ENZYMATIC TREATMENT FOR PREVENTING BIOFILM FORMATION IN THE PAPER INDUSTRY

E. Torres\textsuperscript{a}, G. Lenon\textsuperscript{b}, D. Crapéri\textsuperscript{b}, R. Wilting\textsuperscript{c}, A. Blanco\textsuperscript{a}

\textsuperscript{a}Chemical Engineering Department. Faculty of Chemistry. Complutense University of Madrid. Madrid 28040. Spain. \textsuperscript{b}CTP, F-38044 Grenoble Cedex. France. \textsuperscript{c}Novozymes A/S, Krogshojvej 36, 2880 Bagsvaerd, Denmark

*Corresponding author/Mailing address, Angeles Blanco

Chemical Engineering Department

Faculty of Chemistry. Complutense University of Madrid

Av. Complutense s/n. Madrid 28040

Spain

Tel, +34 91 3944247

Fax, +34 91 3944243

e-mail, ablanco@quim.ucm.es
Abstract
Microbiological control programmes at industrial level should aim at reducing both the detri-
mental effects of micro-organisms on the process and the environmental impact associated to
the use of biocides as microbiological control products. To achieve this target, new efficient
and environmentally friendly products are required. In this paper, seventeen non-specific,
commercial enzymatic mixtures were tested to assess their efficacy for biofilm prevention and
control at laboratory and pilot plant scale. Pectin methylesterase, an enzyme found in the for-
mulation of two of the mixtures tested, was identified as an active compound able to reduce
biofilm formation by 71% compared to control tests.

Keywords: Enzymes, Biofilm, Microbiological Control, Pectin methylesterase, Paper
industry.

Introduction
The formation of biofilms can cause industrial, environmental and health problems often
resulting in operation failure and in a loss of product quality. Process waters in paper mills
present high nutrient concentrations and temperatures from of 25 to 45 ºC, being therefore an
appropriate medium for high microbial growth. In the production of recycled paper most of the
microorganisms are introduced in the system with the raw materials although additives and
water may be also contaminated. In process water microbes find ideal living conditions, which
inevitably lead to massive microbial growth. Volatile compounds produced by microbes (e.g.
organic acids, sulphur compounds and amine compounds) and the development of biofilms
can cause bad odours and/or unacceptable quality defects. Moreover, these problems are
worsened by the increased use of recovered paper and by the closure of the water circuits, both
leading to higher organic loads and higher circuit water temperatures (Blanco et al. 1996;
Blanco 2003; Lahthinen et al. 2006; Rättö et al. 2006).
In general terms a biofilm can be conceived as a structured community of bacteria enclosed in a self-produced polymeric matrix and adhered on an inert or living surface (Van Houdt et al. 2005). Established biofilms can tolerate antimicrobial agents at concentrations of 10-100 times higher than those needed to kill genetically equivalent planktonic bacteria, making them extremely difficult to eradicate (Bardouniotis et al. 2003; Burmolle et al. 2006; Jefferson 2004).

Paper manufacturers have traditionally used biocides such as chlorine, bromine, isothiazolones, glutaraldehyde and others at different points of the process for various purposes, for example to preserve raw materials and filler slurries, to prevent the formation of biofilm deposits and bad odours or to avoid the corrosion of machine components (Torres et al. 2008). Some micro-organisms, however, can develop resistance to these biocides, which justifies the development of new microbiological control aids. In addition, biocides are not always able to penetrate biofilms and to remove them from the surfaces. Furthermore, most of biocides traditionally employed are hazardous substances, whose use is regulated by legal requirements (Directive 98/8 EC; REACH 2006), being a potential source of pollution problems in effluents and in the environment (Blanco et al. 1996; Blanco 2003; Johnsrud 1997; Schenker 1996; Van Haute 1999; Bott 1998).

Current strategies to prevent biofouling should be environmentally friendly and should impede the contact between organisms and surfaces and/or prevent colonizing organisms from building up to problematic levels. One of those strategies is based in the use of dispersants. Dispersants may be applied in process waters as a single agent to prevent the formation of biofilm or they may be complemented with biocide treatments of microbiologically sensitive raw materials (Johnsrud 1997; Schenker 1996). Dispersants do not always kill or inhibit the growth of microorganisms, although some present selective inhibition of biofilm-forming bacteria (Johnsrud 1997). Because of this, neither the determination of their optimal dosage nor the
evaluation of their effect should be based only on cell counts. Dispersants may affect paper chemistry by decreasing resistance to water penetration, increasing moisture retention, or by favouring foaming. Dispersant-based biofilm control systems can also attack other non-biofilm deposits and, therefore, their effects on the whole papermaking process should be considered when they are to be applied.

