engineering and traditional strain improvement technology.

The L-aspartic market is expected to grow 2-3% annually worldwide between now and 2006. New biodegradable specialty polymers (polyaspartic acid and polyaspartates) offer new potential markets as substitutes for polyacrylic acid and polycarboxylates. Applications may include detergents, water treatment systems, corrosion inhibition, and super-absorbent polymers.

9.5 Glucaric acid

9.5.1 Pathways to Building Block From Sugars

Table 23 – Pathway to Building Block From Sugars [Glucaric Acid]

Type of pathway	Technical Barriers	Direct Uses of Building Block
Chemical- One step nitric acid oxidation of starch or Catalytic oxidation of starch with bleach (basic)	Selective oxidation of alcohols (ROH) to acids (RCOOH) Avoiding exotic oxidants in favor of air, oxygen, dilute hydrogen peroxide. Lowering concentrations of oxidants Development of heterogeneous catalyst systems Tolerance to inhibitory elements of biomass based feedstocks	
Biotransformation- Not known		

9.5.2 Primary Transformation Pathway(s) to Derivatives

Table 24 – Family 1 - Dehydration [Primary Transformation Pathway(s) to Derivatives - Glucaric Acid]

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
Lactones	Selective dehydration without side reactions Dehydration to anhydrides or lactones New heterogeneous catalysts (i.e. solid acid catalysts) to replace liquid catalysts and to improve existing catalyst based systems	Solvents

Table 25 – Amination and Direct Polymerization [Primary Transformation Pathway(s) to Derivatives - Glucaric Acid]

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
Polyglucaric esters and amides	Manage rates Selective esterifications to control branching Control of molecular weight & properties	Nylons or different properties (i.e., like Kevlar vs. carpet fiber)

Building Block: Glucaric Acid
Family 1 – Dehydration
Family 2 – Direct Polymerization

9.5.3 Building Block Considerations

Glucaric acid is a member of a much larger family of materials known as oxidized sugars. These materials represent a significant market opportunity. For example, oxidation of glucose to glucaric acid (worldwide consumption, 92 million lb in 1998; F. Dubois, A. DeBoo, A. Kishi, Chemical Economics Handbook, "Chelating Agents", 515.5000, March 2000) can be carried out using chemical or biochemical catalysis in high yield.

In contrast, production of glucaric acid as a building block is more difficult. However, the value to the biorefinery of converting cheap glucose into glucaric acid arises from two features: 1) glucaric acid can serve as a starting point for the production of a wide range of products with applicability in high volume markets and 2) development of efficient processes for production of glucaric acid will also be applicable to efficient oxidation of other inexpensive sugars studied in this evaluation, such as xylose or arabinose. Glucaric can be produced from glucose by oxidation with nitric acid. **Error! Reference source not found.** summarizes the product and its derivative opportunities.

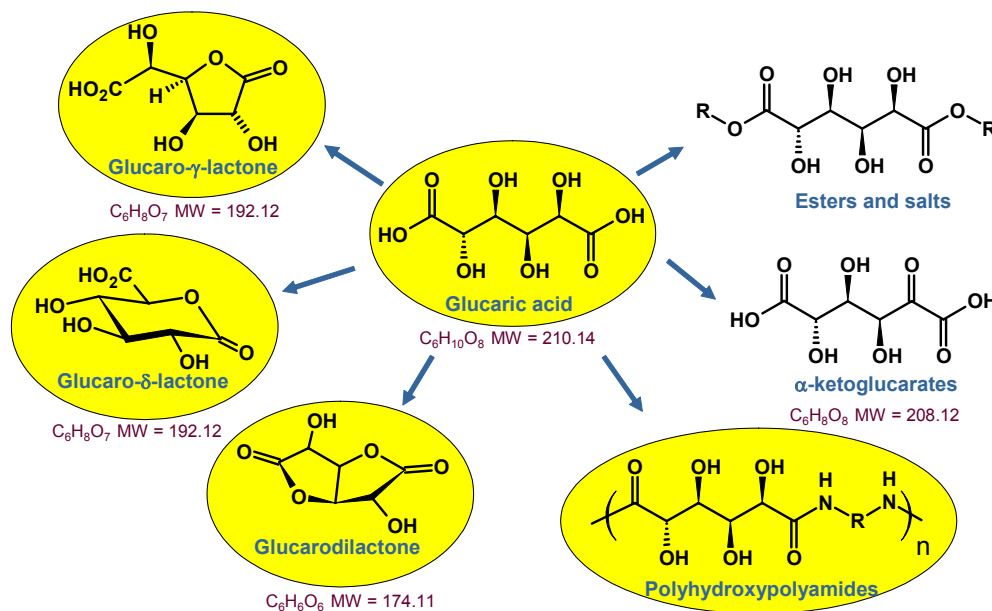


Figure 10- Derivatives of Glucaric Acid

9.5.4 Derivative Considerations

Ready availability of aldaric acids would form the basis of a new family of renewable building blocks derived from carbohydrates. A significant opportunity exists in the production of new nylons (polyhydroxypolyamides). The combination of cheap glucose with currently available diamines could address a market of over 9 billion lb/yr with values between \$0.85 and \$2.20/lb, depending on application. Glucaric acid (and its esters) is also a potential starting

material for new types of hyperbranched polyesters addressing markets of similar sizes to nylons with a similar value structure. Finally, glucaric acid could also address the very large detergent surfactant market, as it should exhibit useful chelating properties for cations. Simple chemical transformation will lead to α -ketoglucarates, starting points for the production additional types of new polymeric materials. It is important to note that success with glucaric acid is not limited to the use of glucose alone. Rather, technology development for this material will be directly applicable to a wide range of other materials evaluated in this analysis, including xylose, arabinose, and glycerol. A very wide range of products and opportunities will be available from an investment in glucaric acid R&D.

The technical barriers for this work include development of efficient and selective oxidation technology for glucose, and eliminating the need to use nitric acid as the oxidant. Recent work indicates that new catalytic processes using inexpensive oxidants may pave the way for high yield production of glucaric acid from glucose. Further technical barriers include development of selective methods for sugar dehydration to transform glucaric into sugar lactones, particularly glucaric dilactone, an analog of isosorbide, another compound included in this evaluation.

9.5.5 Overall Outlook

Selective oxidation of an inexpensive sugar or sugar source to a single compound is analogous to conversion of complex starting petrochemicals to single and much simpler building blocks. Success in development of glucaric acid production and new derivatives will have broad application to the sugar platform and will address high volume and high value markets. Technology specific to glucaric acid will also be applicable to the production of xylaric and arabinaric acids, two other compounds ranked highly within this evaluation, making this portion of the biorefinery R&D effort significant.

9.6 Glutamic acid

9.6.1 Pathways to Building Block From Sugars

Table 26 – Pathways to Building Block From Sugars [Glutamic Acid]

Type of pathway	Technical Barriers	Direct Uses of Building Block
Chemical- None		
Biotransformation- Fermentation product	Improving microbial biocatalyst to 1) reduce other acid coproducts, 2) increase yields and productivities Better control of operating environment Lower costs of recovery process to reduce unwanted salts Scale-up and system integration issues	

9.6.2 Primary Transformation Pathway(s) to Derivatives

Table 27 – Family 1: Reductions [Primary Transformation Pathway(s) to Derivatives – Glutamic Acid]

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
Diols (1,5-propanediol), diacids (1,5-propanediacid), aminodiol (5-amino, 1-butanol)	Selective deamination, reduction and reductive deamination Reduction at mild conditions – atmospheric pressure, low T Managing acid salts Tolerance to inhibitory elements or components of biomass based feedstocks– robust catalysts	Monomers for polyesters and polyamides.

