1 Introduction

The detergent industry has been using enzyme technologies in detergent formulations for the last 30 years. Enzymatic hydrolysis in detergency is usually focused on the fragmentation of the soil or stains into smaller particles, which are then more accessible to the chemical ingredients of the detergent formulation [1–3]. Today, enzymes are commonly used in many industrial applications and 75% of the industrial enzymes are hydrolases, of which carbohydrolases represent the second largest group [4–7]. Recently, the use of novel enzymes that are claimed to effect textile substrate modifications have been reported, and it has been assumed that these represent advances for textile industries [8, 9].

Cellulases comprise a group of different acting enzymes that, in coordinated action, degrade cellulose and their derivatives to glucose (endoglucanase, exoglucanase or β-glucosidase) [10–18]. The complex interactions among endoglucanase [19], cellobiohydrolase and β-1,4-glucosidase change substrate characteristics during the hydrolysis and these changes are represented in Fig. 1. As can be seen from the figure and the bibliography, endoglucanases cut at random amorphous sites of the cellulose chain, producing oligosaccharides of various lengths and new reducing and non-reducing ends on the fibre surface. During the cellulose hydrolysis, the solid substrate characteristics vary, with the number chain ends changing due to the action of endoglucanase and exoglucanase. This alters the cellulose accessibility as a result of sub-
strate consumption and cellulose fragmentation [20–24].

In detergency, cellulase action should be confined to the first stages of hydrolysis to avoid modification of fabric structure, and the cellulose characteristics need to be verified. Two types of advantages can be identified: (i) dirt anti-redeposition benefits, as the partial interaction with single cellulose chains permits an easy release of the trapped soil, and (ii) biopolishing or depilling effect, which will create colour revival and softening effects on the fabric. The colour revival and anti-pilling effect produced by cellulase action are indispensible benefits and cannot be easily achieved in detergency by other detergent ingredients. This effect can also be considered as an indicator for the care and protection of textiles [12, 14]. A proposed mechanism for the release of dirt particles from cotton fibres is shown in Fig. 2. In general, dirt usually adheres to the damaged cotton and endoglucanases hydrolyse the cellulose chains at random amorphous sites. As a result, the fibres become more accessible by chemical ingredients of the detergent formulation.

It has been known for a long time that alkaloophilic Bacillus species are producers of alkaloophilic cellulases, which show pH optima in the neutral to alkaline range. Cellulases from Bacillus species generally show good compatibility with subtilisin proteases, the use of which is interesting for detergent applications. These monocomponent cellulases present reduced anti-pilling properties [17]. The first bacterial alkaline cellulase was produced by Kao and it exhibited endo-β-1,4-glucanase activity (EC 3.2.1.4). The term „alkaline endoglucaanse“ indicates an endoglucanase with an optimum pH above 7 and retaining more than 70% of its optimal activity at pH 10 [25]. These types of enzymes have been advertised as having excellent detergent effects, as shown on sebum-stained cot-

Figure 1. Main stages for degradation of cotton fibres by cellulase are presented. Endoglucanases cut at random amorphous sites of the cellulose.

Figure 2. General mechanism for release of dirt particles from cotton fibres. (A) Damaged cotton fibres. (B) Dirt adheres to the damaged fibres. (C) Cellulases act on the damaged cellulose microfibrils, by hydrolyzing and suspending them in water. Adapted from Husum and Friis-Jensen [28].
ton [26]. Alkaline cellulases contribute indirectly to the removal of soil trapped in the amorphous region of cotton fibres by its reaction with cellulose molecule in the region. Murata et al. [14] proposed a mechanism based on the binding of water, which only occurred at hydroxyl groups of cellulose molecules in the amorphous region of cotton fibres. In fact, it was verified that soil entering the interfibre space of amorphous cellulose in the interior of cotton fibres, was slightly hydrolyzed and removed by alkaline cellulase. In addition, these cellulases cause no damage to the cotton fibre, solving problems relating to tensile strength loss after multiple wash cycles [15, 17, 27–30]. In addition, cellulases were shown to contribute to the whiteness of cotton-containing textiles after several washes [31].

In this work, the alkaline endoglucanase BaCel5 was used, which was commercialized as a preparation developed by Novozyme (Bagvaerd, Denmark) and called Celluclean®. BaCel5 contains a cellulose-binding module (CBM) consisting of two tandem CBMs, one family 17 CBM and one family 28 CBM, both classified as B CBMs, which is supposed to be highly selective regarding amorphous cellulose and to have glucan-binding properties [32, 33].

The main focus of this research was to study the detergency performance of cellulases when several wash cycles of cotton textiles were applied. Cotton garments treated by multiple washings with BaCel5 were characterised in terms of the modification of the accessibility of cotton fibre using iodine sorption value (ISV) tests, reducing-end production on the surface fibre and the interaction between the enzyme and the cellulose fibre using FITC-labelled enzyme. Scanning electron microscopy (SEM) was used to verify the fibre surface morphology.

