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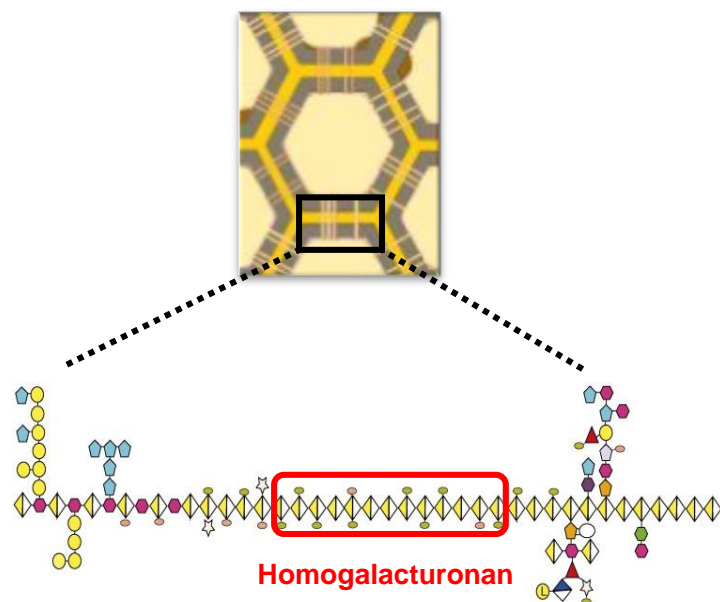
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Corresponding Author:	Fabien Sénéchal, Ph.D UMR1158: Transfrontalière BioEcoAgro Amiens, FRANCE
First Author:	Josip Safran, Ph.D
Order of Authors:	Josip Safran, Ph.D Vanessa Ung Julie Bouckaert, Ph.D Olivier Habrylo, Ph.D Roland Molinié, Ph.D Jean-Xavier Fontaine, Ph.D Adrien Lemaire Aline Voxeur, Ph.D Serge Pilard, Ph.D Corinne Pau-Roblot, Ph.D Davide Mercadante, Ph.D Jérôme Pelloux, Pr. Fabien Sénéchal, Ph.D
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Suggested Reviewers:	Mirjam Czjzek Biological Research Station Roscoff czjzek@sb-roscoff.fr Estelle Bonnin Biopolymers Interactions Assemblies Research Unit estelle.bonnin@inrae.fr

	Suzanna Saez-Aguayo Andrés Bello University susana.saez@unab.cl
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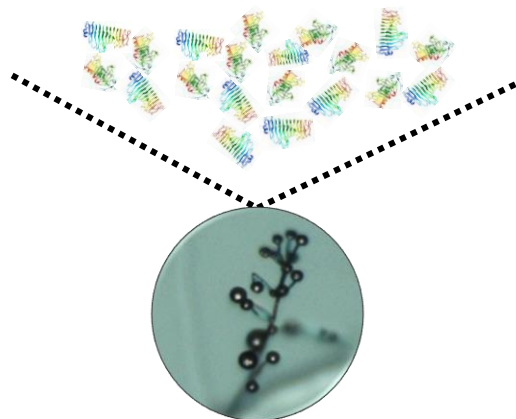
Highlights

- First crystallographic structure of *Verticillium* pectate lyase.
- Identification of key amino-acids for the activity
- Biochemical characterization of the enzyme and LC-MS/MS analysis of the oligogalacturonides (OG) produced
- Susceptible or partially resistant flax cultivars differ in OGs produced

PRIMARY CELL WALL



Pectate lyase like (PLLs) degrading enzymes

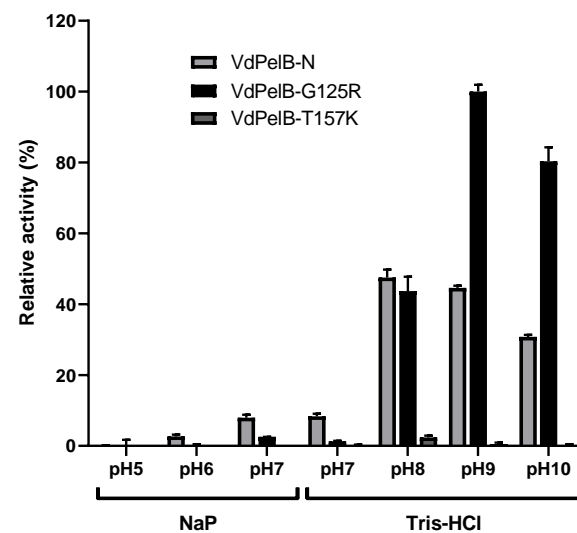


VERTICILLIUM DAHLIAE

VdPeIB CRYSTAL STRUCTURE

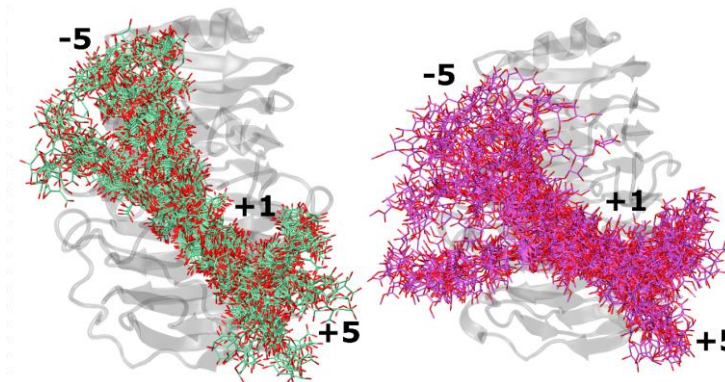


BIOCHEMICAL CHARACTERIZATION

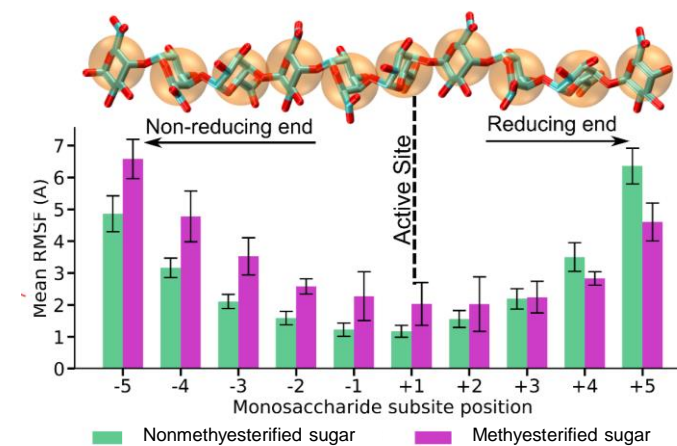


MD SIMULATIONS

VdPeIB substrate dynamics



RMSF analysis



Declaration of interests

☐The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

☒The authors declare the following financial interests/personal relationships which may be considered as potential competing interests:

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Abstract

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Keywords: Pectate lyase, pectins, homogalacturonan, oligogalacturonides, *Verticillium dahliae*.

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Josip Safran¹, Vanessa Ung², Julie Bouckaert³, Olivier Habrylo¹, Roland Molinié¹, Jean-Xavier Fontaine¹, Adrien Lemaire¹, Aline Voxeur⁴, Serge Pilard⁵, Corinne Pau-Roblot¹, Davide Mercadante², Jérôme Pelloux^{1*}, Fabien Sénéchal^{1*}

¹ : UMRT INRAE 1158 BioEcoAgro – BIOPI Biologie des Plantes et Innovation, Université de Picardie, 33 Rue St Leu, 80039 Amiens, France. ² : School of Chemical Sciences, The University of Auckland, Private Bag 92019, Auckland 1142, New Zealand. ³ : UMR 8576 Unité de Glycobiologie Structurale et Fonctionnelle (UGSF) IRI50, Avenue de Halley, 59658 Villeneuve d'Ascq, France. ⁴ : Université Paris-Saclay, INRAE, AgroParisTech, Institut Jean-Pierre Bourgin (IJPB), 78000, Versailles, France. ⁵ : Plateforme Analytique, Université de Picardie, 33, Rue St Leu, 80039 Amiens, France.

