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«Οι σύγχρονες τεχνικές βιο-ανάλυσης στην υγεία, τη γεωργία, το περιβάλλον και τη διατροφή»







Advances in ethanol production Claudia C Geddes, Ismael U Nieves and Lonnie O Ingram

Barriers to the commercialization of lignocellulosic ethanol include the development of more robust biocatalysts, reduction of cellulase costs, and high capital cost associated with a complex process. Improvements have been made in all areas during the past two years. Oxidoreductases, transporters, and regulators have been identified that can increase the tolerance of biocatalysts to inhibitors formed during pretreatment. Biocatalysts are being developed that grow under conditions that are optimal for cellulase activity and others have been engineered to produce glycoside hydrolases. Ethanol yields resulting from most current process configurations are similar, approximately 0.21 g ethanol/g dry cellulosic feedstock. Potentially, this can be increased to at least 0.27 g ethanol/g biomass (83 gal/ton) using simpler processes.

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Introduction

Lignocellulosic biomass (LCB) is an abundant, renewable source of carbohydrates for microbial conversion to chemicals and fuels. However, lignocellulose (cellulose and hemicellulose) is designed by nature to resist depolymerization. Processes that produce fermentable sugars from LCB tend to be complex and capital intensive. Chemical pretreatment is essential to increase cellulase access [1–3]. Pretreatments that hydrolyze hemicellulose into sugar syrups also form side products that retard fermentation. A solid-liquid separation and partial sugar purification are typically included to mitigate toxins, followed by the separate fermentation of C-5 (hemicellulose) and C-6 (cellulose) sugars.

Approximately 200 million dry tons of LCB are produced in the U.S. each year that could be used to produce 16 billion gallons of ethanol (Oak Ridge National Laboratory; URL: http://www.ornl.gov/~webworks/cppr/y2001/ rpt/123021.pdf). Starch-based ethanol (U.S.) and sugarcane-based ethanol (Brazil) are now mature industries but both compete with food uses. Companies such as Abengoa, BP-Verenium, Coskata, Dupont-Danisco and Poet are attempting to commercialize cellulosic ethanol in the next decade (Gigaom; URL: http://gigaom.com/ cleantech/12-companies-racing-to-build-cellulosic-ethanolplants-in-the-us/). Pilot and demonstration plants will serve as platforms to identify bottlenecks and potential barriers to full commercialization.

This review highlights advances in the fermentative production of ethanol from lignocellulose during the past two years. Improvements are noted in the areas of pretreatment, biocatalysts, saccharification and liquefaction, and process simplification.

Advances in pretreatment

Pretreatments using dilute sulfuric acid require reactors made of exotic metals. Although all mineral acids have been explored to some extent, recent studies have proposed the use of phosphoric acid $[4^{\bullet\bullet}, 5]$. As a weaker acid, phosphoric acid pretreatment produces lower levels of toxic side products than sulfuric acid pretreatment and can be used with a stainless steel reactor [4^{••}]. Autohydrolysis produces the lowest levels of side products [2]. With autohydrolysis, hemicellulose components are solubilized as oligosaccharides that require further hydrolysis with enzymes or acid. Ethanologenic Escherichia coli strains have been adapted to phosphoric acid hydrolysates and can now ferment hemicellulose and cellulose derived sugars together in a single vessel, termed simultaneous saccharification and co-fermentation (SScF). The use of phosphoric acid could eliminate the need for separation of hydrolysates from pretreated fiber, detoxification, and reactors of exotic metals resulting in a simpler process quite analogous to that for corn ethanol (Figure 1). Using this process, ethanol yields of up to 0.27 g/g bagasse (dry weight) have been obtained (83 gal/ton) [6^{••}].

The SPORL process (sulfite pretreatment to overcome recalcitrance of lignocellulose) and SO₂ impregnation use sulfur compounds to disrupt the LCB structure [7^{••},8]. SPORL is better suited for biomass with high lignin content and SO₂ impregnation for agricultural residues [8]. SPORL pretreatment was shown to increase sugar yields (from 57% to 88%) and reduce inhibitors by up to 65% compared to dilute sulfuric acid pretreatment of softwood [7^{••},9].