2 Materials and methods

2.1 Enzyme and fabrics

The glycoside hydrolase nomenclature suggested by Henrissat and colleagues [11] has been adopted in this study, where Bacillus spp. cellulases are represented with the prefix Ba. BaCel5 (cellulase type 5; EC 3.2.1.4) was supplied by Novozymes (Bagvaerd, Denmark), under the trademark of Celluclean®, with activity of 325.4 EGU/g.

Unbleached cotton woven fabrics (24/21 ends/picks per cm and area density of 140 g/m²) were used as the cellulosic substrate for the enzymatic treatments. Unbleached cotton was selected to avoid interferences in the fluorescence assays.

The proteins bovine serum albumin (BSA), glucose and fluorescein isothiocyanate (FITC) were obtained from Sigma. All other reagents used were of analytical grade.

2.2 Enzymatic treatments

Cotton fabrics (4 g) were treated with BaCel5 cellulase with an enzymatic dosage of 1 and 0.5 mg protein/L, in 800 mL of a model detergent at a 1:200 fabric to liquor ratio, at 40°C. These studies were performed in a Detergent Tester (Copley Scientific, Nottingham, UK) with orbital agitation of 120 rpm. Two different treatments were performed: one single treatment of about 80 min and four consecutive treatments about 20 min each, adding new enzyme at the beginning of each treatment. Control samples were incubated in the same solution without enzyme and samples were removed after each incubation period. The cotton samples were analysed using an iodine adsorption test and an assay was also performed with FITC-labelled BaCel5. Since these series of experiments were designed to highlight the effect of repeated washing cycles when an alkaline cellulase was used, treatments using several washes were compared to the long single-wash treatments to evaluate the effect on cotton fibre surface of repeated washing using normal household conditions such as temperature, time and protein concentration. BaCel5 is believed to be stable for the whole 80-min washing cycle since sixfold more cellobiose is produced by a 120-min compared with a 20-min treatment under comparable conditions (our unpublished observations).

Additional repeated treatments were performed using the conditions previously described (enzymatic dosage, liquor ratio, agitation and temperature) to characterise the cotton surface. Some samples were treated in ten consecutive cycles of about 20 min each, others for 200 min and, finally, some for 48 h. The cotton samples were analysed using reducing-end measurements and SEM.

It is important to note that, for all repeated treatments performed, new enzyme solution was added for each treatment. After enzymatic treatment, all samples were washed with deionised water for three 15-min cycles of at room temperature and, finally, air dried. BaCel5 protein (2 mL, 0.2 g/mL in a sodium carbonate buffer; pH 9.0) was incubated with 100 µL of FITC solution (1 g/L) at room temperature for 2 h. Unlinked FITC was separated by dialysis until no more FITC was released. Cotton samples were treated in this solution at 40°C, at 125 rpm, for four repeated 20-min treatments and a long treatment of 80 min. The cotton samples were embedded in an
epoxy resin and transverse 15-μm cuts of the fibres were prepared using a microtome (Leitz). Cotton fibre cross-sections were analysed by a fluorescence microscope (Leica DM 5000B) at a magnification of 100× and 40×.

### 2.3 Physical analysis

The accessibility of cotton fabrics after cellulase multitreatment was measured using an ISV assay. Cellulose fabric (0.3 g) was added to 1.2 mL iodine-reactive solution (containing 5 g iodine and 40 g potassium iodide in 50 mL of water). Subsequently, 100 mL 200 g/L sodium sulphate was added and, following a 1-h incubation in the dark, the mixture was filtered. The iodine remaining in solution was titrated with standard sodium thiosulphate (0.01 M). The ISV was calculated for each treated sample (as mg adsorbed iodine/g cellulose), and the differential ISV (ΔISV) for each treated fabric was correlated to the respective ISV value from control fabrics [34, 35].

The number of reducing ends, or reducing power, of cotton fabric samples after enzymatic treatment was measured by the method described by Cavaco-Paulo and colleagues [36, 37], which depends on the complex formed between neocuproine (2,9-dimethyl-1,10-phenanthroline) and reducing groups in the fibre. For the measurement, 0.1 g sample was incubated with 2 mL 2% sodium carbonate and 5 mL reactive aqueous solution of neocuproine containing: 0.2 g/L copper (II) sulphate pentahydrate and 0.4 g/L neocuproine. The mixture was boiled for 5 min and cooled to room temperature. Three analyses were done for each fabric sample. Absorbance was measured at 465 nm in a Helios Gamma UV-Vis spectrophotometer (Thermo Scientific, Waltham, MA, USA). The concentration of reducing sugar (g/L) was determined against glucose standards.