*contributed equally as last authors

Corresponding author: Fabien Sénéchal (fabien.senechal@u-picardie.fr) and Jérôme Pelloux (jerome.pelloux@u-picardie.fr)

UMR INRAE 1158 BioEcoAgro-Biologie des Plantes et Innovation, Université de Picardie Jules Verne, UFR des Sciences, 33 Rue St Leu, 80039 Amiens, France

Abstract

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1. Introduction

Primary cell wall, a complex structure of proteins and polysaccharides, cellulose and hemicelluloses, is embedded in a pectin matrix. Pectins, are complex polysaccharides composing up to 30% of cell wall dry mass in dicotyledonous species [1]. Pectin is mainly constituted of homogalacturonan (HG), rhamnogalacturonan I (RG-I) and rhamnogalacturonan II (RG-II) domains, but its composition can differ between plant organs and among species. The most abundant pectic domain is HG, a linear homopolymer of α -1,4-linked galacturonic acids (GalA), which represents up to 65% of pectins [2]. During synthesis, HG can be O-acetylated at O-2 or O-3 and/or methylesterified at C-6 carboxyl, before being exported at the cell wall with a degree of methylesterification (DM) of ~80% and a degree of acetylation (DA) of ~5-10%, depending on species [3]. At the wall, HG chains can be modified by different enzyme families, including pectin acetyltransferase (PAEs; EC 3.1.1.6), pectin methylesterases (PMEs; CE8, EC 3.1.1.11), polygalacturonases (PGs; GH28, EC 3.2.1.15, EC 3.2.1.67, EC 3.2.1.82), and pectin lyases-like (PLLs), which comprise pectate lyases (PLs; EC 4.2.2.2) and pectin lyase (PNLs, EC 4.2.2.10). All these enzymes are produced by plants to fine-tune pectin during development [4–8], but they are also secreted by most phytopathogenic bacteria and fungi during plant infection [9–13]. PMEs and PAE hydrolyse the O6-ester and O2-acetylated linkages, respectively, leading to a higher susceptibility of HG to PG- and PLL-mediated degradation [14]. PLL are pectolytic enzymes that cleave HG via a β -elimination mechanism leading to the formation of an unsaturated C4-C5 bond [15], and can be divided into two subfamilies depending on their biochemical specificities: i) PLs have a high affinity for non- or low-methylesterified pectins and an optimal pH near 8.5. Their activity requires Ca^{2+} ions. ii) PNLs are most active on high-DM pectins at acidic pH values [16]. Both type of enzymes can degrade HG chains, and release oligogalacturonides (OGs), but their mode of action can differ. For PLs both endo and exo modes of action have been described, while only endo-PNL have been characterised so far [17]. For the latter, it was notably shown that endo-PNLs from *B. fuckeliana*, *A. parasiticus* and *Aspergillus* sp., can first release OGs with degrees of polymerisation (DP) 5–7, that are subsequently used as substrates, generating OGs of DP3 and DP4 as end-products [9,18,19]. Despite having the same DP, the final products can differ in their degrees and patterns of methylesterification and acetylation (DM/DA) depending on the enzymes' specificities; implying potential differences in substrate binding, and therefore in PLLs fine structures. Several crystallographic structures of bacterial and fungal PLL have been reported [15,20–24]. Overall, the PLL fold resemble that of published PME, PGs and

rhamnogalacturonan lyases [25–27], and is composed of three parallel β -sheets forming a right-handed parallel β -helix. The three β -sheets are called PB1, PB2 and PB3 and the turns connecting them T1, T2 and T3 [28]. The active site features three Asp, localized on the PB1 β -sheet and, in the case of PLs, accommodates Ca^{2+} [29]. In PNLs, Ca^{2+} is, on the other hand, replaced by Asp [15]. Additionally, the PL binding site is dominated by charged acidic and basic residues (Gln, Lys, Arg) which can accommodate negatively charged pectate substrates. In contrast, the PNL binding site is dominated by aromatic residues [15,29], which have less affinity for lowly methylesterified pectins. These differences in structure could translate into distinct enzyme dynamics when in complex with substrates of varying degrees of methylesterification.

In fungi, PLLs are encoded by large multigenic families which are expressed during infection. *Verticillium dahliae* Kleb., a soil-borne vascular fungus, targets a large number of plant species, causing Verticillium wilt disease to become widespread among fiber flax, with detrimental effects to fiber quality [30–32]. *V. dahliae* infects plants by piercing the root surface using hyphae. When Verticillium reaches the vascular tissue, hyphae start to bud and form conidia which progress in xylem vessels before germinating and penetrating adjacent vessel elements to start another infection cycle. [33,34]. During its colonization and proliferation Verticillium produces and secretes a number of pectinolytic enzymes, including thirteen PLLs. Considering the role of PLLs in determining pathogenicity, it is of paramount importance to determine their biochemical and structural properties [30,35,36]. This could allow engineering novel strategies to control or inhibit, the pathogen's pectinolytic arsenal. For this purpose, we characterized, via combined experimental and computational approaches, one *V. dahliae* PLL (VdPelB, VDAG_04718) after its heterologous expression in *P. pastoris*. The obtention of the 3D structure of VdPelB after X-ray diffraction and the analysis of enzyme dynamics when in complex with substrates of distinct DM, allowed the identification of the residues favouring pectate lyase (PL) activity. Experiments confirmed the importance of these residues in mediating PL activity showing that VdPelB is a *bona fide* PL, that releases peculiar OG as compared to previously characterized PLs. More importantly, the OGs released from roots of *Verticillium*-partially tolerant and sensitive flax cultivars differed, paving the way for the identification of VdPelB-mediated OGs that can trigger plant defence mechanisms.

2. Material and methods

2.1. Bioinformatical analysis

Verticillium dahliae PLL sequences were retrieved using available genome database (ftp.broadinstitute.org/). SignalP-5.0 Server (<http://www.cbs.dtu.dk/services/SignalP/>) was used for identifying putative signal peptide and putative glycosylation sites were predicted using NetOGlyc 4.0 Server (<http://www.cbs.dtu.dk/services/NetOGlyc/>). Sequences were aligned and phylogenetic analysis was carried out using MEGA multiple sequence alignment program (<https://www.megasoftware.net/>). Homology models used for molecular replacement were created using I-TASSER structure prediction software (<https://zhanglab.ccmb.med.umich.edu/I-TASSER/>) and UCSF Chimera (<http://www.cgl.ucsf.edu/chimera/>) was used for creation of graphics.

2.2. Fungal strain and growth

V.dahliae was isolated from CALIRA company flax test fields (Martainneville, France) and was kindly provided by Linéa-Semences company (Grandvilliers, France). Fungus was grown as described in Safran et al. [37]. Briefly, fungus was grown in polygalacturonic acid sodium salt (PGA, P3850, Sigma) at 10 g.L⁻¹ and in pectin methylesterified potassium salt from citrus fruit (55–70% DM, P9436, Sigma) solutions to induce expression of *PLLs*. After 15 days of growth in dark conditions at 25°C and 80 rpm shaking, mycelium was collected and filtered under vacuum using Buchner flask. Collected mycelium was frozen in liquid nitrogen, lyophilized and ground. Isolation of RNA and cDNA synthesis was realized as previously described in Lemaire et al.[38].

2.3. Cloning, heterologous expression and purification of VdPelB

V. dahliae PelB gene (VdPelB, UNIPROT: G2X3Y1, GenBank: EGY23280.1, *VDAG_04718*) consists of 1 single exon of 1002 bp length. The coding sequence, minus the nucleotides encoding signal peptide, was PCR-amplified using cDNA and gene-specific primers while VdPelB mutants were generated by PCR mutagenesis, a method for generating site-directed mutagenesis, using specific primers carrying mutations (Table S1). Cloning of the PCR-amplified sequences in pPICZαB, sequencing and expression in *P. pastoris* heterologous expression system as well as purification steps were done as previously described in Safran et al. [37].

2.4. Crystallization of VdPelB

Pectate lyase VdPelB was concentrated at 10 mg.mL⁻¹ in Tris-HCl pH-7.5 buffer. Crystallization conditions were screened using the sitting-drop vapor-diffusion method. Pectate

lyase VdPelB (100 nL) was mixed with an equal volume of precipitant (1:1) using Mosquito robot (STP Labtech). The crystals that resulted in best diffraction data were obtained with 0.1 M MIB (Malonic acid, Imidazole, Boric acid system) at pH8.0, with 25 % PEG 1500 as the precipitant (condition B5 from the PACT premier kit, Molecular Dimensions, Sheffield, UK) after 1 month. Optimization was realized using the hanging drop vapor-diffusion method forming the drop by mixing 1 μ L of precipitant solution with 1 μ L of the enzyme. The crystals (120 μ m x 30 μ m) were cryo-protected by increasing PEG 1500 concentration to 35%, before mounting them in a loop and flash-cooling them in liquid nitrogen.

2.5. VdPelB X-ray data collection and processing

X-ray diffraction data were collected at the PROXIMA-2a beamline of the Soleil synchrotron (Saint Aubin, France), at a temperature of -173°C using an EIGER 9M detector (Dectris). Upon a first data collection to 1.3 Å resolution, three more data sets were collected from the same crystal in order to obtain a complete data set. Thereby the kappa angle was tilted once to 30°, once to 60° and finally a helical data set was collected at 1.2 Å resolution. The reflections of each data set were indexed and integrated using XDS [39], scaled and merged using XSCALE [40]. The VdPelB crystal has a primitive monoclinic lattice in the P 1 2₁ 1 space group, with two molecules contained in asymmetric unit [41].

2.6. Structure solution and refinement

The structure of VdPelB was solved by molecular replacement using *Phaser* [42]. The data were phased using pectate lyase BsPelA (PDB: 3VMV, Uniprot D0VP31), as a search model [43]. Model was build using *Autobuild* and refined using *Refine* from PHENIX suite [44]. The model was iteratively improved with *Coot* [45] and *Refine*. The final structure for VdPelB has been deposited in the Protein Data Bank (PDB) as entry 7BBV.

2.7. VdPelB biochemical characterization

Pierce BCA Protein Assay Kit (Thermo Fisher Scientific, Waltham, Massachusetts, United States) was used to determine the protein concentration, with Bovine Serum Albumin (A7906, Sigma) as a standard. Deglycosylation was performed using Peptide-N-Glycosidase F (PNGase F) at 37 °C for one hour according to the supplier's protocol (New England Biolabs, Hitchin, UK). Enzyme purity and molecular weight were estimated using a 12% SDS-PAGE and mini-PROTEAN 3 system (BioRad, Hercules, California, United States). Gels were stained using

PageBlue Protein Staining Solution (Thermo Fisher Scientific) according to the manufacturer's protocol.

