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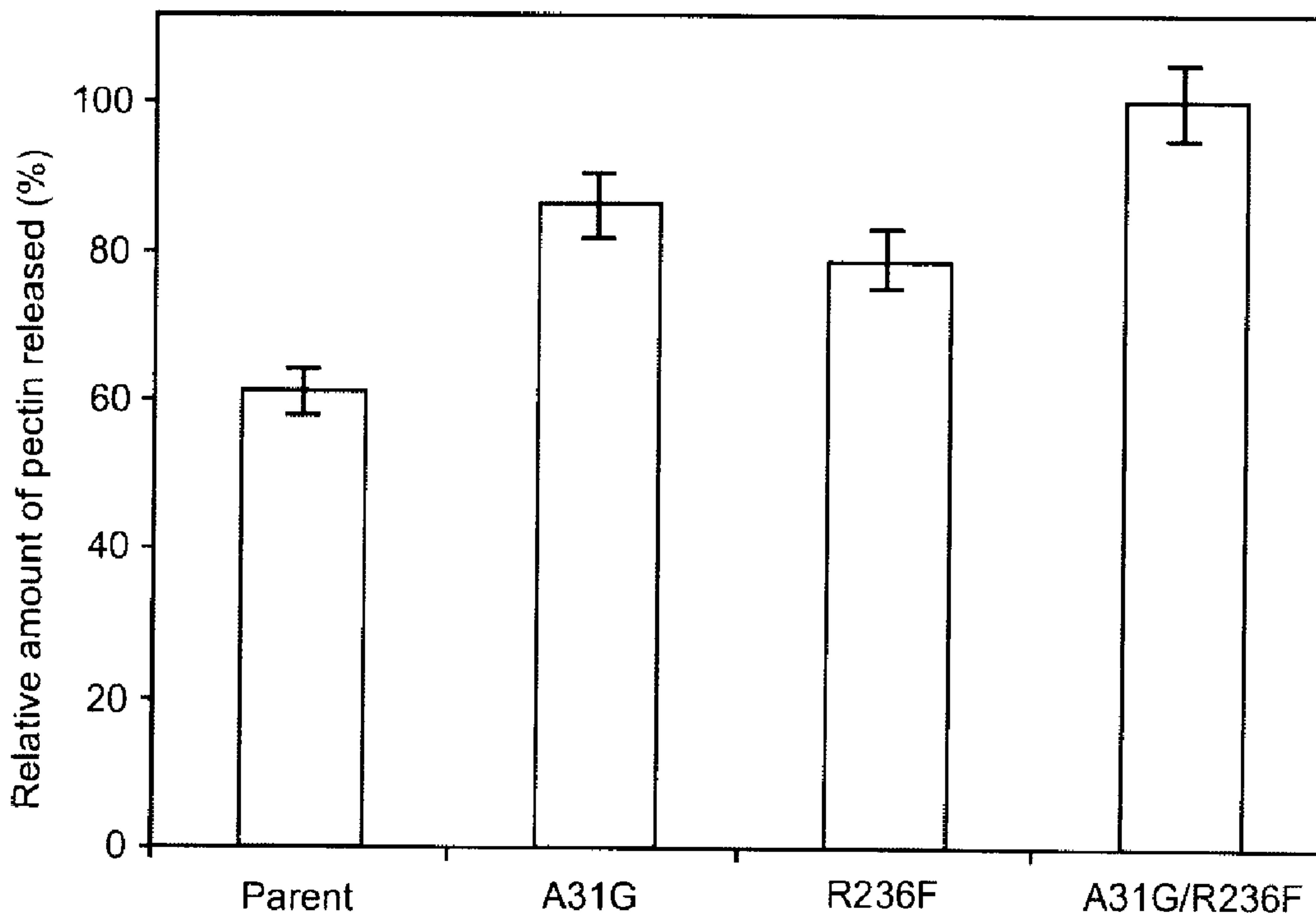


Fig. 6

(57) Abrégé/Abstract:

Using site-directed mutagenesis to mutate the *Xanthomonas campestris* pectate lyase gene, variants of *Xanthomonas campestris* pectate lyase with improved thermostability and/or enzymatic activity have been expressed in *Escherichia coli*, and then isolated

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(57) Abrégé(suite)/Abstract(continued):

and purified. The mutant *Xanthomonas campestris* pectate lyases are more effective than the wild-type enzyme, also expressed in *E. coli*, in removing pectic compounds from natural hemp fiber.

ABSTRACT

5 Using site-directed mutagenesis to mutate the *Xanthomonas campestris* pectate lyase gene, variants of *Xanthomonas campestris* pectate lyase with improved thermostability and/or enzymatic activity have been expressed in *Escherichia coli*, and then isolated and purified. The mutant *Xanthomonas campestris* pectate lyases are more effective than the wild-type enzyme, also expressed in *E. coli*, in removing pectic compounds from natural hemp fiber.

PECTATE LYASES WITH INCREASED THERMOSTABILITY AND/OR ENZYMATIC
ACTIVITY

Cross-reference to Related Applications

5 This application claims the benefit of United States Provisional Patent Application
USSN 60/924,386 filed May 11, 2007, the entire contents of which is herein incorporated
by reference.

Field of the Invention

10 The present invention is directed to pectate lyases, in particular to genetically
modified pectate lyases having increased thermostability and/or enzymatic activity. Such
pectate lyases are useful in detergents and in bioscouring.

Background of the Invention

15 Extraction of fiber from fiber plants would allow its eventual usage. Extraction
primarily involves removal of non-cellulosic material, e.g. pectin and colour-containing
materials, from the fiber. This removal of non-cellulosic material is sometimes referred to
as degumming. Pectin is a polysaccharide which is a polymer of galacturonic acid.
Pectin is not soluble in water or acid. However, it can be removed by strong alkaline
solutions like caustic soda (concentrated sodium hydroxide).

20 General methods for isolation of clean fibers include dew retting, water retting,
and chemical and enzymatic processes, with different variations. In these methods, the
glue that holds the fibers together must first be loosened (or removed altogether) by
retting. In conventional retting, stalks are dew-retted by allowing them to lie in the field
after cutting. These retting approaches depend on digestion of pectin by enzymes
secreted by microbes thriving under favorable conditions. Although water-retting yields
25 more uniform fiber, the process pollutes the water. Dew-retting requires anywhere from
two to six weeks or more to complete, requiring the stalks to be turned at least once for
highest-quality fiber. Dew-retting is thus affected by the weather, which offers no guaranty
of favorable condition.

30 On an industrial scale, chemical scouring is common. Chemical scouring is a
process that improves the water absorbancy and the whiteness of textile by removing the
non-cellulosic substances from cotton, flax or other natural cellulosic fibers, thereby
facilitating the subsequent dyeing and finishing processes (Sawada and Ueda. 2001;
Solbak, et al. 2005; Tzanov et al., 2001). Violent, hazardous chemicals like soda ash,

caustic soda and oxalic acid used in the current scouring processes cause severe environmental problems, besides a loss of fiber strength.

5 In the past several years, considerable attention has been paid to replacing traditional processes, e.g. chemical scouring of raw cotton and flax fabrics and retting of flax and hemp fibers, with more environmentally friendly and economically viable biotechnological processes. Enzyme retting or bioscouring involves the action of a pectinase, especially pectate lyase, with or without other enzymes like xylanase and/or cellulase. Such processes are generally known (Akin et al., 2001; Klug-Santner et al., 2006; Ossola and Galante. 2004; Ouajai and Shanks. 2005; Sawada and Ueda. 2001; 10 Solbak, et al. 2005; Tzanov et al., 2001). Pectate lyases specifically remove non-cellulosic material without damaging the cellulose backbone of the fiber. Further, enzymes used for bioscouring reduce environmental pollution (Solbak, et al. 2005). Pectate lyases widely exist in bacteria (Berensmeier et al. 2004; Klug-Santner, et al. 2006; Kluskens, et al. 2003) and fungi (Benen et al. 2000; Huertas-Gonzalez et al. 1999).

15 Other examples of prior art include United States Patent 6,124,127, WO 2001/079440, WO 2000/042152 and WO 2006/0089283. These documents disclose various pectate lyases for use in detergents or textile processing.

20 Despite the interest in pectate lyases for use in bioscouring processes, many pectate lyases currently in use suffer from a lack of scouring efficiency. Lack of scouring efficiency can arise from thermal instability and/or low enzymatic activity of the pectate lyase. Thermal instability results in reduction of activity over time due to thermally induced changes in enzyme conformation. Higher temperatures accelerate reduction in activity. Low enzymatic activity limits the rate at which scouring can occur. Scouring efficiency can be increased by using pectate lyases having increased thermostability and/or enzymatic activity. 25

Thus, there is a need for enzymes, particularly pectate lyases, having increased thermostability and/or enzymatic activity.

Summary of the Invention

30 It has now been found that certain variants of a *Xanthomonas campestris* pectate lyase have improved thermostability, enzymatic activity or both thermostability and enzymatic activity.

Thus, there is provided a polypeptide comprising the amino acid sequence (SEQ ID NO: 2):

MTSKTLQGALALALSACAAGAIAGPVGYGAX₁TTGGGNKVPVNVATFEAMQSAIDSYSGSGGLVLN
 YTGKDFDGTIKDVCAQWKLPAKTVQIKNKSDVTIKGANGSAANFGIRVVGNAHNVI IQNMTIGLLQ
 5 GGEDADSI SLEGNSSGEP SKIWDHNTVFASLT KC SGAGDASFDGGIDMKKGVHHVTVSYNYVYNY
 QKVALNGYSDSDTKNSAARTTYHHNRFENVESRVPLQRX₂GLSHIYNNYFNNVTTSGINVRMGGIA
 KIESNYFENIKNPVTSRDSSEIGYWDLINNYVGSGITWGT PDGSKPYANATNWISTKVFPE SLGYI
 YTVTPAAQVKAKVIATAGAGKNLAE

wherein X₁ is alanine or glycine and X₂ is arginine or phenylalanine with the proviso that
 10 X₂ is not arginine when X₁ is alanine.