Building Block: Glutamic Acid

Primary Derivatives:

Family 1: Hydrogenation/Reduction

Glutamic acid is a five-carbon amino acid and has the potential to be a novel building block for five carbon polymers. The building block and its derivatives have the potential to build similar polymers but with new functionality to derivatives of the petrochemicals derived from maleic anhydride. These polymers could include polyesters and polyamides. The chemistry of glutamic acid and the routes to the primary families of derivatives is shown in Figure 11.

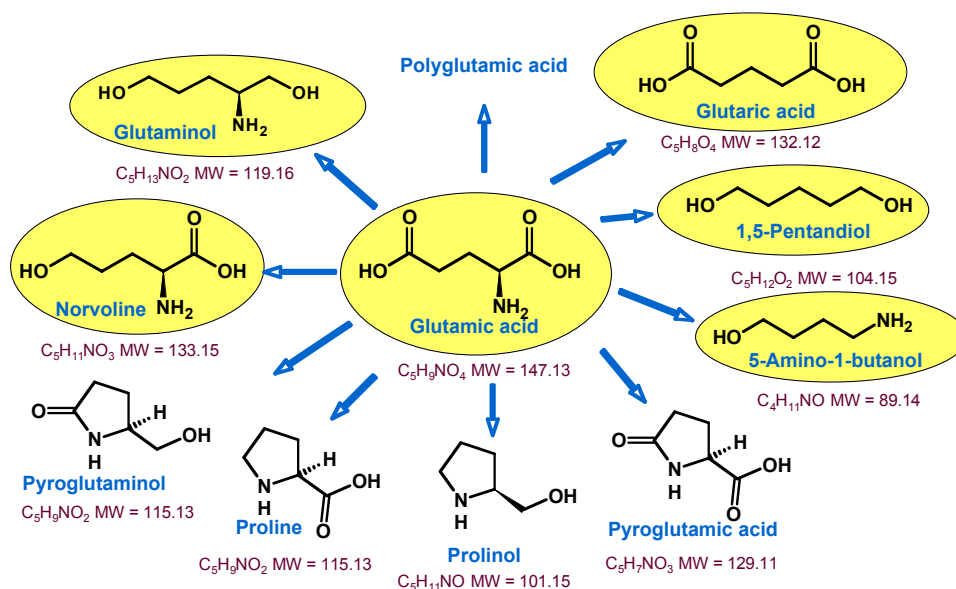


Figure 11 - Glutamic Acid and its Derivatives

The major technical hurdles for the development of glutamic acid as a building block include the development of very low cost fermentation routes. There are currently several fermentation routes for the production of sodium glutamate (MSG). These routes are all based on the production of the sodium salt. One of the major challenges for the development of a low cost fermentation is to develop an organism that can produce glutamic acid as the free acid. This would eliminate the need for neutralization and substantially reduce the costs of purification and conversion of the sodium salt to the free acid. Additional improvements in the fermentation would include increasing the productivity of the organism and improving final fermentation titer.

General considerations for production of the derivatives include the ability to do selective dehydrogenation (reduction) in the presence of other functionalities, specifically the conversion of the acid moieties to alcohols in the presence of amines. This will require the development of new heterogeneous catalyst systems that afford high selectivity, fast reaction rates and moderate operating conditions.

9.6.3 Building Block Considerations

Productivity: Productivity improvements are required to reduce the capital and operating costs of the fermentation. A minimum productivity of 2.5 g/Lhr needs to be achieved in order to be economically competitive on a commodity scale.

Nutrient Requirements: It is essential for commercial fermentations to be run using minimal nutrients. Expensive nutrient components such as yeast extract and biotin must be eliminated. The nutrient requirements should be limited to the use of corn steep liquor or equivalent if possible.

Final Titer: Final titer is also important when considering overall process costs. This is not a showstopper but a high final titer will reduce overall separation and concentrating costs.

pH Considerations: Ideally, the fermentation could be run at low pH, most preferably without requiring any neutralization. The cost of neutralization is not necessarily cost prohibitive, but the conversion of the salt to the free acid does add significant costs. If derivatives are going to be competitive from a cost perspective then low pH fermentation will be essential.

9.6.4 *Derivative Considerations*

Family 1: Glutaminol, 5-amino-1-butanol, 1,5-pentanediol, norvoline

The hydrogenation/reduction chemistry for the conversion of glutamic acid to the analogs of BDO, THF and BGL needs substantial development. The chemistry for doing selective reductions is not well understood, particularly in aqueous media. The technical challenge will be to develop new catalyst systems to obtain high yields of desired products and limit side reactions involving the other functionalities such as the amine. An additional consideration is the development of catalysts that would not be affected by impurities in the fermentation. This is a significant challenge and should be considered early in the development cycle since commercial fermentations for glutamic acid are available.

9.6.5 *Overall Outlook*

There is a significant market opportunity for the development of biobased products from the C5 building block glutamic acid. The major challenges are primarily associated with reducing the overall cost of the fermentation. In order to be competitive with products derived from petrochemicals, the fermentation cost needs to be at or below \$0.25/pound. This is a significant technical challenge but has the potential to be obtained in the midterm since there is substantial experience working with the organism for glutamic acid.

9.7 Itaconic acid

9.7.1 Pathways to Building Blocks from Sugars

Table 28 – Pathways to Building Block from Sugars [Itaconic Acid]

Type of pathway	Technical Barriers	Direct Uses of Building Block
Chemical- Multistep. Not likely a viable option	Costly synthesis. Reducing number of steps	Copolymers with styrene-butadiene polymers
Biotransformation- Fermentation product Aerobic fungal fermentation	Improving microbial biocatalyst to 1) reduce other acid coproducts, 2) increase yields and productivities Better control of operating environment Lower costs of recovery process Scale-up and system integration	Copolymer in styrene butadiene polymers (provides dye receptivity for fibers); Nitrile latex

9.7.2 Primary Transformation Pathway(s) to Derivatives

Table 29 – Family 1: Reductions [Primary Transformation Pathway(s) to Derivatives - Itaconic Acid]

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
Methyl butanediol, butyrolactone, tetrahydrofuran family	Selective reduction of specific functionalities Reduction at mild conditions – atmospheric pressure, low T Tolerance to inhibitory elements or components of biomass based feedstocks– robust catalysts	May confer new useful properties for the BDO, GBL, and THF family of polymers
Pyrrolidinones	Same as above including amination issues	Solvents and polymer precursor

Table 30 – Family 2: Direct Polymerization [Primary Transformation Pathway(s) to Derivatives - Itaconic Acid]

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
Polyitaconic	Manage rates Selective esterifications to control branching Control of molecular weight & properties	New polymer opportunity

Building Block: Itaconic Acid

Primary Derivatives:

Family 1: Reduction

Family 2: Direct Polymerization

9.7.3 Building Block Considerations

Itaconic acid is a C5 dicarboxylic acid, also known as methyl succinic acid and has the potential to be a key building block for deriving both commodity and specialty chemicals. The basic chemistry of itaconic acid is similar to that of the petrochemicals derived maleic acid/anhydride. The chemistry of itaconic acid to the primary families of derivatives is shown in Figure 12.