The substrate specificity of VdPelB was determined using PGA (81325, Sigma) and citrus pectin of various DM: 20–34% (P9311, Sigma), 55–70% (P9436, Sigma) and >85% (P9561, Sigma), with 0.5 μ M CaCl₂ or 5 μ M EDTA (Sigma) final concentrations. Enzyme activity was measured by monitoring the increase in optical density at 235 nm due to formation of unsaturated uronide product using UV/VIS Spectrophotometer (PowerWave Xs2, BioTek, France) during 60 min. The optimum temperature was determined by incubating the enzymatic reaction between 20 and 70°C for 12 min using PGA as a substrate (0.4%, w/v). The optimum pH was determined between pH5.0 and 10.0 using sodium phosphate (NaP, pH5.0 to 7.0) and Tris-HCl buffer (pH7.0 to 10.0) and 0.4% (w/v) PGA as a substrate. VdPelB and VdPelB-G125R kinetic parameters, K_m , V_{max} and k_{cat} , were calculated using GraphPad Prism (8.4.2). The reactions were performed using 0.75 to 15 mg.mL⁻¹ PGA concentrations at 50 mM Tris-HCl pH8.0 (native) and 9.0 (G125R) during 12 min at 35°C. All experiments were realized in triplicate. The statistical analysis was done using the Welch's t-test.

2.8. Digestion of commercial pectins and released OGs profiling

OGs released after digestions by recombinant VdPelB or commercially available Aspergillus PL (named AsPel) were identified as described in Voxeur et al., 2019 [9], using a novel in-house OGs library. Briefly, DM 20–34% (P9311, Sigma) and sugar beet pectin with DM 42% and degree of acetylation (DA) 31% (CP Kelco, Atlanta, United States) were prepared at 0.4 % (w/v) final concentration diluted in 50 mM Tris-HCl buffer (pH8.0) and incubated with either VdPelB or AsPel (E-PCLYAN, Megazyme). For each substrate, enzyme concentrations were adjusted to have enzymes at iso-activities (Table S2). For each substrate two dilutions, were used for analysing OGs released in early, VdPelB-2 and AsPel-2, and late phase, VdPelB-1 and AsPel-1, of digestions. Digestions were performed overnight. Non-digested pectins were pelleted by centrifugation and the supernatant dried in a speed vacuum concentrator (Concentrator plus, Eppendorf, Hamburg, Germany). Separation of OGs was done as previously described using an ACQUITY UPLC Protein BEH SEC column (125Å, 1.7 μ m, 4.6 mm x 300 mm) at a flow rate of 0.4 ml min⁻¹. MS detection was performed using an ACQUITY UPLC H-Class system coupled to a SYNAPT G2-Si-Q-TOF hybrid quadrupole time-of-flight instrument (Waters) equipped with an electrospray ionization (ESI) source (Z-spray) and an additional sprayer for the reference compound (lock spray). Samples were analysed by ESI-high-resolution MS (HRMS) and MS/MS in negative ionization mode. Data

acquisition and processing were performed with MASSLYNX software (v.4.1; Waters). [9,37,46].

The intensities were defined as the area under the curve, for each OG. Peak areas were clustered by hierarchical clustering with complete linkage on the euclidian distance matrix and visualized in the heatmap-package using R version 3.6.0. The statistical analysis was done using the Welch's t-test.

2.9. Molecular Dynamics simulations

Two sets of molecular dynamics (MD) simulations were conducted on the VdPelB protein: one in complex with a fully non-methylesterified polygalacturonate deca-saccharide, and the other with a fully methylesterified polygalacturonate deca-saccharide. Parameters specified by the AMBER14SB_parmbsc1 forcefield [47] were used to create the molecular topologies of the complexes. Each complex was set up in a cubic box with solute-box distances of 1.0 nm and solvated with water molecules specific to the TIP3P water model [48]. Na⁺ and Cl⁻ ions were added to neutralise the system's net charge and reach a salt concentration of 0.165 M. Using a steep-descent algorithm with a step size of 0.01, energy minimisation was performed to resolve clashes between particles, with convergence being established at a particle-particle force of 1000 kJ mol⁻¹ nm⁻¹. Particle-particle forces were calculated by considering van der Waals and electrostatic interactions occurring up to 1.0 nm, as well as long-range electrostatics treated in the Fourier space using the Particle Mesh Ewald (PME) summation method. Solvent equilibration was attained post minimisation in two stages: through the NVT and NPT ensembles, to reach constant temperature and pressure, respectively. Equilibration of the solvent under the NVT ensemble was conducted for 1 ns, integrating the equation of motion at a time step of 2 fs. The target reference temperature was 310.15 K, coupled every 0.1 ps using the V-rescale thermostat3. Based on the Maxwell-Boltzmann distribution [49] at 310.15 K, random velocities were then assigned to each particle in the system. Finally, solvent equilibration under the NPT ensemble was conducted for 1 ns, continuing from the last step of the previous equilibration, in terms of particle coordinates and velocities, at a reference temperature of 310.15 K, coupled every 0.1 ps using the V-rescale thermostat [50]. Here, pressure coupling was isotropically coupled every 2.0 ps, at 1 bar, using the Parrinello-Rahman barostat [51]. Particle-particle interactions were computed by constructing pair lists using the Verlet scheme. Short-range van der Waals and electrostatic interactions sampled through a Coulomb potential, were calculated at a cutoff of 1.0 nm. The PME algorithm [52] was used to compute long-range electrostatic interactions beyond this cut-off in the Fourier space, utilising

a Fourier grid spacing of 0.16 and a cubic B-spline interpolation level at 4. The simulations were then performed on in-house machines, using GROMACS (Groningen Machine for Chemical Simulations) version 2021.37. Each set of simulations were run for 150 ns each, at a time step of 2 fs, with molecular dynamics trajectories written out every 10 ps. Simulations were replicated 7 times for a total production run time of 1.05 μ s per complex. Replicates differed with respect to the random particle velocity sets computed under the NVT ensemble. For analysis, the first 50 ns of each production run were discarded as equilibration time. In-house Python 3 scripts implemented using Jupyter notebooks [53] were used to carry out analyses. Figures were created and rendered with Matplotlib [54] and VMD (Visual Molecular Dynamics)[55].

3. Results and Discussion

3.1. Sequence and phylogeny analysis

In addition to 9 polygalacturonases (PGs) and 4 pectin methylesterases (PMEs), *V. dahliae* encodes 30 putative endo-pectate lyases (EC 4.2.2.2), exo-pectate lyases (EC 4.2.2.9), endo-pectin lyases (EC 4.2.2.10), belonging to PL1-PL3 and PL9 families, respectively [56]. For hierarchical clustering of *Verticilium*'s sequences with other PLLs, fifty-one amino acid sequences encoding putative PLLs, belonging to bacteria, fungi and plants were aligned and a phylogenetic tree was built. Different clades can be distinguished. (**Fig. 1**). *V. dahliae* PelB (VdPelB, VDAG_04718) clustered with VDAG_05344 (59.68% sequence identity) with close relations to VDAG_05402 (57.19% sequence identity) and VDAG_07566 (56.95% sequence identity). VDAG_05402 and VDAG_05344, that are found in the protein secretome, have orthologs in *V. alfalfa* (VDBG_07839 and VDBG_10041), which were shown to possess putative lyase activity [36,57]. Plant PLLs from *A. thaliana* (AtPLL21, AtPLL15 and AtPLL18) formed a separate clade with close connections to *A. denitrificans* (AdVexL), a PLL homologue [58]. *D. dadanti* (DdPelI), *Bacillus Sp.* KSM-P15 (BsPel-P15) and *C. bescii* (CbPel1854) formed separate clades similarly to *B. subtilis* and *D. dadantii* PLs (BsPel, DdPelA and DdPelC). The clade corresponding to PNLs consisting of *A. tubingensis* PelA (AtPelA), *A. niger* (AnPelA, AnPnlA and AnPelB), *A. luchuensis* AlPelB was closely related to VDAG_07238 and VDAG_08154 which were indeed annotated as putative PNLs [18,56]. VDAG_06155 was previously named VdPel1 and previously characterized as a pectate lyase [35].

