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ENZYMATIC AND STRUCTURAL FACTORS LIMITING
HYDROLYSIS OF CELLULOSE BY
TRICHODERMA HARZIANUM E58 CELLULASES

by
Charlene M. Hogan

Supervised by
Dr. D.J. Kushner (University of Ottawa)
and
Dr. J.N. Saddler (Forintek Canada Corp.)

A thesis submitted to the Faculty of Graduate Studies in
partial fulfilment of the requirements for the degree of
Master of Science

Department of Biology
University of Ottawa
Ottawa, Ontario

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ABSTRACT

*Trichoderma harzianum* E58 extracellular cellulases can be applied to a cellulose-to-glucose process. Optimal conditions for long-term hydrolysis were found to be pH 4.8, 45°C. Enzyme adsorption onto the commercial cellulose, Solka Floc BW300 was examined. The enzymes that adsorb to the substrate in the initial stage of hydrolysis had a strong role in governing the overall reaction. The enzyme loading at which the substrate was saturated with the adsorbed cellulases corresponded to the optimal enzyme loading for hydrolysis.

Enzyme-related and structural-related factors responsible for the decline in the hydrolysis rate were studied. Exoglucanase and β-glucosidase were resistant to thermal inactivation at 45°C over a 4 day period, while endoglucanase lost 20% of its activity. Site blockage of adsorptive or catalytic sites on the substrate did not appear to limit hydrolysis. End-product inhibition of β-glucosidase and endo- and exocellulase by glucose and cellobiose, respectively, is a major problem in hydrolysis. Inhibition can be alleviated by supplemental β-glucosidase or use of a combined hydrolysis and fermentation process. *T. harzianum* E58 produced sufficiently high levels of β-glucosidase to cope with glucose inhibition, at low substrate concentrations.

The relationship between initial crystallinity of the substrate and its hydrolysis appeared to be bimodal. No clear
pattern between accessible surface area of the substrate and its hydrolysis emerged. Changes occurring to the crystallinity and the accessible surface area during hydrolysis were followed. The change in crystallinity was negligible, while the accessible surface area increased drastically in the initial phase of hydrolysis, then slowly declined. The extent to which the substrate becomes resistant to enzymatic attack was examined. A 70% drop in its accessibility during 4 days hydrolysis was observed, indicating that the substrate became more recalcitrant during long-term hydrolysis.
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INTRODUCTION

I. HISTORY

Research on optimizing cellulose hydrolysis, ironically, started out in an attempt to prevent microbial decomposition from occurring. During the 1940's, American troops stationed in the South Pacific were dismayed at the rapid deterioration of their cotton goods, so the Army requested help in solving the problem. A long range basic research laboratory was set up at the U.S. Army Natick Development Center to investigate the nature of the rotting, the causal organisms, their mode or action, and a method of control. The laboratory collected deteriorated items, and isolated thousands of microorganisms. The cultures were grown on cotton duck, and the strips tested for loss in tensile strength.

Some interesting facts began to emerge. Most of the cellulolytic fungi grew well on cellulose and were able to degrade it. Filtrates from these cultures contained enzymes that released sugars from amorphous cellulose, but only rarely produced sugars from crystalline cellulose. It was deduced that several enzymes are required to hydrolyze crystalline cellulose, but not all cellulolytic micro-organisms manufacture, or secrete these enzymes.

The strong interest in cellulose degradation waned in the post-war years, but due to the concerted efforts of a small group of scientists, led by Mandels and Reese, research on cellulolytic micro-organisms continued, and much of the ground-work was laid for later researchers.
In the 1970's interest in cellulose degradation was renewed
due to concern over the energy shortage, environmental protection
and waste utilization. Bioconversion of lignocellulosic residues
to their component sugars and then to chemicals was thought to be
the solution to many of the problems, and research into this area
expanded. The prime target for most of this research is the
hydrolysis of cellulose to glucose, which could then be streamed
into a multitude of other products.

Although enzymatic hydrolysis is technically promising, a
better understanding of the mechanisms and limitations involved
in hydrolysis would help to develop and optimize the process.

The major problem in studying enzymatic hydrolysis of
cellulose is the complexity of the system. Wood, the usual
substrate, is insoluble and heterogenous - composed of
cellulose, hemicellulose, lignin - the nature and concentration
depending on the source. Cowling and Kirk (1976) compared the
composition of a number of sources of cellulose (i.e. hardwoods,
softwoods and grasses) with that of cotton, which contains about
90% cellulose. There was a wide variation in the amounts of
cellulose (10-55%), hemicellulose (25-50%) and lignin (10-35%).
"Cellulase" itself is a complex of enzymes acting
synergistically. In addition, both the structural features of
the substrate and the mode of attack of the cellulase complex are
constantly changing during the course of the reaction. Since the
reaction is long-term, inactivation of the enzyme must also be
considered. Given the immense complexity of this system, it is
not surprising that the mechanisms of the reaction are still not
elucidated.
Figure 1.
Antiparallel arrangement of the cellulose chains.

Alberts et al., 1983
(Courtesy of Garland Publishing Inc.)
II. CELLULOSE STRUCTURE

Cellulose is a linear polymer consisting of anhydroglucose units held together by $\beta-1,4$ linkages. This linkage rotates each glucose unit 180 degrees about its main axis in relation to its neighbor (Figure 1). This results in intramolecular hydrogen bonding between adjacent glucose molecules which helps to maintain rigidity of the glucose chain. Glucose chains are arranged in parallel, and are held together by intermolecular hydrogen bonding. Individually, the hydrogen bond is not strong, but this extensive network of inter and intra-hydrogen bonding contributes to the strength of the cellulose molecule.

Many models have been proposed to explain the arrangement of the glucose chain within the cellulose molecule. There are two basic types of models, the extended chain or the folding chain (Figure 2). The most widely accepted model is the Ellefson fibrillized, or folding chain model which proposes that the cellulose molecule folds over on itself.

In the cellulose molecules there are crystalline and amorphous regions. The term, "crystalline regions" relate to the areas of tight hydrogen-bonding, where the chains are arranged in a highly ordered manner (Figure 2). Areas where looser binding occurs are termed "amorphous" or "paracrystalline" regions.

Bundles of the cellulose molecules pack together to form the microfibril. The microfibers are cross-linked with lignin, hemicellulose and protein to form the cell wall of higher plants (Figure 3).
Figure 2.

Chain models of the elementary fibril:
a) fringe-micellar model; b) folding chain model.

Chang and Tsao, 1981
Figure 3.
Organization of cellulose molecules in the cell wall of a higher plant.

Alberts et al., 1983
(Courtesy of Garland Publishing Inc.)
III. STRUCTURAL FEATURES RESTRICTING ENZYMATIC HYDROLYSIS

Native lignocellulosics are virtually inaccessible to enzymatic attack without some form of pretreatment. It is now widely accepted that the crystallinity of a lignocellulosic material and its association with lignin are the major factors restricting enzymatic hydrolysis. These and other structural features that limit its susceptibility to enzymatic attack have been outlined by Cowling (1975): 1) degree of crystallinity of the cellulose, 2) moisture content of the fibres, 3) unit cell dimensions of the crystallites, 4) degree of polymerization, 5) conformation and steric rigidity of the anhydroglucose units, 6) nature, concentration, and distribution of substituent groups and 7) nature of the associated substances.

Currently, Klesov (1986) includes surface area, while Stone et al. (1969), and Grethlein (1985) list pore size of the substrate as major factors limiting hydrolysis. Both these factors relate the size and diffusibility of the enzyme molecule to the size and surface properties of the cellulosic material.

In order to reduce the complexity of the substrate, and to reduce the number of variables involved in examining hydrolysis, this thesis will focus primarily on purified cellulose substrates, with the lignin and hemicellulose fractions extracted.

A. Surface Area

The voids in wood and agricultural crops can be divided into two types: 1) gross capillaries, such as the cell lumina, pit
apertures and pit-membrane pores (these range from 20 to 10,000 nm in diameter); and 2) the cell-wall capillaries, such as the spaces between microfibrils and the cellulose molecules in the amorphous regions. The cell wall capillaries are closed when the material is dry, but can swell up to 20 nm when fully saturated.

The total surface area of the gross capillaries is roughly 2.0 m²g⁻¹, while the total surface area within the cell-wall capillaries is up to 300 m²g⁻¹. Much of the confusion regarding the relationship between surface area and hydrolysis is due to the methodologies employed to determine surface area. Many surface area measurements are made on the basis of the adsorption of much smaller molecules, such as nitrogen gas, which overestimate the area accessible to the enzyme.

The accessible surface area of the cellulose fibers is defined by the size, shape and surface properties of the capillaries within the fiber in relation to the size, shape, and diffusibility of the cellulase enzyme. Some surface area measurements use an enzyme exclusion approach: using an enzyme probe (peroxidase) comparable in size to the cellulase enzyme to measure available surface area. A more direct approach is to measure the amount of cellulase enzyme able to adsorb to the substrate.

Klesov and Sinitsyn (1981) measured the accessible surface area of a wide range of substrates using an enzyme-exclusion technique. They found the initial rate of hydrolysis to be proportional to the surface area of the substrate available to a 4.0 nm spherical enzyme.

On the other hand, Fan et al. (1980) found that the specific
surface area of refined cellulose, as measured by the nitrogen adsorption technique, had only a small effect on the initial rate of hydrolysis.

B. Crystallinity (CRI)

The fraction of crystalline material in the total cellulose is expressed in terms of its crystallinity index. Crystallinity is commonly determined by the method of Segal et al. (1959) with an x-ray diffractometer.

The relationship between crystallinity and enzymatic hydrolysis of refined substrates was first reported in the 1950's in work by Walseth (1952) and Baker et al. (1959). Walseth, using phosphoric acid swollen cellulose, found substrates with low crystallinity were more susceptible to hydrolysis by Aspergillus niger. Baker et al. found an inverse linear relationship between in vitro digestibility and the crystallinity index of cotton linters and wood cellulosics.

Later work using more precise experimental techniques continued to support this relationship. Klesov and Sinitsyn (1981), using an extensive range of substrates and pretreatments found a decrease in the crystallinity of a substrate corresponded with an increase in the initial rate of hydrolysis by Trichoderma reesei and Aspergillus niger cellulases. A strong correlation between increase in surface area and extent of hydrolysis was also demonstrated.

Fan et al. (1980) found an inverse relationship between the crystallinity index and glucose production using T. reesei.
cellulases. They worked out a mathematical relationship between the amount of glucose produced after 8 hours ($X_g$) and the initial crystallinity (CrI) and specific surface area (SSA) of the substrate, as seen in Equation 1:

$$X_g = 0.380(\text{SSA})^{0.195} (100 - \text{CrI})^{1.04}$$  
(Eqn 1)

Crystallinity is held to be a stronger determinant of hydrolysis than specific surface area, as demonstrated by its larger exponent.

Klesov and Sinitzyn (1981), Fan et al. (1980) and Bertran and Dale (1985) noted recrystallization of the pretreated substrate occurs when it is moistened, with the more amorphous substrates recrystallizing to a greater extent. Fan and co-workers obtained a stronger correlation between enzymatic hydrolysis and initial crystallinity index by first moistening, then solvent-drying the pretreated substrates prior to its crystallinity determination.

Information on the effect of enzymatic hydrolysis on the crystallinity of the substrate during long term hydrolysis is conflicting.

Norkrans (1950) speculated that the more amorphous regions of the substrate are selectively hydrolyzed. She hydrolyzed filter paper for 8 hours using Trichoderma nudum enzymes and noted an increase in the crystallinity of the residue.

Similarly, Ohmine et al. (1983) found that the CrI increased during the hydrolysis of Avicel by T. reesei cellulases, and concluded that there was preferential attack of
the amorphous region.

Lee and Fan (1983) found a slight increase in crystallinity during hydrolysis of hammer-milled Solka Floc (from 78.1 to 80.0) with _T. reesei_ cellulases. For more amorphous substances the change in CrI during hydrolysis was more significant. When the substrate was ballmilled to an initial crystallinity of 34.9 the CrI was reported to increase to 71.6 after hydrolysis. This was explained by the presence of a large amount of amorphous material that could be more easily hydrolyzed.

Chang _et al._ (1981) reported finding constant crystallinity during the enzymatic hydrolysis of both native and regenerated cellulose. This led them to conclude that both crystalline and amorphous regions are attacked concurrently.

Caulfield and Moore (1974) using ball-milled cellulose and the commercial cellulase "Onozuka" (from _T. reesei_) noted a relationship between the initial crystallinity and digestibility which they believe to be more a parallel correlation than a causal one. During grinding crystallite size and particle size are reduced, increasing the accessibility of both amorphous and crystalline cellulose to the enzyme molecules. The accessibility of the crystalline component is enhanced to a greater extent than that of the amorphous component.

Schurz (1986), using spruce sulfite pulp, found no significant change in the crystallinity, the degree of polymerization, the specific inner surface, the void fraction or the molecular weight distribution during 48 hours of hydrolysis by _T. reesei_ enzymes. On the basis of these results they proposed an "all or nothing" mechanism for degradation: a chain
is either fully degraded or not at all.

C. **Cellulase Dimensions**

The dimensions of the cellulases produced by various microorganisms have been estimated to range from 2.5 to 7.7 nm for spherical enzymes and 1.3 x 7.9 nm to 4.2 x 25.2 nm for ellipsoids (Cowling, 1975). **Trichodermal** enzymes have dimensions at the upper end of these ranges, and would be primarily limited to attacking the gross capillaries of the fibers. Only a small fraction of the cell-wall capillaries in water-swollen fibers are large enough for the enzymes to penetrate.

Schmuck et al. (1986) determined the molecular structural parameters of the major celllobiohydrolase purified from the culture filtrate of *T. reesei*. The distance distribution function measured by the small angle x-ray scattering technique indicated that the enzyme has a tad-pole like shape. The maximum length is 18 nm and the largest diameter 4.4 nm.

D. **Pore Size**

Pore size distribution is a very refined method of determining the surface area of a substrate. Stone and Scallan (1968) developed the solute exclusion technique, to measure the accessible volume, using a range of dextrans as inert probes.

The importance of pore size distribution of a substrate on its hydrolysis was first noted by Stone et al. (1969), who demonstrated a linear relationship between the initial reaction rate and the surface area within the cellulose accessible to a
molecule of 4.0 nm.

Grethlein (1985) found the initial glucose yield, after 2 hours hydrolysis with \textit{T. reesei}, was linear to the pore volume available to a spherical molecule of 5.1 nm.

Weimer and Weston (1985) found that rate of degradation of native cellulosics by \textit{Trichoderma} cellulases is strongly related to both the total pore volume and the initial crystallinity of the substrate. The rate-limiting component for the fungal cellulase system was found to have a size of 4.3 nm along one axis.

\textbf{E. Degree of Polymerization}

The degree of polymerization (number of anhydroglucose units in a cellulose molecule) varies with the source and pretreatment of the substrate, and ranges up to 10,000-14,000 for wood or cotton. Klesov and Sinitsyn (1981) and Schurz (1986) found no correlation between the degree of polymerization (DP) and hydrolysis.

\textbf{F. Pretreatment}

In nature cellulose is mostly crystalline, and is very resistant to hydrolytic agents, including enzymes. Removal of the lignin and hemicellulose fractions by physical and/or chemical means does not guarantee that the cellulose will become more susceptible to enzymatic attack.

The cellulose can be made more susceptible to the cellulases by reducing its crystallinity. This can be done by subjecting the cellulose to intensive milling or by treatment with cellulose
solvents.

Attrition methods reduce particle size and severely disrupt the crystalline order of the cellulose. Hammer-milling, ball-milling, compression-milling, vibratory-milling have been investigated by various groups.

There are a number of cellulose solvents, such as concentrated mineral acids, nonaqueous organic solvents such as dimethyl sulfoxide, and transition metal complexes e.g. chelating metal caustic swelling solvent (CMCS). These agents have a similar mode of action. They disrupt the hydrogen bonding, causing the cellulose molecule to pass into solution. Excess water is added to re-precipitate the amorphous cellulose.

Most of the attempts to correlate cellulose structure with degradability have involved pretreating cellulosic materials with physical or chemical agents that drastically alter several aspects of the cellulose fine structure. The data from these studies must be interpreted cautiously as several structural features are often altered simultaneously by a treatment.