Another alternative to biocides is the use of enzymatic treatments. Enzymes may affect colonization and adhesion of microorganisms in four different ways. Firstly, they may attack the adhesive of settling organisms, thus preventing the settlement event (Oulahal et al. 2007). Secondly, enzymes may degrade the polymers in the biofilm matrix formed by proliferating, settled organisms. Thirdly, enzymes may catalyse the release of antifouling compounds from the surface. These compounds may be non-toxic or toxic, but they can be much less stable than conventional biocides, what should prevent the problem of bioaccumulation of harmful chemicals. Finally, the intercellular communication during colonization of a surface may be hindered by specific enzymes (Kristensen et al. 2008).

There are two main approaches to develop effective enzymatic products for paper industry application: (1) identifying the polysaccharides present in the biofilm and looking for specific enzymes able to degrade those (Verhoef et al. 2005) and (2) identifying active compounds in different enzymatic products and assessing their effects on biofilm. In this latter approach, the specificity in the enzymes mode action makes it a complex technique, increasing the difficulty of identifying enzymes that are effective against all the different types of biofilms. Therefore, formulations containing several different enzymes seem to be fundamental for a successful biofilm control strategy (Simoes et al. 2010). This second alternative has not yet been considered in published studies and there are only a few works about the effects of specifically designed enzymes on biofilms in papermaking. These have been carried out with a levan
hydrolase and a family of products called Darazyme, developed by Grace Dearborn’s group and were based in the preliminary identification of biofilm components and the subsequent application of an enzyme or a combination of enzymes specifically selected to degrade the identified polysaccharides (Bajpai 1999).

This paper focuses on the second approach and presents a comparative study of the biofilm-preventing and degrading ability of seventeen commercial non-specific enzymatic products on the biofilm formed by isolated bacteria from paper mills.

**Materials and methods**

*Enzymes*

17 commercial enzymatic products not specifically developed for biofilm treatments (from Novozymes® A/S, Krogshoejvej 36, DK-2880 Bagsvaerd, Denmark) were evaluated to assess their effects on biofilm. Table 1 shows the main activities of the different products.

*Continuous flow systems*

The effect of enzymes on biofilm formation was studied using two continuous–flow bioreactors, whose volumes were 300 mL (B300 mL) and 10 L (B10 L). They were designed to allow biofilm growth on coupons, being a modification of the lab scale Pedersen device developed in 1982 (Pedersen 1982). The inflow of the chamber was connected by silicon tubes (B300 mL) or PVC tubes (B10 L) to a peristaltic pump and to a process water tank, and the outflow was recirculated to the tank (Fig1). Each flow chamber contained 10 PVC coupons with spaces of 5 mm between them. The total volumes of liquid in the circuits were 300 mL and 10 L respectively. The centrifugal pump was set up to move this volume of liquid with a flow of 0.29 L/s. Fresh process waters were pumped from the feeding tank to the stirred tank continuously (50 mL/ day for B300 ml and 250 ml/day for B10 L). The total volume of
process water was kept constant through the removal of exceeding process waters from the stirred tank. The flow through the coupon chambers was 0.10 L/s to assure turbulent flow. Since the hydrodynamic conditions affect notably the biofilm build-up and it is very difficult to reproduce the hydrodynamic conditions of the mill at laboratory scale, the effect of the enzymes on biofilm build-up was studied at constant and turbulent hydrodynamic conditions, which allows comparing the results obtained and therefore, establishing the effect of each treatment on the biofilm build-up by comparing it with the control sample.

Experimental procedure

*Preliminary laboratory trials in B300 ml.*

Before starting the experiment, the bioreactors were cleaned with detergent and ethanol and flushed with sterilized distilled water. The coupons were degreased and sterilized with acetone and ethanol and thereafter washed with sterilized distilled water and placed in the flow chamber.