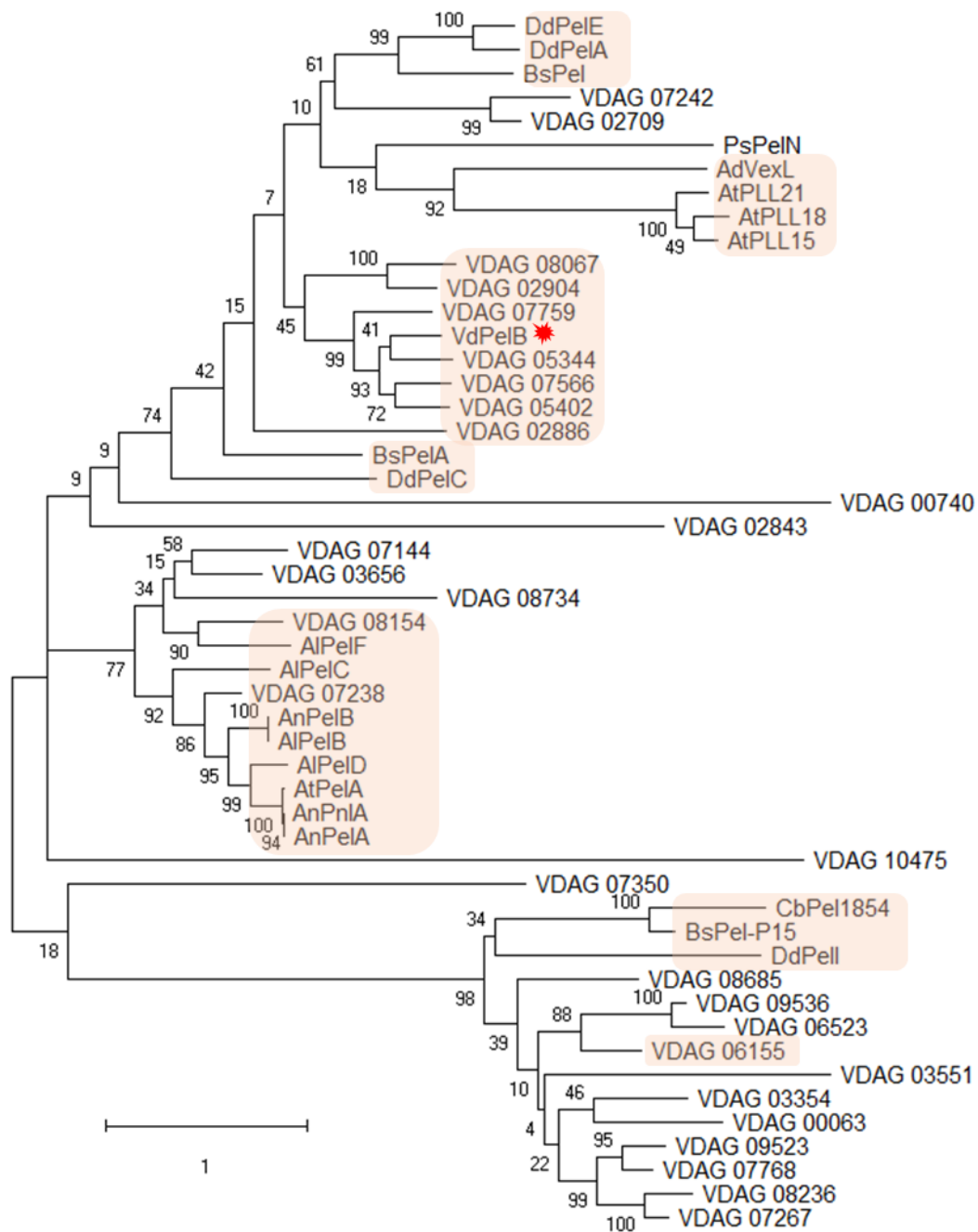


Fig. 1. Phylogenetic analysis of *V. dahliae* pectate lyase VdPelB with selected PLLs

Phylogenetic tree representing *V. dahliae* VdPelB (VDAG_04718, G2X3Y1, red star) amino acid sequence in comparison with PLLs from *Verticillium* [VDAG_00740 (G2WQU8), VDAG_02904 (G2WXC5), VDAG_05344 (G2X628), VDAG_07242 (G2XBA4), VDAG_07759 (G2XC77), VDAG_02709 (G2WWT0), VDAG_02843 (G2WX64), VDAG_02886 (G2WXA7), VDAG_03656 (G2X1P5), VDAG_05402 (G2X597), VDAG_07144 (G2X9U9), VDAG_07238 (G2XBA0), VDAG_07566 (G2XBY8), VDAG_08067 (G2XD35), VDAG_08154 (G2XD2C), VDAG_08734 (G2XF02), VDAG_10475 (G2XJZ3), VDAG_03354 (G2WZB2), VDAG_03551 (G2WZV9), VDAG_07267 (G2XBC9), VDAG_07768 (G2XC86), VDAG_08236 (G2XDK4), VDAG_06155 (G2X8L4), VDAG_06523 (G2X7R5), VDAG_08685 (G2XEV0), VDAG_09523 (G2XH91), VDAG_09536 (G2XHA4), VDAG_00063 (G2WR80), VDAG_07350 (G2XGG7)], *Arabidopsis thaliana* [AtPLL15 (At5g63180), AtPLL18 (At3g27400), AtPLL21 (At5g48900)], *Dickeya dadanti* [DdPelA (P0C1A2), DdPelC

(P11073), DdPelE (P04960), DdPelI (O50325)], *Bacillus subtilis* [BsPel (P39116)], *Bacillus Sp.* KSM-P15 [BsPel-P15 (Q9RHW0)], *Bacillus sp.* N16-5 [BsPelA (D0VP31)], *Aspergillus niger* AnPelA [(Q01172), AnPelB (Q00205), AnPnIA (A2R3I1)], *Achromobacter denitrificans* [AdVexL (A0A160EBC2)], *Aspergillus tubingensis* [AtPelA (A0A100IK89)], *Aspergillus luchuensis* [AlPelB (G7Y0I4)], *Acidovorax citrulli* [AcPel343, (A1TSQ3)], *Paenibacillus sp.* 0602 [PsPelN (W8CR80)], *Caldicellulosiruptor bescii* [CbPel1854 (B9MKT4)]. Maximum likelihood tree was constructed with 1000 bootstrap replicates. Most important clades are indicated in orange squares while VdPelB is marked by red star. Amino acids sequences were retrieved from Uniprot and TAIR.

3.2. Expression and purification of VdPelB

Since the expression was performed using the pPICZαB vector with the yeast alpha-factor directing VdPelB secretion in the culture media, the protein could be easily recovered and purified. The protein is composed of 343 amino acids, including the 6xHIS tag at the C-terminus used for affinity chromatography purification. After purification, VdPelB purity was assessed using SDS-PAGE with an apparent molecular mass of ~38 kDa (**Fig. S1**), higher than what was predicted on the basis of the amino acid sequence (33.8 kDa). However, this shift is likely to correspond to the two tags that are fused to the protein (His and C-myc) and to the presence of 19 putative O-glycosylation sites, as predicted by NetOGlyc 4.0 Server.

3.3. VdPelB has a right-handed parallel β-helix fold

Pectate lyase VdPelB was crystallized and its 3D structure was determined using X-ray diffraction. VdPelB crystallized in monoclinic P 1 2₁ 1 asymmetric unit. Four data sets, collected from the same crystal at 1.2 Å resolution, were integrated, scaled and merged. There are two molecules in the asymmetric units: chains A and B are highly similar with a Cα root mean square deviation (rmsd) value of 0.227 Å (**Fig. S2A**). The VdPelB structure consists of 298 amino acids with 18 at the N-terminus and 27 amino acids at the C-terminus that were not resolved because of poor electron densities, while overall electron densities were well defined. While no N-glycosylation sites could be revealed on the VdPelB structure, six O-glycosylation sites carrying mannose are visible for each molecule: T22, T44, T45, T46, S48 and T54, in accordance with the shift in size previously observed (**Fig. S2A**). During data acquisition no heating of the crystal was observed, as shown by low B factors and good occupancies (**Fig. S3A and B**). The final models' geometry, processing and refinement statistics are summarized (**Table 1**). The VdPelB's structure has been deposited in the Protein Data Bank as entry 7BBV.

Table 1. Data collection, processing and refinement for VdPelB

Characteristics	VdPeIB
Data collection	
Diffraction source	PROXIMA2
Wavelength (Å)	0.980
Temperature (°C)	-100.15
Detector	DECTRIS EIGER X 9M
Crystal-to-detector distance (mm)	115.02
Rotation range per image (°)	0.1
Total rotation range (°)	360
Crystal data	
Space group	P1 21 1
<i>a</i> , <i>b</i> , <i>c</i> (Å)	60.89, 59.69, 93.87
α , β , γ , (°)	90.00, 96.35, 90.00
Subunits per asymmetric unit	2
Data statistics	
Resolution range (Å)	60.52-1.2 (1.243 - 1.2)
Total No. of reflection	8536161 (112943)
No. of unique reflection	207497 (19983)
No. of reflections, test set	10362 (1003)
R_{merge} (%)	0.1036 (4.709)
Completeness (%)	99.30 (96.11)
$\langle I/\sigma(I) \rangle$	18.03 (0.22)
Multiplicity	41.1 (5.6)
$CC_{1/2}$ (%)	1 (0.116)
Refinement	
$R_{\text{crys}}/R_{\text{free}}$ (%)	17.3 / 19.7
Average B – factor (Å ²)	33.18
No. of non-H atoms	
Protein	9097
Ion	2
Ligand	234
Water	1052
Total	10385
R.m.s. deviations	
Bonds (Å)	0.013
Angles (°)	1.39
Ramachandran plot	
Most favoured (%)	95.1
Allowed (%)	4.90
Outlier (%)	-

The VdPelB has a right-handed parallel β -helix fold which is common in pectinases [59]. The β -helix is formed by three parallel β -sheets - PB1, PB2 and PB3 which contain 7, 10 and 8 β -strands, respectively. Turns connecting the PB1-PB2, PB2-PB3 and PB3-PB1 β -sheets are named T1-turns, T2-turns and T3-turns, respectively, according to Yoder and Jurnak (**Fig. 2A, S4A and B**) [60]. T1-turns consist of 2-14 amino acids and builds the loop around the active site on the C-terminus. T2 turns mostly consist of 2 amino acids with Asn being one of the predominant amino acid, forming an N-ladder with the exception of an N245T mutation in VdPelB [15,20]. Pectate lyase VdPelB has a α -helix on N-terminus end that shields the hydrophobic core and is commonly conserved in PLs and PGs [15,61], while the C-terminus end is also protected by tail-like structure carrying one α -helix. Interestingly N- and C- terminus tails pack against PB2 (**Fig. 2A**). There are only two Cys (C25 and C137) that do not form a disulphide bridge.