There is also provided a process of isolating fibers from non-cellulosic material in a fiber plant comprising contacting raw fibers of the fiber plant with a polypeptide of the present invention.

Polypeptides of the present invention are enzymes and are variants of
 15 *Xanthomonas campestris* pectate lyase II (XcPL). Individual variants include polypeptides having an amino acid sequence of SEQ ID NO: 3, SEQ ID NO: 4 or SEQ ID NO: 5.

Enzymes of the present invention are stable in a pH range of 4-10, and function optimally at a pH of about 8.5. They advantageously possess improved thermostability,
 20 enzymatic activity or both thermostability and enzymatic activity, in comparison to the wild-type *X. campestris* pectate lyase. Thermostability may be at least two times greater, for example at least ten times greater, and in some embodiments at least twelve times greater and in other embodiments at least twenty three times greater, than that of the wild-type enzyme. Enzymatic activity may be at least two times greater, and in some
 25 embodiments at least five times greater, than that of the wild-type enzyme. Half-life of enzymatic activity at 45°C for enzymes of the present invention may be 3 hours or more, or even 10 hours or more.

Enzymes of the present invention may be used in detergents for degrading non-cellulosic material, for degrading non-cellulosic material in textile processing, for
 30 degrading non-cellulosic material in cellulose fiber processing, for degrading non-cellulosic material in the treatment of pectic wastewaters, for degrading non-cellulosic material in paper making, for degrading non-cellulosic material in coffee and tea fermentations and for improving the yield and the clarification of fruit juices. They are

particularly useful as a detergent for the treatment of cellulosic material, especially cellulose-containing fibers, yarn, woven or non-woven fabric, and treatment of mechanical paper-making pulps or recycled waste paper, and as a bioscouring agent for retting of fibers. The enzymes of the invention are very effective for use in an enzymatic scouring process in the preparation of cellulosic material e.g. for proper response in subsequent dyeing operations. Further, it is contemplated that detergent compositions comprising the novel enzymes are capable of removing or bleaching stains present on laundry, especially stains resulting from galactan or arabinogalactan containing food, plants, and the like. Treatment with detergent compositions comprising an enzyme of the present invention can prevent binding of certain stains to the cellulosic material.

Bioscouring (enzymatic retting) involves enzymatic removal of non-cellulosic material, especially pectin, from raw plant fibers, for example hemp, jute, flax, ramie and cotton. In accordance with the present invention, enzymatic treatment of raw fibers employs one or more enzymes of the present invention. One or more other enzymes may also be used, for example, proteases, cellulases (endoglucanases), beta-glucanases, hemicellulases, lipases, peroxidases, laccases, alpha-amylases, glucoamylases, cutinases, pectinases, reductases, oxidases, phenoloxidases, ligninases, pullulanases, arabinosidases, mannanases, xyloglucanases, xylanases, pectin acetyl esterases, polygalacturonases, rhamnogalacturonases, galactanases, pectin lyases, other pectate lyases, polygalacturonases, pectin methylesterases, cellobiohydrolases, transglutaminases or mixtures thereof.

Preferably, enzymatic treatment is performed in an aqueous medium at a pH of from about 4-10. More preferably, the pH is from about 8-9. Preferably, the temperature at which enzymatic treatment is performed is in a range of from about 30°C to 55°C, for example in a range of from about 40°C to 50°C. Preferably, the aqueous medium contains salts and/or buffers, for example monosodium citrate. Concentration of any salts or buffers should not be too high as to unduly affect activity of the enzyme. For example, the concentration of monosodium citrate may be in a range of about 3-7 mM, e.g. 5 mM.

Preferably, enzymatic treatment of the fibers is performed for a period of time in a range of from about 0.5-10 hours, for example about 1-6 hours. In some embodiments, 1-3 hour of treatment is sufficient. Stirring or agitation of the aqueous medium may be done. Preferably, the aqueous medium is stirred or agitated constantly during enzymatic treatment. Purified fiber after enzymatic treatment may be rinsed with water.

Specific conditions under which the enzymatic treatment is conducted may depend on various factors, for example the nature and amount of raw fiber being treated, and the use to which the fiber is ultimately destined.

5 In common fiber plants a bark-like layer containing bast fibers surrounds a woody core. Before enzymatic treatment can be performed on the bast fibers, it may be necessary to remove the woody core. Decortication, either manually or mechanically, is a process that separates the bark-like layer from the woody core.

10 Enzymatic treatment of raw fibers may be used in conjunction with other treatment steps, for example chemical scouring. Purified fiber may be subjected to other treatment steps, for example bleaching, dyeing, etc., for its eventual application. Enzymatic treatment advantageously reduces the amount of violent and hazardous chemicals required in the other treatment steps.

Further features of the invention will be described or will become apparent in the course of the following detailed description.

15 Brief Description of the Drawings

In order that the invention may be more clearly understood, embodiments thereof will now be described in detail by way of example, with reference to the accompanying drawings, in which:

20 Fig. 1 depicts multiple alignment of amino acid sequences of thermostable pectate lyases from *Bacillus licheniformis*, *Bacillus sp. TS-47* and *Thermotoga maritima* and a thermolabile pectate lyase from *Xanthomonas campestris* (*X. campestris* pectate lyase II);

Fig. 2 depicts amino acid sequences of parent *Xanthomonas campestris* pectate lyase II and three of its variants (mutants A31G, R236F and A31G/R236F);

25 Fig. 3 depicts graphs illustrating thermostability of parent *Xanthomonas campestris* pectate lyase II and its variants, including mutants A31G, R236F and A31G/R236F;

30 Fig. 4 depicts (left) SDS-PAGE analysis (Coomassie blue stained 10% SDS-PAGE) and (right) a zymogram (SDS-PAGE containing 0.1% PGA stained with ruthenium red) of purified expressed proteins of parent *Xanthomonas campestris* pectate lyase II

and its mutants, in which Lane M = Molecular standard, Lane 1 = parent pectate lyase, Lane 2 = mutant A31G, Lane 3 = mutant R236F, and Lane 4 = mutant A31G/R236F;

Fig. 5 depicts a graph of melting temperatures of parent *Xanthomonas campestris* pectate lyase II and its mutants A31G, R236F and A31G/R236F; and,

5 Fig. 6 depicts a graph of the effects of parent *Xanthomonas campestris* pectate lyase II and its mutants A31G, R236F and A31G/R236F on release of pectin from natural hemp fiber.

10 Fig. 7 depicts a graph of predicted folding free energies versus experimental stabilities of pectate lyase mutants relative to the wild-type protein. Predictions are based on the FOLD-X program using the crystal structure of the wild-type pectate lyase. Experimental stability data is expressed as fold-change in the half-time of inactivation upon mutation converted to logarithmic scale. Arrows indicate larger experimental changes than those plotted in the case of Y66V and V187I mutations.

Description of Preferred Embodiments

15 The term "wild-type enzyme" denotes an enzyme, which is endogenous to a naturally occurring microorganism found in nature.

The term "parent enzyme" means an enzyme in which modifications are being made to produce enzyme variants. A parent enzyme may be an enzyme isolated from a natural source, or an enzyme wherein previous modification(s) have been made while
20 retaining the characteristic activity of the enzyme in question. The parent pectate lyase may be wild-type pectate lyase.

Materials and Methods

Multiple sequence alignment. Three thermostable pectate lyases: CAD56882 from *Bacillus licheniformis* (Berensmeier et al. 2004), BAB40336 from *Bacillus* sp. TS-47
25 (Takao et al. 2001), AAD35518 from *Thermotoga maritima* MSB8 (Kluszens, et al. 2003) with optimal temperatures of 65°C, 70°C and 90°C, respectively and *Xanthomonas campestris* pectate lyase II (NP_638163) (da Silva et al. 2002), a mesophilic protein with optimal temperature of 50°C established in this study, were analyzed by multiple sequence alignment using the European Molecular Biology Laboratory – European
30 Bioinformatics Institute (EMBL-EBI) ClustalW program. Protein sequences in this

alignment are indicated by their accession numbers in the NCBI protein database. Not shown are the putative signal peptide sequences of the respective proteins.

5 *Site-directed mutagenesis.* Site-directed mutagenesis was performed by using the QuikChange[®] Mutagenesis kit (Stratagene, La Jolla, CA) according to its instruction manual. The mutagenic oligonucleotide primers are listed in Table 1. The expected mutations were confirmed by gene sequencing using a Big Dye DNA sequencing kit (Applied Biosystems) with an automated DNA sequencer (Model 377, ABI Prism).

10 *Molecular cloning techniques.* Standard methods were used for isolation of plasmid DNA, cloning and transformation (Sambrook et al. 1989). The *Xanthomonas campestris* pectate lyase II gene was cloned into an IPTG-inducible bacterial-expression vector, pSD80 (Smith et al. 1996), at the *EcoRI* and *HindIII* restriction sites. The recombinant plasmids containing the parent and mutated *X. campestris* pectate lyase gene fragments were separately transformed into the Rosetta pRARE strain, *Escherichia coli* BL21 (DE3) (Novagen).