The surface morphology of cotton fabrics before and after enzymatic degradation was observed in a JEOL 5310 SEM. The samples were previously sputter-coated with gold with an ion sputter JEOL JFC 1100 to increase their conductivity. Free surface micrographs were taken at various magnifications and their microstructures were studied.

### 3 Results and discussion

The benefits of alkaline cellulases in detergent compositions have already been studied [13, 25, 26, 38–40]. Hoshino et al. [12] studied the mechanism of release of trapped soils with the alkaline cellulase, without any significant liberation of hydrolys products. Several benefits for the textiles had been claimed from studies of their interaction with dyes and surfactants and colourimetric investigations after several washes using these alkaline cellulases. The release of hydrolysis products is a possible final step in the degradation process, but no significant release was detected for these alkaline cellulases.

Specific characterisation was also required to evaluate the reproducibility of the action of the enzyme on the fibre under the same detergency conditions and, for example, at lower enzyme concentrations. For this, the investigations of the sorption properties of cellulose were focused primarily on the study of the variation of cotton fibre accessibility (measuring ΔISV) and the definition of enzyme action, by labelling the enzyme fluorescently (with FTIC), and comparing samples after different types of treatment. The fibre accessibility and the ability of cellulose fibres to adsorb low molecular weight compounds give valuable information about the resulting fine structure (crystalline and amorphous regions, microfibrillar structure and morphology) [35]. The actual surface area accessible to the enzyme molecules in the solution also includes the pores, cavities and spaces between microfibrils. Different studies have found that the relative digestibility of cellulosic substrates is directly proportional to their accessibility to enzyme molecules [41]. The accessibility of cotton fabrics after cellulase multitreatment was measured using the ISV test, as the incorporation of iodine provides important information about the assimilation of low molecular weight species into the cotton fibre. First, four consecutive treatments of about 20 min each and then one single treatment of about 80 min were performed. Low enzymatic concentrations, renewed for each repetitive treatment, were used to reproduce the enzymatic levels employed in laundry detergent applications. According to previous work on alkaline cellulases [25], the lack of significant release of reducing sugar from cellulosic fibres under certain conditions could indicate that the fabrics have not been damaged. Therefore, enzyme activity on the cotton fabrics was measured in terms of reducing-end production and fibre accessibility.

The ISV method involves quantifying the iodine adsorbed per gram of cotton. These values allow comparison of samples with morphological differences on the cotton fabric after wash treatments with BaCel5. Control samples undergoing the same treatment but without enzyme were prepared for each treated sample. ΔISVs were calculated and these values led to the visualization of the differences in the accessibility of the fabric after treat-
ment. A positive $\Delta ISV$ is assumed to indicate an increase in the accessibility of cotton fabric; the higher ISV from the enzyme-treated fibres revealed an increased accessibility of amorphous areas. On the other hand, a negative $\Delta ISV$ implies a decrease in accessibility [34, 35].

The results shown in Fig. 3 from treatments using high and low BaCel5 concentrations revealed that several short treatments led to an increase in accessibility on the fibre, and that no significant differences were seen with samples treated for 80 min.

Samples from each successive treatment were collected to correlate each cycle with the final results (Fig. 3). As observed in Fig. 4, an increase in fabric accessibility was found from at least the second consecutive treatment, for both high and low BaCel5 concentrations. Comparing single treatments of 20 min and 80 min, $\Delta ISV$ results indicated that even at the shorter treatment times the enzyme produced an increase in the accessibility. When applying repeated treatments, BaCel5 was able to produce breaks in the cellulose across the fibre surface during a short hydrolysis period, acting as an endoglucanase. The results showed that the action of BaCel5 was enhanced over the repetitive treatments, but that this effect did not appear for a single long treatment. Therefore, BaCel5 appears to acts on the fabric surface after several short treatments, rather than penetrating the fibre body. This fact is interpreted as a positive effect because it is generally accepted that enzymes must act on the fabric surface to avoid fabric damage. Cellulases should act on the fibre by causing fragmentation of cotton fibres. Shorter and weaker fibres are potentially easier to process by low enzymatic concentrations.

The cross-section of a fibre in cotton after labelled enzyme treatment is shown in Fig. 5. After treating cotton fabrics with FITC-labelled enzyme, samples were examined by fluorescence micro-analysis. Only traces of the residual fluorescence of the cotton itself were observed for the control sample, representing the initial state of the cotton fibre. In contrast, green-surrounded surfaces were detected on the surface of the cotton fabric for both treatments (four repeats of about 20 min and one of about 80 min). When several short treatments were applied, a higher intensity around the fibre was achieved compared to long single treatment. Therefore, BaCel5 appears to acts on the fabric surface after several short treatments, rather than penetrating the fibre body. This fact is interpreted as a positive effect because it is generally accepted that enzymes must act on the fabric surface to avoid fabric damage. Cellulases should act on the fibre by causing fragmentation of cotton fibres. Shorter and weaker fibres are potentially easier to process by low enzymatic concentrations.