There is evidence that some structural characteristics may be inter-dependent. Klesov claims a relationship between the degree of crystallinity of cellulose and its specific surface area as they were found to affect the hydrolysis rate to the same extent. The cellulose molecules are packed together more tightly in a crystalline substrate than in an amorphous substrate. Consequently, the specific surface area (i.e. surface area/weight substrate) is smaller for the crystalline substrate, and less surface area is available to the cellulase enzyme.
IV. CELLULASE COMPLEX

A. Characterization of the Enzymes

The three major cellulolytic components produced by *Trichoderma* spp. are: endo-β-1,4-glucanase, exo-β-1,4-glucanase and β-1,4-glucosidases. A multiplicity of each of these enzymes exists – it is not yet known if the individual forms are genetically determined, or if they arise from differential modification by proteolysis or glycosylation. The enzymes are best described by their mode of action, or the products of their reaction (Fig 4).

The endoglucanases (1,4-β-D-glucan glucanohydrolase (EC 3.2.1.4), Cx, CMC'ase), hydrolyze cellulose molecules in a random fashion, with a preference for larger molecules. Water soluble cello-oligosaccharides are the intermediate products of attack, but once formed these are rapidly hydrolyzed to cellobiose and glucose by the exoglucanases and β-glucosidase, respectively. Endoglucanases are typically characterized by their high activity on substituted celluloses e.g. carboxymethylcellulose (CMC), and low specific activity on Avicel (on the order of $10^3$ times lower). These enzymes exist in the largest number of multiple forms, possessing significantly different properties (e.g. in the randomness of their attack, degrees of synergism with the cellobiohydrolase, molecular weight, isoelectric point (Beldman et al., 1985, 1987; Bhikhabhai et al., 1984)).

Two classes of exoglucanase exist, exo-cellobiohydrolase (CBH) and exo-glucohydrolase. The main exoglucanase is exo-
Figure 4.

Schematic representation of the synergistic enzyme hydrolysis of cellulose.

Saddler, 1986
(Courtesy of Forintek Canada Corp.)
cellobiohydrolase (1,4-β-D-glucan cellobiohydrolase (EC 3.2.1.91)
which removes cellobiose units from the non-reducing end of the
cellulose molecule. It interacts synergistically with
endoglucanase to break down crystalline cellulose, against which
neither enzyme acting individually has significant activity.
This CBH is currently equated with the old C1 enzyme by many
researchers. The existence of an exo-glucohydrolase (1,4-β-D-
glucan glucohydrolase, EC 3.2.1.74), able to cleave off glucose
units from the non-reducing end of the cellulose molecule, has
been reported for some fungi, e.g. Tr. reesei (Marsden et al.,
1982) and Penicillium funiculosum (Wood and McCrae, 1982).

Cellobiohydrolases are generally characterized by their
ability to hydrolyze amorphous (phosphoric-acid swollen)
cellulose, with cellobiose as the main product, and by their
inability to attack substituted cellulose (e.g. CMC). Likewise,
glucohydrolases act on amorphous cellulose to release glucose,
but are not active against substituted cellulosics.

β-Glucosidase (EC 3.2.1.21) is responsible for formation of
the final product, glucose. While the terms cellobiase and
β-glucosidase are used interchangeably, β-glucosidase is able to
hydrolyze cellodextrins to glucose.

One of the main difficulties with the assay of cellulolytic
enzymes is the choice of a suitable substrate. As described
above, cellulase is a complex of enzymes. For the complete
hydrolysis of an insoluble cellulose, a synergistic action
between these components is required. Assay of the cellulase
components has usually been tailored to meet the requirements of
the researcher. This has resulted in a large array of
substrates, enzyme actions and nomenclatures (Table 1).

B. **Synergism**

Wood (1975) elegantly demonstrated that the cellulases in the culture filtrate of *T. koningii* act synergistically to hydrolyze fibrous cotton to glucose. The proteins in the culture filtrate were separated on DEAE Sephadex and identified as different enzymes: C₁ (exoglucanase), Cₓ (endoglucanase), and cellobiase. Individually the enzymes had little or no effect on cellulose, but when recombined at levels equal to those of the original filtrate they were able to hydrolyze the substrate to an extent equivalent to that obtained with the original culture filtrate.

When one considers synergism, a description of the substrate must also be included. The structural characteristics of a substrate determine the relative importance of synergism in the enzymes acting against it. Degradation of amorphous cellulose can be effected by either exo- or endoglucanase acting independently, whereas crystalline celluloses require the synergistic interaction of these enzymes (Coughlan, 1985). When incubated with amorphous cellulose the combined pure components show only a simple addition of the independent enzyme activities, rather than a synergism (Wood and McCrae, 1979). Obviously, the co-operative action between the two enzymes is low on amorphous substrates. Synergy operates at a maximum when the components are used in the ratios in which they occur in the fermentation filtrate (Wood, 1975).
Table 1. Cellulase assays.

<table>
<thead>
<tr>
<th>ENZYME</th>
<th>SUBSTRATES</th>
<th>PRODUCT(S) MEASURED</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cellulobiase/ Cellulocyclase</td>
<td>Cellobiose</td>
<td>Glucose</td>
</tr>
<tr>
<td>B-glucosidase</td>
<td>Cellobetaoides</td>
<td>Glucose</td>
</tr>
<tr>
<td>Aryl-B-glucosidase/</td>
<td>O-Nitrophenyl-B-glucoside</td>
<td>O-Nitrophenol or Glucose</td>
</tr>
<tr>
<td>B-glucosidase</td>
<td>Salicin</td>
<td>Saligenin or Glucose</td>
</tr>
<tr>
<td>Exo-B-1,4-glucanase</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cellulobiodylase (CBH, C₁)</td>
<td>Avicel</td>
<td>Cellobiodylase</td>
</tr>
<tr>
<td></td>
<td>Crystalline cellulose (cotton)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>p-Nitrophenyl-B-D-lactoside (pNPL)</td>
<td>p-Nitrophenol or Disaccharide</td>
</tr>
<tr>
<td></td>
<td>p-Nitrophenyl-B-D-cellulobioside (pNPC)</td>
<td>p-Nitrophenol or Disaccharide</td>
</tr>
<tr>
<td></td>
<td>4-Methylumbelliferyl-B-D-cellulobioside (MUC)</td>
<td>4-Methylumbelliferone or Disaccharide</td>
</tr>
<tr>
<td>Glucohydrodolase</td>
<td>Avicel</td>
<td>Glucose</td>
</tr>
<tr>
<td></td>
<td>Crystalline cellulose (cotton)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Cellobetaoides</td>
<td>Glucose</td>
</tr>
<tr>
<td>Endo-B-1,4-glucanase (EG, Cₓ)</td>
<td>Carboxymethylcellulose (CMC)</td>
<td>Reducing sugars or loss in viscosity</td>
</tr>
<tr>
<td>&quot;Complete cellulase&quot;</td>
<td>Amorphous cellulose</td>
<td>Reducing sugars or loss in viscosity</td>
</tr>
<tr>
<td>(FPase, Avicelase)</td>
<td>Filter paper (FP)</td>
<td>Reducing sugars or loss in weight</td>
</tr>
<tr>
<td></td>
<td>Avicel</td>
<td>Reducing sugars or loss in weight</td>
</tr>
<tr>
<td></td>
<td>Cellobetaoides</td>
<td>Reducing sugars</td>
</tr>
<tr>
<td></td>
<td>Cellobetaoides</td>
<td>Release of dye</td>
</tr>
<tr>
<td></td>
<td>Dyed cellulose</td>
<td>Breaking strength</td>
</tr>
<tr>
<td></td>
<td>Thread</td>
<td></td>
</tr>
</tbody>
</table>
The mechanism of synergistic interaction on crystalline substrates is not fully understood.

An endo-exo sequential mode of action has been proposed, with the endoglucanase initiating sites of attack for the exoglucanase. This is supported by the work of Streamer et al. (1975), who demonstrated synergistic hydrolysis is obtained when the substrate is first incubated with endoglucanase, prior to addition of the exoglucanase. Incubation with exoglucanase, followed by the endoglucanase, produced only low levels of sugars.

Ryu et al. (1984) studied the adsorption of the individual cellulolytic components of T. reesei filtrates on Avicel. They postulated that the endoglucanase and cellobiohydrolase components adsorb to different sites on the substrate, corresponding to the sites of hydrolysis. Addition of cellobiohydrolase to a substrate containing bound endoglucanase caused an acceleration in the action of the latter enzyme, and brought about its desorption. Similarly, endoglucanase on binding to cellulose increased the rate of action of the bound cellobiohydrolase and caused its desorption. This phenomenon, termed "competitive adsorption", is believed to explain the synergistic interaction of these components in cellulose hydrolysis. To further support this theory, they found the synergistic interaction was most evident when both components were present in the proportion found in the crude filtrate.

For efficient hydrolysis of amorphous cellulose the quantity of enzyme is the limiting factor. With crystalline substrates it appears that both the "quality" of the endoglucanase (discussed
below) and the "quantity" of the exoglucanase components are important.

Klesov et al. (1986) investigated the adsorption of endoglucanase, produced by different organisms, on a number of pure cellulose substrates varying in their crystallinity and surface area. The endoglucanases fell into two classes; those that adsorbed "strongly" and those that adsorbed "weakly", as determined by the partition coefficient \(K_p\) of the enzyme between the substrate surface and the bulk solution. They found that the decisive factor in effective hydrolysis of amorphous cellulose is the amount and activity of enzyme bound to it. Hydrolysis of the crystalline substrate by the complete system of exo- and endoglucanase depended not only on the extent but also on the "quality" of binding of the endoglucanase. Strongly adsorbed endoglucanases were found in the cellulase systems that are strongly active against crystalline cellulase, while the weakly adsorbed enzymes were found in systems that were relatively inactive against these substrates.

Wood (1982) studied this phenomenon from a different angle. He examined "cross-synergism" of cellobiohydrolase and endoglucanase components, from "strong" and "weak" cellulose degraders. The cellobiohydrolases were purified from fungi capable of degrading crystalline cellulose extensively (e.g. \(T.\ koningii\), \(P.\ funiculosum\), and \(F.\ solani\)), while the endoglucanases were purified from a number of fungal and bacterial sources. The source of endoglucanase had a decisive role in determining the extent of hydrolysis: endoglucanase
derived from "strong" degraders highly solubilized the cotton, while endoglucanase purified from "weak" cellulose degraders caused little or no solubilization. Apparently, there is a lack of synergy between the celllobiohydrolase and the "weak" endoglucanases. The major difference between the two types of endoglucanase is that celllobiohydrolase is not produced by the weak degraders.

In hydrolysis the endoglucanase nicks the cellulose chain generating a non-reducing end. The exoglucanase attacks this site, and then proceeds along the chain cleaving off celllobiose units. From a practical viewpoint, a single endoglucanase generates a large number of sites for a large number of exoglucanase components to act on. It would be expected that for an efficient hydrolysis the ratio of exoglucanase to endoglucanase would be high.

This postulation is supported by the work of Henrissat et al. (1985) who evaluated the action of celllobiohydrolases I and II (CBHII and CBHII) and endoglucanase I and II (EGI and EGII) purified from T. reesei in the degradation of pure cellulose substrates. No synergism between the exoglucanases and endoglucanases was noted on amorphous cellulase. Degradation of crystalline cellulose required the concerted action of both CBHII and EGI or EGII; with the reaction limited by the amount of the exo-enzyme present. Apparently, small amounts of endocellulase provide sufficient chain-ends to saturate the exo-enzymes present. The maximum amount of hydrolysis was observed with an enzyme mixture containing mainly exoglucanase with a small amount of endoglucanase.
A model for the mechanism of hydrolysis of crystalline cellulose was formulated by Coughlan (1985). The first step involves amorphogenesis, whereby the substrate is made more accessible to the cellulases. There is much debate as to the nature of this phenomenon. Reese et al. (1950) proposed that a "swelling factor", C₁, initiates attack on the substrate, facilitating subsequent attack by the cellulase Cₓ. Later researchers identified this C₁ component as cellobiohydrolase, and the idea of a swelling factor lost favour. More recently, Griffin (1984) isolated a "microfibril-generating factor" in the culture filtrate of T. reesei. The factor was characterized as non-enzymatic, due to its low protein content, and low molecular weight. The factor, requiring the presence of iron for its activity, displayed synergism with the cellulase to degrade filter paper.

C. End-Product Inhibition

Both endoglucanase and exoglucanase are strongly inhibited by cellobiose, the product of their concerted action, and moderately inhibited by glucose. There is much variation in the product inhibition models proposed for the exo- and endoglucanases. Exoglucanase purified from T. reesei was competitively inhibited by its products, cellobiose and glucose (Hsu et al., 1980), while a non-competitive product inhibition was reported for the T. viride enzyme system (Howell and Stuck, 1975).

The sensitivity of the cellulase enzymes to the inhibitors varies drastically with the organism under study. Inhibition
constants, $K_m$, ranging from 0.06 to 1.1 mM have been reported for cellobiose acting on exogluconase (Ladisch et al., 1983). Exogluconase, derived from *T. reesei* QM9414 was inhibited 10 times more strongly by cellobiose, than by glucose (Hsu et al., 1980).

β-Glucosidase is strongly inhibited by its product, glucose. Kinetic studies on β-glucosidase purified from *T. reesei* are more consistent than for the cellulases, with non-competitive inhibition reported (Ladisch et al., 1977; Gong et al., 1977). Glucose inhibits cellobiose less strongly than cellobiose inhibits the cellulases with $K_m$ values from 1.2 to 4.3 mM reported for the former (Ladisch et al., 1983).

Nature, in its economy, provided *T. reesei* with sufficient β-glucosidase to support its growth on cellulose, where sugars do not accumulate. *Trichoderma reesei* cellulase preparations have roughly 0.2-0.3 β-glucosidase units per filter paper unit (Ryu and Mandels, 1980). In a hydrolysis reactor, however, glucose levels are high, ranging from 5 to 10%. End-product inhibition is held accountable as one of the major factors limiting hydrolysis. As hydrolysis proceeds glucose accumulates and inhibits the β-glucosidase. In turn, cellobiose builds up and inhibits the cellulase. Therefore, an adequate level of β-glucosidase is critical for cellulose hydrolysis.

There are a number of ways to circumvent end-product inhibition. Supplemental β-glucosidase, derived from a high β-glucosidase producer, such as *Aspergillus phoenicis* can be added (Sternberg et al., 1977). Addition of the *Aspergillus* cellobiose to *Trichoderma* cellulase reduces cellobiose accumulation and
greatly enhances the glucose yield. The effect of supplemental \( \beta \)-glucosidase on enhancing hydrolysis reaches a maximum at a ratio of approximately 1 to 1.25 IU \( \beta \)-glucosidase per filter paper unit (Dekker et al., 1987). A combined hydrolysis and fermentation (CHF) or simultaneous saccharification and fermentation (SSF) has been investigated by several groups (Blotkamp et al., 1978; Saddler et al., 1982; Takagi et al., 1977). The cellulase system is incubated with a microbe that ferments the liberated glucose to the desired product, directly upon its release. The end result is reduction of glucose inhibition, so the enzymes are able to operate at a faster rate for a longer period of time.

D. Enzyme Inactivation

Enzymatic hydrolysis of cellulose is a long term process during which time the enzymes are subject to shearing, and thermal inactivation (Mukataka et al., 1983; Herr, 1980). As the reaction time for hydrolysis increases, the amount of active enzyme decreases, imposing a restraint on efficient hydrolysis. For a practical process the enzyme must remain stable for extended periods of time (at least 2 days, and much longer if recycling is to be considered). But hydrolysis operates at fairly high temperatures (45-50°C) with agitation, factors that are not conducive to longevity of the enzymes.

The effect of long-term incubation on enzyme stability at a wide range of temperatures has been investigated by numerous researchers. There is a wide range in the half-life of the
different components of the cellulase system. The results of these experiments must be interpreted cautiously as biocides, e.g. thimerosal, sodium azide, are sometimes used in these studies. These biocides inactivate the cellulase components to different degrees, depending on the concentration at which they are used (Reese and Mandels, 1980).

The enzyme complex derived from some *Trichoderma* strains are more sensitive to temperature than others. After 2 days incubation at 50°C, at moderate shaking conditions (120 rpm), the *Trichoderma* strains, QM9414, MCG77 and C30 lost 26, 44, and 61% of their filter paper activity, and 0, 0 and 11% of their endoglucanase activity (Mandels et al., 1981). There was little difference in the loss of activity between shaken or unshaken conditions for the above strains.

In general, β-glucosidase of *T. reesei* strains appears to be tolerant of high incubation temperatures under both shaken or unshaken conditions (Mandels et al., 1981).