Ammonia-based AFEX pretreatment (ammonia fiber expansion) is very effective at increasing fiber digestion while producing lower levels of inhibitors than sulfuric



Figure 1

Lignocellulose to ethanol process configurations. The cellulose could be hydrolyzed alone before fermentation (separate hydrolysis and fermentation, SHF) or with the hemicellulose (separate hydrolysis and co-fermentation, SHCF) followed by fermentation of the resulting slurry. Cellulose hydrolysis could also occur simultaneously with fermentation in the presence (SSCF) or absence (SSF) of hemicellulose. In the liquefaction followed by simultaneous saccharification and co-fermentation (L + SSCF) process, there is a cellulose prehydrolysis step in the presence of hemicellulose hydrolysate followed by fermentation but the cellulases continue to hydrolyze the cellulose during fermentation. The consolidated bioprocessing process involves a biocatalyst that is capable of producing all the hydrolytic enzymes required for cellulose hydrolysis and is also capable of fermenting all the resulting sugars in the presence of hydrolysate inhibitors. Adapted from [2].

acid pretreatments [10,11]. Hemicellulose oligomers produced by the AFEX process require further hydrolysis into monomers [2]. After AFEX pretreatment and enzymatic digestion, over 80% of the carbohydrate in the fiber was recovered as soluble sugar [11]. Subsequent studies have identified compounds (4-hydroxybenzaldehyde, lactate, and acetate) formed during AFEX pretreatment that inhibited fermentations with *E. coli* KO11 [12^{••}]. AFEX remains a highly effective pretreatment for grasses.

Advances in biocatalyst

The need for more robust biocatalysts is one of the weakest links in the LCB to ethanol process. These biocatalysts need to be resistant to inhibitors formed during lignocellulose pretreatments, co-utilize a variety of sugars at high yields, secrete cellulase enzymes, and remain active under conditions that are near optimal for cellulase function (pH 5, 50 °C). Much of the complexity in lignocellulosic ethanol processes stems from the need for toxin mitigation (solid liquid separation after pretreatment; sugar cleanup) before fermentation. Additional

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complexity comes from the requirement for external sources of cellulase enzymes. Developing biocatalysts that ferment under conditions that are near optimal for fungal cellulase activity can reduce the requirement for external enzymes. Engineering the fermenting biocatalyst to produce some or all of the cellulase enzymes provides a complementary route to further reduce enzyme cost. The development of a fermentation-based lignocellulose to ethanol industry depends on research advances to minimize these biological impediments.

Developing tolerance to hydrolysate inhibitors

Furans from sugar dehydration, acetate, and soluble products from lignin are the primary inhibitors in hemicellulose hydrolysates from dilute acid pretreatment [13]. Of these, furans appear to be particularly important and have been the focus of many recent papers. Biocatalysts have been developed with increased resistance to furfural [14^{••},15,16^{••},17,18,19^{••}], 5-hydroxymethylfurfural [15,19^{••},20], acetate [21[•]], and to unfractionated dilute acid hydrolysate [6^{••},15,19^{••},22^{••},23^{••}].

Furans can be reduced to less toxic furan alcohols by most organisms using native enzymes. Expression arrays have noted many changes among oxidoreductases in response to furans with yeast [14,15,24,25^{••}], Zymomonas mobilis [19^{••},21[•]], and ethanologenic *E. coli* [16^{••},17,18,20]. In some cases, regulators have been identified [18,19^{••}]. Several of the furfural reductase enzymes in *E. coli* have sufficiently low K_m values for NADPH that growth is inhibited until furan metabolism has been completed. Silencing of these low K_m enzymes (less than 20% of total furfural reductase activity) was beneficial for furfural tolerance in *E. coli* [17]. Other higher K_m enzymes (NADPH), NADH-dependent enzymes [15], and transhydrogenase enzymes have been shown to confer partial resistance to furfural [16^{••},18,20].

Reduced sulfur compounds have long been used to improve the fermentation of dilute sulfuric acid hydrolysates of wood although the mode of action remains unknown [23^{••},26^{••},27]. Saccharomyces cerevisiae fermentation of acid hydrolysates (sugarcane bagasse and spruce wood) was improved by the direct addition of reduced sulfur compounds to slurries containing complex media, termed 'in situ detoxification' [26**]. The use of sodium metabisulfite and sodium hydrosulfite was also shown to improve the fermentation of slurries containing phosphoric acid pretreated sugarcane bagasse (L + SScF) using E. coli in mineral salt medium [23**]. Surprisingly, addition of metabisulfite did not decrease the toxicity of furfural or acetate when each was tested alone. It is possible that metabisulfite neutralizes the toxicity of soluble products from lignin.