15

Table 1

<u>Mutant</u>	<u>Direction</u>	<u>SEQ ID</u>	<u>Nucleotide sequence</u>
V26A	Forward	NO: 6	5'-GATCGCAGGTCCGGCCGGCTACGGTG-3'
	Reverse	NO: 7	5'-CACCGTAGCCGGCCGGACCTGCGATC-3'
A31G	Forward	NO: 8	5'-GCTACGGTGCCGGCACCACCGGCG-3'
	Reverse	NO: 9	5'-CGCCGGTGGTGCCGGCACCCTAGC-3'
L64I	Forward	NO: 10	5'-CGGCGGCCTGGTGATCAACTACACCGGCAAG-3'
	Reverse	NO: 11	5'-CTTGCCGGTGTAGTTGATCACCAGGCCGCG-3'
Y66V	Forward	NO: 12	5'-GCCTGGTGCTGAACGTCACCGGCAAGTTCGACTTC-3'
	Reverse	NO: 13	5'-GAAGTCGAACCTGCCGGTGACGTTTCAGCACCAGGC-3'
K69T	Forward	NO: 14	5'-CTGAACTACACCGGCACGTTTCGACTTCGGC-3'
	Reverse	NO: 15	5'-GCCGAAGTCGAACGTGCCGGTGTAGTTCAG-3'
F70I	Forward	NO: 16	5'-GAACTACACCGGCAAGATCGACTTCGGCACC-3'
	Reverse	NO: 17	5'-GGTGCCGAAGTCGATCTTGCCGGTGTAGTTC-3'
Q123R	Forward	NO: 18	5'-GCACAACGTGATCATCCGGAACATGACCATCGG-3'
	Reverse	NO: 19	5'-CCGATGGTCATGTTCCGGATGATCACGTTGTGC-3'
V187I	Forward	NO: 20	5'-GGCGTGCATCACATCACCGTGTCTACAAC-3'
	Reverse	NO: 21	5'-GTTGTAGGACACGGTGTGTGATGCACGCC-3'
R236F	Forward	NO: 22	5'-CCGCTGCAGCGCTTTGGCTTGAGCCACATC-3'
	Reverse	NO: 23	5'-GATGTGGCTCAAGCCAAAGCGCTGCAGCGG-3'

Protein expression. Single colonies harboring *Xanthomonas campestris* pectate lyase II gene with desired mutations were grown at 37°C on a rotary shaker of 250 rpm in 5 ml of LB medium containing 50 µg/ml ampicillin and 30 µg/ml chloramphenicol. The overnight cultures were diluted 200 fold into the same fresh LB medium, grown to OD₆₀₀ ≅ 0.4 to 0.5 under the same culture conditions, and then induced with 1mM IPTG for 3 hours. The bacterial cells were precipitated and disrupted by French Press. The clear supernatant of cell lysate was used for enzymatic assay after centrifugation at 14000 rpm for 40 min at 4°C.

Protein purification. The supernatant of cell lysate was loaded on to a HiPrep™ 16/10 SP XL column (Amersham Pharmacia). The active peak detected by pectate lyase assay (Collmer et al. 1988) was eluted with a linear NaCl gradient from 0 to 200 mM in 20 mM sodium phosphate buffer, pH 7. Protein concentration was determined by Bradford assay (Bradford 1976).

Gel electrophoresis and zymogram staining. 200 ng of protein samples were heated for 10 min at 45°C in sample loading buffer before being applied to 10% SDS-PAGE containing 0.1% polygalacturonic acid. After electrophoresis, protein bands were stained with Coomassie Blue. For zymogram staining, the gel was soaked in 2.5% Triton™ X-100 for 30 min, washed with 50 mM Tris-HCl pH 8.5 containing 0.5 mM CaCl₂ for 30 min. After 20 h of incubation at 40°C in 50 mM Tris-HCl pH 8.5 containing 0.5 mM CaCl₂ and 0.1% polygalacturonic acid, the gel was stained with 0.05% (w/v) ruthenium red for 20 min and washed with water for 60 min.

Pectate lyase assay. The pectate lyase activity was determined by measuring the increase in absorbance at 232 nm of polygalacturonic acid (PGA) (Collmer et al. 1988). 50 µl of enzyme and 50 µl of 0.5% PGA in 50 mM Tris-HCl pH 8.5 containing 0.5 mM CaCl₂ were incubated at 50°C for 30 min. The reaction was stopped with addition of 20 µl of 0.35 M HCl. 100 µl of reaction mixture was used to determine the absorbance at 232 nm. One unit (IU) of pectate lyase was defined as the enzyme amount producing 1 µmol of unsaturated product in 1 min under the assay conditions. The molar extinction coefficient for the unsaturated product at 232 nm is 4600 M⁻¹ cm⁻¹. All data were average of the triplicate measurements. Kinetic parameters K_m, V_{max}, K_{cat} were determined at the optimal temperature of each selected purified enzymes.

Thermostability measurement. To determine the half-life of inactivation, the pectate lyases were incubated in 50 mM Tris-HCl buffer pH 8.5 containing 0.5 mM CaCl₂ at 45°C. Samples were taken at various time points and followed by the pectate lyase

assay. Purified proteins were adjusted to about 0.5 mg/ml in 10 mM phosphate buffer, pH 7.0 for determining the Circular Dichroism (CD) spectrum and melting temperature. Thermal denaturation experiments were performed at 40°C of temperature increase per hour. The data were collected with JASCO J-710 CD spectrometer at the wavelength of 222 nm, where maximal signal difference was observed.

Protein crystallization and structure determination. Initial crystallization conditions were identified using the Pegs (Nextal, Montreal), Classic Suite I and II (Qiagen) screens, with the sitting drop vapor diffusion set-up at 20°C. Numerous chemical conditions yielded triclinic crystals overnight. The conditions were reproduced and optimized using the hanging drop vapor diffusion method in 24-well Linbro plates (Hampton Research). The best conditions found were 20-30% (w/v) PEG 3350, and various buffers with pH ranging from 5.5 to 8.5. The crystals belong to the primitive triclinic space group $P1$, with the unit cell dimensions of $a = 47.2 \text{ \AA}$, $b = 53.2 \text{ \AA}$, $c = 73.0 \text{ \AA}$, $\alpha = 71.7^\circ$, $\beta = 80.0^\circ$, $\gamma = 69.0^\circ$, and two protein molecules in the asymmetric unit. The Matthews coefficient V_M is $2.20 \text{ \AA}^3 \text{ Da}^{-1}$, corresponding to a solvent content of 44%. The cryoprotectant solution used consisted of mother liquor supplemented with 10-12% (w/v) glycerol and data were collected at 100K. Diffraction data extending to 2, 2.1, 1.9 Å resolution for wild-type, R236F, A31G/R236G proteins, respectively. The images were processed with the HKL2000 program package. The structure of wild-type XcPL was solved by molecular replacement using the MolRep program and the structure of *E. chrysanthemi* pectate lyase (PDB entry 1AIR) as a search model. Free atom density modification was implemented using ARP/wARP to improve the electron density maps, and the protein model was built manually using COOT. The model was refined using REFMAC5 with the TLS option with each of the monomers treated separately. The structures of R236F and A31G/R236F mutants was solved and refined in the same manner as wild-type XcPL whose structure was used as a search model for molecular replacement. Refinement statistics are summarized in Table 5 for the three structures solved. The models were validated with PROCHECK. Coordinates have been deposited in the Protein Data Bank with the codes 2OX3, 2OXZ, 2OY1 corresponding to wild-type, R236F and A31G/R236F, respectively.

Computational method. The stabilities (i.e., the free energies of folding) of pectate lyase mutants relative to the wild type protein were calculated with the FOLD-X program, using the crystal structure of the wild type protein determined in this study (chain A). The FOLD-X program employs a first principle-based energy function that was shown to be able to predict folding free energies with a squared-correlation coefficient of

0.69 and a standard deviation of 0.81 kcal/mol, within a dataset of over 1000 mutations from various proteins. The effect of R236F mutation on stability was further verified by performing a FOLD-X calculation using the crystal structure of the R236F mutant (chain A) and simulating its mutation to the wild-type protein. All crystallographic water molecules and observed phosphate ions were removed. Possible water bridges between protein atoms were implicitly taken into account during FOLD-X calculations. Point mutations were introduced by the FOLD-X program with the mutate function of the WHAT IF program that optimizes the mutated side-chains with respect to rotamer distributions and hydrogen-bond networking. The analysis of relative stabilities in terms of contributions from van der Waals, hydrogen bonding, electrostatic and water bridge interaction energies, hydrophobic and polar desolvation, and main-chain and side-chain entropies, were based on the FOLD-X calculations. Clash energies were not included in the calculation of folding free energies.

Results

15 *Example 1: Characterization of the properties of Xanthomonas campestris pectate lyase II (XcPL)*

The full-length XcPL gene encodes a polypeptide having 353 amino acids, with a molecular weight of 37 kD and pI of 8.7 according to its DNA-predicted amino acid sequences. Optimal pH and temperature of the enzyme were found to be 8.5 and 50°C, respectively. Melting temperature of this enzyme was determined to be 48°C. Half-life of inactivation at 45°C for this enzyme was determined to be 54 min.

Example 2: Selecting mutations on the XcPL polypeptide

Three thermostable pectate lyases, which have been biochemically characterized: CAD56882 from *Bacillus licheniformis*, BAB40336 from *Bacillus* sp. TS-47, AAD35518 from *Thermotoga maritima* MSB8 and thermoliable XcPL (NP_638163) were analyzed by multiple sequence alignment using ClustalW program. Compared with XcPL pairwise, the thermostable pectate lyases share only 25-35% of identical amino acids. Interestingly, 9 amino acid residues in underlined font without bolding in Fig. 1 (corresponding to positions 26, 31, 64, 66, 69, 70, 123, 187 and 236 of the wild-type XcPL (NP_638163)) were found to be highly conserved in all three thermostable pectate lyases, but variable in the thermoliable XcPL (Fig. 1). Residues in bold font without underlining in Fig. 1 are conserved catalytic sites, residues in underlined bold italic font in Fig. 1 are conserved calcium binding sites, the gray shaded region in Fig. 1 is the core structure of the parallel

β -helix vWIDH region, and residues in underlined font without bolding in Fig. 1 are sites conserved in the thermostable pectate lyases (CAD56882 from *Bacillus licheniformis*, BAB40336 from *Bacillus sp. TS-47*, AAD35518 from *Thermotoga maritima MSB8*) but variant in *Xanthomonas campestris* pectate lyase II (NP_638163).