Andren (1978) demonstrated that, for a long-term incubation the faster rate of hydrolysis obtained at high temperatures must be weighed against the effects of thermal inactivation. As the time of saccharification increased, the maximum extent of hydrolysis was found to occur at lower temperatures.

Mukataka et al. (1983) studied the effect of agitation on the overall hydrolysis. They found the rate of hydrolysis was enhanced with agitation up to about 200 rpm, then declined. The drop in the rate was reported due to greater shear at the higher agitation.

These experiments on stability were carried out at pH 4.5-
5.0, and at 45-50°C, but in the absence of substrate, and as such
do not duplicate the hydrolytic conditions. There is some
indication that the enzymes are more stable in the presence of
substrate, due to adsorption onto the substrate (e.g.
immobilization effects). The extent to which thermal
inactivation hinders the enzymatic reaction, however, is mainly
dependent on the source of the enzyme system.
V. Interaction between Enzyme and Substrate

Lee and Pan (1982) divided the hydrolytic reaction into the following steps: 1) mass transfer of the enzyme molecule from the bulk aqueous phase to the surface of the cellulose particle; 2) enzyme adsorption to the surface of the cellulose and formation of the enzyme-substrate (E-S) complex; 3) transfer of water molecules to the active site of the E-S complex; 4) surface reaction between water and cellulose, promoted by the E-S complex; 5) transfer of the soluble products, glucose and cellobiose, from the cellulose particles to the bulk aqueous phase and 6) decomposition of cellobiose into glucose in the aqueous phase by β-glucosidase.

A. Adsorption

As the first step in the enzymatic reaction is adsorption of the enzyme onto the surface of the substrate, it is important to understand the nature of the process and how it may relate to the overall hydrolysis.

Adsorption of cellulase onto a cellulosic material was first observed by Halliwell and Griffin (1973). Enzyme adsorption was accompanied by the rapid liberation of sugars, so it appeared that adsorption was an integral part of the hydrolytic process.

Factors affecting the adsorption of cellulases to cellulose include; 1) the amount of enzyme present and the enzyme/substrate ratio, 2) the accessible surface area and the affinity of the various cellulase components for the substrate,
3) the physical properties of the enzyme (e.g. charge, size, solubility), 4) the environment (pH, temperature and salt concentration), 5) the nature of the substrate (e.g. purity, pretreatment), and the changing topography of the substrate during hydrolysis (Reese, 1977).

1. **Enzyme Concentration**

   Many investigators have found that the extent of adsorption increases as the cellulose concentration or enzyme concentration is increased, the adsorption behaviour obeying the Langmuir adsorption isotherm (Peitersen *et al.* 1977, Lee *et al.* 1982, Goel and Ramachandran, 1983).

   Using *T. reesei* MCG77 Lee and co-workers (1982) demonstrated that the amount of protein adsorbed onto Solka Floc SW40 paralleled its initial rate of hydrolysis.

2. **Accessible Surface Area, Affinity and Crystallinity**

   The initial extent of protein adsorption has been seen to depend on the specific surface area of the cellulose (Lee and Fan, 1982; Mandels *et al.*, 1971; Bisaria and Ghoose, 1977). In contrast, Goel and Ramachandran (1983) found that the substrate particle size i.e. surface area, and crystallinity had only a limited effect on the adsorption of cellulases in *T. reesei* D1-6 culture filtrates.

   Adsorption was reported to be identical on both amorphous and crystalline cellulose (Klesov, 1982). The degree of crystallinity of cellulose did not influence the adsorption of
endoglucanases from *T. reesei*.

Similarly, Lee *et al.* (1982) found that the extent of adsorption onto Avicel (purified cellulose) was similar to that on Sweco-270 (purified cellulose that was milled in a Sweco vibratory mill and passed through a 270-mesh screen), although the respective SSA (specific surface area) and CrI values of Avicel (5.4 m$^2$/g and 80.8%) and Sweco-270 (24.2 m$^2$/g and 12.5%) were very different. As a result of these findings they concluded that the adsorption affinity, as well as the structural properties have a significant effect on adsorption.

The different affinities of the components of the cellulase complex towards cellulose has been the basis for their fractionation and purification (Nummi *et al.*, 1981; Halliwell and Griffin, 1978). The binding mechanism is very specific, with non-cellulolytic proteins such as albumin unable to bind. The endo- and exo-glucanases both adsorb to pure cellulose substrates. Ghose and Bisaria (1979) report that the endo- and exoglucanase components of the *T. reesei* system are adsorbed to the same extent, while Wilke and Mitra (1975) state that the exoglucanase fraction of *T. viride* filtrates adsorb more strongly than the endoglucanase to Solka Floc. Tan *et al.* (1986) report that the cellulbiohydrolase component of *T. harzianum* E58 filtrates adsorbed more strongly to Solka Floc BW300 than endoglucanase in a column reactor, and had much lower levels of leaching.

Ryu *et al.* (1984), using partially purified cellulases, found that the cellulbiohydrolase component of *T. reesei* had a stronger adsorption affinity towards Solka Floc SW40 than the
endoglucanases; with the endoglucanases consisting of adsorbable and non-adsorbable fractions. The role of the unadsorbed endoglucanases in hydrolysis has been debated. One thought is that they hydrolyze the long-chain, soluble cello-oligosaccharides released during saccharification.

Recent work using highly purified enzymes shows the adsorption behavior is very complex. Beldman et al. (1987) studied the adsorption of six endoglucanases and two exoglucanases, derived from T. viride, onto Avicel cellulose. Three of the endoglucanases and one of the exoglucanases adsorbed strongly; adsorption of the remaining glucanases was much lower. It appeared that a portion of the enzyme molecules was irreversibly adsorbed on the cellulose, even after prolonged desorption time. Kinetic measurements indicated that only a part of the adsorbed enzymes were hydrolytically active. This indicates that some of the adsorbed enzymes may not be bound at their catalytic site, or they may not be able to express their hydrolytic action e.g. as an endoglucanase unless another enzyme, such as an exoglucanase, is present.

Woodward et al. (1988) purified the enzymes from T. reesei L27 and found that the binding of CBH protein to cellulose was much stronger than that of endoglucanase.

The β-glucosidase component of T. reesei does not adsorb to pure celluloses (Mandels et al., 1981) whereas it can adsorb to native lignocellulosics (Puls et al., 1985).

The adsorption affinity of the enzyme for the substrate has a significant influence on hydrolysis. This claim is supported
by the work of Castanon and Wilke (1981) who reported that the 
adsorption of filter paper activity on newspaper was hindered by 
Tween 80. The surfactant caused a more reversible binding 
between the enzyme and the substrate resulting in the enhancement 
of the saccharification.

Similarly, Ooshima et al. (1986) reported addition of Tween 
20 decreases the amount of endoglucanase able to adsorb to the 
substrate, while enhancing its hydrolysis. Surfactants may act 
by shifting the adsorption balance between the exoglucanase and 
endoglucanase to the side preferable to the reaction.

Work by Henrissat et al. (1985) has demonstrated the 
relative proportions of endo- and exoglucanase in a reaction 
mixture determines the degree to which they are adsorbed onto the 
substrate, which in turn decides the extent of their synergistic 
action in hydrolysis.

3. Enzyme Properties

The dimensions of the cellulase molecules determine the 
surface area of the substrate available to it, which in turn 
determines the amount of enzyme able to adsorb to the substrate. 
The interrelationship between cellulase size and structural 
features of the substrate have been discussed in Section III.

4. Temperature, pH and Adsorption

There is a wide variation in the reported data on the effect 
of temperature on adsorption. Part of the reason may be the 
rapid rate of hydrolysis as the temperature is increased, 
resulting in a decrease in available surface area per weight
substrate.

Goel and Ramachandran (1983) and Peitersen et al. (1977) reported that adsorption of *T. reesei* cellulases onto a cellulosic substrate was slightly decreased by increasing the temperature from 5 to 50°C. Kim et al. (1988) observed adsorption of *T. viride* cellulases was highest at 5°C and greatly decreased with increasing temperature.

Conversely, Mandels et al. (1971) found the rate of adsorption on Solka Floc increased with temperature, but also reported the maximum levels of adsorption decreased with increasing temperature.

Mandels et al. (1971) and Peitersen et al. (1977) observed adsorption of the cellulases onto cellulose is largely independent of pH in the range of 3.5 to 6.0. Bisaria and Ghose (1977) found adsorption declined slightly at pH's greater than 4.8, but in no way comparable to the drastic drop observed for catalytic activity. Conversely, Kim et al. (1988) found that pH had a significant influence on the adsorption behaviour of cellulase, with maximum adsorption of cellulase on microcrystalline cellulose occurring at pH 4.8.

B. Alteration in Structural Features

During hydrolysis the structural features of the substrate are changing along with the composition and concentration of the adsorbed enzymes. The drastic decrease in the initial rate of hydrolysis is attributed to several factors, including the transformation of the substrate into a less digestible form.
inhibition of the enzyme action, and a decrease in the amount of adsorbed enzyme. As adsorption of the enzyme to the surface of the substrate is a prerequisite for hydrolysis, a decline in the number of binding sites would retard the rate of hydrolysis.

In the initial phase of hydrolysis the specific surface area (SSA) dramatically increased, then slowly declined throughout the remainder of hydrolysis (Lee and Fan, 1983). The initial increase in SSA was explained by endoglucanase attacking the substrate, generating many new adsorption sites for the exoglucanase. As hydrolysis proceeded the number of binding sites decreased, and small portions of the enzyme were released back into solution. Although the SSA decreased during hydrolysis there were sufficient binding sites for the enzyme to adsorb to ensure efficient hydrolysis.

C. Role of Initially Adsorbed Cellulases in Hydrolysis

The enzymes initially adsorbed to the substrate are primarily responsible for the overall hydrolysis. Lee and Fan (1983) compared hydrolysis of Solka Floc with T. reesei QM9414 cellulases in two systems. In the control system the substrate was incubated with enzyme; while in the second system the substrate was recovered after one hour hydrolysis, then resuspended in fresh buffer. The reducing sugars yields were comparable for the two systems, supporting the assumption that the course of hydrolysis is mainly set by the amount of enzyme initially adsorbed.
D. Kinetics of Enzyme Adsorption and Cellulase Action

In cases where the soluble enzyme concentration is much greater than the amount of substrate surface area accessible to the enzyme, Langmuir Adsorption kinetics may be applied (Appendix A). The basic Langmuir assumptions are as follows: 1) there is no interaction between adsorbed molecules 2) all the surface of the solid has the same activity for adsorption 3) all adsorption occurs by the same mechanism and each adsorbed complex has the same structure and 4) the extent of adsorption is less than one complete monolayer on the surface. The applications of the Langmuir equation are limited to use with purified substrate and purified enzymes. The equation will provide information on the maximum amount of enzyme able to adsorb onto the substrate, and the affinity of the enzyme towards a given substrate.

There have been many excellent reviews on the mechanisms and kinetics of cellulase action (Chang et al., 1981, Gilbert and Tsao, 1983, 1985). These are mainly based on studies using purified celluloses as the model substrate and Trichoderma spp. enzymes.

E. Mechanism of Adsorption

The mechanism of adsorption of the cellulases onto cellulose are not fully understood. The adsorbent is the substrate of the enzyme, and there is a preferential adsorption of the cellulases; with the other enzymes left in solution. Ionic bonding, hydrogen bonding, hydrophobic interactions and permeation effects are all involved in the adsorption of cellulases onto cellulose (Reese,
1982; King, 1965; Kim et al., 1988).

The cellulases appear to have both catalytic and adsorptive sites on the substrate, referred to as "productive" and "non-productive" binding (Beldman et al., 1987; Rabinovich et al., 1982).

It has been postulated that the glycosylated regions of the endo- and exoglucanases are involved in the binding to insoluble cellulose. Van Tilbeurgh et al. (1986) partially proteolyzed CBH I from T. reesei into 56 kDa and 10 kDa portions. The core (56 kDa) protein retained its activity against soluble substrates, but completely lost its activity against insoluble cellulose, concomitant with a decrease in adsorption. The 10 kDa peptide was identified as the heavily glycosylated carboxy-terminal of the native CBH I. They proposed a bifunctional organization of the enzyme; one domain containing the active site, while the other, corresponding to the carbohydrate-rich carboxy-terminal containing the cellulose adsorption site.

VI. OBJECTIVE OF THESIS

Cellulases used for enzymatic hydrolysis have been mainly derived from different strains of Trichoderma, especially T. reesei. The yield of β-glucosidase from these systems is low, limiting the usefulness of these cellulases for commercial applications. During the screening of the Forintek Canada Corp. culture collection it was found that Trichoderma harzianum E58 gives higher amounts of β-glucosidase than most other cellulolytic fungi (Saddler, 1982). In addition, the fungus produced high yields of the other components of the cellulase
system, exoglucanase and endoglucanase. In order to use the cellulases derived from *T. harzianum* E58 in a cellulose conversion process the hydrolytic properties of the system must be better defined. This thesis examined the ability of *T. harzianum* E58 cellulases to degrade the commercial cellulose, Solka Floc BW300, to glucose.

Cellulose bioconversion requires high levels of enzyme to counteract factors limiting hydrolysis, such as end-product inhibition, enzyme inactivation, substrate recalcitrancy and enzyme adsorption. The major advantages of high enzyme loading are faster rates of hydrolysis and increased sugar yields—the main drawback is the high cost of the enzyme (Perez *et al.*, 1980; Wilke *et al.*, 1981; Becker *et al.*, 1982). Judicious use must be made of the enzyme to cut costs and avoid waste. The first objective of this thesis was to identify conditions for efficient hydrolysis. Temperature and pH both impact on enzyme activity in a process. The relationship between enzyme loading and hydrolysis was examined, and various methods of determining the minimal amount of enzyme for efficient conversion were studied.

Enzymatic hydrolysis of crystalline cellulose is biphasic: it is characterized by an initially rapid rate of reaction, followed by a dramatic reduction in the rate. This decline is a function of both the changing behavior of the enzyme towards the substrate, and alteration in the substrate. This thesis addresses the factors limiting hydrolysis. My approach has been to examine the relative importance of enzyme-related and structural-related factors in determining a) the initial rate of hydrolysis and
b) long-term hydrolysis.

Enzyme-related factors limiting hydrolysis that were investigated include; thermal stability, site blockage by denatured enzymes and end-product inhibition. The role of the structural features, crystallinity and surface area, in determining the susceptibility of the substrate to enzymatic attack, and in governing long-term hydrolysis were also examined.
MATERIALS AND METHODS

Microorganisms

All the fungi used were obtained from the Fortinet culture collection. *Trichoderma harzianum* E58 was originally isolated from a lumber yard in Ontario, and is a wild type organism. It was identified as *T. harzianum* in 1982 by Dr. Bissett (Agriculture Canada) and later verified as such, by our curator Dr. K. Seifert.

*Saccharomyces cerevisiae* C495 (NRCC 202001) was grown aerobically at 28°C, shaken at 110 rpm, using the medium of Rainbow (1970).

Cellulase Production

Spores maintained on potato dextrose agar (Difco) slants at 4°C were used to inoculate shake flasks (300 mL, 100 mL working volume) containing Vogel's medium (Montecourt and Eveleigh, 1977) with glucose as the carbon source. Each flask was inoculated with a spore suspension to give a final concentration of $5 \times 10^6$ spores/mL of the medium. After two days incubation (28°C, 110 rpm) the culture was centrifuged (10,000 rpm, 40 minutes), the pellet resuspended in 100 mL of distilled water and used as the inoculum. The mycelial inoculum was added to the enzyme production vessels at a final concentration of 0.3-0.6 mg biomass (dry weight)/mL medium. Enzyme production was carried out in shake flasks (500 mL, 150 mL working volume), or in a 30-L New Brunswick Microferm fermentor, containing Vogel’s medium with
1% Solka Floc BW300 as the carbon source. The fermentation vessels were incubated at 28°C, 110 rpm and harvested after peak activity had been reached. The culture filtrate was collected by filtration through Whatman 934-AH glass microfiber filters.

The fermentor-grown culture filtrates were concentrated by ultrafiltration using the Millipore Pellicon Cassette system fitted with a polysulfone membrane, with a molecular weight exclusion limit of 10,000 daltons. The retentate was freeze-dried at -70°C to produce a free-flowing powder which was stored at 4°C. No loss in cellulase activity was noted after one year storage under these conditions.

See Table 2 for information on the cellulase preparations used for the various experiments.

**Purified Exoglucanase and β-glucosidase**

The purified exoglucanases and β-glucosidase used in this study were kindly donated by Dr. L.U.L. Tan.