Sequence and structural alignments show that VdPelB belong to the PL1 family. The VdPelB shares the highest structural similarity with BsPelA (PDB: 3VMV), with 30.06% sequence identity and C α rmsd of 1.202 Å. The second-best structural alignment was with DdPelC (PDB: 1AIR) with 24.20% identity and C α rmsd of 1.453 Å [43,62]. Both of these structures lack the long T3 loop described in *A.niger* pectate lyase (AnPelA, PDB: 1IDJ, **Fig. S2B and S5**) [15]. The putative active site is positioned between the T3 and T2 loops (**Fig. 2A and B**).

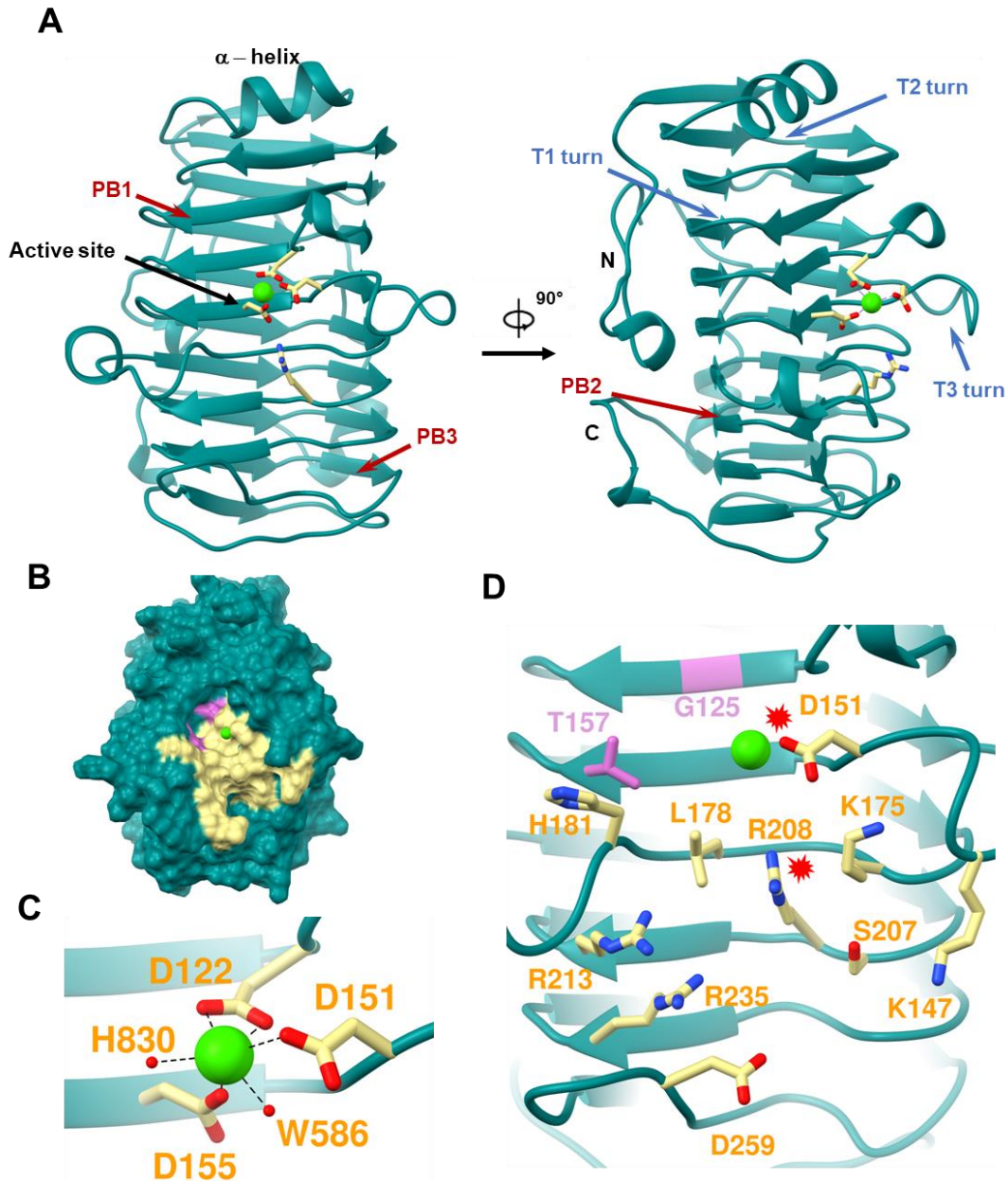


Fig. 2. Structure determination of VdPelB

A) Ribbon diagram of VdPelB crystalized in P1 21 1 space group. VdPelB is a right-handed parallel β helical structure consisting of β strands (red arrows) and turns (blue arrows). VdPelB active site's amino acids are yellow-colored while Ca atom is green. B) Surface representation of VdPelB binding groove. C) Active site of VdPelB highlighting conserved amino acids and atoms interacting with Ca. D) Structure of VdPelB binding groove highlighting amino acids involved in the interaction (yellow) and amino acids not of previously characterized in PLs (plum). Red stars indicate amino acids from the active site.

3.4. Active site harbours Ca^{2+} that is involved in catalysis

The VdPelB active site is well conserved, harbouring strictly conserved acidic and basic amino acids that are required for Ca^{2+} binding. Previously reported structures showed that two Asp (D122 and D155) and one Arg (R208) in VdPelB, are conserved, while D151 can be

mutated to Glu, or Arg in PNLs (**Fig. 2C and D**, Mayans et al., 1997). Other conserved amino acids in VdPelB include K175 and R213, with K175 being responsible for binding the carboxyl oxygen while R213 hydrogen bonds to C-2 and C-3 of GalA (**Fig. 2D**) [29,63]. Mutating any of these amino acids leads to decreased enzyme activity [64]. In VdPelB, Ca^{2+} ion is directly coordinated by D122, two carboxyl oxygen, D151, D155 and two water molecules (W568 and W830, **Fig. 2C**). In addition, mutation of D122T (VdPelB numbering) in BsPelA, is responsible for reduced affinity for Ca^{2+} [43]. In the catalytic mechanism, Ca^{2+} is directly involved in acidification of the proton absorption from C5 and elimination of group from C4, generating an unsaturated product. R208 act as a base, similarly to the hydrolysis in the reaction mechanism of the GH28 family [65,66].

3.5. Structural analysis of VdPelB suggests a PL activity mediated by peculiar specificities

The VdPelB binding groove comprises a number of basic and acidic amino acids including K147, D151, K175 L178, H181, S207, R208, R213 and R235 and D259 (**Fig. 2C and D**), that have previously been shown to be characteristics of PLs. This would suggest an enzyme activity on low DM pectins as amino acids positioned at the binding groove were indeed shown to differ between PNL and PL [15,21,24,29]. These amino acids are indeed mutated in Arg, Trp, Tyr, Gln and Gly in PNLs, which, by reducing their charge, would favour higher affinity for highly methylesterified pectins [15,29]. In VdPelB, as the T3 loop is missing, there are no equivalent to PNL specific W66, W81, W85, W151 amino acids (**Fig. S2B**, AnPelA numbering) and, moreover, W212 and W215 are replaced by K175 and L178 in VdPelB. In BsPelA and DdPelC, these amino acids are replaced by K177/K190 and L180/L193, highlighting the high conservation of amino acids in VdPelB/BsPelA/DdPelC and subsequently in PLs. When a DP4 ligand from DdPelC crystal structure is superimposed to VdPelB, hydrogen bonds and van der Waals interactions are visible with the above-mentioned amino acids (**Fig. S6**) [15,21,29,43].

Interestingly, despite this rather conserved PL-related binding groove, VdPelB harbors, in the vicinity of the active site, G125 and T157 that are not present in the well characterized DdPelC [62]. At these positions, DdPelC, which was shown to be a *bona fide* PL with a high activity on polygalacturonic acid and alkaline pH with Ca^{2+} -dependency, harbours Arg and Lys [15,63,67]. In that respect, the presence of G125 and T157 in VdPelB is similar to that identified in *Bacillus sp.* Pel-22, Pel-66, and Pel-90 and *Bacillus sp.* which showed activity on both PGA and high methylesterified pectins (**Fig. 2B and D**) [22,68,69]. The DdPelC amino acids being

overall positively charged, could explain the binding preference towards non-methylesterified, negatively charged substrates in the vicinity of the active site. In contrast, considering the size of Gly and Thr, they would sterically accommodate the increased size of methylesterified substrate. G125 and T157 could therefore account for a potential dual activity of VdPelB. Moreover, in previously characterized PLs and PNLs there is the presence of a small amino acids, Ser or Ala that replace H181. While H181 interacts directly with the substrate these amino acids do not provide this interaction instead the primary Ca^{2+} in the active site induces a substrate conformation that could be recognized by PLs [43]. Finally, L178 is positioned between the catalytic Ca^{2+} and R208 and is involved in substrate binding making it a perfect candidate to assess its importance (**Fig. 2D**).