Improving hexose and pentose sugar coutilization

The co-utilization of hexose and pentose (xylose and arabinose) sugars remains a challenge for biocatalysts, especially in the presence of hydrolysate inhibitors. Derivatives of *S. cerevisiae* have been previously engineered to ferment xylose and these continue to be improved by additional genetic changes in xylose metabolism $[28^{\bullet\bullet}, 29, 30]$. Although ethanologenic *E. coli* have the native ability to metabolize all sugars from LCB, xylose utilization lags behind glucose and was also improved by further genetic changes [31]. Inhibitors present in acid hydrolysates retarded xylose metabolism during *E. coli* fermentation even with this genetic change. The lag in xylose metabolism was substantially relieved by injection of small amounts of air during fermentation, termed microaeration [6^{••}, 32].

Advances in saccharification Reducing the cost of enzyme production

The cost of cellulase enzymes remains a major concern for the commercialization of LCB ethanol processes. Cost estimation software (*e.g.* Aspen Plus and Aspen Icarus Process Evaluator) has been used to compare the minimum ethanol selling prices of processes involving the purchase of commercial cellulase and on-site cellulase production [33]. International enzyme companies such as Novozymes and Genencor have formed partnerships with Poet LLC and Dupont Danisco Cellulosic Ethanol LLC, respectively, to commercialize lignocellulosic ethanol (The New York Times; URL: http://www.nytimes.com/cwire/2010/02/16/ 16climatewire-economics-improve-for-first-commercialcellu-93478.html?scp=1&sq=cellulosic%20ethanol%20 plants&st=cse). Both have reported that enzymes will cost approximately \$0.50/gallon of ethanol. This represents an 80% price reduction during the last two years.

Efforts continue to reduce the cellulase requirement. Novel cellulases have been isolated from a variety of organisms using improved screening methods [34]. Current research has focused on developing improved cellulase enzyme cocktails, development of biocatalysts with fermentations that match the optimal conditions for cellulase activity, and novel cellulases [34,35^{••}]. *Trichoderma reesei* is currently the primary industrial organism used for the production of cellulase enzymes. Sequencing of the *T. reesei* genome will facilitate further improvements in enzyme production [36].

Improving cellulase performance

Compounds have been identified that increase cellulase effectiveness and enzyme usage [37,38^{••},39]. These include surfactants (Tween 80, cetylpyridinium chloride, and cetyl trimethylammonium bromide) and divalent metals (calcium and magnesium). Up to 35% improvement in saccharification was reported. All are proposed to act by reducing the nonproductive binding of cellulases to lignin. *Bacillus coagulans* is a thermotolerant biocatalyst capable of growth at temperatures and pH (55 °C, pH 5.0) that are optimal for fungal cellulases. Using this organism, the cellulase was reduced to 5 FPU/g cellulose during lactate production [35^{••},40]. Similar benefits would be expected for ethanol production after further metabolic engineering, and for other biocatalysts that can function under these conditions.

Toward consolidated bioprocessing

Consolidated bioprocessing without the need for externally supplied enzymes remains a goal for many scientists $[41^{\circ},42^{\circ},43^{\circ\circ},44,45]$. Expression of endoglucanase I and II genes from *T. resei* QM6a allowed the resulting strain to ferment phosphoric acid swollen cellulose (amorphous) when beta-glucosidase was supplied $[41^{\circ}]$. Tsai *et al.* and Wen *et al.* reported the development of recombinant *S. cerevisiae* strains capable of displaying functional minicellulosomes on their surface exhibiting enzyme synergy and producing 3.5 g/L and 1.8 g/L ethanol, respectively, using phosphoric acid swollen cellulose $[42^{\circ},43^{\circ\circ}]$. Previous studies have demonstrated up to 11 g/L ethanol production from phosphoric acid swollen cellulose using *Klebsiella oxytoca* strain SZ21 expressing endoglucanase



Figure 2

Comparison of ethanol production from lignocellulose and corn. (a) Simplified process using phosphoric acid hydrolysis of hemicellulose and enzymatic hydrolysis of cellulose. Enzymatic liquefaction was added before co-fermentation of hexose and pentose sugars in a single vessel (L + SScF) and (b) enzymatic liquefaction of hydrated corn before simultaneous saccharification and fermentation (L + SSF). Adapted from [21[•]].

genes from *Erwinia chrysanthemi* (*celY*, *celZ*) [44]. Synergies between purified cellulases and xylanases from the thermophilic bacterium *Thermobifida fusca* displayed on 'designer cellulosomes' were found to possess higher activity on wheat straw than the corresponding free enzymes [45].