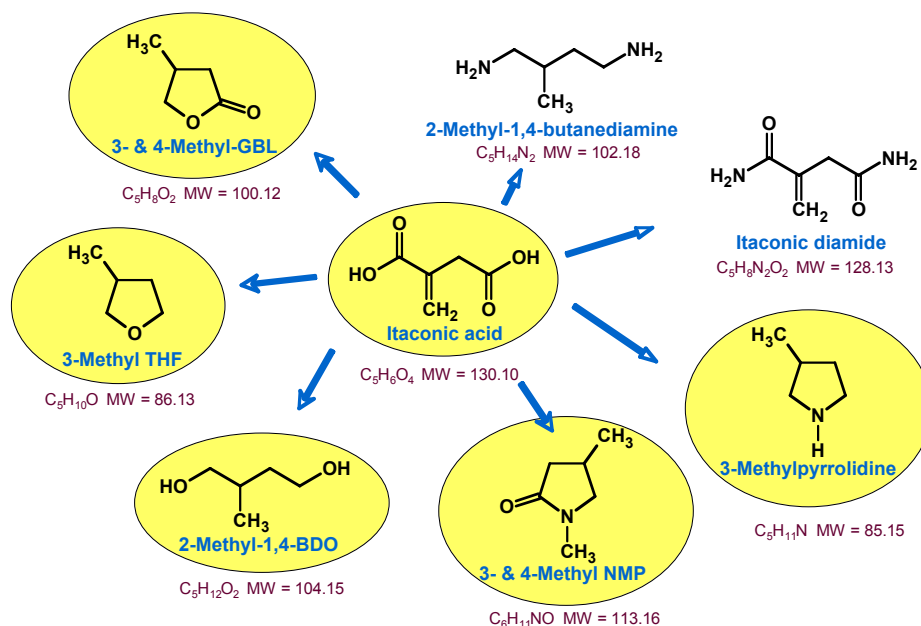


Figure 12- Itaconic Acid Chemistry to Derivatives

Itaconic acid is currently produced via fungal fermentation and is used primarily as a specialty monomer. The major applications include the use as a copolymer with acrylic acid and in styrene-butadiene systems. The major technical hurdles for the development of itaconic acid as a building block for commodity chemicals include the development of very low cost fermentation routes. The primary elements of improved fermentation include increasing the fermentation rate, improving the final titer and potentially increasing the yield from sugar. There could also be some cost advantages associated with an organism that could utilize both C5 and C6 sugars.

Productivity: Productivity improvements are required to reduce the capital and operating costs of the fermentation. A minimum productivity of 2.5 g/Lhr needs to be achieved in order to economically competitive.

Nutrient Requirements: It is essential for commercial fermentations to be run using minimal nutrients. Expensive nutrient components such as yeast extract and biotin must be eliminated. The nutrient requirements should be limited to the use of corn steep liquor or equivalent if at all possible.

Final Titer: Final titer is also important when considering overall process costs. This is not a showstopper but a high final titer will reduce overall separation and concentrating costs.

pH Considerations: Ideally, fermentation could be run at low pH, most preferably without requiring any neutralization. The cost of neutralization is not necessarily cost prohibitive, the conversion of the salt to the free acid does add significant costs. If derivatives such as 2-methyl-1,4-BDO, 3-methyl-THF, 3- or 4-GBL, and 2-methyl-1,4-butanediamine are going to be competitive from a cost perspective then low pH fermentation will be essential.

9.7.4 *Derivative Considerations*

Family 1: 2-Methyl-1,4-BDO, 3-Methyl THF, 3-&4-GBL, 2-Methyl-1,4-butanediamine

The hydrogenation/reduction chemistry for the conversion of itaconic acid to 2-methyl-1,4-BDO, 3-methyl THF, 3- and 4-GBL, and 2-methyl-1,4-butanediamine is not well known but should be similar to that of the hydrogenation chemistry associated with the production of BDO, THF and GBL from maleic anhydride. One technical consideration here is the development of catalysts that would not be affected by impurities in the fermentation.

Family 1: Pyrrolidones

The conversion of itaconic acid to pyrrolidones could follow the same chemistry associated with the conversion of GBL to pyrrolidones. One advantage with the fermentation-derived succinic acid is that the conversion of diammonium itaconate to the pyrrolidones directly could offer a significant cost advantage. This would eliminate the need for low pH fermentation for the direct production of itaconic acid.

Family 2: Direct Polymerization

The potential for direct polymerization would need to be investigated. The similarities of polyitaconic properties to known polymers would need to be ascertained in order to evaluate its use as a polymer. Its novel functionalities make it a potential raw material for polyene type polymers (polyvinylitaconic with side chains).

9.7.5 *Overall Outlook*

There is a significant market opportunity for the development of biobased products from the C5 building block itaconic acid. The major challenges are primarily associated with reducing the overall cost of the fermentation. In order to be competitive with petrochemical-derived products the fermentation cost needs to be at or below \$0.25/pound. This is a significant technical challenge and should be undertaken with a long-term perspective.

9.8 Levulinic acid

9.8.1 Pathways to Building Block From Sugars

Table 31 - Pathways to Building Block From Sugars [Levulinic Acid]

Type of pathway	Technical Barriers	Direct Uses of Building Block
Chemical – One step, acid catalyzed dehydration and decomposition of cellulose and sugars	Selective dehydration without side reactions (high value need for biomass) New heterogeneous catalysts (i.e., solid acid catalysts) to replace liquid catalysts and to improve existing catalyst based systems	
Biotransformation- None		

9.8.2 Primary Transformation Pathway(s) to Derivatives

Table 32 – Family 1: Reductions [Primary Transformation Pathways(s) to Derivatives - Levulinic Acid]

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
Methyl tetrahydrofuran, (γ -butyrolactone (some of this technology is already patented)	Selective reduction of diacids to produce alcohols, lactones and furans Reduction at mild conditions –pressure, low T Tolerance to inhibitory elements or components of biomass based feedstocks (sugar streams) – robust catalysts	Fuel oxygenates, solvents

Table 33 - Family 2: Oxidations [Primary Transformation Pathways(s) to Derivatives - Levulinic Acid]

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
Acetyl acrylates Acetic-acrylic succinic acids	Selective oxidation of alcohols (ROH) to acids (RCOOH) Avoiding exotic oxidants in favor of air, oxygen, dilute hydrogen peroxide. Lowering concentrations of oxidants required (safety issue) Tolerance to inhibitory elements of biomass based feedstocks (sugar streams)	Copolymerization with other monomers for property enhancement

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
	Facile and selective conversion of aldehydes to acids and alcohols to aldehydes Enzymatic oxidation requires cofactors	

Table 34 – Family 3: Condensation [Primary Transformation Pathways(s) to Derivatives - Levulinic Acid]

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
Diphenolic acid	Manage rates Control of molecular weight & properties Control of polymerization processes	Replacement for bisphenol A used in polycarbonate synthesis

Building block: Levulinic acid (LA)

- **Family 1** – Reductions
- **Family 2** – Oxidations
- **Family 3** – Condensations

9.8.3 Building Block Considerations

LA is a product formed by treatment of 6-carbon sugar carbohydrates from starch or lignocellulosics with acid. Five carbon sugars derived from hemicelluloses (viz. xylose, arabinose) can also be converted to LA by addition of a reduction step subsequent to acid treatment. Thus, LA could serve as a valuable building block available from almost all sugars manufactured in the biorefinery. LA is one of the more recognized building blocks available from carbohydrates (and has attracted interest from a number of large chemical industry firms), and for that reason, has frequently been suggested as a starting material for a wide number of compounds. Figure 13 summarizes some possible derivatives of LA.