The bioreactors were inoculated with the flora present in the process water obtained from the sheet forming zone of the wire section of a 100% recycling paper mill, which uses mixed recovered paper as raw material and produces paper for board (Mill 1). The experiments were carried out at a controlled pH of 6.8-7.0 and at 30ºC. Seventeen enzymatic compounds were studied. The dosage of the commercial enzyme preparations was 1000 ppm for all the tests. The experiments with B300 mL were carried out over 4 days and the flow chamber was opened every 24 h to remove two coupons. One coupon was used to quantify biofilm formation, based on its dry weight (mg/cm²) after 6 h at 105ºC, and the other one was used to measure the colony forming units per square centimetre of coupon (CFU/cm²). For this latter measurement, coupons were removed with sterile forceps and flushed with 9 mL sterile physiological solution to remove slightly adhered cells and the excess of aqueous medium.
Biofilms were swabbed from one side of each coupon with sterile cotton wool sticks and transferred to a vial containing a known volume of sterile physiological solution. This suspension was vortexed (20 s) to disperse cells, then, with each obtained sample, a series of dilutions was performed and plated on plate count agar. Plates were incubated at 30ºC for 48 h. Then, the CFU/cm² was determined. The aqueous medium of the bioreactor was aseptically sampled at similar intervals and the number of colony forming units per millilitre (CFU/ml) was determined accordingly. The results of these preliminary trials allowed selecting three treatments for further studies in B10 L.

**Laboratory trials in B10 L.**

The experiments were carried out over 4 days. Water samples and coupons were taken every 24 h and biofilm formation on coupons was determined by counting and weighting, following the procedure explained before for the trials in B300 mL. Process waters from Mill 1 were used for the experiments. This water, rich in fibres, fines and filler was sampled from sheet-forming zone of the wire section and stored at 15ºC. Before being used in B10 mL, process waters were sequentially filtered through 1 mm, 600 µm, 400 µm, 200 µm and 63 µm filters, which reproduce the disc filter treatment employed in the mill to obtain clarified waters.

The colonisation ratio (CR) was calculated from the ratio between the total number of CFU on the coupon (A) and the total number of CFU in contact with a coupon (C) as indicates by the equation (1). The latter (C) was defined as the sum of the total number of CFU in the medium (B) and the total number of CFU on the coupon (A), equation (2). A and B are defined by equations (3) and (4), where CS is the coupon surface (cm²) and TVL is the total volume of liquid in the reactor (mL). In this way, comparison was possible between experiments with different starting conditions (Kielemoes et al. 2002).
Finally, only one commercial enzymatic product was selected for further tests. These were first carried out in the B10 L over 10 days. To ensure that the bacterial population was maintained at a constant level throughout the experiment, a count of microorganisms in the medium was carried out each 24 h observing that the bacterial concentration in process water was maintained between $10^6$ and $10^7$ CFU/mL. Biofilm build-up was assessed every 24 h, through the measurement of the dry weight of biofilm deposited on the coupons.

**Pilot circuit trials**

To identify the active compound having the predominant role against biofilm, two commercial products, containing fractions of the most active product selected from the seventeen commercial products were also studied. This set of experiments was carried out in a pilot circuit simulating the process water circuit in a paper mill that considers fresh water addition, and the removal of exceeding waters.

The pilot circuit consisted of 2 bioreactors BioFlo3000®, manufactured by New Brunswick Scientific (USA) one used to test the effect of the treatments, and the other used as control, allowing bacterial growth under a controlled environment: temperature (45°C), pH (7), dissolved oxygen (up to 20%) and turbulent regime (Reynolds number $> 2000$). High density polypropylene coupons were immersed in each bioreactor to allow the formation of deposits.
Process waters from a 100% recycled paper mill producing writing and printing papers were used for the experiments (Mill 2). Again, this water was sampled from the sheet-forming zone of the wire section and stored at 15°C. Before being used in the pilot circuit, process waters were successively filtered by following the same procedure as explained for trials in B10 L.

All experiments were carried out by duplicate. To test the ability of the enzymatic treatment to prevent the biofilm formation, enzyme was continuously added during 10 days from the beginning until the end of the experiment to keep a constant concentration of enzyme of 1000 ppm. Each day, the total colony number of aerobic bacteria in the waters was assessed by the dry-rehydratable film method (Petrifilm3M™). Counting results are expressed as the number of colony forming units (CFU/mL). At the end of the experiment, the total amount of dry weight of biofilm formed on the coupons was measured.