5 *Example 3: Effect of single amino acid substitution and a double mutation on the thermostability of XcPL*

Individual effects of amino acid substitutions in an enzyme are generally difficult to predict. Enzyme properties may be unaffected, adversely affected or positively affected. To determine the individual effect of each of the conserved amino acids in thermostable pectate lyases on the thermostability of XcPL, nine mutants of the *Xanthomonas campestris* pectate lyase II enzyme were produced, each containing a single substitution with conserved amino acid at a position corresponding to one of the nine sequence differences between the three thermostable pectate lyases and the thermoliable one.

15 Referring to Tables 1 and 2, in the name of each mutant the number refers to the amino acid position in the parent enzyme, the first letter refers to the amino acid present at that position in the parent and the second letter refers to the amino acid present at that position in the mutant.

Table 2

Enzyme	Half-life of inactivation at 45°C (min)	Activity in crude cell extract (IU/mg protein)
Parent	54.2±5.9	0.64±0.01
V26A	38.8±8.5	0.53±0.04
A31G	51.9±8.8	3.37±0.09 (5×)
L64I	44.6±2.8	0.54±0.01
Y66V	NA	No activity
K69T	48.2±7.4	0.87±0.03
F70I	14.0±1.5	0.81±0.03
Q123R	29.0±1.3	0.62±0.08
V187I	<10	0.55±0.08
R236F	1292±139 (23×)	1.42±0.17 (2×)
A31G/R236F	659±41 (12×)	3.48±0.20 (5×)

Fig. 2 provides the amino acid sequences of the parent (wild-type) *Xanthomonas campestris* pectate lyase II (SEQ ID NO: 1) and three of its mutants: A31G (SEQ ID NO: 3), R236F (SEQ ID NO: 4) and A31G/R236F (SEQ ID NO: 5). The amino acid sites that are modified in the mutants are in underlined bold font. Thus, alanine at position 31 in the parent enzyme is converted to glycine in mutants A31G and A31G/R236F. Arginine at position 236 in the parent enzyme is converted to phenylalanine in mutants R236F and A31G/R236F.

One of the single mutants, Y66V, exhibited no enzymatic activity. Four out of nine individual single mutants, F70I, Q123R, V187I and R236F exhibited a remarkable difference in thermostability when compared to the parent enzyme. F70I, Q123R and V187I had remarkably lower thermostability while R236F had remarkably higher thermostability. Notably, the mutant R236F displayed 23 times longer half-life at 45°C than the parent. In addition, the mutant A31G was 5-fold more active than the parent while its thermostability remained similar to the parent. The most active mutant A31G and the most thermostable mutant R236F were combined to produce a double mutant A31G/R236F. This resulted in 5 times more activity and 12 times more thermostability than the parent enzyme (Table 2 and Fig. 3). In Table 2, the numbers in parentheses are fold of improved properties over the parent. It can be seen from Table 2 and Fig. 3 that the stability of wild-type *Xanthomonas campestris* pectate lyase is low. Its half-life of inactivation at 45°C is less than one hour. About 90% of the original activity was lost after 3 hours at 45 °C.

Example 4: Kinetic properties of the improved mutant pectate lyases

To confirm the enzyme properties of the improved mutants derived from *Xanthomonas campestris* pectate lyase II, three mutant enzymes A31G, R236F and double mutant A31G/R236F and the parent enzyme were purified to apparent homogeneity with a single cation-exchange SP-Sepharose column. Those peaks appeared as a single band on SDS-PAGE with a molecular mass of 37 kD, in good agreement with their theoretical molecular weight (37,355 daltons). Furthermore, zymogram analysis of the purified enzymes indicated these single bands were active pectate lyases. The intensity of the clearing band for each enzyme is consistent with its specific activity (Fig. 4). The K_m values of the mutants A31G, R236F and A31G/R236F using polygalacturonic acid (PGA) as substrate are not significantly different from each other and to the parent enzyme. However, the catalytic efficiency for both single mutant A31G and double mutant A31G/R236F increased about 2-fold, compared to parent enzyme. The activity for the most thermostable mutant R236F still retained the same level

of its parent. The specific activities of the purified enzymes are consistent with their K_{cat}/K_m values (Table 3). Similar results were observed when citrus pectin was used as the substrate (Table 4). Comparing the specific activities of the activities in crude cell extract and the purified enzymes (Table 2 and Table 3), the protein expression levels in strains harboring A31G, R236F and A31G/R236F mutant XcPL genes were 2 to 3 fold greater than on the strain harboring the wild-type gene.

Table 3

Enzyme	K_m ($g \cdot l^{-1}$)	K_{cat} (s^{-1})	K_{cat}/K_m ($l \cdot g^{-1} \cdot s^{-1}$)	Specific activity (IU/mg)
Parent	0.98	114	116	196
A31G	0.73	194	266	333
R236F	0.83	121	146	208
A31G/R236F	0.93	216	232	370

Table 4

Enzyme	K_m ($g \cdot l^{-1}$)	K_{cat} (s^{-1})	K_{cat}/K_m ($l \cdot g^{-1} \cdot s^{-1}$)	Specific activity (IU/mg)
Parent	0.80	61	76	105
A31G	0.68	117	172	200
R236F	0.59	67	114	115
A31G/R236F	0.55	139	253	238

Example 5: Determination of melting temperatures of mutant pectate lyases

The melting temperatures of purified parent enzyme, mutant A31G, mutant R236F and mutant A31G/R236F were determined by circular dichroism spectrometry to be 48°C, 47.5°C, 54°C and 53°C, respectively (Fig. 5), which are consistent with their half-lives at 45°C (Table 2).

Example 6: Effect of mutants on efficiency of pectin removal from hemp fiber

The amount of 4,5-unsaturated galacturonate residues liberated from pectin, which is measured by the increasing absorbance at 232 nm (Collmer et al. 1988), was used as an indicator for the effect of pectate lyase on the removal of pectic substance

from untreated hemp fibers (Ouajai and Shanks. 2005). The amount of pectin-degraded products was confirmed with dinitrosalicylic acid assay (Miller. 1959). In this study, to determine the efficiency of the improved *Xanthomonas campestris* pectate lyase variants in the bioscouring process, XcPL mutants A31G, R236F and A31G/R236F were used to treat natural hemp fiber with parent (wild-type) XcPL used as a control. All three of the mutants were found to be more effective in removing pectin from hemp fiber than the parent (Fig. 6). Setting release of pectin by the action of the double mutant A31G/R236F at 100%, the parent XcPL resulted in a pectin release of about 60% while mutants A31G and R236F resulted in pectin releases of about 85% and 80%, respectively.

10 Example 7: Structural analysis of wild-type XcPL and mutant proteins

To understand the structural basis for thermal stability of the mutant proteins, R236F, A31G/R236F and wild-type XcPL were crystallized and their structures solved by molecular replacements. These structures share the same right-handed parallel β -helix architecture as described for *Erwinia chrysanthemi* (now classified as *Dickeya dadantii*) pectate lyase PelC. Crystallographic data collection and refinement is shown in Table 5.

Table 5

	Wild-type	R326F	A31G/R236F
Data Collection			
Resolution range	69.17-2	69.5-2.1	69.3-1.9
Unique reflections	33827	29174	42284
Completeness (%)	83.29	85.3	89.08
Rsym ^a	0.048	0.047	0.059
Average $I/\sigma(I)$	20.1	22	28.3
Refinement Statistics			
Rwork ^b	15.7	15.9	16.1
Rfree ^b	21.1	26.4	18.9
Ramachandran plot			
Allowed (%)	98.9	99.1	99.1
Generously allowed (%)	1.1	0.9	0.9
r.m.s. Deviation			
Bonds (Å)	0.01	0.009	0.006
Angles (°)	1.283	1.180	1.062

$$^a\text{Rsym} = (\sum |I_{\text{obs}} - I_{\text{avg}}|) / \sum I_{\text{avg}}$$

$$^b\text{Rwork/free} = (\sum |F_{\text{obs}} - F_{\text{calc}}|) / \sum F_{\text{obs}}$$

Residue R236 is partially solvent-exposed and located in the T1.6 loop (the 6th loop of turn T1) that confers catalytic activity to the β -helix core. Its guanidinium group is 10Å, 17Å and 22Å away from the putative active sites R235, R230 and K199, respectively. This may explain the similar activities of wild-type and R236 mutant enzymes. The R236F substitution caused no structural change in the enzyme. Although bulkier, the F236 side chain occupies the same region of space as the wild-type R236; no neighboring residues were affected nor major electrostatic interactions interrupted. However, the aromatic side chain in this position allows hydrophobic interactions with several adjacent side chains, particularly M258 from the T1.7 loop, the aliphatic portions of T210 and N212 from the T1.5 loop, and the neighboring R235. Hydrophobic stacking between the T1.5 and T1.7 loops appears to confer higher stability to the T1.6 loop. A structure-based first-principle energetic analysis (Fig. 7 and Table 6) predicts a favorable change in folding free energy of -1.7 kcal/mol for the stabilizing R236F mutation, underscoring the gain afforded by hydrophobic desolvation (-2.8 kcal/mol) as driving the increase in stability. This more than offsets the major destabilizing contribution, i.e., the loss of hydrogen bonding energy upon mutation (1.4 kcal/mol), between the carbonyl group of D207 and the guanidinium group of R236.