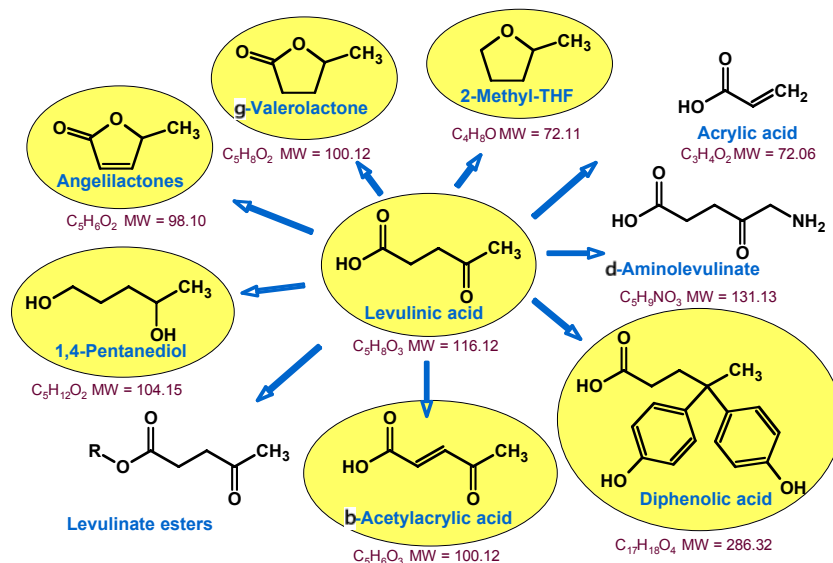


Figure 13- Derivatives of Levulinic Aid

9.8.4 Derivative considerations

The family of compounds available from LA is quite broad, and addresses a number of large volume chemical markets. Conversion of LA to methyltetrahydrofuran and various levulinate esters addresses fuel markets as gasoline and biodiesel additives, respectively. Delta-aminolevulinic acid is a herbicide, and targets a market of 200 – 300 million lb/yr at a projected cost of \$2.00-3.00/lb. An intermediate in the production of δ -aminolevulinic acid is β -acetylacrylic acid. This material could be used in the production of new acrylate polymers, addressing a market of 2.3 billion lb/yr with values of about \$1.30/lb. Diphenolic acid is of particular interest because it can serve as a replacement for bisphenol A in the production of polycarbonates. The polycarbonate resin market is almost 4 billion lb/yr, with product values of about \$2.40/lb. New technology also suggests that LA could be used for production of acrylic acid, whose applicability is described elsewhere in this report. Production of LA-derived lactones offers the opportunity to enter a large solvent market, as these materials could be converted into analogs of N-methylpyrrolidinone. Complete reduction of LA leads to 1,4-pentanediol, which could be used for production of new polyesters.

The technical barriers for this building block include improvement of the process for LA production itself. However, this conversion has been studied, and the LA yield is around 70%. Determination of the value of increasing this yield through more selective dehydration processes would be useful, and an effort in developing new catalysts to facilitate this conversion could be of use. Greater impact could be realized by focusing effort on conversion of LA to various derivatives. Of particular interest are processes to facilitate selective oxidation of LA to succinic and acrylic acid, focusing on the use of simple oxidants such as oxygen or peroxide. Selective reduction of LA to the corresponding lactones and methyltetrahydrofuran will improve access to large volume fuel markets. Finally, development

of diphenolic acid as a bisphenol A replacement, and investigation of the properties of the resulting polymers will provide a body of information to potential industrial partners interested in using the technology for the development of new products.

9.8.5 Overall Outlook

LA offers one of the larger families of potential industrial derivatives among the compounds included in the top 10. Its low cost and ready availability from both 5-carbon and 6-carbon sugars suggests that it could be a building block of central importance within the biorefinery. R&D will be necessary, especially to determine which of the many potential derivatives offer the best opportunities for partnering with industry.

9.9 3-Hydroxybutyrolactone

9.9.1 Pathways to Building Block From Sugars

Table 35 – Pathways to Building Block from Sugars [Pathways to Building Block From Sugars – 3-Hydroxybutyrolactone]

Type of pathway	Technical Barriers	Direct Uses of Building Block
Chemical- One step Oxidative degradation of starch (hydrogen peroxide)	Avoiding exotic oxidants in favor of air, oxygen, dilute hydrogen peroxide. Lowering concentrations of oxidants required (safety issue) Tolerance to inhibitory elements of biomass based feedstocks Ability to use wider range of lignocellulosics for synthesis	Intermediate for high value pharma compounds
Biotransformation- Not likely		

9.9.2 Primary Transformation Pathway(s) to Derivatives

Table 36 – Family 1: Reductions [Primary Transformation Pathway(s) to Derivatives – 3-Hydroxybutyrolactone]

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
Furans. Analogs of pyrrolidones	Selective reduction of specific functionalities, i.e. hydroxyls in presence of C=C. Reduction at mild conditions – atmospheric pressure, low T Selective reduction of aldehydes (RCHO) in presence of alcohols (ROH) Managing acid salts Tolerance to inhibitory elements or components of biomass based feedstocks (sugar streams) – robust catalysts	Solvents

Table 37 – Family 2: Direct Polymerization [Primary Transformation Pathway(s) to Derivatives – 3-Hydroxybutyrolactone]

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
Amino analogs to tetrahydrofuran	Manage rates Selective esterifications to control branching Control of molecular weight & properties	Amino analogs to lycra fibers

Building Block: 3-Hydroxybutyrolactone

- **Family 1** – Reduction
- **Family 2** – Direct Polymerization

9.9.3 Building Block Considerations

3-Hydroxybutyrolactone (3-HBL) is a cyclic C4 compound produced via chemical transformations. Production via fermentation routes is not likely. Chemical synthesis of this compound involves multiple steps and thus, is considered “messy” and/or difficult. One possible route to the building block starts with malic acid (2-hydroxysuccinic). Malic acid can be cyclized to form hydroxysuccinic anhydride, which, via reduction, gives the hydroxybutyrolactone. Malic acid is currently produced from fumaric or maleic acid, both derived from maleic anhydride, which in turn is produced from vapor-phase oxidation of hydrocarbons (particularly butane). The conversion from fumaric to malic is done using fermentation. Thus, if biotechnology advances could lead to malic acid production from sugars, a more cost-effective pathway can be envisioned. Perhaps a direct bioconversion from sugar all the way to 3-HBL may be possible as well. Thus, the major technical hurdles for the development of 3-HBL as a building block include the development of lower cost fermentation routes.

9.9.4 Derivative Considerations

Ring opening reactions can produce hydroxy analogs of succinic acid as potential derivatives. Other potential derivatives include gamma-butenyl-lactone (via dehydration) and acrylate-lactone (via esterification). The chemistry of 3-HBL to the primary families of derivatives is shown in Figure 14.

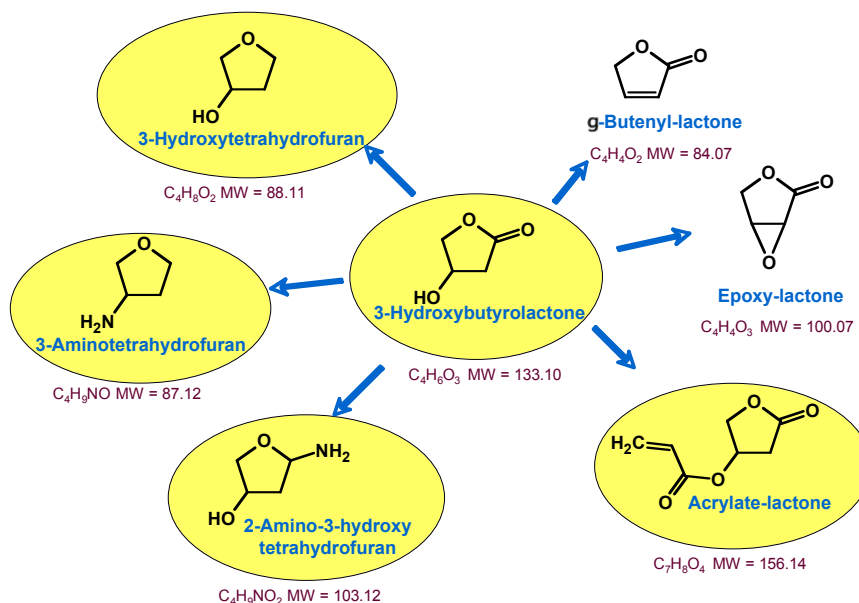


Figure 14 - 3-HBL Chemistry to Derivatives

Dehydration of 3-HBL can yield gamma-butenyl-lactone, and esterification can yield acrylate-lactone. Potential uses for such derivatives may include new polymers. 3-HBL is currently used in pharmaceutical production.