To exclude the effect of non-enzyme components in the commercial product, a test with denatured enzyme was carried out with process water from Mill 2 with a clean circuit. Enzyme inactivation was achieved by heat (121°C, 30 min.). The inactive enzyme was added continuously during the test to keep constant a concentration 1000 ppm and the effects were compared to the biofilm production in a control experiment without enzyme.

Experiments with different doses of the best product were also performed in clean circuits. The best treatment was applied continuously in three bioreactors fed with process waters from Mill 2. Three different concentrations were tested: 100 ppm, 10 ppm and 1 ppm. A fourth bioreactor without treatment was used as control.

Results

Preliminary laboratory trials
The qualitative effectiveness of the treatments for preventing the formation of biofilm is presented in Table 1. From the group of the seventeen treatments assessed, Pectinex Smash® was the most effective in preventing the adhesion of viable colony forming units on the coupons and the biofilm formation. Viscozyme L®, Pulpzyme®, Terminox ® and Bio-feed Beta L® showed a moderate effect. The treatment with Novozyme 863® had no hindering effect on the formation of biofilm, although a significant low number of CFU was measured on the coupon when this product was tested, which was unexpected. Therefore, Novozymes 863 ®, Pectinex Smash® and Viscozyme L®, were chosen for further investigations.

**Laboratory trials with B10 L.**

The results obtained from the test of the selected enzymatic products with B10 L are shown in Fig2. The graphs show the number of colony forming units (CFU/mL) in the medium, the number of viable counts per surface unit of the coupon (CFU/cm²), the dry weight biofilm on the of coupon and the colonization ratio. Biofilm formation was evaluated by an increased dry weight of the biofilm deposited on the coupon and by the colonization ratio. The dry weight increase is caused by bacterial colonization and through the production of EPS. An increasing colonization ratio indicates the formation of biofilm.

The addition of the tested enzymes did not change the value of the CFU/mL in the medium over the time, which remained constant in all cases and similar to the value measured in the control sample. This indicates that the tested enzymatic preparations did not have an undesired biocide effect.

When no treatment was added (control trial), both, the number of CFU on the coupon and the dry weight of material on the coupon increased, whereas the number of CFU in the medium stayed mainly constant, which indicates that bacteria were actively colonizing the surface and
producing biofilm. In fact, the biofilm was already formed after 24 h and increased during the course of the experiment. The colonization ratio increased rapidly after 48 h and approached the value of 100% after 96 h, which implies that nearly all bacteria present in the coupon biofilm were viable.

When an enzymatic treatment was used, the number of viable CFU/cm² of coupon increased during the first 24 h to reach a value higher than that obtained for the control sample. This unexpected effect is in accordance with the observations made by other authors that have proved that the presence of salts, disinfectants or some other compounds, or other bacteria can increase the adhesion of bacteria and even the biofilm maturation (Carpentier et al. 2004; Jensen et al. 2007). However, after the initial increase, the value of CFU/cm² remained constant (see the results for Novozyme 863® and Pectinex Smash® in Fig2) or even decreased slightly after 72 h (in the case of Viscozyme L®) As a result, the colonization ratio was kept below 20% during the trials with enzymes and the final number of bacteria colonizing or growing on the surface was lower than that for the control trial. The effect of Novozyme 863® and Viscozyme L® on the dry weight of biofilm deposited on the coupons was significantly weaker from that of the Pectinex Smash® trial.

The addition of Novozyme 863® or Viscozyme L® did not prevent the build-up of organic material on the coupon, although they limited it and the final values were less than half the value reached at the end of the control trial. However, the addition of Pectinex Smash® prevented the build-up of organic material on the coupon weight maintaining it below 0.2 g. Furthermore, the colonization ratio was kept low during the whole trial, which implies that the treatment reduced the biofilm on the coupon surface both by limiting the number of bacteria colonizing the surface and by reducing the formation of biomass on the coupon. Therefore, Pectinex Smash® treatment was chosen to continue the study with pilot circuit trials.
**Pilot circuit trials**

During the pilot circuit trials, the bacterial concentration in the water remained between $2.3 \times 10^6$ and $7 \times 10^7$ in all the cases. When no treatment was applied, the biofilm formed after 72 h of trial was evident by visual observation of the coupons surface. The dry weight of the biofilm deposited on the coupons in the bioreactor with addition of Pectinex Smash® was between 4 and 5 times lower than that in the bioreactor without enzymatic treatment (Fig3). Therefore, the addition of Pectinex Smash® in a clean circuit limited dramatically the production of biofilm on the polypropylene coupons placed in the bioreactor under turbulent conditions.