Table 6

Mutant	Internal energy				Desolvation		Entropy		Calc ^a	Expt ^b
	vdW	elec	hb	wb	hyd	pol	sc	mc		
R236F	-0.24	0.22	1.40	-0.07	-1.24	-1.54	-0.23	0.01	-1.68	-1.38
A31G/ R236F	-0.09	0.22	1.40	-0.07	-0.97	-1.81	-0.31	0.17	-1.45	-1.08
F70I	0.70	0.01	0.00	-0.02	1.42	-0.75	-0.18	0.18	1.37	0.59
V187I	-0.35	0.00	0.00	0.00	-0.83	0.13	0.25	0.09	-0.70	>0.73
Y66V	1.63	0.00	1.30	0.26	2.89	-2.72	-0.50	-0.33	2.55	NA

^a Total predicted folding free energy change upon mutation, $\Delta\Delta G_{\text{fold}}$ (kcal/mol), calculated as the sum of energetic contributions (kcal/mol), i.e., vdW + elec + hb + wb + hyd + pol + sc + mc, where vdW: van der Waals energy, elec: electrostatic energy, hb: hydrogen bond energy, wb: water bridge energy, hyd: hydrophobic desolvation free energy, pol: polar desolvation free energy, sc: side-chain entropic contribution, mc: main-chain entropic contribution. Calculations were done with the FOLD-X program (Guerios et al. 2002). Absolute contributions larger than 0.81 kcal/mol (the standard deviation of calibrated FOLD-X function) are in bold.

^b Experimental stability data is expressed as fold-change in the half-time of inactivation upon mutation (Table 2), converted to logarithmic scale.

The double mutation A31G/R236F also does not introduce major variations relative to XcPL wild-type and R236F mutant structures. A noticeable change is a peptide flip at the A31G mutation site, bringing the backbone carbonyl of A30 closer to the side chain of K151 side chain that undergoes a concerted conformational change. The A31G mutation resides in the N-terminal region of the enzyme on the opposite side of the β -helix relative to the catalytic site. Thus the 2-fold increase in catalytic efficiency upon the A31G mutation cannot be accounted for by the present structural data. Mutations distant from catalytic sites are documented to be as effective as close mutations in improving enzymatic activity, but the underlying factors are often subtle and elude structural interpretations. Computational predictions yielded a marginal destabilizing effect (0.2 kcal/mol) for the A31G mutation in agreement with the experimental data (Table 2 and Fig. 7).

The remaining single point mutations had either marginal or destabilizing effects on the thermal stability. A retrospective structural and computational analysis can rationalize why these single-amino acid substitutions did not materialize into improved thermostability of the target mesophilic enzyme (Fig. 7 and Table 6).

References: The disclosures of the following references are incorporated herein by reference in their entirety.

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15 Other advantages that are inherent to the structure are obvious to one skilled in the art. The embodiments are described herein illustratively and are not meant to limit the scope of the invention as claimed. Variations of the foregoing embodiments will be evident to a person of ordinary skill and are intended by the inventor to be encompassed by the following claims.

Claims:

1. Polypeptide comprising the amino acid sequence:

5 MTSKTLQGALALALSACAAGAIAGPVGYGAX₁TTGGGNKVPVNVATFEAMQSAIDSYSGSGGLVLN
 YTGKFDGFTIKDVCAQWKLPAKTVQIKNKSDVTIKGANGSAANFGIRVVGNAHNVI IQNMTIGLLQ
 GGEDADSI SLEGNSSGEP SKIWVDHNTVFASLT KC SGAGDASFDGGIDMKKGVHHVTVS YNYVYNY
 QKVALNGYS DSDTKNSAARTTYHHNRFENVESRVPLQRX₂GLSHIYNNYFNNVTTSGINVRMGGIA
 KIESNYFENIKNPVTSRDSSEIGYWDLINNYVGSGITWGTPDGSKPYANATNWISTKVFPE SLGYI
 YTVTPAAQVKAKVIATAGAGKNLAE

10 wherein X₁ is alanine or glycine and X₂ is arginine or phenylalanine with the proviso that
 X₂ is not arginine when X₁ is alanine.

2. The polypeptide according to claim 1, wherein the amino acid sequence is SEQ ID
 NO: 3.

3. The polypeptide according to claim 1, wherein the amino acid sequence is SEQ ID
 NO: 4.

15 4. The polypeptide according to claim 1, wherein the amino acid sequence is SEQ ID
 NO: 5.

5. The polypeptide according to any one of claims 1 to 4 that is isolated.

6. Use of the polypeptide as claimed in any one of claims 1 to 5 for degrading non-
 cellulosic material.

20 7. Use according to claim 6 in detergents, in textile processing, in cellulose fiber
 processing, in the treatment of pectic wastewaters, in paper making, in coffee and tea
 fermentations and for improving the yield and the clarification of fruit juices.

25 8. Process of isolating fibers from non-cellulosic material in a fiber plant comprising
 contacting raw fibers of the fiber plant with the polypeptide as claimed in any one of
 claims 1 to 5.

9. The process according to claim 8, wherein the non-cellulosic material comprises
 pectin.

10. The process according to claim 8 or 9, wherein the fiber plant is hemp, jute, flax,
 ramie or cotton.

11. The process of any one of claims 8 to 10, wherein the contacting is done in aqueous medium at a pH in a range of from 4 to 10.
12. The process of claim 11, wherein the pH is in a range of from 8 to 9.
13. The process of any one of claims 8 to 12, wherein the contacting is done at a
5 temperature in a range of from 30°C to 55°C.
14. The process of claim 13, wherein the temperature is in a range of from 40°C to 50°C.

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NP_638163 -----GPVGYGAATTGGGNKVPVNVATFEAMQSAIDSYSGS----- 59
CAD56882 ASALNSGKVNPLADFLKGFAALNGGTTGGEGGQTVTVTTGDQLIAALKNKNNAN----- 54
BAB40336 -----KELGHEVLKPYDGWAAYGEGTTGGAMASPQNVFVVTNRTELIQALGGNNHTNQY 54
AAD35518 -----ASLNDKPVGFASVPTADLPEGTVGGLGGEIVFVRTAEELKYTTAEGKY----- 49
          . . . * . * * . * . .

NP_638163 --GGLVLNYTGKFDFGTIKDVCAQ----- 81
CAD56882 --TPLKIYVNGTITTS----- 68
BAB40336 NSVPKIIYVKGTIDLNVDDNNQPVGPDFYKDPHFDFEAYLREYDPATWGKKEVEGPLEEA 114
AAD35518 -----VIVVDGTIVFE----- 60
          : * . :

NP_638163 ----WKLPAKTVQIKNKSDVTIKG--ANGSAANFGIRVVGNAHNVI IQNMTIGLLQGGE- 134
CAD56882 ----NTSASKIDVKDVSNSVIVGSGTKGELKGIGIKIWR-ANNIIIRNLKIHEVASG-- 120
BAB40336 RVRSQKKQKDRIMVYVGSNTSIIGVGKDAKIKGGGFLIKN-VDNVIRNIEFEAPLDYFP 173
AAD35518 -----PKREIKVLSDKTIVG-INDAKIVGGGLVIKD-AQNVIIRNIHFEGFYMEDD 109
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NP_638163 -----DADSISLEGNSSGEPSKIWDHNTVFA-----SLTKCSGAGDASF 174
CAD56882 -----DKDAIGIEGPSK----NIWVDHNELYH-----SLN----VDKDYY 152
BAB40336 EWDPTDGTLGEWNSEYDSISIEGSSH-----IWIDHNTFTDGDHPDRSLGTYFGRPFQOH 228
AAD35518 PRGK-----KYDFDYINVENSHH-----IWIDHCTFVN-----GN 139
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CAD56882 DGLFDVKRDAEYITFSWNYVHDGWKSMLMGSSSDSDNYN-----RTITFHHNWFENLNSRV 207
BAB40336 DGALDIKNSSDFITISYNVFTNHDKVTLIGASDSRMADSG--HLRVTLHHNYKNVTQRL 286
AAD35518 DGAVDIKKYSNYITVSWCKFVDHDKVSLVGSSDKEDPEQAGQAYKVTYHHNYFKNCIQRM 199
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CAD56882 PSFRFEGEGHIYNNYFNKIIDSGIN-----SRMGARIRIENNLFEN----- 247
BAB40336 PRVREGQVHIYNNYEFSNLADYD-----FOYAWGVGVFSQIYAQNNYFSFDWDI 336
AAD35518 PRIRFGMAHVFNNFYSMGLRTGVSGNVFPIYGVASAMGAKVHVEGNYFMGYGAVMAEAG- 258
* * * * * : * * * : * . . . : * :

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CAD56882 AKDPIVSWYSSSPGYWHVSNNKFVN-----SRGSMPTTSTTTYNPP-----YSYSL 293
BAB40336 DPSLIIKVWSKNEESMYETGTIVDLPNGRRYIDLVASYNESNTLQLKKEVTWKPMFYHVI 396
AAD35518 IAFLPTRIMGPVEGYLTLGEGDAKNEFYYCKEPEVRPVEEGKPALDPRE-----YYDYTL 313
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NP_638163 TPAAQVKAKVIATAGAGKNLAE----- 353
CAD56882 DNVDNVKSIVKQNAGVGKINP----- 314
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AAD35518 DPVQDVPKIVVDGAGAGKLVFEELNTAQ 341
          . . * * * * :