3.6. Molecular dynamic simulations show higher dynamics of VdPelB in complex with methylesterified substrates

To determine how the structure of VdPelB and the observed differences in amino-acidic composition might influence the affinity with differently methylated substrates, we performed MD simulations on VdPelB in complex with either a non-methylesterified or fully methylesterified decasaccharides, which are able to occupy the entire binding groove (**Fig. 3**). MD simulations show substantially differential dynamic profiles for oligosaccharides with and without methylesterification. Expectedly, the dynamics is lowest in proximity of the catalytic subsite (+1) and increases consistently towards both the reducing and non-reducing ends of the substrate: with the highest dynamics found at the non-reducing end (**Fig. 3A**).

A quantitative estimation of the substrate dynamics was obtained by monitoring the root square mean fluctuations (RMSF) of each sugar residue and shows that a polygalacturonate substrate associates more stably in the subsites of the binding groove placed towards the sugar's reducing end (subsites -1 to -5), with differences between methylesterified and non-methylesterified substrates approaching the obtained standard deviation. In some cases, the dynamics reversed for non-methylesterified sugars, with a higher RMSF for de-methylesterified sugars towards the saccharide's reducing end (**Fig. 3B**). Moreover, and in line with an observed positively charged binding groove, non-methylesterified substrates retain a higher number of contacts than methylesterified sugars, with salt-bridges contributing the most to the observed differences (**Fig. 3C**).

We then additionally and specifically focused on the analysis of the interactions made by T157, which is nested with the -2 subsite of the binding groove. Simulations sample consistently

higher contacts formation between a methylesterified sugar docked in subsite -2 and T157, with distances shifted to lower values when compared to a non-methylesterified sugar (**Fig. 3D**), even the RMSF of non-methylesterified monosaccharides docked in subsite -2 experience on average, significantly lower dynamics. Altogether, the formation of a larger number of contacts between methylesterified saccharides docked in the -2 subsite suggests an active role of T157 in the binding of methylesterified chains: with the butanoic moiety of T157 engaging in hydrophobic interactions with the methyl-ester presented by methylesterified sugar units (**Fig. 3E**). While T157 is seen to have an active role in engaging with the substrate's hydrophobic moieties, an active role of G125, also in the same subsite, was not observed but it is plausible that the minimal size of G125 would increase the accommodability of methylesterified sugars in that position.

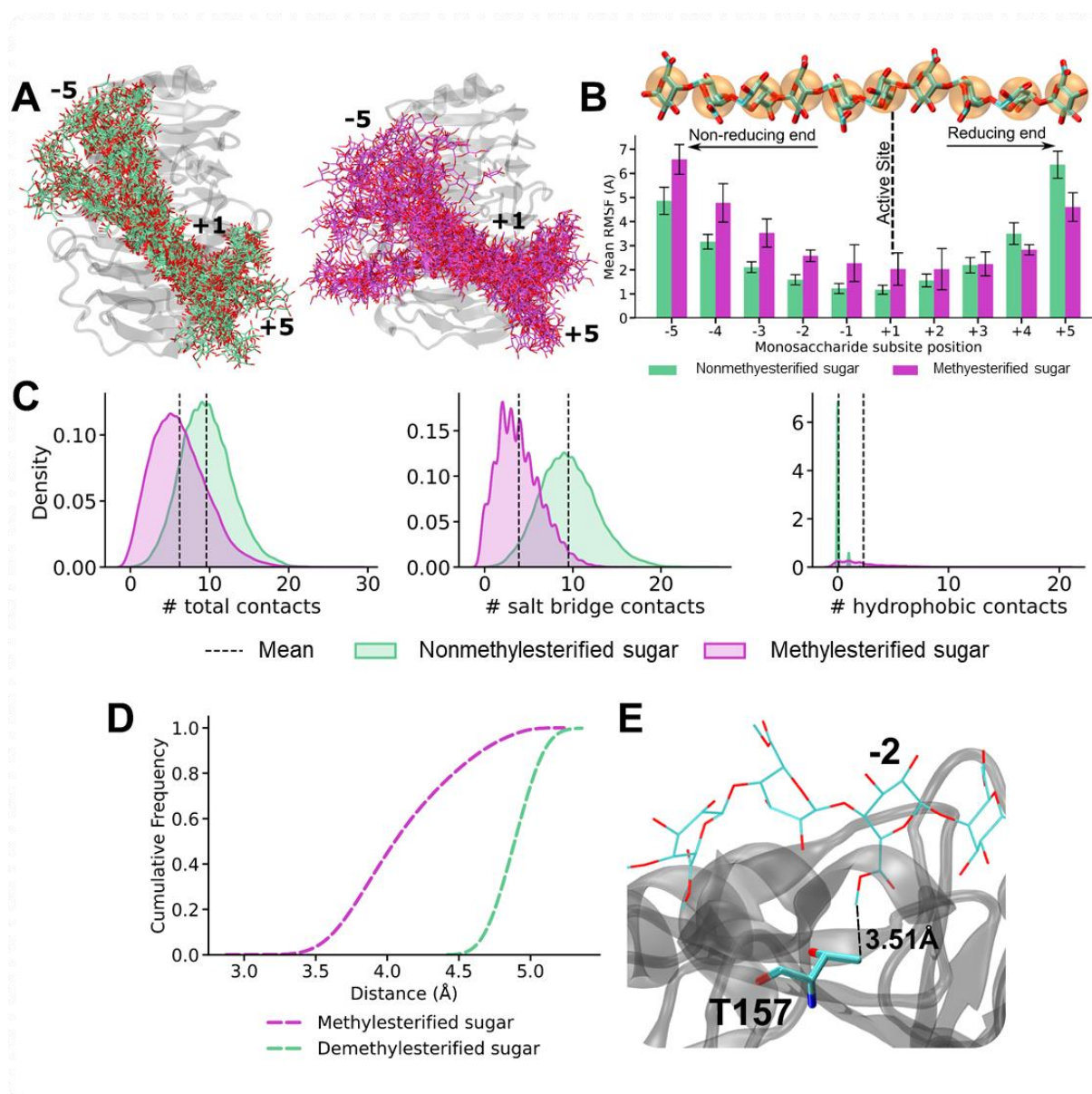


Fig. 3. VdPelB substrate dynamics in complex with fully de-methylesterified and methylesterified complex

A) Ensembles of non-methylesterified (green, left panel) and methylesterified (pink, right panel) HG deca-saccharide at every 100 frames for the simulated VdPelB complexes. Substrate within the enzymes' binding grooves are labelled from -5 (HG-non-reducing end) to +5 (HG-reducing end) B) RMSF of non-methylesterified (green) and methylesterified (pink) HG bound across the binding groove of VdPelB. Numbers indicate the beta sheet position from the active site. C) Analysis of the contacts between VdPelB and non-methylesterified (green) and methylesterified substrate (pink). D) Distance between T157K residue and the substrate residues plotted as cumulative frequency. E) T157 hydrophobic interaction with methyl-ester of methylesterified substrate.

3.7. Biochemical characterization of VdPelB

We first determined the activity of VdPelB by following the release of 4,5-unsaturated bonds which can be detected at 235 nm using UV spectrophotometer. The VdPelB activity was first tested for its dependency towards Ca^{2+} . Using a standard PL assay, with PGA as a substrate, an increase in the VdPelB activity was measured in presence of calcium. In contrast in presence of EDTA, used as a chelating agent, no activity was detected, confirming the calcium-dependent activity of the enzyme (**Fig. S7A**) [15]. Activity measured in absence of added CaCl_2 reflects the presence of calcium from the culture media that is bound to VdPelB during production, and previously identified in the 3D structure. We tested the effects of increasing CaCl_2 concentrations and showed that the maximum activity was already reached when using as low as 0.125 μM (**Fig. S7B**). To test the substrate-dependence of VdPelB, four substrates of increasing degrees of methyl-esterification were used. The VdPelB showed the highest activity on PGA, with less than 10% of the maximum activity measured on the three others substrates (**Fig. 4A**). This shows that, as inferred from above-mentioned structural and dynamical data, The VdPelB act mainly as a PL although it can still show residual activity on high DM pectins. Considering this, PGA was used as substrate to test the pH-dependence of the enzyme's activity in sodium acetate and Tris-HCl buffers (**Fig. 4A**). Pectate lyase VdPelB was most active at pH 8, with only a slight decrease in activity at pH 9 (93%). In contrast, the relative activity at pH 5-7 was close to null. The pH optimum for VdPelB was the same as *B. fuckeliana* Pel (pH 8) [70], close to that reported for *D. dadantii* PelN (pH 7.4) [71], but was lower to that measured for *B. clausii* Pel (pH 10.5) [72]. In contrast, the pH optimum was higher compared to five PNLs from *Aspergillus* sp. AaPelA (pH 6.1), AtPelA (pH 4.5), AtPelA (pH 6.4), AtPelD (pH 4.3) [18] and *A. parasiticus* Pel (pH 4) [19]. The optimum temperature assay showed that VdPelB was most active at 35°C (**Fig. S8**). The VdPelB appeared less heat-tolerant as compared to thermophilic PLLs reported from *Bacillus* sp. RN1 90°C [73], *B. clausii* Pel, 70°C [72], *B. subtilis* Pel168, 50°C [74]. However, its optimum temperature is in the range of that measured

for *X. campestris* Pel [75] and cold-active Pel1 from *M. eurypsychrophila* [76]. The lack of disulphide bridges previously shown in the structure could be responsible for the lower stability of the enzyme at high temperatures, in comparison with previously characterized PLs [77].