9.9.5 Overall Outlook

The research and market opportunities for 3-hydroxybutyrolactone come from its potential to create new derivative compounds. Since it is produced as a specialty chemical for fairly high value uses, little attention has been paid to producing it as a commodity chemical intermediate and all the attendant issues with such development. This would be a challenging product to develop, but one that has some exciting opportunities. Engineering analyses to better define the metrics needed to achieve cost efficient production of this molecule would be required.

9.10 Glycerol

9.10.1 Pathways to Building Block

Table 38 - Pathways to Building Block [Glycerol]

Type of pathway	Technical Barriers	Direct Uses of Building Block
Chemical – Transesterification of oils	Few if any	It is consumed in products such as personal/oral care products, drugs / pharmaceuticals, foods/beverages, and polyether polyols (for polyurethane).
Biotransformation - Enzymatic transesterification	New, robust enzymes active in methanol/water solutions, at costs comparable to chemical esterification	Same

9.10.2 Primary Transformation Pathway(s) to Derivatives

Table 39 –Family 1: Oxidation [Primary Transformation Pathway(s) to Derivatives [Glycerol]

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
PLA Analogs, Glyceric Acid	Alcohols (ROH) to acids (RCOOH) Avoiding exotic oxidants in favor of air, oxygen, hydrogen peroxide Tolerance to inhibitory components of biomass processing streams Oxidation of aldehydes to acids and alcohols to aldehydes	PLA with better polymeric properties Polyester fibers with new properties

Table 40 – Family 2: Bond Breaking (Hydrogenolysis) [Primary Transformation Pathway(s) to Derivatives [Glycerol]

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
Propylene glycol	Specificity for C-C & C-O bonds Increasing rates Tolerance to catalyst poisons (biomass sugar streams)	Antifreeze, humectant, etc.
1,3-propanediol	Same as above	Sorona fiber

Family 3: Direct Polymerization

Table 41 - Family 3: Direct Polymerization [Primary Transformation Pathway(s) to Derivatives [Glycerol]

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
Branched polyesters and polyols	Manage rates Selective esterifications to control branching Control of molecular weight & properties	Unsaturated Polyurethane Resins for use in insulation

Building block: Glycerol

- **Family 1:** Oxidation
- **Family 2:** Bond breaking (hydrogenolysis)
- **Family 3:** Direct polymerization

9.10.3 Building Block Considerations

Glycerol holds the potential of being an extremely versatile building block within the biorefinery. Glycerol is currently a well-recognized item of commerce with an annual world production of 500 – 750x10³ tonnes. In 1945, the work of Leffingwell described over 1500 different uses for glycerol.² The United States is one of the world's largest suppliers and consumers of refined glycerol. Glycerol and various simple derivatives of glycerol (such as glycerol triacetate, glycerol stearate, and glycerol oleate) are currently produced by the chemical industry at companies such as Uniqema, Procter and Gamble, and Stepan.³ Glycerol is produced in two forms: natural glycerol, as a byproduct of the oleochemical and biodiesel industries, and synthetic glycerol, from propylene. Approximately 75% of the U.S. supply of glycerol is derived from natural sources, and the remaining 25% is produced synthetically. Nearly all crude glycerol is refined before its ultimate end use.

Although many uses have been developed for glycerol, most product markets are currently small and fragmented, reflecting glycerol's relatively high price of \$0.60 – 0.90/lb. However, development of a biodiesel market could have a huge impact on the availability and use of glycerol. Since glycerol is a key coproduct of biodiesel manufacture, increasing use of biodiesel will lead to much greater glycerol availability and lower cost. The lowest price that crude glycerol could fall to is \$0.05/lb, because at that value steam reforming to hydrogen, animal feed, and other values will create large markets for crude glycerol. Glycerol prices could fall to \$0.20/lb which is the industry average cost for refining glycerol today although crude, unrefined glycerol, glycerin, may be available for a lower cost³. If prices drop into the \$0.20 - \$0.50/lb range, glycerol can become a major building block for the biorefinery. Small increases in fatty acid consumption for fuels and products can increase world glycerol production significantly. If the United States displaced 2% of the on-road diesel with biodiesel by 2012, almost 800 million pounds of new glycerol supplies would be produced.⁴ Figure 15 describes how glycerol could be used as a building block for the production of a family of derivatives.

² Leffingwell, G.; Lesser, M. Merck Index, 11th edition, p. 705 (1945)

³ Glycerol refining costs for industry provided by Procter & Gamble, July 17, 2003

⁴ Tyson, K. Shaine, Oil Platform Analysis: A Scoping Study for Biorefineries, NREL Technical Report (2004), in press.

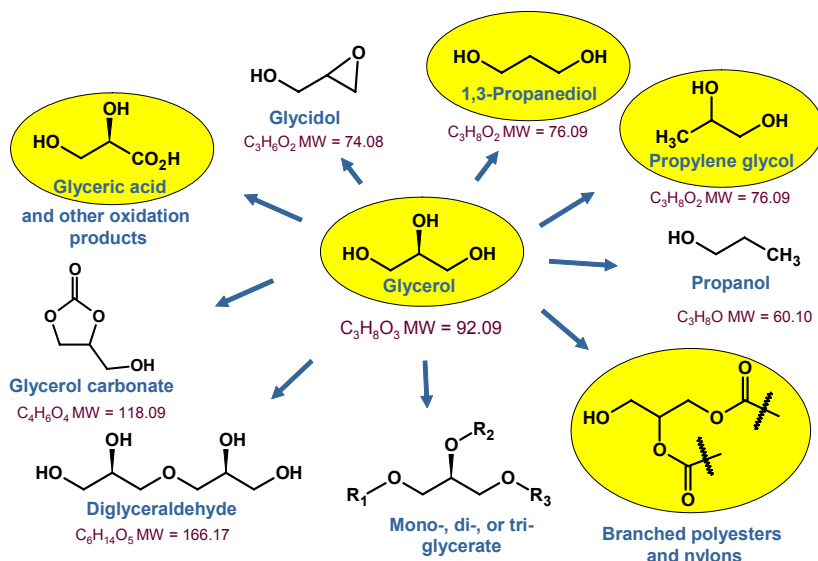


Figure 15 - Derivatives of Glycerol

9.10.4 Derivative Considerations

Historically, the cost of glycerol has meant that it was either used directly, or subjected to very simple structural modifications. Current derivatives include glycerol triacetate, glycerol esters (stearate, oleate), produced through chemical catalysis. At lower projected costs, there is a tremendous potential to develop a variety of new processes and product lines from glycerol, taking advantage of its unique structure and properties. As glycerol is a nontoxic, edible, biodegradable compound, it will provide important environmental benefits to the new platform products. Lower cost glycerol could open significant markets in polymers, ethers, and other compounds. From a technical standpoint, glycerol's multifunctional structure can be exploited by several different means, as shown by the potential glycerol product family in Figure 16. It is important to note that technology developed for glycerol would have broad crosscutting applications throughout the biorefinery. Since glycerol is structurally analogous to sugars, conversion processes developed for glycerol would also be applicable to inexpensive glucose, xylose, etc., greatly increasing the diversity of the biorefinery.