Fractionation of cellulase activities: Crude cellulase from the culture filtrate of *T. harzianum* E58 was fractionated by ion-exchange chromatography on DEAE-Sephadex A50 into 4 major protein peaks. The exoglucanase, with a molecular weight of 67,000 daltons, was found to be homogeneous as judged by a single protein band on SDS-PAGE (Tan et al., 1986).

β-Glucosidase was purified following the method of Tan et al. (1987).
Table 2. Cellulase activity and protein concentration of T. harzianum E50 enzyme preparations.

<table>
<thead>
<tr>
<th>Figure Number</th>
<th>Endoglucanase Activity IU/mg protein</th>
<th>β-glucosidase Activity IU/mg protein</th>
<th>Filter paper Activity IU/mg protein</th>
<th>Protein Concentration mg/mg lyophilized powder or mg/mL enzyme solution</th>
<th>Comments</th>
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<tbody>
<tr>
<td>5</td>
<td>14.91</td>
<td>0.84</td>
<td>0.73</td>
<td>0.06 mg/mg</td>
<td>E_{act} at optimal pH</td>
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<td>6</td>
<td>14.91</td>
<td>0.84</td>
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<td>0.05 mg/mg</td>
<td>E_{L}=15 FPU/g substrate</td>
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<td>7</td>
<td>11.98</td>
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<td>1.20 mg/mL</td>
<td>E_{act} at optimal temperature</td>
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<td>8</td>
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<td>0.74</td>
<td>0.72 mg/mg</td>
<td>E_{L}=30 FPU/g substrate</td>
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<tr>
<td>9</td>
<td>11.65</td>
<td>0.66</td>
<td>0.68</td>
<td>0.91 mg/mg</td>
<td>For pH range 3.0-6.0: E_{L}=11.5 FPU/g substrate.</td>
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<tr>
<td>10</td>
<td>12.32</td>
<td>0.51</td>
<td>0.73</td>
<td>0.79 mg/mL</td>
<td>For pH range 6.0-7.5: E_{L}=13.8 FPU/g substrate.</td>
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<td>11</td>
<td>12.24</td>
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<td>0.04 mg/mL</td>
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<td>0.90 mg/mL</td>
<td>E_{L}=25.6 FPU/g substrate</td>
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<td>13</td>
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<td>0.93 mg/mg</td>
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<tr>
<td>14</td>
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<td>0.73</td>
<td>0.93 mg/mg</td>
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<td>15</td>
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<td>N.D.</td>
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<td>0.93 mg/mg</td>
<td>E_{L}=30 FPU/g substrate</td>
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<td>0.93 mg/mg</td>
<td>E_{L}=30 FPU/g substrate</td>
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<tr>
<td>23</td>
<td>17.00</td>
<td>0.59</td>
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<td>0.93 mg/mg</td>
<td>E_{L}=30 FPU/g substrate</td>
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<tr>
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<td>0.90 mg/mg</td>
<td>E_{L}=30 FPU/g substrate</td>
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<tr>
<td>26</td>
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<td>0.91 mg/mg</td>
<td>E_{L}=30 FPU/g substrate</td>
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<td>0.93 mg/mg</td>
<td>E_{L}=24.7 FPU/g substrate</td>
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<tr>
<td>29*</td>
<td>15.60</td>
<td>0.59</td>
<td>0.73</td>
<td>0.93 mg/mg</td>
<td>E_{L}=20 FPU/g &quot;spent cellulase&quot;</td>
</tr>
</tbody>
</table>

E_{act} = Enzyme activity
E_{L} = Enzyme loading
N.D. = None detected

* Substrate generated in Figure 27 (for hydrolysis) was used for the adsorption experiments (Figures 27, 28) and the recalcitrant study (Figure 29).
Substrates

**Avicel pH 101** (FMC Corp., PA, USA) is a microcrystalline cellulose derived from fibrous plants. It is a particulate, free-flowing powder.

**CF 11 Cellulose Powder** (Whatman medium length fiber)

**Signacell Type 50** (Sigma Chemical Co., St. Louis, MO, USA) is a microcrystalline cellulose with an average particle size of 50 μm.

**Solka Floc AS1040** (Brown and Co., NH, USA) was made from purified cotton linters.

**Solka Floc BW300** (Brown and Co., NH, USA) is manufactured from ball-milled sulfite wood pulp; average fiber length of 22 μm. It was used as the standard cellulose substrate for cellulase production, hydrolysis and adsorption studies.

**Solka Floc BW300** (General Filtration, Division of Lee Chemicals Ltd., Toronto, Ont., Canada) is manufactured from ball-milled sulfite wood pulp; average fiber length of 22 μm.

**Solka Floc SW 40** (Brown and Co., NH, USA) was made from hammer-milled wood pulp and was a fibrous grade of cellulose.

Assays

Soluble protein was determined after precipitation with 7% trichloroacetic acid (TCA) by the method of Lowry et al. (1951), using bovine serum albumin (Sigma Chemical Co.) as a standard.

Total reducing sugars were measured colorimetrically using the dinitrosalicylic acid reagent (Miller, 1959).
Glucose was estimated colorimetrically by the glucostat (Sigma Chemical Co.) enzyme assay (Raabo and Terkildsen, 1960). Ethanol was determined on a gas-liquid chromatograph (Varian 4600) fitted with a 6 ft. x 1/8 in. Chromosorb 101 column (Chromatographic Specialities) and a flame ionization detector. The detector was kept at 250°C, the column oven at 180°C and the injector port at 200°C. Helium, saturated with formic acid (Ackman, 1972), was used as the carrier gas at a flow rate of 30 mL/min.

**Enzyme assays**

The different cellulase activities were measured using the method of Mandels et al. (1976). Enzyme unit values were calculated in the dilution range where approximately 0.3-0.6 mg/mL of reducing sugars were detected at the end of the assay. Therefore, the absorbance read at 575 nm was in the range of 0.4-0.8.

Endoglucanase activity [1,4-β-D-glucan-4-glucanohydrolase, EC 3.2.1.4] was determined by incubating 1 mL enzyme solution with 10 mg carboxymethyl cellulose (Sigma Chem Co.; medium viscosity, D.P. 1100, D.S. 0.7) in 1 mL 0.05 M citrate buffer, pH 4.8, at 50°C for 30 minutes. The reaction was terminated by addition of 3 mL dinitrosalicylic acid reagent. The tubes were placed in a boiling water bath for 5 min then cooled to room temperature and the absorbance read at 575 nm.

Filter paper degrading activity determines the "complete cellulase" (endocellulase and exocellulase) system.
activity was measured by the method of Mandels et al. (1976). One mL of enzyme suspension was added to 1 mL 0.05 M citrate buffer, pH 4.8, containing a 1 cm x 6 cm strip (50 mg) Whatman No. 1 filter paper. After incubation for 1 h at 50°C the reaction was terminated by the addition of 3 mL dinitrosalicylic acid reagent.

β-Glucosidase activity [EC 3.2.1.21] was determined by incubating 1 mL of enzyme suspension with 10 mg salicin (Sigma) in 1 mL 0.05 M citrate buffer, pH 4.8, at 50°C for 30 min. The procedure followed was the same as for the endoglucanase assay described above.

One unit of endoglucanase, filter paper and β-glucosidase activity was defined as 1 μmol glucose equivalents released per minute.

Duplicates

All experiments outlined in Figures 5-29, and Table 3 were performed in duplicate, and the values averaged. The variability between duplicates was within 5%.

Effect of pH on Enzyme Activity

Three parts culture filtrate was combined with one part of 0.2 M citric acid-sodium citrate buffer, pH range 3.0-5.2, or one part of 0.2 M citric acid-Na₂HPO₄ (McIlvaine) buffer, pH range 2.6-7.6 (McKenzie, 1979). The enzyme assays were performed using the same buffer.
Saccharification Studies

The majority of the saccharification studies were carried out using Solka Floc BW300 (Brown & Co.) as the standard substrate.

The substrate and buffer were added to the reaction flasks and autoclaved 15 minutes at 121°C prior to addition of the enzymes. The enzyme solutions were filter-sterilized through a 0.45 μm nylon membrane (Nalgene) to prevent microbial contamination, then aseptically added to the reaction flasks.

The substrates used for hydrolysis in the experiments outlined in Figures 15, 23, 25 and 29 could not be autoclaved, as this would alter the substrate and prevent proper interpretation of the results. To avoid contamination, a bactericide, 0.001% thimerosal, was included in the systems. The enzyme solutions were filter-sterilized, as outlined above.

Hydrolysis was carried out at 45°C, with an agitation speed of 140 rpm. The systems were maintained at pH 4.8 using 0.05 M sodium acetate buffer. The enzymatic reaction was terminated by boiling the systems for 10 minutes.

The following equation was used to determine the percent conversion of cellulose (anhydroglucose) to reducing sugars or glucose:

\[
\% \text{ Conversion} = \frac{\text{reducing sugars or glucose (mg/mL)}}{\text{initial substrate conc. (mg/mL)} \times \frac{180}{162}} \times 100
\]

M.W. Glucose = 180
Base Molecular Weight Anhydroglucose = 162

45
Combined Hydrolysis and Fermentation (CHF) Procedure

The conditions for the saccharification studies were modified slightly in the CHF to satisfy the fermentative requirements of the yeast. Serum vials containing 5% (1.0 g) Solka Floc BW300 and 0.5% (0.1 g) yeast extract (Difco) in 5.0 mL of 0.05 M sodium acetate buffer, pH 4.8, were flushed with N₂, sealed with butyl stoppers and aluminum caps, and autoclaved at 121°C for 15 minutes. The cellulases were dissolved in 0.05 M sodium acetate buffer, filter sterilized, and 14 mL volumes were aseptically injected into the vials. A 5% (1.0 mL) inoculum of a 2 day old S. cerevisiae culture containing approximately 10⁶ cells/mL was added immediately after. The vials were incubated at 37°C, 140 rpm for four days, with samples removed at the designated times. The contents of the flask were assayed for reducing sugars, glucose and ethanol. The percent conversion based on the ethanol yield was calculated as follows:

\[
\% \text{ Conversion} = \frac{\text{EtOH (mg/mL)} \times 180 \times \frac{1}{46.07}}{\frac{2}{\text{Biomass factor}} \times \text{Conc. of initial substrate} \times 180} \times 100
\]

M.W. EtOH = 46.07

*Biomass factor accounts for the fraction of produced glucose which is used to fulfill the energy requirements of the organism. It was determined by running a 5% glucose control system in parallel to the CHF (5% cellulose) system and calculated as follows:

\[
\text{Biomass factor} = \frac{\text{actual ethanol production}}{\text{theoretical ethanol production}}
\]
Adsorption Studies

Routine Methodology

The cellulosic substrates were suspended in 0.05 M sodium acetate buffer at pH 4.8 and left to stand overnight at 28°C to ensure complete water saturation of the substrate. The cellulase enzyme solution, at a temperature of 28°C, was added to the vials. Adsorption experiments were routinely carried out at 28°C, 150 rpm for 10 minutes, after which time the system was vacuum filtered for 15 seconds to remove the cellulose residue. Whatman 934 AH glass microfiber filters were used instead of cellulose filters as the former allowed complete recovery of the cellulases. Longer filtration times, of up to one minute, did not alter the amount, or activity of enzyme recovered from the system (Data not shown). Controls for these experiments consisted of enzyme incubated without cellulose for 10 minutes under the same temperature, pH, and agitation conditions.

Enzyme adsorption could not be measured directly and was determined by difference. The supernatant was assayed to determine recovered protein concentration and enzyme activity. Adsorbed protein and enzyme were calculated by subtracting the recovered protein and enzyme in solution from the control.

Adsorption Studies: Effect of pH

Sodium hydroxide adjustment and the Clark-Lubbs buffer system were used to study the adsorption of the cellulases under various pH's. For the pH range of 4.0-6.0 the culture filtrate
was adjusted to the desired pH and added to the substrate vials. The Clark and Lubbs (KH$_2$PO$_4$-NaOH) system was used for studies ranging from 5.8-7.6. After filtration of the system to separate the adsorbed fraction from the unadsorbed enzymes the filtrates were immediately adjusted back to pH 4.8.

**Adsorption Studies: Effect of Temperature**

The methodology outlined above was used, except the substrate vials and enzyme preparation were preincubated at either 4, 28, 37, or 45°C before mixing.

**Adsorption Studies: Various Proteins**

Bovine serum albumin (Sigma Chemical Co.) and the proteases, chymotrypsin (Sigma Chemical Co.) and trypsin (Sigma Chemical Co.) were dissolved in cold (5°C), 0.05 M sodium acetate buffer, pH 4.8. The routine methodology outlined above was used, except the experiment was run at 4°C to avoid proteolytic activity. The protein preparations were added to the substrate at final protein concentrations of 2.4 and 10 mg/g Solka Floc BW300. After filtration the solutions were immediately assayed for protein content.

**Initially Adsorbed Cellulases**

A 5% concentration of Solka Floc BW300 was incubated in 0.05 M sodium acetate buffer, pH 4.8 for 24 h to allow complete saturation of the cellulose. *T. harzianum* E58 cellulases, at 25.6 FPU/g substrate, were added. The system was incubated at
4°C, 150 rpm, for 1 h to allow the enzymes to adsorb to the substrate. The substrate-enzyme complex was collected by vacuum filtration, rinsed with distilled water to remove the unadsorbed proteins, then lyophilized. A substrate control, with no enzymes added, was prepared in parallel.

The lyophilized substrates at a 2% concentration (see below) were prepared in 0.05 M sodium acetate, pH 4.8:

a) substrate-enzyme complex only

b) substrate-enzyme complex and supplemental purified \( \beta \)-glucosidase at 20 IU \( \beta \)-glucosidase/g substrate

c) substrate (control) and cellulases at 25.6 FPU/g and 20 IU \( \beta \)-glucosidase/g substrate (original preparation).

The systems were incubated for 4 days at 45°C, 150 rpm and assayed for release of sugars.

**Determination of Crystallinity**

The water-swollen cellulose was solvent exchanged in order to preserve its swollen state as closely as possible. Air drying results in collapse of the cellulose fibers, changing its structural characteristics. The expanded capillary structure is maintained if the water is replaced by a polar organic solvent, e.g., methanol, which in turn, is replaced by a nonpolar organic solvent, e.g. benzene. When the nonpolar organic solvent is removed by drying, the water-swollen nature of the cellulose remains largely intact (Fan et al., 1980).

Accordingly, the water-swollen cellulose was recovered by vacuum filtration and immediately mixed with methanol to obtain a 5% (w/v) slurry. The mixture was stored at 4°C for 24 hours, then the substrate was filtered. This methanol wash was repeated
twice, then the methanol was replaced with benzene, using three consecutive 24 hour soakings with fresh benzene. The samples were air dried for 48 hours prior to analysis to remove traces of the benzene.

**Crystallinity Indexes**

Crystallinity indexes were determined on a Norelco diffractometer, under the supervision of R. Hartree, Geology Dept., University of Ottawa, following the methodology of Segal et al. (1959).

The crystallinity was measured using the powder method of x-ray diffraction, and scanning for a range of 2θ from 10° to 29°. From the diffractograms the crystallinity indices (CrI) were determined by the following equation:

$$\text{CrI} = [(I_{002} - I_{am}) / I_{002}] \times 100$$

where $I_{002}$ is the intensity of the 002 peak (at about 2θ=22°), and $I_{am}$ is the intensity at 2θ=18°. The $I_{002}$ peak corresponds to the crystalline fraction and the $I_{am}$ intensity to the amorphous fraction.

**Preparation of Amorphous Cellulose**

**Acid-swollen Cellulose**

Phosphoric-acid swollen cellulose was prepared following the method of Walseth (1952). In a 1 L beaker, 30 g of Whatman crystalline cellulose powder was slowly added with mixing to 400 mL of cold, 85% phosphoric acid. The mixture was worked to a
smooth consistency. The acid-swollen cellulose gel was maintained at 4°C for 2 hours in an ice-water bath, and stirred intermittently. Cold distilled water was added to the sample, with mixing, to disperse the gel. The system was vacuum filtered through a sintered glass filter of coarse porosity. The sample was transferred to a 1.5 L beaker containing 1 L of cold, distilled water and blended to form a suspension using a motor-driven, stainless steel stirring blade. This process was repeated until the washings were neutral. After the last wash the sample was vacuum filtered, lyophilized and stored at -25°C.