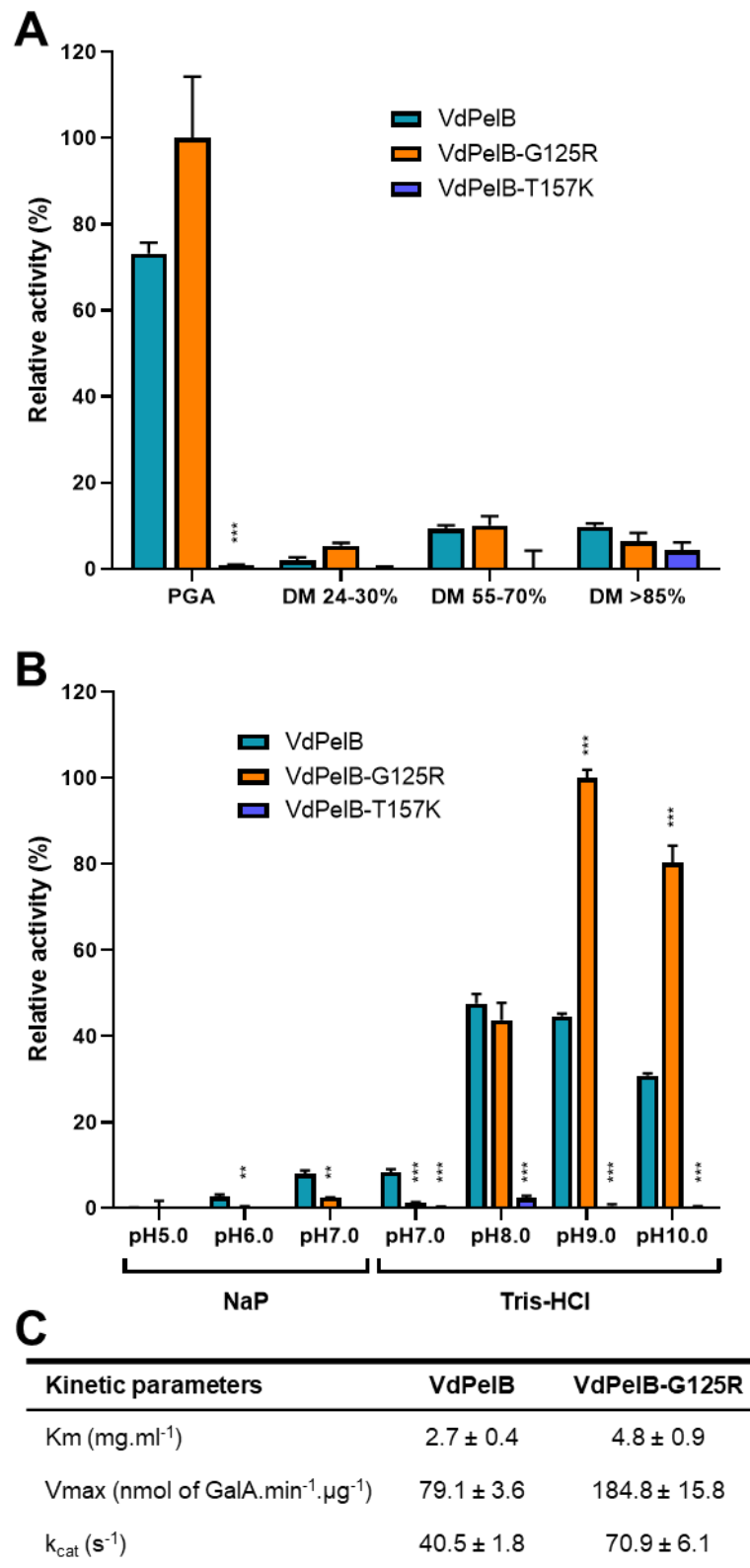


Fig. 4. Biochemical characterization of VdPelB

A) Substrate-dependence of VdPelB, G125R and T157K. The activities were measured after 12 min of incubation with PGA, pectins DM 24-30%, DM 55-70%, DM>85% with addition of Ca^{2+} at 35°C. B) pH-dependence of VdPelB, G125R and T157K activity. The activities were measured after 12 min of incubation with PGA in sodium phosphate (NaP) and Tris-HCl buffer at 35°C. Values correspond to means \pm SD of three replicates. Welch t-test comparing native with mutant VdPelB forms was used for statistical analysis. P value ***<0.001 and **<0.01. C) Determination of K_m , V_{max} and k_{cat} for ADPG2 and PGLR. Activity was assessed using various concentrations of PGA at 35°C and pH8.0 (VdPelB) and pH9.0 (VdPelB-G125R).

3.8. Mutation of specific amino-acids affects VdPelB activity

Considering the structure of VdPelB and our hypothesis related to the role of some amino acids in determining the mode of action of VdPelB, we generated mutated forms of the enzyme for five amino acids that likely to be involved in the catalytic mechanism and/or substrate binding: G125R, D151R, T157K, L178K and H181A. Enzymes were produced in *P. pastoris* and purified (**Fig. S1**). If the importance of some of these amino acids (i.e. D151 and H181) in the catalytic mechanism was previously shown for others PL [43,78], our study highlights the key role of some novel amino acids in the catalytic mechanism. The activities of mutants were tested at iso-quantities of wild-type enzyme. Surprisingly, the G125R mutant was 25% more active on PGA compared to the native enzyme and a shift in the optimum pH was observed (**Fig. 4A**). While native enzyme was most active at pH8.0 with slight decrease in activity at pH9.0, the activity of G125R mutant was approximately doubled at pH9.0 and pH10.0 (**Fig. 4B**). Both enzymes kept the same activity at pH8.0. Moreover, using PGA as a substrate, at 35°C, VdPelB and VdPelB-G125R differ slightly in their K_m (2.7 versus 4.8 mg.ml^{-1}), V_{max} (79.1 versus 184.8 $\text{nmol of GalA.min}^{-1}.\mu\text{g}^{-1}$) and k_{cat} (40.5 versus 70.9 s^{-1} , **Fig. 4C**). The substitution of Gly with Arg, present in a number of previously characterized PL could facilitate the interaction with the substrate and alkalization of the active site thanks to its physio-chemical properties [62]. Despite not knowing the exact rotamer orientation, the presence of Arg with high pK_a allowing strong hydrogen bonding and complex electrostatic interactions within the active site, could change the optimal pH by acting on the protonation state of amino acids involved in catalysis, as previously reported for a number of different enzymes [65,79–82]. The activity of T157K was closed to null when tested on PGA and different pHs. The introduction of a Lys, a chemically different and larger amino acids is likely to introduce steric clashes notably with H181 and L178 that are important for the activity. The H181A mutant lost much of its activity as there are less interaction/recognition with the substrate. No activity for D151R and L178K mutants were observed in line with the fact that D151 is an active site amino acid that binds the Ca^{2+} . Mutation of this amino-acid is negatively

impairing the functioning of the enzyme (**Fig. S7A and B**) [43,78]. We can hypothesize that L178K mutation positioned in between Ca²⁺, R208 and the substrate, induces specific substrate conformation that diminishes the direct interaction between the enzyme catalytic centre and the substrate, which translates to loss of activity (**Fig. S9**).

3.9. Identification of the OGs released by VdPelB from commercial and cell wall pectins

To further understand the specificity of VdPelB on different substrates, we performed LC-ESI-MS/MS to determine the profiles of digestion products (OGs) and to compare with that of commercially available *Aspergillus sp.* Pel (AsPel, **Fig. S10**). To be fully comparable, digestions were realized, for each substrate, at iso-activities for the two enzymes. On the basis of digestion profiles, we identified 48 OGs and created a dedicated library that was used for identification and integration of peaks (**Table S3**). MS² fragmentation allowed determining the structure of some of the OGs (**Fig. S11 and S12**). The OGs released by either of the enzymes mainly corresponded to 4,5-unsaturated OGs, which is in accordance with β -eliminating action of PLLs. When using pectins DM 20-34% and at low enzyme's concentration (VdPelB-2), VdPelB mainly released non-methylesterified OGs of high DP (GalA₅, GalA₆, GalA₇, GalA₈, GalA₉, GalA₁₀) that were subsequently hydrolysed when using more concentrated VdPelB (VdPelB-1, **Fig. 5A**). These digestion products strikingly differed to that generated by AsPel, that are methylesterified OGs of higher DP (GalA₄Me₁, GalA₄Me₂, GalA₅Me₁, GalA₆Me₂, GalA₁₁Me₃...), thus showing distinct enzymatic specificities. Altogether, these first results unequivocally shows both VdPelB and AsPel act as endo-PLs, but they differ in their processivity [9]. When using sugar beet pectins, that are known to be highly acetylated (DM 42%, DA 31%), VdPelB released acetylated OGs (GalA₂Ac₂, GalA₃Ac₁, GalA₄Ac₁, GalA₅Ac₁), while AsPel showed much lower activity and relative abundance of these OGs (**Fig. 5B**). Previous reports have shown that differences exist between PNL, in particular with regards to acetyl substitutions [18]. In contrast, AsPel released mainly methylesterified OGs (GalA₆Me₁, GalA₈Me₁, GalA₁₀Me₃...).