Selective oxidation of glycerol leads to a very broad family of derivatives that would serve as new chemical intermediates, or as components of new branched polyesters or nylons. These products would address very large chemical markets. Targeted polyesters have markets of 2-3 billion lb/yr, at values between \$1.00 – 3.50/lb, while nylons are a 9 billion lb/yr market with values between \$0.85 – 2.20/lb, depending on use. Technical barriers for production of these materials include the need to develop selective catalytic oxidation technology that can operate on a polyfunctional molecule such as glycerol. The processes will also need to use simple oxidants, such as oxygen or air, to carry out the required transformations.

New bond breaking (hydrogenolysis) technology will lead to the formation of a number of valuable intermediates. Propylene glycol (PG) and 1,3-propanediol (PDO) are promising potential derivatives that could be produced from glycerol by development of appropriate catalytic systems. PDO can be produced through aerobic fermentation, however, a direct

route from glucose to PDO (Dupont) is likely to be more cost effective. The conversion to PG would be via chemical catalysis. With the capacity to produce 1.5 billion lb/yr of PG in the US, PG offers a huge potential market for glycerol. A key barrier for this transformation is to make it cost competitive with the current petroleum route. Again, an important technical barrier is the need to develop selective catalysts to carry out these transformations, specifically, catalysts that can differentiate between C-C and C-O bonds.

9.10.5 Overall Outlook

Glycerol offers a very large number of opportunities for chemical production, and with the expected drop in cost, could become one of the more important building blocks for the biorefinery. Given its potential, a preliminary economic screening of the glycerol potential has been carried out as part of the evaluation for the oils platform. This information is shown in Table 42. This evaluation could help to screen some of the many possibilities for glycerol.

Table 42- Preliminary Economic Screening of the Glycerol Potential⁵

<i>Product</i>	<i>Current market size (lb)</i>	<i>Estimated raw material cost from glycerol (\$/lb)</i>	<i>Estimated raw material cost for commercial material (\$/lb)</i>	<i>Comparative commercial product cost (\$/lb)</i>	<i>CEH Source</i>	<i>Opportunity</i>
<i>New polyesters</i>						
<i>Thermoplastic engineering polyesters</i>	924 x 10 ⁶ (PBT); 152 x 10 ⁶ (PET)	0.29 – 0.34	0.379 (PET) 0.59 (PBT)	1.00 – 1.40 (2002)	695.4020	+++++
<i>Polyester film</i>	2209 x 10 ⁶	0.29 – 0.34	0.379 (PET)	1.05 – 3.50 (2001)	580.1170	+++++
<i>Polyester polyols</i>	1276 x 10 ⁶	0.29 – 0.34	0.88 (aliphatic) 0.34 (aromatic)	0.65 – 0.70 (foams) 0.35 – 0.55 (aromatic foams) 1.10 – 1.30 (nonfoam) [2001]	688.2000	+++
<i>Polyester fibers</i>	35864 x 10 ⁶	0.29 – 0.34	0.37 (PET)	0.63 (1999) 0.90 – 1.02 (1996) 0.55 – 1.50 (2001)	541.9000	++
<i>Unsaturated polyester resins</i>	3896 x 10 ⁶	Formulation R&D needed			580.1200	+++
<i>COPE elastomers</i>	152 x 10 ⁶ (2000)	1.20 – 1.62	1.35 – 1.75	3.15 – 3.62	525.8200	++++
<i>Polyurethane foams</i>	8661 x 10 ⁶	0.29 – 0.34		0.22 – 0.33	580.1600	+
<i>Polyurethane elastomers</i>	1276 x 10 ⁶	0.29 – 0.34	1.06	1.85 – 4.72	525.6600	++++
<i>Alkyd resins and coatings</i>						
<i>Resins</i>	1575 x 10 ⁶ (2000)	0.35	0.53	0.75 – 1.35	592.6000	++++
<i>Coatings</i>	4108 x 10 ⁶ (2000)	0.35	0.53	1.22 – 2.60	592.6000	+++++
<i>Glycerol oxidation products</i>	New market; projected initial uses in polymer markets above, and as chemical intermediates	0.30 – 0.45				+++++

⁵ ⁵ Leffingwell, G.; Lesser, M. Merck Index, 11th edition, p. 705 (1945).

⁵ Uniqema - <http://www.uniqema.com/tech/index.htm>

Stapan - <http://www.stapan.com/aboutstapan/about.asp>

Procter and Gamble - [http://www.pg.com/frameset fs.jhtml?frameURL=http%3A//www.pgchemicals.com](http://www.pg.com/frameset_fs.jhtml?frameURL=http%3A//www.pgchemicals.com)

Table 43- Preliminary Economic Screening of the Glycerol Potential (Continued)

<i>Product</i>	<i>Current market size (lb)</i>	<i>Estimated raw material cost from glycerol (\$/lb)</i>	<i>Estimated raw material cost for commercial material (\$/lb)</i>	<i>Comparative commercial product cost (\$/lb)</i>	<i>CEH Source</i>	<i>Opportunity</i>
<i>New nylons</i>	9028 x 10 ⁶	0.73 – 0.87	0.96 – 1.02	1.25 – 1.70 (carpet) 2.15 – 2.20 (textile) 0.85 – 0.12 (staple)	541.7000	++++
<i>PLA analogs</i>	308 x 10 ⁶ for PLA	0.36 – 0.51	New products	0.75 – 1.00 for PLA	670.5000	++++
<i>Surfactants</i>	10721 x 10 ⁶	Formulation R&D needed		Wide range of costs	583.8000	+++?
<i>Acrylic acid</i>	7596 x 10 ⁶	0.26 – 0.44	~0.12	0.87	606.4000	++
<i>Glycerol carbonate</i>	Unknown; propylene carbonate is about 21x10 ⁶ lb/yr	0.20 – 0.35	0.84 – 0.91 for dimethyl carbonate	0.50 – 0.60 for dimethyl carbonate from India		++++

Key: +++++ = good; ++++ = intermediate to good; +++ = intermediate; ++ = poor to intermediate; + = poor

9.11 Sorbitol (Alcohol Sugar of Glucose)

9.11.1 Pathways to Building Block

Table 44 –Pathways to Building Block [Sorbitol]

Type of pathway	Technical Barriers	Direct Uses of Building Block
Chemical – Hydrogenation of glucose	99.7 % yield, few if any technical barriers	
Biotransformation - None		

9.11.2 Primary Transformation Pathway(s) to Derivatives

Table 45 – Family 1: Dehydration [Primary Transformation Pathway(s) to Derivatives - Sorbitol]

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
Isosorbide, anhydrosugars	Selective dehydrations without side reactions Dehydration steps to anhydrides or lactones New heterogeneous catalyst systems (solid acid catalyst) to replace liquid catalyst systems	PET like polymers such as Polyethylene isosorbide terephthalates (bottles, in use for hot-fill)

Table 46 – Family 2: Bond Cleavage (hydrogenolysis) [Primary Transformation Pathway(s) to Derivatives - Sorbitol]

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
Propylene glycol, lactic acid	Specificity for C-C & C-O bonds Increasing rates Tolerance to catalyst poisons (biomass sugar streams)	Antifreeze, PLA

Table 47 – Family 3: Direct Polymerization [Primary Transformation Pathway(s) to Derivatives - Sorbitol]

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
Branched polysaccharides	Selective esterifications to control branching Control of molecular weight & properties	Water soluble polymers (water treatment, etc), new polymer applications

Building Block: Sorbitol

Primary Derivatives:

- **Family 1:** Dehydration
- **Family 2:** Bond cleavage (hydrogenolysis)
- **Family 3:** Direct Polymerization

Sorbitol is the hydrogenation product of glucose. The production of sorbitol is practiced commercially by several companies and has a current production volume on the order of 200 million pounds annually. All of the commercial processes are based on batch technology and the use of Raney nickel as the catalyst. The major reason for using batch technology is to ensure complete conversion of the glucose. This is an important requirement because the majority of the applications for sorbitol are in the food industry and there are very strict requirements for the amount of reducing sugars in the sorbitol product.