**Chelating Metal Caustic Swelling Solvent (CMCS)**

The cellulose solvent, ferric tartrate/sodium hydroxide (FeTNa) was prepared following the methods of Hamilton *et al.* (1984). To a 2.0 L glass beaker, surrounded with aluminum foil 207.1 g sodium tartrate and 400 mL water were added. The mixture was gently heated to dissolve the crystals. The solution was cooled to room temperature, then 81.1 g ferric chloride hexahydrate, dissolved in 160 mL water, were added. These solutions were vigourously mixed for 45 minutes while being protected from the light then cooled in an ice water bath to 10°C. A solution of 98 g sodium hydroxide dissolved in 200 mL water was slowly added, with the temperature of the system maintained below 14°C. During the addition of the sodium hydroxide, the colour of the solution shifted from reddish-brown to yellowish-green. At this point 50 g sodium sulfite was added for stabilization. The solution was made up to 1 L with water.
then filtered to remove any ferric hydroxide precipitate. The solvent was stored in a polyethylene bottle at room temperature until use.

The following cellulose substrates were treated with the cellulose-swelling agent: Solka Floc BW300, Solka Floc AS1040, Solka Floc SW40, Avicel, Sigmacell 50, and CF11. A 2.0 g quantity of the substrate was slowly added, with vigorous mixing to the solvent. The volume of solvent used varied with the solubility of the substrate. Solka Floc AS1040, Avicel, Sigmacell 50 and CF11 dissolved in 50 mL solvent per gram substrate, while Solka Floc BW300 and Solka Floc SW40 required 100 mL of solvent per gram substrate.

The CMCS solvent was vigorously stirred while the substrate was slowly added, to avoid clumping. To avoid oxidation of glucose, a problem inherent in the caustic swelling systems, the solutions were protected from light during the dissolving procedure. Some of the substrates did not completely dissolve as indicated by a "cloudy" appearance; and the solutions were left at 4°C overnight until these fine particles dissolved. The solutions were dark green, translucent and very viscous. The cellulose was reprecipitated by slow addition to a 2 L volume of water, while stirring vigorously. The precipitate was recovered by filtering the mixture through glass filters of medium porosity. The precipitate was washed with 2 L of water, then with 1.0 N HCl until all traces of the iron tartrate complex were removed and a fluffy white residue was left. The precipitate (amorphous cellulose) was washed with distilled water until neutral washings were obtained, lyophilized, and stored in a
dessicator.

**Ball-milling**

A 40.0 g quantity of Solka Floc BW300 (Brown and Co.) was milled dry in a 5 L laboratory porcelain jar following the modified procedure of Brownell (1968). Each jar was charged with 4.5 kg of small burundum grinding cylinders (1.25 cm. in diam., 1.25 cm. in length, 207 to the kg). The jars were rotated at 60 rpm for a two week period.

**Spent Cellulose**

A series of Erlenmeyer flasks containing Solka Floc BW300 were enzymatically hydrolyzed by *T. harzianum* E58 cellulases. Flasks were removed at 0, 2, 4, 8, 12, 24, 48 and 96 hours. The flask contents were vacuum filtered and the residual substrate collected. The supernate was assayed for liberated reducing sugars, and glucose. The residual substrate was treated with 0.1% NaOH for 30 minutes to extract the adsorbed proteins. The substrate was sequentially extracted with methanol and benzene, air-dried, then stored in a dessicator. This substrate, termed "spent cellulose" was investigated in Figures 26-29.
RESULTS AND DISCUSSION

I. ENZYMATIC HYDROLYSIS

A. Enzyme Assays

For the enzymatic process to be optimized, the process parameters for the enzyme system must be defined. Both the cellulase assays and hydrolysis of cellulose substrates are performed under the same conditions of temperature and pH. Most Trichodermal cellulases have a pH optimum of 4.5-5.5 and a temperature optimum of 50-60°C (Coughlan, 1985). The first step was to establish the optimum pH and temperature for detection of cellulase activity in T. harzianum E58.

The components of the cellulase preparation were assayed and found to have pH optima in the range 4.6-5.0 (Figure 5). As a hydrolytic process involves an interaction of the enzyme components, the pH optima for the complete enzyme system acting on crystalline cellulose should also be determined. The pH of the cellulase preparations were adjusted between pH 4.0-6.1, in 0.05 M sodium citrate buffer, and incubated with Solka Floc BW300 for 8 hours. Based on the maximum reducing sugars and glucose yields obtained the optimum pH for hydrolysis was 4.8 (Figure 6).

The optimum temperature for the short-term assays was determined to be 50°C for endoglucanase and filter paper activity and 55°C for β-glucosidase (Figure 7). Cellulase activity is traditionally measured by relatively short-term assays (30-60
Figure 5.

Effect of pH on the endoglucanase (○), β-glucosidase (▲) and filter paper (○) activities of *T. harzianum* E58.

Enzyme activity (IU/mg protein) at the optimal pH was:
Endoglucanase, 14.91
β-glucosidase, 0.84
Filter paper activity, 0.73

Incubation times of 30 minutes (endoglucanase and β-glucosidase) or 60 minutes (filter paper activity) were used for the assays.
Figure 6.

Effect of pH on the hydrolysis of a 5% (w/v) concentration of Solka Floc BW300 to reducing sugars (○) and glucose (■) after 8 hours incubation at 45°C.

Enzyme preparation from Figure 5 was used at an enzyme loading of 15 FPU/g substrate.
Figure 7.

Effect of temperature on the endoglucanase (○), β-glucosidase (▲) and filter paper (○) activities of T. harzianum E58.

Enzyme activity (IU/mg protein) at the optimal temperature was:

Endoglucanase, 11.98
β-glucosidase, 0.66
Filter paper activity, 0.73

Incubation times of 30 minutes (endoglucanase and β-glucosidase) or 60 minutes (filter paper activity) were used for the assays.
minutes) using soluble or fairly amorphous substrates. These assays give a fast estimation for the activity of the individual components of the cellulase system and their concentration. Their major flaw is they can not be extrapolated to measure the ability of the enzymes to act against crystalline cellulose, which requires a synergistic interaction of the enzymes. Crystalline cellulose would be a more representative substrate to examine the phenomena of hydrolysis. Accordingly, a crystalline cellulose, Solka Floc BW300 was selected as the model cellulose substrate to study long-term hydrolysis.

Hydrolysis of crystalline material requires longer incubation times of 1 to 4 days. As the cellulases are fairly labile at high operating temperatures the thermal stability of the enzyme system is a prime consideration. The optimal temperature for enzyme activity must be balanced against greater inactivation at higher temperatures. The optimal temperature for long-term hydrolysis was determined by incubating the cellulases with cellulose at various temperatures and comparing the amount of hydrolysis at different times (Figure 8). Although higher temperatures initially resulted in higher sugar yields, as the incubation time increased the differences in hydrolysis for the different temperatures decreased. This agrees with the work of other researchers (Andren, 1978) who found as the time of saccharification increased the maximum extent of hydrolysis occurred at lower temperatures. The best temperature for long-term hydrolysis appears to be 45°C, based on its fast initial rate combined with its maintenance of high yields throughout the process.
Figure 8.

Hydrolysis of a 2% (w/v) concentration of Solka Floc BW300 by *T. harzianum* E58 cellulases at 28 (--), 37 (---), 45 (----) or 50 (-----)°C to a) reducing sugars and b) glucose.

An enzyme loading of 30 FPU/g substrate was used.
B. Initial Rate

Cellulose hydrolysis by cellulases is characterized by an initially rapid rate, which levels off quite drastically after about 8 hours. The initial rate and the extent of hydrolysis is strongly dependent on the susceptibility of the substrate, and the composition and concentration of the cellulase system. The hydrolysis profile of T. harzianum E58 on Solka Floc BW300 was examined to establish its pattern of degradation. Two different enzyme loadings were investigated and their effects on the initial rate and extent of hydrolysis were compared.

The initial hydrolysis rates were rapid, but these rates declined as hydrolysis proceeded (Figure 9). The initial rate of hydrolysis was higher with the higher enzyme loading, as was the levelling off value. A higher enzyme loading gives the system an added advantage against debilitating factors such as end-product inhibition and thermal inactivation which are serious limitations in long-term hydrolysates. The ratio of the initial rate at high loading to the initial rate at low loading was much greater than the corresponding ratio of final yields. This implies that the enzyme faces an increasingly resistant substrate as hydrolysis proceeds that imposes a limit on the ability of the enzyme to hydrolyze it.

Complete hydrolysis of the substrate was a prolonged process. Although the hydrolysis rate slowed down after 8 hours, there was a steady release of sugar throughout the 4 days of incubation at both enzyme loadings. As the objective of a
Figure 9.

Hydrolysis profile of a 1% (w/v) concentration of Solka Floc BW300 incubated with \textit{T. harzianum} ES8 cellulase at enzyme loadings of 50 (—) or 10 (→) FPU/g.
hydrolytic process is to obtain high (>85%) levels of conversion to glucose as quickly as possible, the region where the reaction slows down is of importance in hydrolysis. By defining the factors responsible for the decline in the rate it may be possible to reduce their effect.

C. Adsorption and Hydrolysis

Contact between the enzyme and the insoluble substrate is essential for hydrolysis. The first step in cellulose hydrolysis is adsorption of the enzyme onto the surface of the substrate (Lee and Fan, 1982). Once adsorbed, the enzyme can hydrolyze the substrate. The substrate has only a given number of binding sites available to the enzyme; so hydrolysis is at first limited by the amount of enzyme able to adsorb to the surface, or penetrate into the substrate. Although hydrolysis is routinely performed at pH 4.8 and 50°C, this may not correspond to the optimal conditions for adsorption. To study the relationship between adsorption and hydrolysis, the effect of pH and temperature on adsorption was determined. The pH in the range of 4.0-7.0 had no effect on the adsorption of endoglucanase, filter paper activity or protein (Figure 10).

Neither β-glucosidase nor the protein standard, bovine serum albumin (at 10mg/g cellulose), adsorbed to the substrate under the pH range studied (Data not given). Mandels et al. (1971) and Peitersen et al. (1977) reported the same observation using the T. reesei enzyme system.

Not all the added endoglucanase or filter paper activities
Figure 10.

Effect of pH on the adsorption of a) endoglucanase and b) filter paper activities and c) protein onto a 5% (w/v) concentration of Solka Floc BW300.

The enzyme loading for the pH range 3.0-6.0 was 11.5 FPU/g substrate.
The enzyme loading for the pH range 6.0-7.5 was 13.8 FPU/g substrate.
The % adsorption at a given pH was based on a pH (minus Solka Floc) control to account for any enzyme inactivation.
bound to the substrate, although low enzyme concentrations were used to avoid saturation of the substrate. Approximately 35 percent of the endoglucanase remained in solution. In general Trichoderma spp. secrete multiple forms of endoglucanase. Beldman et al. (1987), Rabinovich et al. (1983) and Ryu et al. (1984) reported a wide variation in the affinity of the different forms of the enzyme towards cellulose, with some components non-adsorbable. Rabinovich and co-workers (1983) suggest the function of the low-affinity, unbound endoglucanase is to act on the soluble, intermediate hydrolysis products of cellulose.

The effect of temperature on adsorption of cellulases derived from T. harzianum E58 onto Solka Floc BW300 was investigated (Figure 11). There was a slight desorption of the endoglucanase at higher temperatures, probably due to hydrolysis of the substrate concomitant with enzyme release. Filter paper activity also exhibited a slight drop in adsorption with increasing temperature. Adsorption onto cellulosic substrates was reported to be only slightly affected by variations in the temperature from 0-60°C (Goel and Ramachandren, 1983; Mandels et al., 1971; Peiterson et al., 1977). Certainly, the slight changes noted in cellulase adsorption with increasing temperature can not be compared to the drastic increase in activity.

Previous workers have found hydrolysis of the substrate is a major problem in studying adsorption at higher temperatures. Hydrolysis of the substrate alters its structural properties, and interferes with the assessment of cellulase adsorption. The problem of hydrolysis can be reduced by using lower incubation temperatures when examining adsorption. A temperature of 28°C
Figure 11.

Effect of temperature on the adsorption of endoglucanase (◇), and filter paper (○) activities and protein (□) onto a 5% (w/v) concentration of Solka Floc BW300.

Enzyme activity (IU/mg protein) profile was:
Endoglucanase, 16.21
β-glucosidase, 0.61
Filter paper activity, 0.90
Enzyme loading was 17.6 FPU/g substrate

Incubation time for the adsorption study was 10 minutes. Thermal controls (minus substrate) were run in parallel to the test systems. No enzyme inactivation was noted for the above temperature range.
was sufficiently low to avoid hydrolysis of the substrate, and so was used throughout this thesis to examine adsorption.

Neither β-glucosidase nor the protein standard, bovine serum albumin (at 10 mg/g cellulose) adsorbed to the substrate in the temperature range examined. Bovine serum albumin, and the proteases, trypsin and chymotrypsin when incubated with Solka Floc BW300 at 4°C did not adsorb (Data not given). This suggests the substrate is specific as to the nature of the molecules able to bind to it.

The optimal temperature and pH range for adsorption was much broader than for hydrolysis, with high levels of cellulase adsorption onto the substrate occurring at the optimal conditions for hydrolysis. Although an enzyme has to be adsorbed to the substrate for it to effect hydrolysis, this does not guarantee that hydrolysis will occur. The appropriate pH (i.e. 4.8) and thermal (i.e. 37-55°C) conditions must be present for catalysis of the substrate to occur.

The substrate appeared to specific as to the nature of molecule able to bind to it; only enzymes acting on the insoluble substrate could be adsorbed.

Adsorption does not limit hydrolysis, as the optimal conditions for hydrolysis lie within the range for optimal adsorption of the cellulases. Hydrolysis, however, can interfere with the assessment of cellulase adsorption. Based on the above results it was decided to use pH 4.8 to study hydrolysis and adsorption - a pH optimal for both processes. Further hydrolysates were run at 45°C, the optimal temperature for a long-term
reaction. Alternately, adsorption studies were carried out at 28°C to avoid alteration of the substrate due to hydrolysis.

D. **Enzyme Loading**

A hydrolysis curve for *T. harzianum* E58 cellulases at various enzyme loadings was established using Solka Floc BW300 (Figure 12). The amount of hydrolysis increased with the enzyme-loading, so that at sufficiently high enzyme levels hydrolysis was virtually complete. Glucose yields were on average only 10% less than the reducing sugar yields, which is high in relation to other *Trichoderma* species (Saddler et al., 1985). This is due to the high levels of extracellular β-glucosidase produced by *T. harzianum* E58. This suggests that supplemental β-glucosidase may not be necessary in processes converting high levels of cellulose to glucose using *T. harzianum* E58 cellulases.

The prime objective of a cellulose-to-glucose process is to obtain high conversion rates using the least amount of enzyme as possible. As seen in Figure 12, at low enzyme loadings the amount of hydrolysis obtained per filter paper unit was high. At enzyme loadings over 50 FPU/g little additional hydrolysis was achieved with additional enzyme. The maximum amount of enzyme that should be used for hydrolysis appeared to lie roughly in the range of 20-50 FPU/gram.

Why did enzyme loadings over 50 FPU only slightly enhance hydrolysis when there was plenty of substrate left for the enzyme to attack? One possible answer was there were no binding sites available on the substrate. As illustrated in Figure 13, there
Figure 12.

Effect of enzyme loading on the hydrolysis of a 2% (w/v) concentration of Solka Floc BW300 to reducing sugars (●) and glucose (■) after 4 days incubation at 45°C.
Figure 13.

Pictorial representation when a) substrate (soluble)>>enzyme (soluble) and b) enzyme>>accessible substrate.

Gilbert and Tsao, 1985
are a limited number of sites accessible to the enzyme at a given
time due to the insoluble, crystalline nature of the substrate
(Lee and Fan, 1983; Gilbert and Tsao, 1985). Once these sites
are filled the rest of the enzyme remains in solution, until an
accessible site is available. The most efficient amount of
enzyme to add for hydrolysis would be the concentration at which
all the sites accessible to enzymatic attack are filled with
enzyme. The above data suggested the substrate may be saturated
with enzymes at an enzyme loading between 20–50 FPU/gram. Enzyme
adsorption may be a means of determining the optimal enzyme
loading for hydrolysis.

E. **Substrate Saturation**

The critical problem in relating adsorption to hydrolysis is
the difference in the nature of the processes. In adsorption one
captures the substrate at a given moment and examines it; while
in hydrolysis one is looking at a continual process.