**Test with denatured Pectinex Smash®**

During the experiments with the inactivate Pectinex Smash® solution, the total bacterial count in the waters did not vary significantly ($10^7$ - $10^8$ CFU/mL). No significant differences were observed between the dry weight of biofilm deposited on the coupons in the bioreactor with inactive Pectinex Smash® and that in the bioreactor without any treatment (all the measured dry weights obtained on the coupons after 260 h were in the interval $1.4 \pm 0.3$ mg/cm²). These results suggest that the biofilm prevention properties of the Pectinex Smash® solution were not due to non-enzymatic components.

**Effect of Pectinex Smash® dosification**

During the test performed during 190 h, no significant variation in the total bacterial count was observed in the 4 bioreactors. After 24 h, biofilm formation was evident by visual observation of the coupons in the bioreactor without treatment and in that with Pectinex Smash® treatment at 1 ppm. In the bioreactor with 10 ppm and 100 ppm of enzyme, biofilm on the coupons was visually detected only after 48 h. The dry weight of slime on the coupons from the bioreactor
without treatment or with 1 ppm of Pectinex Smash® was very similar, which indicates that a concentration of 1 ppm of Pectinex Smash®, did not affect biofilm production (Fig4). A moderate effect against biofilm build-up was observed with the treatment at 10 ppm concentration, which decreased the weight of slime on the coupons up to 50%. However, added at 100 ppm concentration, Pectinex Smash® hindered dramatically the production of biofilm. Since the weight of biofilm formed on the coupons was 5 times lower with 100 ppm of Pectinex Smash® than that without the enzyme. Therefore, the efficacy of using 100 ppm of this enzyme was comparable by using 1000 ppm of the same enzyme.

Identification of main enzymatic component
Since Pectinex Smash® showed the best performance in the experiments described in the previous assays more tests were carried out to further verify its effect against biofilm formed from the non filtrated water, containing fibers, fines and fillers that can affect the biofilm formation.

Pectinex Smash® is a product obtained from *Aspergillus aculeatus* that presents a wide range of enzymatic activities as various pectinolytic activities between them, a pectin-methylestearase. To identify the active compound having the predominant role against biofilm, two related products, containing fractions of activities of Pectinex Smash® were also tested in the B10 L: Pectinex Ultra SP® (a mixture of mainly pectinolytic enzymes from *Aspergillus aculeatus*) and Novoshape®, (a pectin methylesterase of *Aspergillus aculeatus* as main active enzyme). Fig5 shows the effect achieved with those enzymatic products. Pectinex Ultra SP® had no effect on the formation of biofilm (Fig5), whereas the Pectinex Smash® treatment was effective to control biofilm formation as it shows the weight of dry biofilm on the coupons that was 3 times lower than that in the control experiment. Finally, the results obtained with Novoshape® showed that the dry weight of biofilm on the coupons was 4 times lower than that
for the control experiment. Both Pectinex Smash® and Novoshape® preparations, include a *pectin methylesterase*. It can therefore be concluded that pectin methylesterase was the main enzyme or, at least one of the enzymes, responsible for the efficiency of Pectinex Smash® on the prevention and control biofilm.

**Discussion**

The obtained results showed that among the 19 products studied, Pectinex Smash® and its fraction Novoshape®, were the best treatments to prevent biofilm formation, both have in common that are composed principally by the pectin methylestearase. Pectinex Smash® a mix of pectynolitic activities and Novoshape ® is a liquid enzyme preparation of a microbial pectin methylesterase (PME, E.C.3.1.1.11). The gene encoding the esterase enzyme is derived from the fungus *Aspergillus aculeatus*, and is transferred into a strain of the food-grade organism *Aspergillus oryzae* for commercial production. The NovoShape preparation has a declared activity of 10 PEU/mL and a temperature optimum of \( \sim 50 ^\circ C \) (data provided by the manufacturer). This enzyme belong to carbohydrate esterase family and catalyses the hydrolysis of methyl ester groups and has high specificity for pectin substrates, property that has been widely used in food industry and in plant science (Suutarinen 2002, Micheli 2001,Femenia 1998).