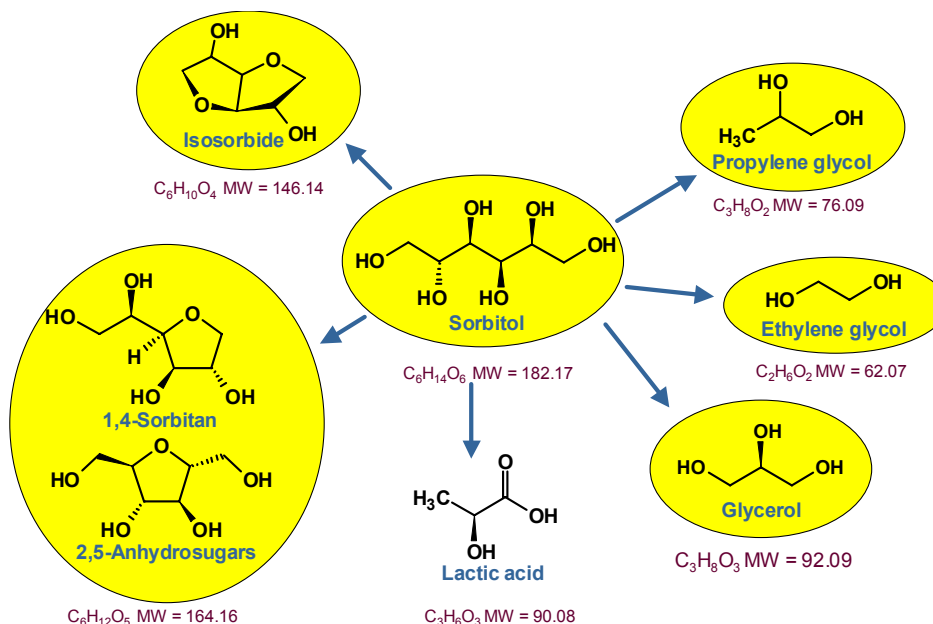


Figure 16 - Sorbitol Chemistry to Derivatives

The use of sorbitol as a building block for derivatives essentially requires no technical development. The only change would be the conversion of glucose to sorbitol in a

continuous process instead of a batch process. Engelhard has demonstrated that the continuous production of sorbitol from glucose can be done continuously using a ruthenium on carbon catalyst. The yields demonstrated were near 99 percent with very high weight hourly space velocity. This would make sorbitol a very inexpensive feedstock for producing derivatives.

Inexpensive sorbitol affords the potential for the production of isosorbide at low costs. Isosorbide has been demonstrated to be a very effective monomer for raising the glass transition temperature of polymers. The major applications are as a copolymer with PET for the use in bottle production. These applications include ridged bottles as well as the ability to use plastic to replace glass in hot fill applications.

9.11.3 *Building Block Considerations*

As was described above there are virtually no building block considerations for sorbitol with the exception of the actual demonstration of a continuous process. Because the hydrogenation of glucose to sorbitol is so straight forward, it is not anticipated that there would be any difficulties in the scale up.

9.11.4 *Derivative Considerations*

Family 1: Isosorbide

The challenge in producing isosorbide from sorbitol is the development of process conditions and dehydration catalysts that afford high yield of isosorbide. The best reported yields are around 76%. In order to increase the use of isosorbide, increased yields to 90% would be desirable. This not only reduces the production cost but also reduces the recovery and purification costs.

Family 2: Glycols

The conversion of sorbitol to glycols, namely propylene glycol is based on the hydrogenolysis of sorbitol. The major difficulty is producing propylene glycol in high yields. The best results in the literature show only about 35 pounds of PG from 100 pounds of sorbitol. In order to be commercially viable, this yield needs to be near 60 pounds per 100 pounds of sorbitol. New catalyst systems that afford high yield of PG are needed for commercial success.

Family 3: Direct polymerization

Copolymerization with other glycols in the unsaturated polyester resin market would be a major opportunity. Other direct polymerizations would yield polyesters that would need to be evaluated for properties and compared with other polyesters.

9.11.5 *Overall Outlook*

Sorbitol has the potential to be an outstanding building block for commodity chemicals. The conversion of sorbitol to isosorbide affords the opportunity for a high value monomer intermediate for several commercial applications. The conversion of sorbitol to glycols affords the opportunity to utilize a renewable resource for the production of a large-scale commodity chemical.

9.12 *Xylitol/arabinitol (Sugar alcohols from xylose and arabinose)*

9.12.1 *Pathways to Building Block From Sugars*

Table 48 – Pathways to Building Block From Sugars [Xylitol/arabinitol]

Type of pathway	Technical Barriers	Direct Uses of Building Block
Chemical- Hydrogenation of sugars or extraction from biomass pretreatment processes.	Very few if any. Commercial processes	Non-nutritive sweeteners, anhydrosugars, unsaturated polyester resins (UPRs)
Biotransformation- Result of pretreatment stream for lignocellulosic processing	Separation from other sugars	

9.12.2 *Primary Transformation Pathway(s) to Derivatives*

Table 49 – Family 1: Oxidations [Primary Transformation Pathway(s) to Derivatives – Xylitol/arabinitol]

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
Xylaric and Xylonic acids Arabonic acid and Arabinoic acid Xylaric and Xylonic acids Arabonic acid and Arabinoic acid	Selective oxidation of alcohols (ROH) to acids (RCOOH) Avoiding exotic oxidants in favor of air, oxygen, dilute hydrogen peroxide. Lowering concentrations of oxidants required (safety issue) Tolerance to inhibitory elements of biomass based feedstocks (sugar streams) Facile and selective conversion of aldehydes to acids and alcohols to aldehydes Enzymatic oxidation requires cofactors	New uses

Table 50 - Family 2: Bond Cleavage (hydrogenolysis) [Primary Transformation Pathway(s) to Derivatives – Xylitol/arabinitol]

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
Polyols (propylene and ethylene glycols) Lactic acid	Specificity among C-O and C-C bonds Increasing rates Tolerance to catalyst poisons (biomass processing streams and purity)	Antifreeze, UPRs,

Table 51 - Family 2: Direct Polymerization [Primary Transformation Pathway(s) to Derivatives – Xylitol/arabinitol]

Derivative or Derivative Family	Technical barriers	Potential use of derivatives
Xylitol, xylaric, xlyonic polyesters and nylons Same sequence for arabinitol	Manage rates Selective esterifications to control branching Control of molecular weight & properties	New polymer opportunities

Building Block: Xylitol/Arabinitol

Primary Derivatives:

Family 1: Oxidation

Family 2: Bond cleavage (Hydrogenolysis)

Family 3: Direct polymerization

Xylitol and arabinitol are hydrogenation products from the corresponding sugars xylose and arabinose. There is limited commercial production of xylitol and no commercial production of arabinitol. Xylitol is used as a non-nutritive sweetener. The technology required to convert the five carbon sugars xylose and arabinose to xylitol and arabinitol can be modeled based on the conversion of glucose to sorbitol. The hydrogenation of the five carbon sugars to the sugar alcohols occurs with one of many active hydrogenation catalysts such as nickel, ruthenium and rhodium. There is no major technical barrier associated with the production of the five-carbon sugar alcohols xylitol and arabinitol.