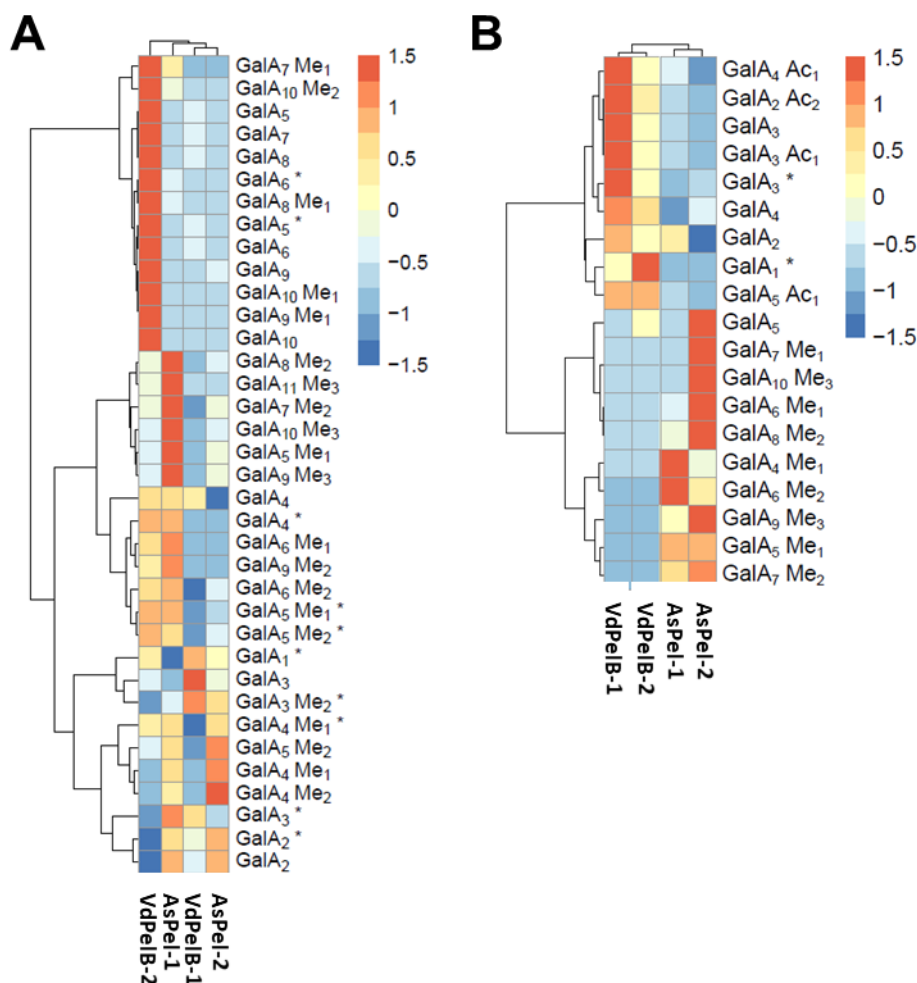


Fig. 5. Analysis of OGs produced by the action of VdPelB and AsPelI on pectins of various degrees of methylesterification and acetylation

OGs were separated by SEC and analysed by MS/MS. A) Pectins DM 24-30%. B) Sugar beet pectins (DM 42% DA 31%). Substrates were digested overnight at 40°C and pH8.0 using isoactivities of VdPelB and AsPel. Enzyme concentrations are stated in Table S2. Subscript numbers indicate the DP, DM and DA. * indicate non-unsaturated OGs. Values correspond to means of three replicates.

V. dahliae and closely related fungus from the same genus are known flax pathogens, where they use their enzymatic arsenal, that includes pectin degrading enzymes, for penetrating the host cell leading to infection [30]. To assess the potential role of VdPelB in flax pathogenicity we digested root cell walls from two flax cultivars, Evea (Verticillium-partially resistant), and Violin (Verticillium-susceptible), and we compared the OGs released. On root cell walls, VdPelB released mainly unsaturated OGs up to DP5 (**Fig. 6**). From similar starting root material, the OG total peak area detected was five times lower for Evea compared to Violin, suggesting that it is less susceptible to digestion by VdPelB. OGs released by VdPelB were mainly non-methylesterified but could be acetylated (GalA₂, GalA₃, GalA₃Ac₁, GalA₄,

GalA₄Ac₁ GalA₅, GalA₅Ac₁), and the abundance of GalA₃ and GalA₄ was five and fifteen times higher in Violin, respectively. These data together with that obtained from sugar beet pectins strongly suggests that VdPelB preference is for non-methylesterified and acetylated substrates. Our data suggest that cell wall structure differ between the two cultivars and that VdPelB could determine Verticillium pathogenicity thanks to a better degradation of the cell wall pectins of sensitive cultivars [37]. Similarly, VdPel1 was previously identified as virulence factor, where the deletion of this gene decreased virulence in tobacco, as compared with the wild-type Verticillium [35].

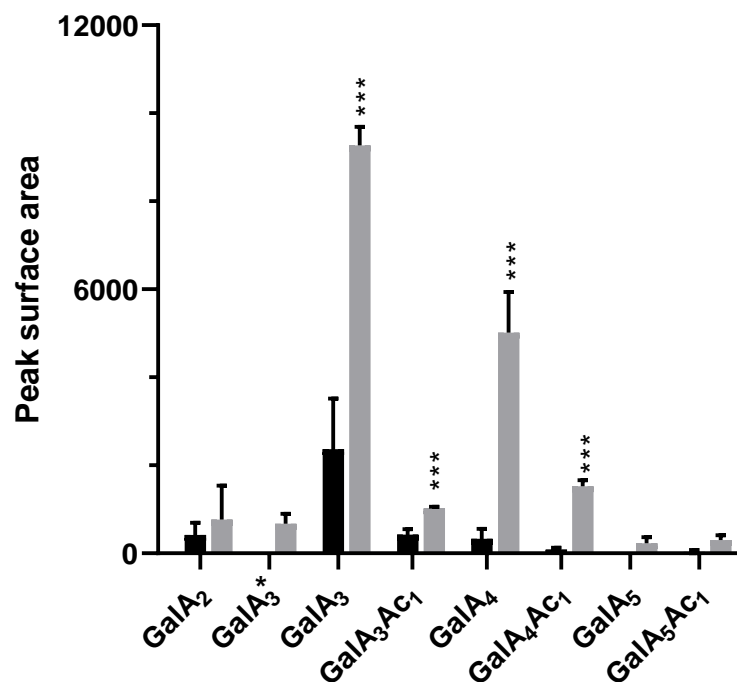


Fig. 6. Analysis of OGs released by VdPelB from flax roots.

The VdPelB was incubated overnight with roots from Évéa (spring flax, partially resistant to Verticillium wilt, black) and Violin (winter flax, more susceptible to Verticillium wilt, grey). Values correspond to means \pm SD of three replicates. Welch t-test comparing Évéa with Violin was used for statistical analysis. P value *** <0.001 . Subscript numbers indicate the DP, DM and DA.

4. Conclusion

We have characterized, by multidisciplinary approaches, a novel pectinolytic enzyme from *V.dahliae*, VdPelB, that belongs to the PLL family. The protein was crystallised and its 3D structure determined at a high resolution. Pectate lyase VdPelB showed a conserved

structure, with parallel β -sheet topology for PLs and the active site harboured three conserved Asp coordinating Ca^{2+} and Arg involved in the β -elimination mechanism. The binding groove of VdPelB reveals conserved amino acids that are characteristic of PLs, with MD simulations confirming the lower dynamics/higher affinity of the enzyme towards non-methylesterified pectins. As inferred from structural and dynamical analyses, VdPelB showed high activity on non-methylesterified substrates, with a maximum activity at pH8.0 and 35°C. The analysis of the structure led the identification, in the VdPelB, of peculiar amino acids that are normally present in PNL. In particular, G125R mutant increased activity on PGA and switch in pH optimum from 8.0 to 9.0 related to amino acids protonation. The analysis of the digestion products showed that VdPelB act as an endo enzyme and that it can release a large diversity of OGs with a preference for non-methylesterified and acetylated products. The OGs generated by VdPelB from pectins extracted from *Verticillium*-partially tolerant and *Verticillium*-sensitive flax cultivars showed that the enzyme could be a determinant of pathogenicity, since these OGs differ as a function of pectins' structure. Thus, our study now paves the way for generating, using dedicated enzymes, OG pools that could be used for protecting plants against phytopathogens.

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Author's contribution

Josip Safran: Conceptualization, Data curation, Formal analysis, Investigation, Methodology, Writing - original draft. **Vanessa Ung:** Data curation, Investigation, Methodology. **Julie Bouckaert:** Data curation, Investigation, Methodology. **Olivier Habrylo:** Formal analysis, Investigation, Methodology. **Roland Molinié:** Data curation, Formal analysis, Investigation, Methodology. **Jean-Xavier Fontaine:** Data curation, Formal analysis, Investigation, Methodology. **Adrien Lemaire:** Investigation, Methodology. **Aline Voxeur:** Investigation, Methodology. **Serge Pilard:** Investigation, Methodology. **Corinne Pau-Roblot** Conceptualization, Methodology. **Davide Mercadante:** Data curation, Methodology, Writing - review & editing. **Jérôme Pelloux:** Funding acquisition, Conceptualization, Writing - review & editing. **Fabien Sénéchal:** Conceptualization, Writing - review & editing.

Conflicts of interest

There are no conflicts of interest.

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