Process simplification

Reducing process complexity remains a major challenge for the commercialization of LCB to ethanol. Current research is focused on eliminating the need for detoxification of hydrolysates, developing robust biocatalysts capable of fermenting pentose and hexose sugars simultaneously, reducing water usage, increasing ethanol yield and titer, and decreasing cellulase usage. Considerable progress has been made during the past two years by developing robust biocatalysts capable of fermenting pentose and hexose sugars simultaneously. Further progress is needed to increase ethanol titers and to decrease water and cellulase usage. Collaborative research projects have focused on comparing pretreatment options for specific biomass types (e.g. corn stover or poplar wood [1,46^{••}]). Various process configurations are shown in Figure 2. These decrease in complexity from separate hydrolysis and fermentation (SHF) to consolidated bioprocessing (CBP). The SHF process involves separation of the cellulose-rich solid from the hemicellulose hydrolysate and separate fermentation trains. L + SScF and SScF processes combine C-6 and C-5 sugar fermentations in a single vessel [6^{••},22^{••},23^{••}]. The consolidation of bioprocessing steps is hindered by the fibrous nature of suspensions at loadings of 10-20% solids [47,48]. Models have been described relating viscosity, solubilized sugars, time, and enzyme loadings for slurries of sugarcane bagasse $[4^{\bullet\bullet}, 48, 49]$. On the basis of these studies, a partial saccharification step using a CSTR (one to six hours residence) was proposed $[4^{\bullet\bullet}]$. This liquefaction step can produce slurries containing 10–15% solids (solids plus solubles) that can be readily pumped and mixed.

SScF of lignocellulosic biomass

Pretreatment processes typically require solid-liquid separations and neutralization of hydrolysate toxins before fermentation. With the development of hydrolysate resistant biocatalysts such as E. coli MM160 [22**,23**] and S. cerevisiae 424A [43^{••},44,45,46^{••},47], comparable yields could be obtained with less process complexity. The development of robust biocatalysts allowed the fiber and liquid from pretreatment to be fermented without separation [6^{••},22^{••},23^{••}]. The resulting process is analogous to the mature corn dry milling ethanol process (Figure 1) that combines all components in a single vessel after an initial liquefaction step (L + SScF process).Ethanol yields for LCB processes have continued to improve during the past two years (Table 1). Despite differences in process complexity, similar ethanol yields were obtained by most researchers, approximately 0.21 g/g (63 gal/ton). Higher yields are obtained when purified cellulose was used (*e.g.* paper sludge [55^{••}]) or starch combined with lignocellulose (e.g. corn silage and whole corn plant [53]). The use of SPORL pretreatment is making similar progress toward process simplifications (e.g. L + SScF) although part of the hydrolysate was removed before fermentation [56]. AFEX treated corn stover supplemented with corn steep liquor was fermented after an initial 96 h prehydrolysis (cellulases and hemicellulases added) to produce 40 g/L ethanol

Comparison of ethanol yields from SScF processes.