The production of xylitol for use as a building block for derivatives essentially requires no technical development. Based on the related chemistry of sorbitol production from glucose it is expected that the conversion of xylose to xylitol would be expected in the 99% yield range. Engelhard has demonstrated that the continuous production of sorbitol from glucose can be done continuously using a ruthenium on carbon catalyst. The yields demonstrated were near 99 percent with very high weight hourly space velocity and should be consistent for xylitol production. If the xylose feedstock is inexpensive then the production of xylitol could be done for very low cost.

9.12.3 Building Block Considerations

As was described above there are virtually no building block considerations for xylitol with the exception of the actual demonstration of a continuous process. Because the hydrogenation of glucose to sorbitol is so straight forward, it is not anticipated that there would be any difficulties for the production of xylitol from xylose and the only need would be in the demonstration of scale up.

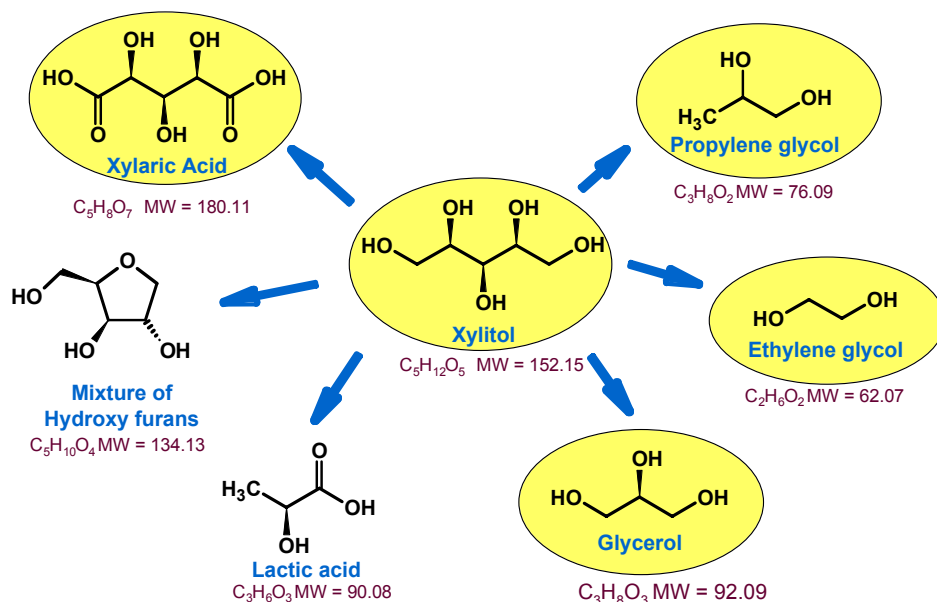


Figure 17 - Chemistry to Derivatives of Xylitol and Arabinitol

9.12.4 Derivative Considerations

Family 1: Xylaric Acid

Xylaric acid requires the selective oxidation of xylitol. The oxidation chemistry has been demonstrated for the oxidation of sorbitol to glucaric acid but with low efficiency. Yields for this reaction are reported at about 60%, which is not a showstopper but the separation of glucaric acid from the product mixture is problematic. New catalysts need to be developed that afford high yields of xylaric acid. In addition, it will be essential to develop the process in a manner that affords the use of oxygen as the oxidizing agent instead of oxidants like nitric acid, or peracetic acid. It is anticipated that the overall yields will need to be 90% or greater for an economic process to be considered for commercial deployment.

Family 2: Glycols

The conversion of xylitol to glycols, namely propylene glycol and ethylene glycol is based on hydrogenolysis. It has been demonstrated that reasonably good yields (80%) of ethylene glycol and propylene glycol can be achieved from xylitol. Improvement in the yields to greater than 90% would further improve the economics. The current challenge is finding a

low cost xylose sugar stream. One interesting concept may be to develop a process from mixed sugars that include xylose, arabinose and glucose for the conversion to propylene glycol as the primary product and ethylene glycol as the secondary product. This scenario could potentially afford a low cost route to glycols. The economics of this need to be demonstrated by some additional detailed modeling.

Family 3: Direct polymerization

Copolymerization with other glycols for the unsaturated polyester resin market would be one of the major opportunities. Other direct polymerizations would yield polyesters that would need to be evaluated for properties and compared with other polyesters. However, polymers with five carbon subunits possibly provide properties that are different from even numbered polymers.

9.12.5 Overall Outlook

C5 sugars such as xylose and arabinose have the potential to be an outstanding building block for commodity chemicals. One challenge will be getting a relatively clean feed stream of these sugars. The conversion of these sugars to the sugar alcohol and subsequent conversion to glycols has been demonstrated and could be a first generation process for the production of propylene glycol and ethylene glycol.

10 Catalog of Potential Chemicals and Materials from Biomass

As described above, the report started with a group of over 300 potential compounds that could be made from biomass. The final version of this report will contain a URL for a searchable database of these compounds.

Bibliography

References Used to Develop Catalog for Potential Biobased Products

Aggressive Growth in the Use of Bio-derived Energy and Products in the United States by 2010. Final Report and Data Volume, Arthur D. Little, October, 2001.

Alternative Feedstocks Program Technical and Economic Assessment: Thermal/Chemical and Bioprocessing Components. Prepared for DOE's Office of Industrial Technologies, J. Bozell and R. Landucci, Editors, NREL 1993.

Biobased Products. Prepared for the Fifth Biomass Conference of the Americas by M. Paster and T. Carole. September, 2001 (personal communication).

Chemical Economics Handbook, Stanford Research Institute International, Palo Alto, CA, 2002

Industrial Bioprocessing Alert. An electronic newsletter published regularly by Technical Insights Alert. Copyright 2003, Frost & Sullivan, San Antonio, TX 78229. ISSN 1056-7194.

Industrial Utilization of Renewable Resources: An Introduction. H.J. Szmant. Technomic Publishing Company, Inc., Lancaster, Pennsylvania, 1986.

Process Economics Program Yearbook, 2002, Volume 3E. SRI Consulting, Menlo Park, California (contains process economics for 934 routes to produce 534 chemicals in the United States, Germany and Japan).

Production of Organic Chemicals via Bioconversion: A Review of the Potential. Prepared for the US DOE's Idaho Operations Office. S. A. Leeper, T.E. Ward, and G. F Andrews. EGG-2645, July 1991.

The Bioproducts Industry: Today and Tomorrow. Prepared for DOE's Office of the Biomass Program by Energetics, Incorporated. M. Paster, J. Pellegrino and T. Carole. July, 2003 <http://www.bioproducts-bioenergy.gov/default.asp>.

References for Assigning Chemical and Biochemical Pathways

Ashford's Dictionary of Industrial Chemicals. 2nd Edition. Wavelength Publications, Ltd., London, England. 2001 (8,360 entries with 7,970 structured diagrams).

Chemical Economics Handbook, Stanford Research Institute International, Palo Alto, CA, 2002

Industrial Organic Chemistry. Klaus Weissermel and Hans-Jurgen Arpe, 3rd Edition, VCH Publishers, NY (Wiley company) 1997.

The Merck Index: An Encyclopedia of Chemicals and Drugs. Merck & Co., Inc. Rahway, NJ.

Kirk-Othmer Encyclopedia of Chemical Technology (online). Copyright © 2003 by John Wiley & Sons, Inc.

Ullmann's Encyclopedia of Industrial Chemistry (online). Copyright® 2003 by Wiley-VCH Verlag GmbH & Co. KGaA.

New Biocatalysts: Essential Tools for a Sustainable 21st Century Chemical Industry. W. Scouten and G. Petersen. Copies available from W. Scouten at wscouten@utsa.edu and G. Petersen at gene.Petersen@nrel.gov

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