NP_638163 = SEQ ID NO: 1
CAD56882 = SEQ ID NO: 24
BAB40336 = SEQ ID NO: 25
AAD35518 = SEQ ID NO: 26

```

FIG. 1

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SEQ ID NO: 1 – parent XcPL

MTSKTLQGALALALSACAAGAIAGPVGYGAAATTGGGNKVPVNVATFEAMQSAIDSYSGSGGLVLNYTGKDFD
GTIKDVCAQWKLPAKTVQIKNKSDVTIKGANGSAANFGIRVVGNAHNVI IQNMTIGLLQGGEDADSI SLEGN
SSGEP SKIWVDHNTVFASLT KC SGAGDASFDGGIDMKKGVHHVTVSYNYVYNYQKVALNGYSDSDTKNSAAR
TTYHHNRFENVESRVPLQRRGLSHIYNNYFNNVTTSGINVRMGGIAKIESNYFENIKNPVTSRDSSEIGYWD
LINNYVGSGITWGTPDGSKPYANATNWI STKVFPE SLGYIYTVTPAAQVKAKVIATAGAGKNLAE

SEQ ID NO: 3 - Mutant A31G

MTSKTLQGALALALSACAAGAIAGPVGYGAGGTTGGGNKVPVNVATFEAMQSAIDSYSGSGGLVLNYTGKDFD
GTIKDVCAQWKLPAKTVQIKNKSDVTIKGANGSAANFGIRVVGNAHNVI IQNMTIGLLQGGEDADSI SLEGN