The fibre properties were also investigated by measuring reducing ends and by SEM. In these cases, treatments were modified by increasing the number and, consequently, the length of treatments: ten consecutive treatments of about 20 min.
were compared with one single treatment of about 200 min, instead of four short repetitions and 80 min. These modifications were performed to observe the effect of a higher number of washing cycles compared with a single cycle. The duration of the longer single treatment was chosen as 200 min to match that of the ten cycles of about 20 min to avoid the variable of the time. Low enzymatic concentrations were added at each repeat treatments and no reducing sugar production was found in the liquor treatment (data not shown).

Reducing ends on cotton were measured and SEM micrographs were evaluated to identify the effect on the fabric surface (Figs. 6 and 7). Cotton samples were subjected to ten consecutive treatments of about 20 min, or one treatment for about 200 min or 48 h to attain measurable changes of cotton surface properties in terms of reducing ends on the cotton surface for longer treatment conditions.

These data indicated that samples treated several times with BaCel5 attain the highest value compared with samples treated for about 200 min or 48 h. The repeated application of enzyme, by replenishing the enzymatic solutions, caused more fragmentation of cellulose end terminals than in the longer treatments. These results are important because they show the impact of using repeated BaCel5 washes, which are not observed in single and longer uses.

The cotton samples without enzymatic treatment appeared mostly as smooth surfaces but some defects were observed (Fig. 7). Micrographs B, C and D in Fig. 7 show the physical changes of the cotton fibres after each treatment. Cotton treated several times showed an significant number of loose fibrils. However, BaCel5 action appears to hydrolyse fibrils at the surface of the fibres, without penetration or damage of the interior of the fibres. In the case of longer treatments, similar numbers of created fibrils were observed in both cases, but fabric damage was also observed in the 48-h samples. The similar results obtained for 200-min and 48-h treatments may be due to the crystalline nature of cotton cellulose. Others [36, 37] have verified that monocomponent enzymes with long measurable enzyme activity do not yield any measurable reducing sugar products after a few hours of treatment. This particular enzyme has been shown to be able to yield sugar from fresh cotton surface with no mechanical agitation. Our enzyme was expected to have a similar stability since it was shown to remain active at least for 120 or 20 min.

As the aim of this study was to analyse the effect of several BaCel5 treatments on cotton fibres, reproducing the household washing, no further analysis regarding the activity in longer treatments.

![Figure 5](image_url)

**Figure 5.** (A) Control samples, treated without enzyme. (1 mg protein/L; 40°C; 125 rpm). (B, C) Fluorescence microphotographs of fibre cross-sections of cotton treated with FITC-labelled BaCel5; (B) four treatments of about 20 min and (C) 80-min treatment.

![Figure 6](image_url)

**Figure 6.** Percentages of reducing ends for indicated incubation periods (1 mg protein/L; 40°C; 125 rpm). Values represent the mean of three independent experiments and error bars the SD.
were carried out. The defects may have resulted from a combined action of inter-fibre friction and mechanical agitation during the treatments.

The first step in enzymatic degradation of the cellulose microfibrils, leading to the formation of even thinner subfibrils, has been proposed by several authors; however, a deeper appreciation of the morphology and structure of cotton fibres is required from the cleaning standpoint [21, 23, 39, 42–52]. In the present studies, the enzymatic action of repeated treatments resulted in the production of short fibres on the cotton fibre. The SEM studies confirmed the short fibre formation on repeated enzymatic treatments. Although release of short fibres was observed within 20 min, an increase of the accessible surface area was verified and was used to monitor the impact of the enzyme. The enzyme might preferentially attack new areas of the fibre, rather than continuing to release the microfibrils that had been cut in the previous washing cycles into the liquor bath. The short fibre formation was assumed to represent the reducing groups on the cotton surface. It is notable that surface modification occurred on repeated treatments and not in longer single treatments. This may be attributed to preferential hydrolysis of short fibres formed by endoglucanase action. Many researchers have assigned the formation of short fibres to a property of endoglucanases [53].

4 Concluding remarks

Cotton fibres can be converted into microfibrillar material by the action of enzyme hydrolysis in short periods of time using an endoglucanase-cellulase system. The production of fibrillation on cotton fibre surface without any release of cellulosic material (reducing sugar generation) proves that BaCel5 tends to be an action of the first stage of cotton hydrolysis, using low concentrations of cellulase and short time treatments. Therefore, the use of BaCel5 helps to avoid fabric damage, thus enhancing fabric care benefits.

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5 References