Composition of the biofilm polysaccharides is still insufficiently known, but available data on planktonic EPS and few biofilm EPS, suggest that some of their monomers are identical or similar to those in plant cell-wall materials (Sutherland 2005). In fact, some bacterial exopolysaccharides have been reported to be acceptable substrates for enzyme mixtures from non-bacterial sources (Sutherland 2001).
In the study performed by Orgaz on biofilm remove of *P. fluorescens* was concluded that the enzyme pectin esterase produced by *Trichoderma viride* (belongin to the same family that pectinese methylestearase), it could possibly deacetylate a polysaccharide in the biofilm matrix, making it softer and possibly more porous (Orgaz et al. 2006; Orgaz et al. 2007). Many microbial exopolysaccharides have different substituent groups as ketal-linked pyruvate or ester-linked acetyl groups. The removal of acyl groups, especially the acetate, can greatly affect the physical properties of polysaccharides. The removal of one of the substituents such as acetyl group, especially the acetate influences the physical properties of exopolysaccharides (Sutherland 1999).

Many fungi can degrade complex plant cell-wall material, by secreting a large variety of enzymes. This versatility makes commercial polysaccharides-degrading enzyme mixtures to have a widespread use in very different fields, such as fruit processing (McKay 1993) or wastewater treatment (Wesenberg 2003). They could possibly be used as well to degrade bacterial biofilm matrices, can be effective to prevent and control the formation of biofilms in the process water circuits of paper mills, a mixture of enzymes may be necessary for sufficient biofilm degradation due to the EPS heterogeneity.

This research opens the way to provide a method of removing biofilms from surfaces submerged in water using an effective enzyme and that could be combined with the use of a biocide. The use of an enzyme which is biodegradable and has a low toxicity is also environmentally acceptable and economically appealing because of its contribution to the minimization of the use of biocides.

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References


## Table 1 Commercial enzymatic products

<table>
<thead>
<tr>
<th>Product</th>
<th>Main enzymatic components</th>
<th>Effect on CFU/coupon</th>
<th>Prevention of biofilm formation</th>
</tr>
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<tr>
<td>Bio-feed Beta L®</td>
<td>Alpha-amylase, beta-glucanase, cellulase, cellulase, hemicellulase, xylanase</td>
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<td>Cerefix 200 L®</td>
<td>Beta-glucanase</td>
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<td>Denilite®</td>
<td>Laccase</td>
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<td>Dextranase 50 L®</td>
<td>Dextranase</td>
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<td>~</td>
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<tr>
<td>Energen W®</td>
<td>Xylanase, beta-glucanase, hemicellulase, pectinolytic activities</td>
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<td>Fructozyme L®</td>
<td>Inulinase</td>
<td>~</td>
<td>~</td>
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<tr>
<td>Gammanase®</td>
<td>Gammanase</td>
<td>~</td>
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<tr>
<td>Glucanex®</td>
<td>Beta-glucanase</td>
<td>~</td>
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<td>Gluzyne 10,000 BG®</td>
<td>Catalase, glucose oxidase</td>
<td>~</td>
<td>~</td>
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<tr>
<td>Kojiyze®</td>
<td>Endo-peptidase, exo-peptidase, aminopeptidase, carboxypeptidase, amylase</td>
<td>~</td>
<td>+</td>
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<tr>
<td>Novozyme 863®</td>
<td>Polymalictraminase, pectolytic and hemicellulolytic activities</td>
<td>++</td>
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<tr>
<td>Pectinex Smash®</td>
<td>Various pectinolytic activities</td>
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<td>++</td>
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~ = no effect; + = slight effect; ++ = moderate effect; +++ = strong effect
Fig. 1 Continuous-Flow Systems: Stirred tank (A), flow chamber with coupons (B), centrifugal pump (C), feeding tank (D), isothermic bath (E) and peristaltic pump (P).

Fig 2 Effects of selected enzymatic treatments on biofilm formation and biofilm build-up in the 10L bioreactor.
**Fig3** Effect of Pectinex Smash® on biofilm prevention

**Fig4** Dose response of Pectinex Smash® on biofilm prevention
**Fig 5** Effect of Pectinex Samsh, Pectinex Ultra SP and Novoshape on biofilm prevention