SSGEP SKIWVDHNTVFASLT KC SGAGDASFDGGIDMKKGVHHVTVSYNYVYNYQKVALNGYSDSDTKNSAAR
TTYHHNRFENVESRVPLQRRGLSHIYNNYFNNVTTSGINVRMGGIAKIESNYFENIKNPVTSRDSSEIGYWD
LINNYVGSGITWGTPDGSKPYANATNWI STKVFPE SLGYIYTVTPAAQVKAKVIATAGAGKNLAE

SEQ ID NO: 4 - Mutant R236F

MTSKTLQGALALALSACAAGAIAGPVGYGAATTTGGGNKVPVNVATFEAMQSAIDSYSGSGGLVLNYTGKDFD
GTIKDVCAQWKLPAKTVQIKNKSDVTIKGANGSAANFGIRVVGNAHNVI IQNMTIGLLQGGEDADSI SLEGN
SSGEP SKIWVDHNTVFASLT KC SGAGDASFDGGIDMKKGVHHVTVSYNYVYNYQKVALNGYSDSDTKNSAAR
TTYHHNRFENVESRVPLQRFGLSHIYNNYFNNVTTSGINVRMGGIAKIESNYFENIKNPVTSRDSSEIGYWD
LINNYVGSGITWGTPDGSKPYANATNWI STKVFPE SLGYIYTVTPAAQVKAKVIATAGAGKNLAE

SEQ ID NO: 5 - Mutant A31G/R236F

MTSKTLQGALALALSACAAGAIAGPVGYGAGGTTGGGNKVPVNVATFEAMQSAIDSYSGSGGLVLNYTGKDFD
GTIKDVCAQWKLPAKTVQIKNKSDVTIKGANGSAANFGIRVVGNAHNVI IQNMTIGLLQGGEDADSI SLEGN