The first step to examine the relationship between the
amount of enzyme adsorbed to the substrate, and the hydrolysis of
the substrate is to study the initial adsorption. The capacity
of the model substrate, Solka Floc BW300, to adsorb *T. harzianum*
E58 cellulases was determined and compared with its hydrolysis at
a range of enzyme loadings (Figure 14). The amount of adsorbed
filter paper activity and protein increased with added enzyme up
to 25 filter paper units (FPU). After this, the amount of
adsorbed protein and filter paper activity levelled off. In
parallel to this, hydrolysis of the substrate to reducing sugars
increased up to 25 FPU, above which the amount of sugars released
Figure 14.

Relation between a) the amount of enzyme adsorbed onto 5% (w/v) Solka Floc BW300 and b) its hydrolysis to reducing sugars (●) and glucose (■).

Enzyme activity (IU/mg protein) profile was:
Endoglucanase, 14.51
β-glucosidase, 0.55
Filter paper activity, 0.68

Incubation time for the adsorption study was 10 minutes.
Incubation time for hydrolysis was 8 h.
with additional enzyme started to level off. The enzyme loading at which saturation of the substrate occurs appears to correlate with the optimum loading for hydrolysis. These results are in accord with those of other researchers, who have demonstrated a relationship between the amount of cellulase enzyme initially adsorbed to the substrate, and the hydrolysis of the substrate. Lee et al. (1982) reported that the adsorption of cellulases onto cellulose is one of the most important parameters governing the enzymatic rate of hydrolysis. Similarly, Wald and co-workers (1984) found glucose yield reached a maximum when the substrate became saturated with enzyme. Additional enzyme did not increase the yield, as the adsorption process was controlling the overall rate of hydrolysis.

F. **Role of InitiallyAdsorbed Cellulases in Hydrolysis**

Can initial enzyme adsorption be used to determine the minimum amount of enzyme needed for hydrolysis? To do so it must be established that the initially adsorbed enzyme has a strong role in the overall hydrolytic process, and is not just representative of the initial phase.

Mandels et al. (1971) found that the enzyme adsorbed initially was sufficient to digest cellulose with no replenishment of enzyme, even though the liquid phase containing the sugars was continually removed during hydrolysis. Similarly, Lee and Fan (1983) obtained comparable sugar yields after either incubating the substrate with the same enzymes for the entire process or removing the liquid phase after one hour and replacing
it with buffer.

In the work cited above the researchers removed the liquid phase and replaced it with buffer, thereby removing the β-glucosidase along with the unadsorbed cellulases. The ability of the bound cellulases to promote hydrolysis could be handicapped by cellobiose inhibition (Ladisch et al., 1983). To measure the extent to which the initially bound cellulases control hydrolysis, cellobiose inhibition should be removed by supplementing the system with β-glucosidase. For comparative purposes, a β-glucosidase loading equal to the control should be added.

The role of the cellulases initially adsorbed to the substrate in effecting hydrolysis was examined. An enzyme loading of 26 FPU/gram cellulose was used, as this appeared to correspond to the maximum protein adsorption level for the substrate as indicated by Figure 14. *T. harzianum* ES8 cellulases were incubated at 4°C for one hour with Solka Floc BW300 to allow maximum enzyme adsorption (See Materials and Methods; Initially Adsorbed Cellulases). After removal of the liquid phase the substrate was washed to remove unadsorbed proteins, then lyophilized. Part of the enzyme-substrate complex was resuspended in buffer; the other part was resuspended in buffer and supplemented with purified β-glucosidase at levels corresponding to those of the original mixture. A control containing the original enzyme activity and substrate was also prepared.

As shown in Figure 15 the reducing sugars and glucose released by the adsorbed cellulases supplemented with β-
Figure 15.

Hydrolysis of Solka Floc BW300 by adsorbed cellulases after removal of the unadsorbed protein remaining in solution: control (---); supernatant removed and replaced with buffer (---); supernatant removed, replaced with buffer and supplemented with purified β-glucosidase (---).

Enzyme loading of 25.6 FPU/g substrate was used.

The enzyme activity (IU/mg protein) profile of the *T. harzianum* E58 cellulase preparation was:

- Endoglucanase, 15.60
- β-glucosidase, 0.56
- Filter paper activity, 0.79
glucosidase was roughly 90% of that obtained by the control. The reducing sugars and glucose yields obtained by the adsorbed cellulases-alone were only 50 and 25% respectively of the control. As no β-glucosidase was present in the latter system the low sugars yields are largely due to end-product inhibition.

It was very interesting to note that the cellulases which initially adsorbed to the substrate were able to hydrolyze the substrate to an extent comparable to the control. The enzymes that do not bind apparently have only a minor role in the hydrolysis of the substrate, or are in excess. This established that the enzymes initially adsorbed to the substrate were primarily responsible for its hydrolysis. The minimal enzyme requirements for effective hydrolysis may then be determined by examining the adsorption of the cellulases to the substrate to be hydrolyzed.

G. **Substrate Saturation/Langmuir Adsorption Isotherm**

A potential problem in using adsorbed enzymes to determine optimal enzyme loading for hydrolysis is the heterogeneous nature of the cellulase system. The components of the cellulase complex are adsorbed in different proportions to the substrate, and have different affinities towards it. For hydrolysis to be feasible, however, all the components of the cellulase system must be present.

Tan and co-workers (1986) have investigated the enzyme system of *Trichoderma harzianum* E58, and studied its pattern of adsorption onto Solka Floc BW300. They found the major enzyme secreted by
this fungus is exoglucanase. In addition, this enzyme has a stronger affinity for the cellulose, than does the endoglucanases. Using culture filtrate the majority of the adsorbed proteins were the exoglucanases.

Purified exoglucanase, derived from *T. harzianum* E58, was used as a binding probe to determine the substrate saturation level (See Materials and Method; Purified Exoglucanase). Varying concentrations of purified exoglucanases were added to Solka Floc BW300, and the amount of adsorbed protein determined (Figure 16). Adsorption increased with added protein up to 15 mg adsorbed protein/g substrate, after which little additional protein was adsorbed. The substrate is saturated with enzyme at this point, any excess enzyme added either remains in solution or is only loosely associated with the substrate.

The adsorption behaviour obeys the Langmuir Adsorption Isotherm, with the amount of enzyme adsorbed increasing with the amount of enzyme added. As a homogeneous macromolecule and purified cellulose were used, the Langmuir Adsorption Isotherm Equation could be applied to predict the maximum amount of protein able to adsorb to the substrate, and the affinity of the enzyme towards the substrate (Appendix A). According to the Langmuir Adsorption equation, the inverse of the free enzyme plotted against the inverse of the adsorbed enzyme will give a measure of the maximum amount of enzyme able to adsorb to the substrate \((1/b)\) and the affinity of the enzyme towards the substrate. The maximum amount of exoglucanase able to adsorb was calculated as 14.4 mg/g substrate. This corresponds to an enzyme loading of 35 mg protein per gram Solka Floc BW300.
Figure 16.

Adsorption of varying loadings of purified exoglucanase, derived from *T. harzianum* E58, onto Solka Floc BW300.

Incubation time for the adsorption study was 10 minutes.
The relationship between the amount of purified exoglucanase adsorbed to the amount of protein adsorbed in the culture filtrate was examined. In Figure 14 varying concentrations of culture filtrate were added to the Solka Floc BW300, and the amounts of adsorbed and unadsorbed proteins determined. These values were plotted using the Langmuir Isotherm Equation, and it was determined that the maximum amount of protein able to be adsorbed was 15.1 mg/g substrate. This value is comparable to that obtained with the purified exoglucanase, suggesting that the maximum amount of enzyme able to adsorb to the substrate could be determined by examining protein adsorption from the culture filtrate.

To determine the minimum amount of enzyme needed for efficient hydrolysis the saturation level of the adsorbed protein (15.1 mg/g substrate) must be correlated with the total amount of protein added. The amount of enzyme which bound at saturation corresponded to an enzyme loading of 23 filter paper units per gram cellulose.

Using \textit{T. harzianum} E58 cellulases in combination with the standard cellulose substrate, Solka Floc BW300 it has been established that: a) a substrate has a limited number of binding sites for the cellulases to adsorb to in order to effect hydrolysis; b) the cellulases initially adsorbed to the substrate have a major role in governing hydrolysis; c) the enzymes that are not adsorbed in the initial phase of hydrolysis have only a minor role in the hydrolysis of the substrate.

The enzyme loading where the substrate is saturated with
enzyme is evidently the most economical enzyme concentration to use for hydrolysis. With Solka Floc BW300 the optimal enzyme loading was determined to be 20-25 FPU/gram.
II. ENZYME RELATED FACTORS LIMITING HYDROLYSIS

A. End-product Inhibition

End-product inhibition is a serious limitation in the hydrolytic process and adequate β-glucosidase levels are essential to good hydrolysis. The cellulases and β-glucosidase in most Trichoderma spp. are strongly inhibited by cellobiose and glucose, respectively (Ladisch et al., 1983). There are a number of ways to circumvent the problem of end-product inhibition. Supplemental β-glucosidase derived from other sources have been shown to alleviate glucose inhibition and increase the efficiency of the process (Bissett and Sternberg, 1978; Sternberg et al., 1977). Most commercial enzyme preparations require the addition of β-glucosidase for effective glucose production. The use of commercial enzymes would add to the expense of the process. T. harzianum E58 is a high producer of β-glucosidase and may be able to achieve high hydrolysis without being handicapped by end-product inhibition.

Using the previously determined "optimal" filter-paper loading of 25 PPU/gram substrate, the effect of supplemental β-glucosidase on hydrolysis was studied. β-Glucosidase, purified from the culture filtrate of T. harzianum E58 (Tan et al., 1987) was used as the source of supplemental β-glucosidase. Varying concentrations of the purified β-glucosidase were combined with a T. harzianum E58 enzyme preparation and incubated with Solka Floc BW300 for a 4 day period (Figure 17). After a 24 hour incubation, hydrolysis to glucose was enhanced 10% by the addition of 4 IU
Figure 17.

Effect of supplemental β-glucosidase purified from T. harzianum E58 on the long-term hydrolysis of a 2% (w/v) concentration of Solka Floc BW300 to a) reducing sugars and b) glucose.

Enzyme activity (IU/mg protein) profile for the T. harzianum E58 cellulase preparation was:
Endoglucanase, 19.07
β-glucosidase, 0.55
Filter paper activity, 0.85
Enzyme loading was 25.7 FPU/g substrate
β-glucosidase/g substrate. Additional β-glucosidase had little effect on hydrolysis. After 48 hours the beneficial effects of supplemental β-glucosidase were slight. Evidently, the β-glucosidase of T. harzianum E58 is present in sufficiently high quantities to handle end-product inhibition for long-term hydrolysis. The enzyme activity of the control (no supplemental β-glucosidase) was 25.7 FPU and 16.5 IU β-glucosidase per gram substrate. For this enzyme system, acting on 2% Solka Floc, a filter paper activity to β-glucosidase ratio of 1.6 FPU: 1.0 IU β-glucosidase is sufficient to cope with end-product inhibition.

B. Combined Hydrolysis and Fermentation

β-Glucosidase, in converting the cellobiose to glucose, removes the more potent inhibitor of the exo- and endocellulases. Glucose is still present, in increasingly greater amounts, as hydrolysis proceeds. Glucose is also an inhibitor of the cellulases, although at a much reduced level (Sternberg et al., 1977; Bissett and Sternberg, 1978; Hsu et al., 1980).

To avoid the problem of glucose or cellobiose inhibition while examining hydrolysis a combined hydrolysis and fermentation (CHF) process can be used. The glucose is continuously removed from the reaction mixture by fermentation to ethanol, greatly stimulating the hydrolysis of the cellulose (Takagi et al., 1977; Ghosh et al., 1982; Saddler et al., 1982). Since end-product inhibition is virtually eliminated, the unhindered action of the endoglucanase and exoglucanase on the substrate can be assessed.

Enzyme loadings ranging from 0-30 FPU/gram cellulose were
examined. The filter paper to β-glucosidase ratio of the cellulase preparation was 1.0 FPU:1.0 IU β-glucosidase. In one system the yeast, *Saccharomyces cerevisiae*, was added concurrently with the cellulases, and the fermentation vials incubated at 37°C. A hydrolysis-only system was run in parallel at 37°C to observe the extent to which glucose inhibition limits the reaction (Figure 18).

In the hydrolysis-only systems, the amount of sugar released increased with the enzyme loading for the entire range studied. The amount of sugars released per enzyme unit was greater in the CHF systems, especially at the lower enzyme loadings. As the enzyme loading increased above 20 FPU the differences in the sugar yield between the CHF and hydrolysis-only systems narrowed. Twenty to twenty-five FPU has been estimated as the minimal value to fulfill the endoglucanase and exoglucanase requirements for efficient hydrolysis, so hydrolysis is not limited by insufficient cellulases. In the hydrolysis-only system it appears that there is insufficient β-glucosidase at an enzyme loading of 20–30 FPU; enhanced hydrolysis at the higher enzyme loadings is due mainly to the alleviation of inhibition by the additional β-glucosidase. The β-glucosidase required to avoid end-product inhibition is higher than estimated in the above section, with the optimal ratio being 1 FPU:1 IU β-glucosidase. In this case hydrolysis was run at 37°C, rather than 45°C, lowering the activity of the β-glucosidase (Figure 7) and increasing the amount of enzyme required.

In the CHF systems the amount of hydrolysis reached a plateau at an enzyme loading of 15–20 FPU per gram of substrate.
Figure 18.

Comparison of a combined hydrolysis and fermentation process (■) and hydrolysis process (●) on the degradation of a 5% (w/v) concentration of Solka Floc BW300 at day 2 (-----) and day 4 (- - -) using a range of enzyme loadings.

In the CHF systems percent hydrolysis was based on the ethanol yield; no glucose was detected on either day 2 or day 4.

Enzyme activity (IU/mg protein) profile was:
Endoglucanase, 19.87
β-glucosidase, 0.84
Filter paper activity, 0.87
at both day 2 and 4. Additional enzyme had no effect on hydrolysis, indicating that for efficient hydrolysis of this substrate the maximum amount of enzyme that should be used is 15-20 FPU per gram substrate. The enzyme loading corresponding to the plateau point in the CHF process provides an alternate method to initial adsorption to determine the most efficient enzyme loading for hydrolysis of the cellulose. Once the enzyme requirements are met, it is the substrate that imposes a limit on hydrolysis.

C. **Thermal Inactivation**

The extent to which thermal inactivation contributes to the drastic slow-down in hydrolysis is unknown. When enzymes are applied to long-term reactions, such as hydrolysis of crystalline cellulose, thermal inactivation becomes an important concern.

An investigation of enzyme stability was carried out under the conditions considered optimal for the enzymatic conversion of cellulose to glucose (45°C, pH 4.8, four days incubation). The enzyme system was buffered and agitated to duplicate the hydrolysis conditions, but in the absence of substrate. The endoglucanase component appeared to be the most thermal-labile, losing 20% of its activity over a 4 day period (Figure 19). The preparation lost only 10-15% of the filter paper and β-glucosidase activities, and 5% of the purified exoglucanase activity after 4 days.

In order to evaluate the effect of thermal inactivation on the hydrolytic efficiency of the cellulases, the ability of the
Figure 19.

Thermal stability of the endoglucanase (O), β-glucosidase (△), and filter paper (○) activities of *T. harzianum* E58 cellulases and purified exoglucanase (Δ) activity after incubation at 45°C, pH 4.8.

**Enzyme activity (IU/mg protein) profile for the cellulase preparation was:**

- Endoglucanase, 13.30
- β-glucosidase, 0.45
- Filter paper activity, 0.67

Protein concentration of the incubated cellulases was 1.12 mg/mL.

Protein concentration of the incubated purified exoglucanase was 1.04 mg/mL.
preincubated cellulases to hydrolyze cellulose was tested. Enzyme samples were recovered at various times during incubation at 45°C (Figure 19), then incubated with fresh crystalline cellulose for 24 hours at 45°C (Figure 20). Enzymes that had been preincubated for 4 days at 45°C had lost only 20% of its ability to degrade Solka Floc BW300 to glucose. The results indicate enzyme stability is a contributing factor to the slow down in long term hydrolysis.

D. Site Blockage

The effect of enzyme loading and incubation time on the rate and extent of substrate conversion was examined (Figure 21). After 5 days of incubation at the lower enzyme loadings the substrate had a slower rate of hydrolysis, and a lower extent of conversion. Some of the factors partially responsible for this lower conversion have been examined, e.g. end-product inhibition, thermal inactivation. These factors limit the reaction at low enzyme loading (<25 FPU/g), but at higher enzyme loadings their effect is negligible. Complete hydrolysis of the substrate was obtained, but only at very high enzyme loadings. It is curious that the substrate is not completely hydrolyzed after 5 days at enzyme loadings over the predetermined minimal amount (>25 FPU/g).