Feedstock	Pretreatment	Biocatalyst	Ethanol		Reference
			Titer (g/L)	Yield (g/g untreated feedstock)	
Rice straw	Dilute acid	M. indicus	11	0.11	[60]
Spruce	SO ₂ impregnation	S. cerevisiae TMB3400	45	Not calculated ^c	[61]
Forage sorghum	Aqueous ammonia	S. cerevisiae	-	0.13	[62**]
Hybrid poplar	Aqueous ammonia ^a	E. coli KO11	16	0.24 calculated ^d	[63]
Rice straw	Aqueous ammonia ^a	S. cerevisiae D5A	12	0.12	[64]
Corn stover	AFEX	S. cerevisiae 424A(LNH-ST)	40	0.20	[51**]
Wheat straw	Steam explosion ^a	K. marxianus CECT 10875	36	0.18 calculated ^d	[65**]
Rice straw	AFEX	S. cerevisiae 424A(LNH-ST)	37	0.21 calculated ^d	[54]
Rice straw	AFEX	P. stipitis FPL-061	30	0.17 calculated ^d	[54]
Rice straw	AFEX	P. stipitis DX-26	28	0.16 calculated ^d	[54]
Switchgrass	Hydrothermolysis	S. cerevisiae D5A	22	0.17 calculated ^d	[66]
Switchgrass	Hydrothermolysis	K. marxianus IMB	19	0.15 calculated ^d	[66]
Barley straw	Steam explosion	K. marxianus CECT 10875	22	0.17 calculated ^d	[67]
Sugarcane bagasse	Dilute phosphoric	E. coli MM160 (KO11 derivative)	29	0.21	[22**]
Sugarcane bagasse	Dilute phosphoric	E. coli MM160 (KO11 derivative)	20	0.20	[23**]
Sugarcane bagasse	Dilute phosphoric	E. coli MM170 (KO11 derivative)	27	0.27	[6**]
Spruce	SO ₂ impregnation	S. cerevisiae (bakers's yeast)	18	0.18 calculated ^d	[68**]
Switchgrass	AFEX	S. cerevisiae 424A(LNH-ST)	36	0.19	[50]
Distillers grains	Liquid hot water	S. cerevisiae D5A	14	0.09 calculated ^d	[69]
Distillers grains	AFEX	S. cerevisiae D5A	14	0.09 calculated ^d	[69]
Corn stover	AFEX	S. cerevisiae 424A(LNH-ST)	40	0.22 calculated ^d	[70]
Corn stover	AFEX	E. coli KO11	31	0.17 calculated ^d	[70]
Corn stover	AFEX	Z. mobilis AX101	32	0.18 calculated ^d	[70]
Forage sorghum	AFEX	S. cerevisiae 424A(LNH-ST)	31	0.17 calculated ^d	[52]
Sweet sorghum bagasse	AFEX	S. cerevisiae 424A(LNH-ST)	42	0.15 calculated ^d	[52]
Forage sorghum	AFEX	S. cerevisiae 424A(LNH-ST)	31	0.18 calculated ^d	[52]
Sweet sorghum bagasse	AFEX	S. cerevisiae 424A(LNH-ST)	29	0.18 calculated ^d	[52]
Rice straw	Dilute acid	C. tropicalis ATCC 13803	20	0.20	[71]
Corn silage	AFEX	S. cerevisiae 424A(LNH-ST)	28	0.31 calculated ^d	[53]
Whole corn plant	AFEX	S. cerevisiae 424A(LNH-ST)	30	0.32 calculated ^d	[53]
Lodgepole pine	SPORL ^b	S. cerevisiae Y5	21	0.21	[7]
Paper sludge	No additional treatments	S. cerevisiae RWB222	45	0.26 calculated ^d	[55**]
Paper sludge	No additional treatments	Z. mobilis 8b	46	0.27 calculated ^d	[55**]
Lodgepole pine	SPORL ^{b, e}	S. cerevisiae D5A	-	0.22	[56]
Corn stover	AFEX	C. phytofermentans	2.8	0.17	[72]
Corn stover	AFEX	S. cerevisiae 424A(LNH-ST)	3.9	0.24	[72]

^a Solids were washed after pretreatment.

^b Solid-liquid separation after pretreatment.

^c Unable to calculate with reported data.

^d Results presented were used to calculate yields on an original biomass basis.

^e Detoxification of hydrolysate before fermentation.

(0.22 g ethanol/g corn stover [51^{••}]) and dilute acid pretreated sugarcane bagasse was fermented to high ethanol yields (0.27 g ethanol/g bagasse) when air was added to the headspace during a L + SScF process [6^{••}].

Dual uses of process residues, chemicals, and water

Beneficial products must be derived from all materials entering LCB to ethanol processes. Vinasse, stillage from sugarcane ethanol processes, has been used for many years as a fertilizer for biomass crops [57]. Pretreatment processes with phosphoric acid offer a similar opportunity by producing an ammonium phosphate fertilizer that includes magnesium sulfate and trace metals [4^{••},22^{••},49]. A phosphoric acid LCB-ethanol process can be viewed as a temporary stop for water and fertilizer *en route* to farms for new crop growth, sharing the cost of these materials. Lignin-rich residues can be used as boiler fuel or converted to higher value products [58,59]. Lignin could also be formed into inert blocks as an effective means for carbon sequestration.

Conclusions

The challenge of producing 36 billion gallons of ethanol by the year 2022 is being met with an expansion of research in the biofuel arena. Major improvements have been made as researchers learn more about the genetic basis of resistance to inhibitors in acid hydrolysates and pentose utilization. Pretreatment processes have been optimized to minimize inhibitor formation and to improve enzymatic hydrolysis of cellulose. The cost of cellulase enzymes remains a concern. Approaches have been proposed to minimize external enzyme usage by producing enzymes in the biocatalyst and by providing conditions that increase the effectiveness of cellulases.

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