SSGEP SKIWVDHNTVFASLT KC SGAGDASFDGGIDMKKGVHHVTVSYNYVYNYQKVALNGYSDSDTKNSAAR
TTYHHNRFENVESRVPLQRFGLSHIYNNYFNNVTTSGINVRMGGIAKIESNYFENIKNPVTSRDSSEIGYWD
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FIG. 2

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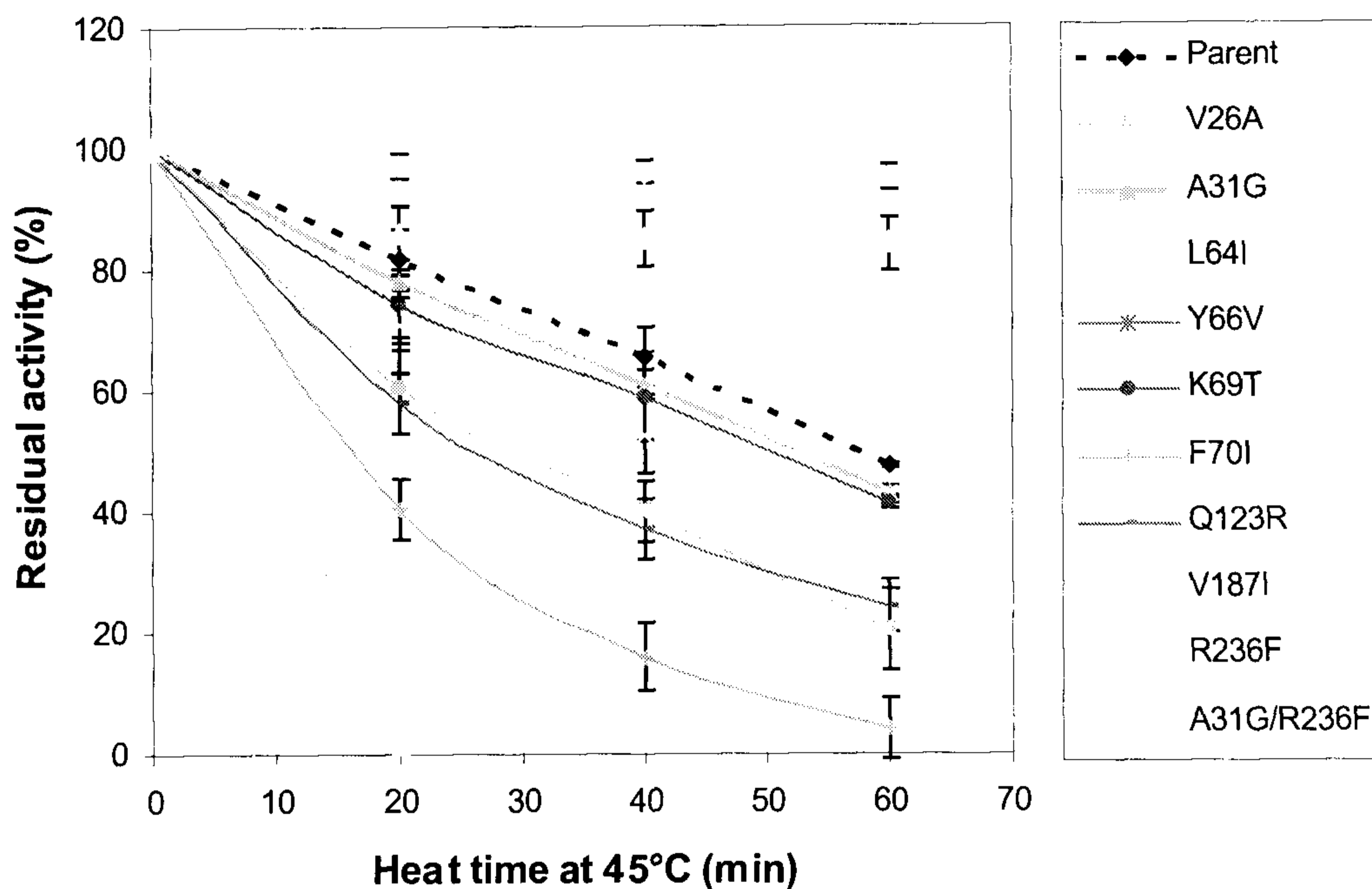


Fig. 3A

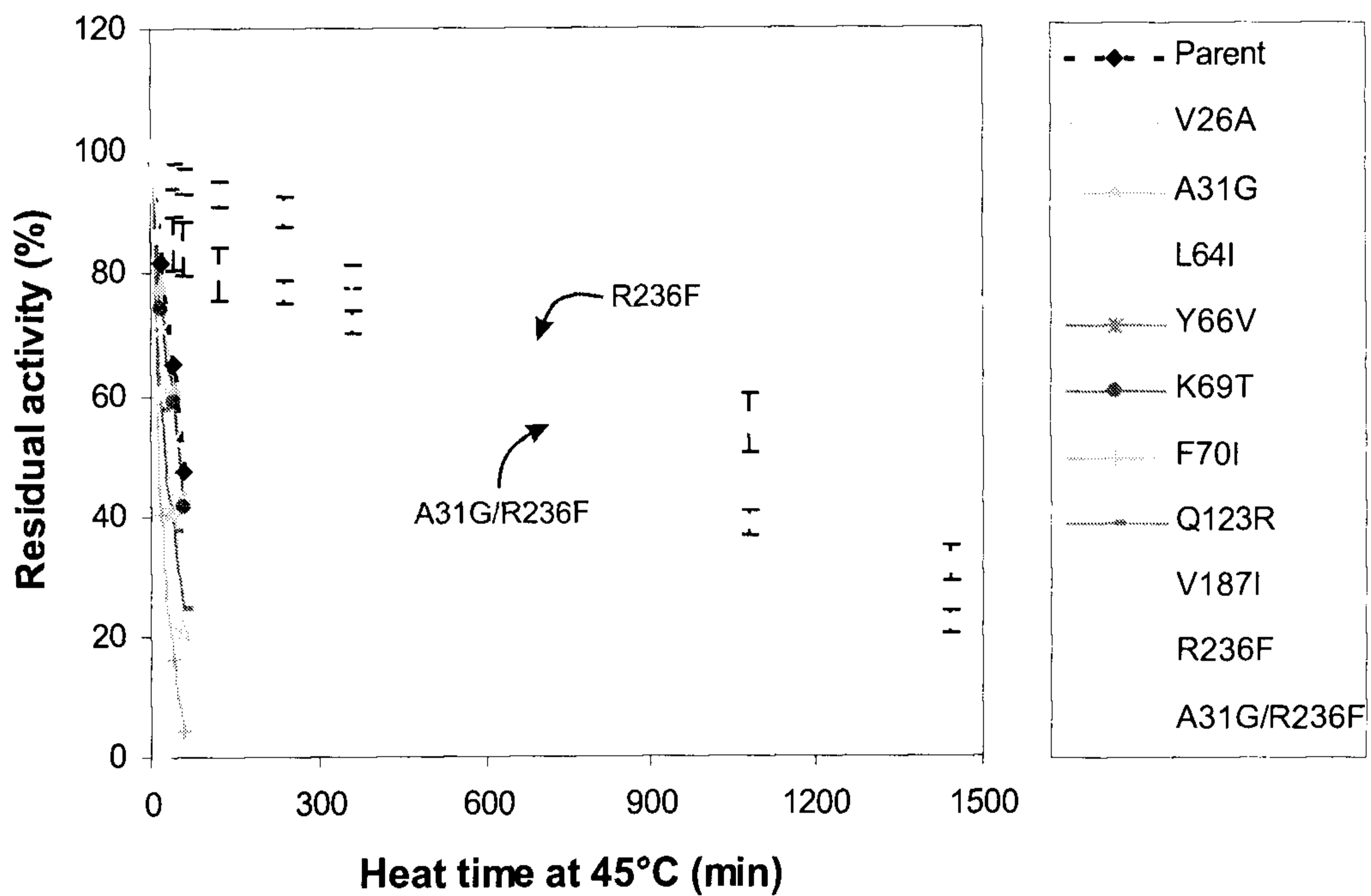


Fig. 3B

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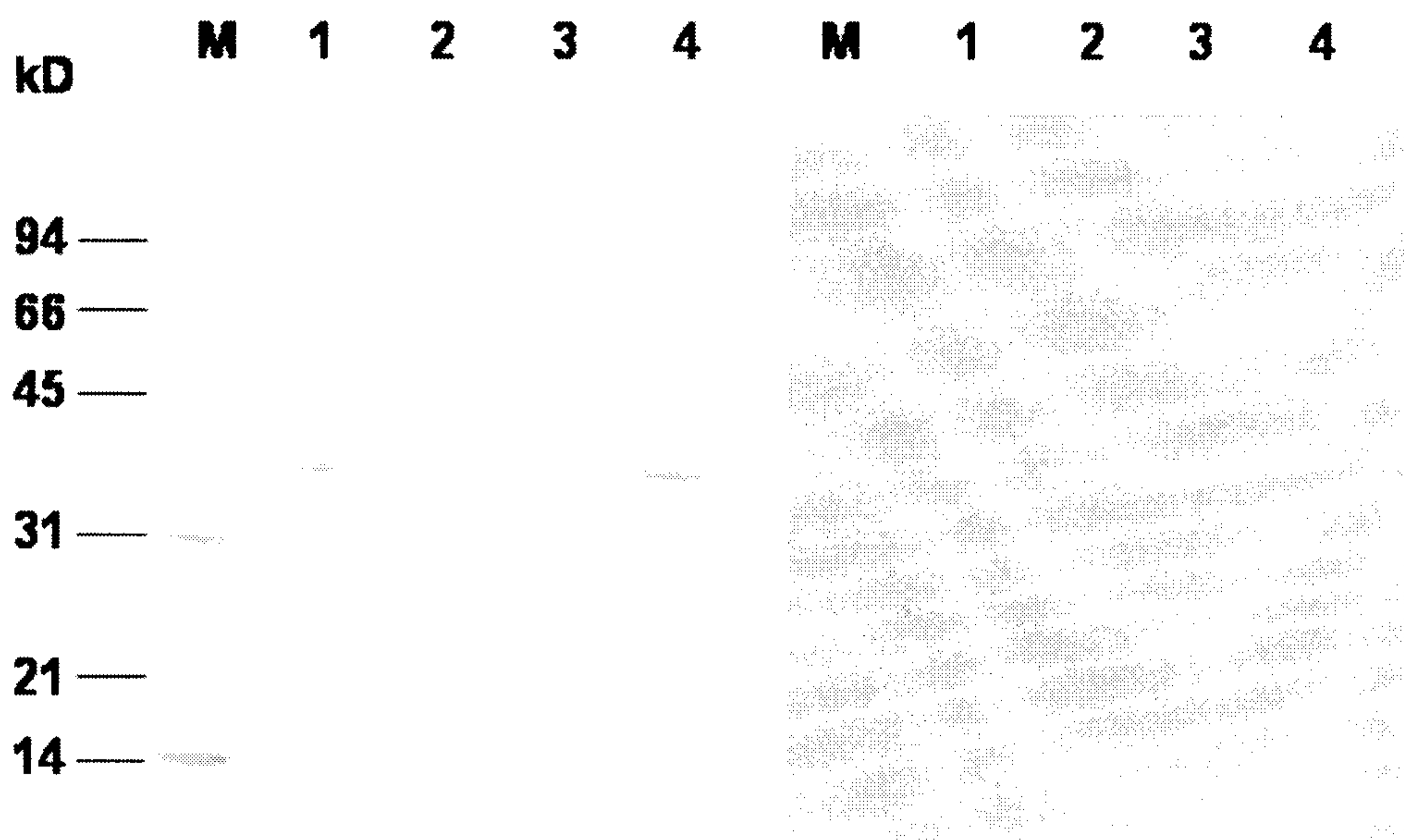


Fig. 4

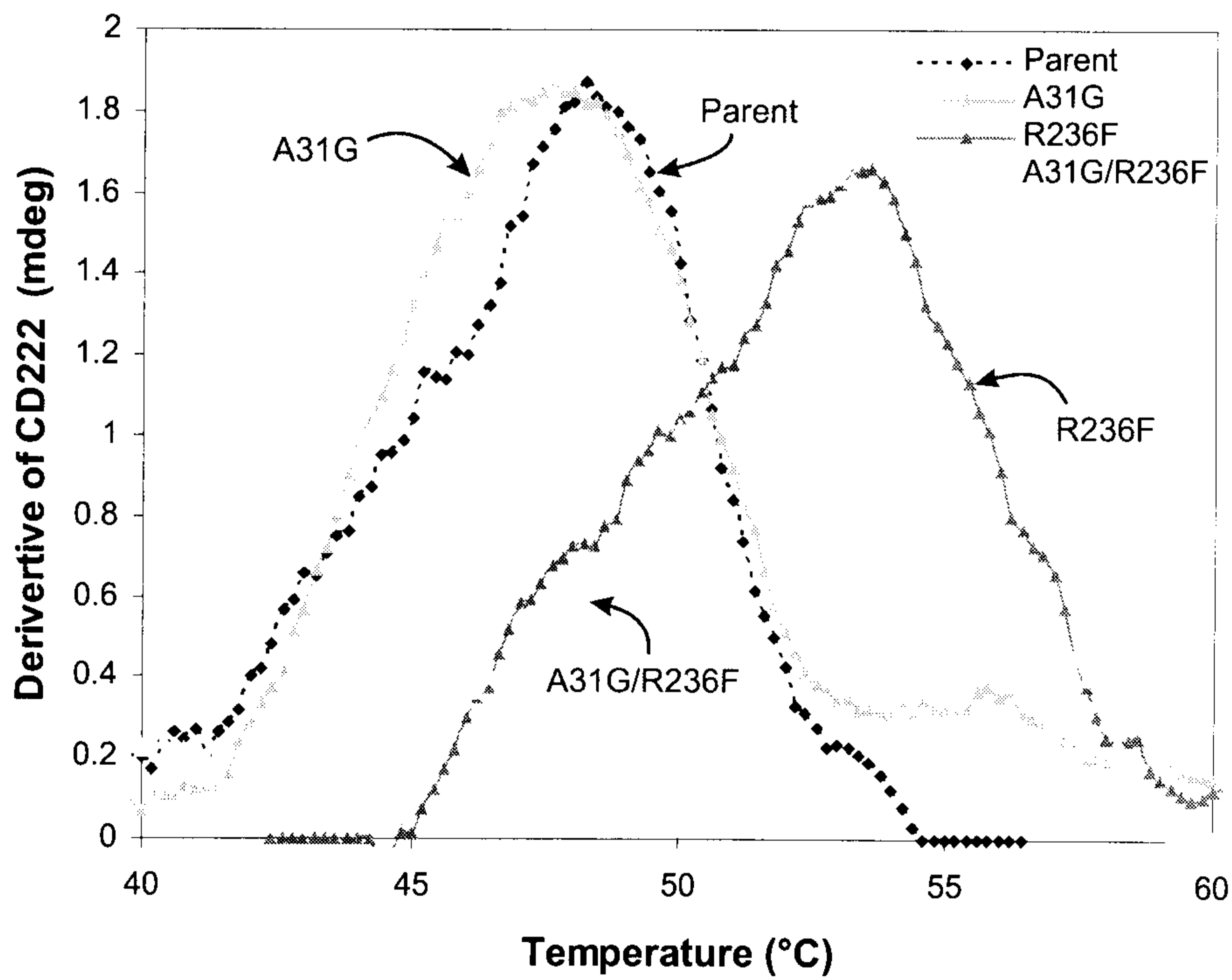


Fig. 5

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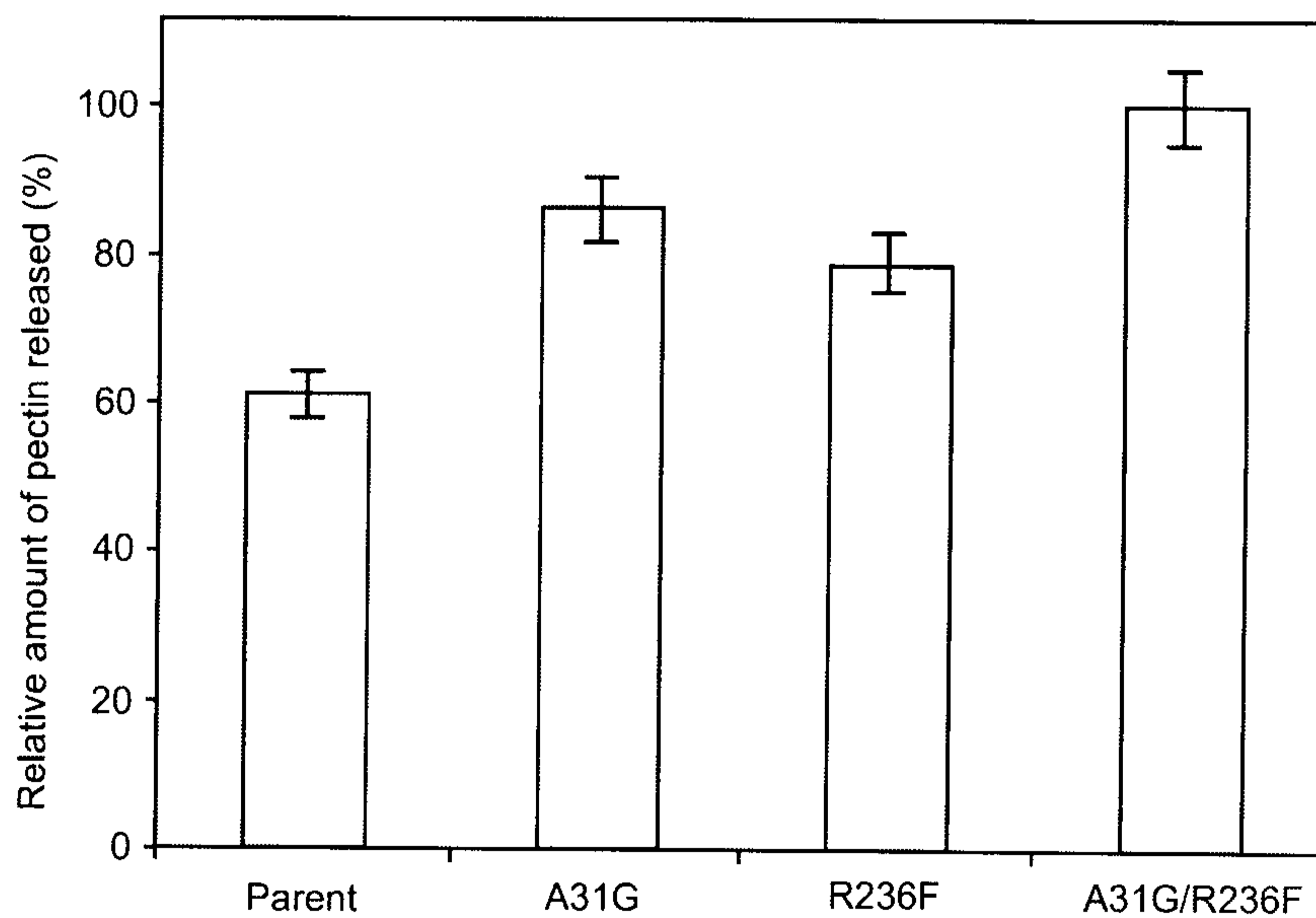


Fig. 6

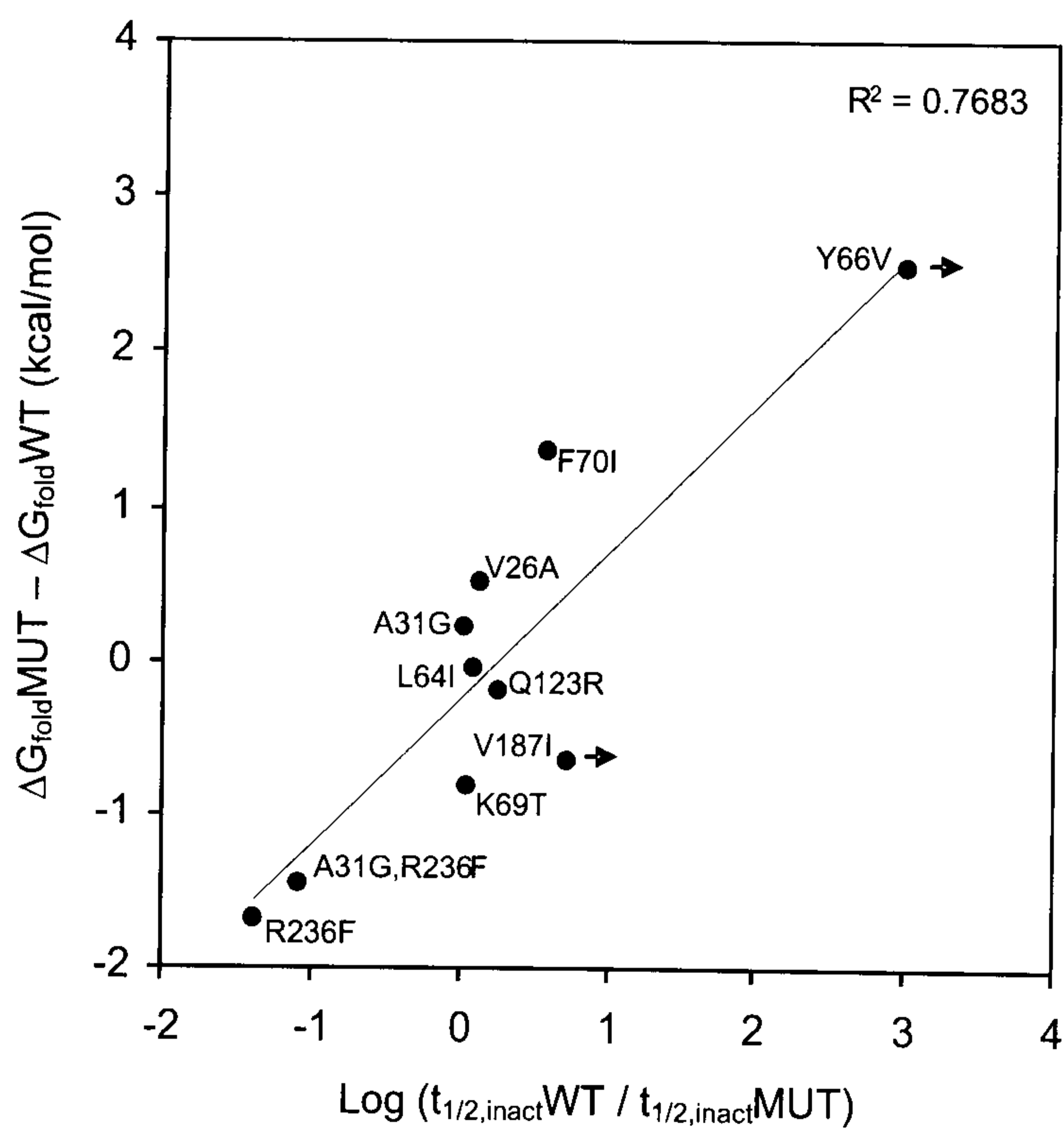


Fig. 7

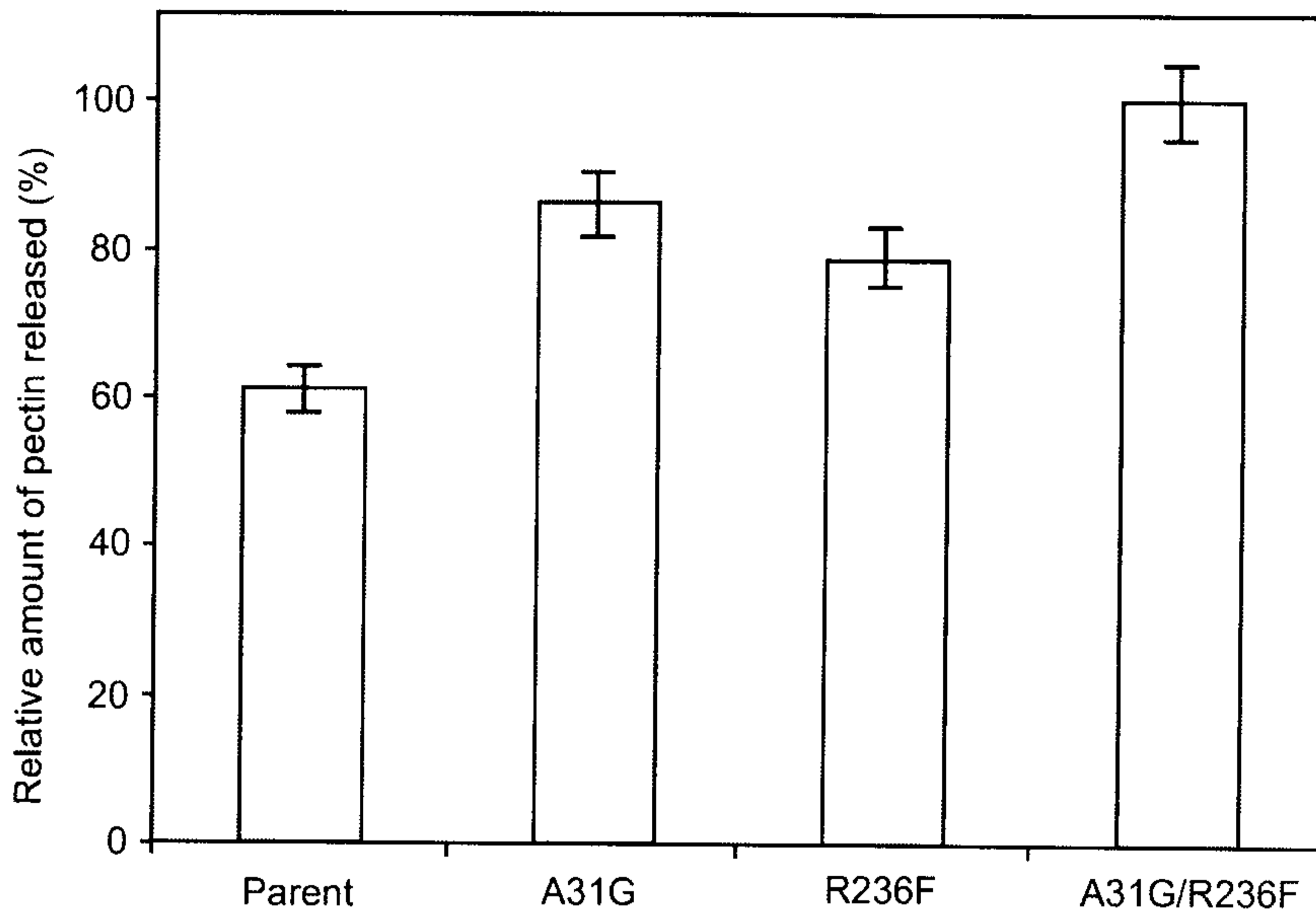


Fig. 6