Site blockage might explain why complete hydrolysis is prevented. During long-term incubation, inactivation of the cellulases occurs, the extent being dependent on the incubation conditions. If these "inactivated" enzyme were irreversibly
Figure 20.

Effect of thermal inactivation on the hydrolytic efficiency of the preincubated cellulases to degrade a 2% (w/v) concentration of Solka Floc BW30O to reducing sugars (●) and glucose (■).

Enzyme fractions were collected at various times during incubation at 45°C (Figure 19). The enzyme solutions were filter-sterilized and added, without dilution, to the hydrolysis systems. After 24 h incubation at 45°C the amount of sugars released by the various systems were compared.

Initial enzyme loading, at t=0 h, was 30 FPU/g substrate.
Figure 21.

Long-term hydrolysis of a 2% (w/v) concentration of Solka Floc BW300 incubated with a range of enzyme loadings.
bound to the substrate, they might block binding or catalytic sites from attack by the active cellulases. This would result in a slow down, or total prevention of catalysis on different sites on the substrate. Site blockage would be more of a problem at low enzyme loadings, as the slower rates of hydrolysis demand longer incubation times, and increase the amount of inactivation.

If adsorption of inactivated enzymes to the substrate is irreversible, then complete hydrolysis of the substrate is not possible, even if fresh additional enzyme is added to the system. To test the existence of site blockage, Solka Floc BW300 was incubated with various levels of enzyme; those encompassing the saturation point, as well as sufficient levels to ensure complete enzymatic hydrolysis (Figure 22). The substrate was incubated for two days, after which time a portion of the enzymes had been inactivated (Figure 19), and should presumably have been in place to block further attack by active enzymes. Complete hydrolysis of the substrate had been achieved at enzyme loadings of 100 FPU/g, so fresh enzyme was added to bring the final concentration of the systems up to this concentration. After 5 days incubation all the systems were 95-98% hydrolyzed, regardless of their original enzyme loadings (Figure 22). If enzyme blockage did occur, the lower enzyme loadings were expected to have lower conversion levels than if all the enzyme had been added directly. These results demonstrate that site blockage by inactivated enzyme is not limiting the extent of hydrolysis. Inactivated enzymes either do not bind to the substrate, or if bound, they are displaced by the active enzyme.
Figure 22.

Measurement of the role of site blockage in hydrolysis. A 2% (w/v) concentration of Solka Floc BW300 was incubated for 2 days with a range of enzyme loadings (Figure 21), then supplemented with enzyme to produce a final enzyme loading of 100 FPU/g. The arrow marks the addition of fresh enzyme.
III. STRUCTURAL FEATURES LIMITING HYDROLYSIS

Once the enzyme requirements are fulfilled it is the substrate that imposes a limit on hydrolysis. The main structural features determining the susceptibility of the substrate to hydrolysis include crystallinity and surface area accessible to the enzyme (Cowling, 1975; Klesov and Sinitsyn, 1981; Fan et al., 1980).

Water has a profound effect on the structure of cellulose. The specific surface area of cellulose increases drastically upon wetting (Browning, 1963), and partial recrystallization of fairly amorphous, dry preparations of cellulose under the action of water occurs (Caulfield and Steffes, 1969). As hydrolysis takes place in an aqueous environment, the changes hydrolysis causes to the substrate should be duplicated prior to any structural analysis.

The initial crystallinity of the substrate has been shown to be inversely related to its rate of hydrolysis. Strong correlation between initial crystallinity and hydrolysis has been reported using both substrate treated in various ways to lower its crystallinity, and substrates derived from different sources. The strong correlation is highly dependent on using water-swollen, dried cellulose rather than unwetted samples, for determination of the crystallinity index (Fan et al., 1980; Klesov and Sinitsyn, 1981).

Klesov and Sinitsyn standardized their cellulose preparations by determining the crystallinity after moistening and air-drying the sample. When cellulose is air-dried from a
water-swollen state the capillary structure collapses and the physical parameters are changed drastically. Alternately, Fan et al. (1980) maintained the water-swollen nature of the cellulose by drying using a solvent-drying technique. The structural features of the water-swollen cellulose are better reflected in the solvent-dried cellulose than in the air-dried cellulose. Accordingly, for these experiments the cellulose sample was water-swollen and solvent extracted prior to determination of its crystallinity index, surface area, or other structural features.

The relationship between structural features and hydrolysis was defined using a range of pure cellulose substrates. All the commercially available cellulose substrates were highly crystalline. Since amorphous cellulose is a useful reference material for studying enzyme-substrate interactions, it was prepared by a number of techniques; ball-milling, solvent-swelling, and phosphoric-acid swelling.

Solka Floc BW300 was ball-milled in several jars for up to two weeks. Based on the literature (Fan et al., 1980) it was predicted that a one week milling was necessary to completely break down the crystalline structure, rendering the substrate amorphous. Five milling jars were charged with the substrate, and milled for 1, 2, 4, 7 or 14 days. As discussed earlier, the milled materials were then water-swollen for 24 hours, then solvent-dried prior to determination of their crystallinity index. Contrary to our predictions, the substrates were found to be completely amorphous after only one day milling, as determined by x-ray diffractometry.
A number of cellulose-dissolving solvents have been developed (Schroeder et al., 1986). Cellulose dissolves in the solvent, losing its crystalline structure. Upon addition of excess water the cellulose is reprecipitated out in an amorphous form, and can be easily hydrolyzed to glucose.

Inorganic acids can be used as solvation agents for cellulose (Walseth, 1952; Tsao et al., 1979). As high acid concentrations are needed for complete solvation, degradation losses are high. Phosphoric-acid swollen cellulose was prepared following the technique of Walseth (1952), producing amorphous cellulose as determined by x-ray diffractometry.

Chelating Metal Caustic Swelling Solvent, CMCS, (Hamilton et al., 1984) was employed as the dissolving agent on a number of crystalline substrates in order to produce amorphous cellulose. The substrates were water-swollen and solvent dried prior to determination of their crystallinity index. X-ray diffractometry of the substrates indicated the substrates had partially recrystallized into the Cellulose II form.

Cellulose has a polymorphic crystallinity. The native cellulose, Cellulose I, exhibits a diffraction pattern different from those of the three "man-made" cellulose crystal forms known as celluloses II, III or IV. Cellulose I occurs as a natural product, a result of biosynthesis. Cellulose II results from mercerization, or solubilization and regeneration of Cellulose I, and once Cellulose II has been generated the transformation remains irreversible. Cellulose III can be made from Cellulose I or II by treatment with liquid NH₃ or some amines, and can be
converted to Cellulose IV by heating in glycerol to 260°C (Sarko, 1986). The x-ray diffractogram of Cellulose II differs from Cellulose I (native cellulose) in having its crystalline peak shifted from 22° to 20° and its amorphous component shifted from 18° to 15°. As this cellulose is radically different from cellulose I, it would likely deviate from the norm in its accessibility to hydrolysis. When these substrates were hydrolyzed concurrently with the other celluloses their conversion values were substantially lower. These substrates were therefore not included in the study of the relationship between crystallinity and hydrolysis.

A. Crystallinity

Once a series of substrates of varying crystallinities had been purchased or prepared, the relationship between hydrolysis and crystallinity could be studied. The substrates were digested with T. harzianum E58 cellulases using an enzyme loading of 30 FPU/gram. Hydrolysis was studied over a four day period to observe the influence of time on the relationship (Figure 23). As expected, the more amorphous substrates were more readily hydrolyzed at all times studied. Surprisingly, a bimodal pattern of hydrolysis was observed; the more crystalline substrate demonstrated a dramatic increase in susceptibility with a small decrease in crystallinity, while the susceptibility of the more amorphous substrates (CrI=65-80) was increased to a lesser extent with a drop in crystallinity. The transition point occurred at approximately CrI=80, and remained constant throughout the 4 days of hydrolysis. These results imply that the relationship between
Figure 23.
Relationship between the initial crystallinity of a substrate and its susceptibility to enzymatic hydrolysis after 8(●), 24(■) and 96(▲) h.

Enzyme loading was 30 FPU/g substrate.
initial crystallinity and hydrolysis may be a reflection of some other factor. When a substrate is pretreated a number of factors are changed simultaneously, i.e. crystallinity, particle size, surface area, degree of polymerization. One, or several, of these structural features may be limiting accessibility in the more crystalline materials, so that a small change in crystallinity may result in a dramatic increase in accessibility.

Caulfield and Moore (1974) also found ball-milling increases the digestibility of both the amorphous and crystalline components of cellulose, especially the latter. This led them to propose that the increase in digestibility is a result of decreased particle size and increased surface area, rather than a result of reduced crystallinity. They suggested coupling crystallinity measurements with surface measurements in order to study more effectively the influence of each structural feature.

The amorphous substrates underwent about 75% conversion to reducing sugars in 8 h, although fairly low enzyme loadings (30 FPU/g substrate) were used. In comparison, the more crystalline substrates were only 5-50% hydrolyzed, depending on their initial crystallinity. It appears that it is the structure of the substrate, not the activity of the enzyme which primarily limits hydrolysis.

B. Surface Area

Cowling (1975), Stone and co-workers (1969), and Greithlein (1985) state the most important structural feature governing enzymatic hydrolysis is the surface area available to the
enzyme. Many effective pretreatments increase surface area either by reduction in particle size (milling), by alkali-swelling, or dissolving and re-precipitating, producing a more susceptible substrate (Fan et al., 1981; Klesov and Sinitsyn, 1981).

There are conflicting reports on the relationship between surface area and susceptibility of the substrate to enzymatic attack. One must consider the technique used to measure accessible surface area when interpreting these results. Nitrogen gas adsorption has been used by several groups (Fan et al., 1980; 1981). As the nitrogen molecules can enter into pores that are inaccessible to the enzyme, surface area available to the enzyme can not be accurately determined.

Direct physical contact between the enzyme and the substrate is necessary for hydrolysis. The amount of cellulase adsorbed is proportional to the accessibility of the substrate to the enzyme (Lee et al., 1982); a substrate able to adsorb more enzyme apparently is more readily hydrolyzable. Adsorption of cellulase, or a protein possessing the same dimensions, provides the most direct measure of surface area available (Mandels, 1985). Berezin and Klesov (1981) using this procedure found the rate of hydrolysis proportional to the surface area.

Direct adsorption of cellulase to the substrate appeared to be both an accurate and quick way to measure the surface accessible to the enzyme. Cellulase adsorption was used to examine the relationship between accessible surface area and hydrolysis of a substrate.

The enzyme adsorption profiles for a number of pure
Figure 24.

Adsorption of varying concentrations of a) endoglucanase and b) filter paper activities and c) protein onto a 5% (w/v) concentration of different cellulosic substrates. Substrates: Avicel (∆), CF11 (O), Solka Floc AS1040 (■), Solka Floc BW300 (●) and Solka Floc SW40 (★).
Figure 24.

Adsorption of varying concentrations of a) endoglucanase and b) filter paper activities and c) protein onto a 5% (w/v) concentration of different cellulosic substrates. Substrates: Avicel (●), CF11 (○), Solka Floc AS1040 (■), Solka Floc BW300 (●) and Solka Floc SW40 (•).

Incubation times for the adsorption study was 10 minutes.
cellulose substrates, incubated with varying concentrations of cellulases, were determined (Figure 24). Avicel and Solka Floc BW300 adsorbed the greatest amount of enzyme, Solka Floc AS1040 and Solka Floc SW40 had intermediate levels of adsorbed enzymes, while CP11 had low levels of adsorption. The pattern of enzyme adsorption onto the substrates was identical for endoglucanase, filter paper activities, as well as for protein. From these results it was predicted that Avicel and Solka Floc BW300, with the highest accessible surface area (i.e. highest amounts of enzyme adsorbed) would have the fastest rates of hydrolysis, and CP11, with the lowest accessible surface area, the slowest.

The digestibility of the substrates was tested using a high and a low enzyme loading. At low enzyme loading equivalent amounts of enzyme were adsorbed to the different substrates, so comparable hydrolysates were expected. At high enzyme loading the benefits of the higher amounts of adsorbed enzyme should come into play, so Avicel and Solka Floc BW300 should be more readily hydrolyzed.

The various substrates were incubated for 4 days, and assayed at various times for reducing sugars and glucose released (Figure 25). Solka Floc BW300 was found to release the highest levels of sugars, at both enzyme loadings, with more sugars produced at the higher enzyme loading. The order of susceptibility of the substrates was the same at both enzyme loadings. Surprisingly, Avicel released only moderate levels of sugars, despite the fact that it had adsorbed the greatest amount of enzyme.
Figure 25.

Hydrolysis of a 5% (w/v) concentration of different cellulosic substrates to reducing sugars and glucose at enzyme loadings of a,c) 5 and b,d) 30 FPU/substrate. Substrates: Avicel (○), CF11 (●), Solka Floc AS1040 (■), Solka Floc BW300 (●) and Solka Floc SW40 (▲).
Figure 25.

Hydrolysis of a 5% (w/v) concentration of different cellulosic substrates to reducing sugars and glucose at enzyme loadings of a,c) 5 and b,d) 30 FPU/substrate.

Substrates: Avicel (●), CF11 (○), Solka Floc AS1040 (■), Solka Floc BW300 (●) and Solka Floc SW40 (▲).
Two facts emerged. The amount of enzyme adsorbed onto a given substrate is related to hydrolysis of that substrate, with more adsorbed enzymes giving higher sugar yields. The effectiveness of the adsorbed enzyme to promote hydrolysis depends on the substrate. Effectiveness can be related to the structural features of the substrate, and the affinity of the enzyme towards the substrate.

IV. STRUCTURAL CHANGES DURING LONG-TERM HYDROLYSIS

Decrease in the hydrolysis rate can be attributed to several factors, including transformation of the structure into a more recalcitrant form, a decrease in the extent of adsorbed enzyme and depletion of the substrate.

There are numerous papers debating the relative importance of the structural features contributing to the increased recalcitrancy of the substrate. Changes in crystallinity (Caulfield and Moore, 1974) and specific surface area (Lee et al., 1980) due to cellulase action have been reported, but the results do not provide a clear picture of the overall hydrolytic mechanism.

A problem in measuring structural changes in the substrate during hydrolysis is the protein remaining adsorbed onto the recovered substrate. These proteins have been inactivated by boiling the substrate to terminate the enzymatic reaction, but the cellulases remain tightly bound to the cellulose, and are not removed by extensive washing. The proteins may interfere in the determination of structural characteristics. While most researchers, investigating structural changes of the "spent"
cellulose, maintain the structural integrity of the water-swollen substrate by solvent extraction, the problem of adsorbed protein is not addressed. To eliminate possible protein interference a method of desorbing the protein from the residual substrate, without altering its structural integrity, had to be devised.

Otter et al. (1984) reported a mild alkali washing removes the bound cellulases from cellulose. A preliminary experiment was done to determine the optimal concentration for enzyme removal (Table 3). A 30 minute incubation in 0.1% NaOH was found to remove 90% of the protein.

Desorption of the adsorbed proteins by alkali, followed by solvent-drying the recovered cellulose, will ensure that a) the structural characteristics of the water-swollen cellulose are retained and b) the sites on the substrate accessible to the enzyme are not blocked by denatured enzymes. These conditions were employed in further studies requiring protein desorption of the spent cellulose (e.g. to study structural changes in the substrate during its hydrolysis).

A. Change in Crystallinity

Conflicting reports are prevalent on the change in crystallinity during hydrolysis. Norkrans (1950), Lee et al. (1983) and Ohmine et al. (1983) found that as hydrolysis proceeds the crystallinity increases, causing the substrate to become increasingly resistant to enzymatic attack. These results suggest that the amorphous portion of the cellulose is preferentially attacked; with the rapid removal of the amorphous
Table 3. Removal of protein adsorbed onto 5% Solka Floc BW300 after 30 minutes incubation in alkali

<table>
<thead>
<tr>
<th>NaOH %</th>
<th>Protein recovered in filtrate, % original</th>
<th>Protein removed by alkali, % original</th>
<th>Total protein recovered, % original</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>47</td>
<td>3</td>
<td>50</td>
</tr>
<tr>
<td>0.02</td>
<td>46</td>
<td>15</td>
<td>61</td>
</tr>
<tr>
<td>0.05</td>
<td>47</td>
<td>30</td>
<td>77</td>
</tr>
<tr>
<td>0.10</td>
<td>48</td>
<td>42</td>
<td>90</td>
</tr>
<tr>
<td>0.20</td>
<td>47</td>
<td>45</td>
<td>92</td>
</tr>
</tbody>
</table>

Protein concentration=0.79 mg/mg powder

$P_L=29.5$ mg/g substrate
portion resulting in a marked increase in the crystallinity during hydrolysis.

In contrast, Lee and Fan (1983) noted only a slight increase in the crystallinity during 72 h hydrolysis of Solka Floc SW40 (From 77.8 to 83.7), with 50% of the substrate hydrolyzed. Other researchers report only a slight (Fan et al., 1980) or no change (Chang et al., 1981) in crystallinity of the substrate throughout hydrolysis.

In light of the conflicting data, we decided to examine the crystallinity index of Solka Floc BW300 at various times during hydrolysis (Figure 26). Crystallinity changed slightly during the initial 24 hours of hydrolysis, increasing from 81.6 to 82.0. During this time, the substrate underwent its greatest period of hydrolysis, releasing 40.6% of its weight in sugars. Although there was a slight increase in crystallinity it is difficult to reconcile this change with the drastic decrease in substrate hydrolyzability.

As the CrI increased only slightly throughout hydrolysis this indicates that a simultaneous hydrolysis of the amorphous and crystalline regions was occurring, rather than a selective attack of the amorphous region. Although the crystalline and amorphous regions are attacked simultaneously the amorphous regions may be more readily hydrolyzed, accounting for a slight increase in crystallinity throughout hydrolysis. These results support an "all-or-nothing" mechanism for cellulose degradation; when attack is initiated on a cellulose chain it goes to completion.
Figure 26.
Change in the crystallinity index of a 5% (w/v) concentration of Solka Floc BW300 during its enzymatic hydrolysis.

Substrate incubated at an enzyme loading of 10 FPU/g substrate was recovered at various times during its hydrolysis at 45°C. The hydrolysate was analyzed for sugars. The residue was deproteinized, solvent-dried then its crystallinity index was determined.
B. Change in Surface Area Accessible to the Enzyme

Increased recalcitrance of the substrate during hydrolysis may be due to a drop in the amount of adsorbed enzyme. If the amount of enzyme able to bind to the substrate declined during hydrolysis, the rate of hydrolysis would be expected to drop. The accessibility of the substrate, Solka Floc BW300, to the enzyme was followed during the course of the hydrolysis (See Methods and Materials; Spent Cellulose). Cellulose recovered at various times during hydrolysis was treated with mild alkali to desorb the bound protens, and solvent extracted. This cellulose was termed "spent cellulose". Adsorption of a fixed concentration of enzyme onto the spent cellulose was measured (Figure 27). During the initial stage of hydrolysis there was a rapid increase in the amount of enzyme adsorbed, signifying a drastic change in the structural features of the cellulose. The amount of enzyme adsorbed almost doubled in the initial 2 hrs of hydrolysis, and remained constant up to 12 hrs, after which the amount of enzyme adsorbed declined gradually. Even after 48 hours of hydrolysis the amount of enzyme adsorbed was greater than for the initial substrate (t=0) even though 50% of the substrate had been hydrolyzed. Lee and Fan (1983) have also observed the protein bound per weight of substrate remained relatively unchanged during hydrolysis. This indicates that the adsorption sites per unit weight of cellulose do not significantly change during hydrolysis, so the amount of enzyme adsorbed onto the surface is not a limiting factor.

The initial rapid increase in the amount of enzyme adsorbed
Figure 27.

Adsorption of T. harzianum E58 cellulases onto a 5% (w/v) concentration of Solka Floc BW300 prehydrolyzed for different lengths of time.

Substrate was recovered at various times during its hydrolysis at 45°C. The hydrolyzate was analyzed for sugars, while the residue was deproteinized then solvent-dried. Cellulase adsorption studies were performed on the dried residue.

Enzyme activity (IU/mg protein) profile of the cellulase preparation used for the adsorption study was:
- Endoglucanase, 11.53
- β-glucosidase, 0.63
- Filter paper activity, 0.73

Enzyme loading for the adsorption study was 24.7 FPU/g substrate
was not surprising. During the early stage of hydrolysis the endoglucanase attacks the substrate, generating a large number of reducing ends for the exoglucanase to adsorb to and cleave. This occurrence was examined further by studying the adsorption of purified exoglucanase onto the spent cellulose (Figure 28). Exoglucanase adsorption onto the spent Solka Floc paralleled the pattern observed using the enzyme mixture. The substrate was drastically altered in the first hour of enzymatic attack, with its accessibility to the enzyme doubling. Apparently, the substrate had been altered to expose more binding sites for the exoglucanase.

Changes are occurring to the substrate that can not be related to either increased crystallinity or lack of surface area accessible to the enzyme. Although the increased crystallinity and changes in the surface area of the substrate during hydrolysis do not appear to limit hydrolysis, they reflect other structural changes.

C. Increased Recalcitrancy of the Substrate

Although there is plenty of substrate, and the amount of enzyme adsorbed to the substrate is high, the rate of hydrolysis drastically drops. The structure of the cellulose is transformed into a more resistant, or less accessible form during hydrolysis.

The extent to which changes in the structural features limit hydrolysis can be determined. Recovery of the unhydrolyzed cellulose throughout the hydrolysis, and testing it for hydrolysis with fresh enzyme will give a measure of the increased
Figure 28.

Adsorption of purified exoglucanase, derived from *T. harzianum* E58, onto a 5% (w/v) concentration of Solka Floc BW300 prehydrolyzed for different lengths of time.

See Figure 27 for description of substrate.

Enzyme loading for adsorption study was 30.65 mg purified exoglucanase/g substrate
recalcitrancy of the substrate. The use of spent cellulose separates changes occurring to the substrate from the enzymatic factors that inhibit hydrolysis e.g. end-product inhibition, thermal inactivation.

Spent cellulose was recovered at various times during hydrolysis (See Materials and Methods; Spent Cellulose). After alkali desorption of the bound enzymes the residue was solvent extracted. A 5% concentration of the spent celluloses were incubated with fresh enzyme for 24 hours (Figure 29). The results indicated the susceptibility of the cellulose to hydrolysis decreased with prolonged incubation or with its extent of hydrolysis. A 70% loss in susceptibility was noted, due only to changes in the substrate.

As discussed earlier, neither an increase in crystallinity, nor a lack of accessible surface area could be held accountable for the reduction in the hydrolysis rate. There are a number of hypotheses as to structural features that may be limiting hydrolysis. Fan and Lee (1983) suggest the cellulose surface area is composed of an active fraction and a less active fraction; the former being easily digested and the latter being the fraction resistant to hydrolysis.

Lee and Fan (1983) claim the transformation of cellulose into the less digestible form cannot be understood by simply examining the individual effects of several major structural features. Several structural features are simultaneously changing during hydrolysis, all contributing to its increased recalcitrancy. It is difficult, if not impossible, to state conclusively that only one factor is limiting hydrolysis. The
Figure 29.

The relationship between hydrolysis of a 5% (w/v) concentration of Solka Floc BW300 and the accessibility of the spent cellulose to further enzymatic hydrolysis.

Substrate was recovered at various times during its hydrolysis at 45°C (See Figure 27 for hydrolysis curve). The hydrolyzate was analyzed for sugars, while the residue was deproteinized then solvent-dried. A 5% (w/v) concentration of the spent cellulose was incubated with fresh enzyme (at an enzyme loading of 20 FPU/g substrate), for 24 hours, 45°C.
synergistic influence of all the structural changes must be assessed.
CONCLUSION

A number of factors affecting and limiting enzymatic hydrolysis using \textit{T. harzianum} E58 cellulases have been investigated. The conditions employed for hydrolysis are not conducive to maintenance of enzyme activity or stability. Factors such as high substrate concentration, high temperature, long incubation times, cellulase adsorption and end-product inhibition, all reduce the activity of the enzyme. The first step was to establish the optimal pH and thermal conditions for detecting enzyme activity. It was found that \textit{T. harzianum} E58 cellulases had a temperature optimum of about 50°C and a pH optimum of 4.8. When applied to a long-term process the stability of the enzymes is a prime consideration. A temperature of 45°C was the best compromise between stability and a fast hydrolytic rate.

As enzyme adsorption is a prerequisite for hydrolysis, the relationship between the two processes was examined. Adsorption has much broader thermal and pH optima than hydrolysis. The optimal conditions for hydrolysis lie within the range for optimal adsorption onto the cellulases, so adsorption does not limit hydrolysis.

The relationship between amount of enzyme adsorbed to the substrate and its hydrolysis was studied. As the concentration of enzyme present in the system was increased the amount of enzyme adsorbed to the substrate increased, up to a saturation point. Hydrolysis of the substrate to reducing sugars reflected this pattern. It was established that the substrate has a
limited number of binding sites for the enzyme to adsorb to. Additional cellulases (i.e. endoglucanase and exoglucanase) remained in solution and had only a minor role in hydrolysis. The enzymes that adsorb to the substrate in the initial stage of hydrolysis (i.e. within the first hour) have a strong role in governing the overall reaction. When working with an insoluble substrate it is the substrate, or the adsorptive process which controls the overall rate. The enzyme loading where the substrate is saturated with the enzyme is apparently the most economical enzyme concentration to use for hydrolysis. This enzyme loading can be determined by studying either adsorption of the cellulase preparation onto the substrate or by employing a combined hydrolysis and fermentation process.

The decline in the hydrolytic rate throughout the reaction can be attributed to changes occurring either to the enzyme or to the substrate.

Exoglucanase and β-glucosidase activity were resistant to thermal inactivation over a 4 day incubation period at 45°C. The endoglucanase component appeared to be the most thermolabile, losing 20% of its activity over the 4 days. Although the cellulases are partly inactivated at 45°C, this does not appear to be a major problem. Site blockage of adsorptive or catalytic sites on the substrate by inactivated enzymes also did not seem to be limiting hydrolysis.

End-product inhibition of the cellulases and β-glucosidase by cellobiose and glucose, respectively, is one of the major factors limiting hydrolysis. Inhibition can be alleviated by
supplemental β-glucosidase or by removing the inhibitors by fermenting them to a less inhibitory product, e.g. ethanol. *Trichoderma harzianum* E58 has high levels of β-glucosidase activity, and is able to achieve high conversion of the cellulose to glucose. Cellulase, added at an enzyme loading of 25 FPU and 17 IU β-glucosidase/g Solka Floc BW300, was able to cope with end-product inhibition of β-glucosidase by the accumulated glucose. Once the problem of end-product inhibition is removed, the major impediment to hydrolysis is the substrate. A faster rate of reaction could be obtained by increasing the susceptibility of the substrate to enzymatic attack.

Crystallinity and surface area accessible to the enzyme were investigated to observe the relationship between structural features of the substrate and its hydrolysis. The relationship between initial crystallinity and hydrolysis appeared to be bimodal. For crystallinities ranging from 80-100, a slight decrease in crystallinity was accompanied by a large increase in its hydrolysability. No clear pattern between accessible surface area of the substrate and its hydrolysis emerged. The changes occurring to the crystallinity and surface area of the substrate, Solka Floc BW300 during hydrolysis were monitored. A slight increase in the crystallinity was found during the hydrolysis period of 4 days. The accessible surface area increased drastically in the initial phase of hydrolysis, then slowly declined through the remainder of hydrolysis. The structural changes could not be correlated to the rapid decline in the rate of hydrolysis during this time. The substrate becomes more resistant to enzymatic attack during hydrolysis due to structural
changes. The extent to which the substrate becomes inaccessible
to enzymatic attack was examined. Cellulose was recovered at
various times during hydrolysis, the bound cellulases desorbed,
and the "spent" cellulose treated with fresh cellulase. A 70% 
drop in the accessibility of the substrate to hydrolysis was
noted; this indicated that the substrate became more recalcitrant
during hydrolysis.

The objective of a cellulose conversion process is to obtain
high yields of glucose, using low enzyme loadings and short
residence times. There are two main routes for further research
into this area: to use a more "active" enzyme system or to
increase the susceptibility of the substrate. The reaction rates
of cellulase on amorphous cellulose are rapid, achieving 90%
conversion to reducing sugars at low (30 FPU/g) enzyme loadings
after 24 hours. So, it is the structure of the substrate, not
the activity of the enzyme which primarily limits hydrolysis. In
addition, the greatest problems in the course of hydrolysis e.g.
adsorption of the celulases to the substrate, mass transfer at
high substrate concentration and long residence times, are due
more to the insoluble and recalcitrant nature of the substrate.
Treatment of the substrate prior to, or during, hydrolysis would
probably be a promising area on which to concentrate further
research.
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APPENDIX A

Determination of Maximum Enzyme Adsorbed and Enzyme Affinity

Many investigators have found that the extent of adsorption increases as enzyme concentration increases - the adsorption behaviour obeying the Langmuir Adsorption Isotherm (Hendels et al., 1977; Lee et al., 1982). Using the data generated in Figures 14 and 16 the maximum amount of enzyme and protein able to adsorb to the substrate, and the relative affinity of the cellulosics to the substrate can be determined using the Langmuir isotherm equation. The Langmuir adsorption isotherm equation relates the adsorbed enzyme or protein concentration to the free enzyme, or protein concentration according to the equation:

\[
\frac{X}{X_m} = \frac{K_rE}{1 - K_rE}
\]

Where
- \( X \) = activity adsorbed/g substrate
- \( X_m \) = maximum activity adsorbed/g substrate
- \( K_r \) = half-saturation constant for enzyme (IU/mL)
- \( E \) = free enzyme activity in filtrate (IU/mL)

The isotherm equation can be rearranged to give:

\[
\frac{1}{X} + \frac{1}{EK_rX_m} = \frac{1}{X_m}
\]

Where a plot of 1/E versus 1/X will give a straight line with the slope \( 1/K_rX_m \) and the intercept \( 1/X_m \). Similar equations are obtained for the protein adsorption, by exchanging \( P \) (mg/mL) with \( E \) (IU/mL) and \( P_{ads} \) (mg/g substrate)
for X (IU/g substrate). The parameters, $K_i$ and $\chi_m$ for Solka Floc BW300 were calculated by linear regression. Although the Langmuir isotherm is a useful equation, it must be kept in mind that other factors are present which the Langmuir equation does not take into account, e.g. formation of a cellulase complex on the substrate during adsorption, the heterogenous nature of the substrate, the multiplicity of the enzyme system.
APPENDIX B

Crystallinity indices for commercial cellulosics and pretreated cellulosics as determined by x-ray diffractometry.

<table>
<thead>
<tr>
<th>Substrate</th>
<th>Crystallinity Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sigmoid 50</td>
<td>90.8</td>
</tr>
<tr>
<td>CF11</td>
<td>95.6</td>
</tr>
<tr>
<td>Avicel</td>
<td>92.8</td>
</tr>
<tr>
<td>Solka Floc AS1040</td>
<td>91.1</td>
</tr>
<tr>
<td>Solka Floc SW40</td>
<td>84.9</td>
</tr>
<tr>
<td>Solka Floc BW300</td>
<td>84.9</td>
</tr>
<tr>
<td>(Brown &amp; Co.) air-dried</td>
<td></td>
</tr>
<tr>
<td>Solka Floc BW300</td>
<td>83.3</td>
</tr>
<tr>
<td>(Brown &amp; Co.), treated with 0.4% NaOH then solvent-dried</td>
<td></td>
</tr>
<tr>
<td>Solka Floc BW300</td>
<td>80.7</td>
</tr>
<tr>
<td>(Brown &amp; Co.), boiled for 5 minutes, then freeze-dried</td>
<td></td>
</tr>
<tr>
<td>Solka Floc BW300</td>
<td>71.4</td>
</tr>
<tr>
<td>(Brown &amp; Co.) General Filtration</td>
<td></td>
</tr>
<tr>
<td>Solka Floc BW300</td>
<td>62.5</td>
</tr>
<tr>
<td>(Brown &amp; Co.) ball-milled 1 day</td>
<td>0.1</td>
</tr>
<tr>
<td>2 days</td>
<td>0.1</td>
</tr>
<tr>
<td>4 days</td>
<td>0.1</td>
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<tr>
<td>7 days</td>
<td>0.1</td>
</tr>
<tr>
<td>14 days</td>
<td>0.1</td>
</tr>
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</table>

Whatman crystalline cellulose phosphoric-acid swollen

Unless otherwise stated the samples were water-swollen then solvents dried prior to the crystallinity determination.
X-ray diffractograms of a) crystalline celluloses  b) amorphous celluloses and c) FeTNa- swollen